Pathogenic bacteria sense and respond to a plethora of signals as they traverse through the host. The integration of these signals into appropriate changes in behavior is an important determinant of a pathogen’s success. The intestine is a particularly complex and dynamic environment, containing signals from the host, the microbiome, and large molecular perturbations from diet. Competition within this environment extends beyond simply colonization of the host, the microbiome, and large molecular perturbations from diet. The molecular environment of the host can have profound effects on the behavior of resident bacterial species. We recently established that the sensing and response of enterohemorrhagic Escherichia coli (EHEC) to D-serine (D-Ser) resulted in down-regulation of type 3 secretion system-dependent colonization, thereby avoiding unfavorable environments abundant in this toxic metabolite. However, this model ignores a key determinant of the success of bacterial pathogens, adaptive evolution. In this study, we have explored the adaptation of EHEC to D-Ser and its consequences for pathogenesis. We rapidly isolated mutated, independent, EHEC mutants whose growth was no longer compromised in the presence of D-Ser. Through a combination of whole-genome sequencing and transcriptomics, we showed that tolerance could be attributed to disruption of one of two D-Ser transporters and/or activation of a previously nonfunctional D-Ser deaminase. While the implication of cytoplasmic transport in D-Ser toxicity was unsurprising, disruption of a single transporter, CycA, was sufficient to completely overcome the repression of type 3 secretion system activity normally associated with exposure to D-Ser. Despite the fact that this reveals a mechanism by which evolution could drive a pathogen to colonize new niches, interrogation of sequenced E. coli O157:H7 genomes showed a high level of CycA conservation, highlighting a strong selective pressure for functionality. Collectively, these data show that CycA is a critically important conduit for D-Ser uptake that is central to the niche restriction of EHEC.

D-amino acids | EHEC | adaptive evolution | metabolism | virulence

Significance

Pathogens ensure infection of favored sites in the body by responding to chemical signals. One chemical abundant in urine, the amino acid D-Ser, is toxic to EHEC and reduces expression of the machinery used for host cell attachment, making the bladder an unfavorable environment. We observed that under D-Ser stress, EHEC acquires genetic changes that lead to blocking D-Ser uptake into the cell or activating a silent enzyme for degrading D-Ser. This prevents growth inhibition and, critically, inhibits the repression of attachment machinery normally caused by D-Ser. These findings highlight the importance of pathogen evolution in determining how host molecules regulate colonization. These interactions underpin a process known as niche restriction that is important for pathogen success within the host.

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The authors declare no competing interest.

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environments with high levels of D-Ser. A number of inner-membrane transporters have been implicated in specific uptake of D-Ser, including DsdX (18), which is encoded in the dsdCX4 locus, CycA (19), and SstT (20).

We recently described how EHEC can sense D-Ser (21) and respond by reducing expression of its primary colonization apparatus, the LEE-encoded type 3 secretion system (10). This allows for restriction of EHEC to a more favorable niche within the gut where D-Ser concentrations are low (10), compared with environments such as the bladder, where D-Ser levels are higher (11). In addition to virulence repression by D-Ser, an SOS-like response is activated, leading to a higher rate of genomic variability (22), likely reflecting the bacterium attempting to overcome the stress induced by D-Ser exposure. Together with the observation that LEE-positive enteric E. coli have evolved to lose the ability to catabolize D-Ser (10), these findings corroborate the role of D-Ser in niche restriction of LEE-positive E. coli.

Here we expand on these studies by investigating if EHEC can adapt to overcome this evolutionary bottleneck. We describe the genetic and phenotypic consequences of repeated in vitro exposure of EHEC to D-Ser. Tolerance to D-Ser convergently evolved by distinct mechanisms and resulted in an ability to overcome not only the growth-inhibitory effects of D-Ser but also the majority of transcriptional alterations associated with D-Ser, including type 3 secretion system-dependent colonization. We discuss these mechanistic insights and the implications that they have for the pathogenic lifestyle of EHEC.

