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**Functional γ-Aminobutyrate Shunt in *Listeria monocytogenes*: Role in Acid Tolerance and Succinate Biosynthesis**

Conor Feehily, Conor P. O’Byrne, Kimon Andreas G. Karatzas

Bacterial Stress Response Group, Microbiology, School of Natural Sciences, National University of Ireland, Galway, Galway, Ireland

*Listeria monocytogenes*, the causative agent of human listeriosis, is known for its ability to withstand severe environmental stresses. The glutamate decarboxylase (GAD) system is one of the principal systems utilized by the bacterium to cope with acid stress, a reaction that produces γ-aminobutyrate (GABA) from glutamate. Recently, we have shown that GABA can accumulate intracellularly under acidic conditions, even under conditions where no extracellular glutamate-GABA exchange is detectable. The GABA shunt, a pathway that metabolizes GABA to succinate, has been described for several other bacterial genera, and the present study sought to determine whether *L. monocytogenes* has this metabolic capacity, which, if present, could provide a possible route for succinate biosynthesis in *L. monocytogenes*. Using crude protein extracts from *L. monocytogenes* EGD-e, we show that this strain exhibits activity for the two main enzyme reactions in the GABA shunt, GABA aminotransferase (GABA-AT) and succinic semialdehyde dehydrogenase (SSDH). Two genes were identified as candidates for encoding these enzyme activities, *argD* (GABA-AT) and *lmo0913* (SSDH). Crude protein extracts prepared from a mutant lacking a functional *argD* gene significantly reduced GABA-AT activity, while an *lmo0913* mutant lost all detectable SSDH activity. The deletion of *lmo0913* increased the acid tolerance of EGD-e and showed an increased accumulation of intracellular GABA, suggesting that this pathway plays a significant role in the survival of this pathogen under acidic conditions. This is the first report of such a pathway in the genus *Listeria*, which highlights an important link between metabolism and acid tolerance and also presents a possible compensatory pathway to partially overcome the incomplete tricarboxylic acid cycle of *Listeria*.

*L. monocytogenes* is a food-borne pathogen that causes listeriosis, a disease with a mortality rate of up to 30% (1), which has recently shown an increase in the number of reported cases across Europe (2). Almost 99% of cases are associated with a contaminated food source (3), and as such, *L. monocytogenes* is of major concern to the food industry. This bacterium is quite resilient to environmental stresses, including an ability to grow under acidic conditions with pH values as low as 4.5 and to survive at pH values as low as 2.0 for extended periods (4–6). This characteristic is important for its pathogenicity because it allows the bacterium to survive the acidic environment in various foods and in the stomach of the potential host.

To counteract this threat, strains of *Listeria* can employ a variety of acid resistance mechanisms, including the arginine deiminase system (7), the adaptive acid tolerance response (ATR) (8), and the glutamate decarboxylase (GAD) system (9). It is clear, however, that there is a degree of strain-to-strain variation in terms of the ability to cope with an acid challenge (9–11). The GAD system operates to maintain the internal pH homeostasis of the bacterium. Upon acid exposure, an extracellular molecule of glutamate (Glu) is taken up by an antiporter (GadT) and decarboxylated intracellularly by a Glu decarboxylase enzyme (GadD) to form γ-aminobutyric acid (GABA). This reaction consumes a proton, thereby increasing the intracellular pH. GABA is then excreted via GadT in exchange for a further molecule of Glu (Fig. 1).

*Listeria monocytogenes* can possess up to three genes encoding Glu decarboxylases (gadD1, gadD2, and gadD3) and two genes encoding antiporters (gadT1 and gadT2), arranged into three operons (gadD1T1, gadD2T2, and gadD3) (12); however, strains of serotype 4 lack gadD1T1. Recent work from our laboratory has demonstrated the existence of the intracellular GAD system (GAD), which converts intracellular Glu to GABA and contributes significantly to acid resistance (11, 13). The activity of this system results in the accumulation of high levels of GABA, under acidic conditions even in the absence of GABA export.

