Evaluating the Contribution of Acid Resistance Systems and Probing the Different Roles of the Glutamate Decarboxylases of *Listeria monocytogenes* Under Acidic Conditions

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Abstract

Listeria monocytogenes is an important zoonotic foodborne pathogen, which can cause a severe invasive illness to susceptible humans and animals with high mortality. As *L. monocytogenes* is widely distributed in natural environments, the bacterium is easy to contaminate food processing facilities and the products to be ingested by host. But during the transition from a saprophyte to intracellular pathogen, one of the biggest challenge *L. monocytogenes* encounters is the acid stress. To combat the acidic environments, the bacterium developed several acid resistance systems, including acid tolerance response (ATR), F0F1-ATPase, glutamate decarboxylase (GAD), arginine deiminase (ADI) and agmatine deiminase (AgDI). In this study, we comprehensively evaluated the contributions of different acid resistance systems and explored the different roles of the three GAD components under acidic conditions. We found that the GadD2 of GAD system made the largest contribution to the survival of *L. monocytogenes* in artificial gastric juice (AGJ) and acidic brain heart infusion (BHI), which was followed by the global stress regulator SigB, GadD3 of GAD system, AguA1 of AgDI system and ArcA of ADI system. Transcription analysis showed that the mRNA level of the three GADs were consistent with their contribution to acid resistance. Similar results were observed in the other three representative strains EGDe, Lm850658 and M7. We further obtained the purified GADs and their poly-antibodies to demonstrate that the contribution of the three GADs were determined by the protein levels in *L. monocytogenes*. Further studies are needed to focus on the regulation of different expression of the GAD system.

Keywords: Listeria monocytogenes, Acid resistance, Glutamate decarboxylase, Survival

Asidik Koşullar Altında *Listeria monocytogenes'*in Glutamat Dekarboksilazlarının Asit Direnç Sistemlerine Katkılarının Değerlendirilmesi ve Farklı Rollerinin Araştırılması

Öz

Listeria monocytogenes, duyarlı insan ve hayvanlarda yüksek ölüm oranı ile seyreden bulaşıcı hastalıklara neden olabilen, önemli bir gıda kaynaklı zoonotik patojendir. L. monocytogenes doğal ortamlarda yaygın olarak bulunduğundan, gıda işleme tesislerinin ve konakçı tarafından tüketilen ürünlerin bakteri ile kontaminasyonu kolaydır. Ancak bir saprofitten hücre içi patojene dönüşmesi sırasında, L. monocytogenes'in karşılaştığı en büyük güçlüklerden biri asit stresidir. Asidik ortamlarla savaşmak için, bakteri, asit tolerans yanıtı (ATR), FOF1-ATPase, glutamat dekarboksilaz (GAD), arginin deiminaz (ADI) ve agmatin deiminaz (AgDI) dahil olmak üzere çeşitli asit direnç sistemleri geliştirmiştir. Bu çalışmada, farklı asit direnç sistemlerinin katkıları kapsamlı bir şekilde değerlendirildi ve üç GAD bileşeninin asidik koşullar altında farklı rolleri araştırıldı. GAD sistemindeki GadD2'nin, L. monocytogenes'in yapay mide sıvısı (AGJ) ve asidik beyin kalp infüzyonunda (BHI) hayatta kalmasına en büyük katkışı yaptığı ve bunu GAD sisteminden global stres regülatörü SigB, GadD3 ile AgDI sisteminden AguA1 ve ADI sisteminden ArcA'nın izlediği belirlendi. Transkripsiyon analizi, üç GAD'nin mRNA seviyesinin, asit direncine katkıları ile tutarlı olduğunu gösterdi. Benzer sonuçlar, diğer üç temsilci suş olan EGDe, Lm850658 ve M7'de de gözlendi. Ayrıca, üç GAD'nin katkısının, L. monocytogenes'teki protein seviyeleri tarafından belirlendiğini göstermek için saflaştırılmış GAD'ler ve bunların poliantikorlarını elde ettik. GAD sisteminin farklı ekspresyonlarının düzenlenme mekanizmasının anlaşılabilmesi için daha fazla çalışmaya ihtiyaç vardır.

