The Cell Wall Lipid PDIM Contributes to Phagosomal Escape and Host Cell Exit of Mycobacterium tuberculosis

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ABSTRACT The cell wall of Mycobacterium tuberculosis is composed of unique lipids that are important for pathogenesis. Indeed, the first-ever genetic screen in M. tuberculosis identified genes involved in the biosynthesis and transport of the cell wall lipid PDIM (phthiocerol dimycocerosates) as crucial for the survival of M. tuberculosis in mice. Here we show evidence for a novel molecular mechanism of the PDIM-mediated virulence in M. tuberculosis. We characterized the DNA interaction and the regulon of Rv3167c, a transcriptional repressor that is involved in virulence regulation of M. tuberculosis, and discovered that it controls the PDIM operon. A loss-of-function genetic approach showed that PDIM levels directly correlate with the capacity of M. tuberculosis to escape the phagosome and induce host cell necrosis and macroautophagy. In conclusion, our study attributes a novel role of the cell wall lipid PDIM in intracellular host cell modulation, which is important for host cell exit and dissemination of M. tuberculosis.

IMPORTANCE Mycobacterium tuberculosis is a major human pathogen that has co-evolved with its host for thousands of years. The complex and unique cell wall of M. tuberculosis contains the lipid PDIM (phthiocerol dimycocerosates), which is crucial for virulence of the bacterium, but its function is not well understood. Here we show that PDIM expression by M. tuberculosis is negatively regulated by a novel transcriptional repressor, Rv3167c. In addition, we discovered that the escape of M. tuberculosis from its intracellular vacuole was greatly augmented by the presence of PDIM. The increased release of M. tuberculosis into the cytosol led to increased host cell necrosis. The discovery of a link between the cell wall lipid PDIM and a major pathogenesis pathway of M. tuberculosis provides important insights into the molecular mechanisms of host cell manipulation by M. tuberculosis.

Mycobacterium tuberculosis is one of the most deadly human pathogens due to its ability to manipulate and evade host innate and acquired immune responses (1–6). These capacities evolved during the longstanding interaction of M. tuberculosis with humans, lasting an estimated 50,000 to 70,000 years (7). One of the mechanisms by which M. tuberculosis establishes a favorable intracellular environment is through the manipulation of host cell death pathways in infected cells (3, 8–10). The prevailing model is that M. tuberculosis inhibits apoptosis at early stages of the infection, favoring replication, but induces necrosis at later stages in order to exit the host cell (8, 9). Nevertheless, the molecular mechanisms by which M. tuberculosis accomplishes these tasks remain poorly understood.

One of the first identified virulence factors of M. tuberculosis is the surface glycolipid PDIM (phthiocerol dimycocerosates), when a PDIM-deficient M. tuberculosis H37Rv mu-
tant was shown to be attenuated in a guinea pig model of infection (11). Subsequent signature tagged-transposon mutagenesis studies identified insertions in the PDIM operon that resulted in a similar attenuation in the mouse model of infection (12, 13). Several roles for PDIM function in pathogenesis have been proposed (for review, see references 2 and 14), including macrophage invasion (15), masking of pathogen-associated molecular patterns (PAMPs) (16, 17), resistance to killing by nitric oxide (16, 18, 19), and preventing the recruitment of activated macrophages to the site of infection (17). Nevertheless, there is still a great deal of uncertainty about the molecular mechanisms of PDIM-mediated virulence regulation of \textit{M. tuberculosis} (2).