In *Escherichia coli*, GABA is used as both carbon and nitrogen sources and is metabolized via the GABA shunt pathway (Fig. 1) (14, 15). This pathway incorporates two further enzymes downstream from Glu decarboxylase. A GABA/α-ketoglutarate aminotransferase (GABA-AT) (GatB) removes the amino group from GABA to form succinic semialdehyde (SSA) and Glu. The SSA is then oxidized through the activity of succinic semialdehyde dehydrogenase (SSDH) (GatD) to form succinate (16). While the GABA shunt pathway has not been extensively studied in bacteria, it is thought to play a role in Glu metabolism, anaplerosis, and antioxidant defense (17, 18). The use of arginine, ornithine, and agmatine as nitrogen sources by *E. coli* relies on the GABA shunt pathway, as these are converted first to putrescine and subsequently to GABA (19). In plants and mammals, the pathway has been described as an alternative route to produce succinate, bypassing two enzymes of the tricarboxylic acid (TCA) cycle, namely, α-ketoglutarate dehydrogenase and succinyl coenzyme A (CoA) synthetase (Fig. 1) (16, 20). For *L. monocytogenes*, the GABA shunt may represent a potential route to compensate for the incomplete TCA cycle identified through sequencing and biochemical analysis (21, 22). Genome sequencing has shown an absence of genes encoding α-ketoglutarate dehydrogenase, succinyl-
FIG 1 Glutamate decarboxylase system and GABA shunt pathway of L. monocytogenes. Shown is the proposed model for the metabolism of GABA in L. monocytogenes. Glu, is decarboxylated to GABA, by GadD. The GABA can either be exported by GadT in exchange for another Glu or enter the GABA shunt pathway. Here GABA, donates its amino group to CoA synthetase, and malate dehydrogenase (Fig. 1). The complete TCA cycle of L. monocytogenes is shown, with the missing steps marked with an “X.” The GABA shunt pathway can provide an alternative source of succinate for the bacteria.

The aim of this study was to determine if L. monocytogenes possesses the two metabolic steps comprising the GABA shunt pathway. Second, strains with disruptions or deletions in genes predicted to be involved in this pathway (argD and lmo0913) were analyzed for effects on the respective activities in order to confirm their roles. Finally, due to the fact that the GAD system and the GABA shunt are linked, the main metabolites of the pathway were quantified in the cell in response to acid treatment.

**TABLE 1** Bacterial strains and plasmids used in this study

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Relevant property(ies)</th>
<th>Source or reference</th>
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</thead>
<tbody>
<tr>
<td><strong>L. monocytogenes</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EGD-e</td>
<td>Serovar 1/2a, wild type</td>
<td>S. Foster</td>
</tr>
<tr>
<td>EGD-e argD::pLSV101 mutant</td>
<td>pLSV101 plasmid insertion into the argD gene</td>
<td>This study</td>
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<tr>
<td>EGD-e Δlmo0913 mutant</td>
<td>EGD-e with an in-frame 1,407-bp deletion of the lmo0913 gene</td>
<td>This study</td>
</tr>
<tr>
<td>EGD-e Δlmo0913 pKAK0913 mutant</td>
<td>Deletion of lmo0913 complemented with a cloned copy of the full lmr07023 gene</td>
<td>This study</td>
</tr>
<tr>
<td>10403S</td>
<td>Serovar 1/2a, wild type</td>
<td>K. Schauer</td>
</tr>
<tr>
<td>10403S Δlmrg_02013 mutant</td>
<td>10403S with an in-frame 1,407-bp deletion of the lmr07023 gene</td>
<td>K. Boor</td>
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<tr>
<td>10403S Δlmrg_02013 pKAK0913 mutant</td>
<td>Deletion of lmr07023 complemented with a cloned copy of the full lmr07023 gene</td>
<td>26</td>
</tr>
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<td><strong>E. coli</strong></td>
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<tr>
<td>K-12 BW25113</td>
<td>Δ(araD-araB)567 ΔlacZ4787(:,rrnB-3) lambda` rph-1 Δ(rhaD-rhaB)568 hsdR514</td>
<td>NBRP (Japan), E. coli</td>
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<tr>
<td>JW2636</td>
<td>BW25113 ΔgabD</td>
<td>NBRP (Japan), E. coli</td>
</tr>
<tr>
<td>JW2636::pKAK0913</td>
<td>pKAK0913 mutant possessing a full copy of lmr07023 from 10403S cloned into pKSV7</td>
<td>This study</td>
</tr>
<tr>
<td><strong>Plasmid</strong></td>
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<tr>
<td>pKAK0913</td>
<td>pKSV7 shuttle vector carrying a full copy of lmr07023 used to complement deletion of lmo0913 or gabD</td>
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</table>

