Enhanced Specialized Transduction Using Recombineering in 
*Mycobacterium tuberculosis*

JoAnn M. Tufariello,a,b Adel A. Malek,a Catherine Vilchèze,a,d Laura E. Cole,a Hannah K. Ratner,a Pablo A. González,a,d Parvas Jain,a,d Graham F. Hatfull,a Michelle H. Larsen,a,b,d William R. Jacobs, Jr.a,c,d

Department of Microbiology and Immunology,a Department of Medicine,b and Department of Genetics,c and Howard Hughes Medical Institute,d Albert Einstein College of Medicine, Bronx, New York, USA; Department of Biological Sciences, University of Pittsburgh, Pittsburgh, Pennsylvania, USAe

* Present address: Pablo A. Gonzalez, Department of Molecular Genetics and Microbiology, Millennium Institute on Immunology and Immunotherapy, Pontificia Universidad Católica de Chile, Santiago, Chile.

**ABSTRACT** Genetic engineering has contributed greatly to our understanding of *Mycobacterium tuberculosis* biology and has facilitated antimycobacterial and vaccine development. However, methods to generate *M. tuberculosis* deletion mutants remain labor-intensive and relatively inefficient. Here, methods are described that significantly enhance the efficiency (greater than 100-fold) of recovering deletion mutants by the expression of mycobacteriophage recombineering functions during the course of infection with specialized transducing phages delivering allelic exchange substrates. This system has been successfully applied to the CDC1551 strain of *M. tuberculosis*, as well as to a *recD* mutant generated in the CDC1551 parental strain. The latter studies were undertaken as there were precedents in both the *Escherichia coli* literature and mycobacterial literature for enhancement of homologous recombination in strains lacking RecD. In combination, these measures yielded a dramatic increase in the recovery of deletion mutants and are expected to facilitate construction of a comprehensive library of mutants with every nonessential gene of *M. tuberculosis* deleted. The findings also open up the potential for sophisticated genetic screens, such as synthetic lethal analyses, which have so far not been feasible for the slow-growing mycobacteria.

**IMPORTANCE** Genetic manipulation of *M. tuberculosis* is hampered by laborious and relatively inefficient methods for generating deletion mutant strains. The combined use of phage-based transduction and recombineering methods greatly enhances the efficiency by which knockout strains can be generated. The additional elimination of *recD* further enhances this efficiency. The methods described herein will facilitate the construction of comprehensive gene knockout libraries and expedite the isolation of previously difficult to recover mutants, promoting antimicrobial and vaccine development.

 Tuberculosis (TB) remains a tremendous global health problem, with recent estimates of almost 9 million new cases annually and 1.4 million deaths (1). In addition, multidrug-resistant (MDR) and extensively drug-resistant (XDR) strains that resist the most effective chemotherapies have become increasingly common, and these strains are particularly lethal in the context of untreated HIV (2).

The development of techniques to genetically manipulate *Mycobacterium tuberculosis*, the causative agent of TB, and to generate defined mutants has revolutionized our ability to explore the biology and physiology of the bacterium and the pathogenesis of the disease. Despite the advances over the past 20 years, the genetic manipulation of *M. tuberculosis* has been limited by several factors, including the slow growth of the organism (doubling time of approximately 24 h), the need for biosafety level 3 (BSL3) containment, and most significantly, the relative inefficiency of existing techniques for targeted gene replacement compared with those used for “model” organisms, such as *Escherichia coli* and yeast (e.g., *Sacharomyces cerevisiae*), exacerbated by the relatively high frequency of illegitimate recombination that has been observed in slow-growing mycobacteria (3).

