A toxin-antitoxin system associated transcription factor of *Caulobacter crescentus* can influence cell cycle-regulated gene expression during the SOS response

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Abstract

Toxin-antitoxin (TA) systems are widespread in bacterial chromosomes but their functions remain enigmatic. Although many are transcriptionally upregulated by stress conditions, it is unclear what role they play in cellular responses to stress and to what extent the role of a given TA homologue varies between different bacterial species. In this work we investigate the role of the DNA damage-inducible TA system HigBA of *Caulobacter crescentus* in the SOS response and discover that in addition to the toxin HigB affecting cell cycle gene expression through inhibition of the master regulator CtrA, HigBA possesses a transcription factor third component, HigC, which both auto-regulates the TA system and acts independently of it. Through HigC, the system exerts downstream effects on antibiotic (ciprofloxacin) resistance and cell cycle gene expression. HigB and HigC had inverse effects on cell cycle gene regulation, with HigB reducing and HigC increasing the expression of CtrA-dependent promoters. Neither HigBA nor HigC had any effect on formation of persister cells in response to ciprofloxacin. Rather, their role in the SOS response appears to be as transcriptional and post-transcriptional regulators of cell cycle-dependent gene expression, transmitting the status of the SOS response as a regulatory input into the cell cycle control network via CtrA.
Importance

Almost all bacteria respond to DNA damage by upregulating a set of genes that helps them to repair and recover from the damage, known as the SOS response. The set of genes induced during the SOS response varies between species, but frequently includes toxin-antitoxin systems. However, it is unknown what the consequence of inducing these systems is, and whether they provide any benefit to the cells. We show here that the DNA damage-induced TA system HigBA of the asymmetrically dividing bacterium *Caulobacter crescentus* affects the cell cycle regulation of this bacterium. HigBA also has a transcription factor encoded immediately downstream of it, named here HigC, which controls expression of the TA system and potentially other genes as well. Therefore, this work identifies a new role for TA systems in the DNA damage response, distinct from non-specific stress tolerance mechanisms which had been proposed previously.
Introduction

Bacterial toxin-antitoxin (TA) systems have been the subject of intensive study since their widespread prevalence in prokaryotic chromosomes, as well as mobile genetic elements, was discovered (1). TA systems consist of two components, a toxin protein which can inhibit some aspect of central cellular metabolism and an antitoxin, either protein or RNA, which can inhibit the toxin activity or production at the post-transcriptional or post-translational level, depending on the type (2-6). The best characterised systems are those of type II where both toxin and antitoxin are small proteins which form a non-toxic complex with each other, of variable stoichiometry depending on the TA system and the organism in which it is found. This complex usually also binds the promoter of the TA system through a DNA binding domain found in the antitoxin and transcriptionally represses it (7). This repressive activity can be modulated by the abundance of the toxin protein by “conditional cooperativity,” a mechanism by which low levels of toxin protein promote TA complex binding to the TA system promoter, while high levels of toxin destabilise TA complex binding resulting in derepression of transcription (8). The mechanisms of toxins are variable but fall into two broad groups: inhibition of DNA replication by inhibiting the activity of DNA gyrase, and inhibition of translation at various levels including cleavage or modification of mRNA, tRNA or rRNA, or phosphorylation of aminoacyl-tRNA synthetases or EF-Tu (9).

However, the physiological relevance of chromosomally encoded TA systems is still an open and hotly debated question. TA systems encoded on plasmids can certainly function as plasmid maintenance systems by elimination of plasmid free cells through post-segregational killing (PSK), where the antitoxin component of the TA complexes in a plasmid-free cell is degraded and not replaced, resulting in elimination of plasmid free cells from the population because they are poisoned by excess toxin (10). It has also been noted that chromosomal TA systems are associated with cryptic prophages (11), superintegrons (12) and transposons (13) and have been suggested to promote stability and/or propagation via horizontal gene transfer of these mobile elements. Other TA systems, of types III and IV, have been shown to have roles in bacteriophage resistance (14, 15). A
third highly popular hypothesis for the role of TA systems was promoting persistence to antibiotics or other stress conditions (3). However, work that purported to show that persistence in *E. coli* was mediated by activation of multiple TA systems was subsequently withdrawn, due to the discovery that the apparent persistence phenotype was due to prophage infection during mutant construction (16). Another study which showed that the type II TA system MqsRA of *E. coli* was an important mediator of the stress response (17) could not be reproduced by other groups (18). Frequently it has been assumed that the transcriptional activation of TA systems in response to various stress conditions is an indication that they are important for the corresponding stress response, and that transcriptional activation could be used as a proxy for activation of the TA system at the functional level, but it has recently been shown that during transcriptional activation of the stress-induced type II TA systems in *E. coli*, their toxins are neither released nor active (19). Hence, it is currently unclear whether type II TA systems have any involvement in persistence at all. Some type I TA systems can indeed induce persistence through their toxins which act as membrane pores and collapse the proton motive force across the membrane, thereby leading to a shutdown of cellular metabolism (20). However, other mechanisms which lead to decreased metabolic rate and slow growth can also induce persistence, independently of TA system induction or activity (21). The current key factor in induction of persistence seems to be slow growth, regardless of the cause, and no TA system has yet been reproducibly found to be necessary or sufficient for this (2).

The persistence-inducing type I TA system mentioned above (*tisAB/istR*) was seen to induce persistence in the presence of antibiotics that cause DNA damage and induce the SOS response, and the presence of type II TA systems in the SOS response regulon has also been observed in *E. coli* (22) and in *Caulobacter crescentus* (23). In these cases, the LexA repressor binds to the promoter of the TA system in addition to the antitoxin. An unusual aspect of the *Caulobacter crescentus* LexA-regulated TA system, HigBA, was that the repression provided by LexA was hierarchically superior to that of the antitoxin, allowing deletion of the antitoxin without loss of viability. Creation of a strain lacking the antitoxin allowed the identification of the targets of the mRNA-degrading toxin HigB in
Caulobacter, which surprisingly included the essential master regulator of the cell cycle, CtrA. This TA system was not transcriptionally induced by any other stress than DNA damaging antibiotics, and appeared to have a role in resistance to these antibiotics because deletion of the higB toxin gene improved viability during growth on ciprofloxacin plates, in particular for the hypersensitized ΔlexA strain (23). However, a role for HigBA in persistence to antibiotics was not explored at that time.

The cell cycle regulator CtrA of Caulobacter is responsible for the asymmetric cell cycle of this bacterium, which can be seen by the division of a predivisional cell into two genetically identical but morphologically dissimilar cell types, the stalked and swarmer cells (24, 25). CtrA is a transcription factor that regulates DNA replication by binding to the origin of replication and silencing it, as well as regulating (positively and negatively) transcription of many other genes, in concert with two other transcriptional regulators MucR and SciP (26-28). CtrA is present and abundant in swarmer cells and keeps these cells in G1 phase, where they are motile but do not replicate. In contrast, it is degraded and dephosphorylated in stalked cells to permit the onset of chromosome replication and cell division (29). The ability of the toxin HigB to cleave the ctrA mRNA suggested that this TA system could potentially influence the cell cycle during DNA damage conditions.