**TABLE 2 Primers used in this study**

<table>
<thead>
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<tr>
<td>lmo0913 A</td>
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<td>lmo0913 B</td>
<td>TGGCCTTGTTCATCAAAATACCACCTCC</td>
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<td>lmo0913 C</td>
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<td>lmo0913 D</td>
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<td>lmo0913 For</td>
<td>GGTTCTATGCTACGG-GAC</td>
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<tr>
<td>lmo0913 Rev</td>
<td>CATCCGTTATGCCCC</td>
</tr>
<tr>
<td>lmo0913 Comp-F</td>
<td>AAGGGTCGCGAGAGTCTTTTTGTAGGCC</td>
</tr>
<tr>
<td>lmo0913 Comp-R</td>
<td>CCGGATCTAAGTGTCCGATCTGTCG</td>
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</table>

**MATERIALS AND METHODS**

**Bacterial strains and growth.** L. monocytogenes wild-type strain EGD-e and its isogenic lmo0913 (in-frame deletion) (Δlmo0913) and argD (insertional disruption with pLSV101) (argD::pLSV101) mutants (24) were used in this study. L. monocytogenes wild-type strain 10403S along with its isogenic 10403S Δlmrg_02013 mutant were also used. Strains were grown on brain heart infusion (BHI) agar plates for 24 h at 37°C. Cultures grown overnight were set up in inoculating a single colony into 25 ml of BHI broth in 250-ml conical flasks with continuous shaking at 37°C. E. coli strains BW25113 and JW2636 were used as reference strains for the GABA shunt. These strains were grown in Luria-Bertani (LB) medium for 18 h at 37°C when needed and stored on LB agar plates. All strains and relevant properties are listed in Table 1.

**Generation of mutant and recombinant strains.** Mutants were generated by the splicing by overlap extension (SOEing) PCR method and allelic replacement, as previously described (25). The deletion of lmr07023 in 10403S was reported previously (26); however, the deletion of lmo0913 in EGD-e was newly generated by using the primers listed in Table 2. Plasmid pKAK0913, as described previously by Abram et al. (26), was used to complement the deletions of lmo0913 in EGD-e, lmr07023 in 10403S, and gabD in JW2636. This plasmid carries a full copy of lmr07023 from 10403S. Positive transformants were selected through growth on BHI agar supplemented with chloramphenicol (10 μg/ml) or LB agar supplemented with ampicillin (100 μg/ml) and confirmed by PCR. PCR amplification of DNA for use in cloning and downstream work was carried out by using the high-fidelity Velocity DNA polymerase (Bioline), while screening was carried out by using Biotaq DNA polymerase (Bioline).
GABA-AT and SSDH assays. Crude protein assays to detect the presence of GABA-AT and SSDH activities were carried out based on an assay described previously by Bartsch et al. (27) but modified in order to run it in a 96-well-plate format.

Protein extraction. To prepare crude protein extracts, a culture of bacteria was grown overnight in 200 ml of BHI (L. monocytogenes) or LB (E. coli) medium at 37°C with continuous shaking. The pellet was retained after centrifugation at 12,000 × g for 10 min, followed by two wash steps in wash buffer (10 mM NaCl, 10 mM Na₂O₅P [pH 7.0]). Pellets were then resuspended in 4 ml of a sonication buffer optimized for their downstream application. For GABA-AT, the sonication buffer comprised 20 mM Na₂O₅P, 0.01 mM pyridoxal phosphate, 5 mM mercaptoethanol, 1 mM phenylmethylsulfonyl fluoride (PMSF), and 1 mM diithiothreitol (DTT) (pH 7.0). For SSDH, it comprised 100 mM sodium phosphate, 1 mM DTT, 1 mM PMSF, and 9% glycerol (pH 9.5). Samples were then sonicated on ice by six 30-s pulses at a 16-μm amplitude, allowing 30 s of rest between each pulse to lyse the cells, followed by centrifugation at 6,000 × g for 10 min and retention of the supernatant. The supernatant was then centrifuged a final time at 15,000 × g for 15 min to remove any remaining cell debris. Protein concentrations for both assays were normalized to 7 mg ml⁻¹ after determining each sample concentration by using the Bio-Rad DC protein assay (Bio-Rad).