Results

D-Ser Exposure Can Induce the Evolution of Tolerant EHEC Isolates. The prototype UPEC and NMEC strains CFT073 and CE10 were chosen for inclusion in this study as, like 85% of UPEC (11) and 97.5% of NMEC (23), they possess complete, responsive, and functional dsd loci (Fig. 1A). The NMEC CE10 isolate is interesting as it carries two copies of the locus. EHEC TUV93 offset (a Shiga toxin-negative derivative of EDL933), on the other hand, is typical of the majority of diarrheagenic LEE-pathogenicity island-carrying E. coli in that it possesses a genomic rearrangement upstream of dsdX leading to loss of dsdCX and hence loss of function (10, 23). EHEC cannot grow on D-Ser as a sole carbon source, and its growth in glucose-containing minimal medium is strongly inhibited by D-Ser (Fig. 1B and C). The specific growth rate (SGR) for EHEC was found to decrease from 0.63 h⁻¹, in agreement with previous reports for M9 + glucose (24), to 0.19 h⁻¹ (P = 0.003) upon inclusion of D-Ser, whereas no significant decrease was observed with UPEC or NMEC (SI Appendix, Fig. 1S). This indicated that D-Ser exerted a sublethal growth-inhibitory effect on EHEC and alluded to the possibility that tolerant mutants could arise by repeated exposure as the bacteria attempt to overcome the inhibition of growth. To test this, we repeatedly batch-cultured EHEC in the presence of D-Ser for a period of 10 d as outlined in Fig. 2A. When subculturing on each day, a sample was serially diluted and spot-plated on solid M9 medium containing glucose and D-Ser (i.e., the same selective conditions as the flask culture). Importantly, this allowed for discrimination of putatively tolerant clones by virtue of their ability to form large colonies. The resulting clones were termed large-colony variants (LCVs) and numbers were assigned from the passage number at which they were isolated (for example, LCV3 was isolated from the third 24-h batch culture). LCV1A and LCV1B represented a second colony selected by distinct mechanisms and resulted in an ability to overcome not only the growth-inhibitory effects of D-Ser but also the majority of transcriptional alterations associated with D-Ser, including type 3 secretion system-dependent colonization. We discuss these mechanistic insights and the implications that they have for the pathogenic lifestyle of EHEC.

The Evolution of Tolerance Alters the Transcriptional Response to D-Ser. Before characterizing the molecular basis of D-Ser tolerance, it was crucial to establish whether this tolerance phenotype had any implications for the distinct transcriptional responses that we have previously associated with exposure of EHEC to D-Ser, namely repression of type 3 secretion-mediated colonization (10) and activation of an SOS-like response via RecA induction (10, 22). We used transcriptional reporters of the LEE master regulator (Fig. 3A) and recA (Fig. 3B). As LCV1 and LCV3 did not exhibit tolerance, one would expect them to behave as the WT with respect to both LEE1 repression and recA induction. This was indeed the case, further supporting the notion that these are false-positive isolates for D-Ser tolerance. The majority of D-Ser–tolerant isolates showed no repression of LEE1 in response to D-Ser (LCV4 to 10) and no significant activation of recA transcription. This group consists of both partially tolerant and completely tolerant isolates (Fig. 2C and D). Two isolates exhibited recovery from recA activation but maintained WT levels of type 3 repression (LCV2 and LCV10B; Fig. 3A and B).

Next, a single representative from these three groupings (no recA induction, complete tolerance: LCV2; no recA induction or type 3 repression with partial tolerance: LCV7; no recA induction or type 3 repression with complete tolerance: LCV10) was tested for infection of host epithelial cells in the presence or absence of D-Ser (Fig. 3C–E). Adherent bacteria formed characteristic A/E lesions with regions of pronounced actin condensation being visible adjacent to the majority of bacterial cells (Fig. 3C). Only WT EHEC showed a significant reduction in adhesion levels in terms of the numbers of bacteria attached per cell (Fig. 3D) and the percentage of cells infected (Fig. 3E). Surprisingly, LCV2 exhibited reduced LEE1 promoter activity in the presence of D-Ser while displaying higher levels of colonization (Fig. 3D); however, we will later discuss further a mechanism by which LCV2 and LCV10B may gain an advantage in the presence of D-Ser. Taken together, these findings illustrate that tolerance starkly affects the transcriptional response to D-Ser and results in overcoming the inhibitory effects of D-Ser on colonization, a factor that could have important implications for niche specificity of such mutants.