It is now well established that during their intracellular life cycle, a fraction of \textit{M. tuberculosis} bacteria escape the phagosome (20–22; see reference 23 for review). Thus, increasing numbers of bacteria can be found in the cytosol over time until a threshold is reached and the host cell undergoes necrosis, allowing for exit of \textit{M. tuberculosis} (21, 24). The process of host cell necrosis induction involves potentially multiple bacterial effectors and host cell signaling pathways that are only beginning to be understood (3, 8–10). Recently, our group described a hypervirulent \textit{M. tuberculosis} mutant resulting from the deletion of the gene \textit{Rv3167c}, a putative member of the tetracycline repressor (TetR)-like family of transcriptional regulators (TFR). Infection of macrophages with the \textit{M. tuberculosis} \textit{Rv3167c} deletion (\textit{ΔRv3167c}) mutant strain resulted in a significant increase in phagosomal escape, autophagy, and host cell necrosis compared to wild-type \textit{M. tuberculosis}-infected cells (25). However, we did not identify the \textit{Rv3167c}-regulated genes, which are responsible for the observed phenotypes. In the present study, we further investigated the virulence regulation by \textit{Rv3167c} and uncovered a novel molecular mechanism for the action of PDIM on the host cell.

**RESULTS**

\textbf{Rv3167c functions as a typical TFR.} TFRs are among the most abundant family of transcriptional regulators in bacteria and as such have been extensively studied (26, 27). \textit{Rv3167c} is annotated as a TFR since 89\% of its protein sequence can be modeled by the Phyre2 software with 99.9\% confidence using the highest-scoring template TFR (SC00332) of \textit{Streptomyces coelicolor}. The predicted structure depicts the N terminus containing 3 \(\alpha\)-helices that form a typical helix-turn-helix DNA binding domain common to TFRs and the additional 6 \(\alpha\)-helices that constitute the C-terminal ligand binding and dimerization domains (Fig. 1a). TFRs typically function as homodimers (26, 27). In order to test whether \textit{Rv3167c} can dimerize, we made use of the mycobacterial protein fragment complementation (M-PFC) assay (28), where bacterial growth is seen on agar containing trimethoprim when the proteins of interest interact. As shown in Fig. 1b, growth was observed when \textit{Rv3167c} was dually expressed, indicating that \textit{Rv3167c} interacts with itself and likely forms a dimer. Depicted are the \textit{Rv3167c} interaction, the positive control (GCN4, a yeast transcriptional regulator known to dimerize), and the negative control (empty plasmids). The full-length image, including all negative controls, can be found in Fig. S6 in the supplemental material. Additionally, we demonstrated that purified \textit{Rv3167c} (molecular mass of 24 kDa) has a retention volume identical to that of ovalbumin (molecular mass of 43 kDa), indicating again that \textit{Rv3167c} forms a homodimer (data not shown). TFRs are typically encoded in the genome in close proximity and in opposing orientation to the genes they regulate (29). Figure 1c depicts the genomic organization of the \textit{Rv3167c} locus with the sequence of the intergenic region between \textit{Rv3167c} and the adjacent operon, \textit{Rv3168-Rv3169}. Based on transcriptome sequencing (RNA-seq) data, \textit{Rv3168} and \textit{Rv3169} are two of the most highly upregulated genes in the \textit{M. tuberculosis }\textit{ΔRv3167c} mutant (see Fig. S1 in the supplemental material), suggesting direct regulation by \textit{Rv3167c}.

\textbf{Rv3167c recognizes a binding site within the upstream region of its own promoter.} TFRs are frequently autoregulatory and bind to palindromic DNA sequences (26, 27). Interestingly, a palindrome (\textit{palRv3167c}) exists within the \textit{Rv3167c-Rv3168} intergenic region (sequence highlighted in red in Fig. 1c). Complementary oligonucleotides corresponding to both strands of \textit{palRv3167c} were annealed to form a double-
stranded product, which was incubated with purified Rv3167c and analyzed by electrophoresis mobility shift assay (EMSA). As shown in Fig. 1d, incubation of increasing concentrations of Rv3167c with palRv3167c resulted in an increased mobility shift indicating binding of Rv3167c to the sequence. Importantly, when Rv3167c was incubated with randomized double-stranded DNA of similar size, no shift was observed, demonstrating that Rv3167c's interaction with palRv3167c is specific. We further char-