GABA-AT assay. One hundred microliters of protein was added to 1,000 μl reaction mix (85 mM Tris base, 15 mM Tris-HCl, 0 to 100 mM GABA, 5 mM α-ketoglutarate, 0.02% bovine serum albumin [BSA]) and incubated at 37°C for 1 h. In a 96-well plate, the SSA produced by the reaction was measured by the use of a GABase assay described previously by O’Byrne et al. (28), using 80 mM 2-aminoethyl hydrogen sulfate for the detection of SSA. Plates were incubated at 37°C in a Sunrise absorbance reader (Tecan, Salzburg, Austria), with optical density at 340 nm (OD340) readings being recorded every 60 s by a Tecan software (Tecan, Salzburg, Austria). Any background signal detected with 0 mM GABA was subtracted to highlight the increase in the level of SSA production.

SSDH assay. NADP to a final concentration of 1 mM was added to 2 ml of a normalized protein preparation (7 mg ml⁻¹), and 150 μl of this mix was added to 100 μl of SSA (0 to 1.0 mM) in a 96-well plate. Plates were incubated at 37°C in a Tecan Sunrise absorbance reader, with OD₃₄₀ readings being recorded every 60 s by Magellan software. GABase assay. GABAₐ, extracellular GABA (GABAₐ), as well as intracellular SSA concentrations were measured as previously described (13, 28). Cultures were grown overnight at 37°C with aeration in BHI medium. Prior to the GABA measurements, the pH of the culture was lowered to 4.0 with HCl. Extractions were made after 1 h of pH treatment. Non-HCl-treated cultures were used as negative controls.

Acid survival assays. Cultures of bacteria were grown overnight at 37°C in BHI medium. The pH of these cultures was lowered to pH 2.5 by diluting 1:1 in phosphate-buffered saline (PBS). Dilutions were plated in triplicate onto BHI agar and incubated overnight at 37°C. Colonies were counted to determine the number of surviving cells.

Measurement of Glu and succinate concentrations. Glu and succinate concentrations were measured in the same samples tested for GABA and SSA. Both methods were performed by the use of enzymatic kits supplied by Roche Bio-Pharm. The methods were adapted to fit a 96-well-plate format. All reagents were scaled down from use in a 3-ml (final volume) reaction mixture to use in a 300-μl (final volume) reaction mixture. Standards of either monosodium glutamate (Sigma) or sodium succinate (BDH) were used to establish a standard curve for the enzymatic reaction. One hundred microliters of sample or standards (0 to 1.0 mM) was added to 200 μl of kit reagents and incubated at 25°C (Glu) or 37°C (succinate) in a 96-well plate in a Tecan Sunrise absorbance reader, with the OD₅₀₀ (Glu) or OD₃₄₀ (succinate) being recorded every 60 s by Magellan software. Concentrations of the metabolites were determined based on the standard curve generated from known sample concentrations.

Statistical analysis of results. Experiments were carried out with three biological replicates and at least two technical replicates for each sample. Significant differences between samples tested were determined by using a paired Student t test. Results were considered significant when they possessed a P value of <0.05. Error bars indicating standard deviations from the means are displayed on graphs.

