The Polyphosphate Kinase Gene ppk2 Is Required for *Mycobacterium tuberculosis* Inorganic Polyphosphate Regulation and Virulence

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ABSTRACT The *Mycobacterium tuberculosis* gene *Rv3232c*/*MT3329* (*ppk2*) encodes a class II polyphosphate kinase, which hydrolyzes inorganic polyphosphate (poly P) to synthesize GTP. We assessed the role of *ppk2* in *M. tuberculosis* poly P regulation, antibiotic tolerance, and virulence. A *ppk2*-deficient mutant (*ppk2*::Tn) and its isogenic wild-type (WT) and complemented (Comp) strains were studied. For each strain, the intrabacillary poly P content, MIC of isoniazid, and growth kinetics during infection of *J774* macrophages were determined. Multiplex immunobead assays were used to evaluate cytokines elaborated during macrophage infection. The requirement of *ppk2* for *M. tuberculosis* virulence was assessed in the murine model. The *ppk2*::Tn mutant was found to have significantly increased poly P content and a 4-fold increase in the MIC of isoniazid relative to the WT and Comp strains. The *ppk2*::Tn mutant showed reduced survival at day 7 in activated and naive *J774* macrophages relative to the WT. Naïve *ppk2*::Tn mutant-infected macrophages showed increased expression of interleukin 2 (IL-2), IL-9, IL-10, IL-12p70, and gamma interferon (IFN-γ) relative to WT-infected macrophages. The *ppk2*::Tn mutant exhibited significantly lower lung CFU during acute murine infection compared to the control groups. *ppk2* is required for control of intrabacillary poly P levels and optimal *M. tuberculosis* growth and survival in macrophages and mouse lungs.

IMPORTANCE *Mycobacterium tuberculosis*, the causative agent of tuberculosis (TB), is a highly successful human pathogen because it has developed mechanisms to multiply and survive in the lungs by circumventing the immune system. Identification of virulence factors responsible for *M. tuberculosis* growth and persistence in host tissues may assist in the development of novel strategies to treat TB. In this study, we found that the mycobacterial enzyme polyphosphate kinase 2 (PPK2) is required for controlling intracellular levels of important regulatory molecules and for maintaining susceptibility to the first-line anti-TB drug isoniazid. In addition, PPK2 was found to be required for *M. tuberculosis* growth in the lungs of mice, at least in part by suppressing the expression of certain key cytokines and chemokines by inactivated lung macrophages.

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Tuberculosis (TB) remains a major global health problem (1, 2). One of the major obstacles to global TB eradication efforts is the prolonged treatment course required to cure active TB, which is believed to reflect the organism’s ability to remain in a nonreplicating persistent state with reduced susceptibility to conventional antituberculous drugs.

Inorganic polyphosphate (poly P), a linear polymer of many tens or hundreds of inorganic phosphate residues linked by high-energy phosphoanhydride bonds, appears to play an important regulatory role in the transition to bacterial persistence (3, 4). Bacteria accumulate poly P intracellularly when they encounter growth-limiting conditions, such as phosphate depletion, amino acid starvation, or osmotic stress (3, 5). Poly P accumulation has been shown to modulate many different bacterial processes, including protein synthesis, nucleotide balance, lipid metabolism, energy utility, and susceptibility to antibiotics (3, 4). Bacterial polyphosphate kinases are of two classes. Polyphosphate kinase 1 (PPK1) is responsible for poly P synthesis through hydrolysis of ATP (3). Although PPK2 enzymes synthesize poly P, they also function as GTP synthases, using poly P as a substrate (3, 6–9). Deletion of the *ppk2* gene in *Corynebacterium glutamicum* reduced PPK activity and intrabacterial poly P content, while overexpression of *ppk2* increased PPK activity and poly P content (10). Conversely, *ppk2* deficiency in *Campylobacter jejuni* led to decreased poly P-dependent GTP synthesis and increased intracellular ATP/GTP ratio (9). Mutation of *ppk2* attenuates bacterial survival under different growth-limiting conditions (7, 9, 10). In addition, PP2K plays an important role in invasion and intracellular survival of *C. jejuni* in human intestinal epithelial cells (9).

