





Development of a Counterselectable Transposon To Create Markerless Knockouts from an 18,432-Clone Ordered *Mycobacterium bovis* Bacillus Calmette-Guérin Mutant Resource

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ABSTRACT Mutant resources are essential to improve our understanding of the biology of slow-growing mycobacteria, which include the causative agents of tuberculosis in various species, including humans. The generation of deletion mutants in slow-growing mycobacteria in a gene-by-gene approach in order to make genome-wide ordered mutant resources is still a laborious and costly approach, despite the recent development of improved methods. On the other hand, transposon mutagenesis in combination with Cartesian pooling-coordinate sequencing (CP-CSeq) allows the creation of large archived *Mycobacterium* transposon insertion libraries. However, such mutants contain selection marker genes with a risk of polar gene effects, which are undesired both for research and for use of these mutants as live attenuated vaccines. In this paper, a derivative of the Himar1 transposon is described which allows the generation of clean, markerless knockouts from archived transposon libraries. By incorporating *FRT* sites for FlpE/*FRT*-mediated recombination and *I-SceI* sites for IScelM-based transposon removal, we enable two thoroughly experimentally validated possibilities to create unmarked mutants from such marked transposon mutants. The *FRT* approach is highly efficient but leaves an *FRT* scar in the genome, whereas the *I-SceI*-mediated approach can create mutants without any heterologous DNA in the genome. The combined use of CP-CSeq and this optimized transposon was applied in the BCG Danish 1331 vaccine strain (WHO reference 07/270), creating the largest ordered, characterized resource of mutants in a member of the *Mycobacterium tuberculosis* complex (18,432 clones, mutating 83% of the nonessential *M. tuberculosis* homologues), from which markerless knockouts can be easily generated.

IMPORTANCE While speeding up research for many fields of biology (e.g., yeast, plant, and *Caenorhabditis elegans*), genome-wide ordered mutant collections are still elusive in mycobacterial research. We developed methods to generate such resources in a time- and cost-effective manner and developed a newly engineered transposon from which unmarked mutants can be efficiently generated. Our library in the WHO reference vaccine strain of *Mycobacterium bovis* BCG Danish targets 83% of all nonessential genes and was made publicly available via the BCCM/ITM Mycobacteria Collection. This resource will speed up *Mycobacterium* research (e.g., drug resistance research and vaccine development) and paves the way to similar genome-wide mutant collections in other strains of the *Mycobacterium tuberculosis* complex. The stretch to a full collection of mutants in all nonessential genes is now much shorter, with just 17% remaining genes to be targeted using gene-by-gene approaches, for which highly effective methods have recently also been described.


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 Closest to genome-wide mutant resource of TB complex strain we've ever got: 83% of nonessential genes hit, >18,000 strains, in WHO BCG Danish. New removable transposon, mutants all publicly available from BCCM/ITM culture collection.

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Tuberculosis (TB) is one of the main threats to public health in large areas of the world (1). Drug resistance is on the rise, and an effective vaccine is still lacking. For many types of *Mycobacterium* research, e.g., into potential drug targets, and for live attenuated vaccine engineering, mycobacterial mutants are required. However, the genetic manipulation of slow-growing mycobacteria has long been problematic due to their slow growth, the low efficiency of classical transformation methods, and the high frequency of illegitimate recombination. The lab of W. R. Jacobs, Jr., has, however, optimized specialized transduction in such a way that we now have a feasible method to create specific deletion mutants in slow-growing mycobacteria (2, 3). Murphy and others developed oligonucleotide-mediated recombineering followed by Bxb1 integrase targeting (ORBIT) for the creation of mutants with targeted deletions, insertions, or fusions in the mycobacterial chromosome (4). Recently, Yan and colleagues optimized a CRISPR-assisted nonhomologous-end-joining (NHEJ) strategy for efficient genome editing in *Mycobacterium tuberculosis* (5).

Nevertheless, even with the improved engineering methods, gene-by-gene mutation of the entire mycobacterial genome still takes a long time and a lot of effort and expense, and no publicly available mutant collections that approach completeness are available, forcing most investigators to invest in the generation of their own mutants of interest time and again. On the other hand, transposon mutagenesis allows the creation of a wide variety of mutants in an isogenic mycobacterial background in a quick and efficient manner. However, it yields a complex mixture of transposon mutants. Such mutant mixtures are useful for analyzing competitive mutant behavior in bulk (6) but are not well suited for extracting single mutants. Ordered transposon insertion libraries can be picked from such random mutant mixtures and screened for insertions in a locus of interest (7, 8), but this is laborious and needs to be repeated for every gene.

In a previous report, we described an easy-to-implement, cost- and time-effective, massively parallel sequencing-based approach (called Cartesian pooling-coordinate sequencing [CP-CSeq]) to characterize ordered transposon libraries (9). These libraries consist of multiwell plates in which each well contains a transposon mutant colony picked from agar plates of the bulk transposon mutant library. The method combines a pooling strategy of a stack of 96-well plates along the Cartesian coordinates (x , y , and z) with multiplex transposon sequencing and reports both on the identity and on the location of the transposon mutants in the plate stack. We illustrated the approach for a large (9,216 [96 × 96]-clone) ordered transposon insertion mutant library of a streptomycin-resistant lab strain of *Mycobacterium bovis* bacillus Calmette-Guérin (BCG) Pasteur. At that time, this library was the largest characterized resource of *M. bovis* BCG mutants and had identified clones with disruptions in 64% of the nonessential *M. tuberculosis* gene orthologues (9). A range of investigators have since made use of this resource to catalyze their research, which motivated us to further innovate to substantially improve on this proof-of-concept resource.

Although transposon insertion mutants are valuable for fundamental mycobacterial research, they also have a few drawbacks. This is mainly due to the presence of transcriptionally active transposon-associated sequences, especially an antibiotic selection marker, increasing the risk for polar effects. We have recently observed such an effect in a vaccine research project in our lab, where transposon insertion in the *sapM* gene resulted in a strong enhancement of expression of the upstream *upp* gene, necessitating a significant effort in complementation experiments and laborious generation of a “clean” *sapM* knockout (KO) to exclude a role for this Upp upregulation in the observed enhanced vaccine efficacy of *sapM* mutation in BCG (10). The presence of antibiotic selection markers also precludes the use of such transposon mutants for further (pre)clinical development as vaccines. Hence, we had a strong incentive to solve

this problem and to investigate how selection marker-free and even heterologous DNA-free mutants could be derived from transposon mutants.

Here, we describe the construction of a derivative of the Himar1 transposon to allow subsequent generation of unmarked KOs from marked transposon mutants. Among several other methodological options that were tested but found impracticable (including CRISPR-Cas9 inspired sequence deletion), we found two effective possibilities to create unmarked mutants: use of *FRT* sites for FlpE/*FRT*-mediated out-recombination of transcriptionally active transposon parts and *I-SceI* sites for I-SceI meganuclease (ISceIM)-based transposon removal. The first approach inherently leaves an *FRT* scar, whereas the last one has the potential of creating clean, markerless mutants. To make this practicable, a focused effort was needed to explore different genetic carriers for the unmarking enzymes, both in fast-growing (*M. smegmatis*) and in slow-growing mycobacteria (*M. bovis* BCG). Finally, we combined the use of this “unmarkable” transposon and CP-CSeq in the clinically used BCG Danish 1331 vaccine strain (WHO reference strain NIBSC 07/270), creating a validated characterized ordered transposon insertion library of 2 96- by 96-well plates, or ~18,000 clones. The library can be used in its entirety or in ordered subcollections, e.g., for chemical genomics screens or *in vitro* tests of interactions with immune cells. Transposon mutants of interest can be retrieved and subsequently unmarked to create “clean” mutants to validate initial findings. Ultimately, these clean mutants can even be used directly for (pre)clinical vaccine development.