RESULTS

L. monocytogenes possesses activity for both enzymes of the GABA shunt. The sequences corresponding to GabT and GabD from E. coli were used to search the L. monocytogenes genome for homologues that might encode GABA aminotransferase (GABA-AT) and succinic semialdehyde dehydrogenase (SSDH) activities, respectively. BLAST analysis showed that ArgD from L. monocytogenes EGD-e shared 52% similarity and 34% identity to GabT. In L. monocytogenes EGD-e, the predicted SSDH is Lmo0913, with 67% similarity and 49% identity to GabD. The corresponding gene product in L. monocytogenes 10403S is LMRG_02013, with 68% similarity and 49% identity. These two proteins, ArgD and Lmo0913/LMRG_02013, appeared frequently throughout the published sequences of L. monocytogenes. Twenty-four out of 25 strains searched possessed a homologue of Lmo0913/LMRG_02013 with >98% identity, while 14 out of 25 possessed an ArgD homologue with >93% identity. Most notable is LO28, which does not appear to have similar proteins. Evidence suggested that lmo0913 produces a monocistronic transcript (29) with a termination site predicted immediately downstream. The transcription of lmo0913 appears to be σ⁷₀ dependent (26).

The first step in the GABA shunt is the conversion of GABA to SSA. Crude protein extracts were tested for their abilities to produce SSA from GABA, as described above. The co-occurrence of these protein preparations (7 mg ml⁻¹) for 1 h with GABA (50 and 100 mM) resulted in the production of SSA (1.79 and 3.94 μM SSA min⁻¹ mg protein⁻¹, respectively) (Fig. 2). In wild-type strain EGD-e, the concentration of SSA produced increased in proportion to the concentration of GABA added. When a mutant with a disruption in a putative GABA-AT gene (argD) was used,
the level of production of SSA was reduced (1.57 and 2.34 μM SSA min⁻¹ mg protein⁻¹, respectively; $P$ value of <0.05), indicating an involvement of this gene in the metabolic step. The effect of the disruption of argD was more apparent at a higher concentration of 100 mM GABA, whereas at 50 mM GABA, the difference did not appear to be significant. The reduced SSA levels in the absence of argD suggest that an alternative GABA-AT activity is also present in the cell extracts. GABA-AT activity was also observed for the 10403S strain but at much lower levels than for EGD-e (0.55 μM SSA min⁻¹ mg protein⁻¹ with 100 mM GABA).

The oxidation of SSA to succinate is the second step in the GABA shunt intermediates. The enzyme that carries out this step, the corresponding gene from 10403S was cloned into the EGD-e protein extract that was incubated with SSA (Fig. 3). The production of NADPH did not occur in the absence of the substrate, and its level of production increased in proportion to the concentration of the substrate (SSA). The deletion of the putative SSDH-encoding gene, lmo0913, resulted in a complete loss of activity for this reaction. To determine if Lmo0913 acts directly as an SSDH, the corresponding gene from 10403S was cloned into pKSV7, generating plasmid pKAK0913, and this plasmid was used to transform two strains of E. coli, BW25113 extracts were found to have SSA activity when incubated with SSA (Fig. 4). The disruption of the SSDH of E. coli, gabD, resulted in a complete loss of this activity. The supplementation of this mutant with lmr_02013 partially restored the SSDH activity.

No SSDH activity was detected in extracts from either strain 10403S or the lmr_02013 derivative of this strain; however, the supplementation of the knockout mutant with lmr_02013 resulted in the production of NADPH. As expected, the supplementation of the lmo0913 mutation in EGD-e restored the SSDH activity in this strain. It is interesting to note that LMRG_02013 from 10403S has six amino acid differences from Lmo0913 in EGD-e (deletions of the V2, F3, and L4 amino acids, as well as the E251/248D, G312/309A, and D352/349E substitutions). These data showed that Lmo0913/LMRG_02013 is responsible for the SSDH activity in L. monocytogenes and further suggest that the baseline SSDH activity varies between strains (Fig. 3 and 4).