Anahtar sözcükler: Listeria monocytogenes, Asit direnci, Glutamat dekarboksilaz, Sağkalım

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INTRODUCTION

Listeria monocytogenes is a facultative anaerobic, grampositive bacterium that is ubiquitous in natural environment as a saprophyte. In field environment, L. monocytogenes is thought to live off decaying plant material. Following ingestion by susceptible humans or animals, L. monocytogenes is capable of making the transition into a pathogen ^[1]. As an important zoonotic foodborne pathogen, L. monocytogenes could cause a severe invasive illness with high mortality in immunocompromised individuals ^[2,3]. Foodborne infection is the most common pathway of both epidemic and sporadic listeriosis, with 99% of human cases caused by consumption of contaminated food products^[4]. Upon entering into the host gastrointestinal tract, L. monocytogenes adhere and invade various types of cells, including the phagocytic cells by the internalins InIA and InIB as well as Lap and InIP^[5]. Following theentry into cell, L. monocytogenes must escape from host cell vacuoles via the pore-forming cytolysin listeriolysin O (LLO) and two phospholipases PI-PLC and PC-PLC quickly ^[6]. If not, the bacteria can be killed by the acidic endosome and digested by enzymes from the fused lysosome (phagolysosome). Entry into the cytosol, L. monocytogenes uses cytosolic nutrient to proliferate, then the bacteria spread to the neighboring cells by usurping actin polymerization as motile force by the bacterial surface protein ActA and with the help of InIC to relieve the cortical tension ^[7]. Then L. monocytogenes need to escape from the double membrane vacuoles to finish cell-to-cell spread with the help of LLO, PC-PLC and PI-PLC once again ^[5]. So it is not difficult to find that during the infection process, one of the biggest challenge L. monocytogenes encounters is acid stresses from stomach and phagolysosomes ^[8].

Listeria monocytogenes contains several enzyme systems including F₀F₁-ATPase, ADI, AgDI, GAD and acid tolerance response, to maintain intracellular pH homeostasis in acidic environments ^[9]. Under acid stress, F₀F₁-ATPase system uses ATP hydrolysis to produce proton motive force to pump cytoplasmic protons, while the ADI and AgDI use arginine and agmatine to produce ammonia to neutralize the cytoplasmic protons, respectively ^[10,11]. The glutamate decarboxylase (GAD) system, which consumes intracellular protons by converting glutamate to γ -aminobutyrate ^[12], also plays a role in acid resistance of L. monocytogenes to protect them in low pH foods. Moreover, pre-exposure of L. monocytogenes to mild acid could induce acid tolerance response (ATR) that improves the survival rate under fatal acid stress. As a global transcriptional regulator, SigB has been reported to positively regulate the ATR to help L. monocytogenes to deal with acid stress [13].

Although all the acid resistance systems were individually demonstrated to play important roles in acid stress and pathogenicity of *L. monocytogenes*, to date, no comprehensive assessment was conducted on these acid resistance systems,

and the relative roles of these systems remain unclear. Moreover, *L. monocytogenes* contains several copies for some acid resistance systems. For example, *L. monocytogenes* 10403S has two AgDI genes (*aguA1* and *aguA2*), and both of them were upregulated in response to acid stress, but only AguA1 contributed to acid resistance and pathogenicity of the bacteria ^[11]. For the GAD system, most of the *L. monocytogenes* strains (lineages I and II) contain three GADs ^[14], but the contributions of different GADs remain unclear. In this study, we tried to evaluate the contributions of different acid resistance systems and to clarify the different roles of the three GAD components under the acidic condition.

MATERIAL and METHODS

Bacterial Strains, Plasmids and Culture Conditions

Listeria monocytogenes 10403S, EGDe, Lm850658 and M7 were used as the wild-type strains. Escherichia coli DH5a was employed as the host strain for plasmids pET30a and pKSV7. E. coli Rosetta was used as expression host. L. monocytogenes and E. coli were cultured in brain heart infusion (BHI, Oxoid, Basingstoke, U.K.) and Luria-Bertani medium (LB, Oxoid), respectively, at 37°C. Stock solutions of ampicillin (50 mg/mL), kanamycin (50 mg/mL) and chloramphenicol (10 mg/mL; Sangong Biotech Co., Ltd, Shanghai, China) were added to the media, when appropriate, at the required concentrations.