(This article was submitted to an online preprint archive [11].)

RESULTS

Himar1 transposon redesigned to enable subsequent mutant unmarking. Several site-specific recombination systems have already been used in mycobacteria to allow the removal of an antibiotic resistance cassette, including the $\gamma\delta$ -resolvase/*res* (2, 12–15), the FlpE/*FRT* (16–18), the Cre/*loxP* (18, 19), and the Xer/*dif* (20, 21) systems. The $\gamma\delta$ -resolvase/*res* and FlpE/*FRT* systems were the first ones shown to be functional in mycobacteria (18). We chose to implement the FlpE/*FRT* system from *Saccharomyces cerevisiae*, as for this system a higher recombination frequency was reported than for the $\gamma\delta$ -resolvase system (>40% versus 3 to 5% in slow-growing mycobacteria) (17).

Another strategy to remove the antibiotic resistance cassette is by using meganucleases. These engineered versions of naturally occurring restriction enzymes typically have extended DNA recognition sequences (12 to 40 bp). The I-SceI meganuclease (ISceIM) is an example; it recognizes a specific 18-bp sequence that statistically occurs only once every 1,010 to 1,011 bp. Most organisms' genomes lack an endogenous *I-SceI* site, including most, if not all, *Mycobacterium* species. The *I-SceI* system has been used to unmark mutants in several organisms (22–26) but not yet for unmarking of mycobacterial mutants. We saw the opportunity to use it for this purpose, as in *Mycobacterium smegmatis*, the *I-SceI* system has already been used to study the repair mechanisms of double-stranded DNA breaks (DSBs) (27, 28). The created DSBs either resulted in death of the cell or were repaired by NHEJ, which added additional nucleotides prior to religation or resulted in deletions (28).

We designed a derivative of the original Himar1 transposon (29) in which we incorporated both the *FRT* and the *I-SceI* system to enable creation of unmarked mutants from marked transposon mutants (Fig. 1a and b). The *FRT* sites are present as direct repeats, so that the DNA sequence in between will be deleted upon site-specific recombination. The *I-SceI* sites are in inverted orientation, so that the overhanging ends are not compatible. This will avoid that a simple ligation can resolve the created DSB, since we want DSBs to be repaired by NHEJ, with its associated exonuclease-mediated shortening of the ends. This exonuclease activity can be extensive, and we hypothesized that it may sometimes degrade all of the transposon sequences prior to religation. In addition, we introduced the negative selection marker *SacB* to allow for direct selection of unmarked clones on sucrose. A transposon plasmid containing this optimized transposon and the hyperactive Himar1 transposase gene was constructed (see

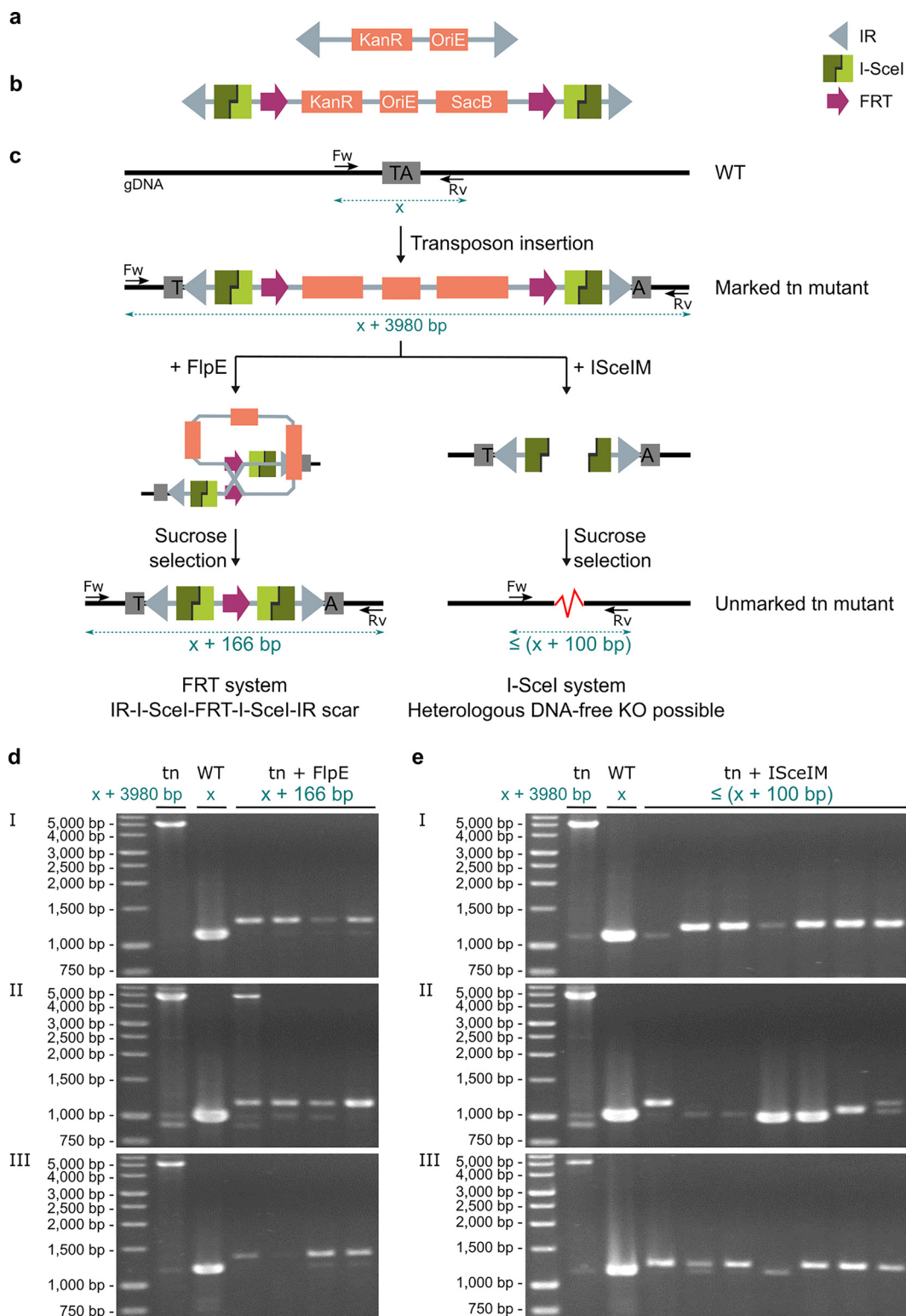


FIG 1 Creating markerless KOs from transposon mutants using an optimized Himar1 transposon. (a) Visual representation of the original Himar1 transposon (2,064 bp), which consists of a kanamycin resistance marker (KanR) (795 bp) and an origin of replication for *E. coli* (OriE) (732 bp) flanked by two inverted repeats (IR) (27 bp). (b) Optimized Himar1 transposon (3,980 bp), which consists of a KanR (795 bp), an OriE (614 bp), and a SacB negative selection marker (1,419 bp) flanked by two *FRT* sites (34 bp), two *I-SceI* sites (18 bp), and two IRs (27 bp). (c) Illustration of the construction of a transposon mutant (marked) using the optimized transposon and subsequent unmarking strategies by bringing FlpE or ISceIM into the cells and selecting the unmarked mutants. FlpE/*FRT*-mediated recombination inherently leaves an *FRT* scar (IR-*I-SceI*-*FRT*-*I-SceI*-IR; 166 bp) in the unmarked transposon mutant, whereas ISceIM-based transposon removal has the potential of creating KO mutants free of (Continued on next page)

Method S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>). Subsequently, we produced an optimized transposon phage using the improved shuttle phAE159 from the lab of W. R. Jacobs, Jr., which has increased cloning capacity and efficient DNA delivery (2). This was essential, as the optimized transposon (~4 kbp) is 2 kbp larger than the original Himar1 transposon (~2 kbp). Furthermore, this larger size of the transposon would be predicted to lead to an ~74% decrease in transposition frequency (30). While we did not formally test this, indeed, transposon mutagenesis of the BCG Danish strain yielded ~25,000 mutants for the new, larger transposon, under conditions that would yield ~100,000 mutants described in the literature for the original Himar1 transposon (31, 32), and as we have indeed observed earlier in our own lab (9).