The Molecular Basis of D-Ser Tolerance. Having observed variable levels of tolerance to D-Ser and distinct patterns of recovery from the transcriptional shift normally associated with D-Ser, we predicted that three distinct mechanisms of tolerance were at play. First, the genomes of each isolate were sequenced to 30-fold coverage and aligned to a resequenced genome of the EHEC parental strain from this study. Single-nucleotide polymorphisms (SNPs) were called where present in 90% of reads. Of the nonsynonymous SNPs present in tolerant isolates, only those in cycA appeared to be strongly selected for, being present in both experimental replicates (i.e., LCV4 to 10 [premature stop at position K280] and LCV1A [I257K]; Table 1), appearing early (batch cultures 4 and 1, respectively), and being maintained in the 10-d experiment until the final day of sampling. CycA is a previously characterized D-Ser/cycloserine/D-alanine/glycine/l-alanine transporter (18, 19). While the cycA mutation alone could putatively be responsible for the tolerance phenotype of

wild-type [WT] D-Ser but lower than WT, 0.31 to 0.38 h⁻¹; Fig. 2C and SI Appendix, Fig. S2). The remaining five clones (LCV2, 8 to 10, and 10B) displayed complete tolerance to D-Ser with growth rates similar to that of WT EHEC cultured in the absence of D-Ser (Fig. 2C and SI Appendix, Fig. S2). As such, D-Ser–tolerant mutants can be isolated and positively selected for by repeated culture in the presence of toxic D-Ser. Strikingly, the observation of two distinct levels of tolerance alluded to at least two independent mechanisms of tolerance.
the majority of the tolerant mutants (LCV4 to 10 and 1A), two questions remained: Why were some cycA mutants more tolerant than others, and what was the mechanism of tolerance for LCV2 and LCV10B that appear to encode WT cycA?

To address the first question, we examined the transcriptome of a representative from the completely tolerant group that possessed a mutation in cycA—LCV10—and compared it with the WT. We hypothesized that a transcriptional alteration in this strain might explain its enhanced tolerance over the partially tolerant group (LCV4 to 7 and 1A). Wild-type EHEC was found to significantly differentially express 369 genes in response to D-Ser (136 up and 233 down; SI Appendix, Table S1), consistent with its inability to tolerate D-Ser due to an insertion that truncates the D-Ser catabolism locus.

Fig. 1. EHEC is unable to tolerate D-Ser due to an insertion that truncates the D-Ser catabolism locus. (A) Schematic depicting the D-Ser catabolism locus in EHEC, UPEC, and NMEC. (B) Growth curves of EHEC, UPEC, and NMEC in M9 + glucose (Glc) ± 1 mM D-Ser. Error bars indicate SEM from a minimum of three experiments. (C) Schematic illustrating the contrasting effects of D-Ser on dsd− and dsd+ strains of E. coli. While D-Ser catabolism promotes the growth of UPEC and NMEC, growth of EHEC is severely restricted by D-Ser. In the case of EHEC, D-Ser is transported into the cytosol by CycA and SstT, where it accumulates and induces an SOS-like response and transcriptional repression of type 3 secretion.

Fig. 2. D-Ser–tolerant EHEC mutants evolve and can be selected for by repeated culture in the presence of D-Ser. (A) Experimental workflow of repeated batch culture for the enrichment of D-Ser–tolerant mutants. (B–D) Growth curves (M9 + glucose + 1 mM D-Ser) of apparent LCV clones obtained after the indicated number of repeat cultures in D-Ser–containing medium. Error bars indicate SEM from a minimum of three experiments.
with our previous reports (10). The majority (247/369) of these differentially expressed genes (DEGs) were oppositely regulated when comparing the transcriptome of WTD-Ser and LCV10D-Ser (SI Appendix, Table S1), highlighting the fact that tolerance can overcome many of the transcriptional effects exerted by D-Ser. A further 361 genes (new DEGs; SI Appendix, Table S2) were differentially expressed in LCV10D-Ser versus WTD-Ser that were not associated with the response to D-Ser (i.e., WT versus WTD-Ser), indicating that the transcriptional alterations in LCV10 extended beyond simply overcoming the normal effects of D-Ser. The most significant of these was sstT (also known as yjgU) with 87.68-fold lower expression (false discovery rate adjusted \( P = 1.86 \times 10^{-107} \) in LCV10 than the WT (Fig. 4A). Reduced transcription of sstT was confirmed by quantitative real-time PCR (LCV10, −147.74-fold; Fig. 4B) and was also observed in a second completely tolerant isolate (LCV9, −92.15-fold), while two partially tolerant isolates (LCV5 and LCV7) and one tolerant isolate that lacked a mutation in cycA (LCV2) showed less than a twofold decrease in transcription. Decreased expression of sstT was found to be constitutive with no significant difference being observed with or without D-Ser (Fig. 4C). SstT functions as a sodium/serine and sodium/threonine symporter (20, 25), and hence transcriptional knockdown had a strong likelihood of facilitating tolerance synergistically with disruption of cycA.