Survival in AGJ or Acidic BHI Broth

Survival assay in artificial gastric juice (AGJ) or acidic BHI broth was conducted as in previous research ^[15]. L. monocytogenes wild-type and mutant strains were grown overnight at 37°C in BHI broth at pH 7.0 with shaking. The cultures were collected by centrifugation at 3000 g at 4°C for 10 min, washed and resuspended in phosphate buffered saline (PBS, 10 mM, pH 7.4) with the OD_{600 nm} adjusted to 1.0. Then 50 µL bacterial suspension was mixed in 950 µL AGJ (8.3 g proteose peptone, 3.5 g D-glucose, 2.05 g NaCl, 0.6 g KH₂PO₄, 0.11 g CaCl₂, 0.37 g KCl, 0.05 g bile salt, 0.1 g lysozyme and 13.3 mg pepsin dissolved in 1 liter distilled water with pH adjusted to 2.5 with HCl, filter sterilized) or BHI broth with indicated pH values (filter sterilized). After 1 h of incubation at 37°C, the mixtures were serially diluted and plated on BHI agar plates. The plates were incubated at 37°C for 24 h and viable bacteria were counted. Survival rate was calculated as percentage of survived bacteria after incubating in the acidic conditions for 1 h relative to the incubated bacteria. Data was reported as the mean ± SD of three independent experiments, each performed in triplicate.

Transcriptional Analysis

Overnight cultures of *L. monocytogenes* strains were inoculated into fresh BHI broth and grown to exponential phase ($OD_{600 nm}$ =0.25) or stationary phase ($OD_{600 nm}$ =0.6) at

37°C. One milliliter of each culture was treated with pH 4.5 BHI for an hour and then pelleted by centrifugation at 4°C. Total RNA was extracted using the Trizol reagent according to the manufacturer's instruction (Sangong Biotech Co., Ltd) and cDNA was synthesized with reverse transcriptase (TOYOBO Biotech Co., Ltd, Shanghai, China). Quantitative real-time PCR (qRT-PCR) was performed in 20 μ L reaction mixtures containing SYBR green qPCR mix (TOYOBO (SHANGHAI) Biotech Co., Ltd) to detect the transcriptional levels of indicated genes on the iCycler iQ5 real-time PCR system (Bio-Rad, Hercules, California, U.S.A.) with specific primer pairs listed in *Table 1*. The housekeeping gene *gyrB* was selected as an internal control for normalization as previous research ^[16].

Construction of Deletion and Complementation Mutants

A homologous recombination strategy was used to construct the deletion mutants of L. monocytogenes 10403S according to the previous research ^[17] using the primer pairs listed in *Table 2*. The homologous fragments of overlapping PCR were purified and ligated to pMD18-T (TaKaRa, Beijing). After sequencing, the inserted fragments were digested with the indicated restriction enzymes, ligated to the temperature-sensitive shuttle vector pKSV7 and transformed into DH5a. Plasmids containing the inserted fragments were subsequently extracted and electroporated into L. monocytogenes competent cells. Transformants were grown at a non-permissive temperature (41°C) on BHI agar containing chloramphenicol (10 µg/mL) to promote chromosomal integration. The recombinants were passed in succession in BHI without antibiotic at a permissive temperature (30°C) to enable plasmid excision and curing. The deletion mutants were identified by PCR and confirmed by sequencing.

For the complementation strains, the encoding sequences of *gadD1*, *gadD2* and *gadD3* were amplified from *L*. *monocytogenes* EGDe with the indicated primer pairs listed in *Table 2*. After restriction digestion with appropriate

