Serine/Threonine Protein Phosphatase PstP of 
*Mycobacterium tuberculosis* Is Necessary for Accurate Cell 
Division and Survival of Pathogen*[^1,5]

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Protein phosphatases play vital roles in phosphorylation-mediated cellular signaling. Although there are 11 serine/threonine protein kinases in *Mycobacterium tuberculosis*, only one serine/threonine phosphatase, PstP, has been identified. Although PstP has been biochemically characterized and multiple *in vitro* substrates have been identified, its physiological role has not yet been elucidated. In this study, we have investigated the impact of PstP on cell growth and survival of the pathogen in the host. Overexpression of PstP led to elongated cells and partially compromised survival. We find that depletion of PstP is detrimental to cell survival, eventually leading to cell death. PstP depletion results in elongated multiseptate cells, suggesting a role for PstP in regulating cell division events. Complementation experiments performed with PstP deletion mutants revealed marginally compromised survival, suggesting that all of the domains, including the extracellular domain, are necessary for complete rescue. On the other hand, the catalytic activity of PstP is absolutely essential for the *in vitro* growth. Mice infection experiments establish a definitive role for PstP in pathogen survival within the host. Depletion of PstP from established infections causes pathogen clearance, indicating that the continued presence of PstP is necessary for pathogen survival. Taken together, our data suggest an important role for PstP in establishing and maintaining infection, possibly via the modulation of cell division events.

Signal sensing and transduction lead to a wide range of cellular responses and thus must be tightly regulated in pathogenic bacteria to allow optimal survival under variable conditions. A common mode of regulation of the cell’s response to external cues is via phosphorylation and dephosphorylation of specific target proteins that lead to cellular responses like altered subcellular localization, protein turnover rates, and protein-protein interactions. Whereas phosphorylation events are mediated by kinases, dephosphorylations are mediated by the action of phosphatases. Bacteria are known to possess the conventional two-component systems involving phosphorylation of His residues in sensor kinases and Asp residues in the corresponding response regulators. These systems are critical for multiple physiological processes and cell survival (1, 2). In most of these systems, kinase/phosphatase activities are possessed by one bifunctional enzyme; for example, in the mycobacterial DevS/DosS system and WalRK system of *Bacillus anthracis*, DevS and WalK were shown to exhibit both kinase and phosphatase activities (3, 4). There are other known bifunctional kinase/phosphatases that do not belong to two-component systems, such as AceK of *Escherichia coli* (5) and HPr of *Bacillus subtilis* (6). Mycobacteria possess 11 two-component systems; besides this, pathogenic bacteria also possess Ser/Thr phosphorylation systems in which a protein that has been phosphorylated by a serine/threonine protein kinase (STPK)[^6] would be dephosphorylated by a serine/threonine phosphatase. Analysis of the *Mycobacterium tuberculosis* whole genome sequence identified 11 eukaryotic-like STPKs (PknA–L, except for PknC), one Ser/Thr phosphatase (PstP), one tyrosine kinase (PtK), and two tyrosine phosphatases (PtpA and PtpB) (7, 8). All of these eukaryotic-like kinases and phosphatases have now been characterized and found to be catalytically active (9–19).

Protein phosphatases belong to distinct families, and members of a family are structurally and functionally conserved enzymes. Phosphatases are classified based on the catalytic domain signature sequence they carry and their substrate preference (the metal-dependent protein phosphatase (PPM) family, the phosphoprotein phosphatase (PPP) family, and the phosphotyrosine phosphatase (PTP) family) (20, 21), and there

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are examples known from all of these families in bacteria (20, 22, 23). PPP phosphatases are predominantly found in euukaryotes and target Ser/Thr residues in their substrates. The PP2C class of Ser/Thr phosphatases belong to PPM family and require Mg\(^{2+}\)/Mn\(^{2+}\) for their activity. In most bacteria, these are usually secreted out of the cell; PTPs are the Tyr phosphatases and are subclassified into class I standard classical PTPs and class II low molecular weight (LMW-PTP), which in many bacteria are secreted into the extracellular milieu (20, 22, 23). *M. tuberculosis* encodes two tyrosine phosphatases, PtpA and PtpB. Deletion of *ptpA*, an LMW–PTP, did not impact *M. tuberculosis* growth either in vitro or in the mouse infection model (24). However, results from competitive co-infection of THP-1 macrophages with *H37Rv* and *MtbΔptpA* strains revealed that PtpA is important for long term infection (25). PtpA is secreted from the bacterial cell and dephosphorylates the host’s VSP33B (vacuolar protein sorting 33B) protein, a subunit of the VPS-C complex, thus causing the inhibition of phago-lysosomal fusion (25). The interaction of PtpA with the host vacuolar H\(^+\) ATPase is necessary for the dephosphorylation of VSP33B and subsequent phagosomal exclusion of V-ATPase (26). PtpB is also secreted into the extracellular milieu, and the disruption of *ptpB* impairs the pathogen’s ability to survive in activated macrophages and guinea pigs (27). Clearance of *M. tuberculosis* from the lungs occurs via endocytic pathways that require the fusion of phagosome with lysosome, mediated by the VPS-C complex (28). Although no direct substrates of PtpB have been identified to date, the expression of PtpB in activated macrophages attenuated IFN-γ and IL-6 production, possibly through ERK2 and p38 dephosphorylation (29).

