Activation of the latent PlcR regulon in \textit{Bacillus anthracis}

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Many genes in \textit{Bacillus cereus} and \textit{Bacillus thuringiensis} are under the control of the transcriptional regulator PlcR and its regulatory peptide, PapR. In \textit{Bacillus anthracis}, the causative agent of anthrax, PlcR is inactivated by truncation, and consequently genes having PlcR binding sites are expressed at very low levels when compared with \textit{B. cereus}. We found that activation of the PlcR regulon in \textit{B. anthracis} by expression of a PlcR–PapR fusion protein does not alter sporulation in strains containing the virulence plasmid pXO1 and thereby the global regulator AtxA. Using comparative 2D gel electrophoresis, we showed that activation of the PlcR regulon in \textit{B. anthracis} leads to upregulation of many proteins found in the secretome of \textit{B. cereus}, including phospholipases and proteases, such as the putative protease BA1995. Transcriptional analysis demonstrated expression of BA1995 to be dependent on PlcR–PapR, even though the putative PlcR recognition site of the BA1995 gene does not exactly match the PlcR consensus sequence, explaining why this protein had escaped recognition as belonging to the PlcR regulon. Additionally, while transcription of major PlcR-dependent haemolysins, sphingomyelinase and anthrolysin O is enhanced in response to PlcR activation in \textit{B. anthracis}, only anthrolysin O contributes significantly to lysis of human erythrocytes. In contrast, the toxicity of bacterial culture supernatants from a PlcR-positive strain towards murine macrophages occurred independently of anthrolysin O expression \textit{in vitro} and \textit{in vivo}.

\section*{INTRODUCTION}

\textit{Bacillus cereus} and \textit{Bacillus anthracis}, members of the \textit{B. cereus} group of bacteria, have high genetic similarity (Ivanova \textit{et al.}, 2003; Read \textit{et al.}, 2003), and may even constitute a single species (Helgason \textit{et al.}, 2000). While \textit{B. cereus} is an opportunistic human pathogen (Drobniewski, 1993), \textit{B. anthracis} is the aetiologic agent of anthrax, a disease with a high lethality in many animal species, including humans (Mock & Fouet, 2001). The strikingly different behaviour of these closely related species is attributed to \textit{B. anthracis} having several additional genetic features, such as the virulence plasmids pXO1 and pXO2. These plasmids encode genes for the anthrax tripartite toxin, composed of protective antigen (PA), lethal factor (LF) and oedema factor (EF), and genes for the biosynthetic enzymes for the antiphagocytic poly-\(\gamma\)-D-glutamic acid capsule (Mock & Fouet, 2001). Another unique genetic trait involves the pleiotropic transcriptional regulator PlcR (Agaisse \textit{et al.}, 1999; Lereclus \textit{et al.}, 1996). In \textit{B. cereus}, PlcR acts by binding to a well-defined site (the PlcR box), consisting of a 16 bp consensus sequence present in promoter regions of PlcR-regulated genes (Agaisse \textit{et al.}, 1999; Okstad \textit{et al.}, 1999). Furthermore, PlcR action requires the participation of a secreted, processed and reimported heptapeptide derived from PapR (Bouillaut \textit{et al.}, 2008; Lereclus \textit{et al.}, 1996; Okstad \textit{et al.}, 1999; Slamti \textit{et al.}, 2004). Comparative protein and RNA expression profiles of PlcR-positive and -negative \textit{B. cereus} and \textit{Bacillus thuringiensis} strains have identified genes regulated by PlcR that encode collagenases, haemolysins, phospholipases and enterotoxins (Gohar \textit{et al.}, 2002, 2008; Slamti \textit{et al.}, 2004). Conversely, in \textit{B. anthracis}, a nonsense mutation in \textit{plcR} results in a premature translational stop (Agaisse \textit{et al.}, 1999; Slamti \textit{et al.}, 2004), and consequently gene expression in \textit{B. anthracis} differs drastically from that in \textit{B. cereus}. Certain phenotypic properties of \textit{B. anthracis}, such as lack of haemolytic activity towards erythrocytes (Burdon, 1956), can be attributed to the non-functional PlcR, even though the genes for at least two haemolysins are present in the \textit{B. anthracis} genome. The best-characterized haemolysin of \textit{B. anthracis} is anthrolysin O, a member of the...
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cholesterol-dependent family of haemolysins, which causes lysis of a variety of cells (Mosser & Rest, 2006; Shannon et al., 2003; Tweten, 2005). Although *B. anthracis* strains are non-haemolytic on blood agar plates, this PlcR-dependent haemolysis is expressed under certain culture conditions as well as *in vivo* (Ross & Koehler, 2006; Shannon et al., 2003).