Introduction of FIpE in the marked transposon mutants was designed to lead to site-specific recombination at the *FRT* sites, resulting in removal of most of the transposon-associated sequence but leaving an inverted-repeat (IR)-*I-SceI-FRT-I-SceI*-IR scar in the genome (Fig. 1c). Every FIpE-unmarked transposon mutant will contain the same scar in its genome. A proof of concept of this strategy was obtained by introducing FIpE in the marked transposon mutants using a standard replicable vector. Colony PCR performed on FIpE-unmarked transposon mutants indeed showed, for all clones, an amplicon of a size that was consistent with a 166-bp insertion compared to the wild-type (WT) amplicon, both for fast-growing *M. smegmatis* (Fig. S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>) and for slow-growing BCG Danish transposon mutants (Fig. 1d).

Introduction of ISceIM in the marked transposon mutants was designed to form a DSB that is repaired by NHEJ (Fig. 1c) (28). Hereby, more extensive exonuclease activity prior to DSB repair will lead to a different outcome. In a small percentage of cases, heterologous DNA-free KOs in which all of the transposon sequences were deleted can be formed. A proof of concept of this strategy using a standard replicable vector to introduce ISceIM in the marked transposon mutants indeed showed that the amplicon length from ISceIM-unmarked clones differed, as shown for ISceIM-unmarked *M. smegmatis* (Fig. S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>) and BCG Danish transposon mutants (Fig. 1e).

Comparison of different genetic carriers for FIpE or ISceIM to unmark the transposon mutants. Standard replicable vectors proved to be sufficient to unmark transposon mutants using FIpE or ISceIM but are known to be more difficult to cure from slow-growing bacteria. Therefore, we also produced several genetic carriers that can be easily cured from the cells or do not replicate in the cells. We compared three different vector systems (standard replicable vectors, replicable GalK vectors, and temperature-sensitive [ts] GalK vectors) and ts phages (Fig. 2). All genetic carriers were tested in both fast-growing (*M. smegmatis*) and slow-growing mycobacteria (BCG Danish) and evaluated for three characteristics: the transformation efficiency, the unmarking efficiency, and the curing efficiency (if applicable). The results are summarized in Fig. 2 and discussed below.

For *M. smegmatis*, we got similar results for a replicable vector with or without counterselectable marker GalK and with or without a ts *oriM*: they all had equal unmarking and curing efficiencies. Neither the negative selection marker GalK nor the ts *oriM* was needed to establish curing; subculturing the clones and plating them on nonselective plates was sufficient to cure the vectors at $\pm 100\%$ efficiency (Fig. 2). The standard replicable vector was the best option for unmarking *M. smegmatis* transposon

FIG 1 Legend (Continued)

heterologous DNA. (d and e) Proof of concept of the unmarking of three *M. bovis* BCG Danish transposon mutants (I, II, and III). Transposon mutants were unmarked via FIpE/*FRT*-mediated recombination (d) or ISceIM-based transposon removal (e). Therefore, replicable vectors expressing FIpE or ISceIM were introduced in the cells via electroporation, after which unmarked mutants (kanamycin sensitive and sucrose resistant) were analyzed via colony PCR. Parental transposon mutants (I, II, and III) and the original wild-type (WT) strain were taken along as controls. Lengths of expected amplicons are indicated in blue in panels c, d and e. x, length of WT amplicon; tn, transposon. Used primers are listed in Table S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

	Genetic carrier	Promoter	Transformation efficiency	Unmarking efficiency	Curing efficiency
<i>M. smegmatis</i>	std replicable vector	const.	+++	~100%	~100%
	replicable GalK vector	const.	++(+) (grew slower)	~100%	~100% (no need to add 2-DOG)
	ts GalK vector	const.	+++ at 32°C	~100%	~100% (no need to add 2-DOG or to apply restrictive temp.)
	suicide vector	const.	NA	~0%	NA
	ts phage	const.	NA	~0% FlpE, 0-27% ISceIM	NA
<i>M. bovis</i> BCG Danish	std replicable vector	const.	+	~100%	35-100%
	replicable GalK vector	tet-ind.	++	~100%	0-40%
	ts GalK vector	const.	-/+ (grew slower)	~100%	80-100% (+ 2-DOG 100%)
	replicable GalK vector	tet-ind.	-/+ (grew slower)	~100%	90-100% (+ 2-DOG 100%)
	ts GalK vector	const.	- at 32°C, ++ at 39°C *	50% FlpE, 100% ISceIM	/
	suicide vector	const.	NA	~0%	NA
	ts phage	const.	NA	0-11% FlpE, 20-80% ISceIM	NA

FIG 2 Unmarking *M. smegmatis* and *M. bovis* BCG Danish transposon mutants using different genetic carriers bringing FlpE or ISceIM to expression. Two or three transposon mutants for *M. smegmatis* (top) or *M. bovis* BCG Danish (bottom) were unmarked by introducing FlpE or ISceIM by different genetic carriers, expressing FlpE or ISceIM from a constitutive (const.) or inducible promoter. The efficiency of transformation (amount of hygromycin-resistant clones) was given a score ranging from - to +++ based on the observed CFU after incubation at 37°C (unless otherwise indicated, e.g., for the ts vectors). The efficiency of unmarking is indicated by the percentage of unmarked clones (kanamycin sensitive and sucrose resistant) among the sucrose-resistant clones (and hygromycin resistant for replicable vectors) obtained after unmarking. The efficiency of curing is indicated by the percentage of cured clones (hygromycin sensitive). All experiments were replicated once or more (e.g., replicable vectors), except the data for the replicable GalK vectors with the Tet-inducible promoters, as no improvement was seen in the transformation efficiency compared to that of the replicable GalK vectors with the constitutive promoters in the first experiment. NA, not applicable (vector or phage cannot replicate in mycobacteria); /, not performed (would not give any added value); std, standard; GalK, negative selection marker galactokinase; ts, temperature sensitive; acet. ind., acetamide inducible; tet-ind., tetracycline inducible; *, problem with the use of the ts vectors in BCG Danish: BCG Danish did not grow at 32°C but grew at 39°C. In gray, we indicate the most preferred tools.

mutants, as the GalK vectors showed a slightly lower transformation efficiency and a slower clone growth.

For BCG Danish, the ts GalK vectors were not feasible to unmark the transposon mutants, because transformed cells showed little to no growth at 32°C. Similar to the situation in *M. smegmatis*, the replicable GalK vector conferred more stress on the cells, resulting in a lower transformation efficiency and smaller colonies. On the other hand, due to apparent stress associated with the incorporation of the negative selection marker GalK, the replicable GalK vectors were more easily cured from the cells than the standard replicable vectors, with or without adding 2-deoxy-galactose (2-DOG) to the agar plates.

