The Vibrio parahaemolyticus ToxRS Regulator Is Required for Stress Tolerance and Colonization in a Novel Orogastric Streptomycin-Induced Adult Murine Model

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Department of Biological Sciences, University of Delaware, Newark, Delaware, USA; Department of Medical Technology, University of Delaware, Newark, Delaware, USA; Discipline of Microbiology, School of Natural Sciences, National University of Ireland, Galway, Galway, Ireland; and U.S. Department of Agriculture, Agricultural Research Service, Delaware State University, Dover, Delaware, USA

Vibrio parahaemolyticus, a marine bacterium, is the causative agent of gastroenteritis associated with the consumption of seafood. It contains a homologue of the toxRS operon that in V. cholerae is the key regulator of virulence gene expression. We examined a nonpolar mutation in toxRS to determine the role of these genes in V. parahaemolyticus RIMD2210633, an O3K6 isolate, and showed that compared to the wild type, ΔtoxRS was significantly more sensitive to acid, bile salts, and sodium dodecyl sulfate stresses. We demonstrated that ToxRS is a positive regulator of ompU expression, and that the complementation of ΔtoxRS with ompU restores stress tolerance. Furthermore, we showed that ToxRS also regulates type III secretion system genes in chromosome I via the regulation of the toxRS gene. We examined the effect of ΔtoxRS in vivo using a new orogastric adult murine model of colonization. We demonstrated that streptomycin-treated adult C57BL/6 mice experienced prolonged intestinal colonization along the entire intestinal tract by the streptomycin-resistant V. parahaemolyticus. In contrast, no colonization occurred in non-streptomycin-treated mice. A competition assay between the ΔtoxRS and wild-type V. parahaemolyticus strains marked with the β-galactosidase gene lacZ demonstrated that the ΔtoxRS strain was defective in colonization compared to the wild-type strain. This defect was rescued by ectopically expressing ompU. Thus, the defect in stress tolerance and colonization in ΔtoxRS is solely due to OmpU. To our knowledge, the orogastric adult murine model reported here is the first showing sustained intestinal colonization by V. parahaemolyticus.
codes a LysR-family transcriptional regulator that is annotated as an LeuO homologue, but it was named CalR in their study as it was shown to respond to extracellular calcium levels (19). In the study of Gode-Potratz et al., a VP0350 deletion mutant exhibited increased cytotoxicity and T3SS-1 expression and is believed to act upstream of the ExsACDE system (19).

The mechanisms controlling global virulence gene regulation in V. parahaemolyticus are still poorly understood. The two-component regulator ToxRS has been shown to be essential for bacterial persistence and virulence during host infection with Vibrio cholerae (10, 49, 50, 75). ToxRS in V. cholerae controls the expression of several outer membrane proteins (Omp), namely, OmpU and OmpT, as well as enhancing the expression of ToxT, a transcription factor that is crucial for the regulation of the expression of the virulence factors toxin coregulated pilus and cholera toxin (37, 66–69). ToxRS also plays some role in V. cholerae survivability in the face of gastrointestinal barriers, such as bile salts, antimicrobial peptides, and acid stress, which are mediated by the ToxT activation of the expression of OmpU (43, 47, 67, 68). In V. cholerae the OmpU protein forms a multimeric porin in the outer membrane. OmpU has been implicated as an important stress response protein in response to extracellular stresses, such as organic acids, antimicrobial peptides, and anionic detergents (43, 47, 48, 67–69). There is some evidence suggesting that OmpU acts as a signal in the activation of RpoE, an alternative sigma factor that is important for the membrane stress response (42). The role of OmpU in in vivo survival is not entirely known, as V. cholerae ompU deletion mutants exhibit only a slight decrease in colonization (67, 68). OmpU has been shown, however, to play an important role in host colonization in Vibrio splendidus (16, 17). Vibrio parahaemolyticus contains a ToxRS homologue, but the role ToxRS plays in the pathogenicity of this organism has not been fully elucidated (38, 67, 68). Data suggesting that ToxRS is involved in regulating TDH expression (38) have been confounded by evidence suggesting that two ToxR-like regulators, virA and virB (located within VpaI–7), control the levels of TDH expressed in V. parahaemolyticus (33).

