Crystal Structure of the Bacillus subtilis Phosphodiesterase PhoD Reveals an Iron and Calcium-containing Active Site*

Received for publication, August 15, 2014, and in revised form, September 9, 2014. Published, JBC Papers in Press, September 12, 2014, DOI 10.1074/jbc.M114.604892

Fernanda Rodriguez1,2, James Lillington3,4,5,6, Steven Johnson4,5, Christiane R. Timmel7, Susan M. Lea8,9, and Ben C. Berks1,2

From the 1Department of Biochemistry, University of Oxford, South Parks Road, Oxford OX1 3QU, the 2Sir William Dunn School of Pathology, University of Oxford, South Parks Road, Oxford OX1 3RE, and the 3Inorganic Chemistry Laboratory, University of Oxford, South Parks Road, Oxford OX1 3QR, United Kingdom.

Background: PhoD family enzymes liberate phosphate from organic compounds.
Results: The structure of PhoD reveals an active site with one Fe3+ and two Ca2+ ions.
Conclusion: PhoD represents a new class of phosphatase related to purple acid phosphatases.
Significance: The requirement of PhoD for iron ions may limit bacterial phosphate acquisition in low iron environments.

The PhoD family of extra-cytoplasmic phosphodiesterases are among the most commonly occurring bacterial phosphatases. The exemplars for this family are the PhoD protein of Bacillus subtilis and the phospholipase D of Streptomyces chromofuscus. We present the crystal structure of B. subtilis PhoD. PhoD is most closely related to purple acid phosphatases (PAPs) with both types of enzyme containing a tyrosinate-ligated Fe3+ ion. However, the PhoD active site diverges from that found in PAPs and uses two Ca2+ ions instead of the single extra Fe2+, Mn2+, or Zn2+ ion present in PAPs. The PhoD crystals contain a phosphate molecule that coordinates all three active site metal ions and that is proposed to represent a product complex. A C-terminal helix lies over the active site and controls access to the catalytic center. The structure of PhoD defines a new phosphatase active site architecture based on Fe3+ and Ca2+ ions.

PhoA family enzymes are predominantly phosphomonoesterases. The active site of the prototypical PhoA enzyme of Escherichia coli contains two Zn2+ and one Mg2+ ion, and the catalytic mechanism involves an enzyme-bound phosphoserine intermediate (3). PhoX enzymes are exclusively phosphomonoesterases, and their complex active site cofactor has recently been shown to contain two Fe3+ ions, three Ca2+ ions, and a μ4-bridging oxo group (4).

Enzymes of the PhoD family are the least well characterized of the three major classes of bacterial extra-cytoplasmic alkaline phosphatase. The archetype of the family is Bacillus subtilis PhoD. This enzyme is a phosphodiesterase but also has significant phosphomonoesterase activity (5). Metal ion reconstitution experiments show that PhoD requires Ca2+ for activity (5). The physiological role of PhoD is proposed to be the release of phosphate from cell wall teichoic acids under conditions of phosphate starvation (6). A second biochemically characterized member of the PhoD family is the secreted phospholipase D (PLD)7 from Streptomyces chromofuscus (7). PLD cleaves the phosphodiester bond that links the head group and diacyl glycerol portions of phospholipids. PLD is reported to contain one iron atom together with substoichiometric amounts of manganese. In addition, the enzyme requires Ca2+ ions for significant activity and to bind to phospholipid membranes.

Sequence analysis suggests that PhoD proteins are unrelated to the PhoA or PhoX phosphatase families but instead form a distinct subgroup within a very broad structural superfamily of hydrolytic enzymes possessing binuclear metal centers at the active site (the “metallophosphatase” superfamily in the Conserved Domain Database (8)). Within this structural superfamily it has been argued that the PhoD family has greatest similarity to eukaryotic purple acid phosphatases (PAPs) (9). In PAPs the active site metal pair are a Fe3+ ion and a divalent ion that can be Fe2+, Mn2+, or Zn2+ depending on the enzyme (10, 11). The Fe3+ ion is ligated by a tyrosine ligand resulting in an intense phenolate-to-Fe(III) charge-transfer transition (λmax = 650–700 nm).

* This article was selected as a Paper of the Week.
\[ \text{Fe}^{3+} \]
Structure of the Phosphodiesterase PhoD

510–560 nm; $\epsilon = \sim 3000 – 4000 \text{ M}^{-1} \text{ cm}^{-1}$), which gives PAPs their characteristic purple color. The occurrence of a tyrosinolate-ligated Fe$^{3+}$ ion is highly characteristic for PAPs, with the only other currently identified examples being found in the structurally unrelated diferric DNA ligase from Ferroplasma acidiphilum (12). It is, thus, striking that $S$. chromofuscus PLD exhibits a PAPS-like visible spectrum. Amino acid residues, including a tyrosine, that could correspond to the Fe$^{3+}$ ligands in PLD have been proposed on the basis of sequence conservation and mutagenesis experiments (9).

In an effort to define the active site architecture of the PhoD phosphatase family we have determined the structure of $B$. subtilis PhoD. Unexpectedly, the active site contains three metal ions, these being identified as a Fe$^{3+}$ ion and two Ca$^{2+}$ ions. Our structural data show that the PhoD active site is distinct from that found in PAPs but that these two enzyme families conserve the position and coordination environment of their Fe$^{3+}$ ions. A phosphate molecule bound to the PhoD active site gives insight into the mechanism of catalysis.

