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Crystal Structure of PhnF, a GntR-Family Transcriptional Regulator of Phosphate Transport in Mycobacterium smegmatis

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Department of Microbiology and Immunology, University of Otago, Dunedin, New Zealand; Laboratory of Structural Biologyb and Maurice Wilkins Centre for Molecular Biodiscoveryc; School of Biological Sciences, University of Auckland, Auckland, New Zealand; Department Biology I, Mikrobiologie, Ludwig-Maximilians-Universität München, Planegg-Martinsried, Germany.

Bacterial uptake of phosphate is usually accomplished via high-affinity transporters that are commonly regulated by two-component systems, which are activated when the concentration of phosphate is low. Mycobacterium smegmatis possesses two such transporters, the widely distributed PstSCAB system and PhnDCE, a transporter that in other bacteria mediates the uptake of alternative phosphorus sources. We previously reported that the transcriptional regulator PhnF controls the production of the Phn system, acting as a repressor under high-phosphate conditions. Here we show that the phnDCE genes are common among environmental mycobacteria, where they are often associated with phnF-like genes. In contrast, pathogenic mycobacteria were not found to encode Phn-like systems but instead were found to possess multiple copies of the pst genes. A detailed biochemical analysis of PhnF binding to its identified binding sites in the phnD-phnF intergenic region of M. smegmatis has allowed us to propose a quantitative model for repressor binding, which shows that a PhnF dimer binds independently to each site. We present the crystal structure of M. smegmatis PhnF at 1.8-Å resolution, showing a homodimer with a helix-turn-helix N-terminal domain and a C-terminal domain with a UbiC transcription regulator-associated fold. The C-terminal domain crystallized with a bound sulfate ion instead of the so far unidentified physiological ligand, allowing the identification of residues involved in effector binding. Comparison of the positioning of the DNA binding domains in PhnF with that in homologous proteins suggests that its DNA binding activity is regulated via a conformational change in the linker region, triggering a movement of the N-terminal domains.

As inorganic phosphorus is an essential and frequently limiting nutrient, the mechanisms of its uptake are important systems for most bacteria. Bacteria possess both high- and low-affinity phosphate transport systems, the former of which are induced under inorganic phosphate (Pi)-limited conditions (1, 2). The high-affinity, ATP-binding cassette (ABC)-type transport system Pst (phosphate-specific transport) consists of four components, PstSCAB, and recognizes free phosphate as its substrate. Pst systems have been well characterized in a number of bacteria, including Escherichia coli (1), Sinorhizobium meliloti (3, 4), and Bacillus subtilis (5, 6). In Mycobacterium tuberculosis, pstS1 and pstS2 have been identified as virulence factors (6, 7). Expression of the pstSCAB operon is generally induced under Pi-limited conditions and is mediated by the two-component regulatory systems PhoBR in Gram-negative microorganisms (8–10), PhoPR in most Gram-positive organisms (11, 12), and SenX3-RegX3 in Mycobacterium smegmatis (13). In E. coli and M. smegmatis, repression of the phosphate starvation response requires the Pst transporter, and mutations in pstS result in constitutive target gene expression (1, 14).

We have previously identified a three-gene operon (phnDCE) in M. smegmatis which encodes a second high-affinity phosphate transport system. Most Phn systems studied to date are responsible for the uptake of alternative phosphorus sources, such as phosphonates (compounds containing a direct carbon-phosphorus bond) or phosphate (15–17). In contrast, the M. smegmatis PhnDCE transporter does not recognize these substrates and is instead specific for Pi (18). Further, we have shown that the transcriptional regulator PhnF controls phnDCE expression by acting as a repressor under phosphate-replete conditions. Induction of phnDCE upon phosphate starvation is further enhanced by the two-component system SenX3-RegX3 (19). The phnF gene is found upstream of and in the opposite orientation to phnDCE, and its expression is autorepressed by PhnF and additionally activated by SenX3-RegX3.