The lone serine/threonine phosphatase of *M. tuberculosis*, PstP, belongs to the PP2C class of PPM family phosphatases and strictly requires the Mn\(^{2+}\) ion for its activity (30). The enzyme localizes to the cell membrane and contains a 240-amino acid intracellular catalytic domain, tethered via a single transmembrane helix to the 196-amino acid-long extracellular domain (18, 31). Although two metal centers are found in the catalytic core in most PP2C phosphatases, the crystal structural of PstP showed the presence of three metal-binding centers (31). The structure of PstP was later refined by the analysis of *M. tuberculosis* from the lungs occurs via endocytic pathways that require the fusion of phagosome with lysosome, mediated by the VPS-C complex (28). Although no direct substrates of PtpB have been identified to date, the expression of PtpB in activated macrophages attenuated IFN-γ and IL-6 production, possibly through ERK2 and p38 dephosphorylation (29).

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**Results**

Impact of Overexpression of PstP—Our laboratory has previously shown that overexpression of PknA or PknB results in cell death, probably due to hyperphosphorylation of substrates critical in modulating cell survival. Because PstP is the sole phosphatase in *M. tuberculosis*, we speculated that overexpression of PstP would also be likely to impact cell survival. To test this hypothesis, PstP was cloned into the IVN-inducible episomal vector pNit-1 (36) and tet-regulatable integrative vector pST-KirT (37). The constructs were suitably prepared and electroporated into the *M. smegmatis* mc^2^155 strain, and the recombinants were analyzed for PstP expression in the presence or absence of inducer. In the case of mc^2^::KirT-pstp strain, we noticed moderate levels of overexpression in the absence of anhydrotetracycline (ATc) (Fig. 1A), which was consistently observed in four independent experiments. The expression was reduced significantly upon the addition of ATc (albeit not to the endogenous levels), indicating that the promoter could be turned off upon the addition of inducer (Fig. 1A). Based on the growth pattern analysis, we concluded that mild overexpression of PstP from the integrative pST-KirT-pstP does not alter the growth profile (Fig. 1B). To quantitate the differences, cfu at 0 and 30 h were enumerated (Fig. 1C). The presence or absence of ATc did not alter the cfu obtained in the case of mc^2^::KirT-pstP strain did not impact the number of cfu (Fig. 1E). Interestingly, even this level of expression led to a statistically significant increase in the cell length, which reverted back upon the addition of ATc (Fig. 2, A and B).

The results obtained for mc^2^::pNit1-pstP varied between two phenotypes. At times, we observed significant cell death upon the addition of IVN. Western blot analysis from such an experiment exhibited significant overexpression of PstP upon the addition of IVN (data not shown). However, on the majority of occasions, we observed a marginal difference in the growth in the absence and presence of IVN (Fig. 1E). Correspondingly, the Western blot analysis indicated marginal differences in the expression pattern of PstP in the absence and presence of IVN (Fig. 1D). To quantitate the impact of PstP overexpression on cell survival, cfu analysis was performed with mc^2^ and mc^2^::pNit1-pstP cultures grown in the presence or absence IVN (Fig. 1F). We observed a marginal but statistically significant
decrease in the cfu upon the addition of IVN (Fig. 1F). Scanning electron microscopy revealed that overexpression of PstP resulted in elongation of cells (Fig. 2, C and D). The presence or absence of ATc or IVN did not alter the cell length of mc2. Interestingly, we observed slight but statistically significant change in mean cell length upon the addition of IVN (Fig. 2D). These results suggest a role for PstP in maintaining normal cell lengths.