EXPERIMENTAL PROCEDURES

Cloning—The sequence coding for the mature domain of PhoD (codons 57–593) was amplified from $B$. subtilis strain 168 chromosomal DNA using the primers 5’-CCAGTGGGTCTC-AGGTGGTGCGCCTAACTAAGC-3’ and 5’-CCGCGTCGACTTATTTCTGATTTTGACG-3’. The resulting ampli-
con was digested with BsaI and Sall and cloned into the same sites in plasmid pCAS97 (13) to produce plasmid pCAS97-mphoD. This plasmid directs the expression of the mature domain of PhoD with a Ulp1-cleavable N-terminal Strep-SUMO tag (where SUMO is small ubiquitin-like modifier). Site-specific mutations were introduced into the phoD coding sequence in plasmid pCAS97-mphoD using the QuikChange™ method (Stratagene).

Protein Purification—$E$. coli strain BL21 (DE3)(14) containing plasmids pCAS97-mphoD and pRARE (Novagen) was grown aerobically at 37 °C in 2×YT medium (Sigma-Aldrich) supplemented with 50 $\mu$g/ml kanamycin. Protein expression was induced at an A$_{600 \text{ nm}}$ of 0.6 with 0.5 mM isopropyl-1-thio-
$\beta$-d-galactopyranoside, and the culture grown was then grown for 15 h at 20 °C.

Cells were harvested by centrifugation and resuspended in resuspension buffer (20 mM Hepes NaOH, pH 7.4, 200 mM NaCl, 1 mM CaCl$_2$, with EDTA-free complete protease inhibitors (Roche Applied Science) and DNase I (Sigma-Aldrich). The cells were broken by three passages through a French press at 8000 p.s.i. The lysate was clarified by successive centrifugation at 10,000 × g for 20 min at 4 °C and at 45,000 × g for 1 h at 4 °C, and then applied to a 5 ml self-packed Strept-Tactin column (IBA, GmbH, Göttingen, Germany). Recombinant PhoD was eluted from the column with resuspension buffer containing 2.5 mM desthiobiotin and then supplemented with 1 mg of His$_6$-Ulp1 protease (13) and dialyzed overnight at 4 °C against resuspension buffer. Cleaved tags and uncleaved mPhoD protein were removed by incubation with Strept-Tactin matrix, and the sample was further purified on a Superdex75 (16/60) prep grade size-exclusion column (GE Healthcare) that had been equilibrated in resuspension buffer. The mPhoD-containing fractions were pooled and concentrated using a 30-kDa cut-off Amicon PLTK Ultracel-PL membrane concentrator (Merck Millipore).

Phosphatase Activity Assays—Alkaline phosphatase activity was assayed spectrophotometrically by measuring the release of p-nitrophenol from p-nitrophenyl phosphate (pNPP) or bis-p-nitrophenyl phosphate (bis-pNPP). Assays were started by the addition of 1 $\mu$g/ml PhoD to 100 mM Tris-HCl, pH 8.0, 2 mM CaCl$_2$, and 1 mM pNPP or bis-pNPP at 25 °C. Reactions were stopped after 5 min by the addition of 1 M final concentration NaOH. p-Nitrophenol production was quantified using $\epsilon_{405 \text{ nm}}$ = 18,000 M$^{-1}$ cm$^{-1}$.

The substrate specificity of PhoD was assessed by measuring phosphate release from test substrates. Each 800-$\mu$l reaction contained 100 mM Tris-HCl, pH 8.0, 2 mM CaCl$_2$, 1 mM test substrate, and 1 $\mu$g of PhoD. After a 5-min incubation at 25 °C the reaction was terminated by the addition of 200 $\mu$l of malachite green reagent (AMS Biotechnology) followed by incubation at room temperature for 30 min. Production of phosphate was obtained from absorbance at 620 nm and quantified using a phosphate standard curve.

Crystallographic Data Collection and Structure Determination—Crystallization trials were performed at 294 K by the sitting drop vapor diffusion method. The best crystals were obtained in 0.2 M NaCl, 0.1 M Na$_2$PO4/KH$_2$PO$_4$, 54% PEG 200, pH 5.6, with protein at 7.8 mg/ml, a total drop volume of 400 nl, and a 50:50 protein-to-mother liquor ratio. The crystals were cryo-protected in mother liquor containing 10% (w/v) ethylene glycol and flash-cooled in liquid nitrogen. X-ray data were collected at beamlines ID23_1 (European Synchrotron Radiation Facility (ESRF), Grenoble, France) and I03 (Diamond Light Source, Oxfordshire, UK). The data were processed in space group I4122 using Process (part of the autoPROC pipeline; Global Phasing Ltd.) (15). The multiple wavelength anomalous diffraction method was used to determine the sites of the three catalytic metals (initially assigned as three iron sites, but later refined as one iron and two calcium sites) using autoSharp (16) and to produce phases that were optimally solvent-flattened to 62% using Solomon (17). The handedness of the sites was determined by the contrast (standard deviation of solvent to standard deviation of protein) falling from 57 to 27% in one hand and from 63 to 39% in the other. Buccaneer (18) built a partial model among a number of other unlikely chains. This model was then manually manipulated in accordance with the kidney bean purple acid phosphatase structure (Protein Data Bank (PDB) 3KBP). Input of this partial model for a second round of Buccaneer building placed 524 residues of a monomer into the map. This was used as the search model for molecular replacement via MOLREP (19), placing the two translational related non-crystallographic symmetry copies at (0.5, 0.5, 0.08) of the unit cell from one another. Iterative refinement and model building were carried out with Refmac (20) and Coot (21). The quality of the model was checked using MolProbity (22), with a final MolProbity score of 1.19 (93th percentile).