While M. tuberculosis possesses multiple copies of genes encoding Pst systems (20, 21), M. smegmatis contains only a single copy of the pstSCAB genes (4, 22). It therefore appears that the PstDCE transporter, regulated primarily by PhnF, offers M. smegmatis a mechanism that is an alternative to that of multiple Pst systems for the efficient uptake of Pi under growth-limiting conditions. The work presented here aims to improve our understanding of the role of PhnF by structural and biochemical characterization and to establish a framework for its interaction with both its target DNA and its activating ligand.
MATERIALS AND METHODS

Sequence analysis. To identify orthologues of the *phnDCE* and *phnF* genes, all actinobacterial genomes available in October 2010 at the NCBI website (http://www.ncbi.nlm.nih.gov) were searched with the BLASTP program (23) using the sequences of *M. smegmatis* PhnD (MSMEG_0649), PhnC (MSMEG_0647), and PhnE (MSMEG_0646) as queries. All genomes containing hits for all three queries encoded by adjacent genes were then subjected to a further BLASTP search using *M. smegmatis* PhnF (MSMEG_0650) as the query. The results from this analysis were updated in August 2013 using the 154 actinobacterial genomes available in the MicrobesOnline database (24).

Cloning and expression. The region encoding the C-terminal ligand binding domain of PhnF (C-PhnF; residues 76 to 244) from *Mycobacterium smegmatis* and the full-length phnF gene were cloned from genomic DNA. The boundary between the N-terminal DNA binding domain and the C-terminal ligand binding domain (C-PhnF) was located by alignment with the gene sequence of PhnF from *Escherichia coli* (25). C-PhnF and full-length phnF were cloned into the Gateway system using the nested two-stage PCR protocol described previously (26, 27) with the following primers: full-length phnF forward primer 5′-GGCGACGGCGCGGGTGAAGGGGGGGGC-3′, C-PhnF forward primer 5′-GGGCAACGGCGGGGATCGACACAAACCCTGCGATG-3′, and full-length and C-PhnF reverse primer 5′-GGGAAGCTTTCGTGACACACAAAAAGATTGGCGG-3′. Clones were verified by sequencing and transferred into the pDEST17 expression vector to produce a His-tagged protein (C-PhnF) or pDEST566 to produce a His-tagged maltose-binding protein (MBP) fusion protein (full-length PhnF). Expression was carried out in *E. coli* BL21(DE3) cells grown in 500 ml of autoinduction medium ZYM-5052 (28) in baffled 2-liter flasks with shaking at 160 rpm. Cultures were grown to 310 K for 4 h and then at 291 K overnight.

Purification. Cells from the expression culture were harvested by centrifugation at 5,000 × g for 20 min at 277 K. These were resuspended in lysis buffer (20 mM HEPES [pH 7.5], 150 mM NaCl) with 5 or 10 mM imidazole (for full-length PhnF and C-PhnF, respectively) and the addition of 0.5 mM dithiothreitol, 4% [wt/vol] glycerol, 50 μg/ml RNase A, 10 μg/ml DNase I, 2 mM MgCl₂, and Roche Complete mini EDTA-free protease inhibitor tablets. Cells were lysed by cell disruptor at 18 kPa. The lysate was centrifuged at 26,000 × g for 20 min at 277 K, and the soluble fraction was filtered through a 0.45-μm-pore-size filter.

C-PhnF was purified by immobilized metal affinity chromatography (IMAC) using a gravity-flow TALON column (Clontech). The clarified supernatant was passed through the column, and the column was washed with lysis buffer containing 20 mM imidazole. Protein was then eluted with lysis buffer containing 150 mM imidazole. This protein was concentrated by using Superox 75 size exclusion chromatography using 20 mM HEPES (pH 7.5), 150 mM NaCl. This protein was then used for crystallization. Full-length PhnF was purified by IMAC as described above, the His₇₆-MBP fusion was removed by proteolysis with recombinant tobacco etch virus protease, and the sample was reapplied to the TALON column. The flowthrough containing PhnF was collected and further purified by gel filtration as described above. Protein purity was monitored throughout by SDS-PAGE.

Crystallization and data collection. Initial crystallization conditions were obtained using a Honeybee (Cartesian Dispensing Systems) nanoliter robotic system in a sitting-drop format with 100 nl of protein solution mixed with 100 nl well solution in Intellilplates (Art Robbins Instruments) and a set of in-house screens covering 480 conditions (27).