In *B. anthracis*, studies of the putative PlcR regulon have been restricted to sequence searches for PlcR boxes and comparisons with *B. cereus*, and have resulted in the identification of about 50 *B. anthracis* genes that are potentially subject to regulation by PlcR (Mignot et al., 2001; Rasko et al., 2005; Read et al., 2003). Several studies report the activation of the PlcR regulon in *B. anthracis* (Mignot et al., 2001; Pomerantsev et al., 2003, 2004). One study shows that expression of PlcR in *B. anthracis* affects sporulation of bacteria containing the virulence plasmid pXO1, but not of bacteria lacking either pXO1 or the pXO1-encoded regulator AtxA alone, and it has been hypothesized that in *B. anthracis*, functional PlcR is incompatible with AtxA-controlled gene regulation (Mignot et al., 2001). However, subsequent reports have described haemolytic *B. cereus* strains having pXO1-like plasmids (Hoffmaster et al., 2004, 2006; Klee et al., 2006), and a recent report describes the presence of two AtxA homologues in a *B. cereus* strain containing an active PlcR regulator (Passalacqua et al., 2009). These observations motivated us to readdress the hypothesis that AtxA cannot coexist with an active PlcR system. We have previously shown that the haemolytic activity of *B. anthracis* can be drastically increased upon introduction of plasmid pFP12 containing a PlcR–PapR fusion protein, leading to growth phase-independent activation of the PlcR regulon in *B. anthracis*. This fusion protein contains a full-length PlcR fused to PapR after activation of the PlcR regulon in *B. anthracis* strain containing an active PlcR regulator (Passalacqua et al., 2009). These observations motivated us to readdress the hypothesis that AtxA cannot coexist with an active PlcR system. We have previously shown that the haemolytic activity of *B. anthracis* can be drastically increased upon introduction of plasmid pFP12 containing a PlcR–PapR fusion protein, leading to growth phase-independent activation of the PlcR regulon in *B. anthracis*. This fusion protein contains a full-length PlcR fused to PapR after deletion of the stop codon and one of three repetitive MKK motifs (Pomerantsev et al., 2004). In this study, using plasmid pFP12, we investigated the effect of a functional PlcR regulon in *B. anthracis* with regard to sporulation, protein expression, anthrolysin-dependent haemolysis, and toxicity towards macrophages *in vitro* and *in vivo*.

**METHODS**

**Bacterial strains, media and growth conditions.** Table 1 lists all strains and plasmids used in this study. Bacteria were routinely grown in Luria–Bertani (LB), Brain–Heart Infusion (BHI; BD Biosciences) or nutrient broth with yeast extract (NBY) containing 0.8 % nutrient broth (Sigma), 0.3 % yeast extract, 9 % heat-inactivated equine serum (HyClone), and 0.6 % NaHCO₃ at 37 °C and 225 r.p.m. in either air or air supplemented with CO₂ regulated at 5 % for media supplemented with NaHCO₃. AtxA functionality was tested for bacteria grown in NBY broth. Spores were prepared as described previously (Sastalla et al., 2009). For haemolytic studies, bacteria were grown on tryptic soy agar with 5 % sheep blood (both BD Biosciences) or BHI broth. When required, the following antibiotics were added to *Escherichia coli* or *B. anthracis* cultures: ampicillin (100 µg ml⁻¹), erythromycin (10 µg ml⁻¹), kanamycin (20 µg ml⁻¹), tetracycline (10 µg ml⁻¹) and spectinomycin (150 µg ml⁻¹). All antibiotics were purchased from Sigma.

**DNA techniques and mutagenesis.** For generating an anthrolysin knockout strain, two *alo*-adjacent fragments were amplified using primers ALL and ALR (fragment *aloLF*), and primer ARL in combination with ARR (fragment *aloRF*), harbouring EcoRI and SpeI sites, respectively (Table 1). Amplified fragments were separately cloned into vector pSC, a plasmid allowing sequential generation of markerless deletions using the Cre/Lox system (Pomerantsev et al., 2009). Briefly, plasmid pSC-*aloRF* was introduced into *B. anthracis* electrocompetent cells as described previously (Park & Leplla, 2000). Transformants were selected at 30 °C on erythromycin and chromosomal integration of the plasmid was achieved by a temperature shift to 37 °C. Plasmid pCrePAS was introduced, resulting in excision of the inserted plasmid backbone by Cre-recombinase activity, leaving a single loxP site. Plasmid pSC-*aloRF* was introduced and the 30 °C/37 °C selection cycle followed by introduction of pCrePAS was repeated, resulting in excision of residual vector and the entire *alo* gene from the bacterial chromosome.

Plasmid pFP12 (Pomerantsev et al., 2004) was introduced by electroporation. To verify the presence (i.e. retention) of pXO1 in *B. anthracis*, bacteria were resuspended in Tris/EDTA buffer (Quality Biological), boiled for 2 min, and supernatants were used as template for the PCR using *Taq* polymerase (Qiagen) and PA (pagA) gene primers PA1 and PA2 (Table 1). For sequencing of *atxA*, primers AtxSeq1 and AtxSeq5 generated a 1742 bp PCR product that was subsequently used as template for sequencing reactions using primers AtxSeq1, AtxSeq2, AtxSeq3, AtxSeq4 and AtxSeq5.

**Sporulation efficacy.** Bacteria were grown on sporulation agar (Sastalla et al., 2009) for 5 days, analysed microscopically using a Nikon Eclipse TE200U microscope, scraped off the plates, and resuspended in 1 ml sterile distilled water. Spore suspensions were heat-treated at 65 °C for 30 min and plated on LB agar. The percentage of viable spores was calculated using the following equation: (c.f.u. after heat treatment/c.f.u. before heat treatment) × 100

**2D SDS-PAGE and protein identification.** Bacteria were grown to the late exponential/early stationary growth phase and centrifuged at 9000 g and filter-sterilized supernatants were precipitated with TCA (10 % final concentration), washed in ice-cold acetone and resuspended in 2D-rehydration buffer [8 M urea, 2 % (w/v) CHAPS, 0.5 % (v/v) ZOOM carrier ampholytes for pH range 3–10 (Invitrogen), 0.002 % bromophenol blue, 20 mM DTT]. Proteins were quantified using 2D Quant (Amersham), and 50 µg was separated on a NuPAGE gel using the IPG Runner system (Invitrogen) according to the manufacturer’s recommendations. Gels were Coomassie-stained and spots of interest were excised, trypsin-digested and analysed by MS. Retrieved data were searched against the *B. anthracis* Ames strain database using Mascot as the search engine. Peptides with a Mascot protein score higher than 60 were considered significant, and the most abundant protein present in each spot is listed in Table 2. MS-MS analysis was performed at the National Institute of Allergy and Infectious Diseases (NIAID) Research Technology Branch (RTB) Core Facility. Protein alignments and sequence analyses were performed using Lasergene MEGALIGN, PSORTb (Gardy et al., 2005), SignalP (Emanuelsson et al., 2007), Prosite (http://expasy.org/prosite) and TMpred (http://www.ch.embnet.org).