The sample for proton-induced x-ray emission (micro-PIXE) analysis was purified in a calcium-free buffer composed of 20 mM Tris-HCl, pH 7.2, 200 mM potassium acetate. Measurements were taken at the Ion Beam Centre (University of Surrey).
EPR Spectroscopy—EPR spectra were collected on an X-band CW EPR spectrometer (Bruker Biospin GmbH) using an X-band Super High Sensitivity Probehead (Bruker) and equipped with a low temperature helium flow cryostat (Oxford Instruments CF935). Data were analyzed using the program EasySpin (23).

RESULTS

Characterization of B. subtilis PhoD—B. subtilis PhoD was heterologously expressed in the cytoplasm of E. coli. Purified recombinant PhoD protein was a monomer in solution as judged by size-exclusion multi-angle laser light scattering (Fig. 1A). The recombinant PhoD exhibited phosphomonoesterase activity with the colorigenic substrate pNPP (black bars) and bis-pNPP (gray bars), respectively. PhoD in 5 mM Tris-HCl, pH 8.0, was treated with 100 μM Na2EGTA for 5 min (bar marked EGTA), and then 2 mM of the chloride salt of the indicated divalent metal cation was added. Error bars indicate the S.D. of three independent experiments. The recombinant enzyme was massively enhanced by Ca2+/H11001 supplementation (Fig. 1B), and so Ca2+/H11001 was subsequently included in all buffers used for the purification and analysis of PhoD. At pH 8.0 and a temperature of 25 °C, the Ca2+/H11001-supplemented recombinant PhoD protein had a \( K_{\text{m}} \) for pNPP of 50 μM and a \( k_{\text{cat}} \) of 1.2 s\(^{-1}\). PhoD was able to release phosphate from compounds containing phosphoester bonds but was not able to cleave phosphorus-nitrogen or phosphorus-carbon bonds (Fig. 2A).

**FIGURE 1.** A, size-exclusion multi-angle laser light scattering analysis of PhoD. The buffer was 20 mM Hepes, pH 7.5, 400 mM NaCl, 2 mM calcium acetate, and the flow rate was 0.4 ml/min. A single species of weight-averaged molecular mass 59,700 Da (± 1%) is observed. B, phosphomonoesterase and phosphodiesterase activities of PhoD measured with pNPP (black bars) and bis-pNPP (gray bars), respectively. PhoD in 5 mM Tris-HCl, pH 8.0, was treated with 100 μM Na2EGTA for 5 min (bar marked EGTA), and then 2 mM of the chloride salt of the indicated divalent metal cation was added. Error bars indicate the S.D. of three independent experiments.

**FIGURE 2.** Characterization of recombinant B. subtilis PhoD. A, phosphate released from different substrates by PhoD. The assays contained 1 μg/ml PhoD and 1 mM of the test substrate in 100 mM Tris, pH 8.0, 2 mM CaCl2. Gly3P is glycerol 3-phosphate; Phoac is phosphonoacetic acid; Glu6P is glucose 6-phosphate; imidodP is imidodiphosphate. Substrates are grouped according to whether the target bond is between phosphorus and oxygen (O–P), phosphorus and carbon (C–P), or phosphorus and nitrogen (N–P). Error bars indicate the S.D. of three independent experiments. B, visible absorption spectrum of 2.3 mg/ml PhoD in 20 mM Hepes NaOH, pH 7.4, 200 mM NaCl, 1 mM CaCl2 in the absence (black) or presence (gray) of 10 mM Na2EGTA. C, EPR spectra of PhoD highlighting the g = 4.3 peak. Spectra are shown for native PhoD, PhoD with 5 mM sodium dithionite, PhoD with 2 mM hydrogen peroxide, and PhoD with 50 mM EDTA. The inset shows a magnified part of the spectrum around g = 9.6. The PhoD concentration was 10 mg/ml in 20 mM Hepes, pH 7.5, 400 mM NaCl, 2 mM calcium acetate with 20% (v/v) glycerol added. A control experiment showed that the glycerol added to the samples did not change the spectral line shape (data not shown). EPR operating conditions: microwave power, 2 milliwatts; microwave frequency, 9.425 GHz; modulation, 5.0 G at 100 kHz; temperature, 10 K. Spectra are buffer-subtracted, with negative controls taken for sodium dithionite, hydrogen peroxide, and EDTA solutions.
Structure of the Phosphodiesterase PhoD

