



# Acid Evolution of *Escherichia coli* K-12 Eliminates Amino Acid Decarboxylases and Reregulates Catabolism

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**ABSTRACT** Acid-adapted strains of *Escherichia coli* K-12 W3110 were obtained by serial culture in medium buffered at pH 4.6 (M. M. Harden, A. He, K. Creamer, M. W. Clark, I. Hamdallah, K. A. Martinez, R. L. Kresslein, S. P. Bush, and J. L. Slonczewski, Appl Environ Microbiol 81:1932–1941, 2015, <https://doi.org/10.1128/AEM.03494-14>). Revised genomic analysis of these strains revealed insertion sequence (IS)-driven insertions and deletions that knocked out regulators CadC (acid induction of lysine decarboxylase), GadX (acid induction of glutamate decarboxylase), and FNR (anaerobic regulator). Each acid-evolved strain showed loss of one or more amino acid decarboxylase systems, which normally help neutralize external acid (pH 5 to 6) and increase survival in extreme acid (pH 2). Strains from populations B11, H9, and F11 had an IS5 insertion or IS-mediated deletion in *cadC*, while population B11 had a point mutation affecting the arginine activator *adiY*. The *cadC* and *adiY* mutants failed to neutralize acid in the presence of exogenous lysine or arginine. In strain B11-1, reversion of an *rpoC* (RNA polymerase) mutation partly restored arginine-dependent neutralization. All eight strains showed deletion or downregulation of the Gad acid fitness island. Strains with the Gad deletion lost the ability to produce GABA (gamma-aminobutyric acid) and failed to survive extreme acid. Transcriptome sequencing (RNA-seq) of strain B11-1 showed upregulated genes for catabolism of diverse substrates but downregulated acid stress genes (the biofilm regulator *ariR*, *yhiM*, and *Gad*). Other strains showed downregulation of H<sub>2</sub> consumption mediated by hydrogenases (*hya* and *hyb*) which release acid. Strains F9-2 and F9-3 had a deletion of *fnr* and showed downregulation of FNR-dependent genes (*dmsABC*, *frdABCD*, *hybABO*, *nikABCDE*, and *nrfAC*). Overall, strains that had evolved in buffered acid showed loss or downregulation of systems that neutralize unbuffered acid and showed altered regulation of catabolism.

**IMPORTANCE** Experimental evolution of an enteric bacterium under a narrow buffered range of acid pH leads to loss of genes that enhance fitness above or below the buffered pH range, including loss of enzymes that may raise external pH in the absence of buffer. Prominent modes of evolutionary change involve IS-mediated insertions and deletions that knock out key regulators. Over generations of acid stress, catabolism undergoes reregulation in ways that differ for each evolving strain.

**KEYWORDS** acid, *Escherichia coli*, experimental evolution, GABA, low pH, RNA polymerase, decarboxylase, *fnr*

Enteric bacteria need to survive a wide range of pH values throughout the human intestinal tract. The pH of the intestinal tract ranges from 1.5 to 3.5 in the stomach, increases to 4 to 7 in the duodenum, reaches 7 to 9 in the jejunum, and decreases to pH 5 to 6 in the cecum (1). On a microscopic scale, extreme gradients of pH occur across

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the intestinal epithelium where enterobacteria adhere. To cope with such conditions, *Escherichia coli* maintains a cytoplasmic pH homeostasis of 7.3 to 7.8 during growth in an external pH range of 5 to 9 and survives extreme acid levels below pH 2.0 (2, 3). *E. coli* uses intricate regulatory networks such as acid resistance systems to survive this wide range of pH values (4). Several acid resistance systems of *E. coli* include decarboxylases of the amino acids lysine, arginine, glutamate/glutamine, and ornithine (5). Each amino acid-dependent decarboxylase catalyzes the decarboxylation of its respective amino acid by consuming a proton and releasing an amine or polyamine, i.e., cadaverine, agmatine, gamma-aminobutyric acid (GABA), and putrescine, respectively. The acid-dependent release of polyamines and of the neurotransmitters GABA and agmatine is of interest in studies of tumorigenesis (6) and the gut-brain axis (7). An important feature of acid-dependent decarboxylases is their different pH ranges for induction and acid consumption: lysine decarboxylase activity (CadA) peaks at approximately pH 6, arginine (Adi) has activity at pH 5, and glutamate (GadA and GadB) has activity below pH 5 (8, 9).

One way to investigate how *E. coli* adapts to contend with stress such as low pH is by experimental evolution (10–12). Experimental evolution of *E. coli* has been conducted under the stress of medium buffered at pH 4.6, the low end of the growth range for this microorganism (13). This pH range is relevant to the digestive tract as bacteria entering the stomach experience the in-between levels upon emerging from the inhibitory condition of stomach acid. Thus, via the fecal-oral route, enteric bacteria naturally undergo periods of selection at pH 4.6 to 5.0. The acid-evolved clones from our experiment (13) out-compete the ancestral strain growing at pH 4.6. Sequence analysis of acid-evolved clones reveals various categories of mutations relevant to pH response, such as single nucleotide polymorphisms (SNPs) in RNA polymerases *rpoB* and *rpoC*, as well as unexplained phenotypes such as loss of lysine decarboxylase activity.

While whole-genome sequencing provides a wealth of information about genetic differences between microorganisms, this type of analysis often fails to reveal complex mutations such as those mediated by insertion sequences (ISs). An updated genome resequencing pipeline, breseq, version 0.27.1, detects split-read alignments, where a single read maps to disjointed areas of the genome (14). In our analysis, breseq, version 0.27.1, revealed evidence of IS5-mediated interruptions and deletions, in particular, those that inactivate regulatory genes such as *cadC*, *gadX*, and *fnr*. Detecting these additional mutation types altered our view of the genomic changes in the acid-evolved clones.

Previously, we hypothesized that mutations in the acid-evolved clones could modulate the expression of acid-induced genes that are energetically wasteful under the conditions of the evolution experiment (13). Under our evolution conditions, the buffered environment would have prevented any rise in pH mediated by acid protection systems such as lysine decarboxylase; thus, the high expression level of CadA near pH 5 (15) would be energetically wasteful. We also hypothesized that the observed mutations in RNA polymerase (RNAP) subunit genes could have modulated the expression of decarboxylase systems. Lysine and arginine decarboxylases are among the most highly inducible pH-regulated genes (16, 17), and thus their overexpression could lead to a substantial energy cost.