In the original paper which defined the SOS regulon of Caulobacter (30) it was indeed seen that higBA was transcriptionally upregulated in a ΔlexA mutant. Moreover, the gene immediately 3’ to higBA, the uncharacterised gene CCNA_03131, was also upregulated to a similar extent, suggesting that it might be associated with higBA in some way. Other type II TA systems with a third component in addition to the antitoxin have been previously characterised, where the third component is a transcription factor (11, 31), an accessory antitoxin (32) or an antitoxin-stabilising chaperone (33). We therefore aimed to characterise whether this factor was involved in regulation or function of higBA, either in terms of the SOS response or cell cycle control via CtrA, and whether it could be considered a true third component of the system. We have also explored further whether either higBA or CCNA_03131 were involved in persistence and whether the cleavage of ctrA mRNA
by HigB results in altered CtrA-dependent gene expression. Our results support a model where CCNA_03131 is a third component of the TA system and is capable of transcriptionally repressing it, but that it also acts independently of it. We find no evidence for a role of HigBA, CCNA_03131 or the LexA SOS response repressor in persistence in Caulobacter. Instead, the main role of the HigBA/CCNA_03131 system appears to be in regulation of cell cycle-dependent gene expression during the SOS response.

Materials and Methods

General growth conditions

Caulobacter crescentus strains were routinely grown in peptone-yeast extract (PYE) medium at 30°C and E. coli strains in LB at 37°C. Antibiotics were used at the following concentrations; tetracycline at 1 μg/ml for Caulobacter and 10 μg/ml for E. coli, gentamicin at 1 μg/ml for Caulobacter and 10 μg/ml for E. coli and kanamycin at 20 μg/ml (solid media) or 5 μg/ml (liquid media) for Caulobacter and 20 μg/ml for E. coli. Ciprofloxacin was prepared as 20 mg/ml stock solution in 0.1 M HCl and in all experiments involving ciprofloxacin, an appropriate volume of 0.1 M HCl was added to the control cultures or plates. Vanillate stock solution was prepared at 50 mM stock solution (adjusted to pH 8.0 with NaOH) and used at 50 or 500 μM as indicated in the figure legends.

Strain and plasmid construction

DNA fragments for cloning were PCR-amplified with Phusion DNA polymerase (New England Biolabs) from stationary phase cultures of wild type (WT) Caulobacter crescentus, using PCR primers listed in Table 1. Products were purified by agarose gel electrophoresis. Cloning of the correct region was confirmed by sequencing and plasmid stocks were maintained in E. coli EC100 or TOP10. Plasmids used in this work are listed in Table 2. Replicating plasmids (pMT335 and plac290 derivatives) were transferred into Caulobacter strains by electroporation while suicide plasmids for generation of deletion mutants (pNPTS138 derivatives) were transferred by conjugation from E. coli S17-1 λpir (34). After integration of suicide vectors by recombination through one of the homologous flanking
regions, secondary recombination events were induced by counterselection on PYE agar containing 3% sucrose and mutants carrying resulting in-frame deletions were screened for by PCR. Double mutants were made by introducing the ΔhigC or ΔhigBAC alleles into the ΔlexA mutant strain. Strain numbers and genotypes are listed in Table 3.

The overexpression plasmid for WT HigC was constructed by amplification of full length higC with primers cc3036_nde and cc3036_eco, digested with Ndel and EcoRI and ligated into correspondingly digested pMT335 to form pMT335-higC. The overexpression plasmid for truncated HigC was constructed by amplification of the C-terminal half of higC with primers cc3036_noTM_nde and cc3036_eco. The forward primer in this reaction (cc3036_noTM_nde) contains a Ndel site (CATATG) which overlaps amino acid 133 (GTG-Val) and replaces it with ATG-Met. This was digested with Ndel and EcoRI and ligated into correspondingly digested pMT335 to form pMT335-higC-noTM.

To construct the higC knockout plasmid pNPTSΔhigC, the flanking regions of higC were amplified with primer pairs 3036_up_bam and 3036_up_hind (606 bp upstream region including the first 6 amino acids of HigC) and 3036_down_bam and 3036_down_eco (556 bp downstream region including the last 6 amino acids and stop codon of HigC). These were digested with BamHI/HindIII and BamHI/EcoRI respectively and ligated simultaneously into EcoRI/HindIII-digested pNPTS138. To construct the higBAC operon knockout plasmid, the pNPTSΔhigBA plasmid backbone (23) was used. This was digested with EcoRI/BamHI to remove the fragment corresponding to the higA downstream region. The vector (including the upstream flanking region of higB) was purified by agarose gel electrophoresis and ligated to the EcoRI/BamHI-digested PCR product of 3036_down_bam and 3036_down_eco (downstream flanking region of higC) to give pNPTSΔhigBAC.

β-galactosidase assay

β-galactosidase assays were performed on strains carrying low copy plasmid-borne transcriptional fusions of the promoters of interest to lacZ. Cultures were grown to early exponential phase (OD₆₀₀ = 0.1 – 0.4) with exposure to ciprofloxacin or vanillate as described in the main text or figure legends,
followed by β-galactosidase assays using the method of Miller (35) on three independent biological replicates.

**RNA extraction and quantitative RT-PCR**

RNA was extracted from 4 ml mid-exponential phase cultures which were treated with 2500 U Ready-Lyse (Bionordika) and homogenized using QiaShredder columns, prior to RNA extraction with the RNEasy Mini Kit (Qiagen) according to the manufacturers’ instructions, including on-column DNase digestion. RNA quality was assessed by agarose gel electrophoresis and concentration was measured in a Nanodrop spectrophotometer. cDNA was prepared using the SuperScript IV Reverse Transcriptase Kit (Thermo Fisher) according to the manufacturers’ instructions, on 400 ng RNA template using random hexamer primers. Quantitative RT-PCR was performed in technical and biological triplicates on a 96-well LightCycler real-time PCR system (Roche) using SYBR Green (Roche). The higB transcript was amplified with primers higB_qrt_fwd and higB_qrt_rev, the higC transcript was amplified with primers CC_3036_qrt_fwd and CC_3036_qrt_rev, and the reference gene rpoD was amplified with primers rpoDfow and rpoDrev. Quantification was by the standard curve method and higB and higC transcript levels were normalized to rpoD.