Following initial success with gene disruption in slow-growing mycobacteria by using linear DNA fragments, additional techniques were developed, including a “suicide vector” approach employing recombinant plasmids unable to replicate in mycobacteria and a two-step method using selectable and counterselectable markers located on replicating or nonreplicating plasmids (4). All of these approaches require electroporation to introduce the constructs into the mycobacterial cell. This can be problematic due to the relatively low efficiency with which DNA can be introduced into slow-growing mycobacteria by electroporation (5). This, together with the inefficiency of homologous recombination, results in significant challenges in recovery of mutants and also limits the technique to those strains that are most amenable to plasmid transformation. The development of methods for introducing the allelic exchange substrates (AESs) into recipient cells with *in vitro* generated specialized transducing phages represented a significant advance in the genetic manipulation of mycobacteria (6).
The specialized transducing phages consist of a cosmid cloning vector containing the AES—that is, upstream and downstream flanks of the targeted gene, separated by an antibiotic resistance cassette and in some instances a counterselectable marker—which is cloned into a conditionally replicating (temperature-sensitive) shuttle phasmid. The shuttle phasmid can replicate in *E. coli* as a plasmid and in mycobacteria as a plasmid and permits transfer of genetic material between the organisms (7). The transducing phage is amplified in *Mycobacterium smegmatis* at the permissive temperature of 30°C to generate a high-titer lysate, which is then used to transduce (infect) recipient mycobacteria at the nonpermissive temperature of 37°C. With appropriate adjustment of the multiplicity of infection (MOI), delivery of the AES by the phage to essentially every mycobacterial cell can be achieved (8). The inclusion of resolvase sites flanking the antibiotic resistance cassette allows the marker to be removed by transient expression of γδ-resolvase within the cell (6), permitting multiple rounds of gene deletion utilizing the same marker. This is important as the availability of only a few selection markers creates difficulties when knockouts of multiple genes are required to study the many redundant gene families in *M. tuberculosi*s.

A recent advance in the genetic manipulation of mycobacteria and the creation of defined deletion mutants came with the successful application of recombineering (recombination-mediated genetic engineering) techniques to both rapidly growing and slowly growing mycobacteria (9). In *E. coli*, transient expression of phage-encoded recombination proteins, including the λ*red system* (Exo, Beta, and Gam proteins) and the Rac prophage RecE/RecT system have been exploited to enhance the efficiency of homologous recombination and the isolation of the desired mutants (10, 11). The exonuclease (Exo or RecE) degrades DNA processively from the 5′ ends of breaks in double-stranded DNA (dsDNA), while the single-stranded DNA annealing protein (Beta or RecT) binds to the 3′ single-stranded tail, preparing the DNA for strand invasion (11). Together these activities facilitate homologous exchange of dsDNA substrates. Through database searches of sequenced mycobacteriophage genomes, van Kessel and Hatfull identified genes 60 and 61 encoding putative recombinases in the genome of the Che9c phage. Although somewhat distant relatives of their Rac prophage counterparts, gp60 possesses a dsDNA-dependent exonuclease activity similar to RecE, while gp61 acts as a single-stranded DNA binding protein similar to RecT (9). Co-expression of the proteins in *M. smegmatis* under control of the inducible acetamidase promoter enhanced homologous recombination sufficiently to permit the efficient recovery of gene replacement mutants following electroporation of linearized DNA substrates, including PCR products with ~500 bp of homology upstream and downstream of the targeted gene (9). Beyond its use in rapidly growing mycobacteria, the recombineering system has been further applied to *M. tuberculosis* for deletion of *groEL1* (9) and additional loci. Subsequently, expression of gp60 and gp61 under control of a nitrile-inducible promoter has been used by other groups for recombineering in *M. smegmatis* (12) and *M. tuberculosis* (13). However, as employed to date, these recombineering techniques are reliant upon introduction of recombination substrates into the bacterial cells by electroporation, which is subject to the limitations of transformability as discussed above.