PhoD purified in the presence of Ca\(^{2+}\) was purple in color with a broad absorbance band in the visible spectrum, indicating the presence of a prosthetic group (Fig. 2B). The addition of the Ca\(^{2+}\)-selective chelator EGTA caused a shift in the visible absorbance maximum from around 540 nm to \(\sim560\) nm (Fig. 2B), suggesting that Ca\(^{2+}\) is associated with the prosthetic group. Proton-induced x-ray emission spectroscopy of PhoD purified in the absence of added Ca\(^{2+}\) ions detected iron and calcium but no cobalt, magnesium, manganese, nickel, or zinc. The presence of iron in PhoD was confirmed by EPR spectroscopy, which detected the \(g=4.3\) and weak \(g=9.6\) peaks characteristic of the middle and lower Kramer’s doublets of an isolated high spin rhombic Fe\(^{3+}\) ion (Fig. 2C). No further signals were elicited by oxidation with hydrogen peroxide. All EPR signals disappeared on reduction with sodium dithionite, as expected of reduction of a monomeric Fe\(^{3+}\) ion to the EPR silent Fe\(^{2+}\) state (Fig. 2C). The Fe\(^{3+}\) signal was unperturbed by the addition of 50 mM EDTA (Fig. 2C), showing that the iron in PhoD is too tightly bound to be removed by chelator treatment. The presence of iron in PhoD has not previously been reported. However, iron is found in the homologous \emph{S. chromofuscus} PLD enzyme (7).

\textbf{Structure of PhoD}—PhoD was crystallized at pH 5.6 in the presence of phosphate. The structure of PhoD was determined to a resolution of 2 Å via a multi-wavelength anomalous diffraction experiment using the iron cofactor as the source of the anomalous signal. Final \(R/R_{\text{free}}\) values at this resolution were 17.8/20.2%. Data reduction, phasing, and crystallographic refinement statistics are shown in Table 1. Every residue in the PhoD mature domain was observed in the electron density.

The core of the PhoD structure is a sandwich of two long \(\beta\)-sheets flanked by \(\alpha\)-helices of varying lengths (Figs. 3A and 4). Three active site metal ions are coordinated by residues on loops on one face of the \(\beta\)-sandwich and are covered by a long C-terminal helix. With the knowledge that PhoD contains Fe\(^{3+}\) and Ca\(^{2+}\) ions (above) anomalous Fourier maps at three wavelengths allowed the assignment of the three active site metal ions as one Fe\(^{3+}\) ion and two Ca\(^{2+}\) ions, which we label Ca\(_A\) and Ca\(_B\) (Figs. 3, B and C, and 5A). Substitution of individual amino acids involved in metal ion coordination at each of the three metal sites either abolished (C180A, D265A, N271A, N272A, and D436A variants) or drastically reduced (D207A and D266A variants) PhoD activity with pNPP as a substrate (Fig. 5B), showing that all three metal ions are required for the phosphomonoesterase activity of PhoD.

PhoD was crystallized in the presence of phosphate, and the structure contains a phosphate molecule bound at the active site (Fig. 3B). Phosphate is a known competitive inhibitor of PhoD (5).

\textbf{PhoD Has Structural Similarity to Purple Acid Phosphatases}—A search of the PDB protein structure database using the DALI server (24) revealed that PhoD has highest structural similarity to PAPs. The closest overall structural match was to red kidney bean PAP (PDB 4DHL) (25) with a DALI \(Z\)-score of 28.6 and a main chain root mean square deviation relative to PhoD of 2.47 Å. The similarity between the two protein families is illustrated in Fig. 3, A and B, where we have chosen to compare PhoD with sweet potato PAP (PDB 1XZW) (26) due to the close structural homology in the way the two proteins bind phosphate (discussed below).

At the Fe\(^{3+}\) site the sole difference in amino acid ligand sets between PhoD and the PAPs is a cysteine-aspartate substitution in PhoD (Cys-180). Cysteine is an unusual ligand to a Fe\(^{3+}\) ion. Indeed, to the best of our knowledge, the structurally unrelated alkaline phosphatase PhoX (4) is the only other protein using a thiolate group to coordinate a redox-inactive Fe\(^{3+}\) ion. The visible absorbance bands of PhoD are likely to arise predominantly from the tyrosine ligation of the Fe\(^{1+}\) ion, as in the PAPs (27). However, additional thiolate-to-Fe\(^{3+}\) charge transfer bands from the cysteine ligand will also contribute to the visible spectrum (28).

The Ca\(_A\) site in PhoD corresponds to the position of the second metal site in the PAPs. Nevertheless, there is little similarity between the metal ion ligands found at this site in the two types of protein (Fig. 3B), consistent with the different coordination preferences of Ca\(^{2+}\) and the divalent metal ions found in the PAPs. Specifically, the Ca\(_A\) site has irregular 8-fold coordination with all amino acid ligation being provide by Asp residues.

The Ca\(_B\) site in PhoD has no analogue in the PAPs. The Ca\(_B\) ion has regular octahedral ligation with Asp and Asn ligands.

All three metal ions in PhoD have bonding interactions with the phosphate group present in the active site (Figs. 3B and 5A). One oxygen atom of the phosphate ligates the iron, one oxygen atom bridges Ca\(_A\) and the iron, and one oxygen atom bridges the two calcium ions. The interactions between the phosphate and the Ca\(_A\)-Fe pair replicate the tripodal \(\mu-\eta2-\eta2\)-geometry observed between phosphate and the two metal ions of sweet potato PAP in PDB structure 1XZW (26)(Fig. 3B). A similar mode of bridging tripodal coordination has also been seen for phosphate bound to the dimetal centers of \emph{Sporosarcina pasteurii} urease (29) and an organophosphate-degrading enzyme from \emph{Agrobacterium radiobacter} (30). In addition to interacting with the active site metal ions the phosphate ion in PhoD has bonding interactions with the amino acid side chains of Arg-562 and His-566 from the C-terminal capping helix (Figs. 5A and 6, A–C).

