High-density transposon mutagenesis in *Mycobacterium abscessus* identifies an essential penicillin-binding lipo-protein (PBP-lipo) involved in septal peptidoglycan synthesis and antibiotic sensitivity

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Abstract

*Mycobacterium abscessus* (*Mab*) is a rapidly growing non-tuberculous mycobacterium (NTM) that causes a wide range of infections. Treatment of *Mab* infections is difficult because the bacterium is intrinsically resistant to many classes of antibiotics. Developing new and effective treatments against *Mab* requires a better understanding of the unique vulnerabilities that can be targeted for future drug development. To achieve this, we identified essential genes in *Mab* by conducting transposon-sequencing (TnSeq) on the reference *Mab* strain ATCC 19977. We generated ~51,000 unique transposon mutants and used this high-density library to identify 362 essential genes for *in vitro* growth. To investigate species-specific vulnerabilities in *Mab*, we further characterized *MAB_3167c*, a predicted penicillin-binding-lipoprotein (PBP-lipo) that is essential in *Mab* and non-essential in *M. tuberculosis* (*Mtb*). We found that PBP-lipo primarily localizes to the subpolar region and later to the septum as cells prepare to divide. Knockdown of *Mab* PBP-lipo causes cells to elongate, develop ectopic branches, and form multiple septa. Dual knockdown of PBP-lipo with PBPs, PbpB, DacB1, and a carboxypeptidase, MAB_0519 lead to synergistic growth arrest. In contrast, these genetic interactions were absent in the *Mtb* model organism, *M. smegmatis*, indicating that the PBP-lipo homologs in the two species exist in distinct genetic networks. Finally, repressing PBP-lipo sensitized the reference strain and 11 *Mab* clinical isolates to several classes of antibiotics, including the β-lactams, ampicillin and amoxicillin by greater than 128-fold. Altogether, this study presents PBP-lipo as a key enzyme to study *Mab* specific processes in cell wall synthesis and importantly positions PBP-lipo as an attractive drug target to treat *Mab* infections.
Introduction

*Mycobacterium abscessus* (*Mab*) is the most common cause of human disease among the rapidly growing non-tuberculous mycobacteria (NTM). It causes a wide range of illnesses including lung, skin and soft-tissue infections, as well as disseminated disease (1–3). While the incidence of *Mab* infections is rising worldwide, treating *Mab* infections remains difficult. The bacterium has intrinsic and acquired resistance mechanisms to many classes of antibiotics, including the standard drugs used to treat tuberculosis (4,5). The current treatment regimen for *Mab* requires taking a combination of multiple antibiotics for up to 18 months and is often associated with severe toxicity and routinely ends in treatment failure (6). Thus, there is an urgent need for new and effective drugs to treat the emerging global public health threat of *Mab* infections (7).

Developing novel antibiotic treatments for *Mab* requires a better understanding of essential biological processes in the bacterium that can be targeted. Identifying essential genes is the first step in this process, and by combining transposon-mutagenesis with massive parallel sequencing, transposon-sequencing (TnSeq) has been an effective tool to determine gene essentiality on a genome-wide scale in other microbes (8). However, while many TnSeq screens have been conducted in *M. tuberculosis* (*Mtb*) (9), up until recently only 2 published TnSeq screens have been reported in *Mab* (10,11). While both *Mab* TnSeq screens have provided tools and important insights into *Mab* biology, neither utilized a mutant library with high enough density to comprehensively identify essential genes. This changed with the recent publication from Rifat *et al*, who generated robust high-density transposon libraries of *Mab* and identified a comprehensive list of genetic elements essential for *in vitro* growth (12). Identification of these essential genes serves as an excellent starting point to classify attractive targets for drug discovery.
One major target of antibiotics is the mycobacterial cell wall whose biogenesis is essential for bacterial growth (13,14). The structure of the mycobacterial cell wall is unique among bacteria because it is composed of three macromolecules: peptidoglycan (PG), arabinogalactan, and mycolic acids. Together, these layers envelop mycobacteria with a thick, waxy coat that forms a permeability barrier to many drugs (15,16). The foundational layer is comprised of peptidoglycan (PG), which consists of glycan sugars cross-linked by short peptides forming a continuous, net-like structure that maintains cell shape and prevents rupture. Mycobacteria build new PG at the cell poles aided by Class A penicillin binding proteins (PBPs) that perform two reactions: a transglycosylation reaction that polymerizes PG monomers to existing glycan strands and a transpeptidation reaction that cross-links glycan strands to each other (17). Class B PBPs solely perform the cross-linking reactions (18,19). A hallmark of PBPs is that they are inhibited by β-lactam antibiotics; however, these antibiotics are typically not used to treat Mab infections due to high levels of resistance and the presence of β-lactamases in the genome (20).

To identify essential genes for in vitro growth and classify potential new drug targets in Mab, we performed TnSeq. We generated high-density transposon libraries of ~51,000 mutants in the reference strain, M. abscessus subsp. abscessus ATCC 19977 and identified 362 genes required for in vitro growth. We next identified essential genes in Mab that were non-essential in Mtb in order to study species-specific vulnerabilities in Mab. From this list of genes, we further characterized MAB_3167c, which encodes PBP-lipo, a conserved Class B PBP. We found that PBP-lipo is required for normal peptidoglycan (PG) synthesis, cell division, and sensitizes Mab to commonly-used β-lactams and other classes of antibiotics. None of these phenotypes were present in the Msm and Mtb PBP-lipo knockouts. Thus, we find that PBP-lipo has a unique essential function in Mab and is a potential drug target for treating Mab infections.
Results

Identification of *M. abscessus* ATCC 19977 essential genes for *in vitro* growth

To identify essential genes in *Mab*, we performed TnSeq on *M. abscessus subsp. abscessus* ATCC 19977 (hereafter referred to as *Mab*) using the ϕMycoMar T7 phage carrying the Himar1 transposon for transduction. This transposon inserts into ‘TA’ dinucleotide sites and disrupts gene function (Figure 1A) (21). We transduced three independently grown cultures of *Mab*, yielding triplicate libraries with saturations of 54.2%, 62.3%, and 64.1% for Library 1, 2, and 3 respectively (Supplemental Table 1). The average Pearson $r^2$ correlation of mean read counts for each gene across 2 libraries was 0.85 indicating good correlation among the libraries (Supplemental Figure 1). The transposon insertions across the three libraries were mapped onto the *Mab* genome as depicted in the chromosome map (Figure 1B). Across three libraries, we generated approximately 51,000 unique transposon mutants, as defined by sites of unique transposon insertions across all three libraries. Of note, the *Mab* libraries shared the same non-permissive ‘TA’ sites for transposon insertion as has been described in other mycobacterial species (Shoudhery *et al*, bioRxiv 2021).

