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Regulation of *Burkholderia cenocepacia* biofilm formation by RpoN and the c-di-GMP effector BerB

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Abstract
Knowledge about the molecular mechanisms that are involved in the regulation of biofilm formation is essential for the development of biofilm-control measures. It is well established that the nucleotide second messenger cyclic diguanosine monophosphate (c-di-GMP) is a positive regulator of biofilm formation in many bacteria, but more knowledge about c-di-GMP effectors is needed. We provide evidence that c-di-GMP, the alternative sigma factor RpoN (σ54), and the enhancer-binding protein BerB play a role in biofilm formation of *Burkholderia cenocepacia* by regulating the production of a biofilm-stabilizing exopolysaccharide. Our findings suggest that BerB binds c-di-GMP, and activates RpoN-dependent transcription of the *berA* gene coding for a c-di-GMP-responsive transcriptional regulator. An increased level of the BerA protein in turn induces the production of biofilm-stabilizing exopolysaccharide in response to high c-di-GMP levels. Our findings imply that the production of biofilm exopolysaccharide in *B. cenocepacia* is regulated through a cascade involving two consecutive transcription events that are both activated by c-di-GMP. This type of regulation may allow tight control of the expenditure of cellular resources.

KEYWORDS
Biofilm, c-di-GMP, rpoN, berB, Burkholderia cenocepacia

1 | INTRODUCTION

*Burkholderia cenocepacia*, a member of the *Burkholderia cepacia* complex (Bcc), is an opportunistic pathogen causing life-threatening infections in immune compromised individuals and in patients with cystic fibrosis (Chiarini, Bevinino, Dalmastari, Tabacchioni, & Visca, 2006; Mahenthiralingam, Urban, & Goldberg, 2005). Biofilm formation is a virulence trait of Bcc strains, and has been associated with the persistence of the infection and an increased tolerance to antibiotics compared with planktonic cells (Caraher, Reynolds, Murphy, McClean,
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2.1 | Strains, plasmids, and growth conditions

The bacterial strains and plasmids used in this study are listed in Table 1. All *B. cenocepacia* and *Escherichia coli* strains were grown at 37°C. Luria broth (LB) medium was used for overnight batch cultivation of all bacteria unless otherwise stated. Solid media were prepared with 2% (w/v) agar. A quantity of 80 μg tetracycline (Tet) mL⁻¹ (liquid medium), 120 μg Tet mL⁻¹ (solid medium), 25 μg gentamicin-sulfate (Gm) mL⁻¹, 100 μg kanamycin-sulfate (Km) mL⁻¹, and 100 μg trimethoprim (Tp) mL⁻¹ were used for *B. cenocepacia* strains, and 20 μg Tet mL⁻¹, 10 μg Gm mL⁻¹, 50 μg Km mL⁻¹, 50 μg Tp mL⁻¹, 100 μg ampicillin (Amp) mL⁻¹, and 25 μg chloramphenicol (Cm) mL⁻¹ were used for *E. coli* strains where appropriate.

2.2 | Standard molecular methods

Basic molecular and microbiological techniques were according to standard protocols (Sambrook & Russel, 2001). Genomic DNA was isolated using the DNeasy Blood and Tissue Kit (Qiagen), the plasmids were isolated using the QiAprep Spin Miniprep Kit (Qiagen), and the PCR fragments were purified and DNA was extracted from agarose gels using Wizard SV Gel and PCR Clean-Up System (Promega). Triparental matings from *E. coli* to *B. cenocepacia* were carried out as described previously (Fazli et al., 2011).

2.3 | Generation of deletion mutants

The deletion mutants investigated in the current study were generated using the method described previously (Fazli, Harrison, Gambino, Givskov, & Tolker-Nielsen, 2015). The primer sequences used in the procedure are available upon request.

2.4 | Phenotypic characterization and biofilm formation assays

Colony morphology on solid agar medium, pellicle formation at the air–liquid interface of standing liquid cultures, and flow-cell biofilm formation assays with *B. cenocepacia* strains were carried out as described previously (Fazli et al., 2013).

2.5 | Protein production and purification

For production and purification of RpoN, BerA, and BerB proteins, the corresponding genes were fused to the glutathione S-transferase (GST) gene. The genes were PCR amplified and the rpoN PCR product was cut with EcoRI and NotI, and the berA and berB PCR products were cut with BamHI and EcoRI and cloned into the EcoRI/NotI or BamHI/EcoRI digested plasmid pGEX-6P-2, respectively. The GST-fusion constructs were transformed into *E. coli* Rosetta (DE3) pLysS cells for protein expression and purification. The *E. coli* strains containing the GST-constructs were grown in 1 L of LB at 37°C, and at an OD₆₀₀ of 0.8, the cultures were transferred to an 18°C water bath, and expression was induced with 0.1 mmol/L IPTG for 16 hr. The cells were pelleted by centrifugation at 3,000 g for 20 min, solubilized in 40 ml lysis buffer (50 mmol/L HEPES pH 7.5, 150 mmol/L NaCl, 10% glycerol, 5 mmol/L DTT, 0.5% CHAPS, 2% Triton X-100, 1× Complete Protease inhibitor cocktail [Roche biochemicals]), and lysed using a French press. Soluble protein lysate was cleared by centrifugation at 20,000 g for 20 min, before being passed through twice on a column with 3 ml of Glutathione-Sepharose FF (GE Healthcare). Thereafter, the resin was washed with 20 column volumes of wash buffer (50 mmol/L HEPES pH 7.5, 150 mmol/L NaCl, 10% glycerol, 1 mmol/L DTT, 0.1% CHAPS, 0.4% Triton X-100, 0.1x Complete Protease inhibitor cocktail), and 10 column volumes of elution buffer.
(50 mmol/L HEPES pH 7.5, 150 mmol/L NaCl, 10% glycerol, 1 mmol/L DTT). The bound GST-fusion proteins were then eluted with elution buffer supplemented with 15 mmol/L reduced glutathione, and fractions of 0.5 ml were collected. Fractions were analyzed by SDS-PAGE, and positive fractions were pooled, dialyzed against elution buffer without glutathione, and protein contents measured.

