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## ORIGINAL RESEARCH The role of cfa gene in ampicillin tolerance in Shigella

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Background: Bacterial dysentery is an intestinal infectious disease caused by Shigella. The resistance of Shigella species to ampicillin has increased rapidly. Besides resistance, bacteria in a state of tolerance to antibiotics can also lead to the failure of infectious diseases therapy. Purpose: The genetic mechanism of antibiotic tolerance remains largely unexplored. The current study aimed to investigate the mechanisms of antibiotic tolerance and to provide novel strategies for the prevention of drug resistance of Shigella.

Methods: We exposed Shigella to lethal doses of ampicillin to obtain tolerant strain. The tolerant strain was sequenced to screen non-single-nucleotide polymorphisms and Indels. We also quantitated the relative expression of gene by RT-PCR.

**Results:** There was one nonsynonymous mutation in the 2252304 loci of the *cfa* gene (G to A/Val to Met) and 10 Indels in the noncoding regions of different genes. The expression of the cfa gene was 7.56-fold higher in the tolerant strain than in the wild-type strain.

**Conclusion:** Our results showed that *Shigella* could be tolerated to ampicillin, and the *cfa* gene might be associated with antibiotic tolerance by increasing its expression. Our data suggest that cfa gene might be a target for overcoming drug tolerance, delaying the occurrence of drug resistance to some extent.

Keywords: Shigella, tolerance, cfa, gene mutation

### Introduction

Bacillary dysentery is a prevailing acute intestinal infectious disease characterized by Shigella bacteria invading colon mucosa and inflammatory of epithelial cells.<sup>1</sup> Approximately 163 million cases of shigellosis were reported annually, particularly in developing countries. In China, its incidence rate accounts for the top three infectious diseases and is still the prevention and control focus of public health.<sup>2</sup> Ampicillin is a broad-spectrum antibiotic that is used to treat Enterobacteriaceae bacteria in humans as well as in animals.<sup>3</sup> Due to the extensive abuse of ampicillin, the resistance of Shigella species to ampicillin has increased rapidly, making it more difficult to control bacillary dysentery.<sup>3</sup>

Besides resistance, bacteria in a state of tolerance to certain antibiotics can also lead to the failure of infectious disease therapy.<sup>4</sup> Tolerance can be defined as bacteria that grow slowly or even do not grow after short exposure to high concentrations of antibiotics, and then grow again after antibiotics are discontinued.<sup>5</sup> However, unlike resistance, the tolerant bacteria can have the same minimum inhibitory concentration (MIC); thus, it is not informative as a metric to evaluate tolerance.<sup>6</sup> In recent years, the minimum duration of killing (MDK) is used to evaluate tolerance,<sup>7</sup> for example, the minimum duration of treatment that is required to kill 99% of bacteria (MDK<sub>99</sub>). Till now, the mechanism of antibiotic

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tolerance is quite complicated and remains largely unclear. Changes in environmental conditions of bacteria often trigger the production of stress-response genes. Studies have shown that antibiotic tolerance may be caused by stress-response genes controlled by *rpoS*.<sup>8</sup> Second messenger guanosine tetraphosphate (ppGpp) can lead to antibiotic tolerance.<sup>9</sup> In addition, Aaron et al have shown that strains are isolated from patients with cystic fibrosis patients, which are biofilm bacteria that are tolerant to antibiotics, compared to plankton.<sup>10</sup>

The genetic mechanism of antibiotic tolerance has not been elucidated. To explore the mechanisms of antibiotic tolerance, in our study, we exposed *Shigella* to high concentrations of ampicillin to obtain tolerant strain. The tolerant strain was sequenced to screen non-single-nucleotide polymorphisms (SNPs) and Indels. Our data suggest that the *cfa* gene may be associated with antibiotic tolerance by increasing its expression and it might be a target for overcoming drug tolerance.