The high density of these libraries allowed us to identify essential *Mab* genes using the Hidden Markov Model (HMM) algorithm implemented by TRANSIT (22). HMM identified 362 genes as essential, 4167 as non-essential, 153 as producing growth defect when disrupted, and 233 as causing a growth advantage when disrupted (Figure 1C, Supplemental Table 2). As expected, many of the genes identified as essential were involved in key biological processes including protein translation, amino acid metabolism, and cell wall biogenesis based on Clusters of Orthologous Groups (COG) analysis (Figure 1E).

Identification and Validation of uniquely essential genes in *Mab*
Understanding differences in essential genes between *Mab* and *Mtb* might uncover new ways of specifically targeting *Mab*. Thus, we identified *Mab*-specific essential genes by comparing gene essentiality between mutual orthologs of *Mab* and *Mtb*. We used *Mtb* essentiality data generated from a comprehensive analysis on *Mtb* essentiality using 14 high-quality TnSeq datasets (9). Of the 2279 mutual orthologs between *Mab* and *Mtb*, 282 genes were essential in both species. We focused on the mutual orthologs that that were essential in *Mab*, but non-essential in *Mtb*. These genes could potentially represent *Mab* specific drug targets. Exactly 20 genes fit into this category, which we refer to as “uniquely essential” (Figure 1D) (Supplemental Table 3).

To understand the quality of predictions by TnSeq, we validated the essentiality of a subset of genes using an anhydrotetracycline (ATc) inducible CRISPRi system developed for use in *Mtb* (23). Expression of a non-targeting sgRNA control in this system has no impact on cell proliferation (Figure 2A). In contrast, repressing canonical essential genes *rpoB*, *gyrB*, and *secY* impaired cell growth in both liquid broth (Figure 2A) and solid media (Supplemental Figure 2A), indicating that the CRISPRi system developed in *Mtb* functions in *Mab*.

We next validated the essentiality of a subset of the 20 uniquely essential genes in *Mab*: *glnA2*, *icd2*, *sdhA*, and *MAB_4471*. CRISPRi knockdown of these uniquely essential genes impaired growth in liquid media (Figure 2B) as well as solid media (Supplemental Figure 2B). These results confirm that at least a subset of uniquely essential *Mab* genes identified by TnSeq are required for optimal *in vitro* growth. The slight rise in OD$_{600}$ later in the time course is consistent with the emergence of escape mutants in the CRISPRi system, which has been previously described (24,25).

**PBP-lipo is uniquely essential in *Mab***
One of the uniquely essential genes we identified was \textit{MAB}_{3167c}, which encodes a Class B penicillin-binding lipoprotein (PBP-lipo). PBP-lipo is predicted to have a single N-terminal transmembrane helix and a transpeptidase domain. From our \textit{Mab} TnSeq data, \textit{MAB}_{3167c} lacked transposon insertions across all three replicate libraries (Figure 3A). This finding corroborates data from previously generated \textit{Mab} transposon libraries that show lack of insertions in \textit{MAB}_{3167c} (10,12). Comparing the essentiality of PBP enzymes in \textit{Mab} with their homologs in \textit{Mtb} revealed that \textit{pbp-lipo} is unique in being the sole PBP whose essentiality differs between the two species. Specifically, \textit{pbp-lipo} is essential in \textit{Mab} and non-essential in \textit{Mtb in vitro} (Figure 3B).

To validate that PBP-lipo is required for \textit{in vitro} growth, we used the CRISPRi system to induce knockdown using constructs encoding 1, 2, or 3 sgRNAs targeting \textit{pbp-lipo} (Supplemental Figure 3A). The single, double, or triple sgRNA constructs allowed for the fine tuning of the magnitude of \textit{pbp-lipo} knockdown, as measured by RT-qPCR (Supplemental Figure 3B). We noted that the empty vector control (0 sgRNA) and single sgRNA construct had no detectable effect on the growth of \textit{Mab} (Supplemental Figure 3C). In contrast, when \textit{pbp-lipo} was targeted with either 2 or 3 sgRNAs, the growth of \textit{Mab} was inhibited, indicating that \textit{pbp-lipo} is indeed required for optimal \textit{in vitro} growth (Figure 3C). We also observed the inhibition of cell growth on solid media where knockdown of PBP-lipo by 2 sgRNA and 3 sgRNA vectors led to a 1000-fold reduction in colony forming units (CFU) (Supplemental Figure 3D). To complement the PBP-lipo knockdown growth defect phenotype, we recoded a version of PBP-lipo with the protospacer adjacent motif (PAM) sequence mutated. This prevents sgRNAs from binding the introduced PBP-lipo copy present on the integrated plasmid. When constitutively expressed, this recoded PBP-lipo rescued growth of the native PBP-lipo knockdown (Supplemental Figure 3D). In contrast, a recoded and catalytically-dead mutant of PBP-lipo did not rescue growth. This data demonstrates that PBP-lipo’s transpeptidase activity is required for \textit{Mab} optimal growth.
In contrast to *Mab*, when we deleted the gene encoding PBP-lipo in *Mtb*, Rv2864c, the growth of the knockout and wild-type strains was indistinguishable (Figure 3D). Similarly, the PBP-lipo homolog in *Msm*, MSMEG_2584c, when knocked out grew identically to wildtype with no growth defect (Supplemental Figure 4). Taken together, these data demonstrate that PBP-lipo is essential for growth in *Mab*, while its homologs in *Mtb* and *Msm* are non-essential.

**PBP-lipo is required to maintain normal cell elongation and division**

A hallmark of PBPs is that they bind β-lactam antibiotics. Therefore, to test whether PBP-lipo is an active PBP in *Mab*, we determined whether PBP-lipo binds bocillin FL, a fluorescent analog of penicillin (26). Using a strain that constitutively expresses a strep-tagged PBP-lipo, we identified a fluorescent bocillin FL band at the same molecular weight as PBP-lipo, which was confirmed to be PBP-lipo by Western blot (Supplemental Figure 5). These data suggest that PBP-lipo binds bocillin FL in vivo and is an active PBP. Interestingly, in wildtype *Mab*, we did not detect an obvious PBP-lipo band at the predicted molecular weight of PBP-lipo, 63 kDa, suggesting that native expression levels of PBP-lipo is either low or PBP-lipo is expressed at distinct times during the cell cycle.