### Table 1  Bacterial strains and plasmids used in the study

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Relevant characteristic(s)</th>
<th>Reference or source</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B. cenocepacia H111</td>
<td>Clinical isolate from a cystic fibrosis patient</td>
<td>Carlier et al., 2014</td>
</tr>
<tr>
<td>E. coli DH5α</td>
<td>Used for standard DNA manipulations</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>E. coli Rosetta DE3 pLysS</td>
<td>Used for protein expression and purification</td>
<td>GE Life Sciences</td>
</tr>
<tr>
<td><strong>Plasmids</strong></td>
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<td></td>
</tr>
<tr>
<td>pBBR1MCS5</td>
<td>Broad-host-range cloning vector; Gm′</td>
<td>Kovach et al., 1995</td>
</tr>
<tr>
<td>pRK404A</td>
<td>Broad-host-range cloning vector; Tet’</td>
<td>Ditta et al., 1985</td>
</tr>
<tr>
<td>pJN105</td>
<td>Cloning vector; araC-PBAD cassette cloned in pBBR1MCS5; Gm′</td>
<td>Newman &amp; Fuqua, 1995</td>
</tr>
<tr>
<td>pRK600</td>
<td>Helper plasmid in matings; ori-ColE1 RK-mob′; RK2-tra′; Gm′</td>
<td>Kessler, de Lorenzo, &amp; Timmis, 1992</td>
</tr>
<tr>
<td>pmini-Tn7-kan-gfp</td>
<td>Delivery vector for mini-Tn7-kan-gfp; Kan′</td>
<td>Norris, Kang, Wilcox, &amp; Hoang, 2010</td>
</tr>
<tr>
<td>pUX-BF13</td>
<td>Mob′ ori-R6K; helper plasmid providing theTn7 transposition functions in trans; Amp′</td>
<td>Bao, Lies, Fu, &amp; Roberts, 1991</td>
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<td>pDAI-Scl-pher5</td>
<td>Cloning vector containing the I-Scl endonuclease; Tet’</td>
<td>Fazli et al., 2015</td>
</tr>
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<td>pGEX-6P-2</td>
<td>Glutathione S-transferase (GST) fusion vector; Amp′</td>
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</tr>
<tr>
<td>pDONRPEX18Tp-Scl-pher5</td>
<td>Gateway compatible gene replacement vector based on Scl and pher5; Tp′</td>
<td>Fazli et al., 2015</td>
</tr>
<tr>
<td>pDONRPEX18Gm-Scl-pher5</td>
<td>Gateway compatible gene replacement vector based on Scl and pher5; Gm′</td>
<td>Fazli et al., 2015</td>
</tr>
<tr>
<td>pYedQ</td>
<td>E. coli yedQ (yhcK) gene cloned into pRK404A; Tet′</td>
<td>Ausmees et al., 2001</td>
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<td>pYedQ2</td>
<td>E. coli yedQ (yhcK) gene cloned into pBBR1MCS5; Gm′</td>
<td>Fazli et al., 2015</td>
</tr>
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<td>pBerA</td>
<td>berA gene cloned in pBBR1MCS5; Gm′</td>
<td>This study</td>
</tr>
<tr>
<td>pBerB</td>
<td>berB gene cloned in pJN105; Gm′</td>
<td>This study</td>
</tr>
<tr>
<td>pRpoN</td>
<td>rpoN gene cloned in pJN105; Gm′</td>
<td>This study</td>
</tr>
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<td>Gene replacement vector for rpoN; Tp′</td>
<td>This study</td>
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<tr>
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<td>Gene replacement vector for berB; Gm′</td>
<td>This study</td>
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<td>pBepB: lacZ</td>
<td>Gene replacement vector used to create bepB::lacZ fusion</td>
<td>This study</td>
</tr>
<tr>
<td>pGEX-6P-2::berA</td>
<td>berA gene fused to GST; Amp′</td>
<td>This study</td>
</tr>
<tr>
<td>pGEX-6P-2::berB</td>
<td>berB gene fused to GST; Amp′</td>
<td>This study</td>
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<tr>
<td>pGEX-6P-2::rpoN</td>
<td>rpoN gene fused to GST; Amp′</td>
<td>This study</td>
</tr>
<tr>
<td>pBK-miniTn7-KmΩSm1</td>
<td>miniTn7 delivery vector, Kan′</td>
<td>Koch, Jensen, &amp; Nybroe, 2001</td>
</tr>
<tr>
<td>pTn7-berA-lacZ</td>
<td>Tn7-based berA-lacZ promoter fusion. Native rpoN-binding site. Kan′</td>
<td>This study</td>
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<tr>
<td>pTn7-berA-lacZ-rpoNdel</td>
<td>Tn7-based berA-lacZ promoter fusion. Deleted RpoN-binding site. Kan′</td>
<td>This study</td>
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<td>pTn7-berA-lacZ-rpoNm1</td>
<td>Tn7-based berA-lacZ promoter fusion. Mutated RpoN-binding site. Kan′</td>
<td>This study</td>
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</tbody>
</table>

### 2.6  c-di-GMP-binding assay

Assessment of c-di-GMP binding was carried out using the surface plasmon resonance (SPR) assay. SPR binding assays were performed using a Reichert SR7500DC instrument optical biosensor (Reichert Technologies, Depew, NY) and SAD500M sensor chips obtained from XanTec (XanTec Bioanalytics, Düsseldorf, Germany). Scrubber 2 software (Version 2.0c, 2008, BioLogic Software) was used for processing and analyzing the data. Changes in refractive index due to DMSO-dependent solvent effects were corrected by use of a calibration curve (seven solutions, 4.25–5.75% DMSO in buffer).