The aim of this study was to determine the function of ToxRS in V. parahaemolyticus. To accomplish this, we examined the effects of a nonpolar in-frame mutation in toxRS compared to those for the wild type under a number of stress conditions. We determined whether or not defects in the stress tolerance response were due to the ToxRS regulation of OmpU. In addition, we developed an adult murine model whereby animals were stably colonized by V. parahaemolyticus following orogastric inoculation. Here, we show that adult animals are susceptible to bacterial colonization following a single oral dose of antibiotic. We examined V. parahaemolyticus colonization throughout the intestine and monitored the adult mice by tracking bacterial shedding in fecal samples and determining V. parahaemolyticus CFU in the intestine. The effect of the deletion of the global regulator ToxRS on the ability of V. parahaemolyticus to colonize the intestine was also examined.

**MATERIALS AND METHODS**

**Bacterial strains, media, and growth conditions.** Studies utilized a spontaneous streptomycin-resistant mutant of the clinical isolate V. parahaemolyticus RMD2210633, an O3:K6 serotype, as a wild-type strain as described previously (44). Briefly, the strain was grown aerobically (250 rpm) at 37°C for 12 h in 5 ml Luria-Bertani (LB) broth (Fisher Scientific, Fair Lawn, NJ) containing 3% NaCl. This 5 ml of overnight culture was centrifuged for 10 min at 4,000 × g, and the pellet was resuspended in 100 μl of LB, 3% NaCl and plated onto LB, 3% NaCl plates containing 1 mg/ml of streptomycin (Fisher Scientific). Plates were incubated for 24 h at 37°C, and streptomycin-resistant colonies were further plated on LB, 3% NaCl plates containing 200 μg/ml of streptomycin. In addition, some experiments utilized a ΔtoxRS derivative of this streptomycin-resistant RMD2210633 strain, generated through splicing by overlap extension (SOE) PCR in a previous study (79). Two additional strains used in this study, ΔvscN1 and ΔvscN2, are T3SS-1 and T3SS-2 defective, respectively, and were previously constructed from RMD2210633 using SOE PCR in a previous study (79). Two additional strains used in this study, ΔvscN1 and ΔvscN2, are T3SS-1 and T3SS-2 defective, respectively, and were previously constructed from RMD2210633 using SOE PCR and allelic exchange (44). All strains and plasmids used in this study are listed in Table 1. All genetic manipulations utilized Escherichia coli strain DH5α and the diaminopimelic acid (DAP) auxotroph λpir and the diaminopimelic acid (DAP) auxotroph λpir. Unless otherwise stated, bacteria were grown on LB broth containing 3% NaCl at 37°C with aeration. An otherwise stated, bacteria were grown on LB broth containing 3% NaCl at 37°C with aeration. An otherwise stated, bacteria were grown on LB broth containing 3% NaCl at 37°C with aeration. An otherwise stated, bacteria were grown on LB broth containing 3% NaCl at 37°C with aeration. An otherwise stated, bacteria were grown on LB broth containing 3% NaCl at 37°C with aeration. An otherwise stated, bacteria were grown on LB broth containing 3% NaCl at 37°C with aeration.