In mycobacteria, the PG layer is critical for maintaining normal cell shape and altering PBP expression often results in abnormal cell morphology. Thus, we studied the morphology of *Mab* when PBP-lipo was repressed and discovered dramatic morphological changes. *Mab* cells elongate, branch inappropriately, and form ectopic poles of active growth, as evidenced by staining with NADA, a fluorescent-D-amino acid (FDAA) that binds areas of active PG synthesis (27) (Figure 4A). PBP-lipo knockdown significantly increased the number of cells with ectopic branches (Supplemental Figure 6B). Furthermore, during normal growth of *Mab*, a small percentage of cells contain one mid-cell FDAA band, indicating active PG synthesis at the site
of septation. However, when PBP-lipo is knocked down, we occasionally find cells with 2 or 3 bands mid-cell, as exemplified in the 2 sgRNA + panel (Figure 4A). PBP-lipo knockdown strains were all significantly longer than their uninduced controls, with the longest average cell length belonging to the strain with the strongest PBP-lipo knockdown (Figure 4B). Of note, while the 0 sgRNA +/- cells are by eye morphologically indistinguishable, there is a statistically significant difference in the mean cell lengths, likely due to the effect of ATc on cells (Figure 4B). Notably, cell width did not change among the strains (Supplemental Figure 6A). In comparison, in \textit{Msm} and \textit{Mtb}, there were no differences in morphology and cell length between the wildtype and PBP-lipo knockout strains (Supplemental Figure 7). The dramatic changes in \textit{Mab} morphology when PBP-lipo is repressed suggests proper expression of this enzyme is required for maintaining normal \textit{Mab} elongation and division.

**PBP-lipo localizes to the subpolar regions and to the mid-cell before cell division**

The morphological defects caused by PBP-lipo knockdown were reminiscent of septal factor depletions observed in \textit{Msm} (28). Thus, we hypothesized that PBP-lipo localizes to the septum. To test this hypothesis, we N-terminally tagged PBP-lipo with mRFP, which has been previously shown to function in the mycobacterial periplasm (29). This mRFP-PBP-lipo construct was constitutively expressed and visualized with fluorescence microscopy. We discovered that PBP-lipo localized to the subpolar regions of the cell and the mid-cell, likely in the growing septum (Figure 5A). The resulting microscopy images were analyzed using OMEGA, which quantified the fluorescent signal across the cell normalized by cell length (Zhu \textit{et al}, unpublished). We then binned the cells into quartiles according to their length and found that the mRFP-PBP-lipo fluorescent signal was concentrated at the sub-polar regions for cells of all lengths, while the mid-cell localization of PBP-lipo was only present in the longest cells in the population (Figure 5B). These data suggest that as cells elongate, PBP-lipo localizes to the septum and is likely involved in septum formation and cell division.
PBP-lipo is incorporated into a unique PG network in *Mab* that does not exist in *Msm*

Given that *Mtb* and *Msm* do not have duplicates of PBP-lipo in their genomes, we hypothesized other reasons for the differential essentiality of PBP-lipo between these species. It is possible that there is a functional difference between PBP-lipo in *Mab* and its homologs in *Msm* and *Mtb*. This possibility intrigued us because an alignment of PBP-lipo from various mycobacteria revealed that PBP-lipo<sub>Mab</sub> has 5 additional amino acids present only in the *Mab* homolog (Supplemental Figure 8A). When *in silico* structures of PBP-lipo<sub>Mab</sub> and PBP-lipo<sub>Mtb</sub> are modeled, these 5 extra residues form an appendage we hypothesized could impart PBP-lipo<sub>Mab</sub> with additional ability to bind proteins (Supplemental Figure 8B).

To test this hypothesis, we constitutively expressed PBP-lipo<sub>Msm</sub> in the 2 sgRNA *Mab* strain (hereafter referred to as the *Msm* complement strain). As expected, knockdown of the native *Mab* PBP-lipo impairs growth. However, PBP-lipo<sub>Msm</sub> expression in the 2 sgRNA strain rescued *Mab* growth (Figure 6A). Furthermore, expressing PBP-lipo<sub>Msm</sub> eliminated the branching phenotype of the native PBP-lipo knockdown and partially reversed the elongated phenotype as well (Figure 6B). The recoded version of PBP-lipo<sub>Mab</sub> also reversed the branching of cells and partially complemented the elongated phenotype (Figure 6B). In contrast, the catalytically-dead version of PBP-lipo<sub>Mab</sub>-S364A did not complement, with cells appearing elongated and branched (Figure 6B, Supplemental Figure 9). These results demonstrate that PBP-lipo<sub>Msm</sub> can rescue knockdown of the endogenous *Mab* enzyme. From these data, we conclude that the essentiality of PBP-lipo<sub>Mab</sub> is not solely due to a unique functional property that is present only in the *Mab* homolog.

An alternative hypothesis is PBP-lipo participates in a PBP interaction network in *Mab* that is distinct from respective PBP networks in *Msm* and *Mtb*. In short, while PBP-lipo is required in...
the *Mab* PBP network, its function is likely redundant in the networks of *Msm* and *Mtb*. This
notion of functionally distinct PBP networks has been previously described in mycobacteria (30).

To interrogate the PBP network of *Mab*, we tested which PBPs genetically interacted with PBP-
lipo. To perform this experiment, we cloned CRISPRi constructs with sgRNAs that would weakly
knockdown each PBP in *Mab* (PBPx) so that growth was not significantly impaired. We also
cloned combinatorial CRISPRi constructs where PBP-lipo and another PBP were weakly
repressed. Using this dual sgRNA vector, we could simultaneously knockdown PBP-lipo and a
PBP of interest.

Using combinatorial knockdowns of each PBP, we compared growth inhibition when a single
PBP was knocked down versus when PBP-lipo and a second PBP were jointly repressed. Of
the 8 combinations we tested, we found that knockdown of three PBPs with PBP-lipo lead to
synergistic growth arrest (Figure 6C). These enzymes were *pbpB*, *dacB1*, and *MAB_0519*,
which encodes a protein with a hypothesized carboxypeptidase domain. This genetic synergy
was also observed in liquid culture for *pbpB* and *dacB1* (Supplemental Figure 10A). The
homolog of PbpB is a well-studied enzyme in both *Msm*, *Mtb*, and the model organisms *E. coli*
and *B. subtilis*, where it is known as Ftsl. In all these organisms, PbpB localizes to the septum
and is largely responsible for septal PG synthesis (31–33). Interestingly, like PbpB and PBP-
lipo, we found DacB1 localizes to the septum (Supplemental Figure 10B). These data suggest
that PBP-lipo genetically interacts with PbpB and DacB1, and that all three enzymes localize to
the same region of the cell and could potentially coordinate septal PG synthesis in a complex
together.

We next tested whether the same PBP-lipo genetic interactions existed in *Msm*. We used
CRISPRi to knockdown *Msm* PBP homologs in the wildtype *Msm* and the *pbp-lipo* knockout
background. In this experiment, genes that interacted with PBP-lipo would show impaired cell
growth when repressed in the PBP-lipo knockout. Interestingly, we did not observe genetic interactions between any of the PBPs tested and PBP-lipo (Figure 6D). This was true for the PBPs tested with homologs in *Mab* and additional PBPs present in the *Msm* genome (Supplemental Figure 11).