**Western blotting.** Filter-sterilized supernatants of bacterial cultures grown in NBY medium to stationary phase were separated on NuPAGE gels (Invitrogen) and blotted onto nylon membranes (Osmonics). Membranes were incubated with 2 µg mouse monoclonal antibodies ml⁻¹ against either PA (PA-05-A-G1) or LF (LF-03-A-G1) (both from the Naval Medical Research Center) in 1 % skim milk, and anti-mouse IgG conjugated to horseradish peroxidase (KPL) was used as secondary antibody. Membranes were developed using a tetramethylbenzidine (TMB) peroxidase substrate (KPL).

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RNA isolation and RT-PCR. Cultures grown in BHI to late exponential/stationary phase were centrifuged, and RNA was stabilized using RNA Protect Bacteria reagent (Qiagen) before further isolation using a Qiagen RNeasy kit. Further preparation of RNA and RT-PCR was performed as described previously (Sastalla et al., 2009). To amplify targeted mRNAs, the following oligonucleotides were used: primers BA1995fw in combination with BA1995rv, primers Sph1 and Sph2, and primers Alo1 and Alo2 (Table 1). Internal control reactions were performed with primers GyrAfw and GyrArv as described previously (Sastalla et al., 2009). Band intensities were quantified using ImageJ version 1.40 g.

Haemolytic assays. Human blood obtained from the National Institutes of Health (NIH) Blood Bank was washed with PBS and resuspended in PBS to the original volume. A 40 μl volume of blood was mixed with 160 μl culture supernatant obtained from bacteria grown in BHI medium to which 20 mM cysteine had been added, incubated at 37 °C for 45 min and centrifuged at 500 g, and the A540 of supernatants was measured in a SpectraMAX 190 plate reader (Molecular Devices). A haemolysis value of 100 % corresponded to haemolysis obtained by addition of 0.1 % SDS in BHI broth. For some experiments, cholesterol at a final concentration of 25 mM was added.

Preparation and use of bone marrow-derived macrophages (BMDMs). Bone marrow cells from 8–14-week old DBA/2J mice (Jackson Laboratories) were cultured in two parts complete Dulbecco’s modified Eagle’s medium (DMEM) [10% fetal bovine serum (FBS), 10 mM HEPES, 50 μg gentamicin ml⁻¹] and one part L929 cell culture

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**Table 1. Bacillus strains, plasmids and primers used in this study**

Restriction sites are underlined.

| Strain, plasmid or primer | Relevant characteristic(s) | Source or reference |
|---------------------------|---------------------------|---------------------|
| **B. anthracis strains**  |                           |                     |
| Sterne 34F2               | pXO1⁺, Plc⁻             | Ivins et al. (1986) |
| SdT                       | 34F2, pXO1⁺, Plc⁻       | Pomerantsev et al. (2004) |
| SdT12                     | 34F2, pXO1⁺, harbouring pFP12, PlcR⁺ | This study |
| Sterne pFP12              | 34F2 harbouring pFP12, PlcR⁺ (pXO1⁺) | This study |
| Sterne AALO               | 34F2, anthrolysin-negative (pXO1⁺) | This study |
| Sterne AALO pFP12         | 34F2, anthrolysin-negative, harbouring pFP12, PlcR⁺ (pXO1⁺) | This study |
| **B. cereus strain**      |                           |                     |
| 569                       |                           | Reddy et al. (1987) |
| **E. coli strains**       |                           |                     |
| TOP10                     |                           | Invitrogen          |
| XL2-Blue                  |                           | Stratagene          |
| SCS110                    |                           | Stratagene          |
| **Plasmids**              |                           |                     |
| pFP12                     | Encodes a PlcR–PapR fusion protein | Pomerantsev et al. (2004) |
| pSC                        | Contains loxP site for B. anthracis mutagenesis | Pomerantsev et al. (2009) |
| pSC-aloLF                  | pSC with ALL and ALR-derived amplicon | This study |
| pSC-aloRF                  | pSC with ARL and ARR-derived amplicon | This study |
| pCrePAS                    | Contains Cre recombinase | Pomerantsev et al. (2009) |
| **Primers**               | **Sequence (5’–3’)**     | Relevant property   |
| ALL                        | ccccgaattcagaagcagaaagccccgcag | Amplification of aloLF |
| ALR                        | cccccactagtggctgtttctgcaaaa | Amplification of aloRF |
| ARL                        | cccccgaattcgtcatgggaatggtggaaga | Amplification of pagA |
| ARR                        | cccccactagtggcccacaaataagctcaa | Used to sequence AtxA |
| PA1                        | gattcaggcagaagtttaaaagcag | Used to sequence AtxA |
| PA2                        | cccccaaaatatcaaaagagaagc | Used to sequence AtxA |
| AtxSeq1                    | tgcatttcctagtaaagccccaaaaaaa | Used to sequence AtxA |
| AtxSeq2                    | tcaagattgaggagatttcg | Used to sequence AtxA |
| AtxSeq3                    | tcacaattcctgctaaacatctc | Used to sequence AtxA |
| AtxSeq4                    | ccttggcgaagtaaaatcc | Used to sequence AtxA |
| AtxSeq5                    | tattatctagctatagctatgtagctc | Used to sequence AtxA |
| BA1995fw                   | cgcgctatctctatctatctg | Used for BA1995 RT-PCR |
| BA1995rv                   | aggctgcttcatctgctctgtgc | Used for BA1995 RT-PCR |
| Sph1                       | tatccgaattcggagaaag | Used for sph RT-PCR |
| Sph2                       | gaaagctgctagcttttgaa | Used for sph RT-PCR |
| Alo1                       | tgggcttgagccgctagatg | Used for alo RT-PCR |
| Alo2                       | cactgctgctgctgctgctgta | Used for alo RT-PCR |
| GyrAfw                     | aaaaacgtgtcccagctggagc | Used for gyrA RT-PCR |
| GyrArv                     | acattagcattggccagcag | Used for gyrA RT-PCR |