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

<table>
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<tr>
<th>Strain or plasmid</th>
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<th>Reference or source</th>
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<tr>
<td>ΔtoxRS</td>
<td>RMD2210633 ΔtoxRS(VP0819-0820)</td>
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<td>WB2467</td>
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<td>pDSΔleuO</td>
<td>pDS132 harboring truncated leuO</td>
<td>(VP0350)</td>
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**RNA extraction and quantitative real-time RT-PCR.** Total RNA was extracted from V. parahaemolyticus strains grown for 4 h in LB containing 3% NaCl using the RNAProtect bacterial reagent (Qiagen, Valencia, CA) and an RNaseasy mini kit (Qiagen) by following the manufacturer’s protocols. The RNA samples were quantified using a Nanodrop spectrophotometer (Thermo Scientific, Waltham, MA) and subsequently treated with Turbo DNase (Invitrogen, Carlsbad, CA), per the manufacturer’s protocol, to remove any contaminating genomic DNA from the samples. cDNA was synthesized using Superscript II reverse transcriptase (RT) (Invitrogen) according to the manufacturer’s protocol, using 500 ng of RNA as a template and priming with 200 ng of random hexamers. cDNA samples were then diluted 1:250 and used for quantitative real-time PCR (qPCR). Real-time PCRs used the HotStart-IT SYBR green qPCR master mix (USB, Santa Clara, CA) and were run on an Applied Biosystems 7500 fast real-time PCR system (Foster City, CA). Gene-specific primers were designed using Primer 3 software according to the real-time PCR guidelines and are listed in Table 2. Data were analyzed using Applied Biosystems 7500 software. Expression levels of each gene, as determined by their threshold cycle (Ct) values, were normalized using the 16S rRNA gene to correct for sampling errors. Differences in the ratio of gene expression were determined using the ΔΔCt method. All assays were performed in duplicate, and all experiments were repeated.
Complementation of \( \Delta toxRS \). The \( \Delta toxRS \) strain was complemented with the \( ompU \) gene (VP2467), the \( leuO \) gene (VP0350), or the \( toxRS \) operon (VP0819-20) using the pBAD33 vector. PCR primers were designed to amplify a promoterless copy of VP2467 encoding OmpU (Table 2). Briefly, the \( ompU \) gene was cloned into vector pJL2.1/blunt and transformed into \( E. coli \) DH5a λpir. The fragment was then subcloned into the vector pBAD33, resulting in pWBW2467, and subsequently transformed into \( E. coli \) BL21 5 Apir. The \( E. coli \) BL21 5 Apir strain harboring pWBW2467 was cross-streaked with the \( \Delta toxRS \) strain onto LB plates containing 0.3 mM DAP, which allows for the conjugative transfer of pWBW2467. The resulting growth was streaked onto LB, 3% NaCl plates containing chloramphenicol and streptomycin (but no DAP) to positively select for \( \Delta toxRS \) cells harboring pWBW2467 and to negatively select against \( E. coli \) BL21 5 Apir. To induce the expression of the complemented genes, strains harboring pWBW2467 were grown in the presence of 0.05% arabinose. This protocol was repeated for the gene VP0350 (leuO) and the \( toxRS \) operon.

Complementation of \( V. parahaemolyticus \) RIMD2210633 VP0350 deletion mutant. An in-frame deletion mutant was created in VP0350, which carries a homologue of \( leuO \), using SOE PCR and allelic exchange (28). Primers were designed using the \( V. parahaemolyticus \) RIMD2210633 genome sequence (40) as the template and purchased from Integrated DNA Technologies (Coralville, IA). These primers were used to perform SOE PCR and obtain a 303-bp truncated version of the VP0350 gene. The \( \Delta VP0350 \) PCR fragment was cloned into the suicide vector pDS132 (28), which was designated pDS\( \Delta VP0350 \). pDS\( \Delta VP0350 \) was subsequently transformed into the \( E. coli \) strain \( \beta 2155 \) Apir. pDS\( \Delta VP0350 \) was conjugated into \( V. parahaemolyticus \) RIMD2210633 via cross-streaking on LB plates containing 0.3 mM DAP. Growth from these plates was transferred to LB plates containing 3% NaCl, streptomycin, and chloramphenicol. The 3% NaCl allowed for optimal \( V. parahaemolyticus \) growth. The absence of DAP from these plates, plus the addition of chloramphenicol, selected only for \( V. parahaemolyticus \) cells that harbored pDS\( \Delta VP0350 \). Exconjugate colonies were cultured overnight in LB, 3% NaCl without chloramphenicol and subsequently serially diluted and plated on LB, 3% NaCl, 10% sucrose to select for cells which had lost pDS\( \Delta VP0350 \). Double-crossover deletion mutants were screened by PCR using the SOEFLVP0350F and SOEFLVP0350R primers and sequenced for verification.

Construction of \( \beta\)-galactosidase-expressing RIMD strain. To coculture mutant strains with the isogenic wild-type strain for competition assays, we developed a blue-white screening assay for \( V. parahaemolyticus \). To accomplish this, we cloned the \( \beta\)-galactosidase gene (VP2403) as our chromosomal insertion point, with the cryptic beta-D-galactosidase subunit alpha gene (VP2403) as our chromosomal insertion point, using a modified SOE PCR protocol. Two regions of VP2403 were amplified using 2 sets of primer pairs (Table 2). These products were then spliced together by homologous EcoRI-KpnI-BamHI sequences incorpo-