**Quantitative PCR-chromatin immunoprecipitation (qChIP)**

Chromatin immunoprecipitation experiments followed by quantitative PCR were performed using an anti-CtrA polyclonal antibody (26) as described previously (36). Quantitative PCR was performed in technical duplicates on two biological replicates using primers pilA_chip_f and pilA_chip_r to amplify the promoter of pilA and primers sciP_chip_f and sciP_chip_r to amplify the promoter of sciP. Quantification was by the standard curve method and results are expressed as fold enrichment of a given product in the ChIP sample relative to the input DNA.

**Efficiency of plating assay**

Resistance to ciprofloxacin and chloramphenicol was assessed by dilution spot plating. Cultures of strains to be tested were grown overnight to stationary phase, then inoculated into new medium to grow to mid-exponential phase. Culture density was measured, normalised to the OD600 of the least
dense culture (OD600=0.5 or less), serially diluted in PYE to $10^{-6}$ and 5 µl spotted onto plates containing PYE medium with sub-inhibitory concentrations of ciprofloxacin (1 µg/ml) or chloramphenicol (0.02 µg/ml). For higC overexpression experiments, the plates contained gentamicin in addition to ciprofloxacin or chloramphenicol in order to maintain selection on the plasmid, and all plates contained vanillate (50 µM) to induce expression of higC. Plates were imaged after 3 days growth at 30°C. Images are representative of three independent biological replicates.

**Persistor assay**

Overnight cultures were diluted into new PYE medium and grown to mid-exponential phase (OD$_{600}$ = 0.4 – 0.6). Ciprofloxacin was added to a final concentration of 10 µg/ml and a 100 µl sample of the culture was immediately taken out for quantification of cfu/ml at zero time. Further 100 µl samples were taken at 2, 4, 6, 24 and 48 hours after ciprofloxacin addition. Immediately after sampling, cells were washed in 1 ml PYE followed by centrifugation at 8000g for 5 minutes, repeated 3 times. Washed cells were serially diluted to $10^{-6}$ and plated in technical duplicates as described in the figure legends, then incubated at 30°C for 3 days. Data are reported as fraction surviving cfu/ml relative to zero time for each time point, for three independent biological replicates.

**Statistical analysis**

All numerical data are reported as mean of all biological replicates performed and error bars indicate the standard deviation unless otherwise stated. Statistical significance was analysed by non-paired equal variance 2-tailed Student’s T test for comparisons between strains or treatment conditions. * signifies p < 0.05 and ** signifies p < 0.01 throughout.

**Results**

*The putative transcription factor CCNA_03131 (higC) is associated with and regulates the higBA toxin-antitoxin system*

We previously observed that loss of the HigB toxin in the ΔlexA background improved viability of the cells, likely because production or activation of this toxin is increased in the absence of LexA.
Unexpectedly, we did not observe this improvement in the ΔlexA ΔhigBA strain even though this also lacks HigB. Moreover, both ΔlexA ΔhigBA and ΔhigBA strains were sensitized to ciprofloxacin relative to the ΔlexA and wild type parent strains, respectively, while the ΔhigB and ΔlexA ΔhigB mutants were not (23). To investigate the reason for this difference, we first measured activity of a P_{higBA} promoter-reporter construct in wild type, ΔlexA and ΔhigBA single mutants compared to ΔlexA ΔhigBA. Although we observed the anticipated increased promoter activity in the absence of LexA or HigA repressors, the absence of both of these in the ΔlexA ΔhigBA double mutant surprisingly led to lower, rather than higher, promoter activity (Fig 1A). Investigating the genomic context of the higBA TA system, we noted that a putative transcription factor gene, CCNA_03131, lies 42 bp downstream of higBA (Fig. 1B). Bioinformatic analysis of promoter and terminator locations (using the bprom (37) and ARNold software (38, 39), respectively) failed to find any putative promoter or terminator sequences between the 5’ end of higB and the 5’ end of CCNA_03131, while the same programs identified a promoter upstream of higBA and a putative rho-independent terminator downstream of CCNA_03131, suggesting that CCNA_03131 is a member of the higBA operon and potentially a third component of this TA system.

Since this gene had been annotated as a LytTR-family transcription factor (40) based on sequence homology, and some TA systems are known to have third components that act as transcription factors (11), we investigated whether it could also regulate higBA. Overexpressing CCNA_03131 from the vanillate-inducible promoter reduced P_{higBA} activity in WT, ΔlexA and ΔhigBA strains relative to the empty vector control, with the most significant effect seen in the ΔhigBA background (Fig 1C). Hence, the product of CCNA_03131 can repress the P_{higBA} promoter, albeit weakly, and seems to have stronger repressive activity when HigA is absent. Due to the likely co-regulation of CCNA_03131 with higBA, and its ability to repress transcription from the higBA promoter, we now consider CCNA_03131 as a part of the higBA TA system operon and name it higC.

We then measured the steady-state mRNA levels for higB and higC in WT, ΔlexA, ΔhigA, ΔhigBA and ΔlexA ΔhigBA strains to confirm whether their expression levels were consistent with the promoter
activity measurements (Fig 1D). Both mRNAs were detectable, but higC appeared to be expressed at a much lower level than higB and no obvious induction of higC in the ΔlexA, ΔhigA, and ΔhigBA strains relative to WT was seen. However, higC was very strongly expressed in the ΔlexA ΔhigBA mutant, in which the in-frame deletion of higBA has placed the higC coding sequence immediately downstream of the higBA promoter, and the LexA and HigA repressors are missing. Taken together, these data show that the product of higC functions as a repressor of the higBAC promoter and is strongly overproduced in the ΔlexA ΔhigBA mutant, providing a plausible explanation for why the $P_{higBA}$ promoter-reporter activity in this strain was unexpectedly low.

The N-terminal helical domain of HigC is required for promoter regulatory activity

Based on sequence homology, HigC belongs to the LytTR family of DNA binding proteins (pfam04397, COG3279), but in addition to the DNA binding domain that is typical of this family, it was previously proposed to contain four transmembrane helices (40). Analysis of the HigC protein sequence by the Dense Alignment Surface (DAS) program (41) agreed with this study, suggesting that the four transmembrane helices were in the N-terminal half of the protein sequence, preceding the DNA binding domain that is predicted to start at amino acid 171 (Fig 2A). Since the existence of transmembrane helices seemed counter-intuitive in a transcription factor, which should be able to localize to the nucleoid rather than the membrane, we constructed a truncated version of HigC in which the transmembrane helices were removed (HigC-noTM). Placing this construct under the control of the vanillate-inducible promoter allowed us to compare its effect on the higBA promoter to wild type HigC or the empty vector control (Fig 2B). In both WT and ΔhigBA strains, the wild type HigC repressed the promoter as before, but the truncated HigC lacking the N-terminal helical domain was completely inactive as a repressor, showing that this domain is required for activity.