The *Mycobacterium tuberculosis* genome contains genes that encode both PPK1 (*Rv2984*) and PPK2 (*Rv3232c*), which play a role in poly P synthesis (7, 11). PPK1 is involved in the stress-induced *mpRAB-sigE-rel* signaling pathway in mycobacteria (11) and appears to be encoded by an essential gene in *M. tuberculosis* (12). *M. tuberculosis* PPK2 is an octamer that catalyzes poly P-dependent phosphorylation of ADP to ATP at a rate >800-fold higher than that of poly P synthesis (13). PPK2 also regulates the intracellular nucleotide pool and is required for survival of...
FIG 1 Complementation of ppp2::Tn. (A) RT-PCR analysis of wild-type *M. tuberculosis* CDC 1551 expression of intergenic regions from *MT3331* to *desA3/MT3326* during late log phase. Chromosomal DNA was used as a control. Primer sets 1 to 5 (Table 1) target specific intergenic areas, as indicated in the figure; primer set 6 targets ppp2. (B) PCR analysis of genomic DNA from the wild-type, ppp2::Tn, and ppp2::Tn Comp strains. Primer set A targets the ppp2 gene (bp 50 to 430), yielding a 480-bp product in the wild type, a 2,547-bp product in the ppp2::Tn mutant, and both products in the ppp2::Tn Comp strain. Primer set B targets bp 108 to 661 of ppp2, yielding a 553-bp product in all strains. (C) Southern hybridization demonstrating binding of the ppp2-specific probe to KpnI-digested fragments of the expected size in the wild type (1,669 bp), ppp2::Tn mutant (2,266 bp), and ppp2::Tn Comp strain (2,266 and 6,095 bp). (D) RT-PCR analysis of ppp2 expression in the wild-type, ppp2::Tn, and ppp2::Tn Comp strains during early stationary phase of growth.
M. smegmatis under acidic, heat, and hypoxic conditions (7). In addition, deficiency of ppk2 or ppk1 is associated with decreased mycobacterial survival during macrophage infection (7, 11). However, the role of PPK2 in M. tuberculosis virulence is unknown.

In this study, we investigated the role of ppk2 on M. tuberculosis poly P homeostasis and susceptibility to the cell wall-acting agent isoniazid. Next, we determined the impact of poly P homeostasis and susceptibility to the cell wall-acting agent known.

RESULTS

ppk2 is coexpressed as part of a three-gene operon. In order to test the role of ppk2 (Rv3232c/MT3331) in M. tuberculosis virulence, a ppk2-deficient mutant (ppk2::Tn) was generated (14). To guide our complementation strategy, we studied potential cotranscription of genes flanking ppk2 using reverse transcription-PCR (RT-PCR). ppk2 is located within a putative operon comprising Rv3234c/MT3331, Rv3233c/MT3330, Rv3232c/MT3329, Rv3231c/MT3328, Rv3230c/MT3327, and Rv3229c/MT3326. As shown in Fig. 1A, the two genes upstream of ppk2 (Rv3234c/MT3331 and Rv3233c/MT3330) are cotranscribed with ppk2, but the three downstream genes are expressed independently in a single transcriptional unit.

A DNA fragment containing the Rv3234c/MT3331, Rv3233c/MT3330, and ppk2 genes, as well as 269 bp of the 5' flanking sequence of MT3331 was cloned into the pMH94H plasmid and then recombined into ppk2::Tn to generate a ppk2 complemented strain (ppk2::Tn Comp). PCR amplification with primers targeting ppk2 (Table 1) yielded products of 487 bp and 2,554 bp in the wild-type and ppk2::Tn strains, respectively, and both products were present, as expected, in the ppk2::Tn Comp strain (Fig. 1B). Southern hybridization, using a specific probe targeting ppk2 (Table 1) after restriction digestion of genomic DNA with KpnI confirmed the generation of a 1,669-bp fragment in the wild-type strain and a 2,266-bp fragment in the ppk2::Tn strain (Fig. 1C).

Southern hybridization, using a specific probe targeting ppk2 (Table 1) after restriction digestion of genomic DNA with KpnI confirmed the generation of a 1,669-bp fragment in the wild-type strain and a 2,266-bp fragment in the ppk2::Tn strain (Fig. 1C). The two expected products (2,266 bp and 6,095 bp) were observed in the ppk2::Tn Comp strain. As confirmation of functional ppk2 complementation in the ppk2::Tn Comp strain, partial restoration of ppk2 expression was observed during early stationary phase (Fig. 1D). RT-PCR demonstrated that expression of the two up-

