Crystal structure of the transcriptional repressor PagR of *Bacillus anthracis*

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PagR is a transcriptional repressor in *Bacillus anthracis* that controls the chromosomal S-layer genes *eag* and *sap*, and downregulates the protective antigen *pagA* gene by direct binding to their promoter regions. The PagR protein sequence is similar to those of members of the ArsR repressor family involved in the repression of arsenate-resistance genes in numerous bacteria. The crystal structure of PagR was solved using multi-wavelength anomalous diffraction (MAD) techniques and was refined with 1.8 Å resolution diffraction data. The PagR molecules form dimers, as observed in all SmtB/ArsR repressor family proteins. In the crystal lattice four PagR dimers pack together to form an inactive octamer. Model-building studies suggest that the dimer binds to a DNA duplex with a bend of around 40°.

**INTRODUCTION**

*Bacillus anthracis* is a Gram-positive bacterium that primarily infects herbivores such as cattle and sheep but can also infect humans, causing the fatal disease anthrax. *B. anthracis* belongs to the Firmicutes and is genetically very closely related to some *Bacillus cereus* strains (Helgason et al., 2000). A major difference of *B. anthracis* from *B. cereus* is that it harbours two virulence plasmids, pXO1 and pXO2. The pXO2 plasmid codes for proteins responsible for capsule formation. The pXO1 plasmid encodes the structural genes required for the production of toxins: *pagA*, which codes for protective antigen PA, *cya*, which codes for edema factor (EF), and *lef*, which codes for lethal factor (LF). The *pag* operon consists of two genes, *pagA* and *pagR*, the latter being a weak autogenous repressor co-transcribed with *pagA* (Hoffmaster & Koehler, 1999).

Sequence comparisons suggest that PagR is closely related to the SmtB/ArsR family of metal-binding repressors involved in the repression of metal-resistance genes (Busenlehner et al., 2003; Xu et al., 1996). The best-characterized members of this family are SmtB from *Synechococcus* (Eicken et al., 2003), and CzxA (Eicken et al., 2003) and CadC (Ye et al., 2005) from *Staphylococcus aureus*. They bind to the DNA either at the promoter region or in the region upstream of the promoter, inhibiting transcription. The binding sites are generally imperfect inverted repeats of 12 bases separated by two bases. In general, these repressors regulate the expression of genes involved in metal resistance, such as those encoding efflux pumps.

All members of the SmtB/ArsR repressor family have helix-turn-helix motifs for DNA binding. The algorithm of Dodd & Egan (1990) predicted a helix-turn-helix motif for the PagR sequence between amino acid residues 42 and 64. PagR probably represses transcription by binding directly to the promoter regions of the controlled gene. DNase I footprinting experiments showed that PagR protects the *pag* promoter region from nucleotide −52 to nucleotide −3 with respect to the transcription start site of the P2 promoter (Mignot et al., 2003). The regions of the *sap* promoter protected by PagR extend from nucleotide −63 to nucleotide −15 with respect to the transcription start site of *sap*. Further, PagR protects the *eag* promoter region from nucleotide +45 to nucleotide +94 with respect to the σ^H^-dependent *eag* promoter (Mignot et al., 2002). The PagR-protected region in *pagAR* and the *sap* promoter overlaps with the σ^A^-recognition elements at the promoter −35 and −10 sites. These results suggest that PagR represses the *pagAR* and *sap* genes by preventing a σ^A^-
complexed RNA polymerase from binding to the *sap* and *pag* promoters (Hoffmaster & Koehler, 1999; Mignot et al., 2003). It was reported earlier that deletion of PagR increases the expression level of AtxA (Hoffmaster & Koehler, 1999), implying that PagR may also bind to the promoter region of *atxA*.

The role of PagR in virulence in animals is untested but there is little doubt of its role in the regulation of S-layer proteins. The S-layer proteins are required for the anchoring of the virulence-plasmid-encoded BslA, which is essential for *B. anthracis* to breach the blood–brain barrier (Kern & Schneewind, 2008). An examination of the sequences of the PagR-protected regions of the *pag*, *sap* and *eag* promoters revealed no specific DNA sequences for PagR binding. To gain a better understanding of PagR function, we crystallized the PagR protein and solved its crystal structure at a resolution of 1.8 Å.

