Acetylation of the CspA family protein CspC controls the type III secretion system through translational regulation of exsA in Pseudomonas aeruginosa

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Received February 27, 2021; Revised May 25, 2021; Editorial Decision May 27, 2021; Accepted June 08, 2021

ABSTRACT

The ability to fine tune global gene expression in response to host environment is critical for the virulence of pathogenic bacteria. The host temperature is exploited by the bacteria as a cue for triggering virulence gene expression. However, little is known about the mechanism employed by Pseudomonas aeruginosa to respond to host body temperature. CspA family proteins are RNA chaperones that modulate gene expression. Here we explored the functions of P. aeruginosa CspA family proteins and found that CspC (PA0456) controls the bacterial virulence. Combining transcriptomic analyses, RNA-immunoprecipitation and high-throughput sequencing (RIP-Seq), we demonstrated that CspC represses the type III secretion system (T3SS) by binding to the 5’ untranslated region of the mRNA of exsA, which encodes the T3SS master regulatory protein. We further demonstrated that acetylation at K41 of the CspC reduces its affinity to nucleic acids. Shifting the culture temperature from 25°C to 37°C in infection of mouse lung increased the CspC acetylation, which derepressed the expression of the T3SS genes, resulting in elevated virulence. Overall, our results identified the regulatory targets of CspC and revealed a regulatory mechanism of the T3SS in response to temperature shift and host in vivo environment.

INTRODUCTION

Upon entering host, pathogenic bacteria encounter numerous challenges, including temperature shift, nutrient availability, antibacterial substances, immune cells, etc. The ability to fine tune gene expression in response to environmental changes is critical for bacterial survival and propagation. Temperatures around 37°C constitute a hallmark of mammal hosts, which serves as a key signal to turn on specific regulatory pathways to orchestrate bacterial virulence gene expression for successful colonization (1,2).

Pseudomonas aeruginosa is a wide-spread Gram-negative opportunistic pathogen of human that causes a variety of community-acquired and nosocomial infections in patients with burn wounds, cystic fibrosis and those with compromised immunity (3–5). A previous transcriptomic study revealed that close to 400 genes were alternatively expressed between 28°C and 37°C, indicating regulatory pathways responding to the temperature shift (6). Genes that were up-regulated at 37°C include those related to the type III secretion system (T3SS) (6) that is a conserved virulence factor in Gram-negative pathogenic bacteria. The T3SS is a syringe-like apparatus that translocates effector proteins directly into the cytosols of a variety of eukaryotic cells, resulting in malfunction or death of the cells (7–9), which plays critical roles in the acute infections of P. aeruginosa (10–12).

Expression of the T3SS genes is highly energy consuming and thus is tightly controlled. ExsA is the master regulator that activates the expression of all the T3SS genes (13). The transcription of exsA is driven by two promoters — a promoter for the exsC-exsE-exsB-exsA operon (PexsA) and a promoter adjacent to the exsA coding region (PexsA). The PexsA is directly regulated by ExsA (14–19), while the PexsA is regulated by Vfr in conjunction with cAMP (20–23). The known T3SS inducing conditions include host cell contact, depletion of extracellular Ca²⁺ and serum albumin (24,25). The upregulation of the T3SS genes at 37°C indicates that the mammalian body temperature serves as one of the signals that coordinate expression of the T3SS genes in response to host environment. Elucidation of the temperature sensing and response pathways is critical to fully understand the bacterial virulence mechanisms.

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Disclaimer: The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.
CspA family proteins are a group of small nucleic acid binding proteins belonging to the cold shock domain protein family (26–28). These proteins share a conserved nucleic acid binding domain that is present in every kingdom of life (29–31). By binding to mRNAs, CspA family proteins control various biological processes and bacterial response to environmental changes (32–39).

Here we show that a *P. aeruginosa* CspA family protein CspC controls the expression of T3SS genes by binding to the 5’ UTR of the *exsA* mRNA. In addition, we demonstrate temperature responsive acetylation of CspC, which affects its nucleic acid binding ability and thus its regulatory effect on the *exsA*.

In combination, our results reveal a global regulatory role of the CspC and a post-translational modification (PTM)-mediated bacterial response to mammalian in vivo environment.

**MATERIALS AND METHODS**

**Bacterial strains and plasmids**

Bacterial strains, plasmids and primers used in this study are listed in Supplementary Table S4. Both *P. aeruginosa* and *Escherichia coli* strains were cultured aerobically in LB medium (10 g/l tryptone, 5 g/l NaCl, 5 g/l yeast extract, pH 7.0–7.5). When needed, the medium for *P. aeruginosa* was supplemented with tetracycline (50 μg/ml) (BBI life sciences, Shanghai, China), gentamicin (50 μg/ml) (BBI life sciences) or carbenicillin (150 μg/ml) (BBI life sciences) and the medium for *E. coli* was supplemented with tetracycline (10 μg/ml), gentamicin (10 μg/ml) or ampicillin (100 μg/ml) (BBI life sciences).

**RNA isolation and quantitative RT-PCR (qRT-PCR)**

Bacteria grown at indicated conditions were harvested by centrifugation at 12,000 × g for 1 min and total RNA were isolated with an RNAprep pure bacteria kit (Tiangen Biotech, Beijing, China). cDNA was synthesized with a reverse transcriptase (TAKARA, Dalian, China). For quantitative reverse transcriptase-polymerase chain reaction (qRT-PCR), indicated specific primers were mixed with diluted cDNA and SYBR Premix Ex Taq II (Takara). *rpsL* that encodes the 30 S ribosomal protein was used as an internal control. The results were determined and analyzed with the CFX Connect Real-Time system (Bio-Rad, USA).