Table 1. Primers used for q-PCR				
Primers	Sequences (5'-3')	Size (bp)		
gadD1-fwd	AGAATATCCACAGACAGCAAAG	140		
gadD1-rev	CATAGCCATTCCACCAAGCAT	142		
gadT1-fwd	CGTTCTCGGTATTACAATTCCT	150		
gadT1-rev	GCAAGCATGAAGATAACAAGAG			
gadT2-fwd	CCCTGTACCACTTATTATGGTT	116		
gadT2-rev	CTACAGTTAAGGAAATTGCGGT			
gadD2-fwd	CCTTTGGAAAGATGAAAGCTAC	- 128		
gadD2-rev	TGTAGTATTGACCGATGATGTG			
gadD3-fwd	ACCAATAATTTGGCTCGCACTA			
gadD3-rev	TTAGTTTATCCGGGTGTTGGTT	- 144		
gyrB-fwd	AGACGCTATTGATGCCGATGA	- 91		
gyrB-rev	GTATTGCGCGTTGTCTTCGA			

enzymes, the PCR fragment was cloned into pIMK2 following the P_{Help} promoter. The recombinant plasmids were then electroporated into *L. monocytogenes* EGDe competent cells. The transformants were plated on BHI agar containing kanamycin (50 µg/mL) and positive clones were picked up and identified by PCR.

Prokaryotic Expression and Purification of GadD1, GadD2 and GadD3

GadD1, GadD2 and GadD3 were expressed as fusion proteins with His-tag using the expression vector pET30a (Invitrogen, U.S.A.) as previously shown ^[18]. The full-length *gadD1*, *gadD2* and *gadD3* were amplified with primer pairs listed in *Table 3*. The amplified fragments were cloned into

Table 2. Prin	ners used for deletion and complement mutants co	nstruction	
Primers	Sequences (5'-3')	Size(bp)	
gadD1-a	AAT <u>AAGCTT</u> ACTACACAGGTTTACAAGCA	515	
gadD1-b	ACTCTCCCATTTTTCATAAATTCCTCCA	515	
gadD1-c	GAAAAATGGGAGAGTGATAAAATTTCTAG	524	
gadD1-d	GCT <u>GAATTC</u> TTTTAATTGAAGTAACGTCA	524	
gadD1-e	AACCAACAGAAACATCGCTTCGTAT		
gadD2-a	ATA <u>GCATGC</u> CACTTATTATGGTTCAAG		
gadD2-b	GATTTTTCCTCCTATAATTTGTCTTGATT	536	
gadD2-c	TAGGAGGAAAAAATCTTCACACATTAA		
gadD2-d	ATA <u>GAATTC</u> GGACTTATTCCGAGTAATG	545	
gadD2-e	GCAGCACTTTGTTACTTTTTGAAGAAG		
gadD3-a	GCA <u>GGATCC</u> AGCTTCTACTCTAACATGGTTCACG		
gadD3-b	TTATAGTGAAGACGACAAGCGAACTTGGATGGT GAGTCCGA	567	
gadD3-c	TTCGCTTGTCGTCTTCACTATAAAGC	605	
gadD3-d	AAC <u>GGTACC</u> CGAGCGTGTCTATCTCACTATTCAT	605	
gadD3-e	GAAATTGTCGATTCCGGTGATGACT		
gadD1-CF	CG <u>GGATCC</u> TATGTTTAAAACAAATGTTGAACAAA	1406	
gadD1-CR	GG <u>GGTACC</u> TTAATGAGTAAAGCCATGTGT	1406	
gadD2-CF	CG <u>GGATCC</u> CATGTTATATAGTAAAGAAAATAA	1412	
gadD2-CR	GG <u>GGTACC</u> TTAATGTGTGAAGCCGTGGA		
gadD3-CF	CG <u>GGATCC</u> GATGCTTTATAGTGAAGACGACA	1421	
gadD3-CR	GG <u>GGTACC</u> TTAGTGCGTAAATCCGTATGAA		
Sequences with underline were restriction enzyme sites			

Table 3. Primers used for expression of the glutamate decarboxylases			
Primers	Sequences (5'-3')	Size (bp)	
gadD1-exp-fwd	GGA <u>GGTACC</u> ATGTTTAAAACAAATGTTGAACAAA	1407	
gadD1-exp -rev	CCA <u>GGATCC</u> TTAATGAGTAAAGCCATGTGT	1407	
gadD2-exp-fwd	GAA <u>GGTACC</u> ATGTTATATAGTAAAGAAAATAAAGA	1413	
gadD2-exp -rev	GCC <u>GGATCC</u> TTAATGTGTGAAGCCGTG	1415	
gadD3-exp-fwd	GGA <u>GGTACC</u> ATGCTTTATAGTGAAGACGACA	1422	
gadD3-exp -rev	TCT <u>GGATCC</u> TTAGTGCGTAAATCCGTATGAA	1422	
Sequences with underline were restriction enzyme sites			