Here we present methods that greatly enhance the efficiency by which targeted mutants can be generated in *M. tuberculosi*s. The technique relies on specialized transduction—which, unlike electro-

trporation, permits efficient phage-mediated delivery of the knockout construct to essentially the entire population of cells—but improves upon the method by taking advantage of the mycobacteriophage recombineering proteins. The plV53 plasmid containing the gp60 and gp61 coding sequences under control of the inducible acetamidase promoter (9, 14) was introduced by electroporation into the *M. tuberculosis* CDC1551 strain, with transformants selected on medium containing kanamycin (see Table S1 in the supplemental material for a list of strains used in this study). The CDC1551 wild-type parental strain and the strain transformed with plV53 ([CDC1551(pplV53-1)], were grown to mid-log phase (optical density at 600 nm [OD₆₀₀] of ~0.5) in 7H9 medium supplemented with 10% oleic acid-albumin-dextrose-catalase (OADC), 0.2% glycerol, 0.05% Tween 80 or tyloxapol, and 0.2% succinate; culture medium for the strain transformed with plV53 also included kanamycin at 25 µg/ml. At this time, the cultures were divided into two aliquots: no additional supplement was added to one aliquot ("uninduced"), while the second aliquot was supplemented with 0.2% acetamide ("induced"). After overnight incubation, bacteria were pelleted by centrifugation, washed with MP buffer (50 mM Tris [pH 8], 150 mM NaCl, 10 mM MgCl₂, 2 mM CaCl₂), resuspended in MP buffer at a 10-fold concentration versus the initial culture volume (~3 × 10⁹ bacteria per transduction), and incubated overnight at 37°C with ~0.5 ml high-titer phage lysate (10⁹ to 10¹¹ PFU/ml). The specialized transducing phages contained cosmid constructs targeting genes whose deletion had been previously demonstrated (15, 16) or would be predicted (17) to result in amino acid or vitamin auxotrophy: *serA1* (*Rv2996c [MT3074]*)*, metA* (*Rv3341 [MT3444]*)*, *panC* (*Rv3602c [MT3707]*)*, *leuD* (*Rv2987c [MT3065]*)*, and *argB* (*Rv1654 [MT1692]*) to induce auxotrophy for serine, methionine, pantothenate, leucine and arginine, respectively (see Table S2 in the supplemental material for sequences of primers used in cloning flanks for construction of AESs by using the p00045 vector [18]). Following the overnight incubation, bacteria were pelleted by centrifugation and resuspended in 7H9 medium supplemented with 0.05% Tween 80, and aliquots were plated in parallel onto 7H10 medium supplemented with hygromycin at 50 to 75 µg/ml alone or with additional supplements, including L-serine at 50 µg/ml, L-methionine at 50 µg/ml, L-leucine at 50 µg/ml, and L-arginine at 50 µg/ml. (The precise combination of supplements was chosen depending upon the panel of phages used in the particular experiment.) In some experiments, the transductions designed to generate amino acid auxotrophs were also plated in parallel onto medium supplemented with 0.2% Casamino Acids, in addition to the plating with the individual amino acid supplements; because numbers of transductants were found to be similar for the two types of media, the results were pooled in calculating the numbers of colonies per transduction.

Results of representative experiments are shown in Fig. 1A. The strain harboring the plV53 plasmid exhibited a dramatic increase in the number of transductants obtained, from ~10⁻⁰-fold up to ~86-fold more than the CDC1551 wild-type strain for the various loci. The loci that were targeted are distributed throughout the bacterial chromosome. Platings done in parallel to media with hygromycin but without other supplements yielded very few colonies—typically <10 per transduction—indicating that the colonies on the supplemented plates were very likely to be auxotrophs. This was verified by "pick and patch" replica plating of colonies

