Interdomain Interaction Reconstitutes the Functionality of PknA, a Eukaryotic Type Ser/Thr Kinase from Mycobacterium tuberculosis*

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Eukaryotic type Ser/Thr protein kinases have recently been shown to regulate a variety of cellular functions in bacteria. PknA, a transmembrane Ser/Thr protein kinase from Mycobacterium tuberculosis, when constitutively expressed in Escherichia coli resulted in cell elongation and therefore has been thought to be regulating morphological changes associated with cell division. Bioinformatic analysis revealed that PknA has N-terminal catalytic, juxtamembrane, transmembrane, and C-terminal extracellular domains, like known eukaryotic type Ser/Thr protein kinases from other bacteria. To identify the minimum region capable of exhibiting phosphorylation activity of PknA, we created several deletion mutants. Surprisingly, we found that the catalytic domain itself was not sufficient for exhibiting phosphorylation ability of PknA. However, the juxtamembrane region together with the kinase domain was necessary for the enzymatic activity and thus constitutes the catalytic core of PknA. Utilizing this core, we deduce that the autophosphorylation of PknA is an intermolecular event. Interestingly, the core itself was unable to restore the cell elongation phenotype as manifested by the full-length protein in E. coli; however, its co-expression along with the C-terminal region of PknA can associate them in trans to reconstitute a functional protein in vivo. Therefore, these findings argue that the transmembrane and extracellular domains of PknA, although dispensable for phosphorylation activities, are crucial in responding to signals. Thus, our results for the first time establish the significance of different domains in a bacterial eukaryotic type Ser/Thr kinase for reconstitution of its functionality.

Signal transduction in living organisms plays a pivotal role in controlling several aspects of cellular processes such as metabolism, cell growth, cell motility, cell division, and differentiation. These processes need to be tightly regulated to ensure signaling fidelity, and this synchronization in eukaryotes is mediated primarily through phosphorylation of serine, threonine, and/or tyrosine residues catalyzed by protein kinases. Although they exhibit similarities in their catalytic domains, different kinases may contain additional regions, allowing a multitude of mechanisms for their control. The role of these Ser/Thr or Tyr protein kinases in eukaryotic signal transduction is well established and widely documented (1). Considerably less is known about the prevalence and role of these protein kinases in bacteria and Archaea, where phosphorylation events are predominantly materialized by two-component His kinases along with response regulators, which do not share any sequence similarity with Ser/Thr or Tyr kinases (2). In fact, in the advent of genome sequencing, the presence of genes encoding eukaryotic type Ser/Thr protein kinases in diverse bacterial species raises the possibility of their indispensable involvement in complex network of signal transduction cascade. Although the phosphorylation of these bacterial kinases has been intensively established, the experimental proof of their physiological function and regulation is scarce. Thus, a detailed study of eukaryotic type kinases in bacteria is indeed essential to have insights on their contribution to signaling events. In this context, we focused on the dreadful pathogen Mycobacterium tuberculosis, the causative agent of tuberculosis. The medical impact of tuberculosis (3), which infects nearly one-third of the world’s human population causing considerable mortality annually, therefore provides the rationale for investigating these regulatory proteins in the signaling process.

The genome of M. tuberculosis has unveiled the presence of a family of 11 eukaryotic type Ser/Thr kinases (4). All of these kinases, except PknG and PknK, encode predicted receptors with a single transmembrane helix dividing the protein into N-terminal intracellular and C-terminal extracellular domains (5). Unlike most of the eukaryotes, the N terminus contains a kinase domain, which is linked to the transmembrane region through a variable length of juxtamembrane linker (6). The C-terminal domain outside the cell presumably binds signaling ligands and is attached to the transmembrane sequence. The architecture of these eukaryotic type kinases is very similar throughout bacteria and typical of receptor-like kinases in plants (7). Majority of them (PknA, PknB, PknD, PknE, PknF, PknG, PknH PknI, and PknL) have been shown to catalyze autophosphorylation and substrates for few of them have been identified (8–18). Besides, the crystal structure of the catalytic domain of PknB was solved, which provided valuable information regarding the regulatory mechanism of this kinase (19, 20).

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In an earlier study, we reported cloning and characterization of PknA from *M. tuberculosis* and indicated its involvement in regulating morphological changes in the process of cell division (9). Our recent study with this kinase further indicated its association in regulating functionality of PtsZ, the protein involved in the process of cytokinesis (21). However, very little structural information of PknA is available as yet, which could aid in elaborative understanding of this kinase, especially the molecular mechanism underlying the regulation of phosphorylation activity as such toward its functionality. In this study, we report the identification of the catalytic core of PknA, which is capable of autophosphorylation as well as substrate phosphorylation. We further demonstrate that the autophosphorylation is a bimolecular reaction. It occurs in trans and follows the universal activation mechanism like PknB. Interestingly, unlike PknB, the juxtamembrane region is the integral part of the kinase domain in constituting the catalytic core of PknA. However, for reconstituting the functionality of PknA, the catalytic core itself is not sufficient. Furthermore, we unambiguously establish here that the catalytic core together with transmembrane and C-terminal extracellular domains is critical for the PknA function. Thus, we provide here for the first time the experimental evidence toward functional significance of various domains of a mycobacterial eukaryotic type Ser/Thr kinase.