(i) C-PhnF. Diffraction-quality crystals were produced by hanging-drop vapor diffusion, in which 1 or 2 μl of protein (at 2 to 5 mg/ml) was mixed with an equal amount of well solution. Initial crystals were produced using well solution containing 1.0 M ammonium sulfate, 0.1 M lithium sulfate, and 0.1 M Tris HCl (pH 8.5). These crystals were used for microseeding of further droplets in 0.65 M ammonium sulfate, 0.1 M lithium sulfate, and 0.1 M Tris HCl (pH 8.5).

(ii) PhnF. The best crystals were produced by hanging-drop vapor diffusion by mixing 1 μl of protein solution (at 5 mg/ml) with 1 μl of well solution. The well solution contained 0.1 M sodium cacodylate (pH 8.0) and 0.8 M sodium acetate. All crystals were incubated in a solution of the appropriate mother liquor with 25% [vol/vol] glycerol and flash cooled in liquid nitrogen. Crystals were maintained at 100 K during X-ray exposure in a stream of cooled nitrogen gas.

Data collection, structure solution, and refinement. X-ray diffraction data were collected to a resolution of 1.8 Å for both PhnF and C-PhnF using an in-house rotating anode X-ray generator (Rigaku MicroMax-007 HF) with an image plate detector (Mar Scanner 345). All other computing was carried out in the CCP4 suite unless otherwise stated (29). For C-PhnF, the structure was solved by molecular replacement using the BALBES system (30) and the structure with PDB accession number 2P19 as the model. Automated model building was performed with the ARP/wARP package (31), and structure refinement was performed with the REFMAC program (32) interspersed with manual rebuilding using the Coot program (33). The structure of PhnF was solved by molecular replacement using PHASER software (34) and C-PhnF as the search model and was refined as described above. Data collection and refinement statistics can be found in Table 1.

Electrophoretic mobility shift assays (EMSAs). A 377-bp fragment containing the intergenic region between *phnF* and *phnD* was PCR amplified using primers PphnFR and PphnDR (19) and end labeled [γ³²P]ATP using T4 polynucleotide kinase (Roche) according to the manufacturer’s instructions. Target DNA carrying mutations in the PhnF binding sites was created as described previously (19). Binding reactions were carried out in binding buffer (20 mM Tris HCl [pH 8], 50 mM KCl, 1 mM dithiothreitol, 4% [wt/vol] glycerol, 5 μl/m³ salmon sperm DNA), and the reaction mixtures contained ca. 2,000 cpm radiolabeled DNA and various concentrations of PhnF, as indicated below. For competition experiments, 100 ng of unlabeled DNA containing either the wild-type sequence (TGGTATAGACCA for both binding sites) or a sequence with mutations in both PhnF binding sites (TGGTATAGACCA in site F1, TGGTATAGACCA in site F2 [the mutated sites are underlined]) (19) was added to the binding reaction mixture. PhnF was diluted in 20 mM Tris HCl (pH 8), 150 mM NaCl, 50 μl/m³ bovine serum albumin. The reaction mixtures were incubated at room temperature for 20 min, followed by electrophoresis on a 6% (wt/vol) acrylamide (19:1 acrylamide/bisacrylamide) gel in 1 × TBE (89 mM Tris base, 89 mM boric acid, 2 mM EDTA) at 300 V for 20 min. Bands were detected using a phosphorimager.

Mathematical model for PhnF-DNA binding. On the basis of the palindromic recognition sequence of PhnF in the intergenic region of *phnD*-*phnD*, it seemed likely that each of the F1 and F2 sites can be bound by a PhnFᵢ dimer. Assuming that proteins dimerize prior to DNA binding with dimerization dissociation constant *Kₛ*, the free dimer concentration, [PhnFᵢ], depends on the total PhnF concentration, [PhnFₜot], as follows:

\[
[\text{PhnF}_{\text{tot}}] = \frac{[\text{PhnF}_{\text{tot}}]}{2} + \frac{Kₛ - \sqrt{Kₛ^2 + 8Kₛ[\text{PhnF}_{\text{tot}}]}}{8} \tag{1}
\]

