Eukaryotic-Type Ser/Thr Protein Kinase Mediated Phosphorylation of Mycobacterial Phosphodiesterase Affects its Localization to the Cell Wall

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Phosphodiesterase enzymes, involved in cAMP hydrolysis reaction, are present throughout phylogeny and their phosphorylation mediated regulation remains elusive in prokaryotes. In this context, we focused on this enzyme from *Mycobacterium tuberculosis*. The gene encoded by Rv0805 was PCR amplified and expressed as a histidine-tagged protein (mPDE) utilizing *Escherichia coli* based expression system. In kinase assays, upon incubation with mycobacterial Clade I eukaryotic-type Ser/Thr kinases (PknA, PknB, and PknL), Ni-NTA purified mPDE protein exhibited transphosphorylation ability albeit with varying degree. When mPDE was co-expressed one at a time with these kinases in *E. coli*, it was also recognized by an anti-phosphothreonine antibody, which further indicates its phosphorylating ability. Mass spectrometric analysis identified Thr-309 of mPDE as a phosphosite. In concordance with this observation, anti-phosphothreonine antibody marginally recognized mPDE-T309A mutant protein; however, such alteration did not affect the enzymatic activity. Interestingly, mPDE expressed in *Mycobacterium smegmatis* yielded a phosphorylated protein that preferentially localized to cell wall. In contrast, mPDE-T309A, the phosphoablative variant of mPDE, did not show such behavior. On the other hand, phosphomimics of mPDE (T309D or T309E), exhibited similar cell wall anchorage as was observed with the wild-type. Thus, our results provide credence to the fact that eukaryotic-type Ser/Thr kinase mediated phosphorylation of mPDE renders negative charge to the protein, promoting its localization on cell wall. Furthermore, multiple sequence alignment revealed that Thr-309 is conserved among mPDE orthologs of *M. tuberculosis* complex, which presumably emphasizes evolutionary significance of phosphorylation at this residue.

Keywords: signal transduction, cAMP, eukaryotic-type serine/threonine kinase, enzyme activity, mutagenesis, intracellular trafficking, phosphodiesterase

Abbreviations: cAMP, 3',5'- cyclic adenosine monophosphate; mPDE, *M. tuberculosis* phosphodiesterase; MBP, Maltose Binding Protein; His6, 6X Histidine tag; OADC, Oleic acid albumin dextrose catalase growth enrichment; LC-MS/MS, Liquid chromatography coupled tandem mass spectrometry; *Msm*, *M. smegmatis*.
INTRODUCTION

Cells respond to various environmental changes by an orchestra of events and transfer information through the process of signal transduction (McDonough and Rodriguez, 2012). The cascade of events involve identification as well as amplification of signals resulting in gene(s) expression that regulates cellular physiology of an organism and ultimately helps to better acclimatize in the adverse environs for its survival. In this direction, we concentrated on bacterial signaling, especially focussing on pathogens like Mycobacterium tuberculosis, the causative agent of the disease tuberculosis (Chevalier et al., 2014). In fact, M. tuberculosis is considered to be one of the most successful pathogen as it is capable of withstanding host defense mechanism(s) for its survival (Chevalier et al., 2014; Dey and Bishai, 2014; Orme, 2014). Such compliance might be attributed to its ability to camouflage within the micro-environment of the host and exhibit atypical or unique signal transduction mechanisms. In this context, signaling involving small molecules such as 3′, 5′-cyclic AMP, cyclic di-AMP, cyclic di-GMP etc., are less explored (Hong et al., 2013; Zaveri and Visweswariah, 2013; Manikandan et al., 2014). These cyclic nucleotide levels are controlled by different phosphodiesterases by promoting their hydrolytic degradation and thus, play a crucial role in the process (Shenoy et al., 2005; Hong et al., 2013; Yang et al., 2014).

Analysis of M. tuberculosis genome sequence revealed the presence of an exceptionally high 3′, 5′- cAMP synthesizing machinery in the form of adenylate cyclases compared to other micro-organisms. While 16 such genes have been identified in M. tuberculosis genome (Shenoy and Visweswariah, 2006; Zaveri and Visweswariah, 2013), there is only one phosphodiesterase (mPDE) enzyme that antagonizes cAMP action through its degradation into 5′-AMP (Shenoy et al., 2005; Chakraborti et al., 2011). Apart from cAMP hydrolysis, mPDE is also involved in altering cell wall permeability (Podobnik et al., 2009). The mPDE has already been characterized and its structure was determined using X-ray crystallographic studies (Shenoy et al., 2007). While its N-terminal catalytic domain is responsible for the enzyme activity, the C-terminal region mediates its localization (Podobnik et al., 2009; Matange et al., 2014).

The presence of ortholog of mPDE in the minimal genome of Mycobacterium leprae further signifies its importance in mycobacterial signal transduction (Shenoy et al., 2005). PDEs in humans are regulated by phosphorylation through Ser/Thr kinases like cAMP- dependent protein kinase A (PKA) or protein kinase B (PKB), thus influencing their catalytic ability (Macphee et al., 1988; Kitamura et al., 1999; Lindh et al., 2007; Bessay et al., 2008). To our surprise, till date there have been no reports of phosphorylation mediated regulation of any bacterial PDE, including that from M. tuberculosis. Although M. tuberculosis possesses highly complex as well as dynamic phosphorylation/dephosphorylation machinery comprising of 11 eukaryotic-type Ser/Thr kinases and the cognate phosphatase (Cole et al., 1998; Chakraborti et al., 2011), their involvement in this process is yet unknown. Recent studies indicated that these kinases have several cellular targets and they regulate a plethora of metabolic events in prokaryotes (Pereira et al., 2011). Bioinformatic analysis of these 11 mycobacterial kinases categorized them in five groups, Clades I-V (Narayan et al., 2007). Among these kinases, we concentrated on PknA from M. tuberculosis belonging to Clade I. Like other eukaryotic-type Ser/Thr kinases, PknA has a characteristic domain structure and it is predominantly phosphorylated at threonine residues (Chaba et al., 2002; Thakur et al., 2008; Ravala et al., 2015). Available literature also established the role of PknA in morphological changes associated with bacterial cell division (Chaba et al., 2002; Kang et al., 2005) by interacting with FtsZ and MurD. Notably, the FtsZ protein and MurD enzyme play a crucial role in bacterial cytokinesis and peptidoglycan respectively (Thakur and Chakraborti, 2006, 2008).