Fig 1 Rv3167c is a TetR-like transcriptional regulator. (a) Phyre2 predicted structure of Rv3167c modeled in PyMol. Helix-turn-helix DNA binding domain is colored blue. The ligand binding and dimerization domain is colored red. The second subunit of Rv3167c homodimer is colored gray. (b) Dilution series of M. smegmatis on trimethoprim agar coexpressing Rv3167c/Rv3167c fusion proteins and the positive-control interaction pair using the M-PFC system. (c) Organization of the Rv3167c genomic locus depicting the intergenic region between Rv3167 and Rv3168. The putative palindromic Rv3167c binding site is in red (palRv3167c) with each arm of the palindrome underlined. Translational start of Rv3167c and Rv3168 are in boldface and italic. (d) EMSA analysis of recombinant Rv3167c (rRv3167c) binding to palRv3167c. The rRv3167c concentration ranged from 2 to 30 μM. The palRv3167c concentration was constant at 200 nM. Randomized DNA serving as a negative control was used at 200 nM with 30 μM rRv3167c. The figure is representative of three independent experiments. (e) ITC analysis of rRv3167c binding to palRv3167c. The figure is representative of three independent experiments. The K_d (dissociation constant) of rRv3167c for palRv3167c is 12.3 ± 3.2 μM.
characterized Rv3167c interaction with \textit{palRv3167c} using isothermal titration calorimetry (ITC). Figure 1e is a representative example of three independent ITC experiments. Rv3167c bound \textit{palRv3167c} in a 1:1 protein-DNA stoichiometry and had a dissociation constant of 12.3 ± 3.2 \(\mu\text{M}\), which is similar to other TFR interactions with DNA. Importantly, Rv3167c demonstrated no significant interaction with randomized control DNA (see Fig. S2 in the supplemental material). Taken together, these data indicate that Rv3167c functions as a typical TFR that represses its own promoter and also negatively regulates the downstream gene(s)—in this case \(\text{Rv3168}\) and \(\text{Rv3169}\).

The \textit{Rv3167c} regulon is enriched for genes involved in PDIM synthesis. In order to characterize the regulon of Rv3167c, we performed RNA-seq on the wild-type \textit{M. tuberculosis}, \textit{M. tuberculosis \textit{\Delta}Rv3167c} deletion mutant, and \textit{M. tuberculosis \DeltaRv3167c} complement (\(\Delta\text{Rv3167c}:\text{Comp}\)) strains grown in standard growth media and harvested during logarithmic growth. Our RNA-seq data were highly reproducible, as evidenced by the high degree of clustering between biological replicates, and indicated that we are analyzing distinct transcriptional states (Fig. 2a). The inclusion of the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}:\text{Comp}\) strain in our analysis provided a valuable internal control for determining the Rv3167c regulon. To do this, we determined genes differentially expressed in the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}\) mutant compared to \textit{M. tuberculosis} as well as genes differentially expressed in the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}\) strain compared to the \(\Delta\text{Rv3167c}:\text{Comp}\) strain. We compared both gene sets (Fig. 2b) and only considered genes differentially expressed in both contrasts as this gave us more confidence Rv3167c was involved in their regulation. In total, 442 genes were considered to be regulated by Rv3167c (see Table S1b in the supplemental material). The differential expression of such a number of genes indicates that Rv3167c has a relatively broad transcriptional impact. Gene Ontology (GO) enrichment analysis identified three GO terms (Table 1). The GO term “DIM/DIP cell wall layer assembly,” which encompasses genes associated with the synthesis and export of PDIM, was enriched in our gene set. Indeed, the entire PDIM operon was upregulated in the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}\) mutant compared to wild-type \textit{M. tuberculosis}, while being mostly unchanged or even downregulated in the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}:\text{Comp}\) strain compared to the wild type, indicating these genes do complement back to wild-type levels (Fig. 2c). Using thin-layer chromatography (TLC), we showed that the transcriptional upregulation manifested in an increase in PDIM lipid in the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}\) mutant (Fig. 2d).