### METHODS

#### Cloning, expression and purification of PagR.

The pagR gene was PCR amplified from chromosomal DNA of *B. anthracis* (34F2 strain) using oligonucleotides BaPagR5 (5’-AGAATTATCATGACGATATTTTGTAGATC-3’) and BaPagR3’Bam (5’-CGCGGATCCCTATATTGGATAGGGTTTAAC-3’). After digestion of PCR-amplified DNA with *Bsp*HI and *Bam*HI, it was ligated with plasmid pET28a (Novagen), generating an ORF of the gene that codes for the nine-auxotrophic strain *E. coli* B834(DE3). The cells were grown in LB medium.

Selenomethionine-containing PagR was expressed in the methionine-defined medium containing 40 mg seleno- DL-methionine ml⁻¹, using oligonucleotides BaPagR5 (5’-AGAATTATCATGACGATATTTTGTAGATC-3’) and BaPagR3’BspH (5’- CGCGGATCCCTATATTGGATAGGGTTTAAC-3’). After digestion of PCR-amplified DNA with *Bsp*HI and *Bam*HI, it was ligated with plasmid pET28a (Novagen), generating an ORF of the gene that codes for the PagR protein without a histidine tag. PagR was expressed in pET28a (Novagen), generating an ORF of the gene that codes for the nine-auxotrophic strain *E. coli* B834(DE3) grown at 30 °C in LB medium. Selenomethionine-containing PagR was expressed in the methionine-auxotrophic strain *E. coli* B834(DE3). The cells were grown in defined medium containing 40 mg seleno-DL-methionine ml⁻¹. The protein purification was carried out in 50 mM HEPES, 50 mM NaCl, pH 8.0 buffer (buffer A), using a two-step chromatographic purification procedure. After sonication and centrifugation, the supernatant was loaded on a cation-exchange column that was washed with a buffer A and eluted with buffer B (buffer A containing an additional 1 M NaCl). The fractions including PagR were pooled, desalted and further loaded onto a heparin column, which was again washed with buffer A and eluted with buffer B. Purified PagR was stored in buffer containing 5 mM Bistris, 50 mM NaCl, pH 6.5. The protein was concentrated using a Centriprep YM-3 concentrator to 6–10 mg ml⁻¹, and DTT was added to a final concentration of 5 mM. The incorporation of selenomethionine was confirmed by MALDI (data not shown).

**Protein crystallization.** PagR crystals were obtained at 22 °C by the hanging-drop method using a reservoir solution containing 0.2 M sodium acetate, 1.6 M NaH₂PO₄ and 0.4 M K₂HPO₄. Crystals grew in 2 weeks to a maximal size of 0.08 × 0.08 × 0.08 mm in a tetragonal space group P4₁2₁2, with unit cell dimensions of a=73.01 Å, c=151.12 Å. The volume of the unit cell suggested that there were four PagR molecules in the asymmetrical unit with a solvent content of 48%. Crystals of PagR containing selenomethionine grew under similar conditions, but the crystal size improved to 0.2 × 0.2 × 0.2 mm after 2 weeks. The crystals were flash-frozen after transferring them into a cryosolution containing 20% ethylene glycol.

**Data collection.** Multi-wavelength anomalous diffraction (MAD) data were collected at the Advanced Light Source (ALS, Berkeley, CA) on beamline 8.3.1 at wavelengths corresponding to the peak (λ₁) and low-energy remote (λ₂) of a selenium MAD experiment. A native dataset (λ₀) to 2.4 Å was also collected on beamline 8.3.1. The datasets were collected at 100 K using ADSC CCD detectors. The MAD data were integrated and reduced using MOSFLM and then scaled with SCALA from the CCP4 program (Leslie, 2006). The native data were integrated and reduced using Denzo and then scaled with the program Scalepack (Otwinowski & Minor, 1997). Data statistics are summarized in Table 1.