Suicide vectors and ts phages expressing FlpE or ISceIM do not need to be cured, as they cannot replicate (they lack an origin of replication for mycobacteria). This is advantageous, as unmarking can be done in a single round of transformation and selection, which is less time-consuming. However, they were not useful to unmark *M. smegmatis* transposon mutants, as their unmarking efficiency was very low. This was due to a high number of SacB escape mutants and the inability to select for uptake of the genetic constructs (the suicide vector or the ts phage). In BCG Danish, the same held true for the suicide vectors. However, the unmarking efficiency for the ts phage was satisfactory in BCG Danish, especially for ISceIM (20 to 80%), allowing their use to unmark transposon mutants in slow-growing mycobacteria. Nonetheless, the unmarking of our transposon mutants with these ts phages was still less efficient than described in an earlier report on unmarking of specialized transduction mutants containing *res* sites with phAE280 (ts phage expressing $\gamma\delta$ -resolvase), which reportedly reached up to 80 to 100% efficiency (reference 2 and our own unpublished observations).

As initial experiments (data not shown) demonstrated that FlpE and ISceIM expression can be toxic in slow-growing mycobacteria, strongly reducing transformation efficiency, we also constructed tetracycline (Tet)-inducible vectors, next to the constitutive vectors. For the tetracycline-inducible vector, we indeed retained more viable transformants than with the respective constitutive vector, reliably obtaining ~100 to

200 clones for one transformation with the Tet-inducible promoter, while in some experiments no unmarked clones were obtained with the constitutive promoters. Most likely because they cause less stress, the tetracycline-inducible vectors were less efficiently cured. We tested the addition of the GalK counterselectable marker to enhance the curing efficiency of these vectors. Unfortunately, when introducing GalK, we lost the positive effect of the tetracycline-inducible promoters on the transformation efficiency. Probably, the toxic stress effect of GalK, which we also observed in *M. smegmatis*, overpowered the reduced stress imparted by low expression of FlpE/ISceIM allowed by tetracycline-inducible promoters. Nevertheless, the curing efficiency of the few unmarked clones that were obtained was indeed enhanced under counterselective conditions.

We conclude that for *M. smegmatis*, the replicable vectors are the best option to unmark transposon mutants. For *M. bovis* BCG Danish, it is difficult to compare the efficacy of the plasmid methods with the unmarking phages, due to the inherent differences between the methods. Some practical advantages of the ts phages are, however, quite clear: (i) transduction was easier to perform than electroporation and was more efficient, and (ii) the carrier did not need to be cured afterwards. However, not all recovered colonies were unmarked, unlike with the replicable vectors, where ~100% of hygromycin- and sucrose-resistant clones were unmarked. Nonetheless, with ts phages we could easily obtain ~200 clones after one transduction, after which a straightforward screening method (streaking on plates with and without hygromycin) allowed the identification of unmarked mutants. Overall, we give preference to the methods that yield a robust number of clones upon transgenesis of the unmarking tool, together with high unmarking efficiency and easy curing of the unmarking vehicle. The Tet-inducible replicable vectors and the ts phages struck the best compromise for BCG.

Different outcomes of the *I-SceI* system. In contrast to the *FRT* system (which always yields the same scar), the *I-SceI* system creates a DSB that is repaired by NHEJ, yielding a range of scar sizes. We classified all obtained ISceIM-unmarked transposon mutants in three categories (Fig. 3a): (i) mutants with two remaining IRs (leaving a transposable transposon in the genome), (ii) mutants with one remaining IR, and (iii) mutants with no transposon sequences remaining in the genome. Several ISceIM-unmarked transposon mutants were analyzed in depth to investigate the different outcomes of NHEJ in fast-growing *M. smegmatis* (test model) (Fig. 3b) and slow-growing BCG Danish (target organism) (Fig. 3c).

For *M. smegmatis*, unmarking via the *I-SceI* system resulted mostly in small indels (>95%) in which two intact IRs remained. Larger deletions occurred only for a small percentage, after which only one IR (<5%) or no transposon sequences (<0.4%) remained. In-depth analysis of the small indels was hard, as the repetitive nature of the IR-indel-IR site (forms an almost perfect hairpin) completely obstructed Sanger sequencing of these indels.

The transformation efficiency in BCG Danish was lower than in *M. smegmatis*, leading to fewer ISceIM-unmarked clones, but with a higher frequency of larger deletions after NHEJ than for *M. smegmatis*. In 21% of the unmarked clones only one intact IR remained in the genome, compared to <5% for *M. smegmatis*. In 2% of the cases, no transposon sequences remained in the genome, compared to <0.4% for *M. smegmatis*. Hence, the *I-SceI* system is a feasible way of generating heterologous DNA-free KOs from BCG Danish transposon mutants. If only inactivation of a transposable transposon is needed, destruction of one of the IRs is sufficient, which occurs at a 10-fold-higher frequency (21% versus 2%).

Note that when no transposon sequences remain in the genome, the size of the deletions varies depending on how the DSB was repaired. Therefore, it is possible that smaller deletions have no, or a moderate, impact on the protein function (e.g., a truncated protein form remains), while bigger deletions can also affect adjacent genes. This can be verified by means of phenotypic analysis of the mutants.