PDIM is involved in regulating host cell necrosis induction by \textit{M. tuberculosis}. PDIM is synthesized in the cytosol and transported to the outer membrane of \textit{M. tuberculosis} by the membrane lipid transporter \textit{mmpl7}. Deletion of \textit{mmpl7} results in the failure of proper PDIM localization and the accumulation of the lipid in the cytosol (12). To assess the role of PDIM in the necrosis induced by the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}\) mutant, we deleted \textit{mmpl7} in the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}\) background, resulting in the \(\Delta\text{Rv3167c} \Delta\text{mmpl7}\) double deletion mutant, which was confirmed by PCR (see Fig. S3 in the supplemental material). Infection of THP-1 macrophages with the \textit{M. tuberculosis} \(\Delta\text{Rv3167c} \Delta\text{mmpl7}\) mutant resulted in a significant decrease in cell death compared to the \(\Delta\text{Rv3167c}\) strain (Fig. 3a). As described above, RNA-seq analysis revealed more than 400 genes differentially regulated in the \textit{M. tuberculosis} \(\Delta\text{Rv3167c}\) strain. In an effort to minimize concerns about the contribution of other genes to the cell death phenotype and to assess whether PDIM alone was sufficient to modulate host cell death, we analyzed an \textit{mmpl7} transposon insertion mutation (\textit{mmpl7::TN}) in the \textit{M. tuberculosis} CDC1551 strain obtained from BEI Resources (30). The deletion of \textit{mmpl7} resulted in significantly less cell death than wild-type \textit{M. tuberculosis} (Fig. 3c; see Fig. S9a in the supplemental material). Complementation of \textit{mmpl7} back into the \textit{M. tuberculosis} \textit{mmpl7::TN} mutant reverted the phenotype to levels similar to those of the wild type (Fig. 3c). A clean deletion mutant generated through a recombination strategy (the \textit{M. tuberculosis} \(\Delta\text{mmpl7}\) mutant) confirmed this finding (see Fig. S7a in the supplemental material). Thin-layer chromatography confirmed that these mutants produce similar levels of PDIM to the wild type (see Fig. S8 in the supplemental material), confirming these mutants are deficient in export of PDIM, not synthesis. Previous work has
implicated PDIM in macrophage invasion (15) and in resistance to killing by an early innate immune response (16, 18, 19). However, we saw similar numbers of bacteria both initially after infection and at 24 h postinfection in the *M. tuberculosis* ΔRv3167cΔmmpl7 (Fig. 3b) and mmpl7::TN (Fig. 3d) mutants compared to wild-type *M. tuberculosis*. These data thus confirm that the bacterial PDIM levels directly impact the capacity of *M. tuberculosis* to induce host cell necrosis and do not just reflect a lower quantity of intracellular bacteria.

**PDIM contributes to phagosomal escape of *M. tuberculosis* and host cell autophagy induction.** Macroautophagy (autophagy) is an evolutionarily conserved cate-
abolistic process that involves sequestration of cytosolic contents into de novo-generated vesicles termed autophagosomes, which then get delivered to the lysosome for degradation. Intracellular pathogens can be targeted and killed by autophagic machinery in a process termed xenophagy (31). Previous work in our lab demonstrated that in addition to increased necrosis, infection of cells with the *M. tuberculosis* ΔRv3167c mutant resulted in increased induction of autophagy but not xenophagy (25). In order to determine whether PDIM played a role in *M. tuberculosis*-induced autophagy in addition to the induction of necrosis, we monitored the conversion of cytosolic LC3I to

### TABLE 1 GO term enrichment of *M. tuberculosis* ΔRv3167c mutant differentially regulated genes

| Category | Term | No. of differentially expressed genes in category | No. of genes in category | Adjusted P value |
|----------|------|--------------------------------------------------|--------------------------|------------------|
| GO:0071770 | DIM/DIP cell wall layer assembly | 9 | 9 | 0.0000062 |
| GO:0006633 | Fatty acid biosynthetic process | 6 | 7 | 0.0048904 |
| GO:0005618 | Cell wall | 123 | 659 | 0.0045409 |