Biotinylated c-di-GMP was obtained as lyophilized sodium salt with a purity of >95% from Biolog (BIOLOG Life Science Institute, Bremen, Germany). The lyophilized powder was resolved in 1 ml PBS buffer (138 mmol/L NaCl, 5.1 mmol/L KCl, 10.6 mmol/L Na2HPO4, 1.8 mmol/L KH2PO4, pH 7.4) to reach a stock concentration of 1 mmol/L. For immobilization, biotinylated c-di-GMP was further
diluted to obtain a final concentration of approximately 40 μmol/L and injected for 10 min using a flow rate of 5 μl/min at 18°C. After quenching by use of a biotin solution (3 μg/ml), immobilization levels of 1,100 μRIU (sensor chip 1) and 2,600 μRIU (sensor chip 2) were obtained.

The BerA protein (28 kDa) was stored in PBS buffer prior to the SPR analysis at a stock concentration of 17.5 μmol/L. The protein was diluted in PBS buffer first by factor 1:10 and for the following dilutions by factor 1:2, leading to a concentration range of 1.7 μmol/L to 3.3 nmol/L. The BerB protein (51.3 kDa) was stored in HEPES buffer at a final concentration of 39 μmol/L. The protein was diluted in HEPES buffer (10 mmol/L HEPES, 150 mmol/L NaCl, pH 7.4, 0.05% Tween (v/v)) by factor 1:2 to obtain a concentration range of 19.5 μmol/L to 1.1 nmol/L. The binding assays were performed using a constant flow rate of 25 μl/min in PBS buffer for BerA and in HEPES buffer for BerB. The protein dilutions were consecutively injected for 120 s association time and 240 s dissociation time.

2.7 Electrophoretic mobility shift assay

A 216-bp DNA probe spanning −280 to −74 base pairs relative to the translational start of the berA gene was generated using PCR. For electrophoretic mobility shift assay (EMSA) analysis, 50 ng of the DNA probe was incubated with purified protein in the presence or absence of 50 μmol/L c-di-GMP in binding buffer (50 mmol/L HEPES, pH 7.5, 150 mmol/L NaCl, 10% glycerol, 0.1% NP40, 5 mmol/L MgCl₂, 1 mmol/L DTT) in a total of 20 μl reaction at room temperature for 40 min. The proteins used were 500 μg RpoN and/or 50 and 500 μg BerB. The DNA-protein complexes were separated by native 5% PAGE in 0.5 × TBE buffer (45 mmol/L Tris-borate/1 mmol/L EDTA) in a Hoefer minigel (GE Healthcare) at 150 V for 45 min. Migration of DNA fragments was visualized by incubating the gel for 60 min in SYBR-GOLD (Thermo Fisher Scientific) diluted 1:10,000 in 0.5xTBE, followed by visualization on a GelDoc (BioRad) system.

2.8 Exonuclease III footprinting

A 5′FAM-tagged 216 bp DNA fragment spanning −280 to −74 base pairs of the berA promoter was prepared using the primers 5′FAM-AGGAGTGTCGCAGAAATGAGA and 5′-ACATTGACAGCGGTTGC. The 5′FAM-tagged 216 bp dsDNA was then gel extraction purified to homogeneity. A quantity of 5 μM BerB or 5 μmol/L bovine serum albumin (BSA) was incubated with 100 nmol/L dsDNA at room temperature for 5 min in 20 mmol/L Tris-HCl, pH 7.4, containing 1.67 mmol/L Mg²⁺ and 150 mmol/L NaCl. The footprinting assay was performed by addition of different amounts (1–10 units) of exonuclease III enzyme to the protein/dsDNA mixtures. After 20 min digestion at room temperature, quenching buffer (final concentration 20 mmol/L Tris-HCl, pH 7.4, 1 mol/L NaCl, 10 mmol/L EDTA) was added to all samples to stop exonuclease III digestion. The remnant fragments of the samples were then analyzed by 4–10% native gradient TBE gel electrophoresis or 10% denaturing urea-TBE gel electrophoresis. Based on preliminary footprinting results with 100 bp and 10 bp DNA ladders, four 5′FAM-tagged ssDNA’s (78 nt, 104 nt, 126 nt, and 146 nt) with corresponding berA promoter sequences were synthesized as the molecular markers to interpret the footprinting results with 10% denaturing Urea-TBE gels. Finally, the potential BerB binding sites were mapped to the berA promoter region.

2.9 Quantitative real-time PCR

RNA was isolated from biofilm grown B. cenocepacia strains on membrane filters placed on AB medium supplemented with 2% agar (w/v), 0.2% glucose (w/v), 0.1% casamino acids (w/v) for 48 hr using the RNeasy Protect Bacteria Mini Kit (Qiagen), and it was DNase treated using the Turbo DNA-free kit (Ambion) according to manufacturers’ instructions. cDNA synthesis and quantitative RT-PCR analysis were carried out using the qScript 1-Step SYBR green qRT-PCR kit (Quanta Biosciences) according to manufacturer’s instructions. As a control, quantitative RT-PCR was similarly applied to analyze the expression of the gyrB gene. The relative expression levels of the target genes were calculated using the threshold cycle (2−ΔΔCT) method (Livak & Schmittgen, 2001).