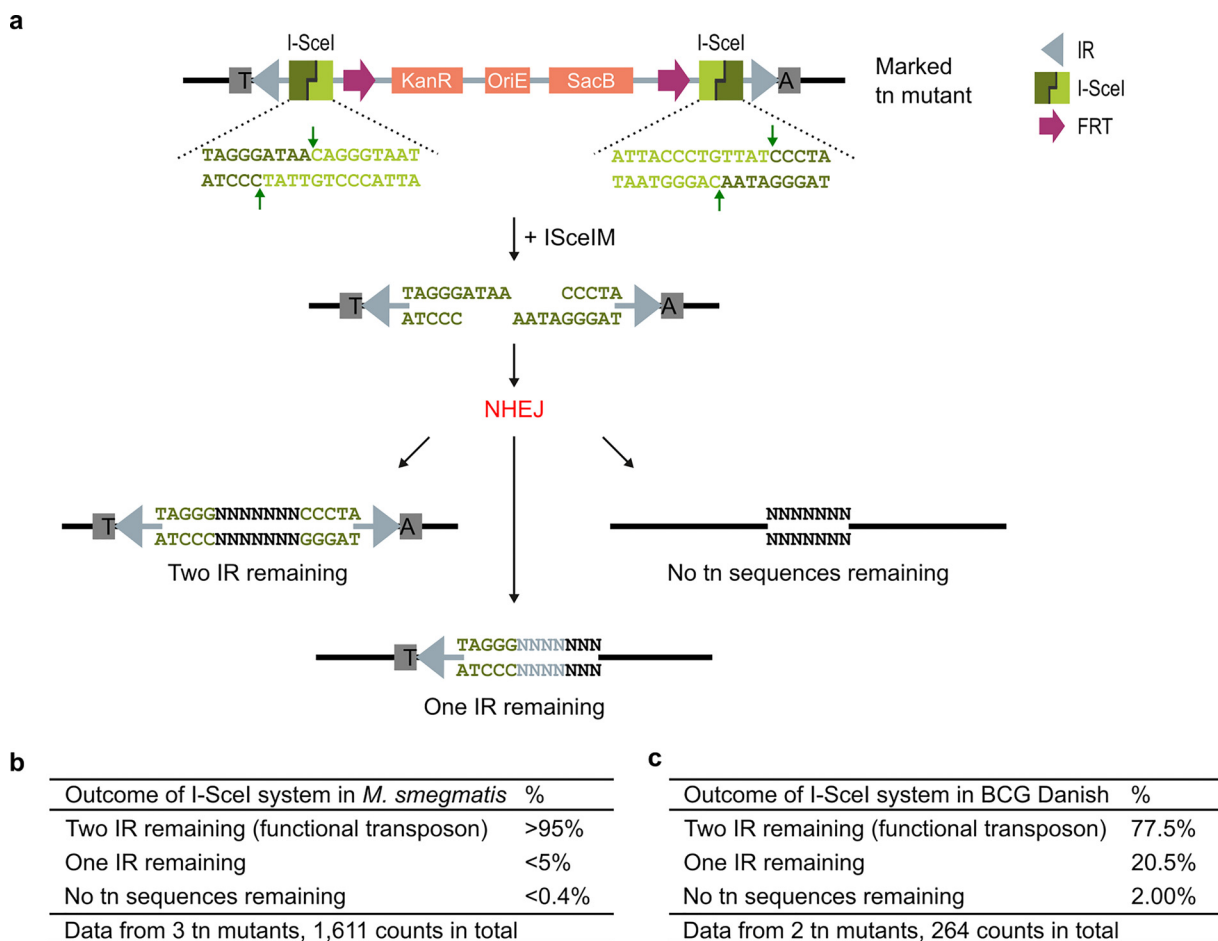


FIG 3 Outcome of unmarking transposon (tn) mutants via the I-SceI system. (a) Graphic illustration of the outcome of unmarking via the I-SceI system. Upon introduction of I-SceIM, the restriction enzyme will cut the I-SceI sites, creating a DSB that will be repaired by NHEJ, resulting in one of three possible outcomes. (b and c) The outcome of the I-SceI system was analyzed for three *M. smegmatis* transposon mutants (b) and two BCG Danish transposon mutants (c) by performing PCR on kanamycin-sensitive and sucrose-resistant clones and Sanger sequencing the PCR amplicons (for *M. smegmatis*, only a subset of the clones was Sanger sequenced). Outcome is given in percentages. Used primers are listed in Table S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

Development of a *Mycobacterium bovis* BCG Danish mutant resource. We have combined the use of this new transposon phage and CP-CSeq (9) in the BCG Danish 1331 vaccine strain (WHO reference 07/270), creating an ordered transposon insertion library of 2 sets of 96-by 96-well plates (Data Set S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>). Each transposon mutant in this library can be unmarked using the methods described above. CP-CSeq characterization of this library provided evidence for transposon insertions at 20,618 sites: 19,481 in unique genome regions and 1,137 in duplicated regions or highly repetitive genes (Fig. 4a). Theoretically, the library could only contain 18,432 mutants (i.e., the total number of wells in the library plates); the 2,186 mutants we have in excess can be explained in two ways. First, due to the clumping of BCG cells, many colonies on agar plates of the transposon library were likely not clonal. While we did the utmost in the clone picking to avoid mixed colonies, it was impossible to avoid this completely. This is also the reason why we stuck to manual picking by experienced scientists, rather than with robots, as these cannot yet achieve the same level of accuracy in distinguishing single colonies. Hence, 14% of the wells for set I and 11% for set II contained more than one clone. Second, due to a minimal aspecific amplification of the genomic DNA (gDNA) (higher background) during amplification of the transposon-flanking regions, a low level of spurious transposon insertion detection occurs. To limit this noise, we excluded transposon insertions

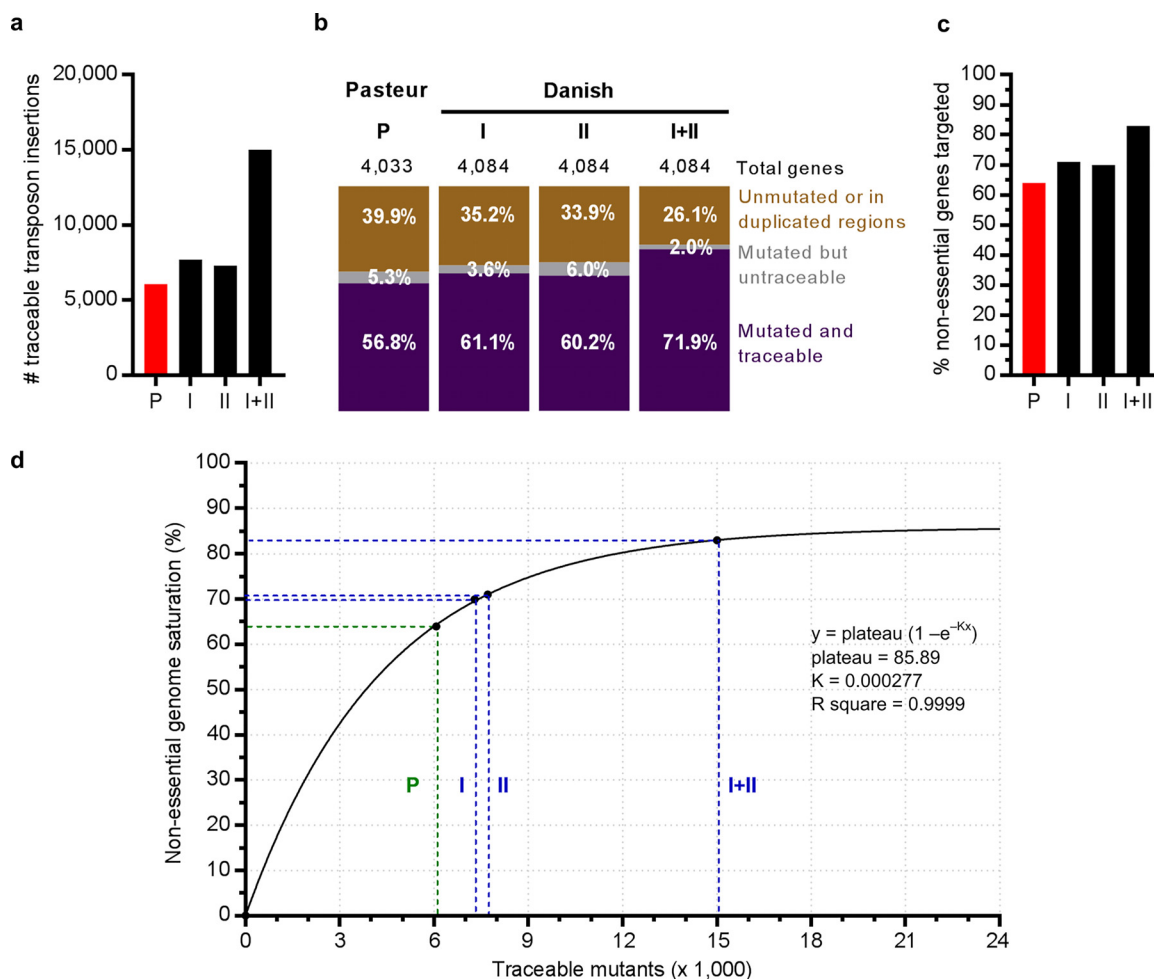


FIG 4 Results of CP-CSeq on a transposon insertion library of *M. bovis* BCG Danish (2 sets of 96- by 96-well plates, set I and set II [this study]) compared to *M. bovis* BCG Pasteur (9). (a) Number of traceable transposon insertions for each library (set). (b) Distribution between disrupted genes that are traceable (purple), disrupted genes for which the locations of their disruption mutants are unknown (“untraceable” [gray]), and untargeted genes or genes in duplicated regions (brown). (c) Percentage of targeted nonessential genes for each library (set). (d) Nonessential genome saturation in function of the amount of traceable mutants. In green are the observed values for the library in *M. bovis* BCG Pasteur (P). In blue are the observed values for the library in *M. bovis* BCG Danish (set I, set II, and set I plus set II). To model what would be gained by further enlargement of CP-CSeq libraries, we fitted a coupon collector’s equation (in black) to the available experimental data from our libraries, which should allow more accurate predictions in the future. The coupon collector’s problem estimates the amount of traceable mutants needed to target a certain percentage of the nonessential genome. The equation for the curve is given in the graph. This curve goes not through 100%, probably due to inherent characteristics of transposon mutagenesis and clone picking of transposon mutants grown on agar.

that were detected with a total Cartesian sequencing read coverage below 60 (the average read depth was $4,709 \pm 3,626$ for the remaining nonnoise transposon insertions).