To investigate the above hypotheses, we updated our genome sequence analysis, and we compared the evolved strains with respect to lysine, arginine, and glutamate decarboxylase activities. We tested constructs with reverted RNAP mutations for decarboxylase activity. We also performed transcriptome sequencing (RNA-seq) to analyze the overall expression patterns of our acid-evolved strains.

## RESULTS

**Genetic analysis of acid-evolved isolates.** Previously, we had conducted a 2,000-generation experimental evolution of 24 bacterial populations under moderately acidic conditions (pH 4.6 to 4.8). Two clones each from four independent bacterial popula-

**TABLE 1** Amino acid decarboxylase mutations and phenotypes of acid-evolved strains

Strain <sup>a</sup>	Lysine decarboxylase		Arginine decarboxylase		Glutamate decarboxylase	
	Activity <sup>b</sup>	Mutation	Activity	Mutation <sup>d</sup>	Activity	Mutation <sup>e</sup>
B11-1	—	<i>cadC::IS5</i>	—	<i>adiY</i> (C→G)	+	
B11-2	—	$\Delta([cadA]-[cadC])^c$	—	<i>adiY</i> (C→G)	+	
F9-2	+		+		—	$\Delta(gadX-slp)$
F9-3	+		+		—	$\Delta(gadX-slp)$
F11-1	—	<i>cadC::IS5</i>	+		—	$\Delta(gadX-slp)$
F11-2	—	$\Delta([ghoS]-cadC)$	+		—	$\Delta(gadX-slp)$
H9-1	—	<i>cadC::IS5</i>	—	<i>adiY</i> ←IS5← <i>adiA</i>	+	
H9-2	—	<i>cadC::IS5</i>	—	<i>adiY</i> ←IS5← <i>adiA</i>	+	

<sup>a</sup>Strains were isolated from the experimentally evolved populations B11, F9, F11, and H9 (13).

<sup>b</sup>Activity is shown as follows: +, the decarboxylase system is active; —, the decarboxylase system activity is decreased or absent.

<sup>c</sup>Brackets indicate that the mutation takes place in the middle of the gene. Delta symbols indicate deletion. En dash indicates that deletion includes all genetic material between indicated genes.

<sup>d</sup>IS5 with double arrows (←IS5←) denotes the insertion of an IS5 element into the intergenic region; the arrows indicate the direction of transcription of flanking genes.

<sup>e</sup>Deletion of most or all of the Gad acid fitness island: *gadA gadX gadW mdtF mdtE gadE hdeD hdeA hdeB yhiD dctR slp*.

tions (B11, F9, F11, and H9) were isolated for genomic analysis. The genomes of our acid-evolved strains (13) were assembled and compared with the *E. coli* K-12 W3110 reference genome using the breseq computational pipeline, version 0.23 (18). Our original results were largely consistent with those of an independent analysis using GATK tools at the Broad Institute that were available in 2014 (<https://software.broadinstitute.org/gatk/>). For the present work, we sought evidence for previously undetected mutations using an updated breseq, version 0.27.1 (19). Our revised analysis identified additional mutations including eight insertion knockouts as well as seven IS-mediated deletions (Table 1; see also Table S1 in the supplemental material). In addition to use of the updated pipeline, our revised analysis includes a less conservative view of mutation “calls” by the pipeline. In some cases, visual inspection of the sequence reads for a low-scoring call showed a mutation clearly present in all reads. We also compared the pairs of sister strains (strains from a shared population of the evolution experiment) for mutations called in one strain but not the other.

The newly identified mutations include IS-mediated knockouts of *cadC* (20), which encodes the pH-sensing transcriptional activator of *cadA* (lysine decarboxylase) and *cadB* (lysine-cadaverine antiporter) (21). The inserted regions consist of an IS5 element, 11 copies of which occur in the reference sequence of strain W3110 (22). The *cadC::IS5* knockouts were found in sequenced strains from three of the four acid-evolved populations (B11, F11, and H9). For population B11, strain B11-1 showed the *cadC::IS5* knockout, whereas B11-2 showed an IS5-mediated deletion that includes *cadA* and *cadB* ( $\Delta([cadA]-[cadC])$ ), where the brackets indicate that the mutation takes place in the middle of the gene. Strain F11-2 showed an IS5-mediated deletion,  $\Delta([ghoS]-cadC)$ , at the same position as a *cadA::IS5* insertion in the sister strain F11-1 (Tables 1 and S1). This more extensive deletion covers *cadABC* as well as the nutritional stress toxin-antitoxin system GhoT/GhoS (23). In all, we found four independent IS-mediated mutations in *cadC* among six strains. Populations B11 and H9 also show point mutations in regulators of arginine decarboxylase (13).

In addition, the strains from two populations (F9 and F11) showed deletions of all or part of the Gad acid fitness island, which includes *gadA* (glutamate decarboxylase) with its regulators *gadX*, *gadY*, and *gadE*, as well as extreme-acid periplasmic chaperones (*hdeA* and *hdeB*) and multidrug transporters (*mdtE* and *mdtF*) (24).

We thus found an overall pattern in which our strains that evolved at a constant narrow range of pH 4.6 to 4.8 experience strong selection pressure to lose or down-regulate one or more of the acid-inducible degradative amino acid decarboxylases (Table 1). Population B11 lost decarboxylases for lysine and arginine, F9 lost glutamate, F11 lost lysine and glutamate, and H9 lost lysine and arginine.

Another major regulator, *fnr*, was lost only in population F9 (strains F9-2 and F9-3) within the IS-mediated deletion  $\Delta([ynaJ]-[ogt])$  (Table S1). The most prominent anaerobic regulator of *E. coli*, Fnr normally activates genes under low-oxygen conditions (25–29).

**Lysine and arginine decarboxylase activities correspond to the acid-evolved genotypes.** While the genetic evidence supports our hypothesis of decarboxylase loss, many mutations in our strains could contribute unknown effects to their phenotypes. We sought to confirm the phenotypes by conducting assays of lysine and arginine decarboxylase under conditions permitting pH rise in the medium, using a modified Møller broth test (13). This pH rise is believed to be the main mechanism by which the lysine, arginine, and glutamate decarboxylases contribute to growth in moderate acid (external pH 4.5 to 6.5) and to survival in extreme acid (external pH 1 to 3) (2, 4, 15).