EXPERIMENTAL PROCEDURES

**Materials**—Restriction/modifying enzymes were obtained from New England Biolabs. All other fine chemicals, including cascin, 3,3’,5,5’-tetramethylbenzidine, glutathione, bovine serum albumin (BSA), Bovine serum albumin; GST, glutathione-S-transferase; HRP, horseradish peroxidase; IPTG, isopropyl thio-β-D-galactosidase; LB broth, Luria-Bertani broth; MBP, maltose-binding protein; PBS, phosphate-buffered saline.

**Construction of Deletion and Point Mutants**—Genomic DNA isolated from *M. tuberculosis* strain H37Rv was used for PCR amplification and subsequent cloning of the *pknA* gene (Rv0015c, ~1.3 kb) in different vectors (pUC-PknA, pMAL-PknA, and p19kpro-PknA) as described elsewhere (9). The deletion mutants, like Δ23–33, Δ17–30, and Δ143–200, were created by restriction digestion (Agel, PstI, and AatII) of pMAL-PknA, followed by re-ligating the backbone (see Fig. 1B). Other deletion mutants (Δ269–431, Δ278–431, Δ339–431, Δ1–252, and Δ1–338) were created employing PCR methods using specific primers (CACAGGAATTCCATGAGCCCCCGAGTTGG/TCATTGCGCTATCTCGTATCGG for Δ1–252; GGGAATTCGGCGCTGCTCTGGGCGGCGCTATCTCGTATCG for Δ1–338; all sequences are 5′ to 3′) and pMAL-PknA as the template. PCR amplified fragments (Δ269–431, Δ278–431, Δ339–431, and Δ1–252) were digested with EcoRI/HindIII and subcloned in pMAL-c2X. The deletion mutants Δ1–252 and Δ1–338, following digestion with EcoRI/Sall and EcoRI/HindIII, respectively, were subcloned at the corresponding sites of pGEX-KG. The Δ339–431 mutant was also subcloned in *Mycobacterium-Escherichia coli* shuttle vector (p19kpro-Core) for constitutive expression in *E. coli* following a similar strategy as described elsewhere (9). Besides these, the deletion mutant, Δ1–252, was digested with EcoRI/Small and subcloned in pGEX-KG to obtain pGEX-253/363 (amino acids 253–363 of PknA) construct. The 363/431 (amino acids 363–431 of PknA) mutant, on the other hand, was obtained by digesting the full-length pUC-PknA with Small/BamHI followed by subcloning at corresponding sites of pGEX-KG.

Two point mutants of PknA, T172A/T174A (threonine substituted with alanine at amino acid residues 172 and 174), were generated using Expand long template PCR system (mixture of Pwo and TaqDNA polymerases; Roche Applied Science) following overlap extension method (22). For each mutation, two external (CATATGAGCCCCGAGTTGG/GGCAAGCTTTCATTGCGCTATCTCGTATCGG, all sequences are 5′ to 3′) and two internal primers (AGCGCGCCGTGCGCCAGCCGCCGCTCATCGATGGCTGGTGGTCGACG/TGCGGCGGCGGCGGCGGTGGTGTCGACG, all sequences are 5′ to 3′ and underlined bases indicate mismatch) were used. To create mutations, two sets of primary (pMAL-PknA as the template) and one set of secondary (mixture of primary reaction products as the template) PCRs were carried out. PCR products containing desired mutations were digested with XhoI/XmaI and incorporated in the corresponding sites of pMAL-PknA. The kinase-dead mutant of PknA, p19kpro-K42N, described earlier (9), was also used in this study. The same mutation was also introduced in Δ339–431 mutant to obtain p19kpro-K42N-Core. Mutations were confirmed by sequencing using an automated DNA sequencer. All constructs and the wild type were transformed in *E. coli* strain DH5α to build up the DNA for further processing.

**Expression of Recombinant Proteins**—Cells harboring MBP-PknA or different MBP fusion constructs were grown in LB broth at 37 °C and induced with 0.3 mM IPTG at an *A* 

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resolved in 8–10% SDS-PAGE and transferred at 250 mA for 45 min. The reaction was stopped by adding SDS sample buffer (30 mM Tris-Cl, pH 7.5, 50 mM NaCl, 10 mM MnCl₂), and the reaction was initiated by adding 2 µCi of [γ-³²P]ATP. Following incubation at room temperature for 20 min, the reaction was stopped by adding SDS sample buffer (30 mM Tris-Cl, pH 6.8, 5% glycerol, 2.5% β-mercaptoethanol, 1% SDS, and 0.01% bromphenol blue). Samples were boiled for 5 min and resolved on 8–12.5% SDS-PAGE. Gels were stained with Coomassie Brilliant Blue, dried in a gel dryer (Bio-Rad) at 70 °C for 2 h, and analyzed in a phosphorimaging device (Molecular Imager FX, Bio-Rad) and also exposed to x-ray films (Eastman Kodak Co.).