**Structure solution and refinement.** PagR is a 99-residue protein containing five methionines. The crystals of selenomethionine-

### Table 1. Crystallographic data

<table>
<thead>
<tr>
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<th>SeMet λ₁ peak inflection</th>
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<th>Native λ₀</th>
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<td>Wavelength (Å)</td>
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<td>1.00344</td>
<td>0.98012</td>
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<tr>
<td>Resolution (Å)</td>
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<td>1.75 (1.75–1.81)</td>
<td>2.40 (2.40–2.49)</td>
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<tr>
<td>No. of total reflections</td>
<td>23 1301</td>
<td>24 7423</td>
<td>121 266</td>
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<tr>
<td>No. of unique reflections</td>
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<td>86.8 (48.9)</td>
<td>99.1 (99.3)</td>
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<td>7.6 (39.7)</td>
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<td>36.0 (2.4)</td>
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<td>Figure of merit</td>
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<td><strong>Refinement statistics</strong></td>
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<td>Model composition</td>
<td>341 residues, 79 H₂O</td>
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<td>Resolution limits (Å)</td>
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<td>Reflection in working/test sets</td>
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<td>R/Refe (%)</td>
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<td>Bond (Å) angle (°) rms</td>
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<td><strong>Ramachandran plot</strong></td>
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<td></td>
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<tr>
<td>Additional allowed regions</td>
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incorporated protein diffracted to a resolution of 1.8 Å. Structure solution was carried out by multi-wavelength anomalous dispersion (MAD) techniques. The initial phase were determined with the 1.8 Å selenium MAD data (l1, l2) using the program SOLVE (Terwilliger, 2003). Automatic model building was performed with RESOLVE (Terwilliger, 2003) and the model was completed using the program O (Jones et al., 1991); refinement of the coordinates was carried out using the diffraction data at the wavelength l2 (Table 1) using program CNS1.0 (Brunger et al., 1998). Refinement statistics are summarized in Table 1. The final model consists of four molecules of PagR and 79 water molecules in the asymmetrical unit. No electron density was observed for residues 1–13 and 97–99 in Mol.A, 1–10 and 97–99 in Mol.B, 1–10 and 97–99 in Mol.C, and 1–12, 73–79 and 99 in Mol.D.

RESULTS AND DISCUSSION

Description of the overall structure


PagR is an α/β protein with the same fold as the members of SmtB/ArsR family of metal-binding repressors. These proteins belong to the class of winged helix–turn–helix (HTH) motif-containing transcription factors. They have a unique topology of five α-helices and a β-hairpin in the following order: αααααββx. The structure of PagR comprises x-helices H1 (residues 14–25), H2 (27–39), H3 (43–52), H4 (55–65) and H5 (86–96) and a two-stranded β sheet with β1 (residues 71–75) and β2 (78–83) (Fig. 1a, b). Two of the helices, H3 and H4, along with the connecting residues form the putative HTH motif seen in many prokaryotic and eukaryotic DNA-binding proteins. The β-strands, β1 and β2, are arranged in an anti-parallel manner to form a hairpin structure that is anchored by hydrophobic and hydrogen-bonding interactions to the rest of the protein. The conserved residues from helices H2 and H3, the termini of the β hairpins and the H5 helix form the hydrophobic core. The classic inverse γ-turn between helices H1 and H2 is a common feature in the SmtB/ArsR family and it is also present in PagR. A GXG motif (where X is an amino acid with a large hydrophobic side chain) seen towards the C-terminal end of the recognition helix in most transcription factors is absent in PagR. The type II’ β-turn in the β-hairpin with the Gly in the second position is also a common feature observed in most members of the SmtB/ArsR family of repressors. The residue Y82, located on strand β2, has been predicted to be a potential tyrosine kinase phosphorylation site (Hoffmaster & Koehler, 1999) and its side chain points towards the recognition helix (Fig. 1a). The residues Y81 and L62 are part of the hydrophobic core between the helical part of the protein and the β-strands (Fig. 1a).

The conserved nature of the residues in the interface between the β-strands and helices indicates that these residues are important for the structural integrity of the protein. The above-mentioned structural features in this fold play an important role in understanding the allosteric mechanism of DNA binding in this family of repressors (Eicken et al., 2003). These conserved features in PagR structure suggest that it might have a mechanism similar to that of the members of the SmtB/ArsR family.