Using these settings for CP-CSeq, for 15,320 insertion events in unique regions of the BCG Danish genome, the well position of the mutant in the plate library could be identified; 7860 for set I and 7460 for set II (Fig. 4a). In the combined resource (set I plus set II), about 85% of these insertions mapped against genes, targeting 83% of all nonessential *M. tuberculosis* orthologues at least once (Fig. 4b). The current library in BCG Danish outperformed the previous library in BCG Pasteur (9), as the separate sets of the new library targeted 71% and 70% of all nonessential genes, while this was only 64% for the old library (Fig. 4c). This improvement was most likely due to experimental improvements in the construction of the libraries (Method S7 at <https://doi.org/10.6084/m9.figshare.c.4507472>).

As Himar1 transposon insertion is a random event that can occur at a finite number of TA sites in the genome, the relation between the “number of attempts” and the

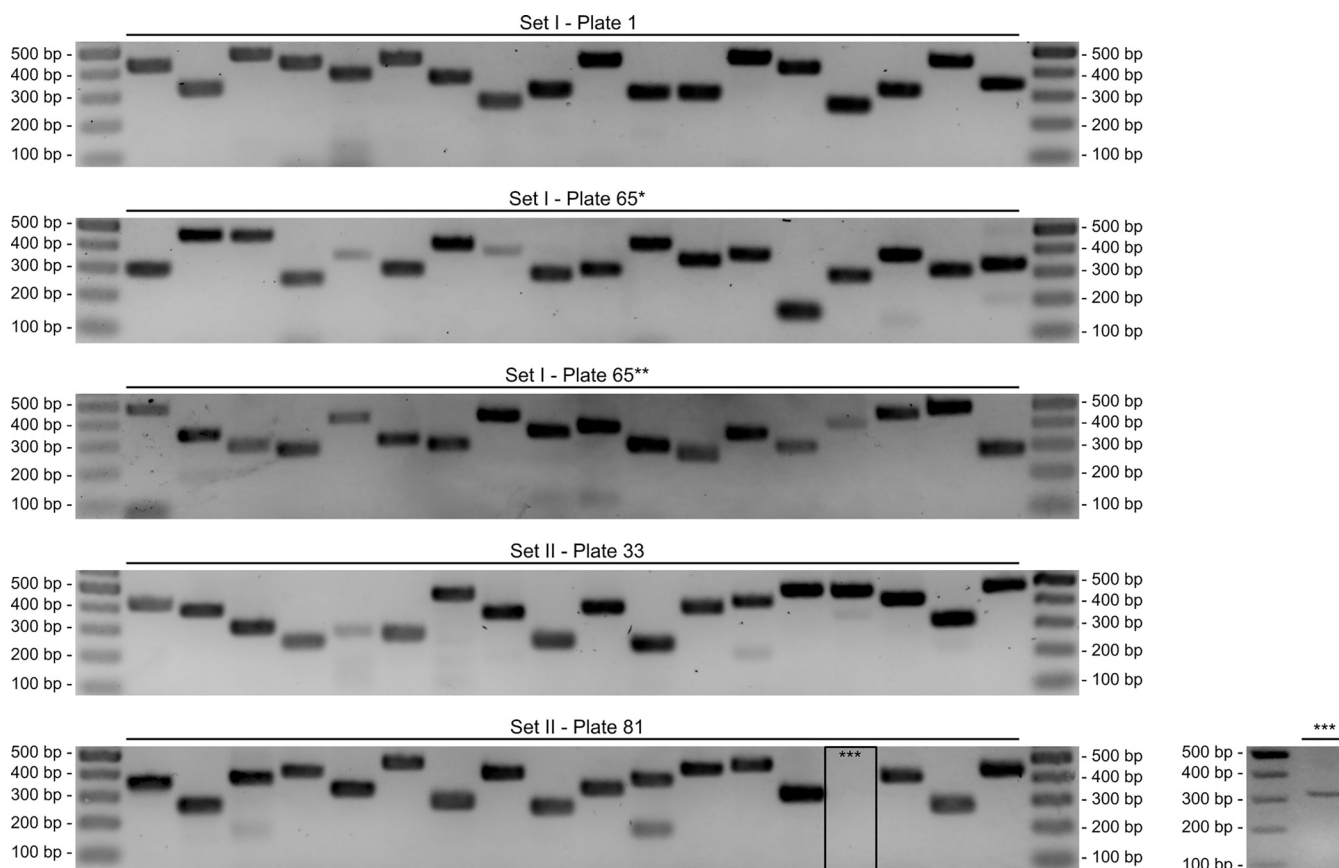


FIG 5 Positional validation of a subset of transposon insertion mutants by PCR. Ninety CP-CSeq assignments for transposon insertion mutants coming from 4 different 96-well plates were verified by PCR with a primer hybridizing to the transposon IR and a gene-specific primer. For plates I-1, II-33, and II-81, containing <96 mutants, we selected 18 transposon insertion mutants with a coverage of >300. For plate I-65, containing >96 mutants, we selected 18 transposon insertion mutants with a coverage of >1,200 (I-65*) and 18 transposon insertion mutants with a coverage of >300 and/or in wells with >1 mutant (I-65**). All CP-CSeq assignments were confirmed, except for 1 from plate II-81 (***). This PCR was successfully redone on a fraction of the glycerol stock (small gel on the right), so that all CP-CSeq assignments were confirmed. The list of mutants and their corresponding primers can be found in Table S3 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

chance of hitting increasing numbers of TA sites is akin to the “coupon collector’s problem” in statistics. To model what would be gained by further enlargement of CP-CSeq libraries, we fitted a coupon collector’s equation to the available experimental data from our libraries (Fig. 4d). As our combined library is very near the plateau of this empirical model, it is clear that further enlargement of the library would not be effective. Indeed, as expected, the nonessential genes that remained untargeted (371 in nonduplicated regions) were short genes and/or contain few TAs (Table S2 at <https://doi.org/10.6084/m9.figshare.c.4507472>), thus having much less chance to be mutagenized during transposon mutagenesis.

To validate the CP-CSeq assignments of transposon insertions to particular wells in the plate library, the positions of 90 transposon mutants from 4 different plates were verified via PCR, and all were correct (Fig. 5). Even though CP-CSeq had identified more than 96 transposon mutants in plate I-65, all 32 selected mutants of which we checked the position for this plate were confirmed via PCR, indicating that the robustness of our approach is high in the face of unavoidable picking of clonally impure colonies. Additionally, we confirmed that we could easily recover mutants in a clonally pure form in all cases by streaking the well contents for single colonies on agar and picking a few colonies for PCR (Fig. S2 at <https://doi.org/10.6084/m9.figshare.c.4507472>). This does not add much supplementary burden in practice, as in any case, it is good microbiological practice to streak for single colonies from culture collection resources prior to commencing further experimentation.