**Mouse acute pneumonia model and histological procedures**

All animal experiments were in accordance with Nankai University and Chinese national laboratory animal care and usage guidelines. The protocol was approved by the animal care and use committee of the College of Life Sciences, Nankai University and the permit number is NK-04-2012. To test the bacterial virulence, the indicated strains were cultured in LB to an OD600 of 1.0, followed by centrifugation at 12,000 × g for 1 min and the pellets were resuspended in phosphate-buffered saline (PBS) to 2 × 10⁸ CFU/ml. 6-week-old female BALB/c mice (Vital River, Beijing, China) were anesthetized by intraperitoneal injection with 85 μl 7.5% chloral hydrate and then intranasally inoculated with ~4 × 10⁶ CFU bacteria. Twelve hours post-infection (hpi), mice were sacrificed with CO₂ and the lungs were removed. For bacterial colonization, the lungs were homogenized in 1% protease peptone (Solarbio, Beijing, China) and bacterial loads were determined by plating. The hematoxylin and eosin (H&E) staining was performed as previously described (40). Briefly, the lungs were fixed with 4% paraformaldehyde (Servicebio, Wuhan, China) overnight, dehydrated in ethanol, embedded in paraffin, sectioned and stained with hematoxylin and eosin.

**Cell culture and lactate dehydrogenase (LDH) release assay**

A549 cells were cultured with the Roswell Park Memorial Institute (RPMI) 1640 medium (Corning, USA) containing 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin (Gibco, USA) at 37°C in 5% CO₂. 2 × 10⁴ cells were seeded into each well of a 96-well plate. The medium was replaced with antibiotic and FBS-free RPMI 1640 3 h before bacterial infection. Bacteria were grown in LB at 37°C to an OD600 of 1.0, and then washed twice and resuspended in PBS. A549 cells were infected with indicated *P. aeruginosa* strains at a multiplicity of infection (MOI) of 50. The plate was centrifuged at 1000 × g for 5 min to synchronize the infection. Two hours post-infection, LDH present in the supernatant was measured by the LDH cytotoxicity assay kit (Beyotime, China). Cells treated with 1% Triton X-100 were used as the control of total release (100% LDH release) and the free RPMI 1640 medium was used as the negative control (0% LDH release). The cytotoxicity of bacteria was calculated following the manufacturer’s instruction.

**β-galactosidase activity assay**

The β-galactosidase activity assay was performed as previously described with minor modifications (41). Bacteria were grown in LB with or without 5 mM EGTA at 37°C until the OD600 reached 1.0. The bacteria were collected and resuspended in 1 ml Z buffer (0.06 M Na₂HPO₄, 0.04 M NaH₂PO₄, 0.01 M KCl, 0.001 M MgSO₄ and 0.05 M β-mercaptoethanol). A total of 500 μl of the suspension was mixed with 10 μl 0.1% sodium dodecyl sulfate (SDS) and 10 μl chloroform by vortex. After the addition of 100 μl of 4 mg/ml onithinotriphenyl-galactopyranoside (ONPG) (BBI life sciences), the mixture was incubated at 37°C immediately. When the color turned light yellow, 500 μl 1M Na₂CO₃ was added to stop the reaction. After recording the reaction time, OD₄₂₀ of the bacterial suspensions and OD₄₂₀ of the reaction mixtures were determined. The β-galactosidase activities (Miller units) were calculated as follows: Miller units = (1000 × OD₄₂₀)/T/500/OD₆₀₀; T, reaction time.

**RNA immunoprecipitation (RIP) and transcriptome sequencing analyses**

Overnight culture of the ΔcspC mutant containing pMMB67EH-CspC-GST was subcultured in fresh LB containing 150 μg/ml carbenicillin at 37°C. 0.5 mM IPTG was added to the medium when the OD₆₀₀ reached 0.5. After 4 h induction at 37°C, the bacteria were harvested.
by centrifugation and the CspC-GST recombinant protein was purified with a GST-tag Protein Purification Kit (Beyotime, China). A total of 1 mM DTT and RNase Inhibitor (Beyotime, China) was added to all liquid reagents in the process. The purified protein was mixed with three times the volume of TE buffer, followed by RNA isolation with an RNAprep pure bacteria kit (Tiangen Biotech, Beijing, China).

The RNA-sequencing was performed by GENEWIZ (Suzhou, China). The RNA samples were quantified using a Qubit 2.0 Fluorometer (Invitrogen, Carlsbad, CA, USA) and qualified by Agilent Bioanalyzer 2100 (Agilent Technologies, Palo Alto, CA, USA) and NanoDrop (Thermo Fisher Scientific Inc.). A total of 1 μg RNA and 10 μg RNA immunoprecipitation (RIP) sample were used for the following library preparation.

Next generation sequencing library preparations were constructed according to the manufacturer’s protocol (NEBNext® Ultra™ Directional RNA Library Prep Kit for Illumina®). The RNA was depleted from total RNA using Ribo-Zero rRNA Removal Kit (Bacteria) (Illumina). The ribosomal depleted mRNA was then fragmented and reverse transcribed. First strand cDNA was synthesized using ProtoScript II Reverse Transcriptase with random primers and Actinomycin D. The second strand cDNA was synthesized using Second Strand Synthesis Enzyme Mix (including dACGTP/dUTP). The double-stranded DNA was purified by AxyPrep Mag PCR Clean-up (Axygen) and then treated with End Prep Enzyme Mix to repair both ends and add a dA-tailing in one reaction, followed by a T-A ligation to add adaptors to both ends. Size selection of the adaptor-ligated DNA was then performed using AxyPrep Mag PCR Clean-up (Axygen), and fragments of ~360 bp (with the approximate insert size of 300 bp) were recovered. The dUTP-marked second strand was digested with the Uracil-Specific Excision Reagent (USER) enzyme (New England Biolabs). Each sample was then amplified by PCR for 11 cycles using P5 and P7 primers as instructed by the Manual of NEBNext® Ultra™ II Directional RNA Library Prep Kit for Illumina® (New England Biolabs). Both the P5 and P7 primers carry sequences that can anneal with flow cell to perform bridge PCR and P7 primer carrying a six-base index allowing for multiplexing. The PCR products were cleaned up using the AxyPrep Mag PCR Clean-up (Axygen), validated using an Agilent 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA, USA) and quantified by Qubit 2.0 Fluorometer (Invitrogen, Carlsbad, CA, USA).