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Fig 1. The CDC1551 strain harboring the pJV53 recombineering plasmid exhibits an enhanced frequency of allelic replacement. (A) Mid-log-phase cultures of the CDC1551 wild-type strain or CDC1551 transformed with pJV53 [CDC1551(pJV53-1)] either were not treated [light blue bars for CDC1551, light red bars for CDC1551(pJV53-1)] or were subjected to overnight treatment with 0.2% acetamide [dark blue bars for CDC1551 and dark red bars for CDC1551(pJV53-1)]. Bacteria were then washed and incubated overnight at 37°C with specialized transducing phages to deliver allelic exchange substrates (AESs) for deletion of the serA1, metA, panC, leuD, or argB genes. Transductions were plated onto 7H10 medium containing 50 to 75 µg/ml hygromycin and the appropriate amino acid or vitamin supplement to allow recovery of auxotrophic deletion mutants. Colonies were counted after ~3 to 4 weeks of incubation. Fold enhancements in colony numbers versus the CDC1551 wild type are indicated. The auxotroph phenotypes of colonies appearing on the supplemented plates were further assessed by “pick and patch” testing, with a portion of each colony streaked onto 7H10 plus hygromycin or 7H10 plus hygromycin plus supplement. Shown are numbers of colonies growing only on supplemented media over the total numbers of colonies tested. In these studies, colonies for the argB mutants were slow to appear on solid media, requiring extended incubation; in a subsequent experiment, the L-arginine concentration of the supplemented media was increased to 200 µg/ml, and colonies then appeared in the typical 3 to 4 weeks. For serA1, the colonies obtained on Casamino Acid plates grew poorly on replica plate testing (even with the supplement), so the “pick and patch” results are for colonies obtained on transduction plates supplemented with L-serine. (B) Additional CDC1551 strains independently transformed with the pJV53 plasmid exhibit enhanced transduction frequencies similar to, or in some cases greater than, those of the original strain. Transductions were performed as in panel A above, with phages targeting the metA, panCD, and leuD loci and plated onto 7H10 medium plus hygromycin (hyg) and with or without L-methionine (met), p-pantothenate (pan), and L-leucine (leu) supplementation. Shown are colony counts for the acetamide-treated CDC1551 wild type (dark blue bars), CDC1551(pJV53-1) (the original [first] transformed strain [dark red bars]), CDC1551(pJV53-2) (the second independent transformant [teal bars]), and CDC1551(pJV53-3) (the third independent transformant [dark orange bars]), as well as CDC1551(pJV53-3) untreated with acetamide (light orange bars). Fold enhancements in colony numbers versus the CDC1551 wild type are indicated, as are proportions of auxotrophs confirmed on “pick and patch” testing for the metA and panC-panD loci in CDC1551(pJV53-1) (original strain) and CDC1551(pJV53-3) (third transformant).

from the original transduction plates (portions of the same colony streaked onto 7H10 plus hygromycin versus 7H10 plus hygromycin plus the single specific supplement required for the auxotroph in question), where the vast majority of the colonies were found to be auxotrophs (Fig. 1A), consistent with successful allelic replacement. Of note, the true enhancement of homologous recombination may be even greater than the 10- to 86-fold range depicted here, because the calculations of increased transduction frequency were determined by comparing numbers of colonies obtained on supplemented media, where a low background frequency of spontaneous hygromycin-resistant colonies can appear. However, “pick and patch” testing showed that the percentage of auxotrophs was in some cases greater for CDC1551(pJV53-1) than for the CDC1551 wild-type strain (see Fig. S1 in the supplemental material for an example with metA). However, this may depend as well on the locus under investigation, as panCD transductants were almost uniformly found to be auxotrophs for pantothenate for both the CDC1551 parental strain and CDC1551(pJV53-1) (data not shown).

The findings indicate that expression of the mycobacteriophage Che9c recombination proteins in the CDC1551 strain of M. tuberculosis promotes a greatly improved efficiency of homologous recombination between the phage DNA and the host chromosome. This may be surprising, as recombineering has been demonstrated between linear substrates (such as PCR products) and circular replicons (such as plasmids or a bacterial chromosome) (19), and more recently, recombineering between two linear DNA substrates has been verified (20), but to our knowledge the process has not been demonstrated between two circular molecules. The mycobacteriophage DNA is expected to circularize...
upon cell entry via cos (cohesive end site) sequences, the complementary single-stranded cohesive ends, by analogy with the phage λ replication cycle (21). However, the mycobacteriophage used in these studies is a temperature-sensitive (ts) derivative of phage TM4, and although the mutations thought to be responsible for the ts phenotype have been defined (22), the functions of the corresponding gene products and the behavior of the ts constructs upon cell entry have not been characterized. In addition, it is also possible that expression of the Che9c recombinases within the bacterial cell may alter the TM4 phage life cycle. This is an area that requires further exploration and may ultimately shed light on mechanisms of both mycobacteriophage replication and mycobacterial recombination.

