Role of cpxa mutations in the resistance to aminoglycosides and β-lactams in salm...

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Introduction

With the overuse and misuse of antibacterial agents, the emergence and spread of multidrug resistant (MDR) Salmonella enterica serovar Typhimurium (S. Typhimurium) has become a clinical and public health threat worldwide (<u>Levy, 2002</u>; <u>Quinn et al., 2006</u>). To adapt to adverse environmental conditions, bacteria have developed many sophisticated signal transduction systems, referred to as two-component systems (TCSs). These systems sense changes in environmental pH, ion concentrations, nutrient levels, quorum signals, and antibiotic concentrations and, in response, regulate the expression of genes involved in processes such as cell growth, virulence, biofilm formation, quorum sensing, and antibiotic resistance (De la Cruz et al., 2015 ; Padilla-Vaca et al., 2017 ; Tiwari et al., 2017 ; Fujimoto et al., 2018). TCSs have been identified as potential drug targets. Unlike conventional antibiotics, these drugs would likely be effective against various drug-resistant bacteria and avoid the emergence of resistant strains (Gotoh et al., 2010; Tiwari et al., 2017). Understanding the regulation mechanisms of TCSs to conventional antibiotic resistance would be beneficial for the reasonable application and low occurrence of drug resistance to these TCSs-targeted antibacterial agents.

The CpxRA system is an important TCS in *Escherichia coli* and *Salmonella*. It consists of an inner membrane-bound histidine sensor kinase, CpxA, and a cytoplasmic response regulator, CpxR (<u>De Wulf et al., 2002</u>). CpxP is a periplasmic chaperone that binds to CpxA and inhibits its kinase activity. The periplasmic protease DegP can disrupt CpxP binding to misfolded proteins (

Isaac et al., 2005). Upon sensing extracytoplasmic stresses, CpxP binds to misfolded proteins, dissociates from CpxA, and is degraded by DegP, and then CpxA autophosphorylates and donates its phosphate group to CpxR at D51 (D51). This allows phosphorylated CpxR (CpxR-P) containing a specific DNA-binding site (M199) to bind to the promoter regions of target genes at the consensus sequence (GTAAAN ₅ GTAAA) (<u>De Wulf et al., 2002</u> ;

DiGiuseppe and Silhavy, 2003.), and to regulate a series of target genes associated with antibiotic resistance and pathogenesis (<u>Nishino et al., 2010</u>; <u>Dbeibo et al., 2018</u>). *cpxA* knock-out mutation can result in constitutive activation of the CpxRA system. This is because *cpxA* deletion abolishes its phosphatase activity for CpxR-P, CpxR then accepts a phosphoryl group from acetyl phosphate, the product of the phosphotransacetylase (Pta)-acetate kinase (AckA) pathway (<u>Wolfe et al., 2008</u>). Similarly, a strain containing the mutant *cpxA* gene (*cpxA* *) encoding a CpxA variant lacking several amino acids in the periplasmic loop, exhibits constitutive CpxRA system activity. This occurs because the mutant CpxA * protein still functions as an autokinase and CpxR kinase but lacks phosphatase activity, resulting in accumulation of CpxR-P and hyperactivation of the CpxRA response (<u>Raivio</u> and <u>Silhavy</u>, 1997).

The CpxRA envelope stress response system is closely related to drug resistance. In *E. coli*, *cpxA* deletion or *cpxA* * mutations will confer the resistance to hydroxyurea, β -lactams, aminoglycosides, and fosfomycin (<u>Mahoney and Silhavy, 2013</u>; <u>Masi et al., 2020</u>). Moreover, overexpression of *cpxR* can confer the resistance to kanamycin, amikacin, deoxycholate,

novobiocin, and β -lactams (<u>Hirakawa et al., 2003a</u>, <u>b</u>). In S. Typhimurium, the *cpxRA* deletion confers susceptibility to ceftriaxone by influencing the expression of *stm1530* and *ompD* (<u>Hu et al., 2011</u>), and overexpression of cpxR in the cpxR deletion mutant increases resistance to AGAs and β lactams as a result of the downregulation of outer membrane protein gene expression (Huang et al., 2016). However, the regulatory effect of CpxA on MDR has not been fully investigated. One study reported that both the cpxRand *cpxA* mutants were more sensitive to amikacin than the wild-type strain, but the *cpxA* * mutation renders *S*. Typhimurium more resistant to the antibiotic amikacin (<u>Humphreys et al., 2004</u>), which suggests that different mutant versions of the CpxA sensor protein exhibit unexpected resistance regulation in S. Typhimurium. Another study reported that the various activation pathways of the CpxRA system determine different sensitivities to β -lactams (Delhave et al., 2016). Therefore, the various mutations of cpxA should be analyzed in concert to understand the full extent of the effects of CpxA on MDR. In this study, we constructed different mutants of the cpxA gene to systematically analyze the regulatory effects of CpxA on AGAs and β lactams resistance and explored the related molecular mechanism. The results suggested that various *cpxA* mutations can activate the CpxRA system and confer resistance to AGAs and β -lactams.

Materials and Methods

Bacterial Strains, Plasmids, and Culture Conditions

The bacterial strains and plasmids used in this study are listed in <u>Table 1</u>. All bacterial strains were mutant derivatives of *S. enterica* serovar Typhimurium strain JS. All strains were cultured aerobically in Luria-Bertani (LB) medium https://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-to-aminoglycosides-and-lactams-in-salmonella-enterica-serovar-typhimurium/

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[1% (w/v) tryptone, 0. 5% (w/v) yeast extract, and 1% (w/v) NaCl, pH = 7. 0] (Takara Bio., Kusatsu, Japan) at 37°C. When necessary, LB medium were supplemented with ampicillin (100 µg/ml), kanamycin (50 µg/ml), chloramphenicol (25 µg/ml), or sucrose (8%, w/v).

TABLE 1

Strains and plasmids used in this study.