2.10 lacZ promoter fusion constructions and β-galactosidase activity assays

The B. cenocepacia H111 berB::lacZ reporter strain, which carries a lacZ gene together with a ribosome-binding site (RBS) downstream of the berB gene, was used to assay the expression of berB. In-frame deletions of target genes were introduced into this reporter strain using the homing endonuclease I-SceI-based method described by Flannagan, Linn, and Valvano (2008). C-di-GMP-overproducing strains were constructed by transformation with pYedQ.

Tn7-based berA-lacZ promoter fusions with different alterations of the putative RpoN-binding site were used to investigate the role of the binding site for transcriptional activity. They were constructed in the following way: Initially, the berA promoter region spanning nucleotides −689 to −41 relative to the berA translational start site was amplified using primers introducing Xhol and HindIII overhangs (sequences are available upon request). The promoter regions with alterations of the putative RpoN-binding site were amplified by SOE-PCR using primers introducing the alterations. The amplified promoter regions were subsequently cloned by Xhol/HindIII digestion and ligation into the broad-host-range lacZ-based promoter probe vector pSU11 (Malott et al., 2009). Finally, blunt-ended XbaI-digested fragments from the pSU11-based vectors encompassing the berA::lacZ fusion preceded by a transcriptional terminator were inserted into the blunt-ended NotI digested Tn7-delivery vector pBK-miniTn7-KmΩSm1. The resulting pTn7-berA-lacZ vectors were introduced into the B. cenocepacia genome through four-parental mating as previously described (Fazli et al., 2013). C-di-GMP-overproducing strains were constructed by transformation with pYedQ2.

For β-galactosidase activity assays, the strains to be tested were grown overnight in LB broth at 37°C with shaking and used...
to inoculate 20 ml of AB medium supplemented with 0.2% glucose (w/v) and 0.1% casamino acids (w/v) to an OD_{600} of 0.05. The cell cultures were incubated with shaking for 24 hr to 48 hr at 37°C. β-Galactosidase activity was measured as described by Stachel, An, Flores, and Nester (1985) with minor modifications. Briefly, 50-200 μl of the cell culture was harvested by centrifugation and resuspended in 500 μl Z-buffer. After addition of 25 μl of CHCl₃ and 25 μl of 0.05% SDS (w/v), the cell suspension was vortexed for 10 s and then incubated at 30°C for 15 min. The reaction was started by adding 200 μl of ONPG (4 mg/ml) and incubated at 30°C. The reaction was stopped by addition of 250 μl of 1 mol/L Na₂CO₃. The cell debris was removed by centrifugation and the absorbance was recorded at 420 nm and 550 nm. β-Galactosidase activity was calculated as Miller Units, using the formula Miller Units = 1000 \times (OD_{420} - (1.75 \times OD_{550})) / (time_{min} \times V_{ml} \times OD_{600}).

3 | RESULTS

3.1 | RpoN is important for biofilm formation by B. cenocepacia

We previously showed that high cellular c-di-GMP levels in B. cenocepacia, achieved through ectopic expression of the E. coli diguanylate cyclase YedQ, lead to wrinkled colony formation on solid medium (Fazli et al., 2011). This phenotype depends on the ability of bacteria to produce extracellular components, which often are important constituents of their biofilm matrix (Branda, Vik, Friedman, & Kolter, 2005; Friedman & Kolter, 2004a,b; Rainey & Travisano, 1998; Spiers, Bohannon, Gehrig, & Rainey, 2003; Spiers, Kahn, Bohannon, Travisano, & Rainey, 2002). We exploited the wrinkled colony morphology of the B. cenocepacia strain with high c-di-GMP levels, and performed a screen of a transposon mutant library for smooth colony formers to identify downstream components of the c-di-GMP signaling cascade. Three of the mutants that did not form wrinkled colonies carried a transposon insertion in the rpoN gene. Figure 1 shows the colony morphology of the wild-type B. cenocepacia, and a wild-type strain harboring the plasmid pYedQ causing high c-di-GMP levels due to overexpression of the YedQ diguanylate cyclase, and one of the rpoN transposon mutants harboring pYedQ.

The smooth colony morphology of the pYedQ-containing rpoN transposon mutants suggested that they are possibly defective in producing biofilm matrix components. To investigate the role of rpoN in B. cenocepacia biofilm formation without overexpression of a diguanylate cyclase, we cured the transposon mutants from the pYedQ plasmid and tested them in a flow-cell biofilm system. The rpoN transposon mutants were not severely affected in initial surface attachment, as they were able to colonize the glass surface and form small microcolonies. However, they were markedly impaired in biofilm maturation compared with the wild-type strain (Figure 2). We were able to restore the biofilm formation ability of the mutants to wild-type levels by complementation with an intact copy of rpoN on a plasmid (data not shown). Moreover, unlike the wild-type strain, the ΔrpoN strain did not form wrinkled colonies when transformed with the pYedQ plasmid (Figure 1).