Then libraries with different indices were multiplexed and loaded on an Illumina HiSeq instrument according to manufacturer’s instructions (Illumina, San Diego, CA, USA). Sequencing was carried out using a 2 × 150 paired-end (PE) configuration. Sequence read analysis was based on the predicted genes of strain PA14 (http://www.pseudomonas.com) (42). Differential expression analysis used the DESeq Bioconductor package, a model based on the negative binomial distribution. The sequencing was performed with each single sample of the wild-type PA14 and the ΔcspC mutant. After adjusted by Benjamini and Hochberg’s approach for controlling the false discovery rate, P-value of genes were setted <0.05 to detect differential expressed ones. The entire transcriptome sequencing data have been submitted in the NCBI SRA, with accession number PRJNA690968.

**Electrophoretic mobility shift assay**

Both dsDNA and ssDNA fragments corresponding to the sequences upstream of the cspa coding region were synthesized by GENEWIZ. A total of 40 ng DNA were incubated with 0 to 2 mM purified GST-CspC protein at 16 or 25°C for 30 min in a 20 μl reaction (10 mM Tris–HCl, pH 7.6, 4% glycerol, 1 mM ethylenediaminetetraacetic acid, 5 mM CaCl2, 100 mM NaCl). The samples were loaded onto an 8% native polyacrylamide gel that had been pre-run for over 1 h. The electrophoresis was performed on ice at 120 V for 1 h or 1.5 h. The gel was stained with 0.5 × TBE containing SYBR® Gold nucleic acid gel stain for ssDNA and 0.5 × TBE containing 0.5 μg/ml ethidium bromide for dsDNA. Bands were visualized with a molecular imager ChemiDoc™ XRS+ (Bio-Rad).

**Western blot**

Protein samples from equivalent amounts of bacterial culture were loaded onto a 12% SDS-polyacrylamide gel electrophoresis (SDS-PAGE) gel. Proteins were transferred to a polyvinylidene difluoride (PVDF) membrane (Millipore, USA). The membrane was incubated with a mouse monoclonal anti-His antibody (CST, USA), a rabbit monoclonal anti-Flag antibody (Sigma, USA), a rabbit monoclonal anti-Glutathione-S-Transferase (GST) antibody (Sigma, USA) or a mouse monoclonal anti-RNA polymerase α antibody (Biologend). For determination of acetylation, equivalent amounts of purified proteins were loaded onto a 12% SDS-PAGE gel, and then transferred to a PVDF membrane. The membrane was incubated with a rabbit monoclonal anti-GST antibody (Sigma, USA) or a mouse monoclonal anti-acetyllysine antibody (Jingjie PTM Biolab, China). Signals were detected with an Immobilon™ Western kit (Millipore).

**RNA extraction and protein purification from in vivo samples**

RNA extraction from in vivo samples was performed as previously described with minor modification (43). Briefly, BALB/c mice were intranasally inoculated with 4 × 10⁷ CFU of indicated strains and sacrificed 6 hpi. The bronchoalveolar lavage fluid (BALF) was collected and the bacteria were harvested by centrifugation. Then the pellets from 20 mice were mixed, followed by purification of the CspC-GST protein with a GST-tag Protein Purification Kit (Beyotime, China).
LC-MS/MS analysis

The LC-MS/MS analysis was performed by the PTM Biolab (Hangzhou, China). The tryptic peptides were dissolved in 0.1% formic acid (solvent A), directly loaded onto a home-made reversed-phase analytical column (15-cm length, 75 µm i.d.). The gradient was comprised of an increase from 6 to 23% solvent B (0.1% formic acid in 98% acetonitrile) over 16 min, 23 to 35% in 8 min and climbing to 80% in 3 min then holding at 80% for the last 3 min, all at a constant flow rate of 400 nl/min on an EASY-nLC 1000 UPLC system.

The peptides were subjected to NSI source followed by tandem mass spectrometry (MS/MS) in Q Exactive™ Plus (Thermo) coupled online to the UPLC. The electrospray voltage applied was 2.0 kV. The m/z scan range was 350–1800 for full scan, and intact peptides were detected in the Orbitrap at a resolution of 70 000. Peptides were then selected for MS/MS using NCE setting as 28 and the fragments were detected in the Orbitrap at a resolution of 17 500. A data-dependent procedure that alternated between one MS scan followed by 20 MS/MS scans with 15.0 s dynamic exclusion. Automatic gain control was set at 5E4.

mRNA stability assay

The mRNA stability assay was performed as previously described (41). Wild-type PA14 and the ΔcspC mutant were grown in LB at 37°C to an OD_{600} of 1.0, followed by treatment with 500 µg/ml rifampin. At indicated time points, equivalent numbers of bacterial cells were collected. Each sample was mixed with equal number of E. coli cells expressing the gfp gene. Total RNA was extracted, and the rpsL and the exsA mRNA levels in each sample were determined by real-time PCR using gfp as the internal control.