the pET30a after restriction digestion. The recombinant plasmids were transformed to expression host E. coli Rosetta competent cells. Positive clones were confirmed by sequencing and then grown in 200 mL of LB medium supplemented with 50 µg/mL kanamycin at 37°C until OD_{600 nm} of the cultures reached 0.6-0.8. Isopropyl β-D-1thiogalactopyranoside (IPTG) was then added into the medium at a final concentration of 0.4 mM to induce expression of GadD1, GadD2 and GadD3 for 12 h at 15°C. Then IPTG-induced cell pellets were collected, resuspended in 50 mM PBS (pH 7.4), and disrupted with 100 cycles of sonication at 300W for 5 sec with intermittent cooling on ice for 10 sec (25 min in total). After centrifugation at 12.000 g for 20 min, the supernatant samples were collected and loaded onto a 2-mL prepacked nickel-chelated agarose gel column (Weishi-Bohui Chromtotech Co., Ltd, Beijing, China). The columns were washed with 50 mM PBS containing 500 mM NaCl and 30 mM imidazole, and the bound proteins were eluted with a linear gradient of 25-500 mM imidazole prepared in the same buffer. Expression and purification of the recombinant proteins were analyzed on a 10% SDS-PAGE gel followed by Coomassie Brilliant Blue staining.

Polyclonal Antibodies Preparation

The purified recombinant protein was used for raising polyclonal antibodies in New Zealand white rabbits according to the previous study^[19]. Rabbits were first immunized with 500 µg protein emulsified by the equal volume of Freund's complete adjuvant (Sigma, St. Louis, U.S.A.) through subcutaneous injection. After two weeks, the rabbit was boosted subcutaneously three times with 250 µg protein emulsified by incomplete Freund's adjuvant (Sigma, St. Louis, U.S.A.) at two-week intervals. Rabbits were bled 10 days after the last immunization and sera were isolated from the whole blood to collect polyclonal antibodies. Animal experiments were approved by the Laboratory Animal Management Committee of Yangtze University (Approval No. 20161212).

Western Blot Analysis

Listeria monocytogenes wild type and mutant strains were cultured with BHI broth to stationary phase at 37°C, then

treated with pH 4.5 BHI broth for an hour. Then bacteria were harvested by centrifugation and pellets were lysed with lysis buffer and then homogenized with a refiner. The supernatant of cell lysis was isolated by centrifugation and analyzed by 10% SDS-PAGE. GadD1, GadD2 and GadD3 were blotted and probed with respective polyclonal antibodies produced in this study. Glyceraldehyde-phosphate dehydrogenase (GAPDH) was used as loading control. Then HRP conjugated goat-anti-rabbit IgG (Sangong Biotech Co., Ltd) was used as the second antibody to probe GadD1, GadD2, GadD3 and GAPDH. The abundance of indicated proteins was evaluated with software Quantity One (Version 4.6.6, Bio-Rad, U.S.A.) to calculate the gray level of specific bands.

Statistical Analysis

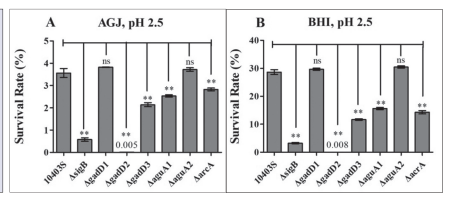
All data comparisons were analyzed using the two-tailed homoscedastic Student's T-test. In all cases, differences with P<0.05 were considered as statistically significant. The GraphPad Prism 5 (Version 5, GraphPad, U.S.A.) software was used to produce the graphs.