The \textbf{C-terminal Helix Occludes Access to the PhoD Active Site}—The C-terminal region of the \emph{B. subtilis} PhoD protein forms an \(\alpha\)-helix (residues 560–577), which lies over the active site (Figs. 3A and 6). This helix is part of a C-terminal extension that is found in PhoD from \emph{B. subtilis} and certain additional \emph{Bacillus} species, but is absent from other bacterial PhoD proteins and from the PAPs (Fig. 6A). The C-terminal helix is held in position by a network of salt bridges, which include interactions with the phosphate ion located in the active site, and with Asp-436, which bridges the Fe and Ca\(_A\) ions (Fig. 6C).

The C-terminal helix blocks access to the catalytic site. Furthermore, as judged by modeling studies with pNPP substituted for the phosphate molecule seen in the crystal structure, steric clashes with the helix would also prevent substrates binding to the active site. As a consequence, the C-terminal helix must move from the position seen in the crystal structure in order for substrates to bind to PhoD. The main chain crystallographic B-factor for the C-terminal helix is high (49.1 Å\(^2\)) relative to the whole protein average (28.2 Å\(^2\)), showing that the C-terminal helix is more mobile than the rest of the PhoD structure (Fig. 6D). This suggests that the helix is relatively weakly bound to...
### TABLE 1
Data collection and refinement statistics

| Data Collection          | ID23_1 | IO3  |
|--------------------------|--------|------|
|                          |        |      |
| Space group              | I4₁,22 |      |
| Molecules in asymmetric unit | 2      |      |
| Unit cell parameters (Å) | 148.05, 148.05, 347.2, 90.0, 90.0, 90.0 |      |
| Solvent content (%)      | 38     |      |
| Resolution range (Å)     | 173.6 – 2.7 | 174.7 – 3.7 |
| Number of Unique reflections | 54642  | 20953 |
| Rmerge(%)                | 11(48) | 16(49)|
| Rpim(%)                  | 12(29) | 18(25)|
| Mean I/σ(I)              | 15.6(4.0) | 11.0(4.2)|
| Completeness (%)         | 100(100) | 100(100) |
| Redundancy               | 8.4(6.8) | 8.5(8.5)|
|                         |        |      |
| MAD Phasing Statistics   |        |      |
| Sharp FOM (acentrics)    | 0.59 (136.19-11.94); 0.22(136.19-2.75); 0.11(2.82-2.68) |      |
| Sharp FOM (centrics)     | 0.20 (136.19-11.94); 0.07(136.19-2.75); 0.07(2.82-2.68) |      |
| Sharp phasing power (iso/ano): |      |      |
| Acentric                 | Peak (-)/0.487; Inflexion 0.086/0.328; Remote 0.023/0.250 |      |
| Centric                  | Peak (-)/(-); Inflexion 0.082/(-); Remote 0.021/(-) |      |
| Rcullis (acentrics)      | Peak (-)/0.947; Inflexion 0.535/0.981; Remote 0.680/0.990 | 0.864 |
| Rcullis (centric)        | Peak (-)/(-); Inflexion 0.490/(-); Remote 0.640/(-) | 0.2933/0.2681 |
| Solvent flattened FOM (overall) | 0.6559/0.7956 |      |
|                         |        |      |
| Refinement summary:      |        | 136.19 (2.0 – 1.9) |
| Resolution (Å)           |        | 17.8 (29) |
| R factor (%)             |        | 20.2 (31) |
| Free R factor (%)        |        | 0.014  |
| R.m.s.d. bond lengths (Å)|        | 1.3    |
| R.m.s.d. bond angles (°) |        | 20841  |
| No. atoms in asymmetric unit |        | 19772  |
| Of which: Protein        |        | 2      |
| Fe                       |        | 4      |
| Ca                       |        | 6      |
| Na                       |        | 420 (73 units) |
| PEG200/EG                |        | 10 (2 units) |
| Phosphate                |        | 627    |
| H2O                      |        | 96     |
| Ramachandram plot:       |        | 4      |
| Preferred (%)            |        | 0      |
| Allowed (%)              |        | 2YEQ   |
| Outliers (%)             |        |      |
| PDB code                 |        |      |
the main body of PhoD and could plausibly be released at some stage of the catalytic cycle. Attempts to observe the predicted movement of the C-terminal helix by co-crystallization with the bulky substrate analogues phosphonoacetic acid and phosphonohexanoic in place of phosphate were unsuccessful.

The function of the PhoD C-terminal helix was investigated using variants possessing truncations of this helix. These experiments show that the whole C-terminal helix is required for catalytic activity and that even the variant with the smallest truncation tested had lost the visible absorbance bands arising from the Fe$^{3+}$ ion (Fig. 7). Thus the C-terminal helix is required to either insert or retain the PhoD active site metal cofactor.