RESULTS

Expression patterns of CspA family proteins in P. aeruginosa

The P. aeruginosa wild-type strain PA14 harbors five cspA homologous genes, namely PA0456, PA0961, PA1159, PA2622 (cspD) and PA3266 (capB) (45). All of the five encode proteins contain a highly conserved oligonucleotide binding (OB)-fold motif (46) (Supplementary Figure S1a). The evolutionary relationships between the CspA family proteins in P. aeruginosa and E. coli were evaluated by constructing a neighbor-joining phylogenetic tree using MEGA X (47) (Supplementary Figure S1b). To examine whether the expression of these genes responds to temperature shift, we cultured wild-type PA14 at 37°C to an OD_{600} of 0.5, followed by shifting to 16°C. After 1 h culture, half of the cells was switched back to 37°C, while the other half was remained at 16°C. At indicated time points along the experiment, the bacterial cells were collected and the mRNA levels of the five csp genes were determined by qRT-PCR (Supplementary Figure S2a). As shown in Supplementary Figure S2b, all of the cold shock genes were upregulated after growth at 16°C for 1 or 4 h. When the culture temperature was shifted back to 37°C, the expression of PA0456, PA0961 and capB was downregulated, whereas the mRNA levels of PA1159 and cspD kept increasing (Supplementary Figure S2b).

The expression of CspA family proteins could be influenced by growth phases (48,49). In the P. aeruginosa strain PA14, our experimental results revealed that the expression of PA1159, capB and cspD was increased in the stationary phase (Supplementary Figure S2c). Thus, PA1159 and cspD might be regulated in response to both temperature and growth phase. In our experiment, the bacterial OD_{600} reached 1.8 after switching from 16°C to 37°C and grown for 1 h, which might result in the increased mRNA levels of the PA1159 and cspD. In combination, all the five CspA homologs in P. aeruginosa are regulated in response to temperature shift, and the expression of PA1159, capB and cspD is also influenced by growth phases.

PA0456 influences the bacterial virulence in a mouse acute pneumonia model

To test whether the five CspA homologs are involved in the virulence of P. aeruginosa, we constructed in-frame deletion mutants of each of the cspA family genes in PA14 and examined the bacterial virulence in a mouse acute pneumonia model. Mutation of PA0456, but not the other four genes, increased the bacterial loads in the lungs (Figure 1A and Supplementary Figure S3a). To examine whether the enhanced virulence of the ΔPA0456 mutant is due to increased growth rate, we monitored the bacterial growth in vitro. In LB medium, the ΔPA0456 mutant grew at the same rate as the wild-type PA14 (Supplementary Figure S3b). These results revealed a role of the PA0456 in modulating the bacterial virulence. Sequence analysis revealed that PA0456 is highly homologous to the E. coli CspC and CspE (Supplementary Figure S1b and Table S1). Complementation with the E. coli cspC or cspE gene restored the mRNA levels of exsA and exsC in the ΔPA0456 mutant (Supplementary Figure S4), indicating a functional similarity between PA0456 and the E. coli cspC or cspE. Since PA0456 is slightly more similar to the E. coli CspC, we designated PA0456 as CspC and conducted further functional studies.

CspC represses the expression of T3SS genes

To understand the mechanism of CspC-mediated regulation on the bacterial virulence, we performed transcriptome sequencing analyses and found that 174 genes were differentially expressed in the ΔcspC mutant (Supplementary Table S2). Of note, all the T3SS genes were upregulated in the ΔcspC mutant (Figure 1B). qRT-PCR results verified that deletion of the cspC resulted in higher mRNA levels of exsA, exsC and perV under both T3SS non-inducing and inducing conditions, which were restored by complementation with an intact cspC gene (Figure 1C). By utilizing a plasmid encoding a 6 × His-tagged exoU driven by its native promoter, we observed that the ExoU-His protein level was also increased in the cspC mutant strain (Figure 1D).

Since the T3SS plays a major role in the bacterial cytotoxicity, we performed a lactate dehydrogenase (LDH) release assay with the human lung epithelial cell A549. As shown in Figure 1E, the cytotoxicity of the ΔcspC mutant was higher than the wild-type PA14 and the complemented strain. In combination, these results demonstrate a negative regulatory role of the CspC on the T3SS genes.
Figure 1. Mutation of PA0456 increases the bacterial virulence in a mouse acute pneumonia model. (A) Mice were inoculated intranasally with $4 \times 10^6$ CFU of the indicated strains. At 12 hpi, the mice were sacrificed and lungs were isolated and homogenized. The bacterial loads were determined by serial dilution and plating. The central bar indicates the mean, and the error bar indicates standard error. *$P < 0.05$ by the Mann-Whitney test. (B) Fold changes of the mRNA levels of the T3SS genes in the ΔcspC mutant. The wild-type PA14 and the ΔcspC mutant were grown to an OD$_{600}$ of 1.0 in LB at 37°C, followed by RNA isolation and RNA-seq analyses. The dark gray, gray and black bars represent the effector, structural and regulatory genes of the T3SS, respectively. (C) Total RNA was isolated from bacteria grown in LB with or without 5 mM EGTA at 37°C and the relative mRNA levels of $exsA$, $exsC$, $pcrV$ were determined by qRT-PCR. Data represent the mean of three independent experiments and error bars indicate standard deviation. ***, $P < 0.001$ by Student’s $t$-test. (D) Bacteria carrying an $exoU$-6×His driven by its native promoter ($P_{exoU}$-exoU-6×His) were grown to an OD$_{600}$ of 1.0 in LB containing 150 μg/ml carbenicillin with or without 5 mM EGTA. The ExoU-6×His levels were determined by western blot. (E) A549 cells were infected with the indicated strains at a MOI of 50 for 3 h. The bacterial cytotoxicity was determined by the LDH release assay. The data shown represent the results from three independent experiments. ***, $P < 0.001$ by Student’s $t$-test.
The regulatory role of CspC on the promoters of exsA and exsC