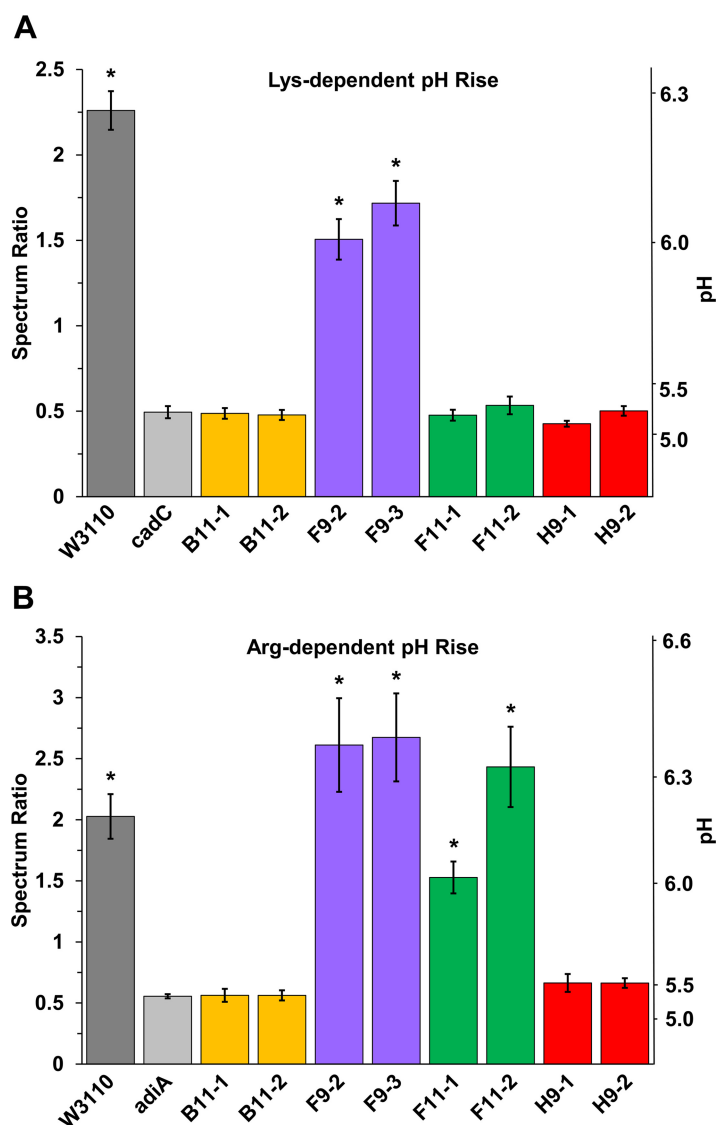
The lysine decarboxylase-associated pH rise was measured for our eight acid-evolved strains (Fig. 1A). Our modified Møller broth for the microplate test allowed precise measurement of the peak spectral intensities for the protonated and deprotonated forms of the dye bromocresol purple. The dye ratios were correlated with pH values by a standard curve (Fig. S1). In all trials, the strains showing *cadC* knockout or deletion showed 24-h endpoint pH values comparable to those of the control strain W3110 *cadC::kan* (Fig. 1A, light gray bar). Only strains F9-2 and F9-3 show decarboxylase activities significantly higher than those of the *cadC* knockout based on Tukey method comparison ( $P < 0.0001$ ).

For the same strains, we measured the pH rise associated with arginine decarboxylase (Fig. 1B). Strains B11-1 and B11-2 contain point mutations in the transcriptional activator *adiY*, which affects expression of arginine decarboxylase *adiA* (30). In addition, our new analysis showed that strain H9-1 contains an IS5 insertion found in the intergenic region between *adiY* and *adiA* (Table 1). Both H9 and B11 populations showed a significantly smaller arginine-dependent pH rise than that seen in W3110 ( $P < 0.001$ ). The spectrum ratios (and inferred pH levels) were comparable to those of the negative-control strain, W3110 *adiA::kan* ( $P > 0.98$ ). In contrast, the strains without *adiY* mutations showed a pH increase comparable to that of W3110. The strains F11-1, F11-2, F9-2, and F9-3 had a pH rise statistically equivalent to that of W3110 ( $P > 0.40$ ).

We also tested the hypothesis that the RNAP gene mutations found in our acid-evolved strains might modulate decarboxylase activities. For strains B11-1, F9-2, and H9-1, we replaced each mutant allele with an RNAP subunit with the reverted parental allele, using cotransduction of a linked Kan<sup>r</sup> marker. Each Kan<sup>r</sup> marker was converted to *frt* scar. The constructs thus obtained were B11-1 *rpoC*<sup>+</sup> (JLSE0083thiG1A::frt) (13), F9-2 *rpoC*<sup>+</sup> (JLSE0137thiH1C::frt), and H9-1 *rpoB*<sup>+</sup> (JLSE0079thiH2C::frt). Of these constructs, B11-1 *rpoC*<sup>+</sup> (JLSE0083thiG1A::frt) showed a modest recovery of arginine decarboxylase activity and pH rise compared to that of strain B11-1 ( $P < 0.0001$ ) (Fig. 2A). Thus, the *rpoC* mutation V507L appears to have some modulatory effect on *adi* expression.

For comparison, we saw no such restoration of *AdiA* activity after *rpo* reversion in F9-2 or in H9-1 (Fig. 2B) ( $P > 0.5$ ). The arginine-dependent pH rise actually increased for strain F9-2 compared to the level in the ancestor W3110; this is the only acid-evolved strain that showed a significant increase in any decarboxylase activity. Population H9 has an IS5 insertion in the intergenic region of *adiA* to *adiY* which may depress *adiA* expression; this insertion effect would be unlikely to be overcome by reversion of an *rpo* mutation. In contrast to H9-1, B11 has no IS5 insertion in the *adi* region, instead having a point mutation in the activator *adiY*. In B11, the *adiY* and *rpoC* point mutations may be exhibiting a combined effect to depress *AdiA*, which would explain the partial restoration we observed with *rpoC* reversion.