**Kinase Assay**—The ability of autophosphorylation and substrate phosphorylation of PknA or its mutants as purified fusion proteins was determined in an *in vitro* kinase assay. Aliquots (usually 1 µg/20 µl reaction volume) of fusion protein were mixed with 1× kinase buffer (50 mM Tris-Cl, pH 7.5, 50 mM NaCl, 10 mM MnCl₂), and the reaction was initiated by adding 2 µCi of [γ-³²P]ATP. Following incubation at room temperature for 20 min, the reaction was stopped by adding SDS sample buffer (30 mM Tris-Cl, pH 6.8, 5% glycerol, 2.5% β-mercaptoethanol, 1% SDS, and 0.01% bromphenol blue). Samples were boiled for 5 min and resolved on 8–12.5% SDS-PAGE. Gels were stained with Coomassie Brilliant Blue, dried in a gel dryer (Bio-Rad) at 70 °C for 2 h, and analyzed in a phosphorimaging device (Molecular Imager FX, Bio-Rad) and also exposed to x-ray films (Eastman Kodak Co.).

**Co-transformation**—*E. coli* (strain DH5α) cells were co-transformed with two incompatible plasmids, pMAL/pGEX-KG and p19Kpro, harboring different constructs. The presence of different antibiotic selections (ampicillin in pMAL/pGEX and hygromycin in p19Kpro) facilitated the co-expression of foreign proteins in *E. coli* using these incompatible plasmids (21, 23). The *E. coli* cells harboring pMAL or pGEX with gene(s) of interest were grown in the presence of ampicillin (100 µg/ml) and made competent using standard methods (24). The cells were then transformed with p19Kpro containing the desired gene(s) and plated on LB agar with both ampicillin (75 µg/ml) and hygromycin (200 µg/ml). Clones obtained were cultured in LB broth in the presence of both the antibiotics and induced with 0.2 mM IPTG (37 °C/3 h). Cells were further processed for microscopy or Western blotting.

**Western Blotting**—Purified fusion proteins (~800 ng) or samples obtained in co-transformation experiments were resolved in 8–10% SDS-PAGE and transferred at 250 mA for 45 min to nitrocellulose membrane (0.45 µm) in a mini-transblot apparatus (Bio-Rad) using Tris-glycine-SDS buffer (48 mM Tris, 39 mM glycine, 0.037% SDS, and 20% methanol, pH ~ 8.3). Primary antibodies (anti-MBP from New England Biolabs and anti-Thr(P) from Cell Signaling Technology) used for different studies, different GST constructs (GST-(253/363), GST-(363/431), and GST-(Δ1–252); 1 µg/well) were coated in microtiter plates for 3 h at 37 °C. Following washing with PBS containing 0.5% Tween (PBS-T), the wells were blocked with 1% BSA in PBS-T for 1 h at 37 °C. After extensive washing with PBS-T, the wells were incubated (1 h at 37 °C) with different concentrations of biotinylated MBP-Core. The interaction was finally monitored by addition of streptavidin-HRP (1 µg/ml for 30 min at 37 °C), and enzyme activity in each well was detected using 3,3’,5,5’-tetramethylbenzidine as the substrate. Binding of biotinylated core with immobilized Δ1–252 was assessed in the presence of different concentrations of nonbiotinylated Δ1–252 to determine the specificity of the interaction. In each case, for controls instead of fusion proteins an equivalent amount of BSA was adsorbed to the wells of microtiter plates.

**Bioinformatic Analysis**—The multiple sequence alignment of the protein sequences retrieved from the mail server at the National Institutes of Health was carried out using the ClustalX1.81 program (26). Secondary structure of the protein was predicted using PSIPRED server (27), and for tertiary structure prediction an automated protein homology-modeling server “Swiss Model” was used (28). Using LSQMAN (Least Square Manipulation; see Ref. 29), the predicted structure of PknA was superimposed with that of the PknB structure, which has been complexed with nucleotide triphosphate analog. For 260 Cα atoms the root mean square deviation was 0.47 Å. The superimposed structure was generated using PyMOL (30).

**Microscopy**—*E. coli* (strain DH5α) cells transformed/co-transformed with different plasmid constructs (p19kpro-Core, pMAL-253/363, pMAL-363/431, and pMAL-Δ1–252) were reinculated at an initial A₆₀₀ of 0.05 and grown further for 12 h. Because there was leaky expression of the MBP fusion proteins (pMAL-253/363, pMAL-363/431, and pMAL-Δ1–252), the cultures were not induced with IPTG. Cells were harvested by centrifugation and subjected to fixation with ethanol after washing with PBS. The fixed cells were collected by centrifugation and resuspended in PBS. The cells were spread on poly-L-lysine-coated glass slides and examined under microscope (Carl Zeiss) following staining with 1% methylene blue solution.