Since the mRNA level of the T3SS master regulator gene exsA was increased in the ΔcspC mutant, we explored the mechanism of CspC-mediated regulation on exsA. The exsA gene is driven by P_{exsA} and P_{exsC} (Figure 2A) that are directly controlled by Vfr/cAMP and ExsA, respectively (23). Thus, the expression of exsA is regulated through an autoactivation loop (Figure 2A). To examine whether CspC affects the transcription of exsA, we constructed three β-galactosidase transcriptional fusions, namely P_{exsC-lacZ}, P_{exsC-A-lacZ} and P_{exsC+120-lacZ} (Figure 2A). The expression levels of β-galactosidase from the P_{exsC-lacZ} and P_{exsC-A-lacZ} were increased in the ΔcspC mutant in the presence or absence of EGTA, indicating an elevated activity of the exsA promoter (P_{exsC}). However, the expression levels of β-galactosidase from the P_{exsC+120-lacZ} were similar in wild-type PA14 and the ΔcspC mutant (Figure 2B). To further verify the role of CspC on the P_{exsC} activity, we inserted two tandem transcriptional terminators (T0T1) between the exsB coding region and the P_{exsA} (Figure 2A), which blocks the read-through transcription from exsC to exsA, leaving the transcription of exsA solely dependent on the P_{exsA}. Insertion of the T0T1 in the chromosome of PA14 or the ΔcspC mutant resulted in similar exsA and exsC mRNA levels in the presence or absence of EGTA (Figure 2C). In combination, these results demonstrate that CspC controls the T3SS by influencing the activity of the exsA promoter but not the exsA promoter.

Identification of the CspC regulatory targets by RIP-seq

To understand the mechanism of CspC-mediated regulation of the T3SS, we overexpressed a C-terminus GST-tagged CspC (CspC-GST) or a GST protein alone in wild-type PA14 and performed RIP-seq analyses. Total bacterial RNA was quantified by RNA-seq and used as the input. Our results demonstrated that CspC represses the T3SS genes under non-inducing condition (without EGTA) (Figure 1C and D), indicating that CspC binds to its targets under this condition. Meanwhile, addition of EGTA retarded the bacterial growth (data not shown), which might interfere with the expression of the cspC-GST driven by the exogenous tac promoter. Thus, we performed the RIP and RNA-seq assays under the T3SS non-inducing condition. The CspC-GST or GST was purified through affinity chromatography under a native condition, followed by RNA extraction. The amount of RNA from the GST RIP sample was too low to perform RNA-seq. Thus, only the CspC-GST RIP sample was subjected to RNA-seq. The relative enrichment fold was calculated as number of reads in the RIP-seq (output) divided by the number of reads of the corresponding RNA fragment in the RNA-seq (input). The enriched RNA fragments were shown in Figure 3A and listed in Supplementary Table S3. Of note, a fragment containing the exsA mRNA and its 5′UTR was enriched 3.63 folds by the CspC-GST (Figure 3A and B). A RIP-coupled qRT-PCR assay confirmed the enrichment of the fragment (Figure 3C). These results indicate a possible post-transcriptional regulation of the exsA mRNA by CspC.

CspC controls the expression of exsA at the post-transcriptional level

To examine whether CspC controls exsA at the post-transcriptional level, we constructed a C-terminus FLAG-tagged exsA driven by an exogenous P_{lac} promoter. Based on the RIP-seq result, a 120 bp fragment upstream of the exsA coding region was included in the construct, resulting in P_{lac-120-exsA-FLAG} (Figure 3D). The ExsA-FLAG level was higher in the ΔcspC mutant under both T3SS inducing and non-inducing conditions (Figure 3E). To narrow down the region that is involved in the post-transcriptional regulation, we reduced the 5′UTR to 74 bp (designated as P_{lac-74-exsA-FLAG}), which displayed the similar expression pattern as the P_{lac-120-exsA-FLAG} (Figure 3E). However, reduction of the 5′UTR to 25 bp (P_{lac-25-exsA-FLAG}) resulted in expression levels of ExsA-FLAG in wild-type PA14 and the ΔcspC mutant (Figure 3E). To examine whether the exsA coding region is involved in the post-transcriptional regulation, we fused the 74 bp 5′UTR to a gst gene, resulting in P_{lac-74-gst} (Figure 3F). The GST level was higher in the ΔcspC mutant than that in the wild-type PA14 (Figure 3G), indicating a critical role of the 74 bp sequence in the CspC-mediated post-transcriptional regulation.

We then performed electrophoretic mobility shift assays (EMSAs) to confirm the binding between CspC and the 5′UTR fragment. Previous studies demonstrated that CSPs bind to ssDNA as efficiently as RNA (38, 50). Therefore, we used ssDNAs in the EMSAs. As shown in Figure 3H, CspC retarded the motility of the 74 nt ssDNA that represents the exsA 5′UTR. However, no obvious retardation was observed with the complementary ssDNA (Figure 3H), a 143 bp dsDNA of the 5′UTR sequence of exsA or a 74 nt ssDNA of the cspC coding region (Supplementary Figure S5a and b). In addition, the exsA mRNA stability in the ΔcspC mutant was similar as that in the wild-type PA14 (Supplementary Figure S5c and d). Taken together, these results demonstrate that CspC controls the translation of the exsA mRNA through its 74 nt 5′UTR.