**GABA production and extreme-acid survival correspond to the Gad genotypes.** Given the large *gad* deletions we found in the F9 and F11 populations (Table 1), we sought evidence for glutamate decarboxylase activity. The *GadA* enzyme is generally expressed at around pH 5 but is most active under extreme-acid conditions such as pH 2 (4). Glutamate decarboxylase converts glutamate into GABA, which accepts a proton in the process, reducing pH stress (4). Glutamate decarboxylase activity can be inferred

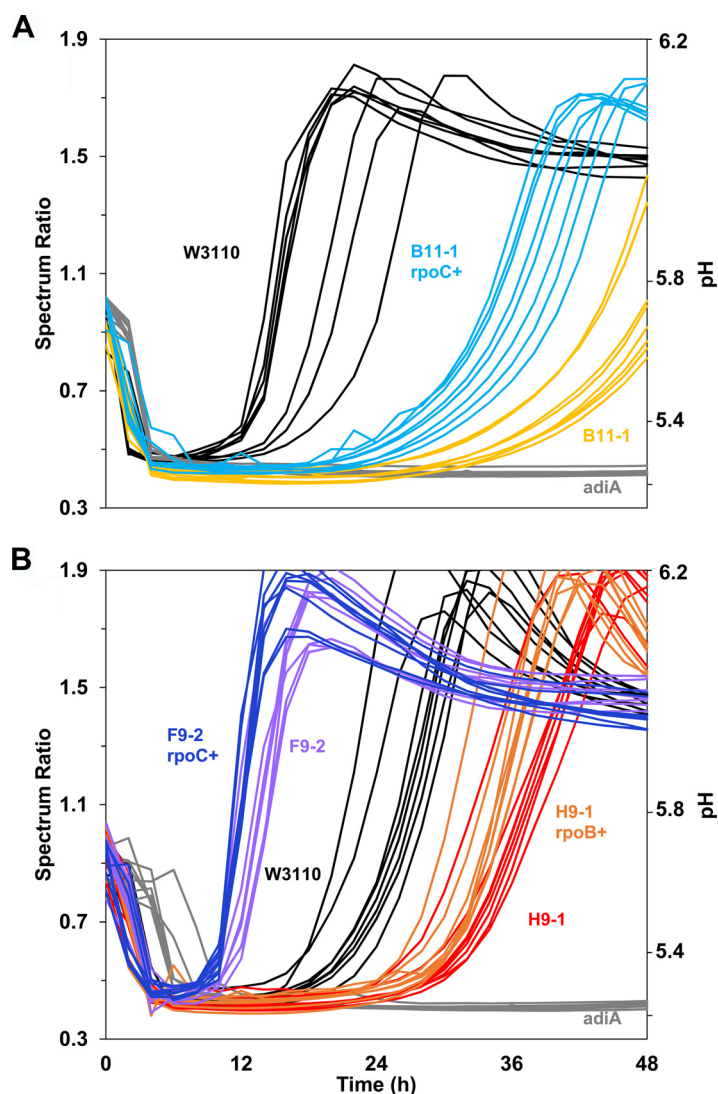


**FIG 1** pH rise of acid-evolved mutants cultured with lysine or arginine. Bacteria were inoculated in microplate wells containing 200  $\mu$ l of modified Møller broth medium at pH 6.8 supplemented with 2.5 g/liter L-lysine (A) or at pH 5.5 supplemented with 2.5 g/liter L-arginine (B). The microplates were incubated for 24 h at 37°C. The terminal spectrum ratio for bromocresol purple absorbance ( $A_{570-590}/A_{400-450}$ ) was converted to indicate pH (scale at right) (see Fig. S1). Error bars indicate standard errors of the means ( $n = 9$ ). One-way ANOVA tests were run with the Tukey comparisons to identify significantly different groups. Asterisks indicate strains for which the results are significantly different from those of the knockout strain ( $P < 0.05$ ).

from GABA production measured by gas chromatography-mass spectrometry (GC-MS) (31). GABA was measured by GC-MS after anaerobic overnight cultures were exposed at pH 2.0. We found that strains F9-2, F9-3, F11-1, and F11-2 produced no detectable GABA (Fig. 3A). However, the other strains produced millimolar quantities of extracellular GABA, comparable to the level of strain W3110 ( $P > 0.05$ ). Thus, we confirmed that the GABA-negative strains are the ones with deletions in the Gad acid fitness island.

Strains that produce GABA should also test positive for the Gad acid survival system (4, 15). All the sequenced populations show a significant loss of survival at pH 2.0 (Fig. 3B). Strains from populations F9 (F9-2 and F9-3) and F11 (F11-1 and F11-2), which have deletions of the *gad* region, showed acid survival rates below our level of detection ( $\log_2$  pH 2/pH 5.5 values below  $-6$ ). Strains B11-1 and B11-2 survived at rates approximately 10% that of the ancestor ( $P < 0.0001$ ). Strains H9-1 and H9-2 survived at

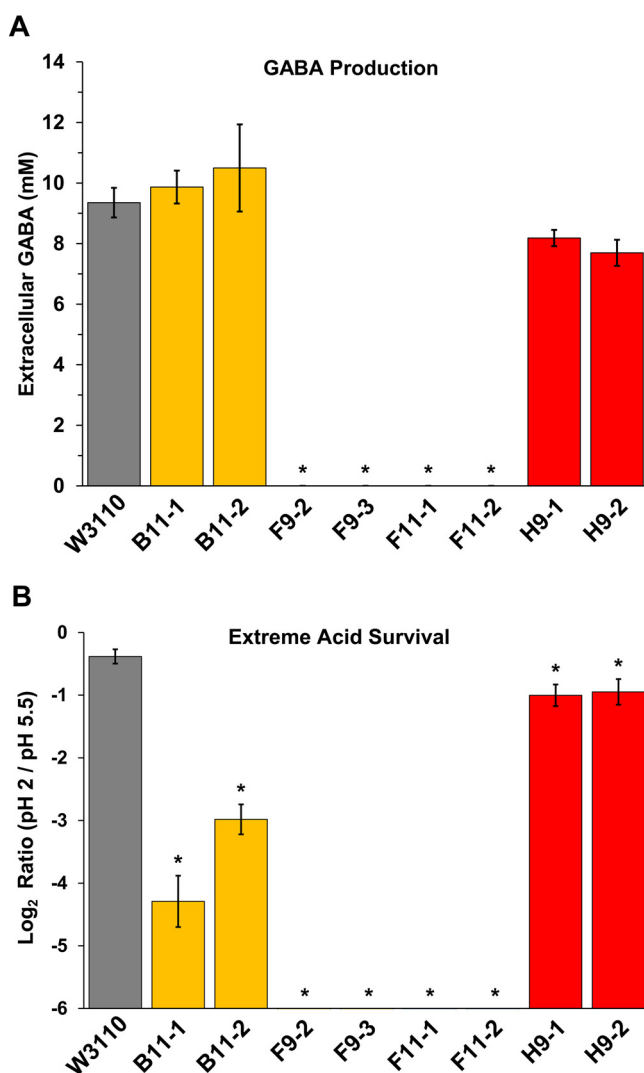




**FIG 2** Arginine decarboxylase assay. *RpoC* modulates arginine-dependent pH rise in strain B11-1 but not in strains F9-2 and H9-1. Unique *rpo* point mutations found in the acid-evolved strains were reverted to represent the original ancestral W3110 sequence for strain B11-1 (A) as well as strains H9-1 and F9-2 (B). The method described in the legend of Fig. 1B was modified for kinetic recording of absorbance ratios. Absorbance readings were taken every 2 h for 48 h. Replicates for each strain are shown as curves of the same color.