| TABLE 1 Primers used in this study | Direction | Sequence |
|-------------------------------------|-----------|----------|
| Primer set 1 F                       | GTAAGGAATTCCGGCCAACCT |
| Primer set 1 R                       | TACATCTGGCGACTGGGTTCC |
| Primer set 2 F                       | CCAAGCTTGCTGGAGAGGCTAC |
| Primer set 2 R                       | CGGGAGTAAGGGCCAAATAAA |
| Primer set 3 F                       | GTGAAAAGCCCAAAGTCAA |
| Primer set 3 R                       | ACCTGTAGCTGCGATGATGG |
| Primer set 4 F                       | ACCAAGACTGCGCATTTCTG |
| Primer set 4 R                       | CACAATTGCTGCGCTTTTAT |
| Primer set 5 F                       | CGAAGTGGGAAAGGGTGAG |
| Primer set 5 R                       | CGGATCTCCATGTTTTGGAT |
| Primer set 6 (ppk2) F                | GTGATATACCATCCGTTGATGTG |
| Primer set 6 (ppk2) R                | Rv0496 |
| Primer set 6 F                       | GCGGAGCTC GTACCTGACGAGGGTTG |
| Primer set 6 R                       | GGGTCTAGACAAACGGACTGGTGCAACA |
| Primer set A F                       | GCATCGAAGTTGTATTTAAAAAACGGCCAGTGAAAT |
| Primer set A R                       | AAGCACTGGAAGCAGCTTCTG |
| Primer set B F                       | Same as primer set 6 |
| Primer set B R                       | GCCTCATCAAGAACGCGATTTG |
| Primer set B F                       | Rv1026 |
| Primer set B R                       | GACAGATCCGCTCTTATCGAC |
| Primer set 2 F                       | AATCTGATGCGCTGTGAT |
| Primer set 2 R                       | TGGTGAAGCTGACAGTCTT |
| Primer set 1 F                       | CTCAGGACGCACTGAAAGAC |
| Primer set 1 R                       | TGGATTTCCAGCACCTTCTC |
| Primer set 4 F                       | TGGTGAAACGTCAGCAGTTC |
| Primer set 4 R                       | TTTCCACCGCTCTGAT |
| Primer set 3 F                       | TTTCCACCGCTCTGAT |
| Primer set 3 R                       | CGGGAAGATGGGCAATATAA |
| Primer set 6 F                       | TCAGATGCGCTGTGAT |
| Primer set 6 R                       | GCACCATCATTCTG |
| sigA primer set F                    | Rv3234c/MT3331, Rv3233c/MT3330, Rv3232c/MT3329, Rv3231c/MT3328, Rv3230c/MT3327, and Rv3229c/MT3326. As shown in Fig. 1A, the two genes upstream of ppk2 (Rv3234c/MT3331 and Rv3233c/MT3330) are cotranscribed with ppk2, but the three downstream genes are expressed independently in a single transcriptional unit. A DNA fragment containing the Rv3234c/MT3331, Rv3233c/MT3330, and ppk2 genes, as well as 269 bp of the 5' flanking sequence of MT3331 was cloned into the pMH94H plasmid and then recombined into ppk2::Tn to generate a ppk2 complemented strain (ppk2::Tn Comp). PCR amplification with primers targeting ppk2 (Table 1) yielded products of 487 bp and 2,554 bp in the wild-type and ppk2::Tn strains, respectively, and both products were present, as expected, in the ppk2::Tn Comp strain (Fig. 1B). Southern hybridization, using a specific probe targeting ppk2 (Table 1) after restriction digestion of genomic DNA with KpnI confirmed the generation of a 1,669-bp fragment in the wild-type strain and a 2,266-bp fragment in the ppk2::Tn strain (Fig. 1C). The two expected products (2,266 bp and 6,095 bp) were observed in the ppk2::Tn Comp strain. As confirmation of functional ppk2 complementation in the ppk2::Tn Comp strain, partial restoration of ppk2 expression was observed during early stationary phase (Fig. 1D). RT-PCR demonstrated that expression of the two up-
**ppk2 is required for control of intracellular poly P levels and susceptibility to isoniazid.** Previous studies have shown that PPK2 hydrolyzes inorganic polyphosphate (poly P) to generate GTP from GDP, contributing to a decreased ATP/GTP ratio in *Mycobacterium smegmatis* (7). Consistent with a role for PPK2 in poly P hydrolysis during exponential growth of *M. tuberculosis*, we found that the ppk2::Tn mutant exhibited significantly higher poly P content relative to the isogenic wild-type (*P = 0.0038*) and ppk2::Tn Comp (*P = 0.0028*) strains (Fig. 2A). Quantitative RT-PCR (qRT-PCR) revealed that ppk1 expression was significantly reduced (*P < 0.001*) and expression of the putative exopolyphosphatase Rv1026 was increased (*P = 0.004*) in the ppk2::Tn mutant relative to the wild type (Fig. 2B), consistent with negative- and positive-feedback mechanisms, respectively. However, expression of the exopolyphosphatase gene Rv0496 was slightly decreased in the ppk2::Tn mutant than in the wild type (*P = 0.028*). Consistent with the hypothesis that poly P accumulation is associated with *M. tuberculosis* tolerance to the cell wall-active agent isoniazid (15), the MIC of isoniazid increased from 0.06 μg/ml for the wild-type and ppk2::Tn Comp strains to 0.24 μg/ml for the ppk2::Tn mutant strain. As in the case of *M. smegmatis* (7), *M. tuberculosis* ppk2 deficiency was associated with a higher ATP/GTP ratio relative to the wild-type (*P = 0.07*) and ppk2::Tn Comp (*P = 0.02*; Fig. 2C) strains. Each experiment was performed using three biological replicates.