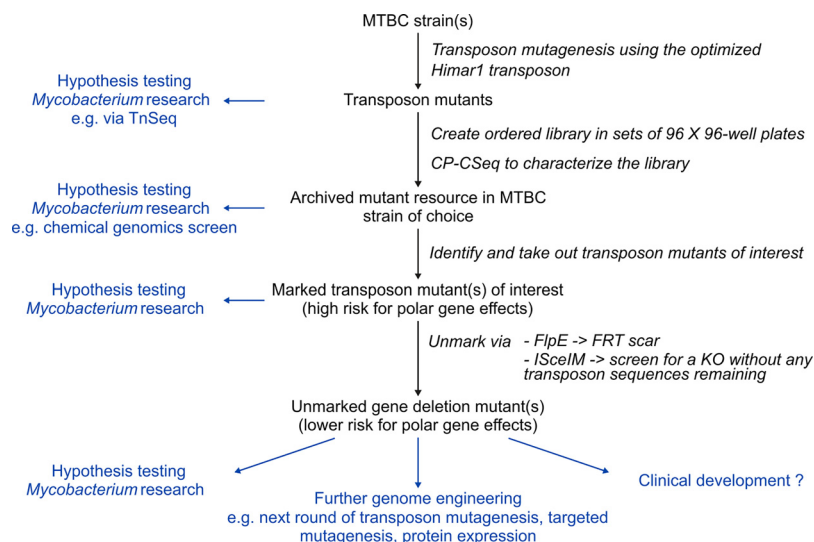


FIG 6 Applications of transposon mutagenesis using the optimized transposon. Shown is an overview of how an archived transposon library can be generated and how transposon mutants of interest can be isolated and unmarked. The scheme includes the different applications (in blue) of the generated products. MTBC, *Mycobacterium tuberculosis* complex.

DISCUSSION

Transposon mutagenesis has been used throughout *Mycobacterium* research to establish genotype-phenotype relationships and to investigate the genetic requirements of *in vivo* persistence of pathogenic bacteria such as *M. tuberculosis*. It is by far the most effective way of generating genome saturating mutant libraries. However, there are two disadvantages: (i) specific mutants cannot be easily retrieved from the mixed libraries, and (ii) the transposon-carried selection genes can trigger polar effects on neighboring genes. We solved the first problem previously by introducing CP-CSeq to characterize large clone-picked ordered 96-well plate collections obtained from transposon mutant libraries (9). By constructing an optimized Himar1 transposon and optimized efficient unmarking tools, we enabled the creation of clean, markerless mutants from marked transposon mutants to resolve the second problem. Combining this new transposon with the CP-CSeq protocol (creating archived mutant libraries) (9) now permits the creation of genome-wide mutant resources, from which markerless KOs can subsequently be efficiently generated (Fig. 6).

By incorporating *FRT* sites for FlpE/*FRT*-mediated recombination and *I-SceI* sites for *I-SceI*M-based transposon removal, we enable two possibilities to create unmarked mutants from marked transposon mutants. The first approach is highly efficient and inherently leaves an *FRT* scar, whereas the *I-SceI* system has the potential of creating clean, markerless mutants but requires clone screening and functional characterization of the unmarked mutants. Once the transposon insertion mutants are created, generating the KO counterparts no longer requires complicated cloning or transduction procedures but only a straightforward introduction of FlpE or *I-SceI*M (via electroporation or transduction). Therefore, this research contributes to the global goal of creating genome-wide KO mutants in multiple strains of the *M. tuberculosis* complex.

The *I-SceI* system, described here, provides a new way to remove the antibiotic resistance cassette in mycobacteria. Although the *I-SceI* system requires screening to identify the clones of interest (in contrast to the site-specific recombination systems, like the *FRT* system), it is the only available strategy that leaves no intact target recognition sites in the genome. As the *I-SceI* sites are cut and cannot be reformed, the unmarked organisms have no intact *I-SceI* sites remaining in the genome, excluding the risk for unwanted genomic rearrangements during iterative construction of multiple mutant strains (33). The *I-SceI* system can also be implemented in other mutagenesis

methods (two-step allelic exchange, specialized transduction [2, 13], or ORBIT [4], for which unmarking of mutants generated with these methods still leaves scars in the genome [34]) to create unmarked mutants.

We constructed an archived library of 2 sets of 96-by 96-well plates of optimized transposon mutants in the WHO reference strain *M. bovis* BCG Danish 1331 (NIBSC 07/270). Each of these transposon mutants can be unmarked to create markerless KOs for genes of interest. The created resource in BCG Danish (15,320 traceable transposon mutants) approaches genome saturation, as it targets 83% of all nonessential genes (excluding the duplicated genes). Applying (enhanced) specialized transduction (2, 3), ORBIT (4), or CRISPR-NHEJ-assisted genome editing (5) to target the remaining nonessential genes would be the most convenient option to complete the mutant resource, as the remaining genes are hard to target via transposon mutagenesis. For this biosafe BCG model, which is also the current TB vaccine and chassis for several of the improved BCG candidates, this goal now comes tantalizingly close. Only 371 nonessential genes remain to be targeted with targeted mutagenesis methods.

As a service to the TB research community, several research groups have made (or are making) the effort to create large mutant resources and make them publicly available in order to catalyze mycobacterial research all over the world. Indeed, transposon insertions resources in *M. tuberculosis* (35, 36) and other mutants were integrated by the TARGET program (Tuberculosis Animal Research and Gene Evaluation Taskforce [<http://webhost.nts.jhu.edu/target/>]), now offering over 4,300 defined transposon insertion mutants in *M. tuberculosis*. Furthermore, the lab of W. R. Jacobs, Jr. (in collaboration with others), took on the effort to construct a comprehensive, ordered library of mutants, each mutated for a nonessential gene, based on specialized transduction (2, 3, 37). Supporting these efforts, we have deposited both of our BCG mutant resources (BCG Pasteur library of 6,072 traceable transposon mutants and this study's BCG Danish library of 15,320 traceable, unmarkable transposon mutants) at the public BCCM-ITM Mycobacteria Collection (<http://bccm.belspo.be/about-us/bccm-itm>), making these mutants easily accessible to the scientific community. For further (pre)clinical development of these BCG mutants as vaccines, an in-depth phenotypic analysis (e.g., including transcriptional and/or proteomic studies) of the (unmarked) mutants is required. Our BCG mutant resources are currently the largest resources of mutants in any strain of the *M. tuberculosis* complex that are publicly available. Replicating the effort for other *M. tuberculosis* complex strains is feasible, and we hope that it will be pursued.

MATERIALS AND METHODS

Bacterial strains, media, plasmids, and phages. The *M. bovis* BCG Danish 1331 substrain (first WHO reference reagent, 07/270; NIBSC, Hertfordshire) and the streptomycin-resistant *M. smegmatis* strain mc²155 (38, 39) (a gift of P. Sander, Institute for Medical Microbiology, Zürich, Switzerland) were grown in Middlebrook 7H9 broth (Difco) supplemented with 10% Middlebrook oleic acid-albumin-dextrose-catalase (OADC; Becton, Dickinson), 0.2% glycerol, and 0.05% Tween 80 (Sigma). For electroporation of *M. bovis* BCG and *M. smegmatis*, cultures were grown in 7H9 broth supplemented with 0.05% Tween 80 and 10% homemade ADS (5 g/liter of bovine serum albumin [BSA], 2 g/liter of dextrose, and 0.85 g/liter of NaCl) but no glycerol. Difco Middlebrook 7H10 agar supplemented with 10% OADC and 0.5% glycerol was used for growth of *M. bovis* BCG on solid culture; LB agar (10 g/liter of BD Bacto tryptone, 5 g/liter of BD Bacto yeast, 5 g/liter of NaCl, and 20 g/liter of BD Bacto agar) was used for growth of *M. smegmatis* on solid culture. A list of all plasmids, shuttle phasmids, and phages used can be found in Table S4 at <https://doi.org/10.6084/m9.figshare.c.4507472>; detailed procedures for the construction of the plasmids, phasmids, and phages generated in this study can be found in Method S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>. Liquid or agar media were supplemented with 50 µg/ml of ampicillin, 50 µg/ml of kanamycin, 75 µg/ml of hygromycin, 2% or 5% sucrose, 0.2% or 0.5% 2-deoxy-galactose (2-DOG), 0.2% acetamide, or 100 ng/ml of tetracycline.