RNA isolation and RT-PCR. Cultures grown in BHI to late exponential/stationary phase were centrifuged, and RNA was stabilized using RNA Protect Bacteria reagent (Qiagen) before further isolation using a Qiagen RNeasy kit. Further preparation of RNA and RT-PCR was performed as described previously (Sastalla et al., 2009). To amplify targeted mRNAs, the following oligonucleotides were used: primers BA1995fw in combination with BA1995rv, primers Sph1 and Sph2, and primers Alo1 and Alo2 (Table 1). Internal control reactions were performed with primers GyrAfw and GyrArv as described previously (Sastalla et al., 2009). Band intensities were quantified using ImageJ version 1.40 g.
supernatant for 7 days. Cells were washed and resuspended in serum-free DMEM, and 80 μl aliquots were transferred to 96-well microtitre plates. Filter-sterilized bacterial culture supernatants (25 μl) harvested from BHI-grown bacteria were added, and plates were incubated for 4 h at 37 °C and 5 % CO₂. For some experiments, supernatants were heat-treated at 95 °C for 10 min before addition to cells. To determine the viability of cells, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT; Sigma) was added at a final concentration of 1 mg ml⁻¹, and pictures of the stained (viable) cells were taken after incubation for 45 min at 37 °Ci n5%C O2. The amount of formazan produced by viable cells was quantified at 590 nm after removal of medium and addition of 50 μl dissolving buffer containing 0.5 % SDS and 25 mM HCl dissolved in 90 % 2-propanol.

Intraperitoneal treatment of mice and isolation of cells. DBA/2J mice (8–14 weeks old, n=2 in at least three independent studies) received intraperitoneal injections of 500 μl filter-sterilized bacterial culture supernatants or BHI alone. After an incubation period of 45 min, intraperitoneal lavages were performed with PBS containing 1 % BSA (PBSA). For some experiments, digital images of lavage samples were taken, or samples were centrifuged, treated twice with ammonium chloride lysing buffer (ACK; Invitrogen) to eliminate red blood cells, washed two more times with PBSA, and resuspended in 600 μl of the same buffer. Animal experiments described in this and the previous section were performed according to NIH and Animal Welfare Act guidelines and approved by the Animal Care and Use Committee of the National Institute of Allergy and Infectious Diseases, National Institutes of Health.

Flow cytometry analyses. Cell suspensions in PBSA were incubated with rat anti-mouse CD32 antibodies (BD Biosciences) for blocking of the Fc receptor, followed by allophycocyanin (APC)-conjugated anti-mouse F4/80 (AbD Serotec) to label macrophages. After incubation for 30 min at 4 °C, cells were washed with PBSA and subjected to flow cytometric analysis using an LSR II flow cytometer (BD Biosciences). Data were analysed either in DIVA software (BD Biosciences) or in FlowJo version 7 (Tree Star).

Statistics. A t test or one-way ANOVA was performed for unpaired data (sporulation efficacy and macrophage retrieval from mice), and a paired t test was performed for paired data (cholesterol inhibition and in vitro treatment of macrophages with heat-inactivated/non-heated supernatant) in GraphPad Prism version 5.01.

Table 2. Proteins identified by 2D gel electrophoresis

| Gene ID | Description                             | Loc.* | Mᵣ  | pI  |
|---------|-----------------------------------------|-------|-----|-----|
| BA0885  | S-layer protein SAP                     | CW    | 86.6| 7.6 |
| BA0887  | S-layer protein EA1                     | CW    | 91.4| 5.8 |
| BA0796  | Conserved hypothetical protein          | E     | 40.8| 9.6 |
| BA1449  | Peptidase, M23/M37 family               | E     | 46.2| 6.7 |
| BA1952  | NLP/P60 family protein                  | CW    | 43.9| 10.2|
| B. anthracis pFP12 |                           |       |     |     |
| BA0267  | GroEL, 60 kDa chaperonin†               | C     | 57.4| 4.5 |
| BA0555  | Putative collagenase†                   | E     | 109.9| 5.0 |
| BA0620  | Putative 8-amino-7-oxononanoate synthase| C     | 43.1| 5.7 |
| BA0670  | Putative transaldolase                  | C     | 23.1| 5.3 |
| BA0672  | InhA metalloprotease††                  | E     | 87.9| 5.7 |
| BA0677  | Phospholipase C††                       | E     | 32.4| 7.9 |
| BA1887  | Enterotoxin††                           | E     | 43.6| 4.8 |
| BA1995  | Putative protease††                     | E     | 43.9| 9.0 |
| BA2032  | Hypothetical protein‡                   | E     | 16.2| 8.9 |
| BA3162  | Putative S⁻ nucleotidase†               | C     | 57.8| 6.5 |
| BA3891  | Phosphatidylinositol diacglycerol-lyase††| E     | 38.1| 7.9 |
| BA3962  | Ribosome recycling factor               | C     | 20.7| 5.5 |
| BA3964  | Translation elongation factor Ts        | C     | 32.4| 5.0 |
| BA4181  | Pyruvate dehydrogenase complex E3 component, dihydrolipoamide dehydratase† | C | 49.5| 5.0 |
| BA4184  | Pyruvate dehydrogenase complex E1 component, alpha subunit | C | 41.4| 5.6 |
| BA4194  | Putative 2,3,4,5-tetrahydropyridine-2-carboxyate N-succinyltransferase | C | 25.7| 5.2 |
| BA4499  | Superoxide dismutase, Mn†               | E     | 22.7| 5.3 |
| BA5130  | Phosphoglucone isomerase                | C     | 50.3| 4.8 |
| BA5364  | Enolase†                               | C     | 46.4| 4.4 |
| BA5470  | Sulfatase                              | MA    | 73.5| 6.4 |
| BA5580  | Fructose-bisphosphate aldolase, class II| C     | 30.7| 4.8 |