Depletion of PstP in M. smegmatis Leads to Cell Death—To decipher the role of PstP in regulating cellular events, we began with creating a conditional gene replacement pstP mutant in M. smegmatis. Previous attempts to generate mycobacterial pstP gene replacement mutants in our laboratory were unsuccessful, possibly due to polarity effects on the expression of the downstream genes in the operon. We therefore modified the strategy for generating the allelic exchange substrate (AES), wherein the carboxyl terminus of the hygr gene was fused with a spacer (encoding four glycine residues and one serine residue) followed by the last 21 nucleotides of the pstP gene (encoding the last seven amino acids of PstP). Because the translational stop codon of PstP overlaps with the translational start codon of RodA, we reasoned that the above strategy would minimally disrupt both transcription and translation of the downstream genes. The mc2-KirT-pstP strain was electroporated with pNitA-ET to generate recombineering-proficient merodiploid strain. This strain was electroporated with blunt-ended linearized AES to replace the pstP at the native locus with a modified hygr gene. Fidelity of recombination at the native pstP locus was confirmed by performing PCR amplification with specific primers across the replacement junctions (Fig. 3, A and B). To determine the impact of PstP depletion on cell growth, we streaked M. smegmatis wild type (mc2), merodiploid (mc2::KirT-pstP), and the mutant (mc2-cd-pstP) on plates in the presence or absence of ATc (Fig. 3C). As expected, both wild type and merodiploid strains grew regardless of the presence of ATc. However, the conditional mutant grew well only in the absence of ATc, because in the presence of ATc we did not detect any growth on plates (Fig. 3C). Western blot analysis of the whole cell lysates, isolated at different time points post-ATc addition, confirmed the depletion of PstP (Fig. 3D). To examine the pos-
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FIGURE 2. Impact of overexpression of PstP on morphology. A, mc^2 or mc^2::KirT-pstP cultures were seeded at 0.1 and allowed to grow in the presence and absence of 50 ng/ml ATc inducer for 12 h. Cells were processed for SEM as described under “Materials and Methods.” Scale bar, 1 μm. B, cell lengths of 150 individual cells from different SEM images were measured using Smart Tiff software, and data were analyzed using GraphPad Prism version 6. Mean lengths are plotted, and significance is calculated using ordinary one-way ANOVA with *p* < 0.0001 (****). Mean lengths of mc^2, mc^2 + ATc, mc^2::KirT-pstP – ATc, and mc^2::KirT-pstP + ATc were 3.74, 3.68, 5.13, and 4.03 μm, respectively. C, mc^2 or mc^2::pNit1-pstP cultures were initiated at an A_0.04 of 0.1 in the absence or presence of 5 μM IVN and grown for 12 h. SEM was performed as described. Scale bar, 1 μm. D, cell lengths of 115 individual cells from SEM images were measured by Smart Tiff software and analyzed as mentioned in B. Mean cell lengths of mc^2, mc^2 + IVN, mc^2::pNit1-pstP – IVN, and mc^2::pNit1-pstP + IVN samples were 3.61, 3.49, 4.60, and 5.61 μm, respectively. ****, *p* < 0.0001, ordinary one-way ANOVA. ns, not significant.

sibility of polarity effects on the expression of downstream genes of the operon, the whole cell lysates were analyzed with anti-PknB antibody. The expression of PknB was found to be similar regardless of the presence of ATc, indicating that the replacement of pstP at its native locus did not alter expression of the downstream genes. A comparison of the growth pattern analysis of the mutant versus the wild type strain revealed that the depletion of PstP significantly reduced the growth (Fig. 3E). The reduced growth observed in the presence of ATc could be due to either bacteriostatic or bactericidal phenotypes. To resolve this question, we enumerated cfu at the 0 and 30 h time points in the presence and absence of ATc. The depletion of PstP resulted in ~3 log-fold decrease in the survival of bacteria, suggesting that depletion of PstP was detrimental to cell survival (Fig. 3F).

Decrease in PstP Expression Compromises M. tuberculosis Growth—A conditional gene replacement of pstP in M. tuberculosis was created with the help of recombineering as described under “Materials and Methods.” Replacement of pstP at the native locus was confirmed by performing PCRs with appropriate primers (Fig. 4A), using the genomic DNA isolated from Rv and Rv-cd-pstP (Fig. 4B). Although the expression of PknB (from the last gene in the operon; Fig. 4A) was similar in the absence/presence of ATc, there was a significant decrease in PstP expression in the presence of ATc (Fig. 4C). Although this decrease in PstP expression was significant when compared with expression in the absence of ATc (compare lanes 2 and 3 of Fig. 4C), the levels of PstP expression in the presence of ATc were only ~1.9-fold lower than PstP expression levels detected in wild type cells (from the native locus; compare lane 3 with lane 1 in Fig. 4C). Nevertheless, we analyzed the mutant for growth defects by determining the cfu/ml of culture grown in the absence or presence of ATc for 0, 2, and 4 days. We observed the impact of PstP decreased expression to be ~1 log-fold (10-fold) compared with the wild type (Fig. 4D). Thus, a tight regulation of PstP expression appears to be critical for optimal growth and survival. Moreover, scanning electron microscopy analysis indicated marginal elongation of cells upon partial depletion of PstP (Fig. 4E).

The Phosphatase Activity of PstP Is Essential for In Vitro Growth—PstP has four distinct domains: an intracellular catalytic domain (amino acids 1–240), a 60-amino acid-long juxtamembrane domain, a transmembrane domain that carries 18 amino acids, and an extracellular domain of 196 amino acids (Fig. 5A). Although the catalytic domain was shown to be functional in vitro (18), the roles played by the remaining domains have not been elucidated either in vitro or in vivo. To decipher the roles of the different domains and the active site residues in the functioning of PstP in vivo, wild type PstP, PstP deletion mutants, and PstP point mutants were cloned into pNitA. The constructs were electroporated into the M. smegmatis PstP conditional mutant mc^2-cd-pstP, and the transformants were assessed for the ability of the mutated PstP proteins (expressed from the plasmids) to complement PstP depletion (in the pres-
ence of ATc). We standardized the conditions for appropriate expression of the PstP deletion fragments by inducing with different concentrations of IVN. Western blot analysis clearly demonstrated the expression of all three deletion fragments (Fig. 5, indicated by arrows). Growth analysis in liquid culture as well as cfu analysis suggested effective complementation by episomally expressed wild type PstP in the presence of ATc (Fig. 5, C and D). Importantly, we observed compromised growth with all three deletion mutants, and the cfu analyses are in agreement with the observed growth patterns (Fig. 5, C and D). Together, analysis suggests that although the deletion fragments could complement, the ability to complement is not equivalent to that of full-length PstP protein.