In this study, we provide evidence that mPDE is transphosphorylated by PknA. Among other mycobacterial Clade I kinases, we noticed PknB and PknL were also able to phosphorylate mPDE albeit with varying magnitude. Utilizing PknA as a representative of mycobacterial eukaryotic-type Ser/Thr kinases, its co-expression with mPDE in Escherichia coli yielded a phosphorylated His-tagged protein. LC-MS/MS identified Thr-309 as the phosphorylatable residue in this protein, which is located within a short but de-structured C-terminal domain of mPDE (Podobnik et al., 2009; Matange et al., 2014). The mPDE upon expression in Mycobacterium smegmatis was recognized by anti-phosphothreonine antibody. Further, our analysis on phosphoablative and phosphomimetic substitutions of the phospho-threonine residue indicated the role of eukaryotic-type Ser/Thr kinase mediated phosphorylation of mPDE for its cell wall association. Interestingly, Thr-309 is conserved amongst mPDE orthologs of M. tuberculosis complex portraying its probable evolutionary significance.

MATERIALS AND METHODS

Materials

Restriction enzymes were procured from New England Biolabs (USA). Other fine biochemicals like ATP, Tris, NaCl, maltose, MnCl₂, MgCl₂, trypsin and dephosphorylated α-casein were procured from Sigma (USA). Antibodies like anti-phosphothreonine (Cell Signalling Technology, USA) and anti-His (GE Healthcare, USA) were commercially available. Phosphodiesterase assay kit was the product of Enzo®Lifesciences (USA). [γ-³²P] ATP was obtained from Board of Radiation and Isotope Technology (BRIT, India). Desalted oligonucleotides were custom synthesized from Sigma.

Cloning, Expression, and Site Directed Mutagenesis

Different genes were isolated through PCR amplification using genomic DNA from M. tuberculosis strain H37Rv as the template. In fact, sequences of mPDE (Rv0805), pknB (Rv0014c), and pknL (Rv2176) were identical at nucleotide level between pathogenic (H37Rv) and non-pathogenic (H37Ra) strains of M. tuberculosis. For amplification of mPDE gene, PCR (denaturation: 5 min at 95°C; reaction: 1 min at 95°C, 0.5 min at 58°C, 0.5 min at 72°C for 29 cycles; final extension: 10 min
at 72°C) was carried out using Herculase DNA polymerase enzyme, primers incorporating restriction sites (CN15: 5′-ACG TACGAAATCCATATGCTAGAATTTAAGG-3′ and CN16: 5′-CTAGCAAGCTTTCGATCGAGGACTTC-3′) and template DNA. Amplified DNA fragments were cloned either in pET-28c/pVV2 (at NdeI/HindIII sites for mPDE) or in pMAL-c2X (BamHI/HindIII for PknB or PknL) and maintained as pET-mPDE/pVV2-mPDE/pMAL-PknB/pMAL-PknL in E. coli strain DH5α. PknB (PCR amplified using 5′-AATTAGGATCCCAT ATGACACCCCTCC-3′ and 5′-ACTGCAAGCTTCTA CTCGGCCGAACCT-3′ primers) and PknL (PCR amplified using 5′-AATCTGATCCATATGCTAGAAGCTGGCACG-3′ and 5′-GGATTAACCTTTACGACAGGCAGCGTCACTGGTG-3′ primers) constructs were available in the lab. The pET-mPDE construct was transformed in E. coli strain BL21(DE3) for protein expression and purification. Besides these, pMAL-PknA/-PknA-core/-PknA-K42N, p19kpro-PknA/-PknA-K42N (Rv0015c or mPDE-T309A/D/E variants were also expressed in E. coli strain BL21(DE3) for protein expression and purification. Besides these, pMAL-PknA/-PknA-core/-PknA-K42N, p19kpro-PknA/-PknA-K42N (Rv0015c or mPDE/T309A/D/E) were also expressed.

mPDE-T309A, mPDE-T309D and mPDE-T309E mutants were generated by employing overlap extension PCR method (Ho et al., 1989) using pET-mPDE DNA as the template. Primary PCR was performed using two sets of primers for each mutant (set 1: terminal forward, CN15 and internal reverse, CN54: 5′-TGCCGAGGACGCAGCCACCAT-3′; set 2: internal forward, CN53: 5′-ATGGTGCTGGACGCCAGCAT-3′ and CN57: 5′-ATGGTGCTGGAATCCTCGGCA-3′ and terminal reverse, CN16 for mPDE-T309A; set1: CN15, and CN57: 5′-ATGGTGCTGGACTCTCCATGGCA-3′; set2: CN58: 5′-TGCCGAGGACTCTCCATGGCA-3′ and CN16 for mPDE-T309D; set 1: CN15 and CN59: 5′-ATGGTGCTGGACTCTCCATGGCA-3′; set 2: CN56: 5′-TGCCGAGGACTCTCCATGGCA-3′ and CN16 for mPDE-T309E). For secondary PCR, two independent fragments generated in this way were mixed at equimolar concentrations (1:1), amplified using primers (CN15 and CN16) and cloned in pET-28c or pVV2 vectors.