**ppk2 is required for *M. tuberculosis* survival during infection of naive macrophages and suppression of proinflammatory cytokine release.** On the basis of previous reports, ppk2 appears to play an important role in *M. tuberculosis* intracellular survival (7). To further investigate the mechanism of defective growth of the ppk2::Tn mutant, we infected naive and activated murine J774 macrophages with each of the three strains. The numbers of intracellular CFU were lower in gamma interferon (IFN-γ)-activated macrophages 7 days after infection with the ppk2::Tn mutant than in the wild type (*P = 0.04*). As in the case of *M. smegmatis* (7), *M. tuberculosis* ppk2 appears to play an important role in *M. tuberculosis* intracellular survival (7).

**FIG 2** ppk2 is required for *M. tuberculosis* control of intracellular poly P levels and ATP/GTP ratio. (A) Intrabacillary poly P levels were measured in the wild-type, ppk2::Tn, and ppk2::Tn Comp strains during mid-log phase using a DAPI-based method and normalized to total protein content of extract lysate. Values that were significantly different (*P < 0.05*) are indicated by a bar and asterisk. Values are means ± standard deviations (error bars) for three biological replicate samples. (B) qRT-PCR analysis of gene expression of ppk2::Tn mutant and ppk2::Tn Comp strain compared to that of the wild type during late log phase of growth in supplemented Middlebrook 7H9 broth, expressed as a change in the normalized cycle threshold (ΔΔCt). (C) Intrabacillary ATP/GTP ratio of the wild-type, ppk2::Tn, and ppk2::Tn Comp strains during late log phase, as measured by HPLC. The values for ppk2::Tn and ppk2::Tn Comp strains were significantly different (*P < 0.05*) as indicated by the bar and asterisk. The values for the ppk2::Tn mutant and the wild-type strain were not significantly different (*P = 0.07*). Three biological replicate samples were used for each strain in each experiment.
(MCP-1) \((P = 0.02)\), macrophage inflammatory protein 1b (MIP-1b) \((P = 0.011)\), and RANTES (regulated upon activation, normal T cell expressed and secreted) \((P = 0.004)\) were more highly induced by the \(ppk2\):Tn mutant than by the wild-type and \(ppk2\):Tn Comp strains, despite similar intracellular CFU values (Fig. 4). Each macrophage CFU and cytokine experiment was performed using three biological replicates.