Construction of Various Mutants of S. Typhimurium

All deletion mutants were generated *via* the Red recombinase system, as reported previously (<u>Datsenko and Wanner, 2000</u>). The respective primers are shown in <u>Supplementary Table 1</u>. The genes cpxA , cpxR , cpxRA , ackA*pta*, *acrB*, and *tolC* were replaced with a kanamycin (*kan*) resistance cassette in the S. Typhimurium strain CVCC541, generating mutant strains $S\Delta cpxA:: kan , S\Delta cpxR:: kan , S\Delta cpxRA: kan , S\Delta ackA-pta:: kan , S\Delta$ acrB:: kan, and JSA tolC:: kan, respectively. Using helper plasmid pCP20, expressing the FLP recombinase, the kan resistance cassette was excised from the $|S\Delta cpxA:: kan$, $|S\Delta cpxR:: kan$, $|S\Delta cpxRA:: kan$, $|S\Delta ackA-pta::$ kan , $|S\Delta \ acrB:: kan$, and $|S\Delta \ tolC:: kan$ mutants, generating mutant strains $S\Delta cpxA$, $S\Delta cpxR$, $S\Delta cpxRA$, $S\Delta ackA-pta$, $S\Delta acrB$, and $\Delta tolC$, respectively. Multiple additional mutants were then constructed from these mutants using the Red recombinase system. From the JS Δ cpxA mutant, additional mutant strains JS Δ *cpxA* Δ *ackA-pta* , JS Δ *cpxA* Δ *acrB* , and JS Δ $cpxA \Delta$ tolC were constructed. From the $|S\Delta cpxR|$ strain, additional mutants JS Δ *cpxR* Δ *acrB* and JS Δ *cpxR* Δ *tolC* were constructed. The main regulon

member deletion mutants JS Δ stm3031:: Kan , JS Δ htpX:: Kan , and JS Δ spy:: Kan were constructed from the wild-strain JS, and JS Δ cpxA Δ stm3031:: Kan , JS Δ cpxA Δ htpX:: Kan , and JS Δ cpxA Δ spy:: Kan were constructed from the JS Δ cpxA mutant. The mutants JS Δ acrD:: Kan , JS Δ cpxA Δ acrD:: Kan , JS Δ acrB Δ acrD:: Kan , and JS Δ cpxA Δ acrB Δ acrD:: Kan were constructed from the corresponding strain JS, JS Δ cpxA , JS Δ acrB , and JS Δ cpxA Δ acrB .

The JS $cpxR_{D51A}$, JS $cpxR_{M199A}$, JS $cpxA_{38}$, and JS $cpxA_{92-104}$ mutants were constructed in our previous study (Jing et al., 2020). Subsequently, mutant strains JS Δc $pxAcpxR_{D51A}$, JS Δc $pxAcpxR_{M199A}$, JS Δ $cpxA_{38}$ cpxR::kan, and JS Δ $cpxA_{92-104}$ cpxR:: kan were constructed from mutant strains JS $cpxR_{D51A}$, JSc pxR_{M199A} , JS $cpxA_{38}$, and JS $cpxA_{92-104}$, respectively, using the Red recombinase system.

Complementation of Various Mutants

To be able to construct the complemented strains of various *cpxA* mutants (JSA *cpxA*, JS *cpxA*₃₈, and JS *cpxA*_{92 - 104}), a strategy to construct the complemented strain in the bacterial chromosome was used according to our previous study (Jing et al., 2020). In order to distinguish *cpxA* complemented strains from wild strains, we changed the terminate code "UAA" of the *cpxA* gene in the wild strain to "UAG" in the *cpxA* complemented strains. First, the *sacBKan* fragment was amplified from plasmid psacBKan using the primers CpxA-H1P1/H2P2, and then integrated into the corresponding *cpxA* mutations locus using the Red recombinase system to obtain the intermediate strain. Positive clones were screened by PCR using primer pairs CpxA-F/R, CpxA-F/sacBKan-k2, sacBKan-k1/CpxA-R,

and sacBKan-k1/k2. Then, using primers CpxA-UF/CpxA-CL-UR and CpxA-CL-DF/CpxA-DR, the wild-type *cpxA* fragment with synonymous substitution of terminates codon was prepared on the *Bam* HI-digested vector pSTV28. The recombinogenic *cpxA* fragment was amplified using primer pair CpxA * -F/R. Finally, the recombinogenic *cpxA* fragment replaced the *sacBKan* cassette and the corresponding *cpxA* complemented strains (JSA *cpxA-CL*, JS *cpxA* 38 -*CL*, and JS *cpxA* 92 - 104 - *CL*) were confirmed by PCR.

The *acrB* and *tolC* complemented strains in the chromosome were constructed using the λ Red recombination system (<u>Datsenko and Wanner</u>, 2000). The corresponding PCR products were obtained using primer pairs AcrB-CL-F/AcrB-CL-R and TolC-CL-F/TolC-CL-F. To distinguish *acrB* complemented strains from wild strains, we changed the 884th codon " GTC" of the *acrB* gene in the wild strain to " GTG" in the *acrB* complemented strains. To distinguish *tolC* complemented strains from wild strains, we changed the terminate code " UGA" of the *tolC* gene in the wild strain to " UAA" in the *tolC* complemented strains. The corresponding PCR products were then used to replace the deletion mutations using the λ Red recombinase system. The complemented strains (JSA *acrB-CL* and JSA *tolC-CL*) were selected on LB agar plates containing cefuroxime (1 µg/ml) and confirmed by Sanger sequencing.

Real-Time Quantitative PCR (qPCR) Analysis of Bacterial Gene Expression Total RNA was isolated from bacterial cultures (OD $_{600} = 1.0$) using a TransZol UP Plus RNA Kit (TransGen Biotech, Beijing, China) according to the manufacturer's instructions. Aliquots (1 µg) of the total RNA were then transcribed into cDNA using a PrimeScript RT Reagent Kit with gDNA Eraser (Takara Bio., Kusatsu, Japan). Quantitative PCR (qPCR) assays were performed using the CFX96 Real-Time System and a C1000 Touch Thermal Cycler (Bio-Rad, Hercules, CA, United States), as well as a QuantiNova SYBR Green PCR Kit (Qiagen, Hilden, Germany). The primers used for qPCR analysis are outlined in <u>Supplementary Table 1</u>. The expression levels of all genes were normalized against that of internal reference gene *rpoD*, and relative fold changes in expression were calculated using the 2 $-\Delta\Delta CT$ method. The expression level of each mRNA in strain JS represents a one-fold change.

Antibiotic-Induced Activation Assays

Briefly, the test strains were grown overnight in LB medium, diluted 1: 100, and subcultured to the mid-exponential phase (OD ₆₀₀ of 0. 4 to 0. 6). When we used LB medium for the antibiotic activation test, it was found that the MICs values of the JS strain against amikacin and cefuroxime was 16 and 8 µg/ml in LB, respectively. So, to observe the relationship between the concentration of amikacin and cefuroxime with the activation level of the CpxRA system, the *S*. Typhimurium wild-type strain was grown in LB medium or LB medium with 4, 8, or 16 µg/ml amikacin or 1, 2, or 4 µg/ml cefuroxime to a stationary growth phase (OD ₆₀₀ ≥ 1. 0). To observe the influence of Δ *cpxA* or Δ *cpxR* cells were grown in LB medium with 16 µg/ml amikacin or 4 µg/ml cefuroxime to a stationary growth phase (OD ₆₀₀ ≥ 1. 0). The mRNA expression levels of the target genes *cpxP* and *degP* were used to measure the activation level of the CpxRA system.