Recombinant proteins expressed in E. coli were purified following protocol described earlier with slight modifications (Chaba et al., 2002). Briefly, fresh culture (1% inoculum) from overnight cultures) using LB broth with appropriate antibiotics was grown till OD₆₀₀ of 0.6–0.8, cells were induced with 0.4 mM IPTG and further incubated at 37°C for 3 h. This was followed by harvesting and resuspension of cell pellet in lysis buffer (50 mM Tris-HCl, pH 7.5 and 150 mM NaCl) along with protease inhibitors (1 mM phenylmethylsulfonylfluoride, 1 µg/ml pepstatin and 1 µg/ml leupeptin). Cells were lysed by sonication (over ice; 10 s “on” and 10 s “off” cycles for 10 min; 20% amplitude) and centrifuged at 18,000 × g at 4°C for 60 min. The supernatant fraction was loaded onto Ni-NTA affinity column equilibrated with 10 mM imidazole in lysis buffer. Column was washed with 20 mM imidazole in lysis buffer and protein was eluted using 50–200 mM imidazole in lysis buffer.

The mPDE or mPDE-T309A/D/E variants were also expressed in M. smegmatis strain mc²155 following its cloning in pVV2 vector (Dhiman et al., 2004; Thakur and Chakraborti, 2008) at NdeI/HindIII sites (pVV2-mPDE or pVV2-mPDE-T309A/D/E). These constructs were transformed by electroporation in M. smegmatis strain mc²155 following standard procedure (Thakur and Chakraborti, 2008). Cells harboring wild-type or mutants in pVV2 vector constructs were inoculated in Middlebrook 7H9 media enriched with OADC (10%) and Tween-80 in presence of kanamycin (25 µg/ml) and grown at 37°C to 200 rpm for 24 h (OD₆₀₀ of ~0.8). Following their inoculation (1% inoculum) in LB media supplemented with kanamycin (25 µg/ml) at 37°C till OD₆₀₀ of 0.6–0.8, cells were induced with 250 mM imidazole (in lysis buffer), aliquoted in small fractions and stored at −80°C till further use. For preparation of sub-cellular fractions, cellular lysis was accomplished by 2-3 passages through French Press at 1500 psi.
followed by sonication (all parameters same as mentioned above except time for 30 min). The lysate was then centrifuged at 8000×g at 4°C for 5 min for removal of unbroken cells/cellular debris. Centrifugation was repeated at 18,000×g at 4°C for 1 h to separate pellet from supernatant. The obtained pellet after suspending in lysis buffer was further extracted with 2% SDS (65°C for 2 h) and the supernatant fraction obtained after centrifugation (18,000×g at 25°C for 45 min) was used as the cell wall fraction (Wang et al., 2000; Dave et al., 2002; Gibbons et al., 2007). On the other hand, supernatant obtained after cell lysis was ultracentrifuged at 110,000×g at 4°C for 2 h. Orange-brown pellet obtained was used as the membrane fraction after resuspending in lysis buffer (400 µl), while the supernatant was the cytosolic part.

**Kinase Assay**

Transphosphorylation of His6-mPDE by PknA (full length construct containing 431 amino acids or N-terminal 338 amino

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**FIGURE 2** | Phosphorylation of mPDE by PknA. (A) In vitro phosphorylation of mPDE. MBP (0.89 µM)/MBP-PknA Core (0.5 µM)/MBP-PknA-K42N (0.44 µM) were briefly incubated with Ni-NTA purified 1 (1.32 µM) or 10 µg (13.2 µM) mPDE. Samples were processed as described under “Materials and Methods.” Coomassie stained radioactive gel provided in lower panel was used as a loading control. (B) In vitro dephosphorylation of mPDE. Phosphorylated proteins were dephosphorylated following incubation with PPP (10 µg or 5 µM) at 37°C for 75 min. Coomassie stained radioactive gel served as the loading control (lower panel). (C) In vivo phosphorylation of mPDE. BL21(DE3) cell lysates expressing His6-mPDE along with either p19kpko-PknA or empty vector (p19kpko) or kinase dead mutant (p19kpko-PknA-K42N) were probed with anti-PknA antibody (left upper panel) while Ponceau-S stained blot (left lower panel) served as a loading control. Anti-PknA antibody used in this study was generated in our laboratory (Thakur and Chakraborti, 2008). His6-tagged mPDE protein purified from these cells was immunoblotted with anti-phosphothreonine (right upper panel) or anti-His (right lower panel) antibody. Notations used: LMW, low molecular weight marker; PPP, phosphoserine/threonine phosphatase.
acid containing catalytic core; Thakur et al., 2008) or kinase dead variant of PknA (PknA-K42N) or PknB or PknL was assessed utilizing an in vitro kinase assay reported earlier (Chaba et al., 2002; Lakshminarayan et al., 2008). Briefly, PknA (0.28 μM for full length or 0.5 μM for catalytic core) or PknB (0.23 μM) or PknL (0.57 μM) or PknA-K42N (0.44 μM) in 1X kinase buffer (50 mM Tris-Cl, pH 7.5 or 6.8 for PknL/50 mM NaCl containing MnCl₂, 10 mM for PknA/PknL/PknA-K42N or 3 mM for PknB, and 10 mM MgCl₂ only for PknL) and 2 μCi of [γ-32P] ATP was incubated in the presence of mPDE (10 μg or 13.2 μM/reaction) at 25°C (30°C for PknL) for 30 min (total reaction volume = 20 μl). To monitor PPP mediated de-phosphorylation, it was added to the reaction mix (10 μg or 5 μM/reaction) at this step and further incubated for 75 min at 37°C. The reaction was stopped by adding 5X SDS buffer. Samples were resolved in 10% SDS-PAGE; gels were stained with Coomassie Brilliant Blue. Gels were analyzed in a phosphoimaging device (Fuji Film model FLA 9000) and/or exposed to Kodak X-Omat/AR film for autoradiography.