FIGURE 1 Colony morphology on AB agar medium of the wild-type (WT) strain, the WT strain carrying pYedQ (WT/pYedQ), the representative rpoN transposon-mutant strain carrying pYedQ (rpoN TnM/pYedQ), and the ΔrpoN and ΔberB strains carrying pYedQ (ΔrpoN/pYedQ and ΔberB/pYedQ)

FIGURE 2 Confocal laser scanning microscope images (CLSM) of SYTO9-stained, 2-day-old flow-cell biofilms formed by the wild-type strain, the representative rpoN transposon- mutant strain (rpoN TnM) and the complemented rpoN TnM (rpoN TnM/pRpoN) strain. The central images show top-down shadow projection views, whereas the flanking images show vertical sections through the biofilms. The scale bars correspond to 50 μm.
3.2 | The promoter region of berA has putative RpoN-binding sites

We carried out a bioinformatics analysis using the publicly available Regulatory Sequence Analysis Tool (Medina-Rivera et al., 2015; http://prokaryotes.rsat.eu) to identify RpoN-dependent genes in B. cenocepacia H111. Based on the results obtained from the biofilm experiments, we hypothesized that RpoN might regulate production of biofilm matrix components in B. cenocepacia. Hence, we chose to analyze the upstream intergenic sequences of the genes predicted to be involved in production of exopolysaccharides, lipopolysaccharides, surface appendages, and adhesins in B. cenocepacia (Holden et al., 2009). In the analysis, we used the position-specific weight matrix built by Dombrecht, Marchal, Vanderleyden, and Michiels (2002) based on a set of 186 characterized RpoN-binding sites from 44 different bacterial species as input (Barrios, Valderrama, & Morett, 1999). We found that the berA gene is preceded by a putative RpoN-binding site, catTGCCACGTAATCGCTATT, located at −196 to −173 base pairs relative to the predicted translational start of berA. It contains the highly conserved GG and GC nucleotides positioned at the −24 and −12 elements, respectively (Barrios et al., 1999). This finding supported our hypothesis because berA encodes for a c-di-GMP-binding protein (Fazli et al., 2011), which regulates the transcription of the bep exopolysaccharide genes in a c-di-GMP-dependent manner (Fazli et al., 2013). We also identified putative −35 and −10 promoter elements located upstream of the RpoN-binding site, which can be recognized by the house keeping sigma factor RpoD. Further analysis of the upstream sequences of berA orthologs from other sequenced B. cenocepacia strains revealed that the RpoN-binding site preceding berA is conserved and aligns with the recently derived consensus sequence for the RpoN-dependent promoters in B. cenocepacia H111 (data not shown) (Lardi et al., 2015). Together these results suggest that RpoN plays a role in B. cenocepacia biofilm formation by positively regulating expression of the berA gene.

3.3 | BerB, a putative bacterial enhancer-binding protein, is important for wrinkled colony formation, and the stability of B. cenocepacia biofilms

Sequence analysis of the genomic region adjacent to the bep exopolysaccharide gene cluster and berA revealed the presence of a gene locus, I35_5193, coding for a putative RpoN-interacting transcriptional regulator. Due to its role in regulating expression of the bep genes described below, we designated the protein encoded by the I35_5193 locus BerB (Burkholderia exopolysaccharide regulator B). Such regulators, called bEBP, typically bind to the DNA sites located upstream of RpoN-dependent promoters, make contact with the DNA-bound RpoN-RNA polymerase closed complex through DNA looping and remodel the closed complex into a transcriptionally active open complex using the energy derived from ATP hydrolysis (Bush & Dixon, 2012). The analysis of the BerB amino acid sequence using NCBI’s conserved domain database revealed that BerB contains a central AAA+-type ATPase domain with the signature GAFTGA motif located at residues 208–213 and a C-terminal DNA-binding domain, but lacks an N-terminal regulatory-sensory input domain. The presence of the GAFTGA motif, which is almost invariant in bEBPs and is essential for their interaction with RpoN (Zhang et al., 2002), suggests that BerB is a bEBP that interacts with RpoN. A map of the berA/berB region on the B. cenocepacia genome is shown in Figure S1.

We hypothesized that BerB together with RpoN may have a role in regulating expression of berA in B. cenocepacia. Accordingly, we generated a ΔberB deletion mutant, transformed the ΔberB strain with the pYedQ plasmid, and investigated the morphology of colonies formed by the ΔberB/pYedQ strain on agar medium. Unlike the pYedQ-containing wild-type strain, the pYedQ-containing ΔberB mutant strain did not form wrinkled colonies on solid medium (Figure 1). On the contrary, when we overexpressed the berB gene from a multicopy plasmid in the wild-type B. cenocepacia strain, the bacteria formed wrinkled colonies on solid medium (Figure 3). The induction of wrinkled colony formation by BerB was dependent on the presence of both the rpoN and berA gene (Figure 3). These results indicate that berB plays a role in the regulation of the bep exopolysaccharide genes in a manner dependent on RpoN, BerA, and c-di-GMP.

We also tested the ability of the ΔberB mutant to form biofilm in flow cells. Similar to what we have reported earlier for a ΔberA strain (Fazli et al., 2011), the ΔberB strain had a defect in biofilm formation, forming very fragile biofilms with increased sensitivity to SDS treatment (Figure 4). This defect was rescued when the ΔberB strain was complemented with an intact copy of berB on a plasmid (Figure 4).