*Cellular location: C, cytoplasmic/unknown; CW, cell wall; MA, membrane-associated; E, extracellular.
†PlcR box present.
‡PlcR box present.
RESULTS

AtxA regulation does not interfere with sporulation in B. anthracis expressing PlcR–PapR

We readdressed the hypothesis that expression of functional PlcR is incompatible with pXO1-encoded, AtxA-controlled gene regulation. Expression of the PlcR–PapR fusion protein encoded by plasmid pFP12 (Pomerantsev et al., 2004) in B. anthracis strain Sterne 34F2 resulted in colonies that were highly haemolytic on blood agar plates (Fig. 1a). Microscopic analyses of the parental and PlcR–PapR-positive (Sterne pFP12) strains surprisingly showed that both strains readily formed spores (Fig. 1b).

Quantification of sporulation efficacy verified that both strains sporulated with similar high efficacies (Fig. 1c).

Growth of B. anthracis at elevated temperatures (above 43 °C) can induce loss of the virulence plasmid pXO1 (Mikesell et al., 1983). To exclude the possibility that pXO1 was lost during growth, we verified the presence of pXO1 by PCR using oligonucleotides specific for the PA gene (pagA) (data not shown).

To determine whether a point mutation in the atxA gene could have caused inactivation of the regulator, we sequenced the entire atxA gene in Sterne 34F2 and Sterne pFP12. The sequences were 100% identical to the annotated Sterne atxA sequence (data not shown), including 470 bp of the upstream promoter region, thereby excluding the possibility that AtxA expression in the PlcR–PapR-positive strain was silenced by mutation.

To rule out the possibility that other unrecognized changes may have inactivated AtxA regulation, we investigated the functionality of AtxA in both the wild-type and pFP12-containing strains. It has been established that expression of B. anthracis toxin genes is enhanced by elevated bicarbonate/CO₂ (Leppla, 1988). This higher expression requires AtxA, even though transcription of the regulator itself is not affected by bicarbonate/CO₂ levels (Dai & Koehler, 1997). Analysis of PA and LF expression in Sterne and Sterne pFP12 grown in the presence of bicarbonate/CO₂ showed that both toxins are expressed at similar levels in both strains, verifying that AtxA is functional in both strains (Fig. 1d). These results show that strain Sterne 34F2 is able to sporulate when PlcR–PapR and AtxA are simultaneously active, and that there is no incompatibility of these two regulators, at least in the strains we investigated here.

Activation of the PlcR regulon in B. anthracis leads to expression of proteins found in the secretome of B. cereus

While the secretomes of B. cereus and B. thuringiensis are highly similar, B. anthracis shows a distinctively different secretome, probably due to the presence of an inactive PlcR (Chitlaru et al., 2006; Gohar et al., 2002). Thus, we assessed the influence of heterologous PlcR–PapR activation in B. anthracis on the composition of the bacterial secretome by 2D SDS-PAGE. Fig. 2 shows that the PlcR-expressing B. anthracis derivative has a secreted protein profile distinctively different from that of wild-type B. anthracis. We selected proteins that appeared unique for MS analysis (Fig. 2, Table 2). We found S-layer proteins SAP/EA1 to be highly abundant in B. anthracis lacking PlcR–PapR, indicating either lower expression or degradation by PlcR-regulated proteases. We identified several proteins found in supernatants of B. cereus, including homologues of an enterotoxin component (BA1887), phospholipases (BA3891 and BA0677), InhA metalloprotease (BA0672) and a putative protease (BA1995). A protein of unknown function.

Fig. 1. Characterization of PlcR–PapR-expressing strains. (a) Haemolysis on sheep blood agar by Sterne strain and Sterne expressing PlcR–PapR. (b) Microscopic analysis of spores formed by Sterne and the isogenic PlcR–PapR-expressing strain. (c) Efficacy of sporulation. Shown are mean ± SD from one representative experiment out of three. (d) Western blot analysis of AtxA activity by determining PA and LF expression in response to bicarbonate/CO₂. Control (CTR) bands represent 1 µg recombinant protein.
BA2032) showed high expression in PlcR-activated B. anthracis, and interestingly the gene is present in all B. anthracis strains sequenced to date, but in only four of the sequenced B. cereus strains (W-8, AH820, E33L and 95/8201). Analysis of the upstream regions of all identified proteins for a putative PlcR binding site showed that five of them indeed harboured a PlcR box (Table 2). These results demonstrate the effect that activation of the PlcR regulon has on the expression profile of B. anthracis, and show that expression of a PlcR–PapR fusion protein causes the secretome of B. anthracis to acquire similarities to that of B. cereus.