### TABLE 2 Primers used in this study

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rated into the 3′ end of the reverse primer in pair 1 and the 5′ end of the forward primer in pair 2. The resulting single product was cloned into the vector pJE112 (pJE1.2-VP2403SOE) and transformed into E. coli DH5α Apir. Primers were also designed to amplify a region in the V. cholerae N16961 genome which encompasses the β-galactosidase gene plus 380 bases of upstream sequence to include a putative ρ′ promoter. The 5′ V. cholerae primer contains an EcoRI site, and the 3′ V. cholerae primer contains a BamHI site, allowing for insertion into the VP2403 SOE product. The V. cholerae β-galactosidase product was also cloned into pJE112 (pJE1.2-VplacZ) and transformed into E. coli DH5α Apir. Both plasmids were extracted from their host strains, and the EcoRI and BamHI fragments from pJE1.2-VplacZ were digested and ligated into the similarly digested pJE1.2-VP2403SOE, resulting in the plasmid pJE1.2-VpVclacZ, and transformed back into E. coli DH5α Apir. This plasmid was then extracted and digested with XbaI and Sall to completely excise the V. parahaemolyticus-V. cholerae lacZ fragment, which was then subcloned into the suicide vector pDS132 (named pDSVpVclacZ), transformed into E. coli B2155 Apir, and transferred to V. parahaemolyticus via conjugation. Transconjugates containing the pDSVpVclacZ plasmid were selected by plating on LB, 3% NaCl containing streptomycin and chloramphenicol. Exconjugate colonies were cured of the pDSVpVclacZ plasmid by replating onto LB, 3% NaCl plates containing 10% sucrose and 5-bromo-4-chloro-3-indolyl-β-D-galactoside product was also cloned into pJET1.2-lof (+)promoter. The 5′ V. cholerae β-galactosidase product was also cloned into pJE112 (pJE1.2-VplacZ) and transformed into E. coli DH5α Apir. Both plasmids were extracted from their host strains, and the EcoRI and BamHI fragments from pJE1.2-VplacZ were digested and ligated into the similarly digested pJE1.2-VP2403SOE, resulting in the plasmid pJE1.2-VpVclacZ, and transformed back into E. coli DH5α Apir. This plasmid was then extracted and digested with XbaI and Sall to completely excise the V. parahaemolyticus-V. cholerae lacZ fragment, which was then subcloned into the suicide vector pDS132 (named pDSVpVclacZ), transformed into E. coli B2155 Apir, and transferred to V. parahaemolyticus via conjugation. Transconjugates containing the pDSVpVclacZ plasmid were selected by plating on LB, 3% NaCl containing streptomycin and chloramphenicol. Exconjugate colonies were cured of the pDSVpVclacZ plasmid by replating onto LB, 3% NaCl plates containing 10% sucrose and 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-Gal), with double-crossover mutants appearing blue due to the functional activity of the β-galactosi-
dase gene.

**Growth and survival analysis.** Wild-type *V. parahaemolyticus*, ΔtoxRS, and ΔtoxRS complemented with either lenO or toxRS (strains WBW2467 and WBW80819-820, respectively) were examined for survival in 4 mM acetic acid, sodium cholate (bile salts) (Sigma), and sodium dodecyl sulfate (SDS) (Fisher Scientific, Fair Lawn, NJ). For survival ass-
ays, strains were first grown overnight in LB broth containing 3% NaCl with aeration at 37°C. Following overnight growth, 100 μl of each culture was inoculated into 5 ml of fresh LB, 3% NaCl and allowed to grow to mid-exponential phase. The cultures were then pelleted and resuspended in LB, 3% NaCl with the pH adjusted to 5.5 with HCl and incubated at 37°C for 30 min. After 30 min, the cultures were pelleted again and sub-
sequently resuspended in fresh LB, 3% NaCl, supplemented with 4 mM acetic acid, and allowed to incubate for an additional 60 min. At specified time points, cultures were serially diluted and plated for CFU to deter-
mine surviving bacteria. This protocol was repeated for conditions of bine stress (15% deoxycholate) or SDS stress (0.5% SDS), with the exception that no preadaption phase was used for these conditions.