The numbers of transductants obtained for CDC1551 (pJV53-1) were similar in the presence and the absence of acetamide (Fig. 1A), suggesting that, although the enhancement of homologous recombination requires pJV53, the inducer of the promoter (acetamide) is dispensable. There are several possible explanations for this observation. First, increased expression of gp60 and gp61 does not necessarily correlate with enhanced recombineering. In fact, in testing several constructs in M. smegmatis with electroporated linear substrates, it was noted that the number of colonies obtained was substantially increased for pJV53 compared with a similar plasmid construct that actually expressed higher levels of gp61 (9). This may be due in part to the toxicity of the recombinases, especially when expression is not tightly regulated (9). Regulation of the acm ammase promoter is complex and in M. smegmatis has been shown to be “leaky,” with a small amount of expression observed in both rich and minimal media in the absence of added inducer (14), while in M. tuberculosis, the situation may be further complicated by plasmid instability, as a vector containing the acm ammase promoter was found to be unstable and subject to deletions in the H37Rv strain (23). Because Parish et al. observed in M. smegmatis that the acm ammase promoter is controlled by a form of catabolite repression (14), we also tested transduction efficiency at the panCD loci in a more minimal medium, lacking the dextrose-containing OADC supplement and including 0.2% glycerol and 0.2% succinate as carbon sources (see Fig. S2 in the supplemental material). Transductions performed with this medium yielded a modest increase in colony numbers compared with the richer (glucose-containing) medium, suggesting a possible influence of catabolite repression on the system and some benefit for approaches that address this complication. In the minimal medium, as in rich medium, acetamide was largely dispensable for the enhanced efficiency of homologous recombination.

Because we observed enhanced specialized transduction in the CDC1551 strain harboring pJV53, even in the absence of acetamide, we sought to further demonstrate that the effect was due to the presence of the recombineering functions and not to a coincidental spontaneous mutation that may have occurred in the transformed strain. Therefore, we subsequently isolated two independent pJV53 transformants of CDC1551 (CDC1551[pJV53-2] and CDC1551[pJV53-3]) and examined their transduction capacities in comparison with the original strain. We found, again using a panel of phages to generate auxotrophs (targeting loci metA, panCD, and leuD), that the new transformants yielded colony counts that were similar to or even greater than the original strain (Fig. 1B). One of the newly created strains [CDC1551(pJV53-3)] was tested in the presence and absence of inducer, and again there was only a relatively modest fold increase in the number of transductants obtained with addition of acetamide, similar to what was observed for the original CDC1551(pJV53-1) strain. The fact that large numbers of colonies were obtained only on supplemented media, and not on 7H10 plus hygromycin alone, indicated that they were likely to be true auxotrophs, which was verified by pick-and-patch testing for the metA and panCD loci (Fig. 1B). Our finding that the three CDC1551 strains harboring pJV53 obtained by independent transformations all share the phenotype of high transduction efficiency makes a coincidental spontaneous mutation an unlikely explanation.

Because we would like to have a more tightly regulated expression system for the mycobacteriophage recombineering functions so as to avoid toxic effects within the cell (10), we conducted similar studies using a plasmid expressing the 60 and 61 genes under the control of a nitrile-inducible promoter (24), a system which has been exploited for recombineering-mediated gene replacement employing linear substrates in M. tuberculosis (13). M. tuberculosis CDC1551 was transformed with the pNIT::ET-sacB plasmid (generously provided by C. Sassetti) to generate strain CDC1551 (pNIT::ET-sacB), with transformants selected on kanamycin and screened by PCR for the presence of 60 and 61 sequences. A transduction assay in comparison with the parental strain for deletion of the leuD locus yielded an increased number of transductants, although the magnitude of the effects was less than for the CDC1551(pJV53-3) strain assayed in parallel (see Fig. S3 in the supplemental material). We once again observed that the effect appeared to be independent of the inducer (isovaleronitrile in this case). Although this remains to be experimentally verified, it is likely that, due to toxicity issues, increased expression levels of the recombineering functions may not lead to enhanced biological effects in M. tuberculosis, as we have observed a similar phenomenon in M. smegmatis (25). This alternative expression system using a different plasmid vector and promoter provides further reassurance that the enhanced recovery of transductants results from expression of the phage recombinases.