approximately half the level of ancestral W3110 ( $P < 0.05$ ). Thus, overall extreme-acid survival did correspond to the presence of the Gad system although other genetic factors may have affected the degree of acid survival. Because the strains were grown under moderate acid stress for 2,000 generations, mutations may have arisen that decreased extreme-acid survival. This loss of acid survival may represent a trade-off for another phenotype that improves growth.

Several mutations of interest that may have decreased acid survival in B11 were found, including *ftsX* (transporter subunit), *hepA* (RNA polymerase-associated helicase protein), and *rpoC* (RNA polymerase beta prime subunit) (Table S1). These genes function in the cell membrane or in transcription, so the mutations may affect the cell membrane permeability to protons, sensitivity to acid, or expression of acid response genes. H9 mutations that may have affected acid survival occurred in *yej*, inner membrane proteins, and in the intergenic region between *rhaA* and *rhaD*, the region within the L-rhamnose degradation pathway (Table S1).



**FIG 3** GABA production and extreme-acid survival of acid-evolved strains. (A) GABA assays of anaerobic overnight cultures after exposure at pH 2.0 for 2 h, as measured by GC-MS (see Materials and Methods). ANOVA tests identified no significant difference in GABA production levels between strains W3110, B11, and H9. (B) Survival of overnight cultures after dilution and incubation at pH 2.0 for 2 h (see Materials and Methods). The y axis shows the log<sub>2</sub> ratios of the colony counts at the indicated pH values. The axis cutoff ( $10^{-6}$ ) represents the lower limit of colony detection. One-way ANOVA tests were run with the Tukey method comparison to identify significantly different groups. Asterisks indicate strains that show significant difference in results from those of the ancestral strain W3110 ( $P < 0.05$ ).

Previously, we reported a similar loss of Gad during evolution in the presence of benzoic acid buffered at pH 6.5 (32). Some of the benzoate-evolved strains show loss of multidrug resistance. We therefore tested our acid-evolved strains for resistance to chloramphenicol but found no significant difference from the ancestral W3110 (data not shown).

**RNA-seq of acid-evolved strains shows downregulated *gad* genes and upregulated anaerobic catabolism.** Many of the mutations in our acid-evolved strains impact regulatory proteins that could modulate transcription of a large number of products. We therefore used RNA-seq to compare the gene expression across the transcriptomes of four acid-evolved strains (B11-1, F9-2, F11-1, and H9-1), each from one of the four independently evolved populations (Tables S2 to S5). Each transcriptome was indexed to that of the ancestor, after culture at pH 4.8 (see details in Materials and Methods).

The four strains all showed unique patterns of differential transcription, with some features in common. Genes showing the greatest upregulation and downregulation are

**TABLE 2** Differential expression of acid-evolved strains compared to ancestral strain W3110

Regulation of expression	Genes in the indicated strain <sup>a</sup>			
	B11-1	F9-2	F11-1	H9-1
Upregulated $\geq 4$ -fold	<i>citCDEFGX</i> , <i>dcuAB</i> , <i>srIABDE</i> , <i>tdcBCDEFG</i> , <i>tttAB</i> , <i>ykgEFG</i>	<i>norVW</i> , <i>ykgEFG</i>	<b>{<i>glpABC glpD glpFK glpTQ</i>},</b> <i>nirBD</i> , <i>ykgEG</i>	<i>fdnGH</i> , <b>{<i>glpABC, glpTQ</i>},</b> <i>ykgEF</i>
Downregulated $\geq 4$ -fold	<b>{<i>ariR, bhsA</i>}, {<i>argO, lysA, lysC</i>},</b> <i>cysACDHIJNPWU</i> , <b>{<i>gadABCE, mdtEF</i>},</b> <i>nrdHI</i> , <i>sufABC</i> , <i>yhiM</i> , <i>ymdF</i>	<i>dmsABC</i> , <i>fdnG</i> , <i>frdABCD</i> , <i><math>\Delta</math>gadBCEW</i> , <i><math>\Delta</math>hdeABD</i> , <i>hyaABCDEF</i> , <i>hybABO</i> , <i>hypABCD</i> , <i>napDF</i> , <i>narJUYZ</i> , <i>nikABCDE</i> , <i>nrfAC</i> , <i>ybaST</i> , <i>yhiM</i> , <i>ynfEFGKO</i>	<b>{<i>argO, lysA, lysC, lysP</i>},</b> <i><math>\Delta</math>gadABCEWX</i> , <i><math>\Delta</math>hdeABD</i> , <i><math>\Delta</math>mdtEF</i> , <i>sufABCD</i> , <i>ybaST</i> , <i>ycaC</i> , <i>yciFG</i> , <i>yhiM</i> , <i>ymdF</i>	<i>appABC</i> , <b>{<i>argO, lysA, lysC</i>},</b> <b>{<i>gadABCEWX, hdeA, mdtEF, slp</i>},</b> <i>hyaABCDEF</i> , <i>osmEY</i> , <i>otsAB</i> , <i>sufABCDES</i> , <i>ybaST</i> , <i>yciFG</i> , <i>yhiM</i> , <i>ymdF</i>

<sup>a</sup>Selected genes and regulons from Table S2 representing all genes upregulated or downregulated  $\geq 4$ -fold relative to expression levels in W3110 and fitting the criterion of a *P* value of  $<0.001$  as assessed by negative binomial tests in DESeq. RNA was isolated from bacteria cultured to log phase in LBK<sub>mal</sub> broth, pH 4.8. Delta symbols indicate gene deletions (Table S1). Genes within braces (boldface) are grouped together based on functional relevance.

summarized in Table 2. Most of the upregulated genes involve anaerobic catabolism and transport. Strain B11-1 showed upregulation of *citCDEFG* (citrate fermentation), *dcuAB* (anaerobic C<sub>4</sub>-dicarboxylate transport), *srIABDE* (sorbitol permease and catabolism), *tdcBCDEFG* (anaerobic threonine catabolism), and *ykgEFG* (D-lactate catabolism). D-lactate catabolism was upregulated in all four strains. Strain F11-1 also showed upregulation of *glp* operons (anaerobic glycerol catabolism), and strain H9-1 showed upregulation of *glp* as well as *fdnGH* (formate dehydrogenase).