Two types of intermolecular associations

The metal-binding repressor proteins in the SmtB/ArsR family (Cook et al., 1998; Eicken et al., 2003) form
elongated dimers, with the recognition helices separated by about 34 Å to interact with the DNA. In the crystal lattice, PagR molecules also form such dimers. Mol.A and Mol.B form a dimer and they are related by a non-crystallographic twofold axis (Fig. 2a), while Mol.C forms a dimer with its symmetry-related partner and so does Mol.D. The association of these two monomers buries a total of 17% of the surface area of the two molecules at the interface. The core of this binding surface is a hydrophobic region formed by the residues of helices H1 and H5 from each monomer, which run antiparallel to each other. Dimerization of the molecule appears to be critical for its function, as it positions the recognition helices suitably for interactions with DNA.

The crystal lattice displays an additional mode of association (Fig. 2b). Mol.B and Mol.C (similarly Mol.A and Mol.D) are also related by a non-crystallographic twofold axis, and they associate through the interactions of helix H1 (Mol.B) with H4 of (Mol.C) and vice versa. The association buries a total of 9% of the surface area of two molecules at the interface and it can be visualized as another mode of dimer formation (Fig. 2b). By using these two modes of associations alternately, the PagR molecules form cyclical octamers in the crystal lattice (Fig. 2c). The second mode of association (Mol.B with Mol.C and Mol.A with Mol.D) is stabilized by two salt bridges between each pair of monomers at their interfaces, indicating that the stability of the octamer may be dependent on the ionic strength of the medium.
While there is evidence for the presence of dimers in solution from mass spectroscopy (data not shown), there is no experimental evidence so far for the existence of higher-order oligomers in solution. Hydrophilic interactions play a major role in the second mode of dimer association, and therefore formation of stable tetramers or octamers would appear unlikely. In the crystal lattice, when the molecules pack themselves to form an octamer, the recognition helices are no longer accessible for interaction with DNA (Fig. 2c). Therefore the formation of the octamer could be thought of as an inactivation mechanism.

**Comparison of PagR with metal-sensing proteins in the SmtB/ArsR family**

In prokaryotes, there are two general classes of transcriptional regulatory proteins with the ability to respond to stress-induced metal toxicity, namely the MerR family and the SmtB/ArsR family (Busenlehner et al., 2003). MerR-like proteins function as repressors in the absence of metal ions, but function as activators upon metal binding. Conversely the SmtB/ArsR family of proteins act exclusively as prokaryotic transcriptional repressors which regulate the expression of genes associated with metal sequestration or efflux in both Gram-positive and Gram-negative bacteria (Busenlehner et al., 2003). PagR is closely related to the SmtB/ArsR family of proteins.

So far, six different crystal structures of the members of SmtB/ArsR family have been reported. They are SmtB in apo form (Cook et al., 1998), SmtB in one and two zinc-bound forms, CzrA in apo and zinc-bound form (Eicken et al., 2003), and CadC in zinc-bound form (Ye et al., 2005). PagR has 28% sequence identity with CzrA and 21% with CadC and SmtB. The structure of PagR has close resemblance to the structures of the proteins in this class. There are however two main differences: (i) PagR does not have metal-binding sites, and (ii) the PagR dimer has a more curved architecture compared to most of the molecules in this class.

While PagR does not possess metal-binding sites, certain members of the SmtB/ArsR family have two distinct metal-binding sites (Cook et al., 1998; Busenlehner et al., 2003). One of them is the α3N site formed by Cys residues of the third α-helix H3 along with Cys residues from the N-terminal region and it binds large, softer metal ions such as Cd(II), Pb(II) and Bi(III). The other is the α5 site; it is formed by carboxylate or imidazole side chains contributed by H5 from each monomer and it binds smaller, harder metal ions such as Co(II) and Zn(II) (Busenlehner et al., 2001, 2002). The two α3N sites are located on opposite ends of the dimer while the two α5 sites are at the dimeric interface (Fig. 3). PagR lacks the residues needed to form any of these binding sites.

In CadC, only the α3N site has functional relevance. On the other hand, in SmtB and CzrA, the N-terminal site is not well conserved and is not known to be involved in any metal binding, but the C-terminal metal-binding site α5 has a regulatory role. As PagR does not have the metal-binding sites, its function must be regulated by some other as yet unknown means.

**B. anthracis** Sterne strain has four additional proteins related to the SmtB/ArsR family. Three of these appear to be cotranscribed with metal-associated functions: BAS0563, in an operon with a heavy-metal-transporting ATPase; BAS2974, in an operon with a protein known to reduce arsenate to arsenite, and BAS3305, in an operon with a zinc-requiring alcohol dehydrogenase. The fourth gene, BAS4321, appears to be transcribed alone. All of these maintain the metal-binding site.