To model the DNA binding probabilities shown in Fig. 1D and E, a thermodynamic framework for protein-DNA interactions was applied (35). In DNA fragments containing only site F1 (Fig. 1E), the probability that site F1 bound by PhnFᵢ would be found (pᵢ) reads

\[
pᵢ = \frac{[\text{PhnF}_{\text{tot}}]}{Kᵢ} \left( \frac{[\text{PhnF}_{\text{tot}}]}{Kᵢ} \right) \tag{2}
\]

where *Kᵢ* is the PhnFᵢ-F1 dissociation constant. Accordingly, the probability that site F1 will be found to be unbound (*pᵢ = 1 – pᵢ*) is given by 1 – *pᵢ*. Expansion of the square root in equation 1 shows that only for small ratios of [PhnFₜot]/Kₛ does the dimer concentration depend quadratically on the total PhnF level, such that in this limit the binding prob-
### RESULTS

**Distribution of Phn systems in actinobacteria.** Previous studies of phosphate transport in mycobacteria have shown that *M. tuberculosis* contains several copies of genes encoding a PstSCAB transporter (21), whereas *M. smegmatis* contains one operon each for a PstSCAB transporter and a PhnDCE transporter (18). To gain a better understanding of the distribution of these transport systems among the actinobacteria, we searched the currently available genomes of these bacteria for orthologues of *M. smegmatis* PhnDCE. Our results showed that none of the pathogenic mycobacteria for which genomes are available encode a PhnDCE-like phosphate transport system but instead have several copies of the *Pst* phosphate transport system (Table 2). In contrast, 15 of the analyzed genomes of environmental mycobacteria and related actinobacteria possess a complete putative *phnDCE* operon, with *Mycobacterium vanbaalenii* and *Clavibacter michiganensis* containing two such operons. None of these bacteria contained multiple copies of *pst* genes (Table 2).

The *phnDCE* operon of *M. smegmatis* is associated with an additional gene, *phnF*, which encodes a transcriptional regulator that acts as a repressor of *phnDCE* expression (19). We therefore searched the actinobacterial genomes encoding a PhnDCE transporter for the presence of orthologues of *M. smegmatis* PhnF. Eleven of the transporter operons possessed a *phnF*-like gene in their genomic neighborhood. In the remaining four genomes, a *phnF*-like gene was encoded elsewhere on the chromosome, and in the two species containing two *phnDCE* operons, only one operon was associated with such a gene (Table 2). PhnDCE transporters were originally identified to be uptake systems for phosphonates, an alternative source of phosphorus that can be utilized by many bacteria, especially inhabitants of the soil (36). Interestingly, only two of the transport operons identified by our analysis were associated with the remaining *phn* genes required for the degradation of phosphonates (Table 2). The PhnDCE system of *M. smegmatis* does not recognize phosphonates as a substrate, consistent with the absence of the degradative genes (18), and this may also be the case for the remaining actinobacterial systems identified here.

Our findings indicate that possession of a PhnDCE phosphate uptake system is a common trait in environmental actinobacteria, where such systems may have originated as phosphonate transporters. The pathogenic mycobacteria appear to have lost these systems and have possibly compensated for this loss through duplication of their *pst* genes. On the basis of the conserved genomic arrangement and co-occurrence of *phnDCE* and *phnF* genes, it appears likely that production of PhnDCE transporters is generally regulated by PhnF-like repressors in the actinobacteria.

**PhnF binds to two sites in the phnF-phnD intergenic region.** We have previously proposed that PhnF binds to two sites in the *phnF-phnD* intergenic region in *M. smegmatis*, based on genetic evidence (19). To determine whether purified PhnF is indeed able to bind to two sites in the *phnF-phnD* intergenic region, we investigated its binding properties using competition experiments and footprinting analysis.

### Table 1

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*Of the modeled residues for C-PhnF, 98.6% lie in the preferred region, 1.4% lie in the allowed region, and no residues lie in the disallowed areas. For PhnF, 98.0% of the modeled residues lie in the preferred region, 1.9% lie in the allowed region, and one residue (Gln 92 on chain C) lies just within the disallowed area; on inspection, this residue lies on a loop region where the electron density for the side chain is weak, aside from that for the terminal amide group, which is clear.