**DISCUSSION**

The structure of *B. subtilis* PhoD confirms earlier suggestions (9) that PhoD family proteins have structural features in common with eukaryotic PAPs. The two protein families share the same protein fold, and both contain a tyrosinate-ligated Fe$^{3+}$ ion at the same position in the active site. Indeed, the sole difference in the Fe$^{3+}$ coordination environment between

---

**FIGURE 3. PhoD has structural similarity to purple acid phosphatases.** A, comparison of the overall fold of PhoD (left) and sweet potato PAP (PDB 1XZW) (right). Both structures are shown in the same orientation with β-strands colored blue and α-helices colored red. The catalytic site metal ions are presented as spheres, with Fe$^{3+}$ shown in orange, Ca$^{2+}$ shown in green, and Mn$^{2+}$ shown in purple. B, comparison of the coordination environment of the PhoD metal ions (left) with those in sweet potato PAP (right). Metal ions are colored as in A, and other atoms are colored as follows: oxygen in red; nitrogen in dark blue; sulfur in yellow; phosphorus in light blue; PhoD carbons in sky blue; and PAP carbons in tan. Metal ion coordination bonds are shown by dashed lines. Alignment between the proteins was carried out with LSQKAB (44). C, electron density at the PhoD active site contoured at 1σ (gray) overlaid with anomalous Fourier maps contoured at 4σ (purple) at the x-ray wavelengths indicated at the top of each panel. The anomalous signals for iron relative to calcium are consistent with the theoretical f values for iron and calcium at these wavelengths calculated using Crossec (45), which are given under the panels.
Structure of the Phosphodiesterase PhoD

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PhoD|B.subtilis

PAPS and _B. subtilis_ PhoD is the substitution of one of the aspartic acid residues in the canonical PAPs ligand set by a cysteine (Cys-180). In this context it is worth noting that PhoD family proteins can be divided on sequence grounds into two subfamilies, one of which possesses the cysteine Fe³⁺ ligand seen in _B. subtilis_ PhoD and also _S. chromofuscus_ PLD, whereas the second has the aspartate ligand found in the PAPS site.

Notwithstanding their similarities, the PhoD and PAPS families clearly form distinct sequence groups and differ in their active site metal complement beyond the tyrosine-ligated Fe³⁺ ion. Although the PAPS bind an additional Fe²⁺, Mn²⁺, or Zn²⁺ ion, _B. subtilis_ PhoD binds two Ca²⁺ ions. Other members of the PhoD family almost certainly bind the same two Ca²⁺ ions because the coordinating amino acids are close to invariant within the family (the residue equivalent to Caₐ in _B. subtilis_ PhoD Caₐ₉ ligand Asn-272 is sometimes conservatively substituted with an Asp, which is also a good Ca²⁺ ligand). Although the position of the Caₐ ion in PhoD is similar to that of the divalent ion in the PAPS, the ligand sets provided by the two protein families are completely distinct. Caₐ in PhoD has five coordination bonds from Asp residues, whereas the divalent ion in PAPS is ligated by an invariant His₂Asp₁Asn₁ set (31). It

FIGURE 4. Amino acid sequence of PhoD. The secondary structure elements are shown above the sequence, with α-helices represented by squiggles, β strands represented by arrows, and turns represented by the letters TT. The signal peptide is highlighted in red, and the C-terminal α-helix is highlighted in blue. Metal ligating residues are indicated by blocks beneath the sequence, with those coordinating the Fe ion in brown, Caₐ in green, and Caₐ₉ in purple. Residues that coordinate two metal ions are indicated by blocks with two colors.
Because PAPs are able to catalyze phosphodiesterase reactions it is unlikely that the presence of CaB is specific to PhoD with the phosphate complex corresponding to the initial product complex formed by the reaction involving this ion.

PhoD is a phosphodiesterase and so must be capable of catalyzing two successive hydrolysis reactions. This process could involve repeating the reaction involving the bridging hydroxide, with the initial product rotating in the active site before the second reaction. Alternatively, the two reactions could utilize the nucleophilic hydroxide between the two metals ions such that one of the oxygen atoms occupies the position normally taken by the bridging hydroxide ion (26). This structure has been interpreted as representing the initial product complex in which the oxygen atom of the hydroxide nucleophile is fully bonded to the substrate phosphorus atom while still maintaining bridging coordination to the metal ions. The position of the phosphate ion in the PhoD structure is highly analogous to that in the sweet potato PAPs structure (Fig. 3B), suggesting that PhoD possesses a nucleophilic hydroxide between the Fe and CaA ions and that the PhoD structure represents the initial product complex formed by the reaction involving this ion.

A striking difference between the PhoD and PAP families is the presence of a third metal ion (CaB) in PhoD. Because PAPs have historically been considered to be true bacterial PAPs. By contrast, our structural analysis shows that the much more restricted substrates have shown that PAPs do possess this capability (38). The inferred phosphodiesterase mechanism in the PAPs is nucleophilic attack by a Fe³⁺-coordinated terminal hydroxide group followed by a second nucleophilic attack by the bridging hydroxide. A similar mechanism is plausible for PhoD with the phosphate complex corresponding to the initial product of the second hydrolytic reaction.