**Disruption of D-Ser Transport as a Means of Conferring Tolerance.**

We hypothesized that the mutations acquired in CycA (K280Stop, I257K) and the marked reduction in sstT transcription led to lack of function or expression at a level that would not allow for functionality, thereby facilitating the observed phenotypes in LCV4 to 10 and 1A. To confirm these mechanisms, we constructed isogenic mutants lacking either CycA, SstT, or both transporters and assayed for tolerance to D-Ser. Deletion of cycA led to enhanced growth in the presence of D-Ser (Fig. 5A), with SGR increasing from 0.15 to 0.35 h\(^{-1}\) (SI Appendix, Fig. S3). While deletion of sstT caused only a modest increase in tolerance (Fig. 5B), the increase in SGR over WTD-Ser was not significant (\( P = 0.122; \) SI Appendix, Fig. S3). Combined disruption of cycA and sstT resulted in complete tolerance to D-Ser (Fig. 5C) with a SGR of 0.61 h\(^{-1}\), statistically equivalent to that of WT\(^{+}\)-Ser (SI Appendix, Fig. S3). Therefore, disruption...
of cyaA alone recapitulates the phenotypes of LCV4 to 7 and 1A, while secondary disruption of sstT in this genetic background recapitulates the phenotype of LCV8 to 10. Sensitivity to D-Ser was restored in all three deletion mutants by overexpression of either transporter on the multicopy plasmid pACYC184 (Fig. 5D–F). Interestingly, overexpression of CyaA led to a hypersensitive phenotype with D-Ser inducing a killing effect, as revealed by negative SGR values being observed in all three mutant backgrounds (Fig. 5D–F and SI Appendix, Fig. S3).

We next tested the ΔcyaA and ΔsstT mutants for transcriptional response to D-Ser using a dual LEE1-gfp and recA-rfp reporter system. Similar to LCV4 to 10 and 1A, mutants lacking cyaA no longer showed repression of LEE1 (Fig. 5G) or activation of recA (Fig. 5H) after 4 h of growth. A 4.97-fold higher level of recA expression was seen with WTD-Ser versus WT (P = 0.003) at 5 h, while D-Ser caused only a 1.07-fold induction of recA (P = 0.006) in the ΔcyaA background. Transport of D-Ser via CyaA appeared to be the predominant factor in determining transcriptional response as deletion of sstT provided no relief from D-Ser–induced type 3 repression (SI Appendix, Fig. S4A) or recA activation (SI Appendix, Fig. S4B). As seen with the corresponding LCV isolate (LCV10; Fig. 3C–E), recovery from repression of LEE1 (Fig. 5G) by disruption of D-Ser transport correlated with increased efficiency of colonization in a HeLa cell infection model (Fig. 5I–K). Unlike WT EHEC, D-Ser no longer caused a reduction in visible A/E lesions (Fig. 5I), bacteria per cell (Fig. 5J), or percentage of cells infected (Fig. 5K). The data obtained from these isogenic transporter deletion mutants corroborate the results obtained with LCV4 to 10 and 1A and further support the idea that disruption of CyaA and SstT can result in tolerance to D-Ser with implications for stress response and virulence of EHEC.

### A Transport-Independent Mechanism of Tolerance?

The data presented thus far suggest that disruption to serine transport systems...

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**Table 1. Mutations detected in D-serine–adapted EHEC isolates**

| Gene/region mutated | Mutation | Alteration in amino acid sequence† | Annotation | Isolate(s) possessing mutation |
|---------------------|----------|-----------------------------------|------------|--------------------------------|
| cyaA                | MNP      | pI257K                            | Transporter| 1A                             |
| tamB                | SNP      | sI358R                            | Membrane assembly | 1A |
| ygFL                | SNP      | kI384V                            | Enzyme     | 3                              |
| cyaA                | SNP      | kI280R                            | Transporter| 4 to 10                        |
| z1097               | INS      | NKNK*I                            | Hypothetical| 9                              |
| z2148               | SNP      | R52G                              | Phage tail | 10 and 10B                     |
| z1364               | SNP      | None                              | Phage portal| 1 and 2                       |
| z1364               | SNP      | None                              | Phage portal| 2                             |
| yqhD                | SNP      | None                              | Enzyme     | 3                              |
| z2148               | SNP      | None                              | Phage tail | 10B                            |
| z2148               | SNP      | None                              | Phage tail | 10B                            |

†Stop codon (*).