**ppk2** enhanced *M. tuberculosis* growth and survival in mouse lungs. In order to test the role of \(ppk2\) in *M. tuberculosis* virulence in the mammalian host, separate groups of BALB/c mice were aerosol infected with the wild type \((1.82 \pm 0.1)\), \(ppk2\):Tn mutant \((2.24 \pm 0.06)\), and \(ppk2\):Tn Comp strain \((1.9 \pm 0.1)\) (all values \(\log_{10}\) number of CFU/lung). During the first 14 days after infection, wild-type and \(ppk2\):Tn Comp strains showed typical exponential growth, increasing by 4.34 \(\log_{10}\) units and 4.15 \(\log_{10}\) units, respectively. On the other hand, \(ppk2\):Tn showed a severe growth defect during the acute phase of the infection relative to wild type, increasing by only 2.29 \(\log_{10}\) units \((P < 0.001)\) (Fig. 5A). After the onset of adaptive immunity (day 28), the wild-type bacillary burden declined to 5.85 \(\pm\) 0.15 \(\log_{10}\) units, after which a relatively stable lung census was maintained. Interestingly, the lung bacillary burden of the \(ppk2\):Tn mutant continued to increase until day 56, when it was equivalent to that of animals infected with the wild type but then declined during the chronic phase of infection, such that by day 119, it was 0.63 \(\log_{10}\) unit lower than that of wild type \((P = 0.098)\). The \(ppk2\):Tn mutant also showed defective growth in mouse spleens, as the numbers of CFU in \(ppk2\):Tn-infected mouse spleens were significantly lower than the numbers in spleens in mice infected with the wild-type and \(ppk2\):Tn Comp strains at days 28, 84, and 119 \((P < 0.001\) for all time points) (Fig. 5B).

Mouse body weights were not significantly different between groups at any time point (see Fig. S3A in the supplemental material). However, by day 28 after infection, the mean normalized lung (see Fig. S3B in the supplemental material) and spleen (see Fig. S3C) weights of \(ppk2\):Tn-infected mice were significantly lower than the corresponding values of mice infected with the wild type \((P < 0.001\) and \(P = 0.023\), respectively). Gross pathology revealed no major differences in the number or size of lung tubercle lesions between groups at any time point (Fig. S4).

After 28 days of infection, histological evaluation revealed moderate inflammation with a predominance of macrophages and bronchiolar obliteration in the lungs of mice infected with the wild-type and \(ppk2\):Tn Comp strains (Fig. 5C). In lung samples from mice infected with the \(ppk2\):Tn mutant, mild inflammation predominantly comprising lymphocytes was noted. Lymphocytic involvement was observed in 15/41 (36.6%) bronchioles in mice infected with the wild type, 8/50 (16%) in mice infected with the \(ppk2\):Tn mutant, and 12/21 (57.1%) in mice infected with the \(ppk2\):Tn Comp strain. Acid-fast staining revealed more bacilli in lungs in mice infected with the wild-type and \(ppk2\):Tn Comp strains than in mice infected with the \(ppk2\):Tn mutant. After 56 days of infection, the lungs of each group exhibited similar degrees of bronchiolitis, bronchiolar obliteration, intra-alveolar inflammation, and perivascular lymphocytic inflammation, with equal numbers of histiocytes and lymphocytes. Each data point from mouse experiments represents data obtained from 5 animals.

**FIG 3** \(ppk2\) deficiency impairs *M. tuberculosis* growth and survival in J774 macrophages. Growth and survival of the wild type, \(ppk2\):Tn mutant, and \(ppk2\):Tn Comp strain after infection of activated (A) and naive (B) J774 murine macrophages. Values that were significantly different \((P < 0.05)\) are indicated by a bar and asterisk. Three biological replicate samples were used for each strain in each experiment.
DISCUSSION

*M. tuberculosis* is a highly adapted pathogen, with the ability to evade host immunity and persist for prolonged periods within infected tissues. The regulatory molecule poly P has been implicated in the bacterial stringent response (16). The presence of poly P in *M. tuberculosis* was reported several decades ago (17), and recent studies have used scanning transmission electron microscopy together with energy-dispersive X-ray spectroscopy to visualize poly P granules in individual *M. tuberculosis* bacilli during nutrient starvation (18). In addition to its role in inducing the stringent response (11, 19), poly P levels have been shown to influence susceptibility of *M. tuberculosis* to certain TB drugs and *M. tuberculosis* virulence in guinea pigs (20). The *M. tuberculosis* polyphosphate kinase PPK2 catalyzes the synthesis of GTP from GDP using poly P as a phosphate donor (7). In this study, we confirm the role of *ppk2* in *M. tuberculosis* poly P hydrolysis in *vivo* by demonstrating that *ppk2* deficiency leads to increased intracellular poly P content. We also found that poly P accumulation was accompanied by tolerance to the bactericidal drug isoniazid, which targets actively multiplying organisms by inhibiting the mycolic acid synthesis pathway (21). These phenotypes are consistent with those observed for another poly P-accumulating *M. tuberculosis* strain deficient in the exopolyphosphatase Rv0496/PPX1 (exopolyphosphatase 1) (15). In addition, we demonstrate that *ppk2* is an important virulence gene required for exponential growth of *M. tuberculosis* during the acute phase of infection in mouse lungs and for bacillary persistence during the chronic phase of infection.