**FIG 3** PDIM contributes to induction of host cell necrosis by *M. tuberculosis*. THP-1 cells were differentiated with PMA for 24 h and were left uninfected (UI) or were infected with the indicated *M. tuberculosis* strains. (a and c) Cells were harvested, stained with propidium iodide (PI), and analyzed by flow cytometry (*n* = 5,000) at 24 h postinfection. (b and d) Bacterial burden was determined by quantifying CFU immediately following infection (0 h) or 24 h postinfection (24 h). Data are representative of three independent experiments. Data are presented as means ± SEM. ****, *P* ≤ 0.0001 (one-way ANOVA).
autophagosome-bound LC3II, a hallmark of autophagy, using LC3-green fluorescent protein (GFP)-expressing THP-1 cells and flow cytometry (32). THP-1 LC3-GFP-expressing cells were infected for 24 h and partially lysed with saponin, and the remaining LC3-GFP was measured using flow cytometry (Fig. S9b for gating and raw data). A significant decrease in autophagy was seen in cells infected with the M. tuberculosis mmpl7::TN mutant compared to wild-type M. tuberculosis (Fig. 4a), a finding confirmed by analysis of the M. tuberculosis Δmmpl7 mutant (Fig. S7b) indicating PDIM plays a role in induction of autophagy in M. tuberculosis.

The M. tuberculosis ΔRv3167c mutant accesses the cytosol earlier and to a greater extent than wild-type M. tuberculosis (25). We hypothesized that PDIM contributes to phagosomal escape and that loss of PDIM would result in decreased access to the cytosol. To test this, we made use of a fluorescence resonance energy transfer (FRET)-based assay described previously (21, 22). Briefly, THP-1 cells were loaded with CCF4-AM. Intact CCF4-AM emits green fluorescence (535 nm) due to FRET between the fluorescent moieties. Cleavage of CCF4-AM by β-lactamase expressed by cytosolic bacteria leads to FRET loss and a shift in the emission wavelength to 450 nm, which can be monitored by flow cytometry (Fig. S9c). Using this assay, we demonstrate that the M. tuberculosis mmpl7::TN mutant accesses the cytosol to a lesser extent than wild-type M. tuberculosis and the mmpl7::TN complemented strain (Fig. 4b).

DISCUSSION

One of the goals of this study was to define the regulon of Rv3167c since we previously documented that this TFR represses virulence of M. tuberculosis, and hence its regulon should contain important virulence factors (25). Our RNA-seq analysis revealed a total of 442 genes differentially regulated in the M. tuberculosis ΔRv3167c mutant suggestive of a broad transcriptional response. TFRs respond to environmental cues through their ligand binding domain (26, 27). We hypothesize that intracellular
stressors such as low pH and increased reactive oxygen species (ROS) and nitrate species could be responsible for generating the Rv3167c ligand, but we failed to detect a stimulus that induces the loss of DNA binding by Rv3167c (data not shown). This is most likely due to the fact that in vitro approaches do not accurately replicate the complex milieu of a macrophage phagosome. The Rv3167c regulon is enriched for genes involved in the synthesis and transport of PDIM. There are several known transcriptional regulators of PDIM biosynthesis, such as rpoB (34) and espR (35), as well as posttranslational regulators such as pkhn (36). However, none of these genes is differentially regulated in the M. tuberculosis ΔRv3167c mutant. The Rv3167c regulon contains several RNA polymerase sigma factors, including sigE and sigB, as well as 21 putative or known transcriptional regulators (Table S1). Differential expression of these regulators could explain the high degree of gene regulation we see in the M. tuberculosis ΔRv3167c mutant and suggests that the control of PDIM gene expression by Rv3167c is indirect.