It has previously been shown that mutating Asp-38 and Asp-229 to glycine resulted in 90% loss of the dephosphorylation activity, whereas mutating Arg-20 to Gly resulted in 60% loss.
of its activity (35). We sought to investigate the role of phosphatase activity of PstP in regulating growth (Fig. 6A). Western blot analysis of lysates prepared from the transformants confirmed the expression of PstP and its active site mutants (Fig. 6B). Growth pattern and cfu analysis suggested that none of the point mutants (D38G, R20G, and D229G) could complement PstP depletion (Fig. 6, C and D). Although these mutants display varying levels of residual phosphatase activity (35), it appears that a 60% loss in phosphatase activity is sufficient to compromise in vitro growth. Thus, our results suggest that the activity of PstP is essential for mycobacterial growth.

Depletion of PstP Results in Multiseptum Phenotype—The pstP gene is located in an operon that encodes for PknA, PknB, RodA, and PbpA. These four genes have all been suggested to play a critical role in cell shape maintenance, cell division, and cell wall synthesis processes (34, 38). PknA and PknB modulate
the activities of a number of proteins that participate in cell division and cell wall synthesis by mediating their phosphorylation (39–41). Given the fact that PstP is the sole serine/threonine phosphatase in mycobacterium and is carried by an operon that carries genes controlling cell shape and cell division (including two kinases), we investigated any possible role that PstP might play in modulating cell shape and cell division by examining the impact of PstP depletion on these parameters using scanning electron microscopy (SEM). Interestingly, significantly elongated and bulged cells with multiple septa were observed in PstP-depleted samples (ATc treatment-mediated 12-h depletion), suggesting a possible role for PstP in modulating cell shape and cell division. To investigate whether stringent regulation of PstP expression is critical for the survival of pathogen within the host, we infected mice with Rv or Rv-cd-pstP, and the mice were either provided or not provided doxycycline (Fig. 8A). Analysis of cfu determined 1 day post-infection showed equivalent implantation of wild type and mutant bacilli in mice lungs (Fig. 8B). To determine the impact of PstP depletion on the survival, cfu were enumerated both in lungs and in the spleen of mice 8 weeks post-infection (Fig. 8, B and C). Gross pathological assessment of lungs at 8 weeks post-infection confirmed the presence of distinct tubercles in the lung tissue of mice infected with Rv and Rv-cd-pstP without ATc, whereas a lesser number of tubercles were observed in the case of mice infected with Rv-cd-pstP that were given doxycycline (Fig. 8D). This was also reflected in the histopathological data and scores, wherein no distinct granulomas were detected in the lungs of mice infected...
with Rv-cd-pstP + doxycycline hydrochloride (Dox) (Fig. 8, E and F). These data corroborated the cfu counts obtained in the lungs as well as the spleen 8 weeks post-infection, which were 1 log-fold (4.0 versus 5.1 and 4.8 for lungs and 2.1 versus 3.3 and 3.3 for spleen cfu) lower in mice infected with Rv-cd-pstP + Dox samples compared with those obtained in Rv and Rv-cd-pstP − Dox-infected mice (Fig. 8, B and C). Taken together, these results underline the importance of PstP for the survival of the pathogen in the host.

Depletion of PstP Decreases the Bacillary Load Even in an Established Infection—To determine the impact of PstP depletion on the survival of bacilli in a well established infection, we infected mice with Rv or Rv-cd-pstP and allowed the infection to progress over 4 weeks. The Rv-cd-pstP-infected mice were then divided into two groups, and only one group was provided Dox. Dox was also provided to Rv infected mice to serve as a control. The bacillary loads in the lungs and spleens of the mice were determined after 1 day, 4 weeks, and 12 weeks post-infection (Fig. 9A, time line diagram). It is apparent from the data in Fig. 9, B and C, that the bacillary load was equivalent after 1 day and 4 weeks in the case of both mice infected with Rv and mice infected with Rv-cd-pstP. Interestingly, depletion of PstP led to ~1 log-fold (3.14 versus 4.49 and 4.08 in lungs and 2.54 versus 3.57 and 3.45 in spleen cfu) decrease in the cfu both in lungs and spleen even in established infection (Fig. 9D). Histopathological analysis of lungs 12 weeks post-infection revealed the presence of a higher number of granulomas in the case of Rv and Rv-cd-PstP. However, in the presence of Dox, we observed a lower number of granulomas and correspondingly lower score in case of Rv-cd-PstP infection (Fig. 9E and F). These data emphasize the need for controlled expression of PstP not only for establishment of infection (Fig. 8), but also for continued maintenance of an established infection.