**Animals and inoculations.** Male C37BL/6 mice, aged 6 to 10 weeks, were purchased from Jackson Laboratory (Bar Harbor, ME) and housed in standard cages under specific-pathogen-free conditions. Mice were caged in groups (3 to 5 per group) and provided standard mouse feed and water *ad libitum*. All experiments involving mice were approved by the University of Delaware Institutional Animal Care and Use Committee. Twenty-eight hours prior to orogastric inoculation, food and water were withdrawn from the animals. Four hours later, mice were orogastrically administered 100 μl of either phosphate-buffered saline (PBS) or streptomycin solution (200 mg/ml PBS) with a sterile 18-gauge gavage needle (Roboz Surgical Instrument Co., Gaithersburg, MD), and food and water were immediately returned. After 20 h, mice were fasted again for 4 h, following which all animals were orogastrically dosed with 100 μl of bacterial suspension in PBS using a sterile gavage needle without anesthetization.

Bacterial inocula of wild-type or mutant strains of *V. parahaemolyticus* were prepared as follows. Glycerol stocks were streaked onto LB media supplemented with NaCl (final concentration of 3%) and streptomycin and grown at 37°C. Streptomycin was included in media at a final concentra-
tion of 200 μg/ml. A single colony was used to inoculate 10 ml of LB, 3% NaCl plus streptomycin and grown overnight at 37°C with aeration (230 rpm). The following day, 25 ml of LB, 3% NaCl plus streptomycin was inoculated with a 2% inoculum of overnight culture and incu-
bated for 4 h at 37°C with aeration (230 rpm). The optical density at 600 nm (OD600) was measured on a 1:10 dilution of bacterial culture to verify that growth was within an expected range, as previously determined by plating cultures to determine CFU per ml (data not shown). Bacteria were pelleted by centrifugation at 2,000 × g for 5 min, washed once with PBS, and centrifuged again. The bacterial cell pellets were resuspended in PBS to a concentration between 5 × 10^8 and 1 × 10^10 CFU/ml. Serially diluted inoculum was plated for each experiment to determine the actual dose administered. Water was provided to animals immediately following orogastric inoculation, with food returned 2 h postinoculation.

**Determination of *V. parahaemolyticus* CFU in intestinal tissues.** Forty-eight h postinfection, mice were sacrificed by carbon dioxide asphyxiation and tissues were aseptically removed. For total intestinal CFU deter-
mation, the entire intestine (encompassing the duodenum to the descending colon) was placed in 4 ml of PBS and mechanically disrupted with a Tissue Tearer (BioSpin Products, Inc., Bartlesville, OK). Homoge-
nized samples were serially diluted, plated on LB, 3% NaCl plus streptomycin, and incubated at 37°C. In indicated experiments, the gastrointestinal tracts from individual animals were analyzed as separate sections of tissue to elucidate regions of the intestine preferentially colonized by bac-
teria. Here, the intestine was separated into the small intestine (duode-
num to ileum), the cecum, and the large intestine. Tissue sections were flushed with PBS to remove the fecal contents, homogenized in 4 ml of PBS, serially diluted, plated on LB, 3% NaCl plus streptomycin, and incubated at 37°C.

**Determination of bacterial CFU shed in fecal samples.** Fresh fecal pellets were collected from individual animals (2 to 4 pellets per animal) at 24-h intervals. Pellets were weighed and placed in 2 ml of sterile PBS for mechanical disruption by homogenization. All samples were serially di-
luted in PBS, plated on LB, 3% NaCl plus streptomycin, and incubated at 37°C. For each experiment, randomly selected colonies were patched onto the highly selective vibrio medium thiosulfate citrate bile salts sucrose (TCBS) agar (Fisher Scientific) and incubated overnight at 37°C to con-
firm that the counts were of *V. parahaemolyticus* and not due to the out-
growth of other intestinal microflora.