### Materials and methods Isolation and identification of bacteria

To study antibiotic tolerance of Shigella strain sensitive to ampicillin, we separated a strain sensitive to ampicillin that was originally isolated from Sui County in Henan province in 2014. This strain was confirmed by biochemical and serological identification. Biochemical identification was performed using API20E (API Systems, SA, Vercieu, France). Briefly, the strain was inoculated into Luria-Bertani (LB) agar using a sterile-inoculating loop and incubated at 37°C for 18 hrs. The colonies were picked up with sterile cotton and resuspended in 0.85% NaCl medium to achieve uniform bacterial turbidity, and the turbidity of the bacterial suspension was adjusted to 0.5 McFarland. According to the instructions of the identification manual, the bacterial resuspension was added to the tube of the test strip using a straw. The test strips of citrate utilization (CIT), acetoin production(VP), and gelatinase (GEL) should be filled with bacterial suspension. The test strips of arginine dihydrolase(ADH), lysine decarboxylase (LDC),ornithine decarboxylase (ODC),H2S production(H2S),urease(URE) should be filled with mineral oil to form an anaerobic environment. Sterile distilled water was added to the culture box, and the test strip was placed in a culture box and placed in a 37°C incubator for 18 hrs. The tryptophan deaminase (TDA) test needed to add 1 drop of TDA reagent. The indole production (IND) test needed to add a drop of JAMES reagent. The VP test needed to add a drop of VP1 and VP2. The test results were observed after 10 mins. The test results were interpreted according to the interpretation table, and the results were numerically coded, and the numerical code was found in the identification manual of API20E to determine the type of the strains.

Serological identification was completed using a slide agglutination method. In brief, a clean slide was taken, and a drop of *Shigella* diagnostic serum was added, and a drop of physiological saline was added as a negative control. Fresh colonies were picked and mixed thoroughly with them. They were allowed to be placed at room temperature for a while, and the flocculation or sand-like agglutination is visible to the naked eye, which is a positive reaction, and vice versa. The results were interpreted according to the test reaction. Strain sensitive to ampicillin was isolated from 39 strains by Kirby–Bauer (K-B) disk diffusion method, as recommended by the Clinical and Laboratory Standards Institute (CLSI, 2017).<sup>11</sup> The laboratory number was mel-ss2014011. *E. coli* ATCC 25922 was used as a control for quality assurance.

### MIC analysis of ampicillin

In order to obtain the induced concentration of *Shigella* on ampicillin, we measured the MIC of *Shigella* to ampicillin by using broth microdilution method, as recommended by the CLSI (2017).<sup>11</sup> A bacteria was suspended in Mueller–Hinton broth (0.5 McFarland), then diluted 1000 times, aliquoted into 96-well plates and incubated at  $37^{\circ}$ C for 16 hrs; the MIC was defined as the lowest concentration of antibiotic at which no visible bacterial growth was observed. Quality control testing was performed using *E. coli* ATCC 25922.

# Construction of ampicillin-induced tolerance model

To acquire a strain tolerant to ampicillin, we exposed *Shigella* strain to the lethal dose of ampicillin, which is 10 times greater than the MIC. In each cycle, a single colony was picked and cultured in Luria-Bertani (LB) broth medium, and then bacterial solution was shaken for 18 hrs to reach the stationary-growth phase at 37°C, 200 rpm, followed by 5 hrs of treatment with 10 MIC of ampicillin at 200 rpm at 37°C. The culture was washed to remove the antibiotic, resuspended in a fresh medium and grown overnight at 200 rpm at 37°C. The nonantibiotic control strains were also grown overnight for 18 hrs in Luria-Bertani (LB)

broth medium, but they were not treated with any antibiotic for 5 hrs, then reinoculated in new medium. After 20 cycles of induction for both wild-type strain and nonantibiotic control strains, we stopped inducing bacteria. The induced strain had the same MIC compared with the wild-type strain, but compared with the wild-type strain, the MDK<sub>99</sub> increased after induction, indicating that the bacteria had been tolerant.<sup>12</sup>