Discussion

Although the eukaryotic-like phosphosignaling systems in prokaryotes are emerging as regulatory systems that are as important to the cell as their more prominent eukaryotic counterparts, the study of their functions is still in its infancy. Sensing the host environment is key to the appropriate execution of pathogen developmental programs, and mycobacterial phos-
Phosphorylation by phosphatases is critical for regulation of serine/threonine/tyrosine protein kinase-mediated cellular signaling (42, 43). In most prokaryotes, the genes encoding STPKs and corresponding protein phosphatases are carried by the same operon, seemingly following the ratio of one phosphatase per kinase (20, 43). For example, in *Streptococcus pneumoniae*, the phosphatase PhpP interacts with the kinase StkP (the genes encoding both lying in the same operon), and the presence of the kinase is essential for appropriate phosphatase localization. Additionally, cells could only survive deletion of phosphatase gene when the kinase gene *stkP* was also deleted (44). Mycobacteria and corynebacteria, however, encode multiple kinases, but they encode for only one serine/threonine phosphatase belonging to the PPM family (7, 45). The physiological purpose of such an arrangement remains undiscovered. The present study targets the investigation of the physiological impact of overexpressing and depleting PstP on growth and survival of the pathogen both *in vitro* and *in vivo*.

Among the criteria used to classify bacteria is cell shape (46). Multiple bacterial proteins are cell shape regulators. The *pstP* is the first gene of an operon that also carries *rodA*, *pbpA*, *pknA*, and *pknB* genes. *RodA*, named for its role in conferring bacteria with a rodlike shape, is a putative lipid II flippase (47, 48). *PbpA*, although not yet characterized, is predicted to be a transpeptidase required for cross-linking peptidoglycans in the periplasm space (49). Previous studies have shown that overexpression of PknA or PknB in *M. smegmatis* or *Mycobacterium bovis* BCG leads to altered cellular morphology. Whereas the overexpression of PknB results in bulged cells, overexpression of PknA is detrimental to *in vitro* growth and cell survival (50).

**FIGURE 7. Depletion of PstP results in multiseptum phenotype.** A, SEM analysis of *mc2* and *mc2-cd-pstP-ATc* strains grown in the absence or presence of ATc (500 ng/ml) for 12 h was performed as described (50). White arrows in the bottom right panel indicate septa. Scale bar, 1 μm. B, FM4-64 labeling and microscopy of *mc2* and *mc2-cd-pstP-ATc* strains grown in the absence or presence of ATc was performed as described under “Materials and Methods.” The arrows indicate the presence of septa. Scale bar, 5 μm. C, 100 individual cells per cell type per FM4-64 labeling experiment were scored for aseptate, uniseptate, or multiseptate phenotypes. The experiment was performed twice, and the average is represented in the graph. D, TEM analysis of *mc2* and *mc2-cd-pstP-ATc* strains grown in the absence or presence of ATc for 12 h. Processed samples were visualized on grids under FEI G2 Spirit. Scale bar, 0.5 μm. Error bars, S.D.
Our efforts to overexpress PstP using inducible nitrile promoter have not been very fruitful. Although on two occasions, we observed significant overexpression that led to cell death, in most of the experiments, we observed marginally higher expression. We do not have a good explanation for this phenomenon. We speculate that the organism found a way to make the nitrile promoter refractory to inducer concentrations. Interestingly, even with this marginal higher expression of PstP, we observed a statistically significant increase in the mean cell length (Fig. 2). Thus, it appears that the entire operon is involved in the modulation and maintenance of cell morphology and cell division.

To decipher the importance of PstP in regulating cellular events, we started with generating *M. smegmatis* and *M. tuberculosis* pstP conditional mutant strains by carrying out gene replacements (Figs. 3 and 4). The *M. smegmatis* pstP conditional mutant (*mc2-cd-pstP*) displayed almost complete depletion of PstP in the presence of ATc. This depletion of PstP was detrimental to cell survival (Fig. 3). Only partial depletion of PstP was observed in the *M. tuberculosis* pstP conditional
mutant (Fig. 4), but despite this, we observed ~1 log-fold difference in cell survival, suggesting that stringent modulation of PstP expression is critical for optimal growth.

PstP and 9 of the 11 STPKs in M. tuberculosis contain a single transmembrane helix, connected to an extracytoplasmic domain (7, 38). The extracellular domain of PknD has been shown to be involved in sensing osmotic stress (51). The extracellular PASTA domains of PknB interact with muropeptides in the periplasmic space and are essential for in vitro growth of M. tuberculosis (50, 52). The membrane-anchoring transmembrane domain, but not the extracytoplasmic domain of PknA, is found to be essential for in vitro growth (53). For most mycobacterial kinases and the sole phosphatase PstP, very little is known about the functional significance of the membrane-anchoring or the extracytoplasmic domains. The phosphatase domain of PstP has been found to be necessary and sufficient for efficient catalytic activity in vitro (18, 35). We observed that active site mutants of PstP completely failed to rescue growth defects of the M. smegmatis pstP conditional mutant in vitro, indicating the importance of catalytically active PstP phosphatase for cell survival (Fig. 6). Similar experimental analyses with multiple other PstP mutants (Fig. 5) revealed that deletion mutants show marginally compromised survival (0.4–0.6 log-fold lower compared with the wild type). Thus, it appears that all of the domains, including the extracellular domain, are necessary for effective functional complementation.