We next used a whole genome approach to identify genes in *Msm* that genetically interact with PBP-lipo. We generated high density transposon mutant libraries in both the wildtype and PBP-lipo knockout strains of *Msm* to identify genes that were conditionally essential in the knockout background. The saturation of the 2 wildtype libraries were 51.9% and 53.4%, while the saturation of the PBP-lipo knockout library was 62.4%. Using the resampling method in TRANSIT, our TnSeq data revealed that there were no conditionally essential genes in the ΔPBP-lipo *Msm* background (Supplemental Table 4). These data strongly suggest PBP-lipo in *Msm* exists in a robust genetic network with multiple functional redundancies. To uncover PBP-lipo’s function in *Msm*, it is possible that double and triple knockout mutants of PG synthesis enzymes would need to be generated.

**Repressing PBP-lipo sensitizes *Mab* to a wide range of antibiotics**

Given PBP-lipo’s essentiality in *Mab* and the genetic interactions it has with other PBPs, we hypothesized that repressing PBP-lipo would affect *Mab*’s sensitivity to antibiotics, including β-lactams. To test this hypothesis, we measured the sensitivity of PBP-lipo knockdown cells to 21 antibiotics that target translation, transcription, and DNA replication in both *Mab* and *Msm*.

Wildtype *Mab* is typically resistant to β-lactams amoxicillin and ampicillin with MICs greater than >256 µg/ml. When PBP-lipo is knocked down, the MIC for both antibiotics dropped to 2 µg/ml, indicating a greater than 128-fold sensitization (Table 1). These results indicate a strong
chemical-genetic interaction between PBP-lipo and both ampicillin and amoxicillin. Interestingly, not all β-lactams had shifts in their MIC when PBP-lipo was knocked down. This indicates that there is a specific chemical-genetic interaction between PBP-lipo and ampicillin and amoxicillin.

MIC changes for other β-lactams were less drastic, with a 4-8-fold reduction observed for faropenem and cefoxitin (Table 1). PBP-lipo knockdown also affects susceptibility to antibiotics that do not target the cell wall. The macrolides, clarithromycin and erythromycin, were 128-fold and 64-fold more potent respectively in the PBP-lipo knock down cells. A 16-fold reduction in MIC was observed for RNA polymerase inhibitor, rifampicin.

The MIC differences in rifampicin and clarithromycin are notable because these antibiotics do not target the cell wall. Thus, the mechanism of synergy between PBP-lipo repression and these antibiotics likely differs from the synergy with ampicillin and amoxicillin. We posited that PBP-lipo repression might increase the permeability of the cell wall, allowing more drug to enter the cell. To test this hypothesis, we measured the accumulation of calcein into Mab, which is proxy for cell permeability. Calcein accumulates more rapidly in PBP-lipo knockdown cells, indicating that PBP-lipo increases the permeability of Mab cells (Supplemental Figure 13). This increased permeability likely leads to increased antibiotic access to the cell, thus sensitizing the bacteria to the antibiotics.

When the MICs of the same 21 antibiotics tested in Mab were measured in the PBP-lipo knockout in Msm, we observed no difference in the MICs (Supplemental Table 5). We also measured the MICs of a subsection of these antibiotics in the pbp-lipo knockout in Mtb and found no difference in MICs between the wildtype and knockout strain (Supplemental Table 6). These data suggest that PBP-lipo’s unique function in Mab’s PG synthesis network likely contributes to the bacterium’s sensitivity to antibiotics when the enzyme is repressed.
Repression of PBP-lipo sensitizes *Mab* clinical isolates to antibiotics

The antibiotic experiments were conducted on the ATCC 19977 reference strain on *Mab*, which is lab adapted, after being isolated from a knee abscess in the 1950s and propagated in the lab since (3,34). Work performed in *Mtb* has demonstrated clinical isolates and lab-adapted strains respond to antibiotics stressors differently (35). As a result, we tested whether knockdown of PBP-lipo in *Mab* clinical isolates led to the same changes in antibiotic sensitivity as observed in the reference strain.

We found that knockdown of PBP-lipo using the 3 sgRNA construct impaired cell growth in all 11 clinical isolates leading to a 100 to 10,000-fold difference in CFUs (Supplemental Figure 12A). We also confirmed in a sub-set of the clinical isolates that PBP-lipo knockdown lead to elongated and inappropriately branched cells, as observed in the reference strain (Supplemental Figure 12B). After confirming the PBP-lipo phenotypes in the clinical isolates, we measured the MICs of ampicillin, amoxicillin, meropenem, rifampicin, and clarithromycin on *pbp-lipo* knockdowns of the clinical isolates.

For amoxicillin and ampicillin, PBP-lipo knockdown reduced the MICs by >32 to >512-fold.

Meropenem served as a negative control and indeed, there was no difference in MICs between uninduced and induced strains (Table 2). Finally, we observed an >8X-32X fold difference in the MICs for rifampicin and 8 to 128-fold difference in the MICs for clarithromycin (Table 3).

Altogether, the antibiotic sensitivity data demonstrate that the phenotypic consequences of repressing PBP-lipo is not only limited to the lab-adapted reference strain, but is also observed in *Mab* clinical isolates.
Discussion

Developing new drugs and drug regimens against Mab requires a better understanding of the bacterium’s essential processes. This includes identifying essential Mab genes for further characterization as potential drug targets. To do this, we generated high-density transposon libraries to identify essential genes in the reference strain, M. abscessus subsp. abscessus ATCC 19977. Importantly, Mab transposon libraries with fewer than 10,000 unique mutants have been previously published, but these studies did not fully identify the list of essential genes in Mab (10,11). Recently, Rifat et al successfully generated high-density libraries and were the first group to identify essential genes and genomic elements in the Mab genome (12). All together, they identified 326 essential genes and compared this list to essential genes identified in Mtb and Mycobacterium avium. Similar to our study, Rifat et al highlighted uniquely essential genes in Mab, which they also state deserve further study as these genes could represent Mab specific drug targets (12).

In addition to identifying essential genes in Mab, our study builds off of Rifat et al’s work by using CRISPRi to efficiently repress gene expression and validate predicted essential genes as required for in vitro growth (23). Furthermore, we found that by targeting genes with multiple sgRNAs, CRISPRi can achieve significant gene repression, allowing for the investigation of essential genes and providing an alternative to gene knockouts. Constructing gene knockouts in Mab can be inefficient (36); thus, CRISPRi provides a convenient tool for genetic manipulation in Mab. Overall, the newly generated Mab transposon libraries can be used to determine genes required for growth in a variety of conditions including infection models, biofilms, and antibiotic stress.

Both Rifat et al’s and our TnSeq data identified PBP-lipo as the sole peptidoglycan synthesis gene that was essential in Mab and non-essential in Mtb (12). Using CRISPRi, we
demonstrated the necessity of PBP-lipo expression for \textit{Mab} to maintain normal morphology and
to divide properly. When PBP-lipo is repressed, cells elongate, branch, and form ectopic poles.
We localized PBP-lipo to the subpolar regions of the cell, and as cells elongate and prepare for
division, PBP-lipo localizes to the septum. This constellation of phenotypes mimic septal-factor
depletions observed in \textit{Msm}, which also causes cells to elongate, branch, and form ectopic
poles (28). These data suggest that PBP-lipo is involved with PG synthesis at the septum and
that depletion of the enzyme likely leads to dysregulated septal PG synthesis and aberrant cell
division.