We further analysed the HigC protein sequence for the presence or absence of a signal peptide, reasoning that if this N-terminal helical domain is genuinely a four-helix transmembrane domain, it should be preceded by a signal peptide to direct it to the membrane for co-translational
insertion. However, using the SignalP software (42), no signal peptide for either the Sec or Tat
secretion pathways was seen (Fig 2C). It is unlikely that this is a false negative, because the same
program could detect the signal peptide of the Caulobacter outer membrane protein ChvT with high
probability (Fig 2D). Therefore, it is possible that this domain was annotated as transmembrane
helices simply because it shares the same helical secondary structure and hydrophobicity of genuine
transmembrane helices, but is not actually targeted to the membrane. Since the domain was
required for promoter repression activity, and bacterial DNA binding proteins frequently function as
dimers or other multimers (43), we hypothesize that this domain may participate in protein-protein
interactions necessary for DNA binding instead, either between HigC monomers or with other
interaction partners.

HigC affects ciprofloxacin resistance independently of the toxin HigB

We then investigated whether HigC overproduction had other phenotypic effects than $P_{higB\Delta}$
repression, by overexpressing it from the vanillate-inducible promoter in WT, $\Delta$lexA and $\Delta$lexA $\Delta$higB
strains and testing its effect on viability in the presence of antibiotics (Fig 3A). At a sub-inhibitory (for
WT) concentration of ciprofloxacin, the viability of the $\Delta$lexA mutant was reduced relative to the WT
and $\Delta$lexA $\Delta$higB strains (all containing empty vector), but the viability of the three strains was
unchanged on the control plate (containing gentamicin to maintain selection of the pMT335 vector)
and on a sub-inhibitory concentration of chloramphenicol. However, on mild overexpression of HigC,
the viability of the $\Delta$lexA strain in the presence of ciprofloxacin was reduced even further, and
strikingly the improved resistance of the $\Delta$lexA $\Delta$higB strain to ciprofloxacin was completely
reversed. This effect was unique to ciprofloxacin, as it was not seen in the control condition or on
chloramphenicol. Viability of a $\Delta$lexA $\Delta$higC strain was slightly improved relative to the $\Delta$lexA parent
strain on ciprofloxacin (Fig 3B), showing that the negative effect of HigC overexpression was not
likely due to non-specific intolerance of producing this protein at higher levels than the cell normally
experiences. Therefore, HigC negatively influences survival in the presence of DNA damaging
antibiotics, especially in the context of constitutively activated SOS response of the ΔlexA mutant.

Moreover, since this effect was observed in a ΔlexA ΔhigB mutant, this effect cannot be ascribed to HigC altering $P_{higA}$ promoter activity and HigB acting as the effector of the response. Rather, HigC must be a direct effector of the ciprofloxacin sensitivity, potentially through regulatory activity on other promoters than $P_{higA}$.

**HigBAC has no effect on formation of persister cells**

Since TA systems had been previously implicated in persister cell formation, we next investigated whether the effect of HigB or HigC on viability in the presence of ciprofloxacin was associated with any change in frequency of persister cell formation. Exposure of WT, ΔhigA, ΔhigBA, ΔhigBAC and ΔhigC cells to a bactericidal concentration of ciprofloxacin followed by dilution spot plating showed that all strains exhibited a biphasic killing curve typical of persister cell formation with the initial rapid killing phase from 0 to 6 hours and with persister cells detectable after 24 and 48 hours (Fig 4A), similar to recent work in which persistence to streptomycin and vancomycin was quantified (44). This timecourse experiment showed that these strains displayed very similar biphasic curve profiles to each other with no difference in the rate of the rapid killing phase or the fraction of persisters recovered at 24 or 48 hours. However, we were unable to consistently recover persisters at 48 hours from the ΔhigBA cultures using the spot dilution plate method, so we repeated the experiment measuring only 48-hour persisters but from a larger number of cells. This showed that all strains reproducibly had a fraction of $10^{-4}$ to $10^{-5}$ surviving persister cells after 48 hours ciprofloxacin, and that there was no significant difference in fraction of surviving persisters between any of these strains (Fig 4B). We also performed the timecourse experiment for the ΔlexA, ΔlexA ΔhigBA and ΔlexA ΔhigB strains relative to WT and found that these strains had similar biphasic curve kinetics and a similar fraction of surviving persister cells at 24 and 48 hours, and again no significant difference between any of the strains was seen (Supplementary Fig S1). Therefore, while *Caulobacter crescentus* is capable of forming persister cells upon bactericidal antibiotic
(ciprofloxacin) treatment, this process is not influenced by the toxin HigB, the transcription factor HigC, or the LexA repressor which controls their expression, and the viability differences observed in our efficiency of plating assays are unrelated to persistence.

**HigB negatively regulates CtrA-dependent gene expression**

We had previously identified the transcript of the cell cycle regulator *ctrA* as a target of the toxin HigB’s mRNA interferase activity (23), and confirmed that increased HigB activity in the Δ*higA* mutant strain was associated both with a decreased proportion of swarmer cells in the population, and with protection against cell cycle arrest caused by overexpression of a dominant-negative CtrA allele (CtrA-DN) that could not be removed from the cells by regulated proteolysis (45), presumably by increased HigB-mediated degradation of the *ctrA*-DN mRNA. To investigate whether this was reflected at the phenotypic level by altered transcription of CtrA-dependent genes, we carried out β-galactosidase assays of CtrA-dependent promoter-reporters in WT, Δ*higA* and Δ*higBA* strains, with and without overexpression of CtrA-DN from the vanillate-inducible promoter. Since this allele forces the cells to arrest in the G1 phase, promoter activity in the presence of the empty vector indicates that of the mixed population, while promoter activity in CtrA-DN-overexpressing cells indicates the level of activity seen specifically in the swarmer cells.

We compared the activity of CtrA-dependent promoters that are expressed in the G1 phase (swarmer cells) and subject to repression by the co-repressors MucR1/2, with CtrA-dependent promoters that are expressed in the late S-phase and in G2 (stalked and pre-divisional cells) and subject to repression by the regulatory protein SciP, which includes the promoter of CtrA itself. In the CtrA-DN-overexpressing but otherwise WT cells, we anticipated that SciP protein levels should be high and that the CtrA-SciP-dependent promoters should be inactive or weakly active compared to the mixed population/empty vector control. Meanwhile, the CtrA-MucR-dependent promoters should be active in both conditions (possibly increased upon CtrA-DN overexpression). Then, any
further differences in promoter activity in the ΔhigA or ΔhigBA backgrounds relative to WT should be accounted for by increased or decreased activity of the toxin HigB against ctrA mRNA. Consistent with our previous result that the loss of HigA had a reduced swarmer cell fraction in a mixed population but no difference in other cell types (23), as observed by FACS, we found reduced activity of the MucR-dependent G1-phase promoter \( P_{p_{i1}} \) in ΔhigA relative to WT both with and without CtrA-DN overexpression (Fig 5A). We did not observe the same effect for \( P_{sc_{i1}} \), suggesting that promoters controlling structural genes are better proxies for this effect than promoters controlling regulatory factors.