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**Fig. 4.** Constitutive knockdown in the serine transporter SstT occurs in isolates that possess a premature stop codon in cyaA and display complete D-Ser tolerance. (A) Representative RNA-sequencing (RNA-seq) tracks of WT and LCV10 cultured for 5 h in MEM-HEPES + 1 mM D-Ser together with a schematic illustrating the gene organization within this locus. (B and C) Quantitative real-time PCR of sstT transcription following 5 h of growth in M9 + Glc ± 1 mM D-Ser. Triplicate experimental values for each isolate were compared with the WT with *P < 0.05, **P < 0.01, and ***P < 0.001 by unpaired two-tailed Student’s t test. ns, not significant.
can contribute to d-Ser tolerance; however, two tolerant isolates remained for which no transport-related SNPs were identified (LCV2 and LCV10B; Table 1). To investigate the underlying tolerance mechanism, we employed transcriptome profiling. Comparing WT+d-Ser and LCV2d-Ser revealed 660 DEGs, 258 of which comprised genes differentially expressed upon WT treatment with d-Ser (SI Appendix, Table S3). Of these, 233 were oppositely regulated, illustrating that, similar to LCV10, LCV2 had overcome the majority of transcriptional alterations exerted on WT EHEC by d-Ser. Comparing WT+d-Ser and LCV2+d-Ser also revealed 401 new DEGs not included in the WT/WT+d-Ser comparison, indicating that the transcriptional alterations in LCV2 extended beyond simply...
overcoming the effects of d-Ser. Among the most significant of these alterations were decreased expression of cscR (z3626 (−147.74-fold, P = 6.50 × 10−10)) and up-regulation of cscA (z3625 (58.29-fold, P = 2.66 × 10−12), cscK (z3624 (155.74-fold, P = 2.89 × 10−20), cscB (z3623 (28.24-fold, P = 2.58 × 10−11), and dsdA (33.05-fold, P = 1.34 × 10−10)).

The first of these genes (cscR) was particularly interesting as no reads were obtained for LCV2-Ser (Fig. 6A), accounting for the apparent large degree of differential expression. We hypothesized that a large genomic rearrangement may have occurred within this region that was not detected by our SNP-calling pipeline. Manual analysis of genome sequencing from this region in LCV2 and LCV10B showed a striking lack of coverage, suggesting that genomic reorganization had led to a loss of genomic content (SI Appendix, Fig. S5A). Primers flanking the putative deletion site of LCV10B (annealing sites by red lines in Fig. 6A) were used to screen all LCVs for similar deletions. Only LCV2 and LCV10B were found to have truncations in this region (SI Appendix, Fig. S5B). Sanger sequencing of the PCR amplicons from LCV2 and LCV10B confirmed the precise coordinates of the deletions. LCV2 possessed a 1,120-bp deletion spanning from 149 bp before the 3′ end of cscA to 8 bp downstream of the 5′ end of cscR (Fig. 6A) with a 2-bp direct repeat of CG flanking the deletion site (red dashed boxes in Fig. 6A). LCV10B possessed a 5,637-bp deletion spanning from 32 bp before the 3′ end of z3622 to 122 bp downstream of the 5′ end of cscR (Fig. 6A) with an 8-bp direct repeat of CCGGATAA flanking the deletion site (blue dashed boxes in Fig. 6A). CscR represses the transcription of the sucrose utilization locus encoded by cscBKA (17). As the deletions in LCV2 and LCV10B resulted in loss of the 3′ end of cscA and the entire csc locus, respectively, these isolates lost their ability to grow on sucrose as a sole carbon source (SI Appendix, Fig. S5C).

As mentioned previously, loss of transcription of cscR in LCV2 coincided with increased transcription of not only cscBKA via derepression but also increased transcription of the d-Ser deaminase-encoding gene dsdA (Fig. 6A and SI Appendix, Table S3). This indicated that LCV2 and LCV10B may have gained an ability to catabolize d-Ser. Both LCV2-Ser and LCV10B-Ser exhibited enhanced growth (SGR, 0.53 and 0.53 h−1) compared with culture in minimal essential media-4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (MEM-HEPES) without d-Ser (0.40 and 0.41 h−1), whereas WT EHEC (0.41 h−1) displayed reduced growth upon inclusion of d-Ser (0.37 h−1; Fig. 6A). Both LCV2-Ser and LCV10B-Ser also achieved a higher maximal OD600 nm than WT-Ser, while no difference was observed when d-Ser was absent (Fig. 6B and C and SI Appendix, Fig. S6B), indicating that d-Ser contributed to increased biomass accumulation in LCV2 and 10B. Quantitation of d-Ser concentrations in this medium confirmed that enhanced growth of these strains correlated with the consumption of d-Ser with significantly lower levels of d-Ser being present in cell-free supernatants from LCV2 and LCV10B at 4, 5, and 6 h compared with WT EHEC (Fig. 6D). An isogenic cscR deletion mutant recapitulated the tolerance phenotype of LCV2 and LCV10B in M9 medium (Fig. 6E) and this phenotype was successfully complemented by reintroduction of cscR in trans (Fig. 6F and SI Appendix, Fig. S7). The ΔcscR mutant also displayed enhanced growth rate and maximal OD600 nm in MEM-HEPES + d-Ser in support of catabolism and assimilation into biomass (Fig. 6G).