**In vivo colonization competition assay.** The inocula for competition assays were prepared as outlined above, with the following modifications. Overnight cultures of the wild-type, WBWlacZ, ΔtoxRS, ΔvscN1, and ΔvscN2 strains were diluted 1:50 into 25 ml LB, 3% NaCl with streptomycin and grown for 4 h. Four-hour cultures of WBWlacZ, wild-type strain RIMD2210633, ΔtoxRS, ΔvscN1, and ΔvscN2 strains were pelleted by cen-
trifugation, washed in PBS, and centrifuged again. The resulting bacterial pellet was resuspended in PBS to a concentration of approximately 1 × 10^10 CFU based on the culture OD_600_. A 1-ml aliquot of each mutant and the wild-type strain WBWlacZ was combined in fresh PBS, yielding an inoculum of 1 × 10^10 CFU, with a ratio of 1:1 mutant cells to WBWlacZ cells. A portion of the inoculum was serially diluted and plated onto LB, 3% NaCl plates with streptomycin (50 μg/ml X-Gal) to determine the exact ratio of CFU in the inoculum. A 100-μl aliquot of the inoc-
ulum was added to 5 ml of LB for *in vitro* competition assays. The cultures were grown overnight at 37°C with aeration. Serial dilutions were plated on LB-Sm supplemented with 40 μg/ml X-Gal (Fischer), and recovered CFU were counted for the *in vitro* competitive index (CI). The *in vitro* experiments were performed in triplicate at least twice. Mice were prepared by following the streptomycin-treated adult mouse protocol outlined above. Twenty-four hours after infection, mice were sacrificed and the entire gastrointestinal tract was harvested. Samples were placed into 8 ml of PBS, mechanically homogenized, and serially diluted in PBS. Samples were then plated on plates of LB, 3% NaCl with streptomycin and X-Gal to determine the output CFU for both strains. The CI was determined using the following formula: CI = ratio out(mutant/wild-type)/ratio in(mutant/wild-type).“Statistical analysis.** Data from replicate experiments were analyzed using either a Kruskal-Wallis one-way analysis of variance (ANOVA) fol-
lowed by a Dunn’s multiple comparison posttest or unpaired Student’s *t
RESULTS

**ToxRS controls the expression of ompU in *V. parahaemolyticus***.

In *V. cholerae*, ToxRS directly regulates the expression of the outer membrane proteins OmpU and OmpT (37, 66–69). *Vibrio parahaemolyticus* contains an OmpU homologue (VP2467) but not an OmpT homologue. We performed real-time PCR to explore whether the ToxRS system controls the expression of *ompU* in *V. parahaemolyticus*. RNA was extracted from the ΔtoxRS and isogenic wild-type strains grown under standard (pH 7) and low-acid (pH 5) conditions. Low pH stress was previously demonstrated to induce the expression of toxR in *V. parahaemolyticus* (79). In our study, in wild-type cells grown in LB, 3% NaCl, pH 5 (low pH), *ompU* showed a significant increase in expression of approximately 17-fold (*P < 0.01) compared to that of wild-type cells grown in LB, 3% NaCl, pH 7 (neutral pH) (Fig. 1). When ΔtoxRS cells were grown at pH 7, the expression ratios of *ompU* to those of wild-type cells grown under the same condition were ~500-fold, a highly significant decrease (*P < 0.001*). Similarly, when ΔtoxRS cells were grown at pH 5, *ompU* expression was ~160-fold (*P < 0.001*) compared to that of the wild-type strain grown under the same conditions (Fig. 1). No statistically significant difference in *ompU* expression was found between growth at pH 7 and that at pH 5 in the ΔtoxRS strain. These results indicate that the deletion of toxRS significantly decreased *ompU* expression and thus plays a role as a positive regulator of *ompU* expression.

**ΔtoxRS controls the expression of T3SS-1 genes via regulation of toxO**.

It has previously been demonstrated that the *V. parahaemolyticus* ΔtoxRS strain is more cytotoxic than the wild-type strain toward Caco-2 cells (79). Similarly, it has been demonstrated that the type III secretion system (T3SS-1) gene encoded by VP0350 (40) in both wild-type and ΔtoxRS cells grown under optimal growth conditions (LB, 3% NaCl, pH 7). Under such conditions, it has been previously demonstrated that wild-type cells do not express T3SS-1 genes (86). We first examined VP1680, which encodes the effector protein VopQ that is secreted via T3SS-1 and plays an important role in cytotoxicity as well as in the host immune response (8, 44, 60, 73). We found an approximately 3-fold increase in VP1680 expression in the ΔtoxRS background compared to that in wild-type cells (*P < 0.05*) (Fig. 2A). We then examined VP1656, which encodes a protein, VopD, that shares homology with the type III secreted proteins PopD and YopD of *Pseudomonas aeruginosa* and *Yersinia enterocolitica*, respectively. VopD and YopD act as translocators that transport effector proteins across the eukaryotic membrane (62). In the ΔtoxRS mutant, the expression levels of VP1656 and VP1695 were not significantly higher than that in wild-type cells (Fig. 2A). VP1654 and VP1695 encode T3SS structural proteins with similarity to the MxiH T3SS secretion needle protein superfamily and the YscD T3SS inner membrane secretion protein superfamily, respectively (14). In the ΔtoxRS mutant, the expression levels of VP1654 and VP1695 were 2.55- and 2.1-fold higher, respectively, compared to those of wild-type cells (*P < 0.05*) (Fig. 2A). These results indicate that genes associated with the T3SS-1 in chromosome I show increased expression in the ΔtoxRS mutant and that ToxRS plays a role in their regulation.