A large number of genes were downregulated in the acid-evolved strains. Most strikingly, the Gad acid resistance genes, as well as multidrug transporters *mdtEF*, were downregulated in B11-1 and in H9-1, our two acid-evolved strains that retained the entire sequence of the Gad acid fitness island. All four strains showed downregulation of *yhiM* encoding a Gad-related acid stress membrane protein (33). Despite downregulation under the low-density culture condition for RNA-seq, stationary-phase cultures of B11-1 and H9-1 produced GABA at levels comparable to the level in the ancestor and showed only a slight decrease of extreme-acid resistance (Fig. 3). We infer that these strains retain Gad activity during stationary phase (the condition that normally shows highest Gad expression, for extreme-acid survival) while downregulating Gad expression during early log phase, where it incurs a fitness cost.

Several operons implicated in the acid stress response were downregulated, including *sufABC* (Fe-S transporter) and *hya* and *hyb* (H<sub>2</sub> consumption hydrogenases Hyd-1 and Hyd-2), all of which are acid-inducible under anaerobic conditions (17). Also downregulated were *cysACD* (sulfate transport) and *ariR* (regulator of acid resistance). Inactivation of these operons is associated with increased cell aggregation, which could contribute to the increased acid fitness in these strains (34, 35).

Strain F9-2, whose genome contains a deletion of the anaerobic regulator *fnr*, showed downregulation of a number of genes normally upregulated by Fnr under low-oxygen conditions (25, 28). The downregulated genes include *dmsABC* (dimethyl sulfoxide reductase), *frdABCD* (fumarate reductase), *hybABO* (Hyd-2), *nikABCDE* (nickel ABC transporter), and *nrfAC* (formate-dependent nitrite reductase). The downregulation of these genes implies that the genes show significant expression in the FNR<sup>+</sup> ancestor W3110 when it is cultured at low pH.

## DISCUSSION

Our analysis of acid-evolved genomes combines genomics, transcriptomics, and pH-modulating enzyme activities to identify the evolutionary selection forces impacting *E. coli* genomes under continual acid stress in semiaerobic microplate culture. The acid stress condition includes nutrient limitation as well as oxygen limitation caused by the physical geometry of the microplate. These conditions are relevant to the mammalian digestive tract, where enteric bacteria experience changes in pH as well as extreme gradients of oxygen across the intestinal epithelium (36).

Surprisingly, the overall picture is that genes highly upregulated by acid incur long-term fitness costs when the acid condition persists across many generations. In



our acid-evolved genomes, different strains showed different combinations of loss or decrease of acid-inducible amino acid decarboxylase activities. Loss of decarboxylase activity occurred via three different mechanisms: IS knockouts of regulatory genes such as *cadC*; IS-mediated large deletions, particularly of the Gad regulon; and regulatory adjustment leading to downregulation of Gad activity at early log phase, such as that seen in the transcriptomes of the Gad-positive strains B11-1 and H9-1. The cause of most of the transcriptome modulation is unknown, but evidence for at least one possible mechanism comes from the modulation of arginine decarboxylase activity by the *rpoC* mutation in strain B11-1; the decarboxylase activity was partly restored in the B11-1 *rpoC*<sup>+</sup> construct (Fig. 2). It is possible for RNAP mutations to offer a way for evolution to quickly fine-tune the expression of a large number of genes (37).

We imagine several possible causes of the loss of acid-induced genes. A comparable result was found in a study of heat-adapted *E. coli* clones (12) that show decreased heat shock gene expression. Genes that confer fitness under a transient extreme stress may actually be detrimental during prolonged exposure; thus, evolution selects for restoration of the nonstress physiology. Another consideration is that “acid stress” actually encompasses a wide range of concentrations of the hydronium ion and that systems conferring fitness at one pH may be less advantageous at a different acid level. Thus, the lysine decarboxylase might be advantageous for cells cultured at pH 6 but unhelpful at pH 4.6. Finally, the high concentration of buffer at low pH negates the advantage of decarboxylases that function to neutralize acid. Under nutrient limitation, without the advantage of neutralizing pH, the catabolic loss of amino acids would be strongly selected against.

The transcriptomes of our acid-evolved clones showed varied patterns of upregulation and downregulation of catabolic pathways. One common result for all four strains (Table 2) is the upregulation of *ykgEFG*, implying an important energy contribution from D-lactate catabolism. The fitness advantage of diverse catabolism implies that in our evolution experiment, carbon source utilization under limited oxygen may have had as great a selective force as pH. The cyclic transition through stationary phase and nutrient limitation may be an underappreciated contributor to fitness in classic evolution experiments where “daily dilution” always includes several hours of stationary phase.

The *fnr* deletion in strain F9-2 represents a divergent alternative path of evolution compared to the paths of our other three populations analyzed. It is unclear how the loss of anaerobic induction helps survival under our conditions of semiaerobic culture at low pH. One possibility is that turning off the consumption hydrogenases (*hya* and *hyb*) eliminates production of H<sup>+</sup> ions that could increase acidification (17, 38, 39). Alternatively, some of the FNR-activated gene expression may have represented a wasteful energy expenditure, i.e., over 2,000 generations of evolution without extreme anaerobiosis. Thus, the loss of anaerobic response under a constant semiaerobic condition may be comparable to the loss of an extreme-acid response under a buffered acid condition.

It would be of interest to further establish the link between loss of decarboxylase activity and increased acid fitness. This could be accomplished with gene complementation although the construct would require testing to rule out construct-specific fitness effects. Alternatively, the mutated decarboxylase genes could be reverted to the wild-type sequence, as we have done with the *rpo* mutations in the acid-evolved strains. We would expect to see a decrease in moderate acid fitness when decarboxylase activity is restored.