A previous study demonstrated that the RNA helicase DeaD controls the translation of the exsA mRNA through the 37 bp 5′UTR and the coding region (51). We thus examined whether CspC interplays with DeaD in the post-transcriptional regulation of exsA. Deletion of the deaD gene reduced the expression level of the ExsA-FLAG from the P_{lac-74-exsA-FLAG} (Supplementary Figure S6a). However, dual deletion of cspC and deaD (ΔcspCΔdeaD) resulted in a similar level of the 74-ExsA-FLAG as that in the wild-type PA14 (Supplementary Figure S6a). We then constructed a FLAG-tagged exsA gene with the 37 bp native exsA 5′UTR driven by the P_{lac} promoter, resulting in P_{lac-37-exsA-FLAG}. Consistent with the previous report, deletion of the deaD gene reduced the expression level of the 37-ExsA-FLAG (Supplementary Figure S6b). However, the 37-ExsA-FLAG levels in the ΔcspC and ΔcspCΔdeaD mutants were similar as those in the wild-type PA14 and the ΔdeaD mutant, respectively (Supplementary Figure S6b). Meanwhile, expression of the 74-GST from the P_{lac-74-gst} was not affected by the deletion of deaD (Supplementary Figure S6c), thus verifying the involvement of the exsA coding region in the DeaD-mediated translational regulation.
Figure 2. The roles of CspC in the regulation of the promoters of exsC and exsA. (A) Schematic diagram of the exsCEBA operon and the P_{exsC}-lacZ, P_{exsA}-lacZ and P_{exsC-A}-lacZ transcriptional fusions. The P_{exsC} and P_{exsA} were indicated by arrows. ExsA binds to the promoter region of the exsCEBA operon and activates its expression. The insertion site of the transcriptional terminators T0T1 was indicated. (B) Bacteria carrying the P_{exsC}-lacZ, P_{exsA}-lacZ and P_{exsC-A}-lacZ were grown at 37°C in LB containing 50 μg/ml tetracycline with or without 5 mM EGTA. The Miller units were the means of results of at least three independent experiments. ***, P < 0.001 by Student’s t-test. (C) The bacteria were grown at 37°C in LB with or without 5 mM EGTA and the mRNA levels of exsA and exsC were determined by quantitative real-time PCR. Data represent the mean of three independent experiments and error bars indicate standard deviation. ***, P < 0.001; NS, not significant by Student’s t-test.
Figure 3. CspC controls the expression of exsA at the post-transcriptional level. (A) Enriched RNA fragments by RIP-seq. Each dot indicates an enriched RNA fragment. The fragment containing the exsA mRNA and its 5′ UTR was shown as a red dot and indicated by a red arrow. (B) Standardized reads distributions of the peak regions of the RIP-seq and RNA-seq (input) results. The open reading frames on the chromosome are indicated in the lowest panel. (C) Fold enrichment of the indicated fragments tested by an RIP-coupled qRT-PCR assay. Data represent the mean of three independent experiments and error bars indicate standard deviation. *** P < 0.001 compared to the other samples by Student's t-test. (D) Schematic diagram of the exsA-FLAG fusions driven by a tac promoter with indicated length of upstream regions. (E) Bacteria carrying the exsA-FLAG with indicated upstream segments (P_tac-120/74/25-exsA-FLAG) were grown to an OD600 of 1.0 at 37°C in LB containing 150 µg/ml carbenicillin with or without 5 mM EGTA. The ExsA-FLAG levels were determined by western blot. (F) Schematic diagram of a gat gene driven by a tac promoter with the 74 bp upstream region of exsA (P_tac-74-gat). (G) Indicated strains carrying the P_tac-74-gat were grown to an OD600 of 1.0 at 37°C in LB containing 150 µg/ml carbenicillin. The amounts of GST and RpoA were determined by western blot. (H) The purified CspC-GST was incubated with a 74 nt ssDNA that represents the 74 nt 5′-UTR of the exsA mRNA and a complementary ssDNA for 30 min at 16°C. The samples were subjected to electrophoresis in a native gel, followed by staining with SYBR Gold Nucleic Acid Gel Stain.
In the ΔcspCΔdeaD mutant, the 74-GST level was similar as that in the ΔcspC mutant (Supplementary Figure S6c). In combination, these results indicate that DeaD and CspC regulate the translation of exsA independently.

The CspC is acetylated in a temperature-dependent manner

Next, we explored the CspC-mediated signaling pathway that controls the T3SS. A previous study demonstrated upregulation of the T3SS genes when the culture temperature was shifted from 28°C to 37°C (6). By growth of the wild-type PA14 at room temperature (25°C) and 37°C, we did confirm the upregulation of the T3SS genes at 37°C (Figure 4B). Utilizing the P_{lac}−74−exsA−FLAG, we found that the ExsA−FLAG level was 2.95-fold higher at 37°C than at 25°C in the wild-type PA14 (Figure 4A). In the ΔcspC mutant, the ExsA−FLAG levels were 4.6- and 1.9-fold higher than those in the wild-type PA14 at 25°C and 37°C, respectively (Figure 4A), indicating that CspC represses the translation of exsA to a greater extent at 25°C.

We then compared the expression levels of CspC at 25°C and 37°C. The mRNA and protein levels of CspC were similar at the two temperatures (Figure 4B and C), which intrigued us to speculate a PTM of the CspC. An LC–MS/MS analysis of the CspC-GST purified from the ΔcspC mutant grown at 37°C revealed an acetylation at position K41 (Figure 4E). In addition, higher amount of acetylated CspC-GST was observed from cells grown at 37°C than that at 25°C (Figure 4D), demonstrating that CspC is acetylated in a temperature-dependent manner.