Reactivation of dsdA via Tolerance-Mediated Adaptive Mutation. There were two plausible explanations for the activation of the previously nonfunctional d-Ser deaminase gene dsdA by upstream genomic deletions. One possibility was that deletion of the transcriptional repressor CscR led to direct derepression of dsdA transcription via its own promoter, dsdAp. The second explanation was that the alteration in genomic context of dsdA could lead to control via read-through from alternative promoters such as cscAp in the case of LCV2 or z3622p in the case of LCV10B.

To explore the first possibility, we examined the effect of heterologous CscR expression in UPEC. UPEC lacks the csc sucrose utilization locus and, as previously mentioned, possesses a functional dsd system for catabolism of d-Ser. The promoter element of dsdA is located 115 bp upstream of the start codon; 114 bp of this sequence are conserved between EHEC and UPEC and, as such, it could be expected that any putative CscR binding in this region would also be conserved. Expression of CscR in UPEC did not result in a decrease in tolerance to d-Ser in M9 minimal medium (SI Appendix, Fig. S8A), as carriage of empty pACYC184 and pscC resulted in similar rates of growth. Qualitative real-time PCR demonstrated that transcription of UPEC dsdCX4 occurred at a low steady-state level but increased markedly within 5 min postaddition of d-Ser and remained at a steady state for at least 20 min (SI Appendix, Fig. S8B). Real-time PCR at 0, 10, and 20 min and quantitative real-time PCR at 0 and 20 min revealed that expression of CscR did not reduce the steady-state level of dsdCX4 transcription and did not affect the ability of dsdCX4 to respond to d-Ser (SI Appendix, Fig. S8B–E). It therefore appeared that CscR was not a direct repressor of d-Ser catabolism.

The activation of dsdA by a new upstream regulatory element seemed more plausible given the data presented in Fig. 6A. The transcript read density for dsdA appears to correlate well with that of cscA, supporting the assertion that dsdA transcription in this strain may be driven by cscAp. The deletion in LCV2 was particularly serendipitous as it not only allowed for control via increasing the proximity of cscAp to dsdA but also via derepression of cscAp due to loss of CscR. To further test how the genomic organization of these regions affected dsdA expression, the relevant chromosomal regions (from cscAp or z3622p to the sixth codon of dsdA) of WT, LCV2, or LCV10B were fused to gfp (Fig. 7A) in a reporter plasmid such that green fluorescent protein (GFP) production could be used as a proxy for DsdA expression. As expected, the fragments that comprised gfp under the control of the canonical DsdA control elements (Fig. 7, fragments i and iii) produced low levels of fluorescence (63 to 134 relative fluorescence units; RFUs) and behaved similarly. Fragments ii and iv that were amplified from LCV2 and 10B and comprised gfp that was predicted to be under the control of cscAp or z3622p, respectively, produced fluorescence significantly higher (2,001 and 257 RFUs) than that of the WT amplified fragments, reflecting the constitutive activation of DsdA in these genetic contexts. Of particular interest was the fact that GFP production from fragment ii was significantly higher in the ΔcscR mutant than the WT whereas no significant difference between WT and mutant was observed with fragment iv. This illustrates that DsdA expression in LCV2 is further facilitated by derepression of cscAp due to lack of CscR, whereas expression via z3622p in LCV10B is not affected by CscR. This also explains why complementation of the ΔcscR mutant tolerance phenotype (Fig. 6F) is possible despite the fact that the change in chromosomal context is the principal driver of DsdA activity (i.e., overexpression of cscR in trans is capable of repressing dsdA in the ΔcscR mutant through interaction with chromosomal cscAp).