* Rmerge = ΣI − ⟨I⟩/2ΣI, where I is the intensity of a reflection and ⟨I⟩ is the average intensity; RMSD, root mean square deviation.

* Values for the highest-resolution shell are shown in parentheses.
to bind to these putative binding sites, we carried out DNA binding studies using electrophoretic mobility shift assays (EMSAs) with a DNA fragment encompassing the intergenic region between phnF and phnD (Fig. 1A). A band shift of this fragment was detected in the presence of PhnF at a concentration of 2 nM or higher, and a supershift occurred at the presence of PhnF at a concentration of 10 nM or higher (Fig. 1B). A complete shift of all DNA was observed from 20 nM PhnF. This is consistent with PhnF binding to two separate sites on the target DNA, where only one site is occupied at low protein concentrations (giving a shift), followed by occupation of both sites simultaneously at higher protein concentrations (resulting in a supershift). Binding of PhnF was specific, as the addition of excess unlabeled target DNA abolished the shift, whereas the addition of unlabeled DNA carrying a single site (site F1) or only the phnD-proximal binding site (site F2). As shown in Fig. 1C, PhnF binds to both of these fragments in a very similar pattern, with a partial shift observed from 5 nM protein and a complete shift observed at 20 nM protein. No supershift occurred, consistent with the presence of only one binding site in the target DNA. In contrast to the more gradual shifting of DNA containing both binding sites, the single-site fragments both showed a steep response between 10 nM and 20 nM protein concentrations, which was also reflected in the sigmoidal curve after quantification of band intensities (Fig. 1E, left).

These results showed that the two sites displayed almost identical binding behavior and, due to the sigmoidal dependency, suggested that binding of PhnF to each individual site is cooperative. In fact, on the basis of the palindromic recognition sequence (TTGTA-N$_4$ACCA) (19), it seemed likely that each site is bound by a dimer of PhnF. In order to examine this, we set up a theoretical model that included protein dimerization prior to binding of DNA. This model indeed produced the same dose-response behavior observed experimentally (Fig. 1E, right). In order to reproduce the sigmoidal shape of the experimental data, the protein dimerization affinity constant was required to be high ($K_D \approx 100$ nM) compared to the total PhnF concentrations used in the band-shift assays. Under these conditions, the PhnF monomer-dimer equilibrium shifted toward the dimer with increasing total PhnF concentration, and together with a high protein dimer-DNA binding affinity ($K_D \approx K_F \approx 1$ nM), this led to an effective Hill coefficient of $n$ nearly equal to 2. Importantly, both the dimerization and the DNA binding affinity constants were chosen from a typical range of in vivo parameters for bacterial transcription factors (37–40), suggesting that the observed sigmoidal binding curves can be explained by dimerization of PhnF alone and do not require further, unknown mechanisms.

Adapting the same model with identical parameters for a DNA fragment containing two binding sites for PhnF could also accurately reproduce the more complex binding behavior observed for the wild-type DNA fragment (Fig. 1D). Here the model predicted that for increasing PhnF concentrations the probability of finding

<table>
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<th>Species</th>
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<th>PhnF orthologue</th>
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* The number of pst5 genes present in the genome is given in parentheses; +, the presence of an additional pseudogene.

* Only locus tags for the first gene (phnD orthologue) are shown for each operon.

* Gene identifiers are given as locus tags.

* PhnF orthologue not directly associated with Phn transporter genes.

* Separated from the phnDCE operon by 10 other phn genes; probably involved in phosphonate metabolism.
either site F1 or site F2 occupied first increases and then decreases again (Fig. 1D, right, dark gray curve). Within the model this decrease is due to occupation of the second site at higher PhnF concentrations, which, accordingly, increases the probability of PhnF binding to F1 and F2 simultaneously (Fig. 1D, right, black curve). Together, these results show that binding of PhnF to each individual binding site is cooperative and that this cooperativity likely arises from dimerization prior to DNA binding. Moreover, the binding of both sites in the intergenic region is consistent with a model without cooperativity between the individual sites, suggesting that PhnF binds sites F1 and F2 independently.