Acetylation of the CspC at K41 modulates the expression of T3SS genes and the bacterial virulence in vivo

To examine whether CspC is acetylated in vivo, we infected mice intranasally with a ΔpscC mutant carrying the cspC-gst driven by the cspC native promoter. The pscC gene encodes a structural component of the T3SS machinery (53, 54), mutation of which attenuates the bacterial virulence, thus allowing a high infection dose for the isolation of the CspC-GST. Compared to the CspC-GST isolated from cells grown in LB at 25°C and 37°C, a higher level of acetylation was observed on the protein isolated from BALF of the infected mice. We then infected mice with the wild-type PA14, the ΔcspC mutant carrying the cspC or the K41R, K41Q variant of cspC. Infection with the ΔcspC mutant expressing cspC(K41R) resulted in reduced bacterial load and host inflammatory response compared to those by the ΔcspC mutant (Figure 6A–C). However, the cspC(K41Q) did not affect the bacterial load of the ΔcspC mutant or the host inflammatory response (Figure 6B and C). In addition, complementation with the cspC(K41R), but not the cspC(K41Q), in the ΔcspC mutant reduced the mRNA levels of exsA, exsC and pcrV in vitro as well as in vivo (Figure 6D). In combination, these results indicate that the in vivo activity of the CspC is influenced by the acetylation of the K41.

DISCUSSION

In this study, we examined the expression patterns of the five CspA family proteins in P. aeruginosa and studied their roles in bacterial virulence. Switching the culture temperature from 37°C to 16°C resulted in up regulation of all the CspA family protein genes, suggesting physiological roles of the CspA family proteins in response to cold shock.

In the acute pneumonia model, mutation of the cspC resulted in higher bacterial load and elevated inflammation. Transcriptome analyses revealed regulation of the T3SS in the ΔcspC mutant. Combining RIP-Seq and transcriptional fusion assays, we show that CspC represses the translation of the exsA mRNA by binding to the 74 nt fragment of its 5′-UTR. The CspC and CspE in Salmonella typhimurium play important roles in the bacterial virulence (55). A previous study in Staphylococcus aureus revealed a role of CspA in the regulation of proteins involved in metabolism, bacterial aggregation, response to oxidative stress and virulence (38). Further studies revealed that the CspA binds to the U-rich region of the 5′-UTR of its mRNA, which melts the stem-loop of the RNA and blocks the processing of RNase III, resulting in repression of its own mRNA translation (38). CspA family proteins have also been found to function as anti-terminators of transcription by altering the mRNA secondary structure, which allows the transcription of downstream genes under low temperatures in various bacteria, such as E. coli and Bacillus subtilis (34). We speculate that the binding of CspC to the 5′-UTR of the exsA mRNA might alter the RNA sec-
Figure 4. The activity of CspC is regulated by acetylation. (A) Indicated strains carrying the P_{	ext{exoA-74}-	ext{exsA-FLAG}} were grown to an OD_{600} of 1.0 in LB containing 150 μg/ml carbenicillin at 25°C or 37°C. The ExsA-FLAG levels were determined by western blot. (B) Relative mRNA levels of exsA and cspC. Total RNA of the wild-type PA14 was isolated from bacteria grown at 25°C or 37°C and mRNA levels of exsA and cspC were determined by qRT-PCR. Data represent the mean of three independent experiments and error bars indicate standard deviation. ***, P < 0.001, NS, not significant by Student's t-test. (C) Wild-type PA14 carrying a cspC-GST driven by its own promoter were grown to an OD_{600} of 1.0 in LB containing 150 μg/ml carbenicillin under 25°C or 37°C. The amounts of GST and RpoA were determined by western blot. (D) Acetylation levels of the CspC-GST protein under 25°C or 37°C. Acetylation and the total amounts of the purified CspC-GST were determined by western blot. (E) MS/MS spectra of the acetylated lysine peptide. The CspC-GST was purified from the ΔcspC mutant and subjected to the LC−MS/MS analysis. The b ion refers to the N-terminal parts of the peptide, and the y ion refers to the C-terminal parts of the peptide.
Figure 5. The acetylation of CspC influences its function in controlling the translation of exsA. (A) Equal amounts of GST, CspC(WT)-GST, CspC(K41Q)-GST, CspC(K41R)-GST, CspC(K48Q)-GST and CspC(K48R)-GST were incubated with the 74 nt ssDNA for 30 min at 16°C. The samples were subjected to electrophoresis in a native gel, followed by staining with SYBR Gold Nucleic Acid Gel Stain. (B) Binding affinities of CspC(K41Q)-GST, CspC(K41R)-GST and GST to the 74 nt ssDNA were determined by an SPR assay. (C) Relative mRNA levels of exsA, exsC and pcrV. Total RNA of indicated strains was isolated from bacteria grown in LB containing 150 µg/ml carbenicillin with or without 5 mM EGTA at 37°C. The mRNA levels were determined by qRT-PCR. Data represent the mean of three independent experiments and error bars indicate standard deviation. *** P < 0.001 by Student’s t-test. (D) Indicated strains carrying the P_tac-74-exsA-FLAG were grown to an OD₆₀₀ of 1.0 in LB containing 150 µg/ml carbenicillin at 25°C or 37°C. The ExsA-FLAG levels were determined by western blot.
Figure 6. Mutations of the K41 residue affected the function of CspC in vivo. (A) The ΔpscC mutant carrying the cspC-gst was grown in LB containing 150 μg/ml carbenicillin to an OD_{600} of 1.0 at 25°C or 37°C. Mice were inoculated intranasally with $1 \times 10^9$ CFU bacteria. At 4 hpi, bacteria from BALF were collected, followed by chromatography purification. Acetylation and the total amounts of the purified CspC-GST were determined by western blot. (B) Mice were inoculated intranasally with $4 \times 10^6$ CFU bacteria of the indicated strains. At 12 hpi, the mice were sacrificed and lungs were isolated and homogenized. The bacterial loads were determined by serial dilution and plating. The central bar indicates the mean, and the error bar indicates standard error. **$P < 0.01$, ***$P < 0.001$ by the Mann–Whitney test. (C) Pathology sections of lungs infected with the indicated strains. Mice were infected with $4 \times 10^6$ CFU of wild-type PA14, the ΔcspC mutant, the ΔcspC mutant complemented with wild-type cspC or the K41Q, K41R variants. Sterilized PBS was used as a negative control. Lungs from the infected mice were removed, fixed, sectioned and stained with hematoxylin and eosin. Images were taken with a 40 × objective lens. (D) Relative mRNA levels of T3SS genes and cspC before or during lung infection. Mice were infected intranasally with $1 \times 10^7$ CFU bacteria of the indicated strains. At 6 hpi, bacteria from BALF were collected, followed by RNA isolation. The mRNA levels of exsA, exsC and pcrV were determined by qRT-PCR. Data represent the mean of three independent experiments and error bars indicate standard deviation. ***$P < 0.001$, **$P < 0.01$, *$P < 0.05$ by Student’s t-test, ND, not detected.
ondary structure, which interferes with the access of ribosomes. Further studies are required to decipher the molecular mechanism on the CspC-mediated regulation on the exsA translation. In addition, the RIP-seq assay was performed under the T3SS non-inducing condition. Under the inducing condition, such as in the presence of EGTA, contact with host cells, additional regulatory targets of CspC might be identified by the RIP-seq assay.