Western Blotting

Purified proteins (usually 1 μg) or whole cell lysates (usually 25 μg) were resolved on 10% SDS-PAGE, transferred (120 V) to nitrocellulose membrane (0.45 μm) using mini-transblot apparatus (Bio-Rad, USA), probed with either anti-His (1:3000) or anti-RpoB (1:3000) or anti-phosphothreonine (1:1000) or anti-PknA (1:1000) or anti-mPDE (1:1000) and anti-mouse (for anti-His and anti-RpoB antibodies) or anti-rabbit IgG (1:5000) as primary and secondary antibodies, respectively. For monitoring in vivo phosphorylation, pET-mPDE or pET-mPDE-T309A was co-transformed with pMAL-PknA or p19kpro-PknA or pMAL-PknB or pMAL-PknL in E. coli BL21(DE3) cells and their expression as a phosphorylated protein was detected employing western blotting using an anti-phosphothreonine antibody as described earlier (Thakur and Chakraborti, 2006). Proteins were finally detected with luminata substrate (Millipore, USA) and developed on X-ray films (Kodak, USA) following manufacturer's recommended protocol.

Phosphodiesterase Enzyme Activity

PDE assays were performed using an assay kit of Enzo® Lifesciences, which employs colorimetric detection of hydrolysis of standard 3’, 5’- cAMP (substrate) by cyclic nucleotide phosphodiesterase. Briefly, the catalytic activity of mPDE (3 μg or 3.96 μM) mediated hydrolysis of 3’, 5’- cAMP (0.05 – 0.8 mM) into 5’AMP was monitored in presence of MnCl₂ (200 μM) at 30°C in a coupled assay following its conversion into adenosine and phosphate catalyzed by 5’ nucleotidase enzyme (0.5 KU/μl). Phosphate such generated was detected by BIOMOL GREEN™ dye at OD of 620 nm at room temperature. Standard curve was prepared by using 5’AMP (0.25 to 3 nmol) in PDE assay buffer that catalyzes its cleavage by 5’ nucleotidase enzyme.

Mass Spectrometry

Samples for mass spectrometry were prepared following the manufacturer's recommended protocol using proteomics grade dimethylated trypsin from porcine pancreas (Sigma, USA). Briefly, Ni-NTA purified proteins in 100 mM ammonium bicarbonate buffer, pH 8.4 was digested in-solution with trypsin at 37°C for 16 h. For cell wall fractions, samples after resolving in SDS-PAGE were eluted in 200 mM ammonium bicarbonate buffer, pH 8.4 containing 40% acetonitrile and then trypsinized (37°C for 16 h) in 40 mM ammonium bicarbonate buffer, pH 8.4 containing 9% acetonitrile. This was followed by running samples on UHPLC system (1290 series, Agilent Technologies) with C8 column coupled to 6550 iFunnel QTOF LC/MS (Agilent Technologies, USA). Electron spray ionization (ESI) was operated in positive ion mode with an MS and MS/MS acquisitions rate of 2 and 4 spectra/second respectively. Peptide sequences were identified utilizing ProteinPilot™ software (http://www.absciex.com/products/software/proteinpilot-software) and Mascot server (http://www.matrixscience.com) as well.

Sequence Analysis and Phylogeny

Reciprocal best blast hits were procured from MycoRRdb database (Midha et al., 2012), multiple sequence alignment was done by Muscle program (Edgar, 2004) and phylogenetic tree was constructed by neighbor joining method (Bootstrap value of 500) using Mega 6.0 software (Tamura et al., 2013).
RESULTS

Phylogenic Placement of mPDE

We initiated our studies to have phylogenetic insights on the placement of mPDE in relation to other bacteria as well as eukaryotes. For this, the gene encoding mPDE (Rv0805) was used as a query to search for the best blast hits in different mycobacterial species. In order to have a wider taxonomic distribution, we extended search in NCBI and Pubmed databases to obtain sequences for few phosphodiesterases present in eukaryotes, gram positive and gram negative bacteria. Multiple sequence alignment using Muscle algorithm followed by phylogenetic tree construction using neighbor-joining method was performed for all the sequences by Mega 6.0 software. Radiation tree shows that mPDE has a unique and distinct taxonomic distribution (Figure 1). Interestingly, orthologs of mPDE are present only in members of M. tuberculosis complex (viz. M. tuberculosis, Mycobacterium africanum, Mycobacterium bovis, Mycobacterium canettii, M. leprae, Mycobacterium ulcerans) along with other slow growing bacilli (Mycobacterium marinum). It is pertinent to mention that phosphodiesterases in Mycobacterium avium and M. smegmatis (MSMEG_2647 and MSMEG_6343), are distantly placed from M. tuberculosis (Figure 1).

Phosphorylation of mPDE by Mycobacterial Eukaryotic-Type Ser/Thr Kinases

A few of the eukaryotic phosphodiesterases (PDE3b, PDE4, PDE5a, and PDE10A etc.) are functionally regulated by PKA/PKB induced phosphorylation (Macphee et al., 1988; Kitamura et al., 1999; Liu and Maurice, 1999; Kotera et al., 2004; Lindh et al., 2007; Bessay et al., 2008; Charych et al., 2010). However, no such phosphorylation mediated regulation of PDE enzyme has ever been reported in any prokaryote. To evaluate this aspect, mPDE protein was purified to substantial homogeneity using Ni-NTA affinity based column chromatography, which on resolving in 10% SDS-PAGE and staining, yielded a major band with molecular mass of 38 ± 6 (Mean ± SD, n = 3) along with slight contamination as discussed previously (Shenoy et al., 2005). We initiated our studies using PknA as a representative of eukaryotic-type Ser/Thr kinases in M. tuberculosis to study phosphorylation of mPDE. Accordingly, in vitro kinase assay was performed wherein mPDE was incubated in presence and absence of MBP-tagged PknA core [amino terminal 338 residues containing catalytic and juxtamembrane domains of the kinase that exhibited enzymatic activity (Thakur et al., 2008)]. As shown in Figure 2A, PknA core (0.5 μM) was capable to trans-phosphorylate mPDE with