Surprisingly, the CtrA/SciP-dependent S/G2-phase promoters were not significantly repressed in the WT background when the CtrA-DN allele was overexpressed. However, loss of HigBA appeared to promote this repression, since the ΔhigBA strain had significantly lower activity of both promoters during CtrA-DN overexpression compared to empty vector. Surprisingly, upon CtrA-DN overexpression in the ΔhigA strain, we observed much stronger repression of \( P_{ctrA} \) than in ΔhigBA or WT (Fig 5B). The bipartite ctrA promoter is subject to complex multi-level regulation by SciP, CtrA itself, the S-phase associated transcription factor GcrA and the methylation state of the promoter DNA (Fig 5C), so the contribution of the multiple regulatory inputs cannot be inferred from the promoter activity measurement alone. However, we can nonetheless conclude that the ΔhigA genetic background influences cell cycle dependent gene expression, in a manner which is consistent with its cognate toxin HigB negatively regulating ctrA at the post-transcriptional level. In support of this function for HigB, we also found by anti-CtrA ChIP followed by quantitative PCR that the CtrA/MucR-dependent promoters \( P_{p_{i1}} \) and \( P_{sc_{i1}} \) had much less CtrA bound to them in non-synchronized populations of the ΔhigA strain compared to WT and ΔhigBA (Fig 6), despite the modest effects observed at the level of promoter activity (Fig 5A). HigB is therefore capable of negatively influencing CtrA binding to and activating its target promoters, regardless of the cell cycle phase they are associated with.
**HigC influences cell cycle gene expression during the SOS response independently of HigB**

Since we had observed that HigC could negatively affect survival in the presence of DNA damaging antibiotics in a HigB-independent manner, we then investigated whether this was associated with cell cycle gene expression by using the P_{pilA-lacZ} construct as a reporter for CtrA-dependent promoter activity in the presence and absence of ciprofloxacin, in strains lacking higBA, higC or lexA separately or together (Fig 7A). There was no difference in P_{pilA} activity between WT and ΔhigBA in the control condition, but its activity was increased in ΔhigBA cells treated with ciprofloxacin. However, this effect was not due to increased HigB activity in ciprofloxacin-treated WT, because the activity in a ΔhigBAC mutant strain treated with ciprofloxacin was reduced down to WT levels again. Hence, HigC must have been responsible for the elevated P_{pilA} activity in the ΔhigBA mutant upon induction of the DNA damage response with ciprofloxacin. In the ΔlexA strain, which has the DNA damage response constitutively activated, we observed similar results. Here, the baseline activity of P_{pilA} was lower, probably because of the LexA-induced cell division block (46, 47) that would prevent normal progression through the cell cycle and the associated pulse of pilA transcription in G1 phase. We did not observe any ciprofloxacin-induced increase in P_{pilA} activity in a ΔlexA ΔhigBA mutant compared to the ΔlexA strain. However, the ΔlexA ΔhigBAC quadruple mutant had decreased activity of this promoter compared to ΔlexA ΔhigBA, both with and without ciprofloxacin. Therefore, in conditions where the SOS response is induced but the HigBA TA system inactive, HigC can promote expression of this CtrA-dependent promoter. This activity must be functionally independent of the HigBA TA system, in the sense that it is not mediated by HigB toxin activity against CtrA via HigC regulation of the higBAC promoter. Overexpression of HigC in WT cells from the vanillate-inducible promoter, under the same conditions in which we saw HigC repression of P_{higBA}, did not result in any alteration of P_{pilA} activity relative to the empty vector (Supplementary Fig S2), suggesting that the effect of HigC on this promoter is either indirect, or undetectable if HigBA is present.
In the present study we report that the HigBA toxin-antitoxin system of *Caulobacter* possesses a transcription factor third component HigC, which participates in auto-regulation of the higBA promoter but which also acts independently of HigBA (Fig 7B). We confirm our findings from previous work that the HigB toxin can target the cell cycle regulator CtrA at the post-transcriptional level, resulting in decreased CtrA-promoter binding and lower CtrA-dependent promoter activity. Moreover, we find that under conditions of SOS response induction, HigC decreases cell viability and can also influence expression of CtrA target genes. The decrease in viability was independent of the higB toxin gene, while the expression of the CtrA-dependent pilA promoter was increased by HigC specifically in the absence of HigBA. Deletion of neither higBA nor higC had any effect on formation of persister cells in the presence of ciprofloxacin, suggesting that the HigBA- and HigC-dependent phenotypes that we observe are unrelated to the persistence phenomenon and instead indicate that HigBAC is acting as a regulatory coupling factor linking regulation of cell cycle genes to the SOS response.

While the close proximity of higC to higBA initially suggested that these genes may be in the same operon and therefore co-regulated, some aspects of whether higC is regulated identically to higBA still remain unknown. Mindful of the recent observation that in the *E. coli mqsRA TA* system, the antitoxin mqsA is transcribed from promoters internal to the mqsR coding sequence (18), we searched for promoters not only in the higA-higC intergenic region but in the entire higBA coding sequence. This bioinformatic analysis did not uncover any cryptic internal promoters. However, *Caulobacter* -10 and -35 promoter sequences do not closely match the canonical -10 and -35 boxes characterised in *E. coli* and other Gram negative bacteria, so it is also possible that this is a false negative. Indeed, in a previous global analysis of genome-wide transcription start sites over the cell cycle (48), it was found that there was a low-frequency transcription start site which corresponded to the A of the start codon of higC, in addition to the high-frequency transcription start site 4 bp
upstream of the start codon of higB. Therefore, it is possible to infer that higBA and higC are transcribed from different promoters, with the higC promoter being much weaker, which would explain our result that the higC transcript is apparently present at much lower levels than higB in WT, ΔhigA or ΔlexA strains (Fig 1D). This is more difficult to reconcile with the similar fold changes (between ΔlexA mutant and WT) for all three genes observed by qRT-PCR by da Rocha et al (30), since the region upstream of the putative higC transcription start site has no LexA binding site. However, if transcriptional readthrough occurred during the high levels of transcription from the higB promoter that would be expected during the SOS response, this could account for the lexA-dependent higC induction. Interestingly, the higC transcription start site was suggested to be cell cycle regulated while the higB transcription start site was not, with RNA-Seq reads corresponding to the higC site peaking at 80 to 100 minutes after synchronisation (48). This correlates closely with the peak time of the transcription start site of the ctrA P2 promoter, but higC is unlikely to be a candidate for direct regulation by CtrA since there is no CtrA binding motif (TTAA-N7-TTAA) in the higA – higC intergenic region and it has not been identified as a CtrA target in any genome-wide analysis (26, 49).