RESULTS

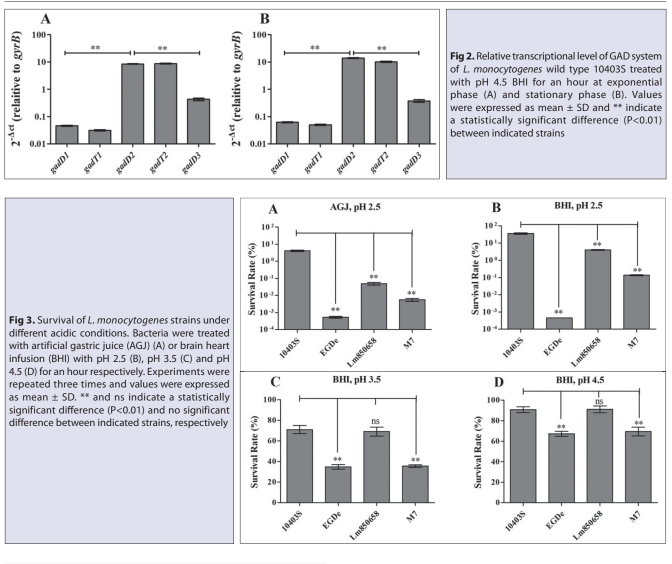
To evaluate the contribution of different acid resistance systems of L. monocytogenes, we knocked out the acid resistance-associated enzymes genes in the background of the reference strain 10403S and then compared the survival rate of the mutants with the reference strain. Our data showed that mutant strain AgadD2 exhibited the lowest survival rate (0.005%) in AGJ, which was followed by strains $\Delta sigB$, $\Delta gadD3$, $\Delta aguA1$, $\Delta arcA$, $\Delta gadD1$ and Δ aguA2 (*Fig. 1A*). Among these mutants, only Δ aguA2 and $\Delta gadD1$ did not show a significant difference on the survival rate of L. monocytogenes in the acidic condition. A similar result was also observed in pH 2.5 acidic BHI broth (Fig. 1B). These data indicated that the contribution of the acid resistance-associated enzymes was quite different. In L. monocytogenes 10403S, GadD2 made the largest contribution to acid resistance, which was followed by SigB, GadD3, AguA1 and ArcA.

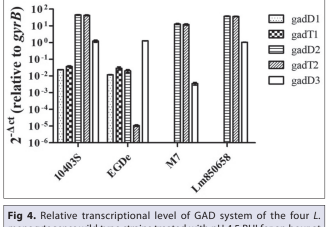
To elucidate the different roles of the components of GAD system in acid resistance, we analyzed the transcriptional

Fig 1. Survival of *L. monocytogenes* wild type 10403S and different mutant strains for an hour in pH 2.5 artificial gastric juice (AGJ) (A) or pH 2.5 brain heart infusion (BHI) broth (B). Experiments were conducted at least three times and values were expressed as mean \pm SD. ** and ns indicate a statistically significant difference (P<0.01) and no significant difference between indicated strains, respectively



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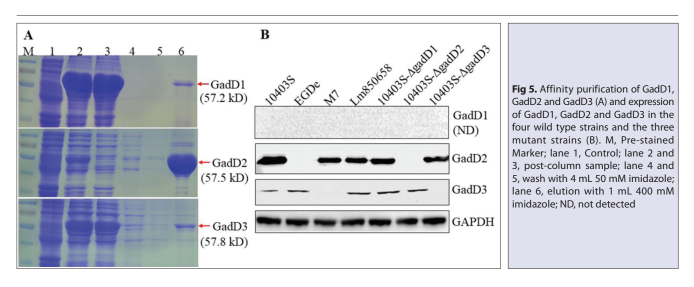


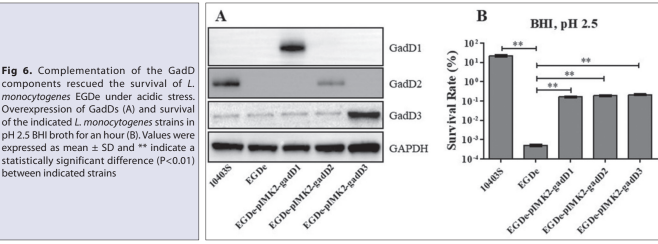
monocytogenes wild type strains treated with pH 4.5 BHI for an hour at the stationary phase

level of the GAD system under acidic conditions. Our data showed that the transcriptional level of *gadT2/gadD2* was significantly higher than that of *gadD3* and *gadD1/gadT1* both at exponential and stationary phases in pH 4.5 BHI broth (*Fig. 2A,B*). Moreover, the mRNA level of

gadD3 was ten-fold more than that of *gadD1/gadT1*. These results suggest that the contribution to acid resistance of different GAD components was correlative to their transcriptional level.