Expression of the putative protease BA1995 is enhanced by PlcR–PapR

One difference between the two B. anthracis secretomes examined in Fig. 2 was a protein band of 40–50 kDa that was highly abundant only in the PlcR–PapR-positive strain (Fig. 2). MS identified this as the putative protease BA1995, and matched peptides from the analysis are shown in Fig. 3(a). This protein belongs to the transglutaminase-like superfamily (PFAM 01841) and a homologue is present in B. cereus (BC1991). PlcR box homology searches had failed to identify BA1995 and BC1991 as likely to be regulated by PlcR, but our data suggested that BA1995 production might be either directly or indirectly regulated by PlcR. Quantification of the BA1995 gene transcript in these two strains relative to the housekeeping gene gyrA showed the amount of transcript for BA1995 and BC1991 as likely to be regulated by PlcR, but examination of the region upstream of the BA1995 gene identified a near-perfect PlcR box 82 bases upstream of the start codon (Fig. 3c). The sequence differed from the consensus sequence by a residue that is generally conserved. The region upstream of the B. cereus BC1991 gene showed the same deviation from the consensus, which might explain why previous sequence searches failed to identify this gene as being PlcR-dependent.

In B. anthracis, transcription of the cytolysins sphingomyelinase and anthrolysin is induced by PlcR–PapR

The genome of B. anthracis encodes fewer active haemolysins than that of B. cereus. The genes for sphingomyelinase (sph) and anthrolysin (alo) are intact, whereas the homologues of B. cereus haemolysins II (hlyII) and III (hlyIII) are truncated (Klichko et al., 2003). Phospholipase C (PC-PLC; BA0677), which we were able to detect in culture supernatants of B. anthracis having an activated PlcR regulon (Fig. 2), is non-haemolytic on its own (Beecher & Wong, 2000). In B. cereus, the sph and clo genes are regulated by PlcR (Gohar et al., 2008; Lereclus et al., 1996; Pomerantsev et al., 2004; Slamti et al., 2004), and we confirmed that expression of PlcR–PapR in B. anthracis results in a haemolytic phenotype. To determine the contribution of each haemolysin, we measured mRNA levels at different time points during growth. In the absence of PlcR, only weak expression of sphingomyelinase and anthrolysin was observed (Fig. 4a). However, constitutive expression of the PlcR–PapR fusion protein greatly enhanced transcription of both haemolysins in the wild-type Sterne strain (Fig. 4a). These results show that both haemolysins are transcribed in the
PlcR–PapR-positive *B. anthracis*, and only weakly in the wild-type strain.

**PlcR–PapR-expressing *B. anthracis* lyses human red blood cells, with anthrolysin being the main contributor**

Although *B. anthracis* is considered non-haemolytic (Burdon, 1956), it has been reported that growth in rich medium causes anthrolysin production that leads to haemolysis (Shannon et al., 2003). To further evaluate the contribution of anthrolysin to haemolysis, we generated an anthrolysin gene knockout mutant of the Sterne strain and investigated the haemolytic behaviour towards human erythrocytes before and after introduction of pFP12. Supernatants of the pFP12-containing strain were highly haemolytic throughout the growth cycle (Fig. 4b). Supernatants of the PlcR–PapR-negative wild-type strain

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**Fig. 3.** Identification and analysis of the PlcR-dependent protease BA1995. (a) Putative protease BA1995 identified by 2D SDS-PAGE in supernatants of Sterne pFP12. Matched peptides are shown in bold type. (b) Semiquantitative RT-PCR of BA1995 transcription in response to PlcR–PapR expression (lower band). The gene for gyrase A (*gyrA*) served as internal control. Fold change indicates *gyrA*-normalized differences in BA1995 transcript between Sterne and Sterne pFP12. (c) Sequence comparisons of the PlcR box present in the promoter region of BA1995 of *B. anthracis* strain Ames and the homologue BC1991 in *B. cereus*. The PlcR consensus sequence is indicated. The nucleotide differing from the consensus is shown in bold type, nucleotides differing between *B. cereus* and *B. anthracis* are shaded, and putative −10, −35 regions and the start codon (ATG) are indicated.

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**Fig. 4.** Transcription and expression of haemolysins in PlcR–PapR-expressing *B. anthracis*. (a) Semiquantitative RT-PCR of *B. anthracis* putative haemolysins anthrolysin (*alo*) and sphingomyelinase (*sph*) in response to PlcR–PapR expression in Sterne and Sterne pFP12. Transcript levels of bacteria harvested at different growth phases were compared with those for gyrase A (*gyrA*), exp., exponential; stat., stationary. (b) Haemolytic activity (as a percentage) towards human erythrocytes of bacterial supernatants derived from Sterne or anthrolysin knockout mutant (*Alo*) in the presence or absence of PlcR–PapR. Supernatants were harvested at different OD$_{600}$. Shown are mean ± SD from one representative experiment out of three. ○, Sterne; ■, Sterne pFP12; ▲, Alo; ▼, Alo pFP12. (c) Haemolytic activity (%) of bacterial supernatants in the presence or absence of 25 μM cholesterol. Shown are mean ± SD from one representative experiment out of three.
showed a much lower degree of haemolysis that was observed only after the onset of stationary phase. In contrast, the anthrolysin knockout strain did not show haemolysis at any point in the growth cycle, and expression of PlcR–PapR in the anthrolysin knockout led to only a very low level of haemolysis (1.9% ± 0.05). Complementation of the knockout by expressing alo on a plasmid under the control of the PA promoter restored haemolysis to the level seen for the Sterne pFP12 transformant (data not shown).