![Figure 4](image.png)

**Figure 4** mPDE is phosphorylated by Clade I mycobacterial Ser/Thr kinases. (A) In vitro phosphorylation of Ni-NTA purified mPDE with different mycobacterial kinases belonging to Clade I. Kinase assay utilizing [γ-32P] labeled ATP was performed by incubating MBP-tagged PknA (0.28 μM) or PknB (0.23 μM) or PknL (0.57 μM) with His-tagged mPDE purified from E. coli strain BL21(DE3) or α-casein (4 μM, internal control) using protocol mentioned in the text. Reactions were stopped by adding 5X SDS dye followed by heating at 90°C for 5 min. Samples were resolved in 10% SDS-PAGE gel and signals were visualized through a phospho-imaging device. Coomassie stained gels were used as loading control. (B) In vivo phosphorylation of mPDE with clade I kinases. Recombinant His-tagged mPDE protein was purified using Ni-NTA affinity chromatography of lysates from BL21(DE3) cells expressing either PknA or PknB or PknL in pMAL-c2 vector. Phosphorylation status of the protein(s) was determined in western blot by probing with anti-phosphothreonine antibody (upper panel). Immunoblot (lower panel) using anti-His antibody as the probe served as a loading control.
an increase in phospho-signal between 1 (1.32 μM) and 10 μg (13.2 μM) of protein (lanes 5 and 6). However, phospho-
signal was lost significantly due to deficit kinase-substrate
interactions when an inactive mPDE (denatured by boiling at
95°C for 5 min) was used in the assay (lane 7). Similarly, no
signal was obtained when it was incubated with kinase dead
mutant MBP-PknA-K42N (lane 8). Furthermore, incubation of
phosphorylated protein (obtained after kinase assay) with only
Ser/Thr phosphatase PPP, at 37°C for 75 min led to loss of
phospho-signal of mPDE (Figure 2B; compare lanes 3 and 4).
Notably, Coomassie stained radioactive gels were used as loading
controls for the experiments (Figures 2A,B, lower panels).

To reveal the existence of phosphorylation of substrate protein
(mPDE) in vivo, mPDE was co-expressed with full length
PknA or kinase-dead variant PknA-K42N in E. coli BL21(DE3)
cells utilizing two incompatible vector systems (pET-28c and
constitutive expression vector p19kpro bearing same origin of
replication). Expression of PknA or PknA-K42N (along with
mPDE) was confirmed in E. coli cell lysates (5 μg protein/lane) by
western blotting using anti-PknA antibody (observed molecular
mass = ~45 kDa, lanes 2 and 4, Figure 2C upper left panel
and also see Figure S1). Ponceau-S staining of same blot served
as a loading control where mPDE bands in all lanes were
stained (Figure 2C, lower left panel and also see Figure S1). Despite loading same amount of total protein in each lane (5 μg),
variation in expression levels of mPDE was observed in Ponceau-
S stained blot, which was very likely the result of differential
selection pressure of two different antibiotics (kanamycin for

![Figure 5](https://example.com/figure5.png)

**FIGURE 5** | Identification of Thr-309 as a phosphorylating residue. (A) Identification of phosphorylated residue by LC-MS/MS. Phosphorylated His<sub>6</sub>-mPDE (obtained when co-expressed with pMAL-PknA in BL21(DE3) cells) along with its unphosphorylated counterpart (8 μg each) were hydrolyzed in ammonium
bicarbonate buffer at 37°C for 16h by in-solution trypsin digestion method. Samples were then used for LC-MS/MS analysis. Phospho-site identification was done using protein pilot. Fragmentation spectrum is shown for the modified peptide bearing threonine-309 residue phosphorylated by PknA. Asterisk (*) indicates modified
Thr residue. (B) Thr-309 is predominantly phosphorylated by PknA in vivo. mPDE-T309A generated by site directed mutagenesis was tested for any alteration in
phosphorylation profile compared to the wild-type in western blotting probing with anti-PknA antibody (upper panel). The same blot following stripping
was probed with anti-His antibody served as loading control (lower panel). mFtsZ and PknA were used as internal controls for the experiment. (C) Quantitative analysis
to determine the arbitrary mean phosphorylation intensities of mPDE-P and mPDE-T309A-P from western blots were calculated using ImageJ (Abramoff et al., 2004;
Schneider et al., 2012) and plotted utilizing Microcal Origin 5.0 software. Western blot shown is a representation of four independent experiments from three different
preparations. Notations used: mPDE-P, phosphorylated mPDE; mPDE-T309A-P, phosphorylated mPDE-T309A.
lane 1 while kanamycin plus hygromycin for lanes 2, 3, and 4 of Figure 2C) and consistent with the previous reports (Yang et al., 2001; Thakur and Chakraborti, 2006; Thakur et al., 2008; Dar and Chakraborti, 2010). Moreover, since the translation machinery of the cell was also engaged in simultaneous expression of two different proteins (mPDE and PknA/PknA-K42N; Figure 2C, lanes 2 and 4), their levels of expression were hard to control. Ni-NTA purification of lysates, followed by immunoblotting revealed that co-expression of mPDE with PknA, but not with the kinase-dead variant PknA-K42N, was a phospho-protein recognized by anti-phosphothreonine antibody (Figure 2C, upper right panel lane 2). The presence of mPDE was also confirmed in western blotting of samples probed with anti-His antibody (Figure 2C, lower right panel).