Discussion

There is a growing appreciation for the role that the metabolic state of the gut plays in determining the behavior of both pathogenic and commensal bacteria. In particular, diet plays a vital role in determining health and disease outcomes, not only through manipulation of the nutritional balance of the host but also through modulation of bacterial species composition and gene expression (26, 27). It has been recently demonstrated that the use of trehalose as a food additive has led to the evolution of new lineages of Clostridium difficile with enhanced virulence and metabolic capacity (28), highlighting the importance of understanding how bacterial adaptation to dietary components can influence disease outcomes.
Many studies have pointed toward important roles for distinct dietary constituents in manipulating virulence and the progression of disease by A/E pathogens. Dietary fiber has been directly implicated in affecting disease severity in EHEC (29), while supplementation of diet with phytonutrients (plant extracts) (30) and pectin (31) reduce the severity of infection by *Citrobacter rodentium*—a natural pathogen of mice that employs a LEE-encoded type 3 secretion system similar to EHEC for intestinal colonization. Fucose has also been shown to repress expression of the LEE in EHEC via the fucose two-component sensor FusKR (32); however, the main source of free fucose is thought to be cleavage of intestinal mucins by commensal bacteria, while the contribution of diet to free fucose levels in the gut has been poorly studied (33). Similarly, biotin was shown to repress the LEE via the biotin sensor BirA which represses transcription of the global transcriptional regulator Fur, preventing it from activating the LEE (34). Accordingly, mice fed a high-biotin diet showed lower levels of intestinal colonization.

Of particular relevance to this study, catabolism of L-serine was recently shown to confer a competitive advantage to adherent invasive *E. coli* (AIEC) under chemically induced murine colitis and also to *C. rodentium* that can naturally induce colitis in mice (35). Pathogenic AIEC were better able to outcompete the non-pathogenic strain MG1655 when germfree mice were provided

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**Fig. 6.** Genomic rearrangements upstream of the inactive EHEC *dsd* locus result in activation of o-Ser catabolism, thereby allowing tolerance via a distinct mechanism. (A) Representative RNA-seq tracks of the WT and LCV2 cultured in MEM-HEPES o-Ser for 5 h. Gene tracks indicate WT, LCV2, and LCV10B organization. Red/blue dashed boxes indicate direct repeats identified flanking the deletion in LCV2/10B, respectively. Annealing sites for primers used to screen for deletions in this region are indicated by red lines above and below each gene track. (B and C) Growth curves of LCV2 and LCV10B cultured in MEM-HEPES ± 1 mM o-Ser. Error bars indicate SEM with a minimum of three experiments being performed. (D) Concentration of o-Ser in cell-free supernatant from MEM-HEPES cultures as determined by o-Ser assay. The experiment was performed in triplicate. Error bars indicate SEM. *P* < 0.05, **P** < 0.01, and ***P** < 0.001 by unpaired two-tailed Student’s t test. (E and F) Growth curves of the WT, ΔcscR (E), and cscR complementation strains (F) cultured in M9 + Glc ± 1 mM o-Ser. (G) Growth curves of the WT and ΔcscR cultured in MEM-HEPES ± 1 mM o-Ser. Error bars indicate SEM with a minimum of three experiments being performed (E–G).
were mutations in the GcvB binding site of

sstT; however, no change in cycA

transcription was observed, nor in gcvB

in LCV8 to 10 compared with WTD-Ser (Fig. 3 iv). In an in vivo context, this could potentially increase metabolic fitness in otherwise unfavorable environments.

The identification of genetically distinct mutants after 10 successive batch cultures is not surprising and is likely reflective of the fact that there are multiple routes to tolerance. Laboratory-based evolution experiments have been reported to result in convergent growth phenotypes via distinct modifications in gene expression states (36). Enteric species often partake in diverse adaptive strategies with genotypic and phenotypic heterogeneity allowing for enhanced versatility in the face of stress and competition (37, 38). In the complex and dynamic environment of the gut, mutations may be advantageous or deleterious depending on specific conditions. As such, heterogeneity represents a savvy route to establishment or maintenance of a successful niche. It is possible that there are further diverse mechanisms by which d-Ser tolerance can arise, each with varying implications for the expression of virulence and metabolism genes. Deep sequencing of a long-term adapted batch culture would yield interesting insight into potential heterogeneity of adaptation within a given population.