To confirm this hypothesis, we analyzed the survival rate of four representative strains, including 10403S, EGDe, Lm850658 and M7, under acidic conditions and the transcriptional level of their GAD system. Our data showed that strain 10403S exhibited the highest survival rate in AGJ for 1 h, which was followed by Lm850658, M7 and EGDe (Fig. 3A). A similar result was also observed in pH 2.5 BHI broth (Fig. 3B). When the pH increased to 3.5 or pH 4.5 in BHI, the survival rate of the four strains were increased, and the survival rate of 10403S and Lm850658 remained significantly higher than that of M7 and EGDe (Fig. 3C,D). This survival rate was consistent with the transcriptional level, as the transcriptional level of *aadT2/* gadD2 in EGDe and M7 were significantly lower than that of 10403S and Lm850658 in pH 4.5 BHI (Fig. 4). The gadD1/ gadT1 mRNA level of EGDe was equal to that of 10403S, and gadD1/gadT1 genes were deficient in lineage III strains





Lm850658 and M7. The mRNA level of *adD3* of 10403S, EGDe and Lm850658 were at the same level in the acidic condition, which was significantly higher than that of

between indicated strains

M7 (Fig. 4).

To confirm whether the role of GAD system was determined by their expression, we purified prokaryotic expressed GadD1, GadD2 and GadD3 (Fig. 5A). Polyantibodies were obtained from immunized rabbit with the indicated purified proteins. Then the protein level of the three components in the acidic condition was detected by Western blot. The results showed that GadD1 was detected neither in wild type strain nor in the mutants (Fig. 5B). GadD2 was not detected in *gadD2* mutant and EGDe, but highly expressed in 10403S and Lm850658 (Fig. 5B). GadD3 was not detected in *gadD3* mutant and M7 (*Fig. 5B*), which was similar to the tendency of its mRNA level (Fig. 4). To confirm the hypothesis that the role of GAD in acid resistance was determined by their expression level, we overexpressed GadD1, GadD2 and GadD3 in EGDe to determine their function in acid resistance individually. Survival assay in pH 2.5 BHI broth showed that the overexpression any of the three GADs significantly improves the acid resistance of EGDe (Fig. 6).

DISCUSSION

Listeria monocytogenes is a bacterium that lives in natural environments as a saprophyte but is capable of making the transition into a pathogen following its ingestion by susceptible humans or animals^[1]. As an important zoonotic foodborne pathogen, this bacterium has the ability to adapt to a variety of environmental conditions ^[20]. Acidic environments such as silage, fermented foods, stomach and phagolysosomes, are the most common conditions that the bacterium encounters. L. monocytogenes contains several enzyme systems including F₀F₁-ATPase, ADI, AgDI and GAD to cope with these unfavorable conditions. Although all of the acid resistance systems had been demonstrated to play important roles in acid resistance in L. monocytogenes^[9,21-24], it's unclear which system plays the major role in acid resistance of *L. monocytogenes*. Here we evaluated the effects of these systems on the survival of L. monocytogenes under acidic conditions. Our data showed that GadD2 of GAD system made the largest contribution to L. monocytogenes 10403S survival in different acidic conditions, which was followed by SigB, GadD3, AguA1, and ArcA. Since the constitution of these systems was complicated, it is difficult to knock out the whole system to

determine its contribution. In this study, we only knocked out the key enzymes of the acid resistance systems, which might be inadequate to evaluate the whole function of an acid resistance system, but we knocked out the whole AgDI system encoding region (Imo0036-Imo0042) at once. No significant difference was observed about the survival rate of strains $\Delta aguA1(\Delta Imo0038)$ and $\Delta AgDI (\Delta Imo0036-$ Imo0042) in acidic broth (data not shown). F₀F₁-ATPase system is essential for L. monocytogenes, in which mutantion will cause a lethal effect to the bacteria^[25]. we didn't evaluate its contribution to acid resistance of L. monocytogenes. Datta et al.^[25] and Cotter et al.^[26] treated L. monocytogenes LS2 and LO28 at the exponential phase with N, N'-Dicyclohexylcarbodiimide (DCCD), an F₀F₁-ATPase inhibitor, which resulted in significantly reducing survival rate of the bacteria under the lethally acidic condition ^[25,26].