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Antibiotic Susceptibility Testing

The MIC values of selected antibiotics against the various bacterial strains were measured in Mueller-Hinton (MH) medium (Hopebio, Qingdao, China) *via* the broth microdilution method as per the Clinical and Laboratory Standards Institute's (CLSI) guidelines (<u>Cockerill et al., 2012</u>). The tested antibiotics included amikacin (AMK), netilmicin (NET), streptomycin (STR), kanamycin (KAN), cefalotin (CEP), cefuroxime (CXM), ceftriaxone (CRO), and cefotaxime (CTX). All assays were conducted in triplicate.

Statistical Analysis

Statistical analyses were performed using GraphPad Prism 6. 0 Windows software and data were compared using the Student's t-test (GraphPad Software, United States). A probability (p) value of <0. 05 was considered statistically significant.

Results

cpxA and *cpxR* Deletion Mutants Have Opposing Roles in the Regulation of Resistance to AGAs and β-Lactams

As described in the introduction, *cpxA* deletion mutations sometimes show different resistance to amikacin. To investigate the effects of *cpxA* deletion mutations on the regulation of drug resistance, *cpxA* and *cpxR* deletion mutants – JS Δ *cpxA* and JS Δ *cpxR* – were generated from the *S* . Typhimurium strain JS. The complemented strain JS Δ *cpxA*-*CL* was also prepared. As shown in Table 2, compared with parental strain JS, two- to eight-fold increases in the MICs of the tested AGA and β -lactam antibiotics against the *cpxA* null mutant were observed; whereas two- to eight-fold decreases in the MICs of the tested AGA and β -lactam antibiotics against the *cpxR* deletion mutant were observed. The MICs of AGA and β -lactam antibiotics decreased by twoto eight-fold for the complemented strain JS Δ *cpxA-CL*, as compared to those for JS Δ *cpxA*. These results indicated that deletions of *cpxA* and *cpxR* have opposing roles in the regulation of AGAs and β -lactams resistance.

TABLE 2

Susceptibility of *S. enterica* serovar Typhimurium to several AGA and β -lactam antibiotics.

AGAs and β-Lactams Activate the CpxRA System

To evaluate whether the CpxRA system is activated by AGAs and β -lactams, a series of concentrations of amikacin (4, 8, or 16 µg/ml) and cefuroxime (1, 2, or 4 µg/ml) were added to the culture medium. The expression levels of *cpxP* and *degP*, two target genes of the CpxRA system (Fujimoto et al., 2018), were measured using a qRT-PCR assay to evaluate the activation level of the CpxRA system of the wild-type strain JS in the presence or absence of the corresponding antimicrobial. As shown in Figures 1A, B, the transcription levels of both *cpxP* and *degP* increased with increasing amikacin and cefuroxime concentrations. Whereas deletion of *cpxR* resulted in significantly reduced expression levels of *cpxP* and *degP* in the presence of amikacin and cefuroxime, compared with the wild-type strain JS (Figures 1C, D). The deletion of *cpxA* resulted in significantly increased expression levels of *degP* in presence of cefuroxime, and no significantly different expression levels of *degP* in presence of amikacin (Figures 1C, D),

compared with the wild-type strain JS. These results suggest that AGAs and β -lactams activate the *S*. Typhimurium CpxRA system dependent on CpxR; however, *cpxA* deletion can activate the CpxRA system independently of amikacin and cefuroxime.

FIGURE 1

CpxRA is activated by amikacin and cefuroxime.(A, B) *S* . Typhimurium wildtype strains were grown in LB medium or LB medium with 4, 8, and 16 µg/ml amikacin or 1, 2, and 4 µg/ml cefuroxime to the stationary growth phase (OD $_{600} \ge 1$. 0), and mRNA levels of CpxRA-regulated genes (*cpxP* and *degP*) were measured by reverse transcription-qPCR.(C, D) *S* . Typhimurium wildtype strains, the JS Δ *cpxA* , or the JS Δ *cpxR* mutant strain were grown in LB medium with 16 µg/ml amikacin or 4 µg/ml cefuroxime to the stationary growth phase (OD $_{600} \ge 1$. 0), and mRNA levels *cpxP* and *degP* were measured by reverse transcription-qPCR. Data were normalized to *rpoD* expression levels. Bars represent means ± standard deviations. *, *P* < 0. 05; **, *P* < 0. 01.

AckA-Pta-Dependent Activation of CpxR Increases Antibiotic Resistance

To investigate the underlying genetic basis of the increased resistance of the *cpxA* deletion mutant to AGAs and β -lactams, the first step was to confirm whether the increased resistance of the *cpxA* null mutant was caused by pleiotropic effects, given that CpxA can regulate downstream target genes by interacting with other regulators rather than CpxR (<u>Nakayama et al.</u>, 2003). For this reason, mutant strain JSA *cpxRA*, lacking both *cpxA* and

cpxR, was constructed. The antibiotic sensitivity of the double mutant was equivalent to that of strain JS Δ cpxR (<u>Table 2</u>). We also constructed point mutations in cpxR ($cpxR_{D51A}$, $cpxR_{M199A}$) and corresponding double mutations based on the *cpxA* deletion mutations (Δ *cpxAcpxR* _{D 51A}, Δ cpxAcpxR_{M199A}). Compared with strain JS, two- to four-fold decreases in the MICs of the tested AGA and β -lactam antibiotics for JS mutant $cpxR_{D51A}$ were observed, and there was no significant change in JS cpxR M 199A. Compared with strain $|S\Delta cpxA|$, 4- to 16-fold decreases in the MICs of the tested AGA and β -lactam antibiotics for mutant strains $|S\Delta cpxAcpxR_{D51A}|$ and $IS\Delta cpxAcpxR_{M199A}$ were observed (<u>Table 2</u>). These results demonstrated that the observed resistance of the *cpxA* deletion mutant to AGAs and β -lactams is dependent on CpxR and is not caused by pleiotropic effects.

CpxA acts as an autokinase, a kinase that activates CpxR, and a phosphatase (7, 8), and *cpxA* null mutations would therefore presumably disrupt all three of these activities. The lack of phosphatase activity leads to the phosphorylation of CpxR via alternative sources, such as acetyl phosphate produced by the AckA and Pta enzymes (14, 28). To investigate whether AckA and Pta participate in the regulation of AGAs and β -lactams resistance in the *cpxA* deletion mutant, double deletion mutant $IS\Delta$ ackA-pta and triple deletion mutant $S\Delta cpxA \Delta ackA-pta$ were constructed. Compared with $JS\Delta$ ackA-pta, no significant differences in the MICs of any of the tested antibiotics against strain $|S\Delta cpxA \Delta ackA-pta$ were observed. However, 2- to 16-fold decreases in the MICs of the tested antibiotics against JS $\Delta c pxA \Delta$ *ackA-pta* were observed compared with the MICs of $JS\Delta cpxA$ (<u>Table 2</u>). This https://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-to-

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result suggests that the increased resistance of the *cpxA* null mutant to AGA and β -lactam antibiotics is associated with AckA-Pta. We then evaluated the activation of the CpxRA system by measuring the expression levels of *cpxP* and *degP*. As shown in Figure 2A, compared with the wild-type strain, significant increases in mRNA levels of both genes were observed in strain JSA *cpxA*. Compared with strain JSA *cpxA*, significant decreases in the expression of both genes were observed in strains JSA *cpxRA*, JSA *cpxAcpxR D* 51A, JSA *cpxAcpxR M* 199A, and JSA *cpxA* Δ *ackA-pta*. It has been suggested

that the knock-out of *cpxA* could confer the resistance to the AGA and β lactam antibiotics. This regulatory mechanism is involved in the activation of the CpxRA system *via* phosphorylation of CpxR by the AckA-Pta pathway.