The crystallographic data of repressors in the SmtB/ArsR family illustrate that the metal binding influences the conformation of the molecule and provides a basis for understanding the allosteric regulation of operator–promoter binding (Eicken et al., 2003). Metal binding at the dimeric interface causes a rigid body reorientation of one subunit with respect to the other. In SmtB, the metal binding causes a dramatic change, a movement of 2.4 Å for recognition helix in each of the subunits (Eicken et al., 2003). The altered conformation is less suitable for DNA binding, resulting in a drastic drop of the binding affinity and causing the release of repressor from the promoter sites and subsequent expression of metal-regulatory proteins.

**Dimer architecture**

Comparison of the PagR dimer with other structures of the SmtB/ArsR family shows that PagR has a more curved architecture. Fig. 4 shows the superposition of CzrA with PagR and it indicates that CzrA is more linear while PagR is more curved, making the recognition helices less parallel. It appears that the curvature of PagR and the concomitant
deviated orientation of the recognition helices would correspond to a significant bend in the DNA.

**Putative DNA-binding residues**

PagR is the transcriptional repressor that regulates the pagAR operon and the S-layer genes *cag* and *sap* in *B. anthracis* by binding to the promoter regions of these genes (Mignot *et al.*, 2003). As the molecule possesses the HTH motif, characteristic of DNA-binding proteins, it appears that helix H4 would function as the recognition helix and interact with the major groove of DNA.

In order to envision the mode of DNA binding, we examined the geometry of the binding of protein dimers with a twofold symmetry like the lambda Cro protein to palindromic DNA (Anderson *et al.*, 1981). The two recognition helices bind to the major groove of DNA separated by 34 Å, corresponding to the pitch of the DNA helix. While some dimers bind to straight B-form DNA (Beamer & Pabo, 1992), certain others bind to bent forms of DNA (Schultz *et al.*, 1991). For example, NarL binding to DNA causes the DNA to bend by 42° (Maris *et al.*, 2002). A comparison of the orientation of the recognition helices in NarL and PagR shows that PagR could have reasonably good interactions with the DNA conformation observed in the NarL–DNA complex. Therefore, we propose that PagR dimer may bind to DNA with a similar bend of around 40° (Fig. 5). The side chains of the residues Lys64, Gln60 and Ser56 of the recognition helix are oriented favourably to interact with the DNA major groove. A recent study (Hadjifrangiskou & Koehler, 2008) on the pag operon using *in vitro* analysis and *in silico* modelling shows that there is intrinsic curvature associated with anthrax toxin promoter genes, in agreement with our proposed model.

Three basic residues, His26, Arg29 and His61, cluster together and are favourably oriented for interactions with the DNA backbone. One of these residues, Arg29, is highly conserved in all the SmtB/ArsR family of proteins (Busenlehner *et al.*, 2003). Modelling of the DNA complex of PagR indicates that Arg29, His26 and His61 could interact with the backbone and major groove of DNA. Sequence alignment (Supplementary Fig. S1, available with the online version of this paper) shows that residues S59, L62 and Y81 are also highly conserved in all the SmtB/ArsR family of proteins. Residue S59 lies on the recognition helix and could have interactions with DNA. As shown in Fig. 1(a), L62 and Y81 lie between the β-hairpin and the α-helices.

DNA footprinting data (Mignot *et al.*, 2003) show that PagR binding protects the region of the DNA from nucleotides −52 to −3, a length representing nearly five helical turns. Additionally, the footprinting data suggest that PagR binds only one operator site at each promoter. The model of DNA–PagR binding (Fig. 5) shows that binding of PagR dimer accounts only for about 2.5 helical turns, suggesting the existence of two tandem binding sites for the promoter.

**Conclusions**

The crystal structure of PagR reveals a fold similar to that of members of SmtB/ArsR family of repressors and shows that it forms a dimer in a similar manner. In the crystal lattice four dimers are packed together to form an octamer. While the members of the SmtB/ArsR family are regulated by metals, the lack of metal-binding sites in PagR points to other modes of regulation. The structure provides a basis for further studies on the role of environmental and host factors in transcription regulation in *B. anthracis*.

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