By analyzing the RIP-seq result, we were unable to identify a consensus CspC binding sequence. In other bacteria, no consensus binding sequence of CspA family proteins has been found so far, which might be due to the short target sequence (estimated to be 5–7 nt) (38, 56, 57).

PTMs have been shown to play vital roles in bacterial metabolism, environmental stress responses and virulence (52, 58–62). Previous proteomic studies on the wild-type PA14 grown in a minimal glucose medium identified 320 acetylated proteins, mainly including virulence factors, the DNA-binding protein HU and those involved in carbon metabolism, oxidative stress response and LPS biosynthesis (63–65). Carbon sources have been shown to influence PTM in bacteria (66–69).

In a study utilizing various carbon sources in minimal medium, it was demonstrated that growth with citrate resulted in a high level of acetylated proteins in PA14 (70). In that study, CspC was found to be succinylated but not acetylated (70). However, our current study revealed acetylation but not succinylation of the CspC. We speculated that the discrepancy might be due to growth conditions, as we grew the bacteria in LB.

Here we found that the K41 is acetylated in CspC, which affects its function in the regulation of exsA. We believe it is the first report on the functional study on the acetylation of bacterial cold shock proteins. Acetylation of the K41 might neutralize its positive charge, thus reduces the binding affinity between CspC and its target nucleotides. Indeed, there was ~10-fold difference in the binding affinity to the exsA 5′-UTR between K41R and K41Q versions of the CspC that mimic unacetylated and acetylated forms, respectively. We further demonstrated a higher percentage of acetylated CspC during lung infection than that grown in vitro. Accordingly, we propose that the acetylation of CspC may play an important role in the bacterial response to host in vivo environment. The K41Q mutation abolished the CspC function in repressing the exsA translation under both in vitro and in vivo conditions. Although the K41R mutation did not enhance the function of CspC in terms of repressing the translation of exsA compared to the wild-type form, it abolished the temperature responsive regulation on the translation of exsA (Figure 5D). We suspect that substitution of the amino acid may have affected the protein structure, thus influenced its function.

Acetylation of lysine can be catalyzed by lysine acetyltransferases or occur nonenzymatically, using acetyl phosphate or acetyl-CoA as the donor (59, 71, 72). Meanwhile, deacetylation is catalyzed by deacetylases (60). Based on genome annotation of the P. aeruginosa strain PA14, there are total 91 acetyltransferases and 13 deacetylases (www.pseudomonas.com) (42). However, the acetyltransferase or deacetylase involved in PTM has not been identified in P. aeruginosa. Temperature or carbon source shift might affect the intracellular level of acetyl phosphate or acetyl-CoA as well as the expression level and activities of the acetyltransferase or deacetylase, which influence the global PTM in the cell. Identification of the acetyltransferases and deacetylases as well as the metabolism under different growth conditions will shed light on the regulatory mechanisms of the PTM of CspC and other proteins in response to environmental signals. Our results demonstrated a higher level of CspC acetylation in the bacteria isolated from BALF compared to that grown in LB at 37°C, indicating that the in vivo environment, presumably the nutrient compositions, might promote acetylation of the CspC.

Overall, our results demonstrate a role of the CspA family protein CspC in the virulence of P. aeruginosa and revealed a novel regulatory mechanism of the T3SS via CspC acetylation. Further studies are warranted for a deeper understanding of the CspC-mediated regulatory pathways in P. aeruginosa.

DATA AVAILABILITY
The datasets generated during the current study are available in the NCBI Short Read Archive (SRA) repository with an accession number PRJNA690968.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

FUNDING
National Key Research and Development Program of China [2021YFE0101700, 2017YFE0125600]; National Natural Science Foundation of China [82061148018, 31970179, 31900115, 31870130]; Tianjin Municipal Science and Technology Commission [19JCYBJC24700]. Funding for open access charge: National Key Research and Development Project of China [2017YFE0125600].

Conflict of interest statement. None declared.

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