### Determination of MDK<sub>99</sub>

To measure the survival under antibiotic treatments, the wide-type strain and tolerant strain and nonantibiotic control strain were inoculated in the Luria-Bertani (LB) broth medium, grown overnight and diluted 1:100 in fresh medium supplement with 10 MIC antibiotic. At the indicated time points, aliquots of the cultures were sampled, diluted to the appropriate dilutions and plated on Luria-Bertani (LB) agar plates. Colony forming units (CFU) were counted after incubation 24 hrs at 37°C to make sure that all CFUs had appeared. All measurements were repeated in at least two independent experiments.

## Homology identification before and after induction

In order to ensure the original source of the tolerant strain, the tolerant strain was identified using 16SrRNA, biochemistry identification. Genetic DNAs from these bacteria were isolated by Bacterial Genomic DNA Kit (Shanghai Lifefeng Biotech, Shanghai, China) and sequenced by 3730XL system at the Beijing Genomics Institute (Shenzhen, China), and then, compared with the BLAST database (http://www.ncbi.nlm.nih.gov/BLAST/).<sup>13</sup> Biochemical identification was performed using API20E (Same as above).

### Sequencing and genomic analysis

To explore the genetic mechanisms of antibiotic tolerance, we extracted bacterial DNA and sequenced it for screening non-SNPs and Indels. DNA was extracted using Bacterial Genomic DNA Kit (Shanghai Lifefeng Biotech). The genome of the wide-type strain was sequenced using a PacBio RSII platform and Illumina HiSeq 4000 system (Illumina, San Diego, CA, USA) and the tolerant strain was sequenced using an Illumina HiSeq 4000 system (Illumina) at the Beijing Genomics Institute (Shenzhen, China). With alignment software MUMmer, each query sequence is aligned with the reference sequence. The variation sites between the query sequence and reference sequence are found out and filtered preliminarily to detect potential non-SNPs and Indel sites.

## RNA extraction and RT-PCR

To detect the gene expression levels, RNA was extracted and RT-PCR was performed. Bacteria were inoculated in a fresh Luria-Bertani (LB) broth medium and grown to mid-logarithmic phase. Total RNA was extracted using RNA pure Bacteria kit (Beijing Com Win Biotech, Beijing, China). PrimeScript<sup>TM</sup> RT reagent Kit (Takara Bio, Beijing, china) was used to eliminate contaminating genomic DNA according to the manufacturer's instruction. The purified RNA was retrieved into the cDNA for the above kit. The PCR reactions (20µl total volume) contained 10 µL TB Green<sup>TM</sup> Premix EX Tap<sup>TM</sup> II, 6 µL RNase-free water, upstream and downstream primers were 0.8 µL, cDNA template 2 µL, ROX reference Dye II (50x) 0.4  $\mu$ L. The PCR condition was as follows: an initial denaturation at 95°C for 30 s, 40 cycles of 5 s at 95°C and 34 s at 60°C.  $2^{-\triangle \triangle CT}$  method was used to evaluate the relative expression of genes.16SrRNA gene expression was used as an endogenous control. All data on gene expression were the mean of triplicate analysis. Gene expression was analyzed using SPSS 21.0. Two independent sample t-test was used for comparison of gene expression.

### Results

### Isolation and identification of strain

To study antibiotic tolerance of *Shigella* strain sensitive to ampicillin, we separated a strain sensitive to ampicillin that was originally isolated from Sui county in Henan province in 2014. This strain was confirmed to be *Shigella* by biochemical and serological identification. This *Shigella* strain sensitive to ampicillin was determined by K-B paper diffusion method (data not shown). As shown in Table 1, GLU, MAN and ARA could be decomposed completely (Table 1). After verification by apiweb TM<sup>M</sup> software, it was identified as *Shigella* sonnei, and we named this strain as melss2014011. These results suggested that *Shigella* strain sensitive to ampicillin was successfully isolated.