Anthrolysin is a cholesterol binding haemolysin, and addition of cholesterol inhibits its activity (Ttweten, 2005). Cholesterol strongly blocked haemolysis by both wild-type Sterne and the pFP12-containing strain (Fig. 4c). No haemolysis was observed for the anthrolysin knockout, and expression of PlcR–PapR in this mutant could not restore the high haemolytic phenotype observed for Sterne pFP12 in the absence of cholesterol. The small residual haemolytic activities of about 3% seen for both the PlcR–PapR-positive anthrolysin knockout and the PlcR–PapR-positive anthrolysin-containing strain show that other PlcR-dependent haemolysins contribute only slightly. In summary, these results demonstrate that anthrolysin is the main contributor to haemolysis of human erythrocytes by PlcR–PapR-positive and -negative B. anthracis and that other PlcR-regulated haemolysins, such as sphingomyelinase, have a negligible effect.

Supernatants of PlcR–PapR-positive B. anthracis kill BMDMs in an anthrolysin-independent manner

In vitro, recombinant anthrolysin lyses a variety of cells, including human leukocytes, monocytes and lymphocytes (Mosser & Rest, 2006). We investigated whether anthrolysin contributes to macrophage toxicity, and for comparison included culture supernatants derived from a B. cereus strain. To exclude the known cytotoxic effects caused by lethal toxin (LT) on certain macrophages, BMDMs from LT-resistant DBA mice (McAllister et al., 2003) were used. Macrophages treated with B. cereus culture supernatants were efficiently lysed, as seen by the lack of purple formazan when compared with cells treated with medium alone (Fig. 5a), and surprisingly, culture supernatants derived from the PlcR–PapR-negative Sterne and anthrolysin-deficient strains were non-toxic (Fig. 5a, b). Treatment of BMDMs with culture supernatants from pFP12-containing bacteria, however, resulted in death of about 50% of macrophages (Fig. 5b). Heat treatment of supernatants prior to incubation with BMDMs abolished the cytotoxic effect (Fig. 5b). These data show that PlcR-dependent proteins other than LT and anthrolysin are responsible for the observed killing.

Intraperitoneal toxicity of PlcR–PapR-positive B. anthracis is not dependent on anthrolysin

Since the PlcR–PapR-positive B. anthracis supernatants efficiently killed murine macrophages in an in vitro setting, we hypothesized that similar cytotoxicity might occur in vivo. Mice having LT-resistant macrophages were injected intraperitoneally with cell-free culture supernatants derived from various strains. After 45 min, peritoneal contents were collected by lavage. Interestingly, in mice that received injections of supernatants from PlcR–PapR-positive bacteria, we observed intraperitoneal bleeding, as indicated by the presence of erythrocytes (Fig. 6a). This haemorrhaging was highly reproducible and was never detected following injection of supernatants derived from PlcR–PapR-negative bacteria. We also observed this bleeding when B. cereus-derived supernatants were injected (results not shown).

The specific cell populations present in the peritoneal washes were assessed by flow cytometry. Two cell populations were missing in the peritoneal washes derived from mice that received injections of supernatants from PlcR–PapR-positive, but not from PlcR–PapR-negative, strains (Fig. 6b). Closer characterization of cell populations using an antibody that binds to the macrophage-specific F4/80 receptor identified the two missing populations as macrophages (Fig. 6c). Quantification of F4/80-positive cells showed a significant (P<0.0001, one-way ANOVA and P<0.05 Tukey test) fivefold reduction in the numbers of macrophages retrieved from mice that received pFP12-containing B. anthracis culture supernatants (Fig. 6d). In concurrence with in vitro results, this reduction does not appear to depend on anthrolysin, since macrophages were also depleted when the anthrolysin gene was absent (Fig. 6b–d). The toxicity appears to depend on one or more proteins, since heat treatment of supernatants prior to injection completely abolished cytotoxicity (results not shown).

DISCUSSION

Besides the presence of the two large virulence plasmids, truncation of the major regulator PlcR is one of the major differences setting B. anthracis apart from other members of the B. cereus group (Kolsto et al., 2009). The nonsense mutation leading to truncation of the PlcR protein is identical in all B. anthracis isolates so far investigated, thereby strongly suggesting that it is a distinctive and defining trait of this species (Slamti et al., 2004). Inactivation of PlcR has a major effect on protein expression (Gohar et al., 2002) in B. cereus; however, little is known about the PlcR regulon in B. anthracis, including the identity of genes that retain the potential to be activated by PlcR. To elucidate the role of PlcR in B. anthracis, we expressed an active PlcR–PapR hybrid protein from a plasmid (Pomerantssev et al., 2004), thereby activating the latent PlcR regulon. An earlier study suggested that expression of PlcR in B. anthracis leads to interference with the pXO1-dependent AtxA regulon, as indicated by the inability of bacteria harbouring both regulators to sporulate (Mignot et al., 2001). Surprisingly, the B. anthracis strain containing both regulators that we describe here readily formed spores. This finding deserves further study in
strains having the PlcR–PapR regulon activated in a manner different from that of the fusion protein encoded by pFP12.