Because the ΔtoxRS strain shared a similar phenotype with the recently described VP0350 deletion mutant with regard to cytotoxicity and T3SS-1 expression (19), we hypothesized that ToxRS indirectly regulates T3SS-1 expression via the regulation of the *toxO* homologue VP0350. To address this, we examined the expression of VP0350 in the ΔtoxRS strain as well as in the isogenic wild-type strain (Fig. 2B). We found that in wild-type cells that were subjected to 30 min in LB, 3% NaCl, pH 5, the expression of VP0350 increased 2.8-fold (*P < 0.05*) compared to that in wild-type cells grown under standard conditions (LB, 3% NaCl, pH 7) (Fig. 2B). However, the expression of VP0350 was significantly decreased (*P < 0.001*) in the ΔtoxRS strain grown in LB, 3% NaCl, pH 7 compared to that of the isogenic wild-type cells examined under the same conditions (Fig. 2B). Similarly, we found that in ΔtoxRS cells that were subjected to 30 min in LB, 3% NaCl, pH 5, the expression of VP0350 was significantly decreased (*P < 0.001*) compared to that of wild-type cells examined under the same conditions (Fig. 2B). Additionally, no significant difference in VP0350 expression was observed between the ΔtoxRS strain in neutral and pH 5 conditions. This indicates that the expression of VP0350 does indeed fall under the regulation of the ToxRS system.

Lastly, we hypothesized that the lack of VP0350 expression seen in the ΔtoxRS strain was the cause of the increased expression of the T3SS-1 genes. To examine this, we constructed a VP0350 deletion mutant and also complemented the ΔtoxRS strain with the VP0350 gene cloned into the expression vector pBAD33. As expected, the ΔVP0350 mutant demonstrated an increase in the expression of the T3SS-1 effector gene VP1680 compared to that of the isogenic wild-type strain (*P < 0.05*) (Fig. 2C). When the ΔtoxRS strain was complemented with the VP0350 gene, the expression of VP1680 was reduced to levels not significantly different from those of the isogenic wild-type strain (Fig. 2C).
ToxRS regulates the expression of T3SS-1 genes and leuO (VP0350). (A) The expression levels of two genes encoding T3SS-1 substrates (VP1680 and VP1695) were examined in the ΔtoxRS and isogenic wild-type strains. An unpaired Student’s t test was used to compare differences in gene expression in the ΔtoxRS strain to expression in wild-type cells. *, P < 0.05. (B) The expression of gene VP0350, which encodes a LysR-family transcriptional regulator, was examined in the ΔtoxRS and wild-type strains and was grown under standard conditions as well as ToxR-inducing conditions (pH 5). An unpaired Student’s t test was used to compare differences in VP0350 expression under acid conditions and in the ΔtoxRS strain to expression in wild-type cells grown in LB, 3% NaCl, pH 7. *, P < 0.05; ***, P < 0.001. (C) The expression of VP1680, which encodes the T3SS-1 effector protein VopQ, was examined in the ΔtoxRS, ΔVP0350, and WBW0350 strains. An unpaired Student’s t test was used to compare differences in VP1680 expression in the ΔtoxRS strain, ΔVP0350, and WBW0350 strains to expression in the wild type. *, P < 0.05. WBW0350 is a ΔtoxRS strain that ectopically expresses VP0350. Bars represent the relative expression of the given gene normalized to the 16S rRNA gene and compared to expression in wild-type cells grown in LB, 3% NaCl, pH 7. RNA was extracted for each condition on two separate occasions, and qPCR was run in duplicate for each sample. Error bars indicate standard deviations.

**FIG 2**

Together, these results indicate that the increase in T3SS-1 expression seen in the ΔtoxRS strain is due to its regulation of the leuO