While the complete tolerance of LCV8 to 10 can be explained by reduced expression of SstT in conjunction with expression of a mutant GcvB, the reason for this reduced rate of transcription is unclear. There appears to be an accumulation of antisense reads in the 5′ untranslated region of sstT, so it is tempting to speculate that control via a regulatory RNA may be at play. Both sstT (39) and cycA (40) have been shown to be regulated by the small RNA GcvB; however, no change in gcvB transcription was observed, nor were mutations in the GcvB binding site of sstT in LCV8 to 10 detected (Table 1). It should be noted that WT EHEC responds to d-Ser by reducing transcription of cycA (−2.38-fold, \( P = 4.09 \times 10^{-3} \)). As such, EHEC may respond to d-Ser by decreasing the transcription of d-Ser transporters; however, the precise regulatory framework through which constitutive knockdown of sstT is achieved remains elusive.

The mechanism by which d-Ser exerts its toxic growth-inhibitory effect on catabolically inactive E. coli has been the subject of some debate. Using a mutant of E. coli K-12 unable to produce d-Ser deaminase, it was shown that addition of l-serine and pantothenate could rescue the tolerance phenotype, thereby implicating a block in l-serine biosynthesis or pantothenate synthesis in d-Ser–induced toxicity (41). This may also explain why d-Ser does not induce growth arrest in more complex media such as MEM-HEPES (Fig. 5B). d-amino acids can be misincorporated into the peptidoglycan of gram-negative bacteria in place of the canonical cell-wall constituent d-amino acids (9, 42). We have previously speculated that the mechanism of d-Ser toxicity may involve a disruption to the peptidoglycan-remodeling machinery that is essential for growth, due to a modified composition of the muramidopeptide chains (10). The results of this study indicate that regardless of the mechanism of toxicity, disruption of d-Ser transport into the cytosol is sufficient to abrogate the inhibition of growth. Analysis of 115 E. coli O157:H7 genomes (National Center for Biotechnology Information BLASTp) reveals strict conservation (>99%) at the amino acid level for CycA and SstT, indicating strong selective pressure. CycA and SstT possess multiple amino acid substrates including alanine, threonine, and glycine. Disruption of transport of these molecules may compromise fitness in certain environments by reducing the efficacy of transport and catabolism of specific carbon and nitrogen sources. The deletions observed upstream of dsdA in this study result in a similar evolutionary tradeoff with such mutants losing the ability to metabolize sucrose. It will be important to analyze the effect of such alterations in the context of an in vivo infection.

Laboratory-based evolution studies have yielded valuable insight into how E. coli adapts to stress (43–45) and can acquire the ability to derive nutrition from noncanonical carbon sources (46, 47). In this study, we have described how multiple genetic mechanisms converge in surmounting d-Ser–induced stress in EHEC with each mechanism having implications for important pleiotropic behaviors including virulence, stress response, and catabolic capacity. We previously showed that the sensing of d-Ser and concordant repression of type 3 secretion-dependent colonization restrict EHEC to the gut (10). The present study aimed to investigate how EHEC might overcome this evolutionary bottleneck. The growth inhibition associated with d-Ser is not linked to type 3 repression as shown by our previous work. The data presented here indicate that through genomic plasticity, EHEC can overcome both the repression of colonization and growth-inhibitory effects of d-Ser. Hence, these adaptations could expand the niche specificity of EHEC, enabling infection in extraintestinal sites such as the bladder, thereby challenging the dogma that EHEC are strictly intestinal pathogens (10, 15). Caution should be exercised with
this assertion as such mutations have yet to appear in the database. This could indicate that environmental substrate sampling via CycA acts as a critical regulator of niche restriction and loss of functionality could lead to a compromise in pathogen success. However, there have been recent reports of EHEC/UPEC hybrid strains (48, 49) with the ability to cause both intestinal and extra-intestinal disease, suggesting that adaptation to alternative niches may already be occurring. Understanding the in vivo ecological implications of the adaptive mechanisms described here will form the basis of our future work.

Materials and Methods

A complete list of bacterial strains, plasmids, and oligonucleotides used in this study can be found in SI Appendix, Tables S4–S6. A detailed description of all methodology is included in SI Appendix, Materials and Methods. This includes isolation of d-Ser-tolerant mutants, engineering of strains and plasmids, HeLa cell infections, DNA and RNA extraction, library preparation and sequencing, quantitative real-time PCR, d-Ser assay, and all data analysis tools.

Data Availability

Raw sequence data reported in this paper have been deposited in the European Nucleotide Archive under accession nos. ER54281497 to ER54281509 (genome sequencing) and ER54281510 to ER54281521 (RNA sequencing). All remaining data are presented in the paper and SI Appendix.

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