PhnF represses transcription of the phnDCE operon and of its own structural gene under phosphate-replete conditions, and repression is relieved when the cells enter phosphate limitation (19). To test whether phosphate had an effect on DNA binding, the EMSAs were repeated in the presence of different phosphate concentrations. Only very high concentrations (above 10 mM) had any adverse effects on binding (data not shown), but as the internal free phosphate concentration of \( M. \) smegmatis is below 0.5 mM (our unpublished data), this is not likely to be of physiological relevance. Because most GntR-family transcriptional regulators repress their target genes in the absence of their ligand (41–46), it appears likely that PhnF responds to a molecule present in phosphate-starved but not phosphate-replete cells. This is discussed in more detail below.

**Structure of PhnF.** In order to better understand the nature of phosphate regulation by PhnF, we set out to determine its structure. PhnF crystallizes as a homodimer with one dimer in the asymmetric unit (Fig. 2A). PhnF is composed of two domains. The N-terminal domain, comprising residues 10 to 73, consists of a canonical winged helix-turn-helix (wHtH) DNA binding domain with an \( \alpha_1-\beta_1-\alpha_2-\alpha_3-\beta_2-\beta_3 \) topology (Fig. 2B) and is connected to the ligand binding domain by a linker region of 21 amino acids (Fig. 2A, pink residues), 5 of which form the \( B_4 \) strand. The C-terminal domain has a chorismate lyase fold (47), comprising a six-stranded \( \beta \)-sheet at its core, which is extended by strands \( B_11 \) and \( B_4 \) from the second monomer. Dimerization is mediated by two equivalent sets of interactions to give a dimer containing two twisted eight-stranded \( \beta \)-sheets. This mode of dimerization is identical to that observed in the crystal structures of other members of the GntR/HutC family of transcriptional regulators, including YvoA, YurK, and YydK from *Bacillus subtilis* and PhnF from *E. coli* (PDB accession numbers 2WVO, 2IKK, 3BWG, and 2FA1) (48, 49). We also independently crystallized the C-terminal
domain of PhnF (C-PhnF). The structure of this domain alone has a fold and a dimerization interface identical to those seen in the full-length protein.

The program DaliLite (50) reports (as of 20 May 2014) 22 different proteins with Z-scores greater than 10 to be near structural neighbors. The nearest structural neighbor was identified as being the GntR/HutC-family transcriptional regulator YvoA from *B. subtilis*, which is involved in the regulation of the N-acetylglucosamine-degrading pathway (PDB accession number 2WV0, Z-score = 21.5, sequence identity = 31%) (Fig. 3A, green strand) (48). The second-nearest neighbor was identified to be an uncharacterized gene product from *Staphylococcus aureus* (PDB accession number 2OO1) with a Z-score of 20.3 and a sequence identity of 12%. The vast majority of matches are results from structural genomics programs and consist of structures encoding putative GntR/HutC-family regulators. This demonstrates that the fold of PhnF is confined to enzymes of the GntR/HutC family of regulators, of which PhnF is a member; Fig. 3A shows a structural comparison between the C-terminal domains of PhnF and three of its related proteins.

**FIG 2** Key structural features of PhnF from *Mycobacterium smegmatis*. (A) Crystal structure of the PhnF homodimer. One monomer is shown in gray (N-terminal domain) and cyan (C-terminal domain), and the other monomer is shown in red (N-terminal domain) and purple (C-terminal domain). The linker region of both monomers is shown in pink. (B) Topology plot of the PhnF monomer. (C) Ribbon representation of the effector domain showing the residues involved in SO$_4^{2-}$ binding in ball-and-stick representation. (Inset) A detailed view of the binding site. Electron density shown is a $2F_o - F_c$ synthesis contoured at approximately 1σ (0.55 electron/Å$^2$). An alignment of the protein sequences for the C-terminal domains of YvoA and PhnF is shown for comparison. Red, amino acids that form the same contacts with SO$_4^{2-}$ through their side chains in both proteins; blue, amino acids that form equivalent contacts through their main chains; orange, amino acids that form similar contacts with SO$_4^{2-}$ but do not occupy similar positions in the binding site. Alignment was performed with the Clustal Omega program (56).
Regulation of PhnF activity. PhnF is a regulator of both the phnDCE operon and of its own transcription (19). When *M. smegmatis* is grown under phosphate-limiting conditions, the repression of the phnDCE operon is relaxed. As mentioned above, other members of the HutC regulator family bind to their target DNA only in the absence of their respective ligands (41–46). Accordingly, our hypothesis for the mechanism of repression by PhnF is that the protein recognizes a molecule present only under phosphate-starved conditions, which would trigger the derepression of the PhnF target genes. In the absence of this ligand, i.e., under phosphate-replete conditions, PhnF remains bound to DNA and the target genes are repressed.