Our findings on acid-influenced evolution of bacteria have implications for future studies of the human gut microbiome, in which ingested bacteria pass through the stomach where they experience an environment at pH 4 to 5 before entering the intestine. Interestingly, IS-inactivated *cadC* alleles are found in enteroinvasive strains of *E. coli* and *Shigella* (40). CadA inactivation is thought to occur in pathogens because cadaverine production lowers virulence. Another relevant observation is that acid-evolved bacteria show altered production of polyamines and GABA, which is of interest in the microbiome for carcinogenesis (6) and the gut-brain axis (7).

**TABLE 3** Strains used in this study

Strain name	Population isolate	Genotype	Source or reference
JLSE0079	H9-1	See Table S1	13
JLSE0080	H9-2	See Table S1	13
JLSE0083	B11-1	See Table S1	13
JLSE0084	B11-2	See Table S1	13
JLSE0091	F11-1	See Table S1	13
JLSE0092	F11-2	See Table S1	13
JLSE0137	F9-2	See Table S1	13
JLSE0138	F9-3	See Table S1	13
W3110	Stock D8		Fred Neidhardt
JW5731-1		$\Delta$ adiA743::kan	45
JW4094-5		$\Delta$ cadC760::kan	45
JW3953-2		$\Delta$ thiH760::kan	45
JW5549-1		$\Delta$ thiG761::kan	45
JLSE0079thiH2C		JLSE0079(H9-1) $\Delta$ thiH760::kan <i>rpoB</i> <sup>+</sup>	This study
JLSE0079thiH2C::frt		JLSE0079(H9-1) $\Delta$ thiH760::frt <i>rpoB</i> <sup>+</sup>	This study
JLSE0083thiG1A		JLSE0083(B11-1) $\Delta$ thiG761::kan <i>rpoC</i> <sup>+</sup>	13
JLSE0083thiG1A::frt		JLSE0083(B11-1) $\Delta$ thiG761::frt <i>rpoC</i> <sup>+</sup>	This study
JLSE0137thiH1C		JLSE0137(F9-2) $\Delta$ thiH760::kan <i>rpoC</i> <sup>+</sup>	This study
JLSE0137thiH1C::frt		JLSE0137(F9-2) $\Delta$ thiH1C760::frt <i>rpoC</i> <sup>+</sup>	This study

## METHODS AND MATERIALS

**Bacterial strains and medium.** *Escherichia coli* K-12 W3110 was the ancestral strain of all acid-evolved populations (13). All strains were cultured in Luria Bertani-potassium (LBK) broth, containing 10 g/liter tryptone, 5 g/liter yeast extract, and 7.45 g/liter KCl. The culture medium was buffered with either 100 mM homopiperazine-1,4-bis(2-ethanesulfonic acid) (HOMOPIPES) ( $pK_a = 4.55, 8.12$ ), 100 mM 2-(*N*-morpholino)ethanesulfonic acid (MES) ( $pK_a = 5.96$ ), or 100 mM 3-morpholinopropane-1-sulfonic acid (MOPS) ( $pK_a = 7.20$ ), depending on the desired pH of 4.4 to 4.8, 5.5 to 6.5, or 7.0, respectively. The pH of the medium was adjusted as necessary with either 5 M HCl or 5 M KOH. LBK medium buffered to the pH range of 4.5 to 4.8 was additionally supplemented with 10 g/liter DL-malic acid ( $pK_a = 3.40, 5.11$ ) (LBK<sub>mal</sub>) as described by Harden et al. (13). As both D- and L-malate may be catabolized by acid-consuming reactions (41, 42), supplementation with malic acid provided a potential resource to enhance acid tolerance.

**Sequence analysis.** The sequence annotations of acid-evolved genes (13) were updated using breseq, version 0.27.1 (18, 19). This breseq version reveals IS-mediated insertions and deletions that are identified by inspection of unassigned new junctions and regions of missing coverage (43). Sequences of acid-evolved strains were compared to the *E. coli* K-12 W3110 reference sequence (NCBI GenBank accession number NC\_007779.1) (22). Variant sequences present in our laboratory strain of W3110, referred to as W3110-D8, were filtered. Assembly and annotations of our evolved isolate sequences were visualized using the Integrative Genomics Viewer (IGV) from the Broad Institute. Sequence identity was confirmed by PCR amplification of selected mutations.

**P1 phage transduction and strain construction.** Mutations were reverted to the ancestral K-12 W3110 sequence by cotransduction of linked markers, using P1 phage by standard procedures (44). Strains with desired deletions or functional insertions were ordered from the Keio collection (Coli Genetic Stock Center, Yale University) (45) and were introduced into the evolved strain of choice or the ancestral strain W3110. Constructs were confirmed by PCR amplification and Sanger sequencing of key alleles of donor and recipient.

**FRT procedure.** Antibiotic resistance was eliminated from strains containing a Keio *kan* insertion allele by a flippase-FRT recombinase procedure as described in Datsenko and Wanner (46) using plasmid pCP20. The designation “frt” (flippase recognition target) is given to strains with the Keio insertion removed and indicates the remaining scar sequence (strains are listed in Table 3).

**Lysine and arginine decarboxylase assays and pH rise.** The pH rise associated with activity of lysine decarboxylase and arginine decarboxylase was measured using a procedure with Møller decarboxylase broth, with modification (47). Lysine decarboxylase broth contained 3 g/liter yeast extract, 1 g/liter D-glucose, 5 g/liter L-lysine, and 0.015 g/liter bromocresol purple, pH  $6.8 \pm 0.2$ . L-lysine was replaced with L-arginine in the arginine decarboxylase broth at pH  $5.5 \pm 0.2$ . Strains were inoculated into a microplate containing aliquoted medium. The plate was sealed for anaerobiosis and incubated at 37°C for 24 h. The plate was inspected visually and verified by absorbance readings from 350 nm to 750 nm using a SpectraMax Plus 384 microplate reader (Molecular Devices). We established an absorbance ratio,  $R = A_{490}/A_{570}$ , that reflects the pH shift as indicated by titration of the dye bromocresol purple. The ratio  $R$  was calculated from mean absorbance values,  $A_{570-590}/A_{400-450}$ . The pH conversion scale is shown in Fig. S1 in the supplemental material.