Given PBP-lipo’s localization and effect on cell division, we posit PBP-lipo acts as an essential
septal PG synthesis enzyme in \textit{Mab}. In \textit{Mtb} and \textit{Msm}, this is accomplished by the enzyme
\textit{pppB}, which is essential for growth. Additionally, when \textit{pppB} is repressed in these species, cells
appear morphologically similar to the PBP-lipo knockdown in \textit{Mab} (28,31). Interestingly, PBP-
lipo knockouts in both \textit{Msm} and \textit{Mtb} are morphologically identical to wildtype cells
(Supplemental Figure 7). These data underscore the notion that biological insights from \textit{Msm}
and \textit{Mtb} should not be de-facto applied to \textit{Mab}. We hypothesize that PBP-lipo is a member of
the \textit{Mab} divisome complex, a collection of structural proteins and enzymes that coordinates cell
division (37). Future studies are needed to determine if PBP-lipo interacts with known septal
factors, such as FtsQ, FtsW, FtsZ, whose functions have been described in \textit{Msm} (38–40).
Future work may also reveal novel PBP-lipo binding factors that coordinate PG synthesis and
cell division in \textit{Mab}.

Phenotypes characteristic of the PBP-lipo knockdown in \textit{Mab} were absent in the \textit{Mtb} and \textit{Msm}
PBP-lipo knockouts. Both \textit{Mtb} and \textit{Msm} do not have a duplicate copy of PBP-lipo in their
genomes. Thus, to explain the differential essentiality of PBP-lipo across these mycobacterial
species, one hypothesis we explored was differences in the genetic networks of PG synthesis
enzymes. Construction of the PG layer is a complex process that involves a collection of enzymes that work in concert to ensure proper synthesis and remodeling. Previous work has shown that PG synthesis enzymes belong to genetic and spatiotemporal networks that do not necessarily replicate across closely related species (30,41). Thus, while the building blocks to construct the PG layer may be similar between species, how the enzymes are networked and subsequently function together may differ. We demonstrated that PBP-lipo genetically interacts with *pbpB*, *dacB1*, and *MAB_0519*. In contrast, TnSeq of the *Msm Δpbp-lipo* strain did not reveal interactions with any other gene in the genome. The TnSeq data demonstrates that despite being homologs, PBP-lipo in *Mab* and *Msm* are incorporated into different PG synthesis networks. In *Mab*, this network renders PBP-lipo’s function as essential for growth, whereas in *Msm*, PBP-lipo is dispensable. Interestingly, in *Mtb* PBP-lipo became more required in a *ponA2* knockout suggesting that PBP-lipo and PonA2 genetically interact *Mtb* (30) – a genetic interaction that was not observed in *Mab*. This observation supports the hypothesis that PBP-lipo is incorporated in different PG synthesis in *Mtb* as well.

In *Mab*, the three genes that genetically interacted with PBP-lipo, *pbpB*, *dacB1*, and *MAB_0519*, are all hypothesized to be septal associated. PbpB is a D,D-transpeptidase that performs 4,3 crosslinking reactions at the septum. In both *E. coli* and *Msm*, depleting PbpB leads to cell elongation and branching, reminiscent of the *Mab* PBP-lipo knockdown phenotypes (28). A potential explanation of this genetic interaction is both PbpB and PBP-lipo perform 4,3-crosslinking at the *Mab* septum. In this model, repressing both enzymes would dramatically reduce 4,3-crosslinking at the septum and lead to cell death. It is possible that PBP-lipo and PbpB have overlapping 4-3 cross-linking function at the septum. However, given that PBP-lipo and PbpB are essential in *Mab*, both enzymes must have distinct functional roles. Their distinct functions could be revealed by identifying unique genetic interactions each enzyme has with...
other PG synthesis enzymes. Why *Mab* would require two PG 4,3 crosslinking enzymes functioning at the septum is an intriguing area of future research.

The two other genes that synergized with PBP-lipo, DacB1 and MAB_0519, are both hypothesized carboxypeptidases that cleave the terminal D-ala-D-ala on the pentapeptide chains of PG monomers (42,43). This reaction is necessary for the subsequent 4,3 crosslinks catalyzed by D-D transpeptidases like PbpB and PBP-lipo. Interestingly, DacB1 localizes to the septum in *Mab* and has been described as septally localized in *Msm* and *Mtb* (44). Furthermore, the homolog of MAB_0519 in *Mtb*, Rv3627c is involved in septal PG synthesis (45). Thus, we hypothesize DacB1 and MAB_0519 are involved in septal PG synthesis and may work with PBP-lipo to ensure proper septum formation. Interestingly, DacB1 and MAB_0519 are not functionally redundant, given that both genes are synergistic with PBP-lipo. Further characterization of the PG enzymes in *Mab* and their genetic networks will uncover species-specific nuances in *Mab* PG synthesis that have the potential to be exploited for drug discovery.

Knockdown of PBP-lipo dramatically sensitized the reference *Mab* strain and 11 clinical isolates to several antibiotics. The PBP-lipo knockdown was greater than 128-fold more sensitive to β-lactams, ampicillin and amoxicillin, while other β-lactams like meropenem, cephalexin, ticarcillin and others showed no difference in MIC. This result indicates a specific chemical-genetic interaction between ampicillin/amoxicillin, and PBP-lipo. Historically, β-lactams have not been used to treat *Mab* infections due to high levels of resistance and the presence of β-lactamases (20). While imipenem and cefoxitin are used with some success, macrolides and aminoglycosides remain the backbone of *Mab* treatment (46). Recently, studies have shown dual β-lactam strategies are effective in killing *Mab* both *in vitro* and *in vivo* (47–49). These synergy data suggest that understanding the specific antibiotic targets of β-lactams in *Mab*
would augment the design of effective dual β-lactam combinations. At the moment, effective combinations are tested empirically with no insight into potential mechanisms (48). To address this, Sayed et al used a combination of a bocillin FL binding assay to stain all the PBPs in *Mab* followed by proteomics to identify the targets of 12 β-lactams and 2 β-lactamases. Interestingly, in their datasets, PBP-lipo was not identified by mass spectrometry of bocillin FL-bounded PBPs (50). This further supports our observation that PBP-lipo is expressed at low-levels in *Mab* and is not identifiable by bocillin FL under native expression conditions.

Nonetheless, our data demonstrate that using a β-lactam or small molecules that target PBP-lipo would not only kill the cell, but also potentiate β-lactams like ampicillin and amoxicillin. In addition, the permeabilizing effect of PBP-lipo inhibition would potentiate drugs by allowing more to enter the cell. Crucially, the observed synergy with PBP-lipo knockdown and β-lactams was not just limited to the reference strain, but also present in 11 clinical isolates of *Mab*, indicating that this synergy can be targeted for use in *Mab* clinical isolates.