Acid exposure affects intracellular concentrations of GABA shunt intermediates. The levels of Glu did not appear to significantly change in response to acid in both EGD-e and pLSV101::argD cells (Table 3); however, the deletion of lmo0913 resulted in a reduction in the levels of Glu from 0.53 mM to 0.45 mM ($P$ value of 0.03). As expected, GABA i concentrations were affected in EGD-e in response to acid. Both EGD-e and the pLSV101::argD strain showed 2.6- and 7.5-fold increases in GABA i, levels, reaching concentrations of 1.1 and 0.6 mM, respectively. However, a dramatic 9.2-fold increase in the GABA i concentration was observed for the EGD-e mutant, reaching a concentration of ~3.5 mM. This, along with a significant decrease in the Glu concentration, suggests a highly active GAD system in this strain. The increase seen in the SSA concentration before and after acid treatment for EGD-e was not significant, and the increase in the concentration seen for the EGD-e::pLSV101 strain was close to the detectable range of the assay, but the increase for the EGD-e mutant was higher, as would be expected for a gene involved in the breakdown of SSA. The succinate concentration increased to a small extent for both EGD-e and the argD::pLSV101 strain after acid treatment; however, the increase for the EGD-e mutant was not significant ($P$ value of 0.165).

Lmo0913 affects acid resistance of L. monocytogenes EGD-e. EGD-e showed a reduction in cell numbers by ~8 log cycles, and the argD mutant strain demonstrated a similar reduction (Fig. 5).
and Lmo0913 via the GABA shunt pathway. As the GABA shunt is not caused by the presence of the acid resistance phenotype (data not shown). This growth decrease was accompanied by 9.2-fold increases in GABAi and argD gene on the plasmid, since strains containing pKSV7 also grew more slowly than the wild type.

**DISCUSSION**

We have previously shown that *L. monocytogenes* possesses a GADi system that leads to GABA accumulation under acidic conditions (11). Here we report that this GABA can be metabolized by ArgD and Lmo0913 via the GABA shunt pathway. As the GABA shunt is coupled to the GAD system, which is the principal mechanism of acid tolerance in *L. monocytogenes*, we investigated the acid tolerance phenotypes of both the argD:pLSV101 and ∆lmo0913 strains. Interestingly, these mutants yielded quite contrasting acid tolerance phenotypes. While the disruption of argD slightly hindered the cells’ ability to cope with acid treatment, the deletion of lmo0913 increased survival by over 5 log cycles (Fig. 5). This significant increase highlights a role for the GABA shunt in acid tolerance. The acid-resistant phenotype of the ∆lmo0913 mutant was accompanied by 9.2-fold and 1.1-fold increases in GABA, and SSA concentrations, respectively (Table 3). The increases in the concentrations of these metabolites are likely to be caused by a diminished capacity for GABA metabolism in this mutant (Fig. 3).

Interestingly, EGD-e does not utilize the GAD system but only the GADi system. Recently, we demonstrated that GAD contributes to acid resistance independently of the antiport carried out by GAD. The fact that the deletion of lmo0913 gene is able to convert EGD-e from one of the weakest strains in terms of acid resistance to a moderate one (2 logs lower than that of 10403S, which is the most acid-resistant strain that we have identified, and 5 logs higher than that of EGD-e) (Fig. 5) through an increase in the GABA concentration underpins the importance of this system in acid resistance.

If the disruption of the GABA shunt by the deletion of lmo0913 is responsible for the GABA buildup and, subsequently, acid resistance, a similar result might be expected for the argD mutant. This, however, was not the case, suggesting that in this mutant, the transamination activity observed is probably carried out by an enzyme that compensates for the loss of argD. GsAB has 46% similarity and 28% identity to GabT, a GABA-AT (30). It was shown previously that for several bacteria, multiple copies of GABA shunt genes exist (31, 32), and this may well be the case for *L. monocytogenes*. BLAST searches for a secondary SSDH also revealed a likely candidate in Lmo0383, which has 54% similarity and 33% identity to the corresponding SSDH (GabD) in *E. coli*. Further work to characterize the roles of both gsaB and lmo0383 are under way to test their roles in the GABA shunt pathway.