Previous work has shown that PPK2 plays an important role in maintaining intracellular GTP levels in *M. smegmatis* by directly synthesizing GTP from GDP and poly P and by interacting with the nucleoside diphosphate kinase Ndk, thereby promoting production of GTP over CTP or UTP (7). Our study corroborates the importance of PPK2 in nucleotide pool maintenance in *M. tuberculosis*, as we found that the ATP/GTP ratio was increased in the *ppk2*-deficient mutant. The significance of such changes in the ratios of nucleotides and their relationship to growth arrest and antibiotic tolerance remain to be determined. Similar to our findings, *ppk2* deficiency in *Campylobacter jejuni* is associated with significant survival defects under osmotic, nutrient, aerobic, and antimicrobial stresses and within human intestinal epithelial cells (9).

Although the *ppk2::Tn* mutant was implanted at a higher dose in mouse lungs relative to the control strains, the number of lung CFU in mutant-infected animals was ~1.5 log10 units lower than those infected with the control strains after 14 days of infection. Our macrophage data shed some light on the impaired growth of the *ppk2::Tn* mutant during the acute phase of infection in mouse lungs. Although the number of intracellular CFU of the *ppk2::Tn* mutant was similar or even slightly lower than the numbers of intracellular CFU of the control strains following a multiplicity of infection (MOI) of 1:1, naive macrophages infected with the mutant strain showed significantly greater induction of the proinflammatory cytokines IL-12(p70) and IFN-α than those infected with the control strains after 14 days of infection. Our macrophage data shed some light on the impaired growth of the *ppk2::Tn* mutant during the acute phase of infection in mouse lungs. Although the number of intracellular CFU of the *ppk2::Tn* mutant was similar or even slightly lower than the numbers of intracellular CFU of the control strains following a multiplicity of infection (MOI) of 1:1, naive macrophages infected with the mutant strain showed significantly greater induction of the proinflammatory cytokines IL-12(p70) and IFN-γ, which serve to control *M. tuberculosis* infection in the lungs (22, 23). Furthermore, the *ppk2::Tn* mutant induced significantly greater expression of the chemotactic factors MIP-1β, MCP-1, eotaxin, and RANTES by naive macrophages (24, 25). Finally, *M. tuberculosis* deficiency of *ppk2* was also associated with increased macrophage expression of IL-2, which can control inflammation by interfering with IL-6-dependent Th17 differentiation, and IL-10, which limits immune system-mediated damage to the host following infection (26). The net effect of these changes on macrophage function may have contributed to improved control of the *ppk2::Tn* mutant in mouse lungs, as well as more limited lung histopathology in mutant-infected animals 28 days after infection, as manifested by milder, predominantly lymphocytic infiltration with relative preservation of bronchiolar architecture. Interestingly, the *ppk2::Tn* mutant continued to grow in mouse lungs to wild-type levels by day 56 after infection. Whether continued growth after the onset of adap-
tive immunity is due to compensatory metabolic changes in the mutant (e.g., poly P hydrolysis through increased expression of exopolyphosphatase [PPX]) and/or defective host immunity remains to be determined.

We observed incomplete restoration of all wild-type phenotypes following complementation of \( \text{ppk2} \). Despite our efforts to preserve the native promoter of \( \text{ppk2} \) in the \( \text{ppk2::Tn Comp} \) strain, it is possible that regulation of the gene may be altered at the recombination site, which is supported by the RT-PCR data showing mildly decreased expression of \( \text{ppk2} \) in the \( \text{ppk2::Tn Comp} \) strain relative to the wild type during early stationary phase (Fig. 1D). Another difference between the wild-type and complemented strains is the presence of two copies of the upstream genes \( \text{Rv3233c}/\text{MT3330} \) and \( \text{Rv3234c}/\text{MT3331} \) in the latter. However, the expression levels of these genes did not differ significantly between these two strains by RT-PCR (see Fig. S1 in the supplemental material). Incomplete complementation may be attributed to polar effects of the transposon insertion in \( \text{ppk2} \); however, expression of the two upstream and two downstream genes was not disrupted, as revealed by RT-PCR (Fig. S1).