Overall, this work employed TnSeq to identify the uniquely essential gene PBP-lipo in *Mab*. We present PBP-lipo as a promising drug target given both its essentiality and its role in sensitizing *Mab* to a range of antibiotics, including commonly used and accessible antibiotics, ampicillin and amoxicillin. Future studies exploring the PBP networks of mycobacteria can expose further species-specific vulnerabilities in peptidoglycan synthesis. Finally, identifying chemical inhibitors of PBP-lipo may help form the foundation for new treatments against this emerging and difficult to treat pathogen.
Materials and Methods

A full list of the strains, plasmids, and primers is available in Supplemental Tables 7, 8, and 9 respectively.

Bacterial Strains and Culture Conditions: All *M. abscessus* (*Mab*) and *M. tuberculosis* (*Mtb*) strains were grown shaking at 37°C in liquid 7H9 media consisting of Middlebrook 7H9 salts with 0.2% glycerol, 0.85 g/L NaCl, OADC (oleic acid, 5 g/L albumin, 2 g/L dextrose, 0.003 g/L catalase), and 0.05% Tween80. *Mab* and *Mtb* were plated on Middlebrook 7H10 agar supplemented with 0.5% glycerol. *M. smegmatis* mc2 155 (*Msm*) was grown shaking at 37°C in liquid on 7H9 media consisting of Middlebrook 7H9 salts with 0.2% glycerol, 0.85 g/L NaCl, ADC, and 0.05% Tween80 and plated on LB agar. Antibiotic selection concentrations for *Mab* cultures were kanamycin 50 µg/µl and zeocin 100 µg/µl. For transposon mutant selection, 100 µg/ml of kanamycin was used. Antibiotic selection concentrations for *Msm* and *Mtb* cultures were kanamycin 25 µg/µl, zeocin 20 µg/µl, and gentamicin 5 µg/µl. For *E. coli* (TOP10, XL1-Blue and DH5α), all antibiotic concentrations were twice those of the concentrations used for *Msm* and *Mtb*. Induction of all CRISPRi plasmids was performed with 500 ng/ul anhydrous tetracycline (ATc).

Strain Construction

Construction of 0, 1, 2, 3 sgRNA PBP-lipo knockdown strains and PBP-lipo complement strains

To build the 0 sgRNA (empty sgRNA) *Mab* strain, we transformed the pCT296 vector containing an empty sgRNA into *Mab*. To build the 1, 2, and 3 sgRNA strains, first we built individual CRISPRi plasmids containing single sgRNAs that targeted PBP-lipo. These plasmids were cr25, cr26, and cr56. We then designed primers that amplified the sgRNA targeting region with the appropriate flanking sequences and performed Golden Gate cloning to combine the inserts into one vector. This method was previously described by Rock et al (23). For the 1 sgRNA strain,
we transformed cr25 into the Mab strain. For the 2 sgRNA, we transformed the vector pCRC1, which contained the cr25 and cr26 sgRNAs on one backbone. Finally, for the 3 sgRNA strain we transformed pCRC2, which contained the cr25, cr26, and cr56 sgRNAs on one backbone. To build the PBP-lipo complement strain, we designed primers to amplify PBP-lipo with modified protospacer adjacent motif (PAM) sequences that were no longer homologous to the 1, 2, and 3 sgRNAs. This recoded PBP-lipo was then cloned into a constitutive expression vector resulting in the pCA323 construct, which was transformed into the 2 sgRNA strain. The catalytically dead mutant version of the recoded PBP-lipo was generated by using primers to generate the S364A mutation. This engineered PBP-lipo was cloned into a constitutive expression vector and the resulting construct, pCA324 was transformed into the 2 sgRNA strain.

Msm and Mtb ΔPBP-lipo strains

To generate PBP-lipo knockout mutants in Msm and Mtb, we used a recombineering approach to replace the endogenous copy of the gene with a zeocin resistance cassette as previously described (51). First, 500 base pairs of upstream and downstream sequence surrounding PBP-lipo (Rv2864c and MSMEG_2584c) were amplified by PCR KOD Xtreme™ Hot Start DNA polymerase (EMD Millipore, Billerica, MA). These flanking regions were amplified with overlaps for the zeocin resistance cassette. The 2 flanking regions and zeocin resistance cassette were then assembled via isothermal assembly (52) into a single construct, which was then amplified by PCR. Each deletion construct was transformed into either Mtb or Msm expressing inducible copies of RecET for recombination (53). The resulting colonies were screened for PBP-lipo deletion by PCR.

PBP-lipo_{Mab-strep expression strain}
PBP-lipomab was amplified using primers that added a C-terminal strep tag to the protein. The resulting insert was Gibson stitched to the vector backbone that drove expression of PBP-lipomab-strep from the constitutively active MOP promoter.

mRFP-PBP-lipomab

We amplified PBP-lipomab in two fragments. The first fragment contained the N-terminal signal sequence of PBP-lipo. The second fragment contained the remainder of the protein. We then amplified mRFP with overhangs that were compatible with the two halves of PBP-lipo. The resulting three fragments were Gibson assembled to a vector backbone that constitutively drove expression of the mRFP-PBP-lipomab fusion protein. A gly-gly-ser-gly-ser-gly linker was used in between mRFP and the new N & C terminus of PBP-lipo.

DacB1-GFPMut3

DacB1 lacking a stop codon was amplified along with GFPMut3 containing overlaps with DacB1 and a gly-gly-ser linker. The two fragments were Gibson assembled to a vector backbone that constitutively drove expression of the DacB1-GFPMut3 fusion protein.

Growth Curve: Mab, Msm, and Mtb cultures were grown to log phase and diluted to OD600 0.05. Growth curves were performed in a 96 well format at 37°C with shaking. OD600 was measured every 15 min for 24-72 hr using the TECAN plate reader

Colony Forming Units: Mab and Msm cultures were grown to mid-log phase and then serially diluted in their respective media in 96 well plates. Dilutions were then spot plated onto solid media +/- 500 ng/ul ATc. Plates were incubated at 37°C for 4 days (Mab) or 3 days (Msm). After incubation, the resulting CFUs were calculated by counting the number of colonies at the appropriate dilution. The fraction survivability was calculated as CFUs +ATc / -ATc.
Minimum Inhibitory Concentration Determination: Mab, Msm, and Mtb were grown to mid-log phase and diluted to an OD600 = 0.003 in each well of non-treated 96-well plates (Genesee Scientific) containing 100 μL of antibiotic serially diluted in 7H9 + OADC + 5 μg/mL clavulanate (Sigma Aldrich). For MICs on Mab knockdown cells, cultures were induced for knockdown 18 hr prior with 500 ng/ul ATc. Wells with knockdown bacteria also contained 500 ng/ul ATc. Msm media contained ADC rather than OADC. Cells were incubated with drug at 37°C with shaking for 1 day (Mab, Msm) or 7 days (Mtb). Afterwards, 0.002% resazurin diluted in ddH2O (Sigma Aldrich) was added to each well. Plates were then incubated for 24 hr. MICs were determined by the concentration of antibiotic that turned wells blue signifying no metabolic activity (54).

Generation of Transposon Libraries: To create transposon mutant libraries of Mab ATCC 19977 and Msm (wildtype and ΔPBP-lipo), 100 ml of bacterial cultures were grown to an OD600 of 1.5 – 2.0. For Mab, cultures were grown in biological triplicate. For Msm, the wildtype strain was grown in duplicate while one replicate of the ΔPBP-lipo strain was grown. The cells were then washed twice with 50 ml of MP Buffer (50mM Tris-HCl pH 7.5, 150 mM NaCl, 10 mM MgSO4, 2 mM CaCl2) and resuspended in 1/10th the culture volume in MP Buffer. Afterwards, 2 x 1011 pfu of temperature sensitive φMycoMarT7 phage carrying the Himar1 transposon were added to bacteria. Phage and bacterial cultures were incubated at 37°C for 4 hr with shaking. The transduced cultures were spun down at RT 4000 rpm for 10 min and resuspended in 12 ml of PBS-Tween. The cultures were then tittered by plating on 7H10 plates (Mab) or LB (Msm) supplemented with kanamycin 100 μg/μml (Mab) or 20 μg/μl (Msm) before freezing the aliquots for future plating. After determining the titer, 150,000 bacterial mutants of each transduced culture were plated onto kanamycin selective plates and grown for 4 days at 37°C. The resulting mutant libraries were harvested and stored in aliquots with 7H9 + 10% glycerol at -80°C.
Genomic DNA Extraction: To prep libraries for sequencing, we isolated the gDNA of the mutant libraries using a bead-beating protocol developed in the lab. 2-6 ml of transposon mutant libraries were thawed and spun down to remove excess glycerol in the sample. Pellets were then resuspended in TE Buffer (10 mM Tris HCl pH 7.4, 1 mM EDTA pH8) and placed in bead-beating tubes with 600 µl of phenol:chloroform. The samples were then bead-beat 4X for 45 s at 4000 rpm. Samples were cooled on ice for 45 s between each successive bead-beat round. After bead-beating, the samples were spun down for 10 min at 13,000 rpm. The supernatant was transferred to a new falcon tube and incubated with 1:1 volume phenol-chloroform for 1 hr rocking. Samples were then spun down in MaXtract High Density phase-lock tubes (Qiagen) at 1500 g for 5 min. This separated the aqueous and organic layers. Afterwards, ½ volume of chloroform was added to the tube and the contents were spun down at 1500 g for 5 min. The aqueous phase was transferred to a new tube and RNase A (Thermo Fisher) was added to a final concentration of 25 µg/µl. The tubes were incubated at 37°C for 1 hr with shaking. Finally, the supernatant was washed with 1:1 volume of phenol-chloroform followed by ½ volume of chloroform. The aqueous phase was transferred to a new tube and genomic DNA was precipitated with 1 volume of isopropanol alcohol and 1/10th volume of 3 M sodium acetate pH 5.2. The library gDNA was washed twice with fresh 70% ethanol and resuspended in nuclease free ddH₂O. This protocol consistently generated high yields of Mab gDNA.

Transposon sequencing, mapping, and analysis: Sequencing libraries were prepared from the extracted gDNA by amplifying chromosomal-transposon junctions following the protocol as described by Long et al, 2015 (21). These amplicons were then sequenced using the Illumina Hi-Seq platform. The resulting reads were mapped onto the Mab or Msm genomes. The data were analyzed using the TRANSIT pipeline (22). Insertion counts at each TA site were normalized using TTR normalization and averaged across replicates. Essentiality categories of genes were determined by the Hidden Markov Model (HMM) algorithm in TRANSIT.
Resampling analysis was used to compare insertion accounts between genes in the wildtype and ΔPBP-lipo Msm strains.

**Cloning of CRISPRi Constructs:** To clone sgRNAs onto the CRISPRi vector backbone, complimentary targeting oligos were annealed and ligated (T4 DNA ligase, NEB) into the BsmBl digested CRISPRi vector backbone. To clone multiple sgRNAs into the same vector, the promoter, sgRNA handle, and sgRNA sequences were amplified with the appropriate flanking sequences to perform Golden Gate cloning using the SapI cloning site in the CRISPRi vector (23). All cloning was performed using DH5a cells and sequences were verified before transformation into Mab and Msm.

**Transformation of M. abscessus:** To generate electrocompetent Mab cells, 50 ml of Mab cultures were grown to mid-log phase. Cells were spun down at 4000 rpm for 10 min and washed twice with pre-chilled 10% glycerol. Cell pellets were resuspended in 1/100th of the starting volume and kept on ice for immediate use or stored at -80°C for future use. For transformations ~100 ng of plasmid DNA was incubated with electro-competent cells for 5 min before transforming the cells with 2500 V, 125 Ω, 25 μF. Afterwards, cells were recovered in 1 ml of 7H10 with OADC without selection for 3 hr at 37°C with shaking. Cells were then plated on 7H10 with the appropriate selection marker and incubated for 4 days at 37°C.

**Microscopy and Image Analysis:** All imaging was performed on an inverted Nikon TI-E microscope at 60X and 100X magnification. For PBP-lipo knockdown experiments, cultures were induced for knockdown with 500 ng/ul ATc for 18 hr prior to imaging. All Mab cultures were fixed with 7H9 + 3% paraformaldehyde (PFA) for 1 hr and resuspended in PBS + 0.05% Tween80 prior to imaging. Cellular features including cell length, width, and fluorescence signal...
were analyzed using the OMEGA image analysis pipeline developed in the lab (Zhu et al, unpublished).

**Fluorescent D-Amino Acid Staining:** NADA (3-[(7-Nitro-2,1,3-benzoxadiazol-4-yl)amino]-D-alanine hydrochloride) was synthesized by Tocris following the published protocol (27). To stain *Mab* with NADA, 0.1 mM of FDAA was added to 1 ml of exponentially growing cells and incubated for 3 min before washing in warm 7H9 twice. For *Mab* imaging, after the second wash, cells were fixed in 7H9 + 3% paraformaldehyde for 1 hr. The PFA was then washed off and cells were resuspended in PBS + 0.05% Tween80 prior to imaging.

**RNA Extraction and RT-qPCR:** For each strain, cultures were grown in biological triplicate to mid-log phase and then diluted back in +/- 500 ng/ml ATc and grown for 18 hr to achieve target knockdown. Afterwards, 2 OD600 equivalents of cells from each culture were harvested by centrifugation, resuspended in TRIzol (Thermo Fisher), and lysed by bead beating (Lysing Matrix B, MP Biomedicals). Total RNA was extracted using the RNA miniprep (Zymo Research). Residual genomic DNA was digested with TURBO DNase (Ambion), and samples were cleaned with RNA clean-up columns (Zymo Research). cDNA was prepared using random hexamers following manual instructions (Life Technologies Superscript IV). Alkaline hydrolysis was then used to remove RNA. cDNA was purified by spin column (Qiagen) and then quantified by real-time quantitative PCR (RT-qPCR) on a Viia7 light cycler (Applied Biosystems) using iTaq Universal SYBR Green Supermix (BioRad). All qPCR primer pairs were confirmed to be >95% efficient. The masses of cDNA used were experimentally validated to be within the linear dynamic range of the assay. Signals were normalized to the *sigA* (*MAB_3009*) transcript and quantified by the ΔΔCt method. Error bars are standard deviations from three biological replicates.
Bocillin FL staining: *Mab* cultures were grown to mid-log phase and 10 OD units of bacteria were spun down and resuspended in 1 ml of fresh 7H9 media with 5 µg/ml of clavulanate. Bocillin-BODIPY or FITC were added to cultures to a final concentration of 10 µM. Cells were then incubated at 37°C for 2.5 hr, with aluminum wrapping to prevent light exposure. After labeling, cells are washed with PBS + 0.05% Tween80 three times and resuspended in 200 µl of lysis buffer (50 mM Tris-HCl pH = 7.4, 50 mM NaCl, protease inhibitor (Roche)). Samples were bead beat 4X at 4000 rpm for 1 min with 1 min of rest on ice in between rounds of bead-beats. CaCl$_2$ was added to a final concentration of 1 mM along with 4U of DNAse I and 10 mg/ml of fresh lysozyme. Samples were incubated at 37°C for 30 min. Afterwards, supernatants were normalized by A280 protein concentration, diluted with 6X Laemmli buffer, and run on 4%–12% NuPAGE Bis Tris precast gels (Life Technologies). To visualize fluorescence, we used a BioRad Gel Doc system. To detect the presence of PBP-lipo-strep, we performed a Western Blot. Membranes were blotted with rabbit α-Strep (GenScript) at 1:1000 in TBST + 3% BSA.

Calcein Staining: For each strain, cultures were grown in biological triplicate to mid-log phase and diluted back in +/- 500 ng/ml ATc and grown for 18 hr to achieve target knockdown. Afterwards, 3 OD units of bacteria were washed twice with PBS + 0.05% Tween80. Cells were resuspended in PBS + 0.05% Tween80 and 100 µl of cells at OD600 = 0.4 was added to 96 well plates. The plates were incubated in the plate reader for 30 min at 37°C with shaking. Afterwards, calcein was added at a final concentration of 1 µg/ml. Fluorescence signal was measured every minute using the Tecan plate reader.
Acknowledgements

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References


Figure 1. Overview of TnSeq Analysis. (A) Schematic of TnSeq protocol. Phage carrying Himar1 transposon is used to transduce *Mab* ATCC 19977 cultures. Over 51,000 independent transposon mutants were generated across three libraries (B) Location of transposon insertions in *M. abscessus* subsp. *abscessus* ATCC 19977 genome. Transposon insertions represent an average of the three replicates. Black bars represent number of insertions at each TA site. (C) Breakdown of gene essentiality categories. Essentiality calls were determined using the Hidden Markov Model (HMM) in TRANSIT (D) Essentiality comparison of mutual orthologs between *Mab* and *Mtb*. Venn Diagram shows 282 mutual orthologs are essential in both species. 20 essential *Mab* genes have mutual orthologs in *Mtb* that are non-essential. (E) COG (Clusters of Orthologous Group) categories of essential genes in *Mab*. 

**Figures and Legends**
Figure 2. Validation of Essential Genes. (A) Growth curve of *Mab* strains with sgRNAs targeting canonical essential genes. (B) Growth curve of *Mab* strains with sgRNAs targeting uniquely essential *Mab* genes.
Figure 3. PBP-lipo is uniquely essential in Mab and non-essential in Mtb. (A) Transposon-insertion profile of PBP-lipo across three replicate transposon libraries. Red bars indicate the number of transposon insertions. Horizontal black lines indicate ‘TA’ sites, where the transposon inserts. Genes are schematized with gray arrows, with PBP-lipo colored in blue. Dotted box demarcates lack of insertions in PBP-lipo. (B) Comparison of essentiality for PBP orthologs between Mtb and Msm. (C) Growth of Mab cultures transformed with CRISPRi plasmid carrying either 2 or 3 sgRNAs targeting PBP-lipo. “-” symbol indicates cultures were grown without ATc. “+” symbol indicates cultures were grown with ATc. (D) Growth of wildtype Mtb and PBP-lipo knockout strains.
Figure 4. Knockdown of PBP-lipo disrupts cell morphology. (A) Microscopy images of PBP-lipo knockdown cultures. *Mab strains carrying CRISPRi plasmids with either 0, 1, 2, or 3 sgRNAs targeting PBP-lipo. Arrows indicate sites of multiple septa formation. (B) Cell lengths of uninduced and induced strains. Measurements were obtained by image pipeline, OMEGA (Zhu et al, unpublished). Student’s t test used to calculate the statistical difference in mean cell lengths. *** = p-val < 0.0001. * = p-val < 0.05.
Figure 5. PBP-lipo localizes to the subpolar region and the septum. (A) Microscopy images of PBP-lipo N-terminally tagged with mRFP. White arrows demarcate subpolar and septal localization of PBP-lipo. (B) Quantification of mRFP signal along the length of the cell using OMEGA imaging pipeline (Zhu et al., unpublished). Signal was normalized by cell length and cells were binned into quartiles based on length.
Figure 6. PBP-lipo network in Mab does not exist in Msm. (A) Growth curve of 2 sgRNA strain targeting PBP-lipo and 2 sgRNA strain with a constitutively expressed copy of PBP-lipo in Msm. (B) Microscopy images of 2 sgRNA and PBP-lipo complement strains. (C) Genetic synergy of PBP-lipo with other PBPs in Mab. Solid bar represents CFU of strains where a single PBP was knocked down. Open bar represents strains where PBP-lipo was knocked down in combination with the listed PBP. (D) Genetic synergy of PBPs and PBP-lipo in Msm. CRISPRi plasmids carrying sgRNAs targeting each of the PBPs in Msm were transformed into wildtype Msm (closed bar) and the ΔPBP-lipo mutant (open bar).
Table 1: Minimum Inhibitory Concentration (MIC) (µg/ml) of ATCC 19977 +/- PBP-lipo Knockdown

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Table 2: MIC (µg/ml) of *Mab* Clinical Isolates +/- PBP-lipo knockdown (Cell Wall Antibiotics)

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### Table 3: MIC (µg/ml) of Mab Clinical Isolates +/- PBP-lipo Knockdown (Rifampicin and Clarithromycin)

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