**RESULTS**

**Kinase Domain Together with the Juxtamembrane Region Constitutes the Catalytic Core of PknA**—Analysis of the nucleotide-derived amino acid sequence of PknA indicated the presence of a highly conserved catalytic domain, followed immediately by a juxtamembrane region rich in alanine and proline. The juxtamembrane region leads to a hydrophobic stretch of 23 amino acids constituting the putative transmembrane and the
mutants were purified as MBP fusion proteins, and expression of whether the catalytic domain alone of PknA could perform a homology with regions of these two kinases was considerably less (PknB). However, the homology between the juxtamembrane regions of PknA (using the “Swiss-Model”) predicted tertiary structure of PknA (20). As can be seen in Fig. 2B, the superimposed structure of PknA with PknB as template, it was found that only the ∆339–431 protein among all mutants and MBPβ-galactosidase control (Fig. 1C, lower panel). Thus our results argue that the catalytic domain along with the juxtamembrane region (residues 1–338) are required for PknA autophosphorylation, and hereafter we designate the mutant as the “core” of PknA.

The boundary of the catalytic domain was identified on the basis of primary sequence alignment. However, on careful analysis of the predicted secondary structure and tertiary structures of PknA, using PknB as template, it was found that the ∆269–431 deletion construct missed more than half of the α1 helix, which has been found to be involved in a four-helix bundle in PknB (20). As can be seen in Fig. 2A, the superimposed structure of PknA with PknB highlights the presence of a helix (α1) toward the end of the catalytic domain. The boundary of the helix was further marked by analyzing the predicted secondary structure of PknA (Fig. 2A, inset), and it extended to VRAG (ending at residue 277) corresponding to VHNG (ending at residue 279) in PknB (Fig. 1A). Therefore, to analyze the role of this region in the activity of PknA, another deletion mutant, ∆278–431 (retaining the helix), was constructed. The mutant was tested for its autophosphorylation ability by in vitro kinase assay as well as immunoblotting with anti-Thr(P) antibody. As shown in Fig. 2B, no incorporation of γ32P occurred even with 10 μg of the ∆278–431 mutant; however, 2.5 μg of the ∆339–431 mutant (core) exhibited phosphorylation (compare lanes 2, 4, and 6). This observation was further supported by immunoblotting with anti-Thr(P) antibody, which did not recognize the ∆278–431 mutant (Fig. 2C, lane 2). These observa-
Phosphorylating itself in a concentration-dependent manner, phosphorylation assays were carried out. The core was capable of phosphorylation of the core with 5 μg of heat-inactivated protein (lane 1) in the presence of 10 mM Mn²⁺. B, effect of divalent cations. Autophosphorylation of core (5 μg of protein/reaction) was carried out in the presence of 10 mM of Zn²⁺ (lane 1), Ni²⁺ (lane 2), Co²⁺ (lane 3), Ca²⁺ (lane 4), Mn²⁺ (lane 5), Mg²⁺ (lane 6), or absence (lane 7) of ions and boiled protein (lane 8) in the presence of 10 mM Mn²⁺. C, effect of different inhibitors. Lane 1, boiled protein; lane 2, no inhibitor; lane 3, 2.5 mM ammonium molybdate; lane 4, 5 mM ammonium molybdate; lane 5, 2.5 mM sodium tungstate; lane 6, 5 mM sodium tungstate; lane 7, 2.5 mM sodium vanadate; lane 8, 5 mM sodium vanadate.

these results therefore strongly suggest that the autophosphorylation behavior of the core mimics the MBP-PknA (wild type) protein.

Thr-172 and Thr-174 in the Activation Loop of PknA Are the Phosphorylating Residues—Protein kinases exhibit a multitude of mechanisms for their regulation. The best understood aspect of regulation reconciled in recent years is phosphorylation on a residue(s) located in a particular segment in the center of the kinase domain, which is termed the activation segment or T-loop (32). The activation loop in several kinases has been found to be highly disordered in the crystal structure and is capable of undergoing large conformational changes when the kinase switches between active and inactive states (33). The crystal structure of PknB has highlighted the importance of the activation loop in supporting the universal activation mechanism of the kinase (18). The loop thus identified consists of two threonines actively participating in the activation of the protein (34). We therefore examined if PknA undergoes activa-

A.  
B.  
C.  

FIGURE 2. Model of PknA catalytic domain superimposed on PknB. A, amino acid residues 1–273 of PknA modeled using Swiss-Prot were superimposed on the structure of PknB (residues 1–279). Inset, secondary structure prediction of the last helix α-helix (a) of the catalytic domain of PknA and PknB highlighting the boundary of the helix. B, γ³₂P labeling of deletion mutants of the catalytic domain of PknA. A representative experiment showing Coomassie-stained gel (upper panel) and the autoradiogram (lower panel) of MBP-β-gal (lane 1, 5 μg), MBP-(Δ278–431) (lane 2, 2.5 μg, and lane 6, 10 μg), MBP-(Δ269–431) (lane 3, 2.5 μg, and lane 5, 10 μg) and MBP-Core (lane 4, 2.5 μg) following incubation with [γ³₂P]ATP as described under “Experimental Procedures” under kinase assay. C, Western blotting of MBP-Core (lane 1), MBP-(Δ278–431) (lane 2), MBP-(Δ269–431) (lane 3), and MBP-β-gal (lane 4) using anti-MBP-PknA (upper panel) and anti-pThr (lower panel) antibodies. In each case 1 μg of protein was loaded. Lane numbers are shown at the top and bottom. Position of molecular mass markers is indicated.