The role of HigC in repressing the HigBAC promoter is consistent with that observed for other three-component type II TA systems encoding a transcription factor (11, 31), but we also observe some unique differences. In those studies, the transcription factor was primarily responsible for repression of the system, either alone or together with the cognate TA complex acting as co-repressor. Meanwhile, for HigBAC, the HigC repression appears much less important than the repression provided by LexA and HigA. Based on our β-galactosidase data, HigC seems to exert the strongest repressive effect when HigA is absent, suggesting that it might act as a negative feedback mechanism to bring higBAC transcription back under control during the late SOS response, if LexA and/or HigA have been absent from the promoter. It will be intriguing to investigate how the 4-helix N-terminal domain of HigC is involved in promoter regulation, since removal of this domain completely abolished its activity as repressor of the higBA promoter. While we find that it is unlikely
to be a true membrane protein, on account of the lack of signal peptide, one possibility is that it could mediate protein-protein interactions between HigC monomers or between HigC and other proteins. Pull-down assays of WT and truncated HigC could identify binding partners of this protein and differentiate between ones that depend on the presence of the 4-helix domain and ones that do not. Moreover, since these helical domains were identified in proteins of this family from other alpha-proteobacteria, not only *Caulobacter* (40), this domain may represent a novel conserved mediator of DNA binding protein interaction in this class of bacteria. It will also be important to define the regulon, either direct or indirect, of HigC in order to fully characterise the role of HigC in the SOS response based on which other genes it regulates in addition to *higBA*.

Our genetic approach, in which we have characterised the effect of HigB in the absence of the antitoxin, and HigC in the absence of the HigBA TA system, has allowed us to gain valuable insight into the functions of these two proteins. However, it is also important not to infer too much from studies of mutant strains about the physiological roles of these proteins in wild type cells. A criticism which is often levelled at studies of TA systems is that phenotypes of antitoxin mutant strains are not equivalent to phenotypes of wild type cells experiencing high levels of toxin production and therefore not physiologically relevant (2, 19, 21), and therefore a phenotype associated with a given TA system should only be postulated if a phenotype can be observed for a toxin or whole TA system mutant. We do indeed observe such a phenotype for HigBA, since the loss of the toxin in the ΔlexA background substantially improved its resistance to ciprofloxacin. However, in this work we have also found that the difference in this ciprofloxacin resistance phenotype between our ΔhigB and ΔhigBA strains was due to the polar effect of the higBA deletion on higC (specifically, placing it immediately downstream of the strong higBA promoter leading to much stronger higC expression than would normally occur). This underscores the importance of taking genetic context into account and not assuming that in-frame deletions are free of polar effects. Nonetheless, we can still conclude that the HigB toxin is likely to be active to some extent during the SOS response, based on the ciprofloxacin resistance phenotype of the ΔlexA ΔhigB strain, and that
when active it should inhibit CtrA at the post-transcriptional level resulting in lower expression levels of CtrA-activated genes. In addition, the effect of HigC overexpression or deletion on ciprofloxacin resistance of the ΔlexA strain shows that it can exert its effect when expressed at relatively low levels and when higBA is still present. Taken together, our data show that the TA system HigBAC of \textit{Caulobacter crescentus} is a uniquely acting three-component TA system in which the toxin HigB and the transcription factor HigC exert SOS-responsive gene regulation activities at transcriptional and post-transcriptional levels, on genes involved in the cell cycle regulatory network.

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**Author contributions**

KG, KAB and CLK performed experiments. KG and CLK wrote the paper. CLK conceived and designed the study.

**Figure Legends**

Figure 1. The TA system higBA of \textit{Caulobacter crescentus} has a third component, higC. (A) Beta-galactosidase activity of P_{higBA}~lacZ transcriptional reporter in NA1000 (WT), ΔhigBA, ΔlexA and ΔlexA ΔhigBA strains. (B) Cartoon of operon structure, approximately to scale, of higB, higA and CCNA_03101/higC. higB and higA are translationally coupled with a 4 bp overlap and higC lies 42 bp downstream of the 3’ end of higA. The relative positions of LexA and HigA binding at the higBAC
promoter are based on the position of the known LexA box in the promoter (30) and on previously published ChIP-Seq data for HigA (23). (C) Beta-galactosidase activity of $P_{higBA}$-lacZ transcriptional reporter in NA1000 (WT), $\Delta$lexA and $\Delta$higBA strains carrying either the pMT335 empty vector (ctrl) or the higC overexpression construct pMT335-higC (+ higC) after 3 hours treatment with 50 µM vanillate to induce higC expression. * indicates p < 0.05 for comparison of higC overexpression to empty vector control in $\Delta$higBA. (D) Quantitative RT-PCR for the higB (grey) and higC (black) coding sequences performed on cDNA prepared from NA1000 (WT), $\Delta$higA, $\Delta$higBA, $\Delta$lexA and $\Delta$lexA $\Delta$higBA strains. For every sample, higB and higC quantity values were normalised to the values obtained for the housekeeping gene rpoD. Error bars for this graph indicate standard error of the mean.

Figure 2. The N-terminal helical domain of HigC is needed for its activity. (A) Cartoon of HigC domain structure showing the positions of the DNA binding domain as predicted by pfam and the putative transmembrane helices as predicted by DAS, alongside the truncated variant HigC-noTM. (B) Beta-galactosidase activity of $P_{higBA}$-lacZ transcriptional reporter in NA1000 (WT), $\Delta$lexA and $\Delta$higBA strains carrying the pMT335 empty vector (pMT335), the truncated higC overexpression construct pMT335-higC-noTM (+ higC-noTM) or the WT higC overexpression construct pMT335-higC (+ higC) after 3 hours treatment with 50 µM vanillate to induce higC expression. * indicates p < 0.05 for comparison of higC overexpression to empty vector control. (C) Signal peptide prediction using the SignalP-5.0 program set to detect signal peptides of Gram-negative bacteria, on the first 70 amino acids of HigC. The yellow line indicates that all of the provided sequence corresponds to non-signal peptide sequence with a probability of 1. (D) Signal peptide prediction using the SignalP-5.0 program as in (C) on the first 70 amino acids of ChvT. The red line indicates a high-probability Sec-dependent signal peptide and the peaks in the green line indicate two possible cleavage sites where the signal peptide could be cleaved from the rest of the protein after export.

Figure 3. HigC influences cell viability during the SOS response. (A) Efficiency of plating assays of NA1000 (WT), $\Delta$lexA and $\Delta$lexA $\Delta$higB strains carrying the empty vector pMT335 or the higC
overexpression plasmid pMT335-\textit{higC}, on plates containing the indicated antibiotics and 50 µM vanillate to induce \textit{higC} expression. (B) Efficiency of plating assays of \textit{NA1000 (WT), ΔlexA and ΔlexA ΔhigC} strains. All images are representative of three independent biological replicates.