A striking difference between the PhoD and PAP families (and, indeed, other members of the metallophosphatase superfamily) is the presence of a third metal ion (CaB) in PhoD. Because PAPs are able to catalyze phosphodiesterase reactions at a similar rate to phosphomonoester hydrolysis with only two metal ions (38) it is unlikely that the presence of CaB is specifically linked to the phosphodiesterase functionality of PhoD.

should be noted that the earlier predictions for the non-iron ion ligands in the PhoD family are incorrect (9). A few bacteria encode proteins that conserve all PAP metal ligands (32), and these proteins can be considered to be true bacterial PAPs. By contrast, our structural analysis shows that the much more widely distributed enzymes of the PhoD family should be regarded as forming a distinct but closely related sister group to all other members of the PhoD family, analogous “cap domains” are a feature of certain other enzymes in the metallophosphatase superfamily (33–36). In most cases the role of these cap domains is unclear. Intriguingly, however, the cap domain of a mycobacterial metallophosphatase has recently been implicated in cell wall binding (33). An equivalent function for the PhoD C-terminal helix is an attractive hypothesis given the proposed role of PhoD in teichoic acid metabolism.

The way in which phosphate is bound to the active site of PhoD in our crystal structure is likely to mimic the substrate-enzyme interaction at some point in the catalytic reaction. We can get considerable insight into the likely mechanism of PhoD catalysis by interpreting the phosphate complex in the light of the extensive body of work on the mechanism of PAPs. In PAPs the two active site metal ions are bridged by a hydroxide ion. The current view is that phosphomonoester substrates coordinate both PAP metal ions. The bridging hydroxide ion then undertakes a nucleophilic attack on the phosphorus atom in-line with the bond to the leaving group. This results in the hydroxide ion being incorporated into the phosphate product (10). A structure has been determined for sweet potato PAP in which a phosphate ion is bound in a tripodal μ–η2–η2-geometry between the two metals ions such that one of the oxygen atoms occupies the position normally taken by the bridging hydroxide ion (26). This structure has been interpreted as representing the initial product complex in which the oxygen atom of the hydroxide nucleophile is fully bonded to the substrate phosphorus atom while still maintaining bridging coordination to the metal ions. The position of the phosphate ion in the PhoD structure is highly analogous to that in the sweet potato PAPs structure (Fig. 3B), suggesting that PhoD possesses a nucleophilic hydroxide between the Fe and CaA ions and that the PhoD structure represents the initial product complex formed by the reaction involving this ion.

B. subtilis PhoD has a flexible C-terminal helix overlaying the active site (Fig. 7). Although this helix is not present in most of the PAPs. Because PAPs are able to catalyze phosphodiesterase reactions at a similar rate to phosphomonoester hydrolysis with only two metal ions (38) it is unlikely that the presence of CaB is specifically linked to the phosphodiesterase functionality of PhoD.
Instead, Ca\textsubscript{a} is appropriately placed to stabilize the developing charge on the leaving group during nucleophilic attack by the bridging hydroxide ion. More generally, the additional metal ion may compensate for the weaker polarizing ability of Ca\textsuperscript{2+} relative to the divalent metal ions found in the PAPs (Fe\textsuperscript{2+}, Mn\textsuperscript{2+}, Zn\textsuperscript{2+}).

The combination of Fe\textsuperscript{3+} and Ca\textsuperscript{2+} ions used by PhoD is unusual and to the best of our knowledge has only otherwise

---

**FIGURE 6.** The C-terminal helix of PhoD occludes the active site. **A**, multiple sequence alignment of the C termini of PhoD family proteins showing the extension seen in the PhoD protein of *B. subtilis* and some other *Bacillus* species. Secondary structure elements from the *B. subtilis* PhoD structure are shown above the alignment with the \( \alpha \)-helix shown as a coil, \( \beta \)-strands shown as arrows, and \( \beta \)-turns shown by the letter T. Conserved amino acids are highlighted in red, and similar amino acids are highlighted in yellow. The black box indicates the C-terminal \( \alpha \)-helix of *B. subtilis* PhoD. Residues in the C terminus involved in salt bridges with the main structure (colored blue) are indicated with blue dots above the alignment. **B**, electrostatic surface model (\( \sim 5 \frac{kT}{e} \)) of intact PhoD (left) and with the C-terminal helix removed (right). Regions of negative charge are shown in red, and regions of positive charge are shown in blue, but with the C-terminal helix in gray. Electrostatic calculations were performed with APBS (47) and include the charges on the active site metal ions. **C**, the salt bridge network between the C-terminal helix, the bound phosphate ion, and the active site. The portion of PhoD shown corresponds to the region of the PhoD structure highlighted in panel B. **D**, B-factor putty diagram of PhoD generated by PyMOL. Flexibility is indicated by increasing ribbon girth and color scaling from blue to green.
been seen in the structurally unrelated PhoX alkaline phosphatase (4). Remarkably, therefore, two out of the three major classes of alkaline phosphatases in bacteria use the same, rare combination of metal ions. Both PhoD and PhoX are extracytoplasmic bacterial enzymes, and so the selection of Ca\(^{2+}\) as the divalent cation may simply reflect the high availability of Ca\(^{2+}\) ions in most terrestrial and marine environments (39). By contrast, the bioavailability of iron is low in many environments (40) and may limit the activity of PhoD family enzymes in such locations, as has also recently been argued for PhoX (4). Export of both PhoD and PhoX family proteins uses the Tat protein translocation system, which is dedicated to the transport of proteins that need to fold before export to bind cofactor molecules (41–43). The Tat pathway is often used by prokaryotes to export folded, iron-containing proteins taking place via the Tat apparatus. In summary, the PhoD structure reveals that members of this widely distributed family of bacterial enzymes represent a new class of Fe\(^{3+}\)- and Ca\(^{2+}\)-dependent phosphatase.