FIGURE 3. Autophosphorylation of PknA core. Autophosphorylation of the PknA core was monitored by incubating purified protein with [γ³₂P]ATP followed by separation of the reaction products by SDS-PAGE. The labeled proteins were visualized in a phosphorimaging device or by autoradiography of the dried gel (see “Experimental Procedures”). A, autophosphorylation with 0 (lane 2), 1 (lane 3), 2.5 (lane 4), 5 (lane 5), and 10 μg (lane 6) of purified MBP-Coe along with 5 μg of heat-inactivated protein (lane 1) in the presence of 10 mM Mn²⁺. B, effect of divalent cations. Autophosphorylation of core (5 μg of protein/reaction) was carried out in the presence of 10 mM of Zn²⁺ (lane 1), Ni²⁺ (lane 2), Co²⁺ (lane 3), Ca²⁺ (lane 4), Mn²⁺ (lane 5), Mg²⁺ (lane 6), or absence (lane 7) of ions and boiled protein (lane 8) in the presence of 10 mM Mn²⁺. C, effect of different inhibitors. Lane 1, boiled protein; lane 2, no inhibitor; lane 3, 2.5 mM ammonium molybdate; lane 4, 5 mM ammonium molybdate; lane 5, 2.5 mM sodium tungstate; lane 6, 5 mM sodium tungstate; lane 7, 2.5 mM sodium vanadate; lane 8, 5 mM sodium vanadate.
tion through the same mechanism. Notably, on aligning the sequence of PknA with that of PknB, two threonines corresponding to the phosphotheonines in the activation loop of PknB were located (Fig. 1A). The modeled three-dimensional structure of PknA also showed the presence of disordered activation loop (data not shown). Interestingly, comparable threonines have also been found to exist in other mycobacterial Ser/Thr kinases, like PknD, PknE, and PknF (30).

To analyze the role of these two mapped theonines of the putative activation loop (DFGIKAVDAAPVTQDATMVMGTAQYIPE) of PknA or its core (Δ339–431) on kinase activity, they were mutated one at a time to alanine (T172A and T174A) (Fig. 5A). These single mutants were purified as MBP fusion proteins, and their autophosphorylating abilities were monitored. Both the mutations affected the autophosphorylating ability of PknA (Fig. 4A). However, the effect of T172A was predominant over the T174A, as evident by the difference in signal intensity (Fig. 4A, compare lanes 2, 4, and 6 or 3, 5, and 7). Both the mutations, on the other hand, affected the autophosphorylating ability of the core confirming the results obtained with the full-length protein (Fig. 4B). Interestingly, the profound effect of both the mutations on the core, compared with that of the PknA, presumably point toward difference in the stability between the proteins. In PknB, replacement of both the threonines to alanine equally affected the autophosphorylating ability of the protein (33). Thus, our results establish the direct regulatory role of threonines of the activation loop in autophosphorylation of PknA and favor the universal activation mechanism of this kinase.

PknA Core Exhibits Autophosphorylation in Trans—We have reported earlier that PknA is an active eukaryotic type Ser/Thr protein kinase, and it predominantly phosphorylates at threonine residues (9). However, the biochemical mechanism underlying phosphorylation is poorly understood. Available literature indicates that autophosphorylation of protein kinases occur either through an intramolecular (cis) or intermolecular (trans) association (35). To resolve the underlying mechanism of autophosphorylation of PknA, we tested the ability of the protein to phosphorylate its inactive version. For this purpose, full-length protein (MBP-PknA) was mixed with increasing concentrations (2–8-fold excess) of a kinase-inactive version of PknA core carrying altered Mg\(^{2+}\)-ATP orienting lysine to asparagine (MBP-Core-K42N), and the samples were subjected to kinase assay. As shown in Fig. 5A, the K42N mutant, which is unable to autophosphorylate itself (lane 5), undergoes phosphorylation on incubation with wild type protein (lanes 2–4). Interestingly, a decrease in the phosphate content of wild type protein on incubation with increasing concentrations of the K42N indicates that the mutant was capable of suppressing the intrinsic autophosphorylation ability of the PknA in a dose-dependent manner. However, no such compromise in the PknA phosphorylation ability could be observed when the wild type protein was incubated with an excess of the MBP-βgal (Fig. 5A, lane 6), which further confirmed the authenticity of this experiment.