FIGURE 2

Relative mRNA levels of the CpxRA-regulated genes (*cpxP* and *degP*) were measured by real-time PCR.(A) Relative mRNA levels of the were measured in the WT *S* . Typhimurium JS strain and its isogenic JS Δ *cpxA* , JS Δ *cpxRA* , JS Δ *cpxAcpxR* ₅₁ , JS Δ *cpxAcpxR* ₁₉₉ , JS Δ *cpxA* Δ *acka-pta* , and JS Δ *acka-pta* . (B) Relative mRNA levels of the were measured in the WT *S* . Typhimurium JS strain and its isogenic JS *cpxA* ₃₈ , JS *cpxA* _{92 - 104} , JS *cpxA* ₃₈ Δ *cpxR:: kan* , JS Δ *cpxA* _{92 - 104} Δ *cpxR:: kan* . The expression level of each mRNA in strain JS represents one-fold. Data were normalized to *rpoD* expression levels. Bars represent means ± standard deviations. **, *P* < 0. 01.

AGAs and β-Lactams Resistance Can Be Conferred by CpxA-Mediated Ativation of CpxR

It has been confirmed that an in-frame deletion mutation that removed amino acids 92–104, or a site-directed mutation resulting in a Leu38Phe substitution in the periplasmic domain of CpxA, both constitutively activated the CpxRA system (collectively referred to as a cpx^* mutation) (Pogliano et al., 1998; Humphreys et al., 2004). Here, to determine whether the CpxAmediated activation of CpxR also confers AGAs and β -lactams resistance in S . Typhimurium, constitutively active mutants of cpxA (JS cpxA ₃₈ and JS cpxA $_{92-104}$) were constructed and the MICs of AGAs and β -lactams were measured. Moreover, the corresponding complemented strains IS cpxA 38 -CL and JS cpxA 92 - 104 - CL were constructed. As shown in Table 2, the MICs of $|S cpxA_{38}|$ and $|S cpxA_{92}| - 104$ were the same as that of the cpxA null mutant. Compared with strain JS cpxA 38 and JS cpxA 92 - 104, two- to eight-fold decreases in the MICs of the tested AGA and β -lactam antibiotics against the complemented strain JS cpxA 38 -CL and JS cpxA 92 - 104 -CL were observed. In the strains that do not produce cpxR (JS Δ cpxA ₃₈ Δ cpxR: kan , JS Δ cpxA ₉₂ $_{-104} \Delta cpxR; kan$), the MICs of all of the drugs except for kanamycin were the same as that of the cpxR deletion mutant (JS $\Delta cpxR$). As shown in Figure <u>2B</u>, compared with the wild-type strain, significant increases in mRNA levels of both target genes cpxP and degP were observed in strains IS cpxA 38 and JS $cpxA_{92-104}$, whereas significant decreases in the expression of both genes were observed in strains IS $cpxA_{38} \Delta c pxR$: kan and IS $cpxA_{92} - 104 \Delta c$ pxR: kan compared with strains JS cpxA 38 and JS cpxA 92 - 104. These results demonstrate that resistance to AGAs and β -lactams can be conferred by

activated CpxA, which is dependent on the phosphorylation of CpxR. https://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-toaminoglycosides-and-lactams-in-salmonella-enterica-serovar-typhimurium/

CpxRA System Can Modulate Resistance to AGAs and β-Lactams Independent of the AcrAB-TolC Efflux Pump

Drug efflux mechanisms are ubiguitous among Gram-negative bacteria and contribute significantly to MDR. Among the mechanisms reported to date, the AcrAB-TolC efflux pump plays a significant role in resistance to various antibiotics (<u>Baucheron et al., 2004</u>; <u>Du et al., 2014</u>; <u>Li et al., 2015</u>). To investigate whether AcrAB-ToIC affects CpxRA-mediated regulation of drug resistance, we constructed several double and triple deletion mutant strains from single mutants $|S\Delta cpxA|$, $|S cpxA|_{38}$, $|S cpxA|_{92-104}$, $|S\Delta cpxR|$, $|S\Delta$ acrB, and ISA toIC, as shown in Table 2. Compared with strain IS, 2- to 32fold decreases in the MICs of all of the drugs except for netilmicin, kanamycin and ceftriaxone against strain $IS\Delta$ acrB were observed. Meanwhile, the MICs of all the tested antibiotics against complemented strain $|S\Delta \ acrB-CL$ were the same as strain |S. Compared with strain $|S\Delta \ acrB$, two- to four-fold increases in the MICs of all of the drugs against strain JSΔ $cpxA \Delta acrB$, |S $cpxA_{38} \Delta acrB$, and |S $CpxA_{92-104} \Delta acrB$ were observed, whereas two- to four-fold decreases in the MICs of all of the drugs except for cefalotin against strain JS Δ *cpxR* Δ *acrB* were observed. Compared with strain IS, 2- to 32-fold decreases in the MICs of all of the drugs except for netilmicin and ceftriaxone against strain $IS\Delta$ to/C were observed. Meanwhile, the MICs of all the tested antibiotics against complemented strain ISA toIC-CL were the same as strain JS. Compared with strain JS Δ tolC, two- to eight-fold increases in the MICs of all of the drugs against strains $JS\Delta cpxA \Delta tolC$, JS*cpxA* ₃₈ Δ *tolC*, and |S *cpxA* _{92 - 104} Δ *tolC* were observed, whereas two- to eight-fold decreases in the MICs of all of the drugs except for cefalotin against strain JSA $cpxR \Delta$ tolC were observed (<u>Table 2</u>). As shown in Figure https://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-toaminoglycosides-and-lactams-in-salmonella-enterica-serovar-typhimurium/

3., the expression levels of *acrB*, *tolC*, and the transcription factor genes *marA*, *soxS*, and *ramA* were measured in strains JS, JS Δ *cpxA*, JS *cpxA*₃₈, JS *cpxA*₉₂₋₁₀₄, and JS Δ *cpxR*. In all *cpxA* mutants, a significant decrease in the expression levels of *marA* and *soxS* was observed, along with a significant increase in the expression level of *ramA*, and no significant difference in the expression levels of *acrB* and *tolC* were observed compared with strain JS. We then constructed mutants JS Δ *ramA:: kan* and JS Δ *cpxA* Δ *ramA:: kan*. Compared with strain JS Δ *ramA:: kan*, two- to four-fold increases in the MICs of all of the drugs except for kanamycin against strain JS Δ *cpxA* Δ *ramA:: kan* were observed. These results demonstrate that AcrB and TolC confer resistance to some AGAs and β -lactams, but both activation and inactivation of the CpxRA system can modulate resistance to AGAs and β -lactams in both an *acrB* or *tolC* background and a Δ *acrB* or Δ *tolC* background.