In summary, our data show that \( \text{ppk2} \) (\( \text{Rv3232c}/\text{MT3329} \)) is required to maintain \( \text{M. tuberculosis} \) homeostasis of poly P and nucleotide pools. In particular, dysregulation of poly P balance in the \( \text{ppk2::Tn} \) mutant may have contributed to phenotypic tolerance to isoniazid. Finally, we show for the first time that PPK2 plays an important role in \( \text{M. tuberculosis} \)-host interactions, perhaps by suppressing macrophage production of proinflammatory cytokines and chemokines, thereby promoting bacillary growth during the acute phase of infection in mouse lungs.

**MATERIALS AND METHODS**

**Bacterial strains.** An \( \text{M. tuberculosis} \) strain deficient in \( \text{ppk2/Rv3232c}/\text{MT3329} \) (\( \text{ppk2::Tn} \)) was generated by transposon mutagenesis of the wild-type \( \text{M. tuberculosis} \) strain CDC 1551 (14). The point of insertion of the \( \text{himar1} \) transposon, which contains a kanamycin resistance cassette, was determined to be at nucleotide position 8 in the \( \text{ppk2} \) gene (27). A DNA fragment containing \( \text{MT3329} \) and the coregulated genes \( \text{MT3330} \) and \( \text{MT3331} \), along with 269 bp of upstream and downstream flanking sequences, was cloned into the mycobacterial integrating vector pMH94H.
using the BspHI and AclI sites to yield a single-copy plasmid conferring hygromycin resistance (28, 29). The complementation plasmid was introduced into the pck2::Tn mutant by electroporation, and transformants were selected on hygromycin-containing 7H10 plates. The complemented strain (pck2::Tn Comp) was confirmed by PCR and Southern blotting. Briefly, genomic DNA from the wild-type, pck2::Tn, and pck2::Tn Comp strains was isolated, digested with KpnI (NEB), separated by 1% agarose gel electrophoresis, and blotted onto positive-charge nylon membranes. The DNA templates were hybridized with PCR-generated digoxigenin-11-dUTP-labeled gene probes recognizing the region of pck2 from 108 to 601 bp. After cross-linking, probe binding was detected with an antidigoxigenin alkaline phosphatase conjugate antibody and developed with diosodium 3-(3,4-methoxydiphenyl)-1,2-dioxetane-3,2’-(3’-chloro)tricyclo[3.3.1.13,7]decan-4-yl) phenyl phosphosphate (CSPD) substrate (Roche Diagnostics).

Determination of intracellular inorganic polyphosphate and nucleotide content. A 4’,6-diamidino-2-phenylindole (DAPI)-based method was used to determine intracellular polyphosphate (poly P) content in each strain (15, 30). Briefly, the bacteria were lysed by bead beating in 5 M guanidinium thiocyanate (GITC; Sigma)–50 mM Tris-HCl (pH 7.0) buffer. The total protein levels of the lysates were determined by colorimetric protein assay (Bio-Rad) and used to normalize poly P content. Poly P was harvested by glass milk from Geneclean III kit (MP Biomedical LLC) and then treated with DNase (Ambion) and RNase (NEB) before elution with distilled water (95°C, pH 8.0). Each elution was further diluted 1:10 in distilled water and mixed with DAPI (Roche Diagnostics) at a final concentration of 100 μg/ml. The poly P concentration was determined by fluorescence of the DAPI-poly P complex following excitation at 415 nm and emission at 525 nm by FLUOstar OPTIMA (BMG Labtech). The fluorescence intensity of DAPI-containing solution was used as a background signal for blanks. Increasing concentrations of poly P (type 65; Sigma-Aldrich) ranging from 2.5 μg/ml to 312.5 ng/ml were used to generate a standard curve. For nucleotide measurements, bacilli were pelleted by centrifugation at 8,000 rpm at 4°C for 10 min. The pellets were resuspended in ice-cold buffer (acetonitrile-methanol-water [40:40:20]) and incubated on ice for 15 min. The cell suspension was heated at 95°C for 10 min and cooled on ice for 5 min. The cell debris was removed by centrifugation, and the supernatant was concentrated by evaporation and reconstituted in distilled water. ATP and GTP (Sigma) were used to generate the reference peak. Cell lysate (50 μl) was injected into a reverse-phase high-performance liquid chromatograph (HPLC) (Waters 2690) fitted with a Sunfire C8 column and UV detector (Waters 2487) to study ATP and GTP contents at the same time. The resulting peaks were analyzed with Waters Empower software, and the area of ATP and GTP peaks was used to determine the ratio for further comparison.