To substantiate the intermolecular mechanism of autophosphorylation of this kinase, we carried out in vivo co-expression experiments in E. coli by transforming PknA or K42N (in p19kpro vector) along with the core (in pMAL-c2X vector). Because PknA is predominantly phosphorylated at threonine residues (9), we observed that an anti-Thr(P) antibody recognizes PknA or its core (Fig. 5B, left upper panel, lane 1) (31) but not its kinase-inactive variant K42N (Fig. 5B, left upper panel, lane 3). Loading of both the proteins was confirmed by the anti-MBP-PknA antibody in the same blot following its strip-
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![Image of gel with bands labeled 1-5](image_url)

**FIGURE 6.** Substrate phosphorylation of PknA core. Casein or GST-mFtsZ was incubated (25 °C for 20 min) with MBP-Core in kinase buffer along with [γ-32P]ATP. The reaction was stopped by adding 5× SDS buffer, and samples were resolved in 10% SDS-PAGE and processed for autoradiography as described under "Experimental Procedures." Lane 1, MBP-Core (5 μg); lane 2, casein (10 μg); lane 3, casein (10 μg) with MBP-Core; lane 4, mFtsZ (2 μg); lane 5, mFtsZ (2 μg) with MBP-Core. The position of molecular mass markers (in kDa) is indicated. The lane numbers are shown at the bottom. Arrowheads represent casein or mFtsZ.

Thr protein kinases interact with each other (36). Although this aspect has not yet been elucidated in any prokaryotes, being a sensor kinase interaction between domains of PknA is expected. We therefore focused our attention to evaluate such association between the core and the C-terminal domains (transmembrane and extracellular regions) of PknA. For this purpose, although core (Δ339–431) was tagged with MBP (MBP-Core), other domains (juxta-transmembrane, 253/363; extracellular, 363/431; and juxta-transmembrane-extracellular, Δ1–252) were expressed as GST fusion proteins. In vitro interactions between them were examined in pulldown assays. The GST-tagged proteins (GST-(253/363), GST-(363/431), and GST-(Δ1–252)) were incubated (one at a time) with MBP-Core and passed through amylose resin. As shown in Fig. 7A, Western blotting of the samples eluted from the column with anti-GST antibody highlighted GST-(253/363) as well as GST-(Δ1–252) (upper panel) and anti-MBP antibody recognized the MBP-Core (lower panel). Surprisingly, anti-GST antibody recognized an ~31-kDa band in addition to GST-(Δ1–252) (~53 kDa), which seems to be a cleaved product of the fusion protein (Fig. 7A, lane 2). However, no band corresponding to the GST-(Δ1–252) could be detected on incubating the protein with amylose resin in the absence of the MBP-Core, indicating the specificity of the interaction (Fig. 7A, upper panel, compare lanes 1 and 2). GST-(363/431), on the other hand, showed negligible interaction with the MBP-Core protein (Fig. 7A, upper panel, lane 3). The results of the in vitro interaction studies were further confirmed in solid phase binding assays (see "Experimental Procedures"). As shown in Fig. 7B, following affinity purification, binding of biotinylated MBP-Core and GST-(253/363) or GST-(Δ1–252) exhibited a saturation kinetics (half-maximal binding = 4 ng/ml; dissociation constant = 0.08 ± 0.01 nM). On the other hand, GST-(363/431) exhibited hardly any binding. The binding of GST-(253/363) or GST-(Δ1–252) could not be observed when BSA replaced the MBP-Core. Because both GST-(253/363) and GST-(Δ1–252) indicate the interactions of PknA core with its transmembrane domain, we further monitored the ability of unlabeled GST-(Δ1–252) to inhibit its binding of biotinylated MBP-Core (Fig. 6C). The 50% inhibition of binding was achieved at ~2 μg/ml of GST-(Δ1–252), although use of MBP-βgal (pMAL-c2 vector as negative control) had no significant effect (Fig. 7C). Although the involvement of overlapping 87 amino acids (residues 253–339 spanning the juxtamembrane region) among the constructs (core, MBP-(Δ339–431) and GST-(253/363) or GST-(Δ1–252)) used in this study could not be ruled out, all the lines of evidence strongly argue in favor of interaction between core and the transmembrane domain of PknA.