We also compared the secretomes of PlcR–PapR-expressing strains with those of their PlcR–PapR-negative counterparts. The secretomes of *B. anthracis* having active or inactive PlcR were distinctively different, and in the former we identified five proteins harbouring a PlcR binding site in their promoter region. It is likely that analysis of all protein spots present in the secretome of PlcR-active *B. anthracis* would have identified even more such proteins. Furthermore, we detected many proteins in the secretome of PlcR-expressing *B. anthracis* that are found in culture supernatants of PlcR-active *B. cereus* (Gohar et al., 2002, 2005). For example, the *B. cereus* homologue of the putative protease BA1995 was present in supernatants of PlcR-expressing *B. anthracis*, but absent in supernatants derived from the PlcR-negative parent strain. The database annotations available for BA1995 note similarities to a transglutaminase-like protease (Makarova et al., 1999), although the function of the protein is unknown. Our hypothesis that this protein is dependent on PlcR was confirmed by semiquantitative RT-PCR. Closer analysis of the BA1995 promoter region revealed a PlcR box having one mismatch at position 11, resulting in the exchange of a highly conserved nucleotide. The protease InhA2 of *B. thuringiensis* also shows an aberrant PlcR consensus sequence (Fedhila et al., 2003), and the effect of base exchanges within the PlcR box on regulator-dependent activity has been investigated. It was found that mutations within the PlcR consensus lead to lower transcription depending on their location (Gohar et al., 2008). However, the substitution that we identified in the BA1995 PlcR box was not included in the earlier study. Our results showing that BA1995 is still transcribed at a low level in the absence of PlcR imply that changes in the more conserved residues lower the specificity of PlcR for its target sequence, as shown by Gohar et al. (2008). Additionally, the ability of PlcR to recognize an aberrant recognition sequence suggests that the PlcR regulon is larger than anticipated.

One striking phenotype of bacteria harbouring a functional PlcR is their haemolytic behaviour on blood agar plates. PlcR–PapR-expressing bacteria were clearly haemolytic, and anthrolysin, the best-characterized haemolysin of *B. anthracis* (Mosser & Rest, 2006; Shannon et al., 2003), was found to be the main contributor, causing lysis of human erythrocytes in PlcR-positive and -negative bacteria. Although sphingomyelinase is transcribed at a low level, its contribution to haemolysis of human erythrocytes appears to be negligible. However, the content of sphingomyelin in the membrane of human red blood cells is low when compared with other species. In sheep, sphingomyelin can make up to 50% of the phospholipids present in the red blood cell membrane (Ikezawa et al., 1980); therefore, for detection of sphingomyelinase activity, erythrocytes of other species are expected to be a better indicator.

Macrophages are key players during anthrax pathogenesis, functioning as vehicles for transport of phagocytosed...
spores to peripheral lymph nodes. Mosser & Rest (2006) have shown that recombinant anthrolysin and supernatants derived from *B. anthracis* are able to kill a variety of cells, including macrophages derived from peripheral human blood. Surprisingly, we found no toxic response in macrophages treated with supernatants from *B. anthracis* Sterne. However, the final concentration of 25% supernatant in our cell assay may be lower than the concentrations used by Mosser & Rest (2006). In contrast, macrophages treated *in vitro* with culture supernatants derived from *B. anthracis* Sterne. However, the final concentration of 25% supernatant in our cell assay may be lower than the concentrations used by Mosser & Rest (2006). In contrast, macrophages treated *in vivo* with culture supernatants derived from *B. cereus* expressing PlcR–PapR were killed, and deletion of anthrolysin expression did not have an effect, indicating that toxin(s) other than anthrolysin were responsible for the killing. For *B. cereus*, the PlcR-dependent expression of multiple toxins has been described, including the non-haemolytic enterotoxin (Nhe), which is a three-component toxin associated with food poisoning (Lund & Granum, 1996). *B. anthracis* is known to express NheA in a PlcR-independent manner (Mendelson et al., 2004), yet it is likely that activation of PlcR increases expression, since its promoter harbours a PlcR box. Concomitantly, we found NheA (BA1887) in the supernatant of *B. anthracis* expressing PlcR. Many other toxins and virulence factors, including proteases and phospholipases, are dependent on PlcR (Gohar et al., 2002), and all of them could contribute to toxicity towards macrophages. The specificity with which the macrophage population was targeted in our *in vivo* studies was surprising, indicating a crucial role for these cells. PlcR is truncated in all known *B. anthracis* strains to date, and silencing of these unknown toxin(s) that mediate macrophage killing could be the reason why mutation of PlcR was favoured in these strains. The exact mechanism by which PlcR-reconstituted *B. anthracis* attacks macrophages and the identity of the toxin(s) involved deserve further investigation, especially in strains harbouring both virulence plasmids.

Fig. 6. *In vivo* macrophage toxicity of proteins secreted by Bacillus. (a) Intraperitoneal (IP) washes of mice (*n*=2) treated with 500 μl filter-sterilized bacterial culture supernatants derived from *B. anthracis* or *B. anthracis* anthrolysin knockout strain in the presence or absence of PlcR–PapR. Control animals were treated with BHI broth alone. (b) Flow cytometry analysis showing forward and side scatter of IP washes derived from mice treated with cell-free bacterial culture supernatants. Arrows indicate macrophage (F4/80+) populations lacking in washes treated with supernatants derived from PlcR–PapR-expressing bacteria. (c) Chart of macrophage (F4/80+) populations present in IP washes of mice treated with bacterial supernatants or with BHI alone. (d) Quantification of macrophages (%) present in the IP washes of mice treated with bacterial culture supernatants. Percentages represent the number of F4/80+ cells relative to the entire cell population present in the IP wash. Shown are mean±sd from one representative experiment out of three.
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