## Induction of tolerance to ampicillin by Shigella

We tested whether the long-term treatment with ampicillin would develop a *Shigella* strain that was tolerant to ampicillin. To this end, wild-type strain was exposed to a Luria-Bertani (LB) broth medium for 18 hrs to reach the stationary-growth

Biochemical	Reaction results			
reaction name	mel-ss2014011	mel-ss2014011/20		
ONPG	-	-		
ADH	-	-		
LDC	-	-		
ODC	-	-		
CIT	-	-		
H2S	-	-		
URE	-	-		
TDA	-	-		
IND	-	-		
VP	-	-		
GEL	-	-		
GLU	+	+		
MAN	+	+		
INO	-	-		
SOR	-	-		
RHA	-	-		
SAC	-	-		
MEL	-	-		
AMY	-	-		
	L _	L		

 Table I Biochemical reaction results between wild-type strain and tolerant strain

Abbreviations: ONPG, Beta-galactosidase; ADH, Arginine dihydrolase; LDC, Lysine decarboxylase; ODC, Ornithine decarboxylase; CIT, Citrate utilization; H2S, H2S production; URE, Urease; TDA, Tryptophan deaminase; IND, Indole production; VP, Acetoin production; GEL, Gelatinase; GLU, Glucose fermentation; MAN, Mannitol fermentation; INO, Inositol fermentation; SOR, Sorbitol fermentation; RHA, Rhamnose fermentation; SAC, Sacharose fermentation; MEL, Melbiose fermentation; AMY, Amygdalin fermentation; ARA, Arabinose fermentation.

phase, followed by 5 hrs of treatment with 10 MIC of ampicillin. But nonantibiotic control strains were not treated with any antibiotic for 5 hrs. After 20 cycles of such treatment, bacteria could not be killed effectively and were tolerant to ampicillin, and we named tolerant strain as mel-ss2014011/20. The tolerant strain, wild-type and nonantibiotic control strain had the same MIC, which was 2  $\mu$ g/mL (Figure 1). The bacterial concentration of the wild-type strain decreased from



Figure I The comparison of MIC between wild-type strain and tolerant strain and nonantibiotic strain. Notes: (A) The MIC of wild-type strain. (B) The MIC of tolerant strain. (C) The MIC of nonantibiotic strain. Abbreviation: MIC, minimum inhibitory concentration.

nonantibiotic control strain decreased from  $1.75 \times 10^9$  CFU/mL to  $1.5 \times 10^7$  CFU/mL during 10 mins (MDK<sub>99</sub> was 10 mins), but tolerant strain only reduced from  $1.8 \times 10^9$  CFU/mL to  $2.6 \times 10^8$  CFU/mL in 60 mins (Figure 2). In order to detect the MDK<sub>99</sub> of the tolerant strain, we extended the time to 10 hrs and sampled every 2 hrs. The results showed that both wild-type strain and nonantibiotic control strain decreased from  $1.2 \times 10^{10}$  CFU/mL to 0 in 4 hrs, while the bacterial concentration of tolerant strain decreased from  $2.8 \times 10^9$  CFU/mL to  $5 \times 10^7$  more than 10 hrs (MDK<sub>99</sub> exceeds 10 hrs) (Figure 2). The nonantibiotic control strains were consistent with the wild-type MIC and MDK<sub>99</sub>, indicating that *Shigella* tolerance would be generated under antibiotic pressure.

 $6.5 \times 10^{10}$  CFU/mL to  $2.9 \times 10^8$  CFU/mL during 10 mins

(MDK<sub>99</sub> was 10 mins), and the bacterial concentration of

## Homology identification before and after induction

To confirm that the tolerant strain was derived from a wild-type strain, 16SrRNA and biochemical was used to identify its homology. The 16SrRNA was identified, and the homology of the tolerant strain and wild-type strain to *Shigella* was 99% (data not shown). API20E was used to identify the tolerant (mel-ss2014011/20) strain as *Shigella*.spp. (Table 1). The results showed that tolerant strain was derived from the wild-type strain and the induction process was not contaminated.