Cell division is a complex process comprising cell elongation, septum formation, and subsequent cytokinesis, involving a myriad of proteins. Interestingly, several proteins involved in cell division and cell wall synthesis processes have been identi-
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fied to be phosphorylated in phosphoproteomic studies (54–57). Among the 301 proteins identified in a high throughput phosphoproteomic study from *M. tuberculosis* H37Rv, 69 proteins were linked to cell division and cell wall synthesis processes (54). In a recent study, 398 *M. bovis* BCG proteins were identified to be phosphorylated, and 19.5% of these phosphoproteins were involved in various cellular processes (57). Similarly, phosphoproteomic analysis of *M. smegmatis* mc^2_155 and *M. bovis* BCG has revealed the phosphorylation of a number of cell division proteins, such as FtsQ, FtsW, CwsA, FtsK, and FtsY (56). Although all these proteins are important for cell division, FtsZ, a homolog of eukaryotic tubulin harboring GTPase activity, acts as a primer for divisome assembly (58). PknA phosphorlates FtsZ, thus impairing its GTPase activity and polymerization functions (59). Phosphorylation at the carboxyl terminus of FtsZ also impairs its interaction with other divisome proteins, FipA and FtsQ (59). Taken together, these studies confirm the significant role of phosphorylation in modulating the cell division process. Hence, we hypothesized that the depletion of PstP would have an impact on the cell division process. Data from scanning electron microscopy, fluorescence microscopy, and transmission electron microscopy experiments presented in Fig. 7 show the formation of multiple septa upon PstP depletion. Thus, it appears that appropriate dephosphorylation of cell division proteins is critical to orderly cell elongation and cytokinesis.

Phosphatases play an important role in determining the virulence of the pathogen. PtpA and PtpB, secretory tyrosine phosphatases of *M. tuberculosis*, have been shown to be crucial for intracellular survival of the pathogen (25, 27). However, to date, the impact of PstP depletion on mycobacterial survival in the host has not been assessed. Data from mouse infection experiments that we have carried out indicate that even partial depletion of PstP compromises pathogen survival by ~10-fold (Fig. 8). Interestingly, partial depletion of PstP from an established infection (4 weeks post-infection) also led to a ~10-fold decrease in the survival, signifying the need for continued expression of PstP for the maintenance of *M. tuberculosis* infection (Fig. 9). Specific inhibitors of PtpA and PtpB with useful IC\_50 values have been recently identified (60, 61). Standard inhibitors of PP2C phosphatases, such as cyclosporine and sodium fluoride, are not very effective against PstP. Taken together, these findings underline the importance of PstP both for *in vitro* growth and *in vivo* survival and suggest that PstP could be an effective target for therapeutic intervention.

**Materials and Methods**

**Bacterial Strains and Reagents—** A list of constructs generated and strains used in the study is presented in Table 1. Sequences and descriptions of the oligonucleotides used in the study are provided in supplemental Table 1. All medium components were purchased from BD Biosciences. Restriction endonucleases and DNA-modifying enzymes were procured from New England Biolabs and MBI Fermentas. Oligonucleotide primers and analytical grade chemicals were procured from Sigma-Aldrich or GE Healthcare. α-PstP, α-PknB, and α-GroEL-1 antibodies for immunoblotting were raised in the laboratory. Electron microscopy reagents were procured from Electron Microscopy Sciences. Doxycycline hydrochloride was purchased from Biochem Pharmaceutical (Mumbai, India). The pENTR/directional TOPO cloning kit was purchased from Invitrogen. Mycobacterial shuttle plasmids pNit-1 (Kan\(^\prime\)) (36), pJV53 (Kan\(^\prime\)) (62), and pNit-ET (Kan\(^\prime\)) (63) were kind gifts from Dr. Christopher Sassetti, Dr. Graham Hatfull, and Dr. Eric Rubin, respectively.

**Generation of Plasmid Constructs—** The pstP gene was amplified from *M. tuberculosis* genomic DNA with adaptor primers carrying Ndel and HindIII sites at the 5′-ends of the forward and reverse primers, respectively, using Phusion DNA polymerase (New England Biolabs). The amplicon was digested with Ndel-HindIII and cloned into the corresponding sites in pNit1 and pST-KirT (64) to generate pNit1-pstP and pST-KirT-pstP constructs, respectively. pNit1, pJV53, and pNit-ET constructs were modified by replacing the *kan\(^\prime\)* gene with the *apra\(^\prime\)* gene from pMV261-apa (a kind gift from Dr. William Jacobs), to generate pNitA, pJV53A, and pNitA-ET, respectively. PstP deletion mutants were created by amplifying specific fragments of the gene with adaptor primers carrying Ndel-HindIII sites and cloning the amplicons into pENTR vector. This was followed by subcloning the Ndel-HindIII fragments into the corresponding sites in pNit-apa vector. PstP point mutants created in a previous study (35) were subcloned into pNitA vector. Details of constructs used in the study are provided in Table 1.