When the SSDH activities of EGD-e and 10403S were compared, it was interesting that no SSDH activity was detectable under the conditions used for 10403S. The difference between the two strains may suggest differing degrees of importance of the GABA shunt between strains (although both belong to lineage II and share the same serotype, serotype 1/2a). Strain-to-strain variation is also exemplified by the fact that previously, a deletion of *lmg_02013* in 10403S did not confer acid resistance but instead resulted in a mild sensitivity to acid (26). As mentioned above, the main difference between the two strains is that EGD-e utilizes only

### TABLE 3 Intracellular concentrations of intracellular metabolites in stationary-phase cells

<table>
<thead>
<tr>
<th>Strain</th>
<th>Glutamate pH7.0 (mM)</th>
<th>Glutamate pH4.0 (mM)</th>
<th>GABA pH7.0 (mM)</th>
<th>GABA pH4.0 (mM)</th>
<th>SSA pH7.0 (mM)</th>
<th>SSA pH4.0 (mM)</th>
<th>Succinate pH7.0 (mM)</th>
<th>Succinate pH4.0 (mM)</th>
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</thead>
<tbody>
<tr>
<td>EGD-e</td>
<td>0.55 (±0.06)</td>
<td>0.53 (±0.05)</td>
<td>0.43 (±0.05)</td>
<td>0.61 (±0.08)</td>
<td>0.41 (±0.41)</td>
<td>0.52 (±0.52)</td>
<td>0.24 (±0.05)</td>
<td>0.44 (±0.07)</td>
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<tr>
<td>argD:pLSV101</td>
<td>0.59 (±0.02)</td>
<td>0.48 (±0.06)</td>
<td>0.51 (±0.02)</td>
<td>0.60 (±0.01)</td>
<td>0.24 (±0.02)</td>
<td>0.34 (±0.02)</td>
<td>0.17 (±0.01)</td>
<td>0.34 (±0.04)</td>
</tr>
<tr>
<td>∆lmo0913</td>
<td>0.53 (±0.01)</td>
<td>0.45 (±0.02)</td>
<td>0.35 (±0.06)</td>
<td>3.24 (±1.83)</td>
<td>0.53 (±0.53)</td>
<td>0.61 (±0.61)</td>
<td>0.24 (±0.05)</td>
<td>0.54 (±0.24)</td>
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</table>

*Statistically significant difference between wild-type and mutant strains under the conditions tested. A difference was identified to be statistically significant by using a paired Student’s t test (P < 0.05).*

**FIG 5** Acid survival of *L. monocytogenes* EGD-e and GABA putative shunt system mutants. Stationary-phase EGD-e and GABA shunt mutants were challenged at pH 2.5. Cell counts were taken every 20 min. Values are the means of data from three individual cultures, with the cell counts for each culture being the means of counts from three platings. The detection limit of the assay is highlighted by the dotted line, while the downward-pointing arrow indicates that subsequent counts dropped below this level.
GAD, and therefore, unlike 10403S, which utilizes both GAD and GAD, it must rely solely on the GABA shunt to metabolize the GABA generated during acid stress (11). Thus, it appears that the elevated levels of SSDH activity in EGD-e might be related to the larger GABA pools that exist in this strain during acid stress.

The existence of the GABA shunt pathway has significant implications for our understanding of the overall metabolism of L. monocytogenes. The genome sequence of L. monocytogenes lacks a complete set of enzymes for the classical TCA cycle model (21), and this is consistent with biochemical measurements (22). Two of the enzymes missing from L. monocytogenes are α-ketoglutarate dehydrogenase and succinyl-CoA synthetase, which are responsible for the conversion of α-ketoglutarate to succinate (Fig. 1). The GABA shunt pathway appears to serve different species of bacteria in a range of ways, from sporulation (33) to nitrogen metabolism (30). For L. monocytogenes, it is possible that the pathway can in part substitute for the lack of a functional TCA cycle. It is important to note, however, that the GABA shunt does not provide a means of completing the TCA cycle, since malate dehydrogenase, required to convert malate to oxaloacetate, is also absent from L. monocytogenes (21, 23). According to data reported previously by Trivett and Meyer (34), L. monocytogenes lacks a functional succinate dehydrogenase, required to convert succinate to fumarate. This presents yet another disruption in the classical TCA cycle and would indicate a dead end for the succinate produced by the GABA shunt pathway. However, their experiments did show activity for fumarate reductase. Evidence was presented previously which showed that fumarate reductase can compensate for a loss of succinate dehydrogenase (35, 36). This could putatively convert fumarate both secondary enzymes in the pathway and the regulatory mechanisms involved, we have shown that the GABA shunt pathway is functional in L. monocytogenes, providing this pathway with a means to synthesize the TCA intermediate succinate.

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