### Sequencing and genomic analysis

To elucidate the genetic basis of the *Shigella* tolerance for ampicillin, the wild-type strain was subjected to whole-genome sequencing, and the tolerant strain was sequenced. Whole-gene sequencing results showed that mel-ss2014011 contained a 4814885bp chromosome. The GC -content of chromosome is 51.99%, 5203 predicted protein-coding



Figure 2 Time-dependent killing of Shigella by ampicillin.

Notes: Wild-type strain and tolerant strain and nonantibiotic strain in stable growth stage were exposed to 10 MIC of ampicillin and CFUs were recorded. (A) Wild-type strain, nonantibiotic control and tolerant strain were exposed to ampicillin within 60 mins, and samples were taken every 10 mins and CFUs were recorded. (B) Wild-type strain, nonantibiotic control and tolerant strain were exposed to ampicillin within 10 hrs, and samples were taken every 2 hrs and CFUs were recorded. (B) Wild-type strain, nonantibiotic control and tolerant strain were exposed to ampicillin within 10 hrs, and samples were taken every 2 hrs and CFUs were recorded. (B) Wild-type strain, monantibiotic control and tolerant strain were exposed to ampicillin within 10 hrs, and samples were taken every 2 hrs and CFUs were recorded.

genes, 95 tRNA genes and 54 sRNA genes (Figure 3). Compared with the wild-type strain (mel-ss2014011), tolerant strain (mel-ss2014011/20) had one nonsynonymous mutation, and 10 Indels were located in the noncoding region. This nonsynonymous mutation was found at 2252304 loci of the GL002314 gene (G to A/Val to Met) in mel-ss2014011/20 (Figure 4). The annotation in the Kyoto Encyclopedia of Genes and Genomes (KEGG)database to the GL002314 gene is *cfa*, encoding cyclopropane-fattyacyl-phospholipid synthase. Its Cluster of Orthologous Groups of Proteins (COG) function is classified as I, which is interpreted as lipid transport and metabolism (Figure 3). In addition, there were 10 single base insertions in the noncoding region of different genes (Table 2). The annotations for both GL000777 gene and GL004546 gene in the KEGG database are K07495, a putative transposase. Its COG function is classified as X, which is interpreted as Mobilome: prophages, transposons. The annotation for GL003859 gene is insB, encoding IS1 protein. Its COG function is classified as X, which is interpreted as Mobilome: prophages, transposons (Table 2). The remaining seven genes are not explained in the KEGG database (Table 2). These results indicate that it is possible that the cfa gene may be associated with Shigella tolerance to ampicillin.

#### cfa expression analysis by real-time PCR

Next, we used RT-PCR to detect the expression of cfa gene. Our results showed that the relative expression levels of cfa genes in mel-ss2014011 and mel-ss2014011-20 were 1±0.14 and 7.56±0.90, respectively

(Figure 5). The two independent sample *t*-test analysis showed a difference significant among the expression of wild-type and tolerant strains (t=12.46, P<0.01) (Figure 5). These results suggested that the *cfa* gene expression was increased in the tolerant strain. It is possible that the mutation of the *cfa* gene leads to higher stability of its expression.

#### Discussion

The failure of antibiotic treatment is largely attributed to drug resistance. Many mechanisms of drug resistance have been elucidated, such as the role of bacterial efflux pumps, the decrease of cell membrane permeability, the change of antibiotic targets and the modification of enzymes.<sup>14</sup> However, another mechanism employed by bacteria to combat antibiotics is tolerance.<sup>15</sup> Thus, it is very important to find tolerance in the clinic, delaying the occurrence of drug resistance to some extent.