**Generation of *M. smegmatis* and *M. tuberculosis* Conditional Gene Replacement Mutants—** *M. tuberculosis* H37Rv and *M. smegmatis* mc^2_155 strains were electroporated with integration-proficient pST-KirT-pstP (with PstP containing an amino-terminal FLAG tag) construct, to generate merodiploid strains H37Rv::KirT-pstP and mc^2::KirT-pstP. These strains were then electroporated with pNitA-ET and pJV53A, respectively, to generate recombinocering proficient strains. These strains were electroporated. The hygromycin resistance gene (*hyg*) along with its promoter was amplified from pYUB1474 construct (65) (a kind gift from Dr. William Jacobs) using adaptor primers carrying PflMI sites compatible with those found in pYUB1474. The reverse primer of *hyg* was designed such that the stop codon would be replaced with an amino acid spacer (GGSGG). The amplicon obtained was cloned into pENTR vector to generate pENTR-*hyg*. The apramycin resistance gene (*apra*) was amplified from pMV261A and cloned into pENTR vector to generate pENTR-*apra*. Upstream and downstream flanks (~1 kb on either side) of *pstP* were amplified from genomic DNA of *M. tuberculosis* and *M. smegmatis*. The forward primer of the downstream (3′) flank was designed to carry a PflMI site along with the last seven amino acids of PstP, such that these amino acids would be in frame with the *hyg* gene. We also introduced an EcoRV site in the 5′-flank forward and 3′-flank reverse primers after the PflMI site. Upstream, downstream flanks digested with PflMI were ligated with the *hyg* gene from pENTR-*hyg*, the apramycin resistance gene from pENTR-*apra*, and oriE and Acos fragment from pYUB1474 to generate allelic exchange substrates. The AES cassette containing 5′-flank-*hyg*-3′-flank was released with EcoRV digestion, and the cassette was electroporated into recombinocering-proficient merodiploid strains. Induction of recombinocering genes before competent cell preparation was performed as described earlier (62). Con-
Plasmids and strains used in the study

| Description | Source |
|-------------|--------|
| pST-KirT | Integrative MTB expression vector with N-terminal FLAG tag and r-tetR cloned in SmaI site; Kan’ |
| pNit-1 | IVN inducible MTB expression vector; Kan |
| pNit-ET | Construct containing p60 and p61 genes from Cfe9c phase under nitrile-inducible promoter; Kan’ |
| pV3 | Construct containing p60 and p61 genes from Cfe9c phase under acetamide inducible promoter; Kan’ |
| pMV-261A | E. coli-mycobacterium shuttle vector, Ap’ |
| pNitA | pNit-1 construct wherein kan’ gene with the apra’ gene from pMV261A; Apra’ |
| pNitA-ET | pNit-ET construct wherein kan’ gene was replaced with the apra’ gene from pMV261A; Apra’ |
| pV53A | pV53 construct wherein kan’ gene was replaced with the apra’ gene from pMV261A; Apra’ |
| pKirT-pstP | pstP cloned into NdeI-HindIII sites of pST-KirT; Kan’ |
| pNit1-pstP | pstP cloned into NdeI-HindIII sites of pNit-A; Kan |
| pNitA-pstP | pstP cloned into NdeI-HindIII sites of pNit-A; Apra’ |
| pNitA-pstP-TM | pstP-TM cloned into NdeI-HindIII sites of pNitA; Apra’ |
| pNitA-pstP-JM | pstP-JM cloned into NdeI-HindIII sites of pNitA; Apra’ |
| pNitA-pstP-PstP | pstP cloned into NdeI-HindIII sites of pNitA; Apra’ |
| pNitA-pstP20G | pstP20G cloned into NdeI-HindIII sites of pNitA; Apra’ |
| pNitA-pstP222G | pstP222G cloned into NdeI-HindIII sites of pNitA; Apra’ |

Strains

| Description | Source |
|-------------|--------|
| DH5a | E. coli strain used for cloning experiment |
| mc | M. smegmatis mc155 strain |
| Rv | M. tuberculosis H37Rv strain |
| Rv-pNitA-ET | Rv strain electroporated with pNitA-ET construct |
| mc::pV53A | mc strain electroporated with pV53A construct |
| mc::pKirT-pstP | mc strain electroporated with pST-KirT-pstP construct |
| Rv::pKirT-pstP | Rv strain electroporated with pST-KirT-pstP construct |
| mc-cd-pstP | pstP conditional mutant in mc2::KirT-pstP merodiploid strain |
| Rv-cd-pstP | PstP conditional mutant in Rv::KirT-pstP merodiploid strain |
| mc-cd-pstP-pstP | mc-cd-pstP strain complemented with pNitA-pstP construct |
| mc-cd-pstP-pstP-TM | mc-cd-pstP strain complemented with pNitA-pstP-TM construct |
| mc-cd-pstP-pstP-JM | mc-cd-pstP strain complemented with pNitA-pstP-JM construct |
| mc-cd-pstP-pstP-PstP | mc-cd-pstP strain complemented with pNitA-pstP-PstP construct |
| mc-cd-pstP-pstP20G | mc-cd-pstP strain complemented with pNitA-pstP20G construct |
| mc-cd-pstP-pstP222G | mc-cd-pstP strain complemented with pNitA-pstP222G construct |

Additional mutants mc2-cd-pstP and Rv-cd-pstP were screened by PCR amplification using specific primer sets to confirm that genuine recombination had occurred at the native locus (Figs. 3 and 4).