FIGURE 3

The influence of various *cpxA* mutations on the expression levels of the efflux pump AcrAB-TolC.(A) Relative mRNA expression levels of the efflux pump genes *acrB* and *tolC*.(B) Relative mRNA expression levels of the transcription factor genes *marA*, *soxS*, and *ramA*. The expression level of each mRNA in strain JS represents one-fold. Data were normalized to *rpoD* expression levels. Bars represent means ± standard deviations. **, *P* < 0. 01.

Effects of Different CpxA Mutations on the Expression Levels of Cpx Regulons

The CpxRA system can alter the expression levels of a series of published Cpx regulon members to mediate drug resistance, such as those encoding outer membrane proteins OmpF, OmpC, OmpD, OmpW, and STM3031; efflux pump AcrD; proteases PpiA and HtpX; protein folding factors Spy and YccA, and respiration-related proteins CyoA, NuoA, and SdhC (Hu et al., 2011; Mahoney and Silhavy, 2013; Raivio et al., 2013; Raivio, 2014; Huang et al., 2016). To identify whether different *cpxA* mutations have a consistent effect on the expression levels of these Cpx regulons, we detected the relative mRNA expression levels of these genes. As shown in Figure 4, compared with strain IS, the mRNA expression levels of *ompW*, *acrD*, *spy*, *ycca*, *ppia* , *htpX* , and *stm3031* in all three *cpxA* mutants changed significantly (P < 0. 01 or P < 0.05). In detail, the mRNA expression level of *ompW* was decreased, whereas the expression levels of acrD, spy, ycca, ppia, htpX, and stm3031 were increased. The mRNA expression levels of ompD, cyoA, and *acrF* showed no significant differences in these three *cpxA* mutants. The mRNA expression levels of *ompF* and *ompC* were increased in JS Δ *cpxA* and $|S cpxA_{92} - 104|$, but showed no significant difference in $|S cpxA_{38}|$. The mRNA expression levels of *nuoA* and *sdhC* were decreased in JS $cpxA_{92-104}$, but showed no significant difference in $|S\Delta cpxA|$ and $|S cpxA|_{38}$. In mutant strain JS Δ cpxR, significant decreases (P < 0.01 or P < 0.05) in mRNA expression were only seen for *spy*, *ycca*, *ppia*, *htpX*, and *stm3031*. Then, we individually constructed null mutations in sharply upregulated genes stm3031, htpX, and spy in a cpxA -deficient background and WT (JSA) stm3031:: kan , $|S\Delta htpX:: kan , |S\Delta spy:: kan , |S\Delta cpxA \Delta stm3031:: kan ,$ https://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-toaminoglycosides-and-lactams-in-salmonella-enterica-serovar-typhimurium/

JS Δ *cpxA* Δ *htpX:: kan*, and JS Δ *cpxA* Δ *spy:: kan*). However, compared with JS Δ *cpxA*, none of these genes showed a decrease in the resistance conferred by the *cpxA* deletion mutation (<u>Table 2</u>). To further study the impact of efflux pumps on the resistance conferred by the *cpxA* deletion mutation, we also constructed some corresponding *acrD* null mutations (JS Δ *acrD:: Kan*, JS Δ *cpxA* Δ *acrD:: Kan*, JS Δ *acrB* Δ *acrD:: Kan*, and JS Δ *cpxA* Δ *acrB* Δ *acrD:: Kan*). As shown in <u>Table 2</u>, the MICs of JS Δ *acrD:: Kan* were the same as that of the JS. Compared with JS Δ *cpxA*, the MICs of JS Δ *cpxA* Δ *acrD:: Kan* were the same as that of the *cpxA* null mutant. Compared with *JS* Δ *acrB* Δ *acrD:: Kan*, two- to four-fold increases in the MICs of all of the drugs except for kanamycin against strain JS Δ *cpxA* Δ *acrB* Δ *acrD:: Kan*.

FIGURE 4

Relative mRNA levels of a series of MDR-related genes were measured by real-time PCR.(A) Relative mRNA expression levels of the outer membrane proteins genes *ompF*, *ompC*, *ompW*, and *ompD*;(B) Relative mRNA expression levels of protease genes *ppiA* and *htpX* and protein folding factors genes *spy* and *yccA*;(C) Relative mRNA expression levels of the genes involved in respiration *cyoA*, *nuoA*, and *sdhC*;(D) Relative mRNA expression levels of the efflux pumps genes *acrD*, *mdtA*, and *acrF*;(E) Relative mRNA expression levels of the outer membrane proteins genes STM3031; The expression level of each mRNA in strain JS represents onefold. Data were normalized to *rpoD* expression levels. Bars represent means \pm standard deviations. *, *P* < 0. 05; **, *P* < 0. 01.

Discussion

In this study, all of the tested *cpxA* mutants of *S*. Typhimurium were found to confer resistance to AGAs and β-lactams. The previous studies showed the strain $cpxA * (IS cpxA_{92-104})$ was more resistant to amikacin than its wildtype strain SL1344, but the *cpxA* deletion mutant was more sensitive to amikacin than its parent strain (Humphreys et al., 2004). Such contradictory findings suggested that although *cpxA* deletion can active the CpxRA system and display numerous phenotypes, whether null mutations of *cpxA* regulating resistance are caused solely by hyperphosphorylation of CpxR in S. Typhimurium still needs to be determined. On the basis of studies involving *cpxR* deletion and site-directed mutagenesis, we ruled out the pleiotropic effects of CpxA and provided evidence that the increased resistance of *cpxA* deletion mutants to AGAs and β -lactams is directly mediated by CpxR-P, which is activated *via* the acetyl phosphate pathway. This finding is consistent with previous reports showing that the deletion of cpxA eliminates its phosphatase activity, causing CpxR to accept phosphate groups from acetyl phosphate generated by Acka and Pta, activating the CpxRA system and contributing to some phenotypic changes (<u>Danese and</u> Silhavy, 1998; Buelow and Raivio, 2005; Lima et al., 2012).