To obtain a tolerant strain, we induce *Shigella* tolerant to ampicillin by exposing them with high-level antibiotic. Studies have reported that bacteria undergo two mutations from susceptibility to resistance: one is tolerance mutation, and the second mutation is resistance mutation on the basis of tolerance mutation, and bacterial tolerance is prior to the occurrence of drug resistance.<sup>16</sup> Preventing the evolution of tolerance may offer a new strategy for delaying the emergence of resistance. Tolerance was assessed by measuring MDK, for example, the minimum duration of treatment that is required to kill 99% of bacteria (MDK<sub>99</sub>). A high tolerance translates to a longer MDK<sub>99</sub>, that is, a treatment of longer duration is needed to reach the same level of killing. Mechler et al<sup>17</sup> showed that



Figure 3 mel-ss2014011 chromosome gene map.

**Notes:** A total of 8 circles from the outside to the inside, the first circle represents the size of the genome; each large grid is 0.8 M. The second and third circles are CDS on the positive and negative chains, and the different colors indicate the functional classification of the different COGs of the CDS. The fourth and fifth circles are positive and negative chain ncRNAs. The sixth circle is a repeating sequence and the seventh circle is the GC content. The innermost circle is the GC skew value. The specific algorithm is G-C/G+C. When the value is positive in the biological sense, the positive chain is more inclined to transcribe CDS. When it is negative, the negative chain is more inclined to transcription CDS.

S. aureus was exposed to a lethal dose (100 MIC) of daptomycin. After 6 cycles, the drug could not effectively kill the bacteria, and the tolerant strain D6 was obtained. The MDK<sub>99</sub> was  $\geq$ 96 hrs. In our study, mel-ss2014011 (wild-type strain) was exposed to a lethal dose of ampicillin (10 MIC). After 20 cycles, bacteria could not be killed by ampicillin effectively and were tolerant to ampicillin, and we named this strain as mel-ss2014011/20. The MIC of two *Shigella* strains remained unchanged. The MDK<sub>99</sub> of wild-type strain was 10 mins (from  $6.5 \times 10^{10}$  CFU/mL to  $2.9 \times 10^{8}$  CFU/mL); the MDK<sub>99</sub> of induced strain was  $\geq$ 10 hrs (from  $2.8 \times 10^{9}$  CFU/mL to  $5 \times 10^{7}$  CFU/mL). These results suggest that the effectiveness of antibiotics was reduced and the time to treat tolerant bacteria was prolonged.

The mechanism of bacterial tolerance to antibiotics remains largely unknown. To discover the genetic basis of tolerance, wild-type strain was subjected to whole-genome sequencing and the tolerant strain was sequenced to screen non-SNPs and Indels. Compared with mel-ss2014011, mel-ss2014011-20 had one nonsynonymous mutation at *cfa* gene, encoding cyclopropane-fatty-acyl-phospholipid synthase, and 10 Indels were located in the noncoding region. Cyclopropane