Figure 4. HigBAC does not influence persister cell formation. (A) Biphasic killing curve of \textit{NA1000 (WT), ΔhigA, ΔhigBA, ΔhigBAC and ΔhigC} strains treated with 10 µg/ml ciprofloxacin. Samples were taken over time and spot plated (5 µl) in technical duplicates after washing and dilution to calculate surviving cfu/ml. Data were expressed as fraction surviving persisters normalised to cfu/ml at zero time, from three independent biological replicates. (B) Measurement of surviving persisters in the same strains as in (A) after 48 hr treatment with 10 µg/ml ciprofloxacin. Cfu/ml values were calculated from plating 100 µl samples of $10^{-5}$ and $10^{-6}$ dilutions of the zero-time sample and $10^{0}$ and $10^{1}$ dilutions of the 48 hr ciprofloxacin treated sample and normalised to fraction surviving persisters at 48 hr relative to zero time.

Figure 5. HigB activity affects expression of \textit{CtrA}-dependent cell cycle genes. (A) Beta-galactosidase activity of the \textit{CtrA/MucR}-dependent promoters $P_{\text{pilA}}$ and $P_{\text{scip}}$ in \textit{NA1000 (WT), ΔhigBA and ΔhigA} strains carrying either the empty vector pMT335 (white) or the non-proteolysable \textit{CtrA overexpression plasmid pMT335-ctrA-DN} (black), after 3 hours treatment with 500 µM vanillate to induce \textit{CtrA-DN} to a sufficient level to induce cell cycle arrest in the G1 phase (23). (B) Beta-galactosidase activity of the \textit{CtrA/SciP}-dependent promoters $P_{\text{flgB}}$ and $P_{\text{ctrA}}$ in \textit{NA1000 (WT), ΔhigBA and ΔhigA} strains carrying either the empty vector pMT335 (white) or the non-proteolysable \textit{CtrA overexpression plasmid pMT335-ctrA-DN} (black), after 3 hours treatment with 500 µM vanillate as in (A). * and ** indicate \textit{p < 0.05} and \textit{p < 0.01} respectively for within-strain comparisons of empty vector control to \textit{ctrA-DN} overexpression. § and §§ indicate \textit{p < 0.05} and \textit{p < 0.01} respectively for between-strain comparisons of \textit{ΔhigA or ΔhigBA} to WT cells carrying the same plasmid. (C) Cartoon of the \textit{ctrA} promoter and factors that regulate it. The dotted line indicates transcriptional repression of the \textit{gcrA} promoter by \textit{CtrA}, while solid lines indicate direct activation or repression.

* and ** indicate \textit{p < 0.05} and \textit{p < 0.01} respectively for within-strain comparisons of empty vector control to \textit{ctrA-DN} overexpression. § and §§ indicate \textit{p < 0.05} and \textit{p < 0.01} respectively for between-strain comparisons of \textit{ΔhigA or ΔhigBA} to WT cells carrying the same plasmid. (C) Cartoon of the \textit{ctrA} promoter and factors that regulate it. The dotted line indicates transcriptional repression of the \textit{gcrA} promoter by \textit{CtrA}, while solid lines indicate direct activation or repression.
Figure 6. Increased HigB activity reduces binding of CtrA to its target promoters. Binding of CtrA to its target promoters P_{pilA} (A) and P_{sciP} (B) measured by ChIP with anti-CtrA antibodies followed by quantitative PCR and expressed as fold enrichment in anti-CtrA ChIP over input DNA. Data are expressed as average and standard deviation of two biological replicates, with technical duplicates performed in each experiment.

Figure 7. HigC affects cell cycle-regulated gene expression independently of HigB, but only during induction of the SOS response. (A) Beta-galactosidase activity of P_{pilA}-lacZ in NA1000 (WT), ΔhigBA, ΔhigBAC, ΔlexA, ΔlexA ΔhigBA and ΔlexA ΔhigBAC, treated with vehicle (white) or with 5 µg/ml ciprofloxacin (black) for 2 hours. * indicates p < 0.05 and ** indicates p < 0.01 for between-strain comparisons of isogenic ΔhigBAC with ΔhigBA strains under identical treatment conditions (ciprofloxacin or vehicle). (B) Graphical summary of the regulation and function of the HigBAC 3-component TA system.

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Table 1. Oligonucleotide sequences used in this study. Restriction enzyme sites incorporated for cloning purposes are underlined.

| Oligo name          | Sequence (5’ – 3’)                  |
|---------------------|-------------------------------------|
| cc3036_nde          | AAAACATATGAGCGCTGACGCTTCCAAG        |
| cc3036_noTM_nde     | AAGCATATGGGCGCTACTTTCTCTGCCG        |
| cc3036_eco          | AAAAGTAATTCACTCGGCTGCTCGCCTCAGCCA  |
| 3036_up_bam         | AAGGATCCGGAAGCGTGCTGACGCTATGG      |
| 3036_up_hind        | AAAAGCTTGGTGATCGGTGCTCGATTGG       |
| 3036_down_bam       | AAGGATCCCTGGGCCGACGACGGAGTA        |
| 3036_down_eco       | AAAAATTCTGCCAGATCGACGCTTCT         |
| CC_3036_qrt_fwd     | GTTGACGCAAGGATGGGGAT               |
| CC_3036_qrt_rev     | CATGGTTGGGCCAGTGCCT                 |
| higB_qrt_fwd        | CCGACATGGAGCGCAATTTC               |
| higB_qrt_rev        | GTTGCAAGGCTGGCTAGGG                |
| rpoD_fwd            | GAAGAAGCGGGGCAAAGGTT               |
| rpoD_rev            | CGTCTTTGCTCTGGAGGTC                |
| pilA_chip_f         | TTTAGGGTCTGCAAAGCGTT               |
| pilA_chip_r         | GATCGGGAGCTGGAGGGGTT              |
| sciP_chip_f         | AGGCGTAAGCCATGGTTG                 |
| sciP_chip_r         | TTGTAACGATCGACGCTTC               |
Table 2. Plasmids used in this study