JS Δ *cpxR* showed no significant differences in the mRNA expression levels of the major tested genes in the absence of antibiotics, which is consistent with a previous report in *S*. Typhimurium (<u>Huang et al., 2016</u>). It is also consistent with a previous study in *E. coli* that showed that a *cpxR* mutant had few differentially regulated genes compared with wild-type uropathogenic *E. coli*, which indicated that the CpxRA system is only https://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-to-

nttps://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-toaminoglycosides-and-lactams-in-salmonella-enterica-serovar-typhimurium/ minimally active in the wild-type strain under the experimental growth conditions (Dbeibo et al., 2018). This raises the question of why the wild-type strain shows more resistance to AGAs and β -lactams. Here, we confirmed that amikacin and cefuroxime can activate the CpxRA system and that the activated CpxRA system increases the resistance to amikacin and cefuroxime. Therefore, it is proposed that the presence of antibiotics activates the CpxRA response system in wild-type strains, which in turn, leads to an increase in drug resistance compared with strain JS Δ *cpxR*. The results also suggest that the phenotypic changes caused by *cpxR* deletion mainly depend on whether the CpxRA pathway can be activated by a corresponding external stimulation.

In Gram-negative bacteria, RND-family multidrug efflux pumps play the most prominent roles in drug resistance (Poole, 2001; Becker and Cooper, 2013; Du et al., 2018). Among the RND efflux pumps, AcrD is mainly responsible for resistance to the hydrophilic class of drugs such as AGAs in *E. coli* (Rosenberg et al., 2000). Moreover, some reports suggest that this efflux pump confers resistance to some other compounds, such as tetracycline, novobiocin, norfloxacin, Fosfomycin, and some β -lactams in *E. coli* (Nishino and Yamaguchi, 2001; Nishino et al., 2003). By contrast, AcrB confers resistance to practically all types of antibacterial agents, except aminoglycosides in *E. coli* (Elkins and Nikaido, 2002; Li et al., 2015). In the current study, a significant increase in the expression level of *acrD* was observed in the *cpxA* mutants, which is consistent with previous reports and supports the important role of the AcrAD-TolC efflux pump in resistance regulation of the CpxRA system (Hu et al., 2011; Huang et al., 2016). However, the *acrD* deletion mutation did not decrease the resistance conferred by the *cpxA* deletion mutation, which is consistent with previous

conferred by the *cpxA* deletion mutation, which is consistent with previous research and suggests that no single factor is necessary of the resistance to β-lactams and AGAs conferred by the *cpxA* mutation (<u>Mahoney and Silhavy</u>, 2013). AcrAB-TolC contributes to the resistance to some aminogly cosides (amikacin and streptomycin) in *S. enterica* serovar Typhimurium JS, which is consistent with the previous result where the deletion of *acrB* decreased resistance to aminoglycosides (amikacin and neomycin) (Huang et al., 2016). To our knowledge, it is unclear why the mutant strain lacking *acrB* is more susceptible to some aminoglycosides in *S. enterica* serovar Typhimurium JS and a further study is needed. We did not find any significant change in the mRNA expression levels of *acrB* and *tolC*, this differed from a previous report showing that overexpression of *cpxR* results in a significant decrease in the mRNA expression levels of *acrB* and *tolC* compared with the wild-type strain JS (<u>Huang et al., 2016</u>). This difference may be explained by the paradigm that, for the tested global regulatory factors, the three cpxA mutations all led to a significant reduction in the levels of marA and soxS, thereby downregulating the expression levels of the AcrAB-TolC efflux pump, but led to a significant increase in the levels of *ramA*, thereby upregulating the expression of the AcrAB-TolC efflux pump. Moreover, in the *acrB* and *tolC* deletion mutants, which showed decreased resistance to AGAs and β lactams, both activation and inactivation of the CpxRA system could modulate resistance. To investigate further, we constructed strain $|S\Delta cpxA\Delta|$ ramA:: kan and JSA ramA:: kan . Compared with strain JSA ramA:: kan , the MICs of the tested antibiotics also showed an increase. These results suggest

that the efflux pump AcrAB-ToIC does not play a decisive role in the CpxRAmediated AGAs and β -lactams resistance of *S*. Typhimurium.

Membrane proteases is one of the important mechanisms of AGAs resistance (<u>Becker and Cooper, 2013</u>). When bacteria are exposed to AGAs, there is an increase in the abundance of misfolded and mistargeted proteins within cells. Whereafter, genes that are involved in protein turnover are upregulated, as well as those encoding membrane proteases. If the membrane proteases cannot keep up with the increasing number of faulty proteins, their surplus accumulation eventually destroys the integrity of the membrane and kills the bacteria (<u>Magnet and Blanchard, 2005</u>; <u>Becker and</u> <u>Cooper, 2013</u>). In this study, the selected protease genes, *ppiA* and *htpX*, and the protein folding factor genes, *spy* and *yccA*, were all upregulated in various *cpxA* mutants, suggesting that the function of maintaining membrane integrity of the CpxRA envelope stress response system, is important for the resistance to AGAs.

One theory proposes that all bactericidal antibiotics may act through a common mechanism involving the production of reactive oxygen species (ROS), which is dependent on metabolism-related NADH depletion and the electron transport chain (<u>Van Acker and Coenye, 2017</u>). It has been shown that *cpxA* or *cpxR* deletion mutations confer antibiotic resistance by reducing the production of ROS (<u>Kohanski et al., 2008</u>). A study also demonstrated that compared with the parent strain, the expression levels of genes encoding succinate dehydrogenase (*sdh*), NADH dehydrogenase (*nuo*), and cytochrome oxidase (*cyo*) were significantly downregulated under

multiple conditions in the transient NIpE overexpression strain, and the elimination of these genes (*cyoA*, *nuoA*, and *sdhC*) conferred resistance to amikacin (Raivio et al., 2013). However, we found that this common factor was inconsistent among the tested *cpxA* mutants. The mRNA expression levels of *nuoA* and *sdhC* were only found to be decreased in strain JS *cpxA* 92 – 104 , with no significant differences detected in strains JS Δ *cpxA* and JS *cpxA* 38 . These results suggested that the decrease in ROS may explain the accompanied tolerance to aminoglycosides of the activated CpxRA system in certain *cpxA* mutants, particularly JS *cpxA* 92 – 104 . Furthermore, the factors involved in drug resistance regulation appear to differ between the tested *cpxA* mutants.

Conclusion

In conclusion, this study analyzed the mechanism involved in the CpxAmediated regulation of resistance to AGAs and β -lactams in *S*. Typhimurium. It was revealed that various *cpxA* mutations show the same resistance phenotype relying on phosphorylated CpxR. These findings broaden our understanding of the complex regulatory network governing CpxRAmediated antibiotic resistance.

Data Availability Statement

The original contributions presented in the study are included in the article/ <u>Supplementary Material</u>, further inquiries can be directed to the corresponding author/s. WJ, JL, and SW designed the experiment. YL was responsible for funding acquisition and project supervision. WJ, XL, and YL contributed to manuscript writing. All authors contributed to the article and approved the submitted version.