Query 1	MSSSCIEEVSVPDDNWYRIANELLSRAGIAINGSAPADIRVKNPDFFKRVLQEGSLGLGE	60
	${\tt MSSSCIEEVSVPDD}{\tt NWYRIANELLSRAGIAINGSAPADIRVKNPDFFKRVLQEGSLGLGE}$	
Subject 1	${\tt MSSSCIEEVSVPDD}{\tt NWYRIANELLSRAGIAINGSAPADIRVKNPDFFKRVLQEGSLGLGE}$	60
Query 61	${\tt SYMDGWWECDRLDMFFSKVLRAGLENQLPHHFKDTLRIAGARLFNLQSKKRAWIVGKEHY}$	120
	${\tt SYMDGWWECDRLDMFFSKVLRAGLENQLPHHFKDTLRIAGARLFNLQSKKRAWIVGKEHY}$	
Subject61	${\tt SYMDGWWECDRLDMFFSKVLRAGLENQLPHHFKDTLRIAGARLFNLQSKKRAWIVGKEHY}$	120
Query 121	eq:dlgndlfsrmldpfmqyscaywkdadnlesaqqaklkmiceklqlkpgmrvldigcgwgg	180
	DLGNDLFSRMLDPFMQYSCAYWKDADNLESAQQAKLKMICEKLQLKPGMRVLDIGCGWGG	
Subject121	eq:dlgndlfsrmldpfmqyscaywkdadnlesaqqaklkmiceklqlkpgmrvldigcgwgg	180
Query 181	${\tt LAHYMASNYDVSVVGVTISAEQQKMAQERCEGLDVTILLQDYRDLNDQFDRIVSVGMFEH}$	240
	LAHYMASNYDVSVVGVTISAEQQKMAQERCEGLDVTILLQDYRDLNDQFDRIVSVGMFEH	
Subject181	${\tt LAHYMASNYDVSVVGVTISAEQQKMAQERCEGLDVTILLQDYRDLNDQFDRIVSVGMFEH}$	240
Query 241	VGPKNYDTYFAVVPRNLKPEGIFLLHTIGSKKTDLNVDPWINKYIFPNGCLPSVRQIAQS	300
	VGPKNYDTYFAV+DRNLKPEGIFLLHTIGSKKTDLNVDPWINKYIFPNGCLPSVRQIAQS	
Subject241	VGPKNYDTYFAVMDRNLKPEGIFLLHTIGSKKTDLNVDPWINKYIFPNGCLPSVRQIAQS	300
Query 301	${\tt SEPHFVMEDWhnfgadydttlmawyerflaawpeiadnyserfkrmftyylnacagafra}$	360
	${\tt SEPHFVMED} Whn {\tt Fgadydttlmawyerflaawpeiadnyserfkrmftyylnacagafraa}$	
Subject301	${\tt SEPHFVMEDWhnfgadydttlmawyerflaawpeiadnyserfkrmftyylnacagafra}$	360
Query 361	RDIQLWQVVFSRGVENGLRVAR 382	
	RDIQLWQVVFSRGVENGLRVAR	
Subject361	RDIQLWQVVFSRGVENGLRVAR 382	

Figure 4 The comparison of amino acid sequences of cfa genes between wild-type and tolerant strains. Note: Subject represents the wild-type strain and Query represents the tolerant strain.

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lable	2	Indels	ot	non-coding	region	are	screened	by	sequencing
	_							-/	

Gene ID	KEGG name	KEGG defination	Indel location	Base site	COG	Annotation
GL000257	NA	NA	225401	G	NA	NA
GL000777	K07495	Putative transposase	752208	G	x	Mobilome: prophages, transposons
GL000906	NA	NA	868766	G	x	Mobilome: prophages, transposons
GL003130	NA	NA	3040569	с	NA	NA
GL003141	NA	NA	3050041	G	х	Mobilome: prophages, transposons
GL003859	insB	ISI protein	3787181	G	х	Mobilome: prophages, transposons
GL003920	NA	NA	3847162	С	NA	NA
GL004546	K07495	Putative transposase	4501422	А	х	Mobilome: prophages, transposons
GL004596	NA	NA	4557986	С	NA	NA
GL004740	NA	NA	4706946	G	x	Mobilome: prophages, transposons

Abbreviation: NA, not available.



Figure 5 Relative expression levels of cfa gene in wild-type and tolerant strains. Note: Relative expression levels of cfa gene detected by real-time PCR (compared with wild-type strain, \*\*\*P<0.01). The results were repeated three times and then averaged. The melting curve of cfa gene showed only one major peak, indicating a good purity.