3.4 | Mutations of rpoN and berB do not affect c-di-GMP levels in B. cenocepacia

One explanation for the negative effects of the rpoN and berB mutations on wrinkled colony and biofilm formation could be that the mutations affect the c-di-GMP level in the bacteria, which in turn influence cellular functions involved in biofilm formation. To test this, we measured c-di-GMP concentrations in the wild-type and mutant strains growing in biofilms or in liquid cultures. Consistent with the previously published results (Fazli et al., 2011), we observed a substantial increase in the cellular c-di-GMP levels in the pYedQ-containing wild-type strain compared with the wild-type strain under both growth conditions (Figure S2). We found that the deletion of neither rpoN nor berB substantially affected the elevated cellular c-di-GMP levels caused by ectopic expression of the diguanylate cyclase YedQ. Both the pYedQ-containing ΔrpoN and ΔberB strains had increased levels of c-di-GMP similar to the pYedQ-containing wild-type strain (Figure
ΔberB transcript levels in the ΔberA vector control strain (Figure 6a). In contrast, we saw no significant increase in 10-fold increased level of the berA proteins on solid medium (Fazli et al., 2013). In the present study, we found that overproducing c-di-GMP or BerA in the wild-type strain leads to increased transcription of the exopolysaccharide biosynthesis gene bepB (Bcam1331). In the present study, we examined the involvement of rpoN and berB on activation of bepB transcription in response to high levels of c-di-GMP. Contrary to the results obtained with the wild-type, we observed no significant increase in bepB transcript levels when c-di-GMP levels were elevated in the ΔrpoN and ΔberB strains (Figure 6a). We also measured β-galactosidase activity from a chromosomally encoded reporter in which a promoterless lacZ gene together with a ribosome-binding site was fused to the bepB gene. β-Galactosidase activity was increased when c-di-GMP levels were elevated in the wild-type strain but not in the ΔrpoN and ΔberB strains (Figure 6c). Taken together, these findings indicate that RpoN and BerB control the transcription of berA in response to high cellular c-di-GMP levels.

3.5 | RpoN and BerB promote the transcription of berA in response to high c-di-GMP levels

We previously showed that overproduction of c-di-GMP or the BerA protein in the wild-type strain causes the formation of wrinkled colonies on solid medium (Fazli et al., 2013). In the present study, we found that ΔrpoN and ΔberB strains with increased c-di-GMP levels do not form wrinkled colonies (Figure 1). In contrast, overproducing BerA from a plasmid in the ΔrpoN and ΔberB strains resulted in formation of wrinkled colonies (Figure 5). These findings suggest that BerA is crucial for wrinkled colony formation, and that the ΔrpoN and ΔberB strains with high c-di-GMP levels lack sufficient amounts of BerA protein required to form wrinkled colonies. It also indicates that RpoN and BerB exert their regulatory effect upstream of BerA. To test this further, we carried out qRT-PCR analyses and observed an induction of berA transcription in response to high c-di-GMP levels, which was dependent on the presence of both the rpoN and berB genes. The high c-di-GMP levels in the pYedQ-containing wild-type strain caused a 10-fold increased level of the berA transcripts compared with the vector control strain (Figure 6a). In contrast, we saw no significant increase in berA transcript levels in the pYedQ-containing ΔrpoN and ΔberB strains (Figure 6a). We also measured β-galactosidase activity in c-di-GMP-overproducing (pYedQ2-containing) bacteria harboring a promoterless lacZ gene fused to the berA promoter with or without mutated putative RpoN-binding sites. Unlike the native berA-lacZ fusion, the berA-lacZ fusions with a mutated or deleted RpoN-binding site were only expressed at a low level (Figure 6b).

In a previous study (Fazli et al., 2013), we showed that overproduction of c-di-GMP or BerA in the wild-type strain leads to increased transcription of the exopolysaccharide biosynthesis gene bepB (Bcam1331). In the present study, we examined the involvement of rpoN and berB on activation of bepB transcription in response to high levels of c-di-GMP. Contrary to the results obtained with the wild-type, we observed no significant increase in bepB transcript levels when c-di-GMP levels were elevated in the ΔrpoN and ΔberB strains (Figure 6a). We also measured β-galactosidase activity from a chromosomally encoded reporter in which a promoterless lacZ gene together with a ribosome-binding site was fused to the bepB gene. β-Galactosidase activity was increased when c-di-GMP levels were elevated in the wild-type strain but not in the ΔrpoN and ΔberB strains (Figure 6c). Taken together, these findings indicate that RpoN and BerB control the transcription of berA in response to high cellular c-di-GMP levels.

3.6 | BerB binds to berA promoter DNA and interacts with RpoN

Most bEBPs bind to DNA at sites typically located ~80–150 bp upstream of the RpoN-dependent promoters they control. To test if BerB directly binds to the berA promoter DNA, we carried out an electrophoretic mobility shift assay (EMSA) using purified BerB and a 216-bp DNA fragment spanning −280 to −74 base pairs relative to the translational start of berA. BerB caused a shift in the mobility of the DNA fragment in a concentration-dependent manner (Figure 7), indicating that BerB binds to the berA promoter DNA in vitro. We observed a substantial difference in berA transcript levels in the ΔberB strain compared with the wild-type strain when the cellular c-di-GMP levels were elevated (Figure 6a). Hence, we hypothesized that c-di-GMP and BerB regulate berA transcription in concert and that
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Our gene expression data demonstrated that RpoN and BerB induce berA transcription when the cellular c-di-GMP levels are high (Figure 6a). This made us hypothesize that BerB may sense the levels of c-di-GMP by directly binding to the signaling molecule. Accordingly, we investigated binding of BerB to c-di-GMP using SPR. As a positive control in our SPR assay, we included the BerA protein, which is a known c-di-GMP-binding protein (Fazli et al., 2011). In accordance with our previous results (Fazli et al., 2011), BerA was found to bind the biotinylated c-di-GMP in a concentration-dependent manner with an estimated $K_D$ value of 3 µmol/L (Figure 9). Subsequently, we tested the purified BerB protein in different concentrations for its ability to bind c-di-GMP. We observed that BerB was able to bind to the biotinylated c-di-GMP in a concentration-dependent manner with an estimated $K_D$ value of approximately 3 µmol/L (Figure 9). The binding affinity of BerB falls within the physiological affinity range of previously characterized c-di-GMP-binding proteins (Pultz et al., 2012). In addition to the SPR assay, binding of c-di-GMP to BerB was also demonstrated by the use of a differential scanning fluorimetry (DSF) assay. DSF measures the melting temperature of a protein, and exploits that the melting point of the protein may shift in the presence of small c-di-GMP might have an effect on the DNA-binding ability of BerB. However, the addition of c-di-GMP to the reaction mixtures did not change the DNA-binding activity of BerB in our assay (Figure 7). The addition of RpoN to the reaction mixtures together with BerB shifted the mobility of the berA promoter DNA more than did BerB alone (Figure 7). We also tested the ability of RpoN to bind to the berA promoter DNA without BerB. In the absence of BerB, RpoN did not cause a detectable shift in the mobility of the berA promoter DNA fragment (Figure 7). Together, these results indicate that BerB and RpoN physically interact on the berA promoter DNA. Binding of RpoN to the berA promoter DNA in the absence of BerB may require the presence of the RNA polymerase. We subsequently investigated binding of BerB to the berA promoter region by the use of an exonuclease III footprinting assay (Figure 8). Unlike BSA, BerB was found to protect distinct sequences in the berA promoter region. The use of ad hoc synthesized FAM-tagged ssDNA fragments allowed us to determine the size of the protected DNA fragments, and map putative BerB binding sites on the promoter DNA as shown in Figure 8.
interacting ligands (Niesen, Berglund, & Vedadi, 2007). In the presence of increasing concentrations of c-di-GMP, we observed a shift in the melting temperature of BerB beginning at 1 μmol/L c-di-GMP, and increasing up to around 10 μmol/L c-di-GMP (Figure S3). These data indicate a binding of c-di-GMP to BerB with a $K_d$ value between 1 and 10 μmol/L. Together, the SPR and DSF experiments suggest that BerB is a c-di-GMP-binding protein.