M. tuberculosis was long thought to reside exclusively within a vacuole that has the characteristics of an early endosomal compartment (1, 5, 37). However, it is now evident that M. tuberculosis also escapes the phagosome to reach the host cell cytosol (2, 38). After gaining access to the cytosol, M. tuberculosis induces necrosis in order to exit the infected cell and to disseminate (20, 21, 24, 39). Our previous findings indicated that the Rv3167c mutant consistently induced more necrosis and escaped the phagosome earlier and to a higher degree than wild-type M. tuberculosis (25). We hypothesized that the increased PDIM in the ΔRv3167c mutant may play a role in the associated increase in necrosis induction and escape. This hypothesis was confirmed in the present study via deletion of the PDIM transporter mmpl7 in the background of the Rv3167c mutant in M. tuberculosis H37Rv (Fig. 3a) and by using an mmpl7 transposon mutation in the clinical strain CDC1551 (Fig. 3c). We propose that PDIM primarily contributes to the escape of M. tuberculosis from the phagosome and that the increase of host cell necrosis is merely a consequence of the increased number of cytosolic bacteria. However, it cannot be ruled out that PDIM has a direct role in induction of necrosis in addition to its function in phagosomal escape. Our previous findings demonstrated that induction of necrosis by the ΔRv3167c mutant was dependent on increases in mitochondrial reactive oxygen species (ROS) (25). M. tuberculosis lipids can traffic within endosomal membranes of host cells (40). While it has not been shown that M. tuberculosis lipids can also localize to host cell mitochondria, phagosomes do interact with mitochondria in order to optimize ROS production for defense against pathogens (41). Consequently, it is possible that lipids of M. tuberculosis inserted into the phagosomal membrane will be shared with mitochondrial membranes. PDIM may interact directly with mitochondrial membranes, leading to increased ROS production, which may initiate host cell necrosis.

The phagosomal escape of M. tuberculosis is dependent on the pore-forming activity of the ESX-1-secreted substrate EsxA (20, 22, 33). PDIM may influence M. tuberculosis escape through direct effects on EsxA secretion. Nevertheless, we show here that loss of PDIM does not affect secretion of EsxA (Fig. S5). Furthermore, while the deletion of esxA results in a complete lack of M. tuberculosis escape into the cytosol (20, 22, 33), the deletion of mmpl7 leads to only an impaired escape (Fig. 4b). This suggests that while EsxA is absolutely necessary for M. tuberculosis to gain access to the cytosol, PDIM may work synergistically with EsxA to help M. tuberculosis escape the phagosome. Nevertheless, we are comparing these results with caution since the esxA mutant data were generated previously by a different lab and in a different genetic background. It was shown previously that EsxA favors association with liposomes containing phosphatidylcholine and cholesterol (42). While these lipids are not essential for insertion of EsxA (43, 44), the idea that EsxA may show preference for certain membrane compositions is enticing. Cholesterol is a critical component of mammalian cellular membranes as it provides the membrane rigidity necessary for signaling events (45). Certain biophysical properties, such as membrane fluidity, may promote insertion of EsxA into host membranes. PDIM has the ability to insert into host membranes, decreasing their
fluidity (15). While not necessary for EssA pore formation, insertion of PDIM into phagosomal membranes may alter the biophysical properties of the membrane to favor further EssA insertion. There is also the possibility that EssA and PDIM work independently toward the same goal. While EssA is essential for gaining access to the host cell through membrane pore formation, PDIM may insert and weaken the phagosomal membrane, favoring bacterial escape.