We further expressed mPDE as His-tagged protein in M. smegmatis, which has endogenous sensor kinases but no orthologs of mPDE (Etienne et al., 2005; Narayan et al., 2007; Podobnik et al., 2009). In fact, M. smegmatis is often used as a model in carrying out genetic studies with M. tuberculosis (Thakur and Chakraborti, 2008; Baronian et al., 2015; Zhou et al., 2015). Western blotting using anti-phosphothreonine antibody recognized His-tagged mPDE protein that phosphorylated in cellular extracts (10 μg/lane, Figure 3, lanes 2 and 3) and in purified form (2.5 μg, Figure 3, lane 4). Authenticity of mPDE protein was confirmed utilizing polyclonal anti-mPDE antibody (Figure 3, middle panel). The Ni-NTA purified mPDE protein exhibited a dose-dependent increase in band intensity (inset, Figure 3). Thus, our results strongly indicated mPDE as a substrate of eukaryotic-type Ser/Thr protein kinases.

To elucidate, if transphosphorylation of mPDE is PknA specific, we assessed the ability of other kinases belonging to the same clade (PknB and PknL) for such activities utilizing in vitro kinase assay. For this, we used full length PknA and a basic protein like α-casein as internal controls. As shown in Figure 4A, both PknB and PknL, like PknA were able to transfer [γ-32P] to either α-casein or mPDE. However, magnitude of transphosphorylation ability of different kinases, as has been reflected in phosphorylated band intensity of the substrates was found to be varied (Figure 4A, compare lanes 2, 5, and 8 for α-casein or 3, 6, and 9 for mPDE). Surprisingly, despite using same amount of kinase (PknA or PknB or PknL) in transphosphorylation assay, the addition of mPDE in reaction mixture always resulted in increased band intensity for phospho-PknB compared to that of the PknB alone (Figure 4A, compare upper and lower panels, lanes 4 and 6). Of note, autorad/ phosphoimaging scans with kinases especially in fusion with MBP often exhibited bands at around ~45 kDa besides specific signals and this was consistent with previous reports (Thakur and Chakraborti, 2006; Zhou et al., 2015). These bands could either be phosphorylated protein without fusion tag or degradation product(s) or non-specific entities within the reaction mix. We further validated this observation by immunoblotting of Ni-NTA purified mPDE from E. coli BL21(DE3) cell lysates co-expressed with either PknB or PknL using anti-phosphothreonine antibody. As observed with PknA, mPDE protein was recognized by anti-phosphothreonine antibody in all cases (Figure 4B, upper panel; see also Figure 2C, right panel). Ponceau-S stained blot served as a loading control for the same (see Figure S2) and authenticity of the protein was confirmed by probing with anti-His antibody (Figure 4B, lower panel). Interestingly, despite the loading of same amount of mPDE, the phospho-mPDE as reflected in band intensity varied with the kinase used (Figure 4B, compare upper and lower panels). Thus, these findings argued in favor of in vitro phosphorylating ability of mPDE by mycobacterial eukaryotic-type Ser/Thr kinases belonging to Clade I.

It was further intriguing to identify the residues of mPDE phosphorylated by PknA. For this purpose, lysate from E. coli BL21(DE3) cells co-expressing His6-mPDE as well as MBP-PknA was purified through Ni-NTA column, dialyzed, trypsin-digested and finally used for LC-MS/MS analysis along with unphosphorylated protein. Protein pilot and MASCOT search led to the identification of Thr-309 of mPDE as the phosphorylated residue with 99% confidence (see Table S1). To confirm this observation, we generated a mutant mPDE-T309A by site directed mutagenesis that replaced threonine with alanine. Immunoblot analysis with anti-phosphothreonine antibody showed that the mutant protein (mPDE-T309A) was less phosphorylated than the wild-type (Figure 5B, compare lanes 2 and 4). His6-mFtsZ, which is also a substrate of PknA was used here as a control. Authenticity of different proteins used in this experiment was confirmed by western blotting with anti-His antibody. Quantitation of phosphorylation intensity of bands from western blots recognized by anti-phosphothreonine antibody through ImageJ analysis (in arbitrary units; n = 4) revealed ~8-fold decrease in transphosphorylation of mPDE-T309A compared to that of the wild-type mPDE (Figure 5C).

**Phosphorylation of mPDE is Associated with Its Cell Wall Localization**

The crystal structure of mPDE revealed that the active site comprising of metal and cAMP-binding sites are confined within N-terminal 265 amino acid residues (Shenoy et al., 2007; Podobnik et al., 2009). The C-terminal 20 amino acids, on the other hand, is disordered as no electron density was observed for this region (Podobnik et al., 2009; Matange et al., 2014). Available reports indicated that the C-terminal region has patches, which support protein localization to cell wall (Podobnik et al., 2009; Matange et al., 2014). Thus, the presence of phosphorylating Thr-309 residue (Figure 6A) fascinated us in evaluating the role of phosphorylation on the protein. Kinetic analysis of enzyme activities of mPDE and mPDE-T309A proteins exhibited a typical Michaelis-Menten curve (Figure 6B) with comparable catalytic (Km and Kcat) parameters (Inset, Figure 6B).