Transposon mutagenesis. Bacteria were grown to an optical density at 600 nm (OD₆₀₀) of 0.8 to 1.0 in supplemented 7H9 broth. Cells were washed 3 times with mycobacteriophage (MP) buffer (50 mM of Tris, pH 7.5, 150 mM of NaCl, and 2 mM of CaCl₂) and prewarmed to 39°C. Next, warm phage solution (39°C), or MP buffer as a negative control, was added to the warm bacilli. The transduction mixture was incubated for 4 h at 37°C. After incubation, the transduction mixture was plated on prewarmed (37°C) selective plates containing 50 µg/ml of kanamycin for incubation at 37°C. Additional details can be found in Method S2 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

Unmarking of transposon mutants via electroporation of unmarking plasmids. Bacteria were grown to mid-log phase (OD_{600} of 0.7 to 0.8) in 7H9 medium supplemented with Tween 80 and ADS. For *M. bovis* BCG cultures, 1.5% glycine was added to the cultures the day before electroporation. On the day of electroporation, cells were centrifuged and washed 3 times with 0.05% Tween 80 (in water). The cells were resuspended in the same buffer, after which the cells were mixed with the plasmid/phasmid (1 μ g) in a prewarmed 2-mm electroporation cuvette and electroporated (2.5 kV, 25 μ F, and 800 Ω). After recovery of the cells without antibiotics, the transformation mixture was plated on selective LB agar plates and incubated at 37°C (or at 32°C or 39°C when working with ts plasmids). The number of hygromycin-resistant clones was calculated as a measure for the efficiency of transformation. Details can be found in Method S3 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

Unmarking of transposon mutants via transduction of unmarking phages. Bacteria were grown to an OD_{600} of 0.8 to 1.0 in supplemented 7H9 broth. Cells were centrifuged, washed 3 times with MP buffer, and resuspended in the same buffer. Next, 500 μ l of $\sim 10^{10}$ PFU of warm phage suspension, or MP buffer as a negative control, was added to the bacilli. After incubation, the transduction mixture was plated on selective plates for incubation at 37°C. Details can be found in Method S4 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

Analysis of unmarked transposon mutants. The colonies obtained after unmarking were analyzed for the presence or absence of resistance markers by streaking the clones on agar plates containing different supplements. The percentage of unmarked clones (kanamycin sensitive and sucrose resistant) among the obtained sucrose-resistant colonies was calculated as a measure of the efficiency of unmarking. Unmarked transposon mutants were subsequently analyzed via colony PCR and/or Sanger sequencing. The used primers are listed in Table S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>, the primer sequences in Table S5 at <https://doi.org/10.6084/m9.figshare.c.4507472>, and additional details in Method S5 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

Curing mycobacteria of unmarking plasmids. Unmarked mycobacteria carrying an unmarking plasmid, containing a hygromycin-positive selection marker and/or a Galk-negative selection marker, were inoculated in 7H9 broth and subcultured before making serial dilutions and plating them on agar plates with or without 2-DOG. After incubation, single colonies (at least 20) were streaked on agar plates with or without hygromycin to check if the unmarking plasmid was cured. The percentage of cured clones (hygromycin sensitive) was calculated as a measure for the efficiency of curing. Details can be found in Method S6 at <https://doi.org/10.6084/m9.figshare.c.4507472>.

Construction and validation of the archived mutant library. The construction of the archived mutant library of two sets of 96- by 96-well plates in the *M. bovis* BCG Danish 1331 strain is based on the protocol by Vandewalle and others (9). For creating the transposon library, the medium was extra supplemented with 1% MD-VS vitamin supplement (ATCC), 0.2% Casamino Acids (Bacto), and 50 μ g/ml of L-Trp, to allow the inclusion of auxotrophic mutants in the library. The 192 96-well plates containing the transposon mutants were processed in two sets of 96- by 96-well plates. We recently established a reference genome sequence for *M. bovis* BCG Danish 1331 (NIBSC 07/270 [<https://doi.org/10.6084/m9.figshare.c.4489496>] [40, 41]), which was used to map the sequencing reads. A detailed procedure for the construction of the archived mutant library can be found in Method S7 at <https://doi.org/10.6084/m9.figshare.c.4507472>. The library size calculations are based on calculations described by Vandewalle and colleagues (9).

To verify the CP-CSeq assignment of the transposon insertion sites, frozen aliquots (from set I plates 1 and 65 and set II plates 33 and 81) were heated to 98°C for 45 min to release gDNA in the medium. After centrifugation, the supernatant was used in a PCR with a primer hybridizing to the transposon IR (Rv primer) and gene-specific primers (Fw primer) (Table S3 at <https://doi.org/10.6084/m9.figshare.c.4507472>). To analyze the clonal purity for a subset of transposon mutants, mutants were streaked on 7H10 agar plates supplemented with kanamycin, after which single clones were picked and grown in liquid medium. A PCR on supernatant of the BCG cultures and/or on gDNA of these single clones was performed with a unique primer set for the mutant (Table S6 at <https://doi.org/10.6084/m9.figshare.c.4507472>). PCR was performed using Phusion high-fidelity polymerase (New England BioLabs [NEB]), after which PCR products were run on a 1.2% or 2% agarose gel, stained with ethidium bromide, and visualized under UV light.

Data availability. Supplemental material accompanies this paper at the Figshare repository at <https://doi.org/10.6084/m9.figshare.c.4507472>, also including the Galaxy workflows for the bioinformatics data analysis and the updated BioPerl Script for the automated mutant location assignment (Zip File S1 at <https://doi.org/10.6084/m9.figshare.c.4507472>). The BCG Danish 1331 genome assembly as used during the data analysis can be found on Figshare at <https://doi.org/10.6084/m9.figshare.c.4489496> (40). The novel sequencing data (raw Illumina reads) generated in this study have been deposited at NCBI's Sequence Read Archive under project code [PRJNA528037](https://doi.org/10.6084/m9.figshare.c.4507472). The plasmids and phasmids generated in this study have been deposited in the BCCM/GeneCorner Plasmid Collection (<http://bccm.belspo.be/about-us/bccm-gene-corner>), their LMBP accession numbers are listed in Table S4 at <https://doi.org/10.6084/m9.figshare.c.4507472>. The remaining data that support the findings of this study are available on request from the corresponding author.

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We declare no conflicts of interest.

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K.B. designed and performed the experiments, analyzed the data, and cowrote the manuscript. K.V. assisted in experimental design and interpretation and constructed some vectors. G.M. constructed some plasmids and performed some unmarking experiments and curing experiments. A.V.H. aided in the transduction, growing, and pooling of the transposon insertion mutant library. E.P. aided in the plating after transduction. A.V.H., E.P., L.V.S., N.F., and S.V. assisted in picking the colonies. The Illumina library preparation and sequencing were performed by the VIB Nucleomics Core Facility. L.S. refined the BioPerl algorithm for automated location assignment. N.C. and N.F. assisted in experimental design and interpretation and cowrote the manuscript. N.C. conceived and supervised the project.

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