However, the identity of the ligand to which PhnF responds is unknown. Unlike *E. coli*, *M. smegmatis* is unable to utilize phosphonates or phosphite as a phosphorus source (18). Furthermore, repression of phnF and phnDCE expression by PhnF in *M. smegmatis* is relieved by phosphate starvation alone, regardless of the presence of any alternative phosphorus compounds. It is therefore unlikely that *M. smegmatis* PhnF responds to a phosphonate ligand. A simpler hypothesis—that PhnF binds phosphate directly and that this allows it to bind DNA—is also unlikely to be correct, because DNA binding was not affected by the absence or presence of physiological concentrations of phosphate. It therefore appears likely that *M. smegmatis* PhnF responds to a small molecule that is produced or accumulates in the cell as it enters phosphate starvation. Due to the large-scale metabolic rearrangements of cellular metabolism upon phosphate limitation, including changes in energy metabolism and adaptations that reduce the cell’s phosphorus demand (52, 53), it is difficult to predict candidate molecules. The presence of sulfate in the ligand binding pocket of C-PhnF may suggest that the true ligand is a phosphate-containing species, based on the structural similarity between sulfate and phosphate groups, but we have no experimental evidence to date.

Structural mechanism of DNA binding by PhnF. PhnF crystallized as a homodimer, with the N-terminal DNA binding domains being positioned on opposite sides of the effector domains (Fig. 3B, purple strand). This position means that in their current orientation the two domains would be unable to simultaneously contact the same DNA molecule. As described above, we have shown that each dimer of PhnF binds to one DNA binding site and that PhnF dimerization occurs before binding to the DNA. Thus, the structure observed in the crystal must undergo a rearrangement that brings the two N-terminal DNA binding domains closer together in order to allow PhnF to functionally bind DNA. The structurally related GntR/HutC transcription factor YvoA from *M. smegmatis* has a similar arrangement of its DNA binding domains (Fig. 3B, green strand) (48). In this case, the authors postulated that the structure that they determined is that of the non-DNA binding form of the protein, which then undergoes a shift in N-terminal domain orientation in order to allow the formation of a 1:1 DNA-protein complex. In addition, they observed a sulfonate ion bound to the effector domain, which they suggest mimics the true ligand and so triggers the protein to adopt a non-DNA binding conformation, thus explaining the relative positioning of the N-terminal domains.

The existence of an alternative orientation of the N-terminal DNA binding domains is supported by the report of the

DISCUSSION

The C-terminal domain of PhnF was crystallized in a high concentration of sulfate ions, as was YvoA from *B. subtilis* (48), and a sulfate ion is found in the structures of both these proteins. This ion is bound in the same position in both C-PhnF and YvoA, being contacted by residues from strands β5 and β6, the loop region between helices α7 and α8, and helix α4 (Fig. 2C). This sulfate ion forms hydrogen bonds with the side chains of R129, S162, R98, and S84 and with the main chain peptide NH groups of L163 and Y164. The sulfate-binding site of PhnF shows a great degree of similarity with that of YvoA, with several of the residues being conserved. R129, S162, and Y164 form contacts equivalent to those made by R133, S165, and Y167 in the *B. subtilis* protein, and L163 forms the same contact through the backbone as I166 of YvoA. There are, however, some differences in binding that likely allow the two proteins to have different substrate specificities. Whereas in the *B. subtilis* protein R135 and T90 make side chain hydrogen bonds with the sulfate ion, these interactions are replaced in C-PhnF by hydrogen bonds with R98 and S84. The interaction between the sulfate ion and R135 is also observed in the related trehalase repressor from *B. subtilis* TreR (R128; PDB accession number 2OGG) (51). Although anions commonly bind to the N termini of two or more α-helices, mutations in the residues involved in binding the sulfate ion effectively obliterated effector binding in YvoA (48), suggesting that this is indeed the region of the protein involved in effector binding.