fatty acids (CFAs) are an important component of phospholipids in many bacteria<sup>15,18</sup> and are mainly produced at the end of the exponential growth phase and right before the stable growth phase.<sup>19</sup> Bacteria have a complex regulatory network that recognizes and regulates many physiological stresses. In Gram-negative bacteria, this regulation is through the RpoS. The RpoS is a sigma factor that can positively regulate many genes in a stationary phase,<sup>20</sup> one of which is the cfa gene<sup>21</sup> (Figure 6), which encodes a cyclopropane-fatty-acyl-phospholipid synthase. Murakami et al<sup>8</sup> showed that the *rpoS*-knockout strain and the wild-type strain were exposed to biapenem, and the survival rate of the rpoS-knockout strain was lower than that of the wild-type strain in the stationary-growth stage, proving that the *rpoS* is involved in antibiotic tolerance (Figure 6), but the specific mechanism is still unclear. RpoS mainly regulates transcription, translation and post-translational levels through regulatory factors such as ppGpp, H-NS and  $clpP^{22,23}$  (Figure 6). Dalebroux et al showed that ppGpp is generated by relA and SpoT<sup>24</sup> (Figure 6). Published studies have shown that the rpoS and ppGpp are interrelated, and when bacteria are exposed to antibiotics, they can keep the bacteria alive.<sup>25</sup> The increase in RpoS mRNA translation is through ppGpp.<sup>26</sup> In addition, the *hipA* mutation (G22S, D291A) can greatly increase the tolerance level in E. coli. It was found that *hipA* mutation could lead to the synthesis of ppGpp. The increase of ppGpp level can significantly increase





**Notes:** ppGpp is generated by relA and spoT. ppGpp can positively regulate RpoS. Activation of ppGpp can produce antibiotic tolerance. RpoS is involved in the formation of antibiotic tolerance. RpoS (sigma factor) can positively regulate many genes in a stationary phase, one of which is *cfa*, which encodes a cyclopropane-fatty-acyl-phospholipid synthase.

the level of bacterial tolerance<sup>27</sup> (Figure 6). In our study, the results of comparative genomics showed that the tolerant *Shigella* strain had one nonsynonymous mutation at *cfa* (G to A/Val to Met) gene, and 10 Indels were located in the noncoding region of different genes. Since the Indels are located in the noncoding region and none of these genes carry any resistance gene, these Indels are less likely to affect the tolerance of ampicillin to *Shigella*. It is possible that the *cfa* gene under the regulation of *cfa* gene may increase ppGpp, thereby increasing the tolerance level of *Shigella* to ampicillin.

We used RT-PCR to detect the expression of cfa gene. Studies have shown that cfa increases in bacteria when it responds to low pH values, as well as high temperatures, and osmotic imbalance.<sup>28</sup> Shabala et al showed that in cfa-deletion mutations, *E. coli* increased proton permeability and reduced the ability to excrete H<sup>+</sup> and they confirmed that cfa can protect *E. coli* from acid stress by reducing cell membrane

permeability to H<sup>+</sup>.<sup>29</sup> Macdonald et al have confirmed that an increase in cfa leads to an increase in cell membrane stability and rigidity, and an increase in cell membrane rigidity is associated with a decrease in proton permeability.<sup>30</sup> Studies have shown that the decrease of cell membrane permeability of bacteria will lead to the decrease the concentration of antibiotics entering biofilms, the slow growth of bacteria, which will lead to bacterial tolerance to antibiotics.<sup>31–33</sup> In our study, by measuring the tolerance level of the mutant strain, the  $MDK_{99}$ was 60-fold higher than the wide-type strain, indicating that the tolerance level has been reached. Real-time PCR results showed that the relative expression levels of cfa were 7.56fold higher in the tolerant strain than in the wild-type strain. Based on the above studies, we propose a hypothesis that tolerance may be related to cfa gene expression. It is possible that the mutation of the *cfa* gene leads to higher stability of its expression. Increased relative expression of cfa gene in the tolerant strain may increase the rigidity of the cell membrane and decrease the permeability of the cell membrane, resulting in the inability of ampicillin to effectively kill Shigella bacteria and prolong the treatment time to make the bacteria tolerant.

cfa gene may be related to tolerance, and the mutation of the cfa gene leads to higher stability of its expression. Moreover, the cfa gene may be a target for overcoming drug tolerance, delaying the occurrence of drug resistance to some extent, and provide novel strategies for the prevention of drug resistance of *Shigella*.

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### Disclosure

The authors report no conflicts of interest in this work.

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