The role of different domains of PknA toward its functionality was further analyzed. For this purpose, a heterologous expression system was utilized. Earlier studies have already established that constitutive expression of PknA results in elongation of E. coli cells (9). We therefore utilized this distinguishing property of the full-length protein (p19kpro-PknA) to elucidate the interaction between different domains of PknA. Because the core emulates the catalytic activities of the wild type protein, we examined the effect of its expression on the morphology of the E. coli cells. Surprisingly, we found that con-
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Constitutive expression of p19kpro-Core did not alter the cell phenotype (Fig. 8, compare a and b). This observation together with the identification of possible in vitro interaction between the core and the C-terminal region, especially the transmembrane domain, led us to speculate their self-association for the functionality of the protein. Despite their in vitro interaction (see Fig. 7, A and B), co-transformation of the p19kpro-Core and pMAL-253/363 in E. coli did not restore the morphological phenotype (Fig. 8d). On the other hand, elongated morphology of the cells could be restored on co-expression of p19kpro-Core and pMAL-/H9004-1–252 constructs (Fig. 8f). Furthermore, p19kpro-Core when transformed with pMAL-363/431 did not affect the cell shape (Fig. 8e), suggesting the importance of both transmembrane and extracellular domains in exhibiting functional reconstitution of PknA. The altered phenotype of the cells thus supported our hypothesis that the domains associate in trans to reconstitute a functional protein in vivo. However, there could still be the possibility that the genetic interaction or reconstitution was mediated by the overlapping cytoplasmic portion of the kinase, i.e. juxtamembrane region. To test this possibility, nonoverlapping construct pGEX-/H9004-1–338 (transmembrane-extracellular domains) was co-transformed with the p19kpro-Core. Interestingly, the phenotype was restored as shown in Fig. 8f, inset. We therefore concluded that the transmembrane domain could be involved in mediating the interaction; however, signaling by complete molecule cannot take place in the absence of the sensory cues from the extracellular domain.

DISCUSSION

Protein kinases play a cardinal role in phosphorylation of proteins and in the process regulate a variety of crucial activities in prokaryotes as well as in eukaryotes. In recent years, genes for eukaryotic type phospho-signaling system were identified in bacteria, and several studies are now focused on unraveling their physiological roles (37). In this scenario, we concentrated on PknA, one of the 11 such Ser/Thr protein kinases present in the genome of the dreadful pathogen M. tuberculosis. PknA has been implicated in playing a role in regulating morphological changes associated with cell division (9, 21). Like any other bacterial eukaryotic type Ser/Thr protein kinases, although the

FIGURE 7. Interaction of PknA core and other domains. A, pulldown assay was carried out by incubating purified GST fusion proteins (GST-(253/363), GST-(363/431), and GST-(Δ1-252)) with purified MBP-Core and amylose resin in binding buffer with 10 mM glutathione. The proteins were extracted from the beads using sample buffer as described under “Experimental Procedures” and analyzed by immunoblotting with anti-glutathione S-transferase (anti-GST; upper panel) and anti-MBP (lower panel) antibodies. Lane 1, GST-(Δ1-252) incubated with amylose resin in the absence of MBP-Core; lane 2, GST-(253/363) with MBP-Core; lane 3, GST-(363/431) with MBP-Core; lane 4, GST-(Δ1-252) with MBP-Core. B, binding of GST fused domains with MBP-Core. Different deletion constructs of PknA (GST-(253/363), GST-(363/431), GST-(Δ1-252); 1 μg/well) were coated on microtiter plates, washed, and incubated in the presence of the indicated concentrations of biotinylated MBP-Core (bCore). Following addition of streptavidin-HRP, enzyme activity of each well was monitored to quantitate binding. The relative enzyme activity was calculated after subtracting the blank (well coated with 1 μg of BSA) and expressed as mean ± S.D. from three independent experiments. C, competition of MBP-Core binding. The binding of biotinylated core (25 ng/well) with immobilized GST-(Δ1-252) was determined. The HRP activity in the absence of any competitor was taken as 100%, and the results (mean ± S.D., n = 3) are expressed as percentage of biotinylated core bound to GST-(Δ1-252). Arrowheads represent GST-(Δ1-252) or GST-(253/363).
Self-association of PknA Reconstitutes Its Functionality

presence of catalytic, juxtamembrane, transmembrane, and extracellular domains has been revealed in PknA through bioinformatic analysis, how the domains collaborate with each other toward its functionality is not yet known. We therefore concentrated on the structure-function analysis of this kinase, especially concentrating on the identification of region responsible for the catalytic activity, its mechanism of activation, and finally to evaluate interaction between different domains for the functionality of PknA.