Growth Pattern Analysis—M. smegmatis mc2 wild type and mc2-cd-pstP mutant strains were electroporated with pST-KirT-pstP, pNit1-pstP, pNitA-pstP, and pNitA-pstP mutant (deletion mutants or point mutants) constructs to generate recombinant strains. The cultures were grown in 7H9 medium supplemented with 10% ADC (albumin, dextrose, and catalase) or in 7H10 agar medium supplemented with 0.6% OADC (oleic acid, albumin, dextrose, and catalase). To analyze growth on plates, cultures were diluted to an A600 of ~0.1 and streaked on 7H10 agar plates containing either no antibiotics or ATc (50 ng/ml). To analyze growth patterns in liquid culture, cultures were initiated in triplicates at an A600 of ~0.02, either in the presence or absence of ATc, and absorbance at 600 nm was measured every 3 h for 30 h. The data were plotted using GraphPad Prism version 6 software. To enumerate cfu at 0 and 30 h, cultures were harvested, washed twice with PBST (PBS containing 0.05% Tween 80), resuspended, and serially diluted, and containing OADC. In the case of M. tuberculosis (Rv) or M. tuberculosis mutant (Rv-cd-pstP), cultures were seeded at an A600 of ~0.1, either in the presence or absence of ATc, and cultures were grown for 4 days. Cfu were determined on day 0, day 2, and day 4 on 7H10 agar plates in the absence of antibiotics.

Western Blotting Analysis—Cultures of M. smegmatis mc2, mc2::KirT-pstP, mc2::pNit1-pstP, mc2-cd-pstP, or mc2-cd-pstP: pstP wt/mutant, strains seeded at an A600 of ~0.1 were grown for different times in the absence or presence of ATc or IVN or ATc + IVN. In the case of M. tuberculosis strains Rv or Rv-cd-pstP, cultures seeded at an A600 of 0.1 were grown in the absence or presence of ATc for 4 days. Whole cell lysates were isolated from harvested cells and analyzed by Western blotting as described earlier (53).

Scanning Electron Microscopy—Cultures of M. smegmatis mc2 or mc2-cd-pstP strains were initiated at an A600 of 0.1 and grown for 12 h in the absence or presence of ATc in filtered 7H9 medium. Similarly, Rv and Rv-cd-pstP cultures were initiated at an A600 of ~0.1 and grown for 4 days in the absence or presence of ATc in filtered 7H9 medium. Cells were harvested, and SEM analysis was performed as described earlier (50).

Transmission Electron Microscopy—M. smegmatis mc2 and mc2-cd-pstP strains were grown until mid-log phase in the absence or presence of ATc. The cells were harvested at 4,300 × g and washed three times with 100 mm sodium phosphate buffer (pH 7.4). Transmission electron microscopy was performed as described earlier (66).

Immunofluorescence Analysis—FM4-64 labeling methodology was used to visualize mycobacteria by fluorescence. For this, 7H9-agarose pads were prepared on frosted slides (Corning Microslide Frosted; 75 × 25 mm) using an AB gene frame (Thermo Scientific; 17 × 54 mm). The 7H9-agarose pads comprised high resolution low melting agarose (Sigma; 1.5%) in 7H9 medium supplemented with ADC. mc2 or mc2-cd-pstP cultures at an A600 of ~0.6 were diluted to an A600 of ~0.1, and 1 μl of diluted culture of either mc2 or mc2-cd-PstP was spread on the
agarose pad (either with or without ATc (50 ng/ml)). The agarose pads were supplemented with FMM4-64 (2 μg/ml) (67). Image acquisition was performed on a Leica TCS SP8 confocal laser scanning microscope at ×63 oil immersion with 3 × optical zoom (68).

Infection of Mice with Mycobacteria—Rv and Rv-cd-PstP cells were grown to mid-log phase and processed, and mice were infected with these cells, as described (50). BALB/c mice of either sex that were 6–8 weeks old (obtained from the National Institute of Immunology breeding facility) were infected by the aerosol route. The lung bacillary loads were enumerated 24 h post-infection to determine the implantation dosage. For reducing PstP levels in mice infected with Rv-cd-PstP, Dox was supplied at a concentration of 1 mg/kg with 5% sucrose in the drinking water. Mice were dissected at the desired time point, bacillary loads were determined, and histopathology of lung sections was performed as described earlier (50).

Author Contributions—A. K. S., V. K. N., and Y. S. conceived and designed the experiments. A. K. S., D. A., A. G., and L. K. S. performed the experiments. V. K. N., Y. S., A. K. S., D. A., V. M., and A. S. analyzed the data. V. K. N., Y. S., A. S., and A. K. S. wrote the paper.

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