**GABA assays.** The procedure for measuring GABA production via glutamate decarboxylase was modified from that of Creamer et al. and Ma et al. (32, 48). Strains were cultured overnight in LB medium (10 g/liter tryptone, 5 g/liter yeast extract, 100 mM NaCl, 10 mM glutamine) buffered with 100 mM MES,

pH 5.5. Glutamine is converted to glutamate, the substrate of glutamate decarboxylase (49). Closed 9-ml screw-cap tubes nearly full of medium were incubated for 18 h at 37°C. The pH of each sample was lowered with HCl to pH 2.0 and incubated for 2 h with rotation. The supernatant from 1 ml of culture was filtered and prepared for GC-MS by EZfast amino acid derivatization (50). GABA from the culture fluid was identified using National Institute of Standards and Technology (NIST) library analysis.

**Extreme-acid survival assays.** The conditions for testing extreme-acid survival were based on the aerobic assay from Riggins et al. and Lin et al. (38, 51). Cultures were grown overnight (16 to 18 h) in LBK broth, buffered with 100 mM MES at pH 5.5 to upregulate acid response systems (4). Overnight cultures were diluted 1:200 in LBK broth at pH 2.0 and incubated at 37°C for 2 h. After exposure, cultures were serially diluted in M63 minimal medium [0.4 g/liter  $\text{KH}_2\text{PO}_4$ , 0.4 g/liter  $\text{K}_2\text{HPO}_4$ , 1 g/liter  $(\text{NH}_4)_2\text{SO}_4$ , 7.45 g/liter KCl] buffered with 100 mM MOPS at pH 7.0 to a final dilution of 1:400,000. Control cultures at pH 5.5 were serially diluted to a final ratio of 1:400,000 in M63 buffered with 100 mM MOPS, pH 7.0. Fifty microliters of each final dilution was spread onto LBK agar plates and incubated at 37°C. Colony counts for each replicate were log transformed. The means of  $\log_2$  values from each replicate are reported.

**RNA purification and extraction.** Bacteria were cultured to stationary phase (16 to 18 h) in LBK<sub>mal</sub> broth (pH 4.8), diluted 1:50 into fresh medium at pH 4.8, and grown to early log phase as indicated by an optical density at 600 nm ( $\text{OD}_{600}$ ) of 0.2. Cultures were diluted 6:1 by volume into a 5% buffer-saturated phenol-ethanol solution and centrifuged (4 min at 5,000 rpm and 4°C). The supernatant was decanted, and centrifugation was repeated. The pellet was resuspended in Tris-EDTA (TE) buffer (100  $\mu\text{l}$ ) with 3 mg/ml lysozyme. To isolate RNA, a Qiagen RNeasy minikit was used according to the manufacturer's instructions, with the addition of one RNase-free DNase treatment for 15 min at room temperature. A second DNase treatment using MoBio DNase-Max was conducted according to the manufacturer's instructions.

**Construction and sequencing of Illumina RNA-seq libraries.** The mRNA population from all RNA samples was enriched by depleting rRNA sequences, and strand-specific libraries were constructed as described earlier (52). Starting with 8 to 50 ng of rRNA-depleted RNA, random-primed cDNA synthesis was done using a ScriptSeq, version 2, RNA-seq library preparation kit (Epicenter, WI). cDNA was purified using an Agencourt AMPure XP system (Beckman Coulter, NJ). Libraries were amplified using a FailSafe PCR enzyme kit (Epicenter, WI). Typically 12 (50 to 100 ng of starting RNA) or 15 (8 ng of starting RNA) PCR cycles were used, and reverse primer from the kit was replaced with one of the ScriptSeq Index primers. After PCR amplification, libraries were purified and size selected (~280 bp) using the Agencourt AMPure XP system (Beckman Coulter, NJ). Profiles of library insert sizes were verified on an Agilent Technology microfluidic platform using a DNA 1K chip (Agilent Technologies, Wilmington, DE). Libraries were sequenced on an Illumina NextSeq 500 (150-cycle paired-end mode).

**Mapping and assembly of raw reads.** Sequences were initially analyzed using CLC Genomics software, version 6.0. Sequences with a quality score of less than 30 were discarded, the remaining sequences were trimmed, and sequences of less than 36 bp were discarded. Sequences were mapped to the *E. coli* W3110 genome, (NCBI accession number NC\_007779.1) (22), and mapping was done using the following CLC genomics mapping parameters: mismatch, 1; insertion, 3; deletion, 3; length, 0.9; similarity, 0.95; auto-detect paired distances on and map randomly. The CLC RNA-seq was performed using the following parameters: mismatch, 2; length fraction, 0.9; similarity fraction, 0.95; strand specific selected; maximum 3 hits, 3; paired settings, 36 to 500; broken pairs counting selected. Only unique counts generated for individual genes were used as the starting data for all subsequent analyses.

**RNA-seq analysis.** Differential expression analysis was performed using the R package DESeq (53). A multifactor design was used to control for batch effects. For each comparison, two models were fit to generalized linear models (GLM), one regressing the gene expression on both batch and strain and one regressing only on the batch. *P* values were determined by a chi-square test that compared the two GLM fits. Samples were normalized by size factor, and dispersions were estimated as described previously (53). Comparisons resulting in a *P* value of <0.001 with a  $\log_2$  fold change of  $\geq 2$  or  $\leq -2$  were considered significant. Quality assessment was conducted using heat maps with variance-stabilized data and principal component analyses.

**Statistical analysis.** Analysis of variance (ANOVA) with *post hoc* Tukey honestly significant differences (HSD) tests were used for between-strain statistical comparisons of Lys- and Arg-dependent pH rise, extreme-acid survival, and GABA production. The terminal spectrum ratios of bromocresol purple absorbance ( $A_{570-590}/A_{400-450}$ ) for each strain were used for comparisons of Lys- and Arg-dependent pH rise. The time required to reach a spectrum ratio of 0.7 after 4 h was used for statistical comparisons in the kinetic decarboxylase assays. The  $\log_2$  colony count ratios pre- and post-acid exposure were the values used to compare extreme-acid survival. Extracellular GABA levels (in millimolar amounts), as measured by GC-MS, were compared to assess differences in GABA production between strains.

**Accession number(s).** Genomic sequence and alignment data have been deposited in the NCBI Sequence Read Archive (SRA) under accession number SRP041420. RNA-seq data have been deposited in the NCBI SRA under accession number SRP095404 (see Table S6 for sample descriptions).

## SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/AEM.00442-17>.

**SUPPLEMENTAL FILE 1**, XLSX file, 2.1 MB.

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