MIC determination. Logarithmically growing wild-type, pck2::Tn, and Comp cultures (5 × 10⁴/ml) were inoculated in 15-ml conical tubes containing 2 ml of Middlebrook 7H9 broth without Tween 80 and with increasing concentrations of isoniazid, as follows: 0, 0.03, 0.06, 0.12, 0.24, 0.48, and 0.96 μg/ml. Bacterial growth was determined by the presence of visible pellets after 10 days of standing culture at 37°C. The MIC was recorded as the lowest concentration of antibiotic for which there was no visible pellet.

Quantitative polymerase chain reaction and RT-PCR. Total RNA was extracted from 50-ml M. tuberculosis cultures and treated with DNase, and cDNA was generated using Superscript III (Invitrogen) (31). cDNA corresponding to each transcript was subjected to 40 cycles of PCR for quantification using gene-specific primers (Table 1) and an iCycler 5.0 (Bio-Rad). The cycle threshold (Ct) value obtained for each gene was normalized with that of the housekeeping gene sigA (32), and the normalized change in Ct (ΔCt) was calculated (15). Reverse transcription-PCR (RT-PCR) was used to determine genes coexpressed with pck2/Rv3232c/MT3329 using the intergenic primers listed in Table 1. RT-PCR was also used to determine the expression of genes upstream and downstream of the pck2 transposon insertion in the mutant and complemented strains (see Fig. S1 in the supplemental material). Primers used for the expression of upstream and downstream genes include the following: for Rv3232c/MT3331, forward (F) primer CAATTGATGTGCGGTTGCTG and reverse (R) primer TCGGATGCCTGCACTACC; for Rv3232c/MT3330, F primer CTGGTCTGAGCCAGGACTAT and R primer AGGTCTCCAGCACCTGGTA; for Rv3321c/MT3328, F primer TACGATGATCCGAGTGGTTGC and R primer CAGAAATGGCAGCTTGTG; for Rv3320c/MT3327, F primer GCTTCGTCACCCCATGTT and R primer GTTTCGAATCGACAT; and for sigA, F primer TGAGGTGATCACAAGACTG and R primer TGATTTCGACCTCCTTC.

Murine infections and virulence endpoints. BALB/c mice (4 to 6 weeks old, female; Charles River Laboratories, Wilmington, MA) were housed in a biosafety level 3, specific-pathogen-free facility and fed water and chow ad libitum. All procedures followed protocols approved by the Institutional Animal Care and Use Committee at Johns Hopkins University. Separate groups of mice were infected with wild-type, pck2::Tn, or pck2::Tn Comp strains (~100 bacilli implanted per animal lung) via the aerosol route using a Glas-Col inhalation exposure system. Five mice in each group were sacrificed at days 1, 4, 28, 56, 84, and 119. Organs were homogenized and plated for CFU determination, and randomly selected sections were fixed for formalin for histological analysis, which was performed by a pulmonary pathologist blind or unaware of the specimen identity (D. A. Belchis). Lung and spleen samples harvested from pck2::Tn-infected mice at the last time point were plated on Middlebrook 7H11 agar plates with or without kanamycin. No difference in CFU was observed, indicating stability of the transposon insertion throughout the entire experiment.

Macrophage infections and cytokine assays. The mouse macrophage-like cell line J774.1 was maintained and infected with each M. tuberculosis strain (33). Macrophages were activated with treatment with mouse gamma interferon (IFN-γ) (Invitrogen) overnight and lipopolysaccharide from Escherichia coli O26:B6 (Sigma) for 3 h before infection. Naïve macrophages were divided 1 day before infection without any treatment. At day 0, 10⁴ macrophages were infected with an equal number of logarithmically growing bacilli of the wild-type, pck2::Tn, and pck2::Tn Comp strains. Intracellular M. tuberculosis was recovered and plated on days 0, 1, 3, 5, and 7. At days 1 and 3, the supernatant of each culture was collected and frozen at −80°C until analysis. Macrophage-secreted cytokines were analyzed by immunobead cytokine assays (mouse cytokine 23-plex assay; Bio-Rad).

Statistical analysis. Data from at least three biological replicates were used to calculate the means and standard deviations (SDs) for graphing purposes. Statistical analysis employed the unpaired Student’s t test, and a P value of <0.05 was considered significant.

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.00039-13/-/DCSupplemental.

Figure S1, TIF file, 0.1 MB.
Figure S2, TIF file, 0.1 MB.
Figure S3, TIF file, 0.3 MB.
Figure S4, TIF file, 1.4 MB.

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