The proposed role of PDIM in *M. tuberculosis* virulence is multifaceted (14), but there is still some uncertainty over what the molecular mechanisms are (2). For example, the impact of PDIM on cell entry seems to be marginal since it could only be demonstrated under very limited experimental conditions (15), and also our own experimental results did not detect any differences in uptake between wild-type *M. tuberculosis* and *mmpl7* mutants (Fig. 3b and d). PDIM is involved in the recruitment of activated macrophages to the site of infection in zebrafish (17). In light of our novel findings, one could hypothesize that the differential recruitment of macrophages may also be explained due to differences in host cell necrosis induction and dissemination mediated by wild-type *Mycobacterium marinum* versus the *mmpl7* mutant, which was not analyzed in the study (17). Indeed, the recruitment of neutrophils to infected areas of the lung is stimulated by the increase in host macrophage necrosis (46). Altogether, our findings establish a new and prominent role for PDIM in the phagosomal escape of and host cell necrosis induction by *M. tuberculosis*. This novel mechanism for manipulation of the host cell provides an additional answer for the long-standing puzzle of how PDIM contributes to the virulence of *M. tuberculosis*.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** All *M. tuberculosis* cultures were grown at 37°C in 7H9 broth or on 7H11 agar medium with 1× ADC enrichment (albumin-dextrose-catalase), 0.5% glycerol, and 0.05% Tween 80 (broth). *Escherichia coli* cultures were grown in LB medium. Kanamycin, hygromycin, and zeocin were used at 40, 50, and 100 μg/ml, respectively, for *M. tuberculosis* cultures and 40, 150, and 100 μg/ml, respectively, for *E. coli*.

**Cell culture and infections.** THP-1 cells from ATCC were maintained at 37°C and 5% CO2 in RPMI (ATCC) supplemented with 10% heat-inactivated fetal calf serum (FCS) (growth medium). THP-1 LC3-GFP-expressing cells were grown in growth medium plus 100 μg/ml G418 sulfate (Cellgro). For infections, THP-1 cells were treated with 20 ng/ml phorbol 12-myristate 13-acetate (PMA) (Sigma) for 30 min. The supernatant was applied to columns packed with His-Pur cobalt Superflow resin (Thermo-Fisher) equilibrated with PBS plus 200 mM NaCl plus 10 mM imidazole and incubated at 4°C for 1 h with rocking. The column was washed with 10 volumes of PBS plus 200 mM NaCl plus 30 mM imidazole. PDIM was eluted from the column with PBS plus 200 mM NaCl plus 250 mM imidazole. Ten percent glycerol was added, and the purified protein was stored at 4°C until fast protein liquid chromatography (FPLC) purification.

**FPLC.** All FPLC experiments were carried out on a GE AKTA Explorer chromatography system. Gel filtration chromatography was carried out on a Superdex 75HR 10/30 column (GE Healthcare Life Sciences). Eluent was then further purified with ion-exchange chromatography on a MonoQ 5/50 Gl column (GE Healthcare Life Sciences). Purified protein was confirmed to be Rv3167c by mass spectrom-
at the indicated time points, cells were harvested and washed once in PBS and then resuspended in PBS.

Randomized oligonucleotides served as the negative control. All primer sequences are listed in Table S1a.

at 140°C.

by spraying the plate with a solution of 3% cupric acetate in 8% phosphoric acid followed by baking

The plate was resolved in a mobile phase of 9:1 petroleum ether-diethyl ether. Lipid spots were revealed

formed into

escape from the phagosome, the CCF4 FRET assay was performed as described previously (BD Accuri C6).

Mycobacterial protein fragment complementation (M-PFC) was conducted as described in reference 28. Rv3167c was cloned into both pUBA300 and PUAB400. Constructs were then cotransformed into Mycobacterium smegmatis. Transformants were then spotted with appropriate controls on 7H11 medium supplemented with 50 μg/ml trimethoprim (Fisher) and grown at 37°C for 3 days.

FRET assay for translocation of M. tuberculosis in host cell cytosol. To detect mycobacterial escape from the phagosome, the CCF4 FRET assay was performed as described previously (21). Briefly, cells were stained with 8 μM CCF4 (Invitrogen) in EM buffer (120 mM NaCl, 7 mM KCl, 1.8 mM CaCl2, 0.8 mM MgCl2–5 mM HEPES [pH 7.3]) containing 2.5 μM probenecid (Sigma-Aldrich) for 1.5 h at room temperature. Live populations were distinguished from dead ones by addition of LIVE/DEAD fixable red stain (Invitrogen) for 30 min at room temperature. Following staining, cells were fixed with 4% paraformaldehyde (PFA) overnight and analyzed by flow cytometry (BD FACSCanto).