Acknowledgments—We thank Oliver Zeldin and Elspeth Garman for micro-PIXE measurements, Janet Lovett for assistance with the EPR measurements, Jeff Errington for providing B. subtilis 168 genomic DNA, and Shree Chien Yong, Magda Atilano, Petros Ligoxygakis, and Richard Daniel for discussions. We also thank the staff of beamline ID23_1 of the European Synchrotron Radiation Facility, Grenoble, France. We acknowledge the Diamond Light Source for time on beamline I03 under proposal MX9306.

REFERENCES

1. Rusch, D. B., Halpern, A. L., Sutton, G., Heidelberg, K. B., Williamson, S., Yooseph, S., Wu, D., Eisen, J. A., Hoffman, J. M., Remington, K., Beeson, K., Tran, B., Smith, H., Baden-Tillson, H., Stewart, C., Thorpe, J., Freeman, J., Andrews-Pfannkoch, C., Venter, J. E., Li, K., Kravitz, S., Heidelberg, J. F., Utterback, T., Rogers, Y. H., Falcón, L. I., Souza, V., Bonilla-Rosso, G., Eguiarte, L. E., Karl, D. M., Sathyanarathan, S., Platt, T., Bermingham, E., Gallardo, V., Tamayo-Castillo, G., Ferrari, M. R., Strauberg, R. L., Nelson, K., Friedman, R., Frazier, M., and Venter, J. C. (2007) The Sorcerer II Global Ocean Sampling expedition: northwest Atlantic through eastern tropical Pacific. PLoS Biol. 5, e77

2. Luo, H., Benner, R., Long, R. A., and Hu, J. (2009) Subcellular localization of marine bacterial alkaline phosphatases. Proc. Natl. Acad. Sci. U.S.A. 106, 21219–21223

3. Lassila, J. K., Zalatan, J. G., and Herschlag, D. (2011) Biological phosphophoryl-transfer reactions: understanding mechanism and catalysis. Annu. Rev. Biochem. 80, 669–702

4. Yong, S.C., Roversi, P., Lillington, J., Rodriguez, F., Krehenbrink, M., Zeldin, O. B., Garman, E. F., Lea, S. M., and Berks, B. C. (2014) A complex iron-calcium cofactor catalyzing phosphotransfer chemistry. Science 345, 1170–1173

5. Yamane, K., and Maruo, B. (1978) Purification and characterization of extracellular soluble and membrane-bound insoluble alkaline phosphatases possessing phosphodiesterase activities in Bacillus subtilis. J. Bacteriol. 134, 100–107

6. Eder, S., Shi, L., Jensen, K., Yamane, K., and Hulett, F. M. (1996) A Bacillus subtilis secreted phosphodiesterase/alkaline phosphatase is the product of a Pho regulon gene, phoD. Microbiology 142, 2041–2047

7. Zambonelli, C., and Roberts, M. F. (2003) An iron-dependent bacterial phospholipase D reminiscent of purple acid phosphatases. J. Biol. Chem. 278, 13706–13711

8. Marchler-Bauer, A., Zheng, C., Chitsaz, F., Derbyshire, M. K., Geer, L. Y., Geer, R. C., Gonzalez, N. R., Gwadz, M., Hurwitz, D. I., Lanzczky, C. J., Lu, F., Lu, S., Marchler, G. H., Song, J. S., Thaini, N., Yamashita, R. A., Zhang, D., and Bryant, S. H. (2013) CDD: conserved domains and protein threedimensional structure. Nucleic Acids Res. 41, D348–D352

9. Zambonelli, C., Casali, M., and Roberts, M. F. (2003) Mutagenesis of putative catalytic and regulatory residues of Streptomyces chromofuscus phospholipase D differentially modifies phosphatase and phosphodiesterase activities. J. Biol. Chem. 278, 52282–52289

10. Schenk, G., Mitić, N., Gahan, L. R., Ollis, D. L., McGeary, R. P., and Gud- dat, L. W. (2012) Binuclear metallohydrolyses: complex mechanistic strategies for a simple chemical reaction. Acc. Chem. Res. 45, 1593–1603

11. Mitić, N., Smith, S. J., Neves, A., Guddat, L. W., Gahan, L. R., and Schenk, G. (2006) The catalytic mechanisms of binuclear metallohydrolyses. Chem. Rev. 106, 3338–3363

12. Ferrer, M., Golyshina, O. V., Belouki, A., Bottger, L. H., Andreu, J. M., Polainia, J., De Lacey, A. L., Trautwein, A. X., Timmis, K. N., and Golyshin, P. N. (2008) A purple acidophilic di-ferric DNA ligase from Ferroplasma. Proc. Natl. Acad. Sci. U.S.A. 105, 8878–8883

13. Andréasson, C., Fiaux, J., Rampelt, H., Mayer, M. P., and Bukau, B. (2008)

Structure of the Phosphodiesterase PhoD

FIGURE 7. The C-terminal helix of PhoD is functionally important. A, phosphatase activity of C-terminally truncated PhoD variants. PhoD (57–583) is the intact protein. Activities were measured using pNPP as substrate. Error bars indicate the S.D. of three independent experiments. B, visible absorption spectra of wild-type PhoD (57–583)(black line) and the C-terminal truncation variant (57–549)(gray line).