In the λ red recombineering system, Gam facilitates homologous recombination by interacting with the host RecBCD complex and inhibiting its various catalytic activities, including the exonuclease activity (26), preventing degradation of the incoming DNA. The RecBCD enzyme of E. coli possesses a complex set of activities and promotes recombination preferentially at short recognition sequences known as Chi sites (5’ GCTGTTGG 3’). Null mutations in recB and recC cause recombination-deficient phenotypes and sensitivity to DNA-damaging agents, while null mutations in recD yield cells which not only are recombination proficient but actually display enhanced recombination activity in the absence of Chi (27). It is thought that RecD functions as an inhibitor of recombination, perhaps due to interference with RecD with the loading of RecA (28), until the D subunit dissociates from the RecBCD enzyme upon encountering a Chi site (29).

Gam-related proteins do not appear to be encoded in mycobacteriophage genomes (9) and so are not available for testing in concert with gp60 and gp61; although not absolutely required for λ red-mediated recombination with linear double-stranded DNA substrates, Gam does improve the efficiency of this process (11). Therefore, we decided, as an alternative to including Gam in these studies, to examine the behavior of a recD null mutant of M. tuberculosis CDC1551 in our transduction assays. We targeted this..
component of the RecBCD complex due to the E. coli literature on the role of RecD as an inhibitor of homologous recombination. Specialized transduction was used to delete recD from the chromosome of the CDC1551 wild-type strain and also from the CDC1551 strain harboring pJV53, with deletion confirmed by Southern blot analysis (see Fig. S4 in the supplemental material); for the CDC1551(pJV53-1) strain, selection with kanamycin was included throughout to maintain the plasmid. The hygromycin resistance–sacB cassette was then removed by introducing resolvase via phage transduction (18), followed by plating on medium containing 10% sucrose, and the unmarking was confirmed by replica plating to demonstrate hygromycin sensitivity and by PCR analysis across the deleted region (see Fig. S4). The CDC1551 ΔrecD strain exhibited an enhanced transduction efficiency on the order of ~5- to 10-fold greater than that of the wild-type strain (Fig. 2A) for several loci tested; again, auxotrophy was confirmed by “pick and patch” testing. The experiment was repeated with deletion of panCD and yielded similar results (not shown). Furthermore, deletion of recD from the CDC1551 strain also expressing the phage recombineering functions [CDC1551(pJV53-1)] yielded a remarkably enhanced efficiency of gene replacement in this strain (Fig. 2A). Colony numbers increased substantially, from typically 10 or fewer per transduction for the wild-type strain to ~700 to 1,000 for deletion of the serA1 and metA loci in the presence of acetamide, for an approximate 100- to 150-fold enhancement of transduction efficiency.

In interpreting these results, it should be noted that Chi sites have not been identified in mycobacteria, and although mycobacteria do encode homologues of RecBCD, deletion of recBCD from M. smegmatis did not result in increased sensitivity to UV damage (30), in contrast to the findings for E. coli. The situation for mycobacteria is complex, with three distinct DNA double-strand break repair pathways identified, and it appears that RecBCD is involved in the single-strand annealing (SSA) pathway and not the homologous recombination pathway, which instead requires the AdnAB helicase-nuclease (31). However, Gupta et al. noted that while the SSA pathway of double-strand break repair was abolished for an M. smegmatis ΔrecBCD mutant, the homologous recombination activity was actually significantly enhanced versus the wild type, indicating that mycobacterial RecBCD actually suppresses homologous recombination (31) via mechanisms yet to be defined. Therefore, this could provide an alternative explanation for the enhanced gene replacement that we observe in the CDC1551 recD deletion mutant, in both the absence and presence of recombineering functions.