| Plasmid name       | Description                                                                 | Source   |
|--------------------|-----------------------------------------------------------------------------|----------|
| pP<sub>higBA</sub>-lac290 | plac290 (oriV, Tet<sup>R</sup>, lacZ low copy transcriptional fusion vector) with promoter of higBA inserted upstream of lacZ | (23)     |
| pP<sub>pilA</sub>-lac290 | plac290 with promoter of pilA inserted upstream of lacZ                    | (26)     |
| pP<sub>sciP</sub>-lac290 | plac290 with promoter of sciP inserted upstream of lacZ                    | (26)     |
| pP<sub>flgB</sub>-lac290 | plac290 with promoter of flgB inserted upstream of lacZ                    | (26)     |
| pP<sub>ctrA</sub>-lac290 | plac290 with promoter of ctrA inserted upstream of lacZ                    | (26)     |
| pMT335             | pBBR1 ori, rep, mob, Gent<sup>R</sup>, medium copy plasmid for vanillate-inducible gene expression | (50)     |
| pMT335-higC        | pMT335 derivative, P<sub>van</sub>-higC                                    | This work|
| pMT335-higC-noTM   | pMT335 derivative, P<sub>van</sub>-higC-noTM (Δ1-132 amino acids, Val133Met) | This work|
| pMT335-ctrA-DN     | pMT335 derivative, P<sub>van</sub>-ctrA-DN (DS1E CterDD)                   | (23)     |
| pNPTSΔhigC         | pNPTS138 (colE1 ori, M13 ori, oriT, Km<sup>R</sup>, sacB, suicide vector for in-frame deletions) derivative to introduce ΔhigC allele | This work|
| pNPTSΔhigBAC       | pNPTS138 derivative to introduce ΔhigBAC allele                             | This work|
### Table 3. Strains used in this study

| Strain number | Genotype                  | Source       |
|---------------|---------------------------|--------------|
| NA1000        | Wild type                 | (51)         |
| CLK891        | ΔhigA                     | (23)         |
| CLK113        | ΔhigBA                    | (23)         |
| CLK1203       | ΔlexA                     | (23)         |
| CLK1204       | ΔlexA ΔhigB               | (23)         |
| CLK1205       | ΔlexA ΔhigBA              | (23)         |
| CLK1658       | ΔlexA ΔhigC               | This work    |
| CLK1659       | ΔhigC                     | This work    |
| CLK1660       | ΔhigBAC                   | This work    |
| CLK133        | NA1000 pP<sub>higBA</sub>-lac290 | (23)         |
| DM217         | ΔhigBA pP<sub>higBA</sub>-lac290 | (23)         |
| DM223         | ΔlexA pP<sub>higBA</sub>-lac290 | (23)         |
| DM224         | ΔlexA ΔhigBA pP<sub>higBA</sub>-lac290 | (23)         |
| CLK196        | NA1000 pP<sub>higBA</sub>-lac290 pMT335 | (23)         |
| CLK1508       | NA1000 pP<sub>higBA</sub>-lac290 pMT335-<i>higC</i> | This work    |
| CLK1514       | ΔlexA pP<sub>higBA</sub>-lac290 pMT335 | This work    |
| CLK1512       | ΔlexA pP<sub>higBA</sub>-lac290 pMT335-<i>higC</i> | This work    |
| CLK234        | ΔhigBA pP<sub>higBA</sub>-lac290 pMT335 | (23)         |
| CLK1510       | ΔhigBA pP<sub>higBA</sub>-lac290 pMT335-<i>higC</i> | This work    |
| CLK1661       | NA1000 pP<sub>higBA</sub>-lac290 pMT335-<i>higC-noTM</i> | This work    |
| CLK1662       | ΔhigBA pP<sub>higBA</sub>-lac290 pMT335-<i>higC-noTM</i> | This work    |
| CLK113        | NA1000 pMT335              | (23)         |
| CLK158        | ΔlexA pMT335              | This work    |
| CLK1663       | ΔlexA ΔhigB pMT335         | This work    |
| CLK1664       | NA1000 pMT335-<i>higC</i> | This work    |
| CLK1665       | ΔlexA pMT335-<i>higC</i>   | This work    |
| CLK1666       | ΔlexA ΔhigB pMT335-<i>higC</i> | This work    |
| CLK1667       | NA1000 pP<sub>pilA</sub>-lac290 pMT335 | This work    |
| CLK1668       | NA1000 pP<sub>pilA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1669       | ΔhigA pP<sub>pilA</sub>-lac290 pMT335 | This work    |
| CLK1670       | ΔhigA pP<sub>pilA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1671       | ΔhigBA pP<sub>pilA</sub>-lac290 pMT335 | This work    |
| CLK1672       | ΔhigBA pP<sub>pilA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1673       | NA1000 pP<sub>sciP</sub>-lac290 pMT335 | This work    |
| CLK1674       | NA1000 pP<sub>sciP</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1675       | ΔhigA pP<sub>sciP</sub>-lac290 pMT335 | This work    |
| CLK1676       | ΔhigA pP<sub>sciP</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1677       | ΔhigBA pP<sub>sciP</sub>-lac290 pMT335 | This work    |
| CLK1678       | ΔhigBA pP<sub>sciP</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1679       | NA1000 pP<sub>higBA</sub>-lac290 pMT335 | This work    |
| CLK1680       | NA1000 pP<sub>higBA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1681       | ΔhigA pP<sub>higBA</sub>-lac290 pMT335 | This work    |
| CLK1682       | ΔhigA pP<sub>higBA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1683       | ΔhigBA pP<sub>higBA</sub>-lac290 pMT335 | This work    |
| CLK1684       | ΔhigBA pP<sub>higBA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work    |
| CLK1685       | NA1000 pP<sub>ctrA</sub>-lac290 pMT335 | This work    |
| Strain number | Genotype | Source |
|---------------|----------|--------|
| CLK1686       | NA1000 pP<sub>ctrA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work |
| CLK1687       | Δ<i>higA</i> pP<sub>ctrA</sub>-lac290 pMT335 | This work |
| CLK1688       | Δ<i>higA</i> pP<sub>ctrA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work |
| CLK1689       | Δ<i>higBA</i> pP<sub>ctrA</sub>-lac290 pMT335 | This work |
| CLK1690       | Δ<i>higBA</i> pP<sub>ctrA</sub>-lac290 pMT335-<i>ctrA-DN</i> | This work |
| CLK1322       | NA1000 pP<sub>pilA</sub>-lac290 | This work |
| CLK1326       | Δ<i>higBA</i> pP<sub>pilA</sub>-lac290 | This work |
| CLK1691       | Δ<i>higBAC</i> pP<sub>pilA</sub>-lac290 | This work |
| CLK1692       | Δ<i>lexA</i> pP<sub>pilA</sub>-lac290 | This work |
| CLK1642       | Δ<i>lexA</i> Δ<i>higBA</i> pP<sub>pilA</sub>-lac290 | This work |
| CLK1693       | Δ<i>lexA</i> Δ<i>higBAC</i> pP<sub>pilA</sub>-lac290 | This work |
Figure 3. Analysis of antibiotic sensitivity in strains with and without higC expression.

A. Control empty vector pMT335 and higC overexpression.

- Gentamicin
- Ciprofloxacin
- Chloramphenicol

B. PYE no antibiotic
- Ciprofloxacin
- Chloramphenicol

Strains:
- NA1000
- ΔlexA
- ΔlexAΔhigB
- ΔlexAΔhigC