We further expressed both His6-mPDE and His6-mPDE-T309A in M. smegmatis, prepared sub-cellular fractions (cytosol, membrane, and cell wall) and compared protein profiles in western blotting using anti-His or anti-mPDE antibody. Although probing the protein with anti-His antibody yielded non-specific bands, we observed band of expected molecular mass of mPDE at membrane and cell wall fractions (Figure 7A top and bottom panels, lanes 2 and 3). Such non-specific band(s) is not unusual considering the fact that the cellular
fractions might possess several proteins bearing stretch of histidine residues, which would have been recognized by anti-His antibody. The intensity of the band, however, at the expected molecular size of mPDE in western blotting was more in cell wall compared to that of the membrane fraction (Figure 7A, compare lanes 2 and 3 of upper and lower panels). Comparative analysis on the localization of mPDE-T309A vs. mPDE revealed that there was no significant alteration in the distribution pattern of the protein in membrane, but its intensity was significantly lower in cell wall fraction (Figure 7A, see lanes 2 and 3 of top and bottom panels). Considering the unaltered intensity of the non-specific band at ∼70 kDa, variation seen in Figure 7A is independent of expression artifact or loading defect in SDS-PAGE gel. Subsequent mass spectrometry experimental results indicated that the ∼70 kDa band seem to arise from a mixture of proteins (GroEL/GntR family transcription regulator/phosphoglycerate mutase/glyceraldehyde-3-phosphate dehydrogenase etc.) but not mPDE (please see Table S2). In fact, these proteins were classically considered to be restricted to cell interiors but have later also shown to be cell surface/cell wall associated [please see http://tuberculist.epfl.ch/ and references (Mawuenyega et al., 2005; He and De Buck, 2010; Boradia et al., 2014)]. However, intensity of this band varied from preparation to preparation despite loading of the same amount of protein. The identity of mPDE in M. smegmatis cell wall was also confirmed by mass spectrometry (inset, Figure 7A; also see Table S3). To validate our findings, we monitored the localization of mPDE/mPDE-T309A in sub-cellular fractions using anti-mPDE antibody following their overexpression in M. smegmatis. Our findings clearly demonstrated that mutation of phosphorylating residue (Thr-309) to alanine affected the localization of protein to cell wall (Figure 7B, compare lanes 3 and 6 of upper panel), whereas the presence of other non-specific bands obtained in Figure 7A were not seen with mPDE specific antibody. Western blotting using anti-RpoB antibody [predominantly a cytosolic marker, see reference (Chao et al., 2013)] was performed to rule out the possibility of cross-contamination between cellular fractions prepared (Figure 7C, upper panel).

Available literature suggested that phosphorylation renders negative charge to a protein, which in turn influences its functionality (Wagner et al., 2004). To have an insight on this aspect, we compared protein localization profiles of wild-type, mPDE-T309A mutant, and phosphomimics (mPDE-T309D and mPDE-T309E) in cell wall fractions following their expression in M. smegmatis (Figure 8A, upper panel). Western blotting using anti-mPDE antibody revealed that cell wall localization of the protein was affected only in mPDE-T309A while both the phosphomimics behaved like wild-type mPDE (Figure 8B). Thus, our results established that eukaryotic-type Ser/Thr kinase mediated phosphorylation of mPDE at C-terminal Thr-309 residue portrayed negative charge to the protein, which in turn endorsed its cell wall localization.

Thr-309 is Conserved in M. tuberculosis Complex

We further considered reciprocal best blast hits for mPDE using MycoRRdb database (Midha et al., 2012) and protein sequences for those whose functional annotation has been done were procured from NCBI (ncbi.nlm.nih.gov). Multiple sequence alignment revealed a high sequence identity between the mPDE orthologs in different mycobacterial species. Interestingly, Thr-309 residue of mPDE is conserved in all the orthologs (Figure 9; PDE sequences from M. smegmatis and M. avium were used as outgroups), illustrating its probable evolutionary significance.

DISCUSSION

Signal transduction in mycobacteria involves highly efficient post translational modification machinery and secondary messengers. Cyclic AMP is one such secondary messenger, which is produced in an exceptionally high amount compared to other bacteria (Shenoy and Visweswariah, 2006). However, only single phosphorydiesterase (mPDE) is identified and characterized, which regulates mycobacterial cAMP levels within cells (Shenoy et al., 2005). Interestingly, its overexpression led to reduction in 3′, 5′-cAMP level in E. coli or M. smegmatis by 30%, while it is of 50% in the case of M. tuberculosis indicating a negative feedback control is operational (Agarwal et al., 2009). Thus, existence of an

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![Figure 6](image-url) | **Comparison of catalytic activities of mPDE and mPDE-T309A.** (A) Schematic representation of domain architecture of mPDE. Position of Thr-309 is indicated by asterisk. N and C denote amino- and carboxy-terminal ends of mPDE respectively. (B) Kinetic analysis. H164-mPDE and H164-mPDE-T309A proteins were purified through Ni-NTA columns. Commercially available PDE assay kit was used to determine kinetic parameters of mPDE and mPDE-T309A proteins as mentioned under “Materials and Methods.” Insets show Coomassie stained gel with purified wild-type (mPDE) and mutant (mPDE-T309A) proteins (3 µg/lane) used for determining catalytic parameters (table). Notation used: LMW, low molecular weight marker.
Phosphorylation of Mycobacterial Phosphodiesterase

FIGURE 7 | mPDE phosphorylation affects its cell wall localization. Phosphorylation influences sub-cellular localization of mPDE. pVV2-mPDE or pVV2-mPDE-T309A were expressed in M. smegmatis. Following sub-cellular fractionation, samples (10 µg protein/slot) were resolved in 10% SDS-PAGE and subjected to immunoblotting. (A) Western blotting with anti-His antibody. Upper and lower panels represent M. smegmatis sub-cellular fractions expressing wild-type and mPDE-T309A proteins probed with anti-His antibody. Ponceau-S staining of this blot served as a loading control (see Figure S3). Inset represents the identity of mPDE protein in M. smegmatis cell wall fraction through in-gel trypsin digestion followed by LC-MS/MS and analyzed by protein pilot software. (B) Western blotting with anti-mPDE antibody to confirm sub-cellular localization of wild-type and mPDE-T309A proteins following expression in M. smegmatis. Reproducibility of western blot results was verified at least three times from two different protein preparations. Numbers indicate molecular weight markers in kDa. (C) Sub-cellular fractions did not show any cross contamination. Upper panel indicates western blotting of cellular fractions of M. smegmatis expressing mPDE/mPDE-T309A utilizing anti-RpoB antibody to probe localization of cytosolic marker RpoB in cells. Numbers indicate pre-stained molecular weight markers in kDa. Lower panels of (B,C) depict Ponceau-S-stained blots as loading controls for the experiments. Notations used: WCL, whole cell lysate; Msm, M. smegmatis; PM, pre-stained marker.