The existence of an alternative orientation of the N-terminal DNA binding domains is supported by the report of the
structure of YydK, which is structurally closely related to PhnF (Z-score = 19.7) (PDB accession number 3BWG). The N-terminal domains in this structure are positioned closely together in such a way that they could make contact with the same DNA binding site (Fig. 3B, blue strand). That the structure of YydK shows the authentic DNA binding orientation of the domains is lent weight by the observation of how the GntR/HutC-family regulator FadR from E. coli binds to its DNA binding site (54). The DNA binding domain dimer of FadR can be overlaid with that of YydK, with a reported C-α root mean square deviation of 5.3 Å, while differences between the single DNA binding domains are much smaller. This confirms that the orientation of the domains is very similar in both instances. The structure of YydK does not contain an anion in the effector domain, unlike the structures of YvoA and the related proteins YurK and TreR (described above), which all contain sulfate ions bound in equivalent positions and show DNA binding domains in the non-DNA binding orientation on opposing faces of their effector domains. Comparison of these structures with the structure of PhnF shows that the N-terminal domains in the latter occupy a position between the positions seen in YvoA and YydK (PDB accession number 3BWG) (48).

Although the structure of full-length PhnF described in this work does not contain a bound anion and would be postulated to be in the DNA binding form, this does not seem to be the case. Nevertheless, a small movement in the linker region between the N- and C-terminal domains would be sufficient to bring the N-terminal domains into the position occupied by those of YydK and thus render them capable of binding DNA. Comparison of this region of the two structures shows that there is a turn in the backbone of YydK at residues 71 to 73, which orients the N-terminal domain to lie above the dimerization interface. In contrast, this turn is absent in PhnF, but the equivalent residues, 73 to 75, form only one hydrogen bond with the rest of the protein, between the backbone of Lys75 and the side chain of Gln92, and this might easily be broken to allow movement of the N-terminal domain about this hinge point. Use of the web server HINGEPROT (55) identified a hinge region for the PhnF dimer at residue 74, providing further support for the possibility of movement in this region of the protein.

**Effector binding site of PhnF.** The identity of the effector molecule for PhnF is currently unknown, but it seems likely that the endogenous ligand is a phosphate-containing molecule, as is postulated to be the case for YvoA. While there are large similarities in the binding sites of M. smegmatis PhnF and B. subtilis YvoA, the differences in side chain orientation are likely to be responsible for the different specificities of the proteins (51). It is possible that any allosteric change that would take place on ligand binding is not observed in PhnF in the case of binding sulfate, because the anion is too small in comparison to the size of the physiological effector molecule to make the contacts necessary to drive a conformational change.

**Conclusions.** The crystal structure of PhnF shows that the protein consists of an N-terminal winged helix-turn-helix DNA binding domain and a C-terminal ligand-binding domain of the Ubic transcription regulator-associated (UTRA) fold, as predicted by sequence alignment (19). The protein forms a homodimer, mainly through interactions between the C-terminal domains. Comparison with structurally related proteins suggests that a conformational change in the linker region upon ligand release is likely to facilitate DNA binding in the absence of the ligand by triggering movement of the N-terminal domains. We have demonstrated that phnDCE genes in environmental actinobacteria are generally associated with a phnF gene and are therefore likely to be regulated by PhnF-mediated repression. In most cases there are no associated phosphate degradation genes, which implies that phosphate specificity in these actinobacteria is likely to be similar to that in M. smegmatis. Moreover, we showed that PhnF binds to two independent binding sites in the phnD-phnF intergenic region and that binding to each site is cooperative and compatible with a thermodynamic model assuming PhnF dimerization prior to DNA binding. The hypothesis that dimerization occurs in the absence of the effector ligand is further supported by the observation of a dimer in the crystal structure where no ligand is present. While the identity of the physiological effector ligand for PhnF remains elusive, it is likely to be a small, phosphate-containing molecule produced in phosphate-starved cells.

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