Given the large numbers of deletion mutants we were able to obtain in M. tuberculosis using strains expressing the Che9C recombinases, we performed small-scale transductions using reduced amounts of mycobacteria and specialized transducing phage, in volumes that could be accommodated in a 96-well-plate high-throughput format. We plated the transductions onto agar deposited into wells of a 24-well plate to illustrate the capacity for...
screening for synthetic lethal or, as in this case, conditional essential genes (Fig. 2B). Using the CDC1551(pJV53-3) strain and a phage to target deletion of the panCD locus, colonies were recovered in abundance on 7H10 medium supplemented with hygromycin and pantothenate but not on 7H10 medium supplemented with hygromycin alone. Similar studies with panels of phages should provide valuable information as to gene essentiality under different growth conditions.

Implications. Gene disruption is a basic tool for the geneticist and allows gene function to be elucidated. Inactivation of genes in a targeted fashion has been challenging for the slow-growing mycobacteria, including M. tuberculosis. We demonstrate here methods that greatly enhance the efficiency by which deletion mutants of M. tuberculosis can be generated. These methods will facilitate the construction of comprehensive mutant libraries. For example, in an effort to extend the potential of mycobacterial genetic analysis, a consortium of laboratories at the Albert Einstein College of Medicine (W. R. Jacobs, Jr.), Johns Hopkins University (W. Bishai), and University of Pittsburgh (G. Hatfull) in collaboration with the Genomic Institute of the Novartis Research Foundation (GNF) (A. Orth) seeks to generate a comprehensive set of specialized transducing phages that can be used to create bar-coded single-gene deletions in multiple strains of M. tuberculosis, such as the virulent H37Rv laboratory and CDC1551 clinical strains, as well as corresponding auxotrophic strains approved for use at biosafety level 2. The aim is to make these valuable resources openly available to the TB research community. Although the established methods for generating individual deletions in slowly growing mycobacteria are often successful, more facile methods amenable to high-throughput approaches would greatly facilitate the process of generating the complete bar-coded mutant collection. Such mutant libraries will make possible analyses that have been difficult or impossible to perform in a truly comprehensive fashion for M. tuberculosis. These include fitness profiling, or defining growth defects under specific environmental conditions (32), including growth in various in vivo models. The bar-coded library will offer a significant advantage over transposon site hybridization (TraSH) screens (33), where candidate genes can be identified, but the transposon insertions responsible for given phenotype are not recovered as bacteria, requiring the experimenter to separately construct and test specific mutants in the implicated region. The library of specialized transducing phages also provides a framework for undertaking systematic analyses of genetic interactions (34), such as synthetic lethal analysis to identify combinations of mutations in two or more genes that lead to loss of viability, often by affecting a common function or pathway. It should be possible to create an (unmarked) deletion mutant in the gene of interest, in the CDC1551(pJV53) strain, and then perform transductions with either the full complement of specialized transducing phages or a desired subset. Due to the greatly enhanced efficiency of gene replacement in this context, any transductions which yield colonies only with the CDC1551(pJV53) parental strain, but not with the strain with the gene of interest deleted, would be candidates for synthetic lethal interactions. Ultimately, examining the phenotypes of double mutants on a genome-wide scale should help to elucidate the components of essential biological pathways and their interactions within the cell and facilitate the development of novel antimycobacterial agents.

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at http://mbio.asm.org lookup/supp/doi:10.1128/mBio.01179-14/-/DCSupplemental.

Figure S1, EPS file, 6.7 MB.
Figure S2, EPS file, 0.8 MB.
Figure S3, EPS file, 3.1 MB.
Figure S4, EPS file, 3.4 MB.
Table S1, DOCX file, 0.1 MB.
Table S2, DOCX file, 0.1 MB.
Table S3, DOCX file, 0.1 MB.

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