4 | DISCUSSION

In contrast to other alternative sigma factors, RpoN is widely represented in the bacterial kingdom (Merrick, 1993). It was originally identified as a sigma factor required for transcription of the genes involved in nitrogen assimilation (Hirschman, Wong, Sei, Keener, & Kustu, 1985; Reitzer & Schneider, 2001). However, a great deal of evidence has now accumulated that RpoN also controls many other biological activities in bacteria, ranging from utilization of alternative carbon sources (Lundgren, Harris, Serwer, Scheel, & Nomura, 2015) and biofilm formation (Saldias et al., 2008) to nitrogen assimilation (Merrick, 1993; Reitzer & Schneider, 2001). However, a great deal of evidence has now accumulated that RpoN also controls many other biological activities in bacteria, ranging from utilization of alternative carbon sources (Lundgren, Harris, Serwer, Scheel, & Nomura, 2015) and biofilm formation (Saldias et al., 2008) to nitrogen assimilation (Merrick, 1993; Reitzer & Schneider, 2001).

It has been shown that RpoN controls flagellum-mediated motility in B. cenocepacia, which was found to be essential for biofilm formation in microtiter trays (Saldias et al., 2003; Saldias, Lamothe, Wu, & Valvano, 2008; Totten, Lara, & Lory, 1990), and biofilm formation (Saldias et al., 2008; Thompson, Webb, Rice, & Kjelleberg, 2003; Wolfe, Millikan, Campbell, & Visick, 2004). It has been shown that RpoN controls flagellum-mediated motility in B. cenocepacia, which was found to be essential for biofilm formation in microtiter trays (Saldias et al., 2003; Saldias, Lamothe, Wu, & Valvano, 2008; Totten, Lara, & Lory, 1990), and biofilm formation (Saldias et al., 2008; Thompson, Webb, Rice, & Kjelleberg, 2003; Wolfe, Millikan, Campbell, & Visick, 2004). It has been shown that RpoN controls flagellum-mediated motility in B. cenocepacia, which was found to be essential for biofilm formation in microtiter trays (Saldias et al., 2003; Saldias, Lamothe, Wu, & Valvano, 2008; Totten, Lara, & Lory, 1990), and biofilm formation (Saldias et al., 2008; Thompson, Webb, Rice, & Kjelleberg, 2003; Wolfe, Millikan, Campbell, & Visick, 2004). It has been shown that RpoN controls flagellum-mediated motility in B. cenocepacia, which was found to be essential for biofilm formation in microtiter trays (Saldias et al., 2003; Saldias, Lamothe, Wu, & Valvano, 2008; Totten, Lara, & Lory, 1990), and biofilm formation (Saldias et al., 2008; Thompson, Webb, Rice, & Kjelleberg, 2003; Wolfe, Millikan, Campbell, & Visick, 2004).

FIGURE 8  Mapping of BerB binding sites at the berA promoter region using exonuclease III footprinting. Protection of distinct DNA sequences by BerB, but not by BSA, was visualized by native gradient TBE gel electrophoresis (a). The size of the protected DNA fragments was estimated by denaturing Urea-TBE electrophoresis with synthesized ssDNA ladders (b). Thereby putative BerB binding sites at the berA promoter region could be mapped as shown (c) with these previous studies, our flow-cell experiments indicate that B. cenocepacia ΔrpoN strains have biofilm formation defects.