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Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary Material

The Supplementary Material for this article can be found online at:

https://www. frontiersin. org/articles/10. 3389/fmicb. 2021.

604079/full#supplementary-material

Supplementary Table 1 | Primers used in this study.

References

Baucheron, S., Tyler, S., Boyd, D., Mulvey, M. R., Chaslus-Dancla, E., and Cloeckaert, A. (2004). AcrAB-TolC directs efflux-mediated multidrug resistance in *Salmonella enterica* serovar typhimurium DT104. *Antimicrob. Agents Chemother.* 48, 3729–3735. doi: 10. 1128/AAC. 48. 10. 3729-3735. 2004

Becker, B., and Cooper, M. A. (2013). Aminoglycoside antibiotics in the 21st century. *ACS Chem. Biol.* 8, 105–115. doi: 10. 1021/cb3005116

Buelow, D. R., and Raivio, T. L. (2005). Cpx signal transduction is influenced by a conserved N-terminal domain in the novel inhibitor CpxP and the periplasmic protease DegP. *J. Bacteriol.* 187, 6622–6630. doi: 10. 1128/JB. 187. 19. 6622-6630. 2005

Cockerill, F., Wikler, M., Alder, J., Dudley, M., Eliopoulos, G., Ferraro, M., et al. (2012). Methods for dilution antimicrobial susceptibility tests for bacteria that grow aerobically: approved standard. *Clin. Lab. Stand. Inst.* 32, M07– M09.

Danese, P. N., and Silhavy, T. J. (1998). CpxP, a stress-combative member of the Cpx regulon. *J. Bacteriol.* 180, 831–839.

Datsenko, K. A., and Wanner, B. L. (2000). One-step inactivation of chromosomal genes in *Escherichia coli* K-12 using PCR products. *Proc. Natl. Acad. Sci. U. S. A.* 97, 6640–6645. doi: 10. 1073/pnas. 120163297 Dbeibo, L., van Rensburg, J. J., Smith, S. N., Fortney, K. R., Gangaiah, D., Gao, H., et al. (2018). Evaluation of CpxRA as a Therapeutic Target for Uropathogenic *Escherichia coli* Infections. *Infect. Immun.* 86, 798–717e. doi: 10. 1128/IAI. 00798-717

De la Cruz, M. A., Perez-Morales, D., Palacios, I. J., Fernandez-Mora, M., Calva, E., and Bustamante, V. H. (2015). The two-component system CpxR/A represses the expression of *Salmonella* virulence genes by affecting the stability of the transcriptional regulator HilD. *Front. Microbiol.* 6: 807. doi: 10. 3389/fmicb. 2015. 00807

De Wulf, P., McGuire, A. M., Liu, X., and Lin, E. C. (2002). Genome-wide profiling of promoter recognition by the two-component response regulator CpxR-P in *Escherichia coli* . *J. Biol. Chem.* 277, 26652–26661. doi: 10. 1074/jbc. M203487200

Delhaye, A., Collet, J. F., and Laloux, G. (2016). Fine-Tuning of the Cpx Envelope Stress Response Is Required for Cell Wall Homeostasis in *Escherichia coli* . *mBio* 7, 47–16e. doi: 10. 1128/mBio. 00047-16

DiGiuseppe, P. A., and Silhavy, T. J. (2003). Signal detection and target gene induction by the CpxRA two-component system. *J. Bacteriol.* 185, 2432–2440. doi: 10. 1128/jb. 185. 8. 2432-2440. 2003

Du, D., Wang-Kan, X., Neuberger, A., van Veen, H. W., Pos, K. M., Piddock, L. J. V., et al. (2018). Multidrug efflux pumps: structure, function and regulation. *Nat. Rev. Microbiol.* 16, 523–539. doi: 10. 1038/s41579-018-0048-46

Du, D., Wang, Z., James, N. R., Voss, J. E., Klimont, E., Ohene-Agyei, T., et al. (2014). Structure of the AcrAB-TolC multidrug efflux pump. *Nature* 509, 512– 515. doi: 10. 1038/nature13205

Elkins, C. A., and Nikaido, H. (2002). Substrate specificity of the RND-type multidrug efflux pumps AcrB and AcrD of *Escherichia coli* is determined predominantly by two large periplasmic loops. *J. Bacteriol.* 184, 6490–6498. doi: 10. 1128/jb. 184. 23. 6490-6499. 2002

Fujimoto, M., Goto, R., Haneda, T., Okada, N., and Miki, T. (2018). *Salmonella enterica* Serovar Typhimurium CpxRA Two-Component System Contributes to Gut Colonization in *Salmonella* -Induced Colitis. *Infect. Immun.* 86, 280–218e. doi: 10. 1128/IAI. 00280-218

Gotoh, Y., Eguchi, Y., Watanabe, T., Okamoto, S., Doi, A., and Utsumi, R. (2010). Two-component signal transduction as potential drug targets in pathogenic bacteria. *Curr. Opin. Microbiol.* 13, 232–239. doi: 10. 1016/j. mib. 2010. 01. 008

Hirakawa, H., Nishino, K., Hirata, T., and Yamaguchi, A. (2003a). Comprehensive studies of drug resistance mediated by overexpression of response regulators of two-component signal transduction systems in *Escherichia coli*. *J. Bacteriol.* 185, 1851–1856.

Hirakawa, H., Nishino, K., Yamada, J., Hirata, T., and Yamaguchi, A. (2003b).
Beta-lactam resistance modulated by the overexpression of response
regulators of two-component signal transduction systems in *Escherichia coli*. *J. Antimicrob. Chemother.* 52, 576–582. doi: 10. 1093/jac/dkg406

https://assignbuster.com/role-of-cpxa-mutations-in-the-resistance-toaminoglycosides-and-lactams-in-salmonella-enterica-serovar-typhimurium/ Hu, W. S., Chen, H. W., Zhang, R. Y., Huang, C. Y., and Shen, C. F. (2011). The expression levels of outer membrane proteins STM1530 and OmpD, which are influenced by the CpxAR and BaeSR two-component systems, play important roles in the ceftriaxone resistance of *Salmonella enterica* serovar Typhimurium. *Antimicrob. Agents Chemother.* 55, 3829–3837. doi: 10. 1128/AAC. 00216-211

Huang, H., Sun, Y., Yuan, L., Pan, Y., Gao, Y., Ma, C., et al. (2016). Regulation of the Two-Component Regulator CpxR on Aminoglycosides and beta-lactams Resistance in *Salmonella enterica* serovar Typhimurium. *Front. Microbiol.* 7: 604. doi: 10. 3389/fmicb. 2016. 00604

Humphreys, S., Rowley, G., Stevenson, A., Anjum, M. F., Woodward, M. J., Gilbert, S., et al. (2004). Role of the two-component regulator CpxAR in the virulence of *Salmonella enterica* serotype Typhimurium. *Infect. Immun.* 72, 4654–4661. doi: 10. 1128/IAI. 72. 8. 4654-4661. 2004