Equilibrium between the levels of cellular cAMP synthesized and degraded or exported out of the cells has already been postulated (Bai et al., 2011). Apart from cAMP hydrolysis, mPDE also plays a key role in maintaining cell wall permeability (Podobnik et al., 2009; Matange et al., 2014) and regulating downstream signaling independent of that mediated by cAMP receptor protein (Matange et al., 2013). Similarly, a well-orchestrated post translational machinery involving cascade of events through phosphorylation and dephosphorylation is mediated by 11 eukaryotic-type Ser/Thr kinases and a cognate phosphatase in M. tuberculosis which have been identified to regulate plethora of cellular functions (Molle et al., 2006; Molle and Kremer, 2010; Pereira et al., 2011). Accordingly, in the current study we assessed the ability of mycobacterial kinases belonging to clade I (PknA/PknB/PknL) to transphosphorylate mPDE. Since, mPDE is phosphorylated by all these kinases with varying degree, our results shown in Figures 2, 4, offer first clue for it being a probable substrate of mycobacterial eukaryotic-type Ser/Thr kinases. Furthermore, mPDE is not a basic protein (calculated pI = 6.1) so that it would be non-specifically phosphorylated.
by these kinases. At this juncture, it needs to be emphasized here that recent genome-wide phosphoproteome mapping of *M. tuberculosis* did not identify mPDE as a phosphorylated protein (Prisic et al., 2010; Prisic and Husson, 2014). However, this is not unlikely considering the fact that experimental conditions might influence the results of mass spectrometric data.

Like eukaryotes, where phosphorylation is one of the important mechanisms to regulate phosphodiesterase activity as well as its functionality (Omori and Kotera, 2007), such a control has not yet been reported in prokaryotes. In this scenario, our results argue that phosphorylation mediated regulation is not confined to eukaryotes only. Interestingly, single threonine residue (Thr-309) at extreme C-terminus of the protein was predominantly phosphorylated with a significant number of modified b and y ions generated in our mass spectrometric study (Figure 5A). Moreover, when this residue was mutated to alanine and co-expressed along with PknA in *E. coli* BL21(DE3) cells, mPDE-T309A was rarely recognized by anti-phosphothreonine antibody (Figure 5B). This highlights the importance of Thr-309 as the phosphorylating residue. Conversely, mPDE-T309A protein did not show any significant alteration in its enzymatic activity compared to that of the wild-type mPDE (Figure 6B), which did not seem to be inconsistent considering the mutation at a non-catalytic site. Thus, in the absence of any Ser/Thr phosphatase use of *E. coli* system as a surrogate host for the expression of mycobacterial Ser/Thr kinases aided us in studying occurrence of phosphorylation of mPDE.

We further introduced *M. tuberculosis* phosphodiesterase gene under the control of an inducible promoter in non-pathogenic saprophyte *M. smegmatis*, where intrinsic kinases such as PknA, PknB and PknL are present but there is no close orthologs of mPDE (Etienne et al., 2005; Narayan et al., 2007; Podobnik et al., 2009). Monitoring the phosphorylation status of the mPDE in such a scenario revealed it as a phospho-protein (Figure 3), though the intensity of phospho-signal was quite low compared to the protein purified from *E. coli* BL21(DE3) cells (Figures 2, 4). This might be an effect of phosphatase present in *M. smegmatis* along with multiple eukaryotic-type Ser/Thr kinases. Surprisingly, we noticed an alteration between wild-type and mPDE-T309A while monitoring sub-cellular localization pattern of these proteins (Figures 7A,B, 8B). On the other hand, Thr-309 when substituted with phosphomimics (Asp or Glu) behaved similar to that of the wild-type (Figure 8B). Such an observation illustrated the importance of the negative charge of the protein (rendered by its phosphorylation) in cell wall anchorage. Our findings are consistent with a previous report demonstrating that phosphorylation at Thr-16 of PDE10A in humans is necessary for its localization from membrane to cytosol (Kotera et al., 2004; Charych et al., 2010). It is therefore logical to infer that phosphorylation mediated control of phosphodiesterases as evident in eukaryotes is also a reality in mPDE despite distinction in their phylogenetic placement (Figure 1). Furthermore, alignment of mPDE with other mycobacterial PDE sequences also explicates Thr-309 as a conserved entity in all the orthologs within *M. tuberculosis* complex (Figure 9) suggesting its phosphorylation to be an evolutionarily important phenomenon. Alternatively, the presence of proline or serine at +1 position of Thr-309 of mPDE orthologs (Figure 9) might be crucial to maintain the structural dynamicity, thereby governing phosphorylation (at Thr-309) in these proteins. Unraveling these aspects of mycobacterial signal transduction mechanisms will therefore open up the new horizon towards understanding of *M. tuberculosis* basic biology.

**CONCLUSION**

Our results established that phosphorylation of mycobacterial phosphodiesterase, which is mediated by eukaryotic-type Ser/Thr kinases like PknA, leads to its cell wall localization. Thus,
this finding indeed insinuates the probable functional interplay between two distinct signaling cascades, one catalyzing post translational modification, and the other involving small molecule trafficking. Further studies in this direction would certainly be expedient to develop potential anti-mycobacterial compounds in future.

**AUTHOR CONTRIBUTIONS**

PC and NM conceived the idea; planned experiments, analyzed the results and wrote the manuscript; NM carried out experiments.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2016.00123

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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