Sequence alignment, as well as the modeled three-dimensional structure of PknA, has indicated its high order of homology with the catalytic domains of PknB (Figs. 1A and 2). However, in the juxtamembrane regions of the two kinases, the similarity is considerably less (Fig. 1A). Full-length PknA (431 amino acid residues) has been shown to exhibit autophosphorylation as well as substrate phosphorylation abilities (4). The truncation of the protein to cytosolic region harboring catalytic and juxtamembrane domains did not hamper such activities and behaved similarly in respect to the effect of divalent cations (Fig. 3B) or inhibitors (Figs. 1C, 3, and 6). However, unlike PknB and PknF (30), further trimming the protein to the catalytic domain alone completely abolished the enzymatic activity of PknA (Figs. 1C and 2B). Strikingly, the homology shared by the juxtamembrane region rich in alanine and proline with that of the Ala/Ser/Thr/Pro/Gly region of Pkn2, the Ser/Thr protein kinase from Myxococcus xanthus (38), and its requirement for the catalytic activity, corroborate well with our phylogenetic prediction earlier (9). Thus, from this study it is apparent that the catalytic domain along with the juxtamembrane region are the minimum spans required for autophosphorylation as well as substrate phosphorylation abilities of PknA and together constitute the "catalytic core." Although such an indispensability of the juxtamembrane region so far has not been reported for any mycobacterial Ser/Thr kinase, its involvement in activation of several eukaryotic kinases is well known. For instance, in receptor tyrosine kinases and transforming growth factor-β receptor Ser/Thr kinase, the juxtamembrane region serves as a key auto-inhibitory element regulating kinase activity (39). In human epidermal growth factor receptor, the juxtamembrane region has been shown to be indispensable for allosteric kinase activation and productive monomer interactions within a dimer (40). In this scenario, our observation may point toward a dimerization interface in the juxtamembrane region of PknA, distinct from that already predicted in its catalytic domain (15), imparting the kinase activity. Furthermore, it can also be hypothesized that the juxtamembrane region constituting part of the core could be involved in providing structural integrity or stability to the kinase. Nonetheless, the indispensability of the juxtamembrane region seems to be a distinctive feature of PknA, which led us to elucidate the regulation of catalytic activity as well as its functionality.

It is well known that most of the kinases control phosphorylation status through the activation loop. Sequence comparison has identified certain structural determinants explaining the reason behind their activation through phosphorylation. It has been suggested that kinases containing a catalytic aspartate preceded by an arginine residue, termed as RD kinases, are known to be activated by phosphorylation in the activation segment (33). Analysis of the PknA sequence revealed the presence of RD motif, suggesting the existence of a parallel mechanism of activation as has been observed with PknB (20). Interestingly, we also identified the activation segment carrying the two predicted phosphorylating threonines on the basis of sequence alignment with PknB (Fig. 1A). We therefore mutated these two threonines and as expected could identify Thr-172 and Thr-174 as the phosphorylating residues (Fig. 4). Thus, it could be predicted that activation of PknA occurs by a universal activation mechanism like other eukaryotic kinases (32).

Autophosphorylation of Ser/Thr kinases in eukaryotes is known to occur either by the cis or trans mechanism (41, 42). In fact, very little information is available regarding the mechanism of autophosphorylation of bacterial Ser/Thr kinases. To have an insight on this aspect, in vitro phosphorylation assays were performed utilizing the kinase-active and -inactive versions of full-length and PknA core, respectively. It is apparent from the experiment depicted in Fig. 5 that the autophosphorylation occurs in trans. Furthermore, in vivo phosphorylation of the full-length kinase-inactive mutant (p19Kpro-K42N) by the co-expressed kinase-active core (pMAL-Core) emphasized the prevalence of intermolecular autophosphorylation of PknA.
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(Fig. 5B). Thus, our results ostensibly established that the PknA exhibits bimolecular autophosphorylation like the majority of histidine kinases in bacteria and most of the eukaryotic receptor kinases (43–45).

Previously we have reported that PknA, when expressed in E. coli under a constitutive promoter, resulted in cell elongation (9). Surprisingly, the core, which is enzymatically active, upon transformation in E. coli was unable to bring about phenotypic changes (Fig. 8b) as has been observed with the full-length protein. It was therefore hypothesized that there could be a prevalence of interaction/association between different domains of PknA, which may lead to phenotypic changes. To test this possibility, the core was co-transformed with either transmembrane or extracellular domain. Both the core and transmembrane domains were found to interact in vitro (Fig. 7); however, co-transformation of the two did not restore the phenotype, indicating the insufficiency of the transmembrane domain alone in reconstitution of the functionality of PknA (Fig. 8d).

On the other hand, co-transformation of the extracellular domain along with core neither showed any interaction (Fig. 7) nor any phenotypic change (Fig. 8e). Interestingly, co-expression of core with the transmembrane and regulatory regions exhibited interaction (Fig. 7) as well as restored the elongation phenotype (Fig. 8f, inset), suggesting association of these domains in trans and reconstitution of a fully active protein in this heterologous setting. All these observations therefore indicated the indispensability of the domains for reconstitution of the functionality of PknA and therefore the prevalence of a distinct mechanism for this kinase. Such self-association and functional reconstitution have also been reported for a Ser/Thr protein kinase fused (Fu), a component of Hedgehog signaling complex in Drosophila (36). Alternatively, the interdomain interaction/association might have restored the ability of PknA to recognize regulatory cues that lead to its transformation from inactive to active state through conformational changes as has been reported for transforming growth factor-β receptor Ser/Thr kinase (39).

Finally, our results for the first time signify the importance of each domain of a bacterial eukaryotic type Ser/Thr kinase toward reconstitution of its functionality. All these lines of evidence underscore the molecular mechanism of regulation of PknA and hence may provide an insight into the mechanism of signal transduction in mycobacteria.

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Interdomain Interaction Reconstitutes the Functionality of PknA, a Eukaryotic Type Ser/Thr Kinase from Mycobacterium tuberculosis
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