Isaac, D. D., Pinkner, J. S., Hultgren, S. J., and Silhavy, T. J. (2005). The extracytoplasmic adaptor protein CpxP is degraded with substrate by DegP. *Proc. Natl. Acad. Sci. U. S. A.* 102, 17775–17779. doi: 10. 1073/pnas. 0508936102

Jing, W., Liu, J., Wu, S., Chen, Q., Li, X., and Liu, Y. (2020). Development of a Method for Simultaneous Generation of Multiple Genetic Modification in *Salmonella enterica* Serovar Typhimurium. *Front. Genet.* 11: 563491. doi: 10. 3389/fgene. 2020. 563491 Kohanski, M. A., Dwyer, D. J., Wierzbowski, J., Cottarel, G., and Collins, J. J. (2008). Mistranslation of membrane proteins and two-component system activation trigger antibiotic-mediated cell death. *Cell* 135, 679–690. doi: 10. 1016/j. cell. 2008. 09. 038

Levy, S. B. (2002). The 2000 Garrod lecture. *Factors impacting on the problem of antibiotic resistance*. *J. Antimicrob. Chemother.* 49, 25–30. doi: 10. 1093/jac/49. 1. 25

Li, X. Z., Plesiat, P., and Nikaido, H. (2015). The challenge of efflux-mediated antibiotic resistance in Gram-negative bacteria. *Clin. Microbiol. Rev.* 28, 337-418. doi: 10. 1128/CMR. 00117-114

Lima, B. P., Thanh Huyen, T. T., Basell, K., Becher, D., Antelmann, H., and Wolfe, A. J. (2012). Inhibition of acetyl phosphate-dependent transcription by an acetylatable lysine on RNA polymerase. *J. Biol. Chem.* 287, 32147–32160. doi: 10. 1074/jbc. M112. 365502

Magnet, S., and Blanchard, J. S. (2005). Molecular insights into aminoglycoside action and resistance. *Chem. Rev.* 105, 477–498. doi: 10. 1021/cr0301088

Mahoney, T. F., and Silhavy, T. J. (2013). The Cpx stress response confers resistance to some, but not all, bactericidal antibiotics. *J. Bacteriol.* 195, 1869–1874. doi: 10. 1128/JB. 02197-2112

Masi, M., Pinet, E., and Pages, J. M. (2020). Complex Response of the CpxAR Two-Component System to beta-Lactams on Antibiotic Resistance and

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Envelope Homeostasis in *Enterobacteriaceae*. *Antimicrob. Agents Chemother.* 64: 32229490. doi: 10. 1128/AAC. 00291-220

Nakayama, S., Kushiro, A., Asahara, T., Tanaka, R., Hu, L., Kopecko, D. J., et al. (2003). Activation of hilA expression at low pH requires the signal sensor CpxA, but not the cognate response regulator CpxR, in *Salmonella enterica* serovar Typhimurium. *Microbiology* 149, 2809–2817. doi: 10. 1099/mic. 0. 26229-26220

Nishino, K., Yamada, J., Hirakawa, H., Hirata, T., and Yamaguchi, A. (2003). Roles of TolC-dependent multidrug transporters of *Escherichia coli* in resistance to beta-lactams. *Antimicrob. Agents Chemother.* 47, 3030–3033. doi: 10. 1128/aac. 47. 9. 3030-3033. 2003

Nishino, K., and Yamaguchi, A. (2001). Analysis of a complete library of putative drug transporter genes in *Escherichia coli* . *J. Bacteriol.* 183, 5803-5812. doi: 10. 1128/JB. 183. 20. 5803-5812. 2001

Nishino, K., Yamasaki, S., Hayashi-Nishino, M., and Yamaguchi, A. (2010). Effect of NIpE overproduction on multidrug resistance in *Escherichia coli* . *Antimicrob. Agents Chemother.* 54, 2239–2243. doi: 10. 1128/AAC. 01677-1679

Padilla-Vaca, F., Mondragon-Jaimes, V., and Franco, B. (2017). General Aspects of Two-Component Regulatory Circuits in Bacteria: Domains. *Sign. Roles. Curr. Prot. Pept. Sci.* 18, 990–1004. doi: 10.

2174/1389203717666160809154809

Pogliano, J., Dong, J. M., De Wulf, P., Furlong, D., Boyd, D., Losick, R., et al. (1998). Aberrant cell division and random FtsZ ring positioning in *Escherichia coli* cpxA * mutants. *J. Bacteriol.* 180, 3486–3490. doi: 10. 1128/JB. 180. 13. 3486-3490. 1998

Poole, K. (2001). Multidrug resistance in Gram-negative bacteria. *Curr. Opin. Microbiol.* 4, 500–508.

Quinn, T., O'Mahony, R., Baird, A. W., Drudy, D., Whyte, P., and Fanning, S. (2006). Multi-drug resistance in *Salmonella enterica* : efflux mechanisms and their relationships with the development of chromosomal resistance gene clusters. *Curr. Drug Targets* 7, 849–860.

Raivio, T. L. (2014). Everything old is new again: an update on current research on the Cpx envelope stress response. *Biochim. Biophys. Acta* 1843, 1529–1541. doi: 10. 1016/j. bbamcr. 2013. 10. 018

Raivio, T. L., Leblanc, S. K., and Price, N. L. (2013). The *Escherichia coli* Cpx envelope stress response regulates genes of diverse function that impact antibiotic resistance and membrane integrity. *J. Bacteriol.* 195, 2755–2767. doi: 10. 1128/JB. 00105-113

Raivio, T. L., and Silhavy, T. J. (1997). Transduction of envelope stress in *Escherichia coli* by the Cpx two-component system. *J. Bacteriol.* 179, 7724–7733.

Rosenberg, E. Y., Ma, D., and Nikaido, H. (2000). AcrD of *Escherichia coli* is an aminoglycoside efflux pump. *J. Bacteriol.* 182, 1754–1756. Tiwari, S., Jamal, S. B., Hassan, S. S., Carvalho, P., Almeida, S., Barh, D., et al. (2017). Two-Component Signal Transduction Systems of Pathogenic Bacteria As Targets for Antimicrobial Therapy: An Overview. *Front. Microbiol.* 8: 1878. doi: 10. 3389/fmicb. 2017. 01878

Van Acker, H., and Coenye, T. (2017). The Role of Reactive Oxygen Species in Antibiotic-Mediated Killing of Bacteria. *Trends Microbiol.* 25, 456–466. doi: 10. 1016/j. tim. 2016. 12. 008

Wolfe, A. J., Parikh, N., Lima, B. P., and Zemaitaitis, B. (2008). Signal integration by the two-component signal transduction response regulator CpxR. *J. Bacteriol.* 190, 2314–2322. doi: 10. 1128/JB. 01906-1907