To better deal with acid stress, L. monocytogenes might employ several systems simultaneously. Moreover, the AgDI and GAD systems encode multiple isoenzymes, as two AgDI (aguA1 and aguA2) and three GADs (gadD1, gadD2 and gadD3) genes are encoded in most of L. monocytogenes [14,27]. The roles of these isoenzymes might make different contributions to the acid resistance of *L. monocytogenes*. As our previous study showed that aguA1 and aguA2 were both significantly up-regulated in response to acid conditions, only AguA1 but not AguA2 contributed to survival and growth under acidic environments and was involved in the pathogenicity of L. monocytogenes 10403S, in which Glycine 157 determined the activity of AguA1 and AguA2^[11]. In this study we found that the three GADs in L. monocytogenes 10403S also made different contribution (GadD2 > GadD3 > GadD1) to the survival of bacteria in acid stress. We further demonstrated that the function of the three GADs were determined by the expression level instead of their enzyme activity, which were different from AugA1 and AguA2^[11]. As our results showed that L. monocytogenes EGDe with little GadD1 and GadD2 was quite sensitive to acid stress, while overexpressed GadD1, GadD2 or GadD3 in EGDe with the *pHelp* promoter of plasmid pIMK2 significantly improved its survival rate in pH 2.5 BHI broth (Fig. 3, 4, 6). Previous studies showed the difference of GAD system in L. monocytogenes and divided them into two groups, the outside GAD system (GAD_o, including GadD1/GadT1 and GadT2/GadD2) and the inside GAD system (GAD_i, GadD3)^[23], and GAD_o played the major role in LO28 and 10403S, while acid resistance of EGDe was dependent on GAD_i^[22,28]. These studies found the different roles of GAD system in various strains, but did not clarified the determinants that involved in the contribution in acid resistance of the GAD components. In this study, we demonstrated that the contribution of the three GADs was determined by their expression level for the first time.

The pH values and substrates that could be used by the bacterium to combat with low pH were various in different conditions ^[29], which might promote the bacterium to

choose suitable acid resistance systems to cope with specific acidic environment. Whether these acid resistance systems perform the same in different conditions need further investigate. For instance, acid resistance in the phagosome might be complicated. On one hand, the bacteria need to initiate acid resistance system to cope with the acidic phagosome, on the other hand, the activation of LLO, which mediate L. monocytogenes escape from phagosome, need acidic compartment ^[30]. But food products contain glutamates that tend to benefit for the GAD system to deploy acid resistance [8]. Moreover, the molecular mechanisms involved in the different expression remained unclear. Kazmierczak et al.^[31] found that *gadD3* was positivly regulated by SigB, and Cotter et al.[32] also found that gadD1/gadT1 partially regulated by SigB. Bowman et al.^[33] found that *qadT2/qadD2* operon was constitutively expression in L. monocytogenes LO28 by proteomic analysis. We found that the expression of gadT2/gadD2 in the four representative strains were not in response to the acidic treatment, but the sequence of this operon (including the promoter region) was quite conserved between 10403S and EGDe or between Lm850658 and M7 (data not shown). It is suggested that different expression of gadT2/gadD2 in these strains might be regulated in an undiscovered manner. Taken together, we demonstrated that the different contribution to acid resistance of GAD components were determined by the expression levels. The mechanisms that mediate the expression difference of GAD system need further investigation.

STATEMENT OF AUTHOR CONTRIBUTIONS

CF and XWF designed and conducted experiments, analyzed data, and they were contributed equally to the work. CF, XWF, XYC, XYL, CW, YFG, WHF and YYY were involved in study design and data collection. YYY supervised the study and critically read the manuscript.

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COMPLIANCE WITH ETHICAL STANDARDS

Disclosure of potential conflicts of interest: All authors declare no conflict of interest.

Research involving Human Participants and/or Animals: This article does not contain any studies with human participants. Rabbits used for antibodies preparation were approved by the Laboratory Animal Management Committee of Yangtze University (Approval No. 20161212).

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