Here, we provide evidence that RpoN and the putative bEBP BerB regulate expression of the berA gene, coding for a c-di-GMP-responsive transcriptional regulator previously shown to activate the expression of bep exopolysaccharide biosynthesis genes in B. cenocepacia (Fazli et al., 2013). We identified a putative RpoN-binding site in the promoter region of the berA gene, and sequence analysis of the berA gene, located in the vicinity of berA, suggested that it encodes an RpoN-interacting transcriptional regulator. Accordingly, our gene expression data demonstrated that both rpoN and berA are required for the expression of berA and, by extension, bep exopolysaccharide genes under high cellular c-di-GMP conditions. Evidence has been provided that Bep exopolysaccharide is important for the stability of flow-cell grown B. cenocepacia biofilms, and for the formation of wrinkled colonies on solid medium in response to high levels of c-di-GMP (Fazli et al., 2013). Our flow-cell biofilm experiments indicated that the ΔberB strain forms biofilm with reduced stability as previously reported for the ΔberA strain (Fazli et al., 2011) and Δbep exopolysaccharide-deficient strains (Fazli et al., 2013). Moreover, contrary to the wild-type, but similar to what has previously been reported for the ΔberA (Fazli et al., 2011) and Δbep strains (Fazli et al., 2013), the ΔberB strain did not form wrinkly colonies in response to high levels of c-di-GMP.

The domain architecture of BerB and the presence of the signature GAFTGA motif, essential for interaction with RpoN, indicate that BerB is a bEBP. We provide evidence that BerB directly binds to c-di-GMP, and our data further indicate that this binding is important for c-di-GMP-mediated induction of berA transcription and exopolysaccharide production in B. cenocepacia. Several bEBPs have been shown to activate transcription of genes involved in exopolysaccharide production in other bacteria including Pseudomonas aeruginosa and Vibrio species. In Vibrio fischeri, the bEBP-type response regulator SypG controls the transcription of symbiosis polysaccharide (Syp) genes and biofilm formation in a manner dependent on RpoN (Visick, 2009; Yip, Grublesky, Hussa, & Visick, 2005). Whether SypG is able to bind to c-di-GMP is
not known. In contrast, the bEBP-type transcriptional regulator VpsR from Vibrio cholerae controls expression of Vibrio polysaccharide (Vps) genes and biofilm formation independent of RpoN (Yildiz, Daiganov, & Schoolnik, 2001). VpsR binds to c-di-GMP and in turn activates the transcription of the vpsT gene (Srivastava, Harris, & Waters, 2011), coding for a LuxR-type transcriptional regulator required for the expression of Vps biosynthesis genes (Casper-Lindley & Yildiz, 2004). In P. aeruginosa, the c-di-GMP-binding protein FleQ is an unusually
versatile bEBP. While the regulation of flagella biosynthesis genes by FleQ depends on RpoN (Dasgupta et al., 2003; Jyt, Dasgupta, & Ramphal, 2002), FleQ controls the expression of the Pel polysaccharide genes independent of RpoN in response to changing cellular c-di-GMP levels (Baraquet, Murakami, Parsek, & Harwood, 2012; Hickman & Harwood, 2008). It was initially demonstrated that FleQ binds to the pel promoter DNA and represses pel transcription in the absence of c-di-GMP. Upon c-di-GMP binding to FleQ, this inhibition is relieved and the transcription from the pel promoter is activated (Hickman & Harwood, 2008). However, it was later found that, in the presence of c-di-GMP, FleQ also functions as an activator when bound to a second site on the pel promoter DNA (Baraquet et al., 2012), which indicates that c-di-GMP converts FleQ from a repressor to an activator of exopolysaccharide gene transcription. Our data suggest that BerB activates gene transcription upon c-di-GMP binding, and unlike VpsR and FleQ, requires RpoN. BerB therefore represents the first example of a bEBP that functions together with both RpoN and c-di-GMP.

We showed that BerB binds to the berA promoter DNA in vitro. However, the addition of c-di-GMP did not change the DNA-binding activity of BerB, suggesting that binding of c-di-GMP may instead have effects on other functions of BerB such as its ATPase activity. Recent studies have suggested that c-di-GMP binds to AAA+ ATPase-domain containing proteins from diverse bacteria and, except for only one case, inhibits their ATPase activity (Baraquet & Harwood, 2013; Roelofs et al., 2015; Trampari et al., 2015). For example, c-di-GMP competes with ATP to bind to the ATP-binding site on FleQ and inhibits FleQ’s ATPase activity (Baraquet & Harwood, 2013), but in the case of MshE from V. cholerae, binding of c-di-GMP to the N-terminal domain of MshE enhances its ATPase activity (Roelofs et al., 2015). In our case, BerB appears to activate RpoN-dependent transcription of the berA gene. Hence, if c-di-GMP binding has an effect on BerB’s ATPase activity, then it should be stimulation rather than inhibition, possibly by affecting the oligomerization of the AAA+ domain on BerB, which is essential for the ability of bEBPs to hydrolyze ATP to drive the RpoN-RNA polymerase closed complex into a transcriptionally active open complex. Alternatively, c-di-GMP may affect BerB’s interaction with RpoN.

Exopolysaccharides are a major component of the biofilm matrix with importance for the mechanical stability of biofilms. However, their biosynthesis is an energy-intensive process and therefore requires tight regulation. The requirement of an activator protein by the RpoN-RNA polymerase closed complex to initiate transcription of genes related to exopolysaccharide biosynthesis, such as the regulation of berA and by extension bep genes described here, allows tight control of gene expression in the off-state until the activating signal is present. Furthermore, our work suggests that c-di-GMP regulates Bep exopolysaccharide production in B. cenocepacia both by stimulating berA transcription and by stimulating transcriptional activation by BerA. This results in a cascade regulation of a biofilm exopolysaccharide in which two consecutive transcription events are both activated by c-di-GMP (Figure 10). Such sustained c-di-GMP-mediated regulation may allow tight control of expenditure of cellular resources. The BerB protein represents the first example of a bEBP, whose regulatory function depends on both RpoN and c-di-GMP. However, mechanistic questions relating to how c-di-GMP modulates BerB-RpoN-dependent transcription of the berA gene remain to be answered.

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CONFLICT OF INTEREST

None declared.

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