Representative examples of the flow cytometry gating strategy to determine loss of FRET in infected cells can be found in Fig. S9c.

RNA isolation. M. tuberculosis cultures were grown in 7H9 until they reached an OD_{600} of 0.6 to 0.8. Cultures were pelleted and resuspended in 1 ml TRIzol (Ambion). RNA was extracted with chloroform, precipitated with 100% isopropanol, and washed with 70% ethanol. Purified RNA was treated with Turbo DNase (Ambion) for 1 h. RNA was used right away or stored at −80°C.

cDNA synthesis. The Maxima first strand cDNA synthesis kit (Thermo Scientific) was used to synthesize cDNA as per the manufacturer’s instructions.

RNA-seq library preparation and analysis. Ribosomal RNA was removed from samples using the Epicentre Gram-positive Ribo-Zero rRNA magnetic removal kit. RNA-seq libraries were prepared using the Illumina ScriptSeq v2 library preparation kit according to the manufacturer’s protocol. Library quality was assayed by Bio-analyzer (Agilent) and quantified by quantitative PCR (qPCR) (Kapa Biosystems). Libraries were run on an Illumina HiSeq1500, generating 100-bp paired end reads. RNA-seq read quality was verified using FastQC, and low-quality base pairs were trimmed using Trimmomatic. The M. tuberculosis H37Rv reference genome was downloaded from TB Database, and reads were mapped to the genome using TopHat2. Count tables were generated using HTSeq along with a custom GFF (Kapa Biosystems).

Total lipid extraction and TLC. Cultures were grown in 7H9 broth in a volume of 50 ml to an OD_{600} of 0.8. Cultures were pelleted and resuspended in 2:1 methanol-chloroform and left at 4°C overnight. Samples were spun down, and supernatant was transferred to a glass beaker. The pellet was resuspended in 1:1 methanol-chloroform and left at room temperature for an hour. Samples were spun down, and the supernatant was removed. The process was repeated again with 1:2 methanol-chloroform. All three fractions were pooled, and the methanol and chloroform were allowed to evaporate overnight in a fume hood. The total lipid pellet was weighed, and 300 μg of total lipid was spotted onto a TLC plate. The plate was resolved in a mobile phase of 9:1 petroleum ether-diethyl ether. Lipid spots were revealed by spraying the plate with a solution of 3% cupric acetate in 8% phosphoric acid followed by baking at 140°C.
Immunoblotting. Bacteria were pelleted and resuspended in 25 mM Tris HCl, followed by bead beating to lyse cells. Lysate was cleared by spinning at 12,000 rpm for 5 min. The Pierce bicinchoninic acid (BCA) protein assay kit (Thermo Scientific) was used to measure protein concentrations. Antibody against EsxA was purchased from Santa Cruz Biotechnology, Inc., and used at a 1:200 dilution. Antibodies against Fap and GroEL were obtained from BEI Resources and used at 1:7,500 and 1:50 dilutions, respectively.

Statistical analysis. Statistical analysis was performed using GraphPad Prism version 6.0 software. Data are presented as means ± standard errors of the means (SEM) from three independent experiments, and one-way analysis of variance (ANOVA) with Tukey’s posttest was used unless mentioned otherwise in the figure legends. P value significance is indicated on the figures as follows: *, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001; and ****, P ≤ 0.0001.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.00148-17.

FIG S1, JPG file, 1 MB.
FIG S2, JPG file, 0.1 MB.
FIG S3, JPG file, 0.1 MB.
FIG S4, JPG file, 0.1 MB.
FIG S5, JPG file, 0.1 MB.
FIG S6, JPG file, 0.1 MB.
FIG S7, JPG file, 1.6 MB.
FIG S8, JPG file, 0.1 MB.
FIG S9, JPG file, 2.9 MB.
TABLE S1, XLSX file, 0.1 MB.

ACKNOWLEDGMENT

All authors were supported by grant AI114269 from the NIH/NIAID. Jeff Quigley was supported by the Training Program in Cell and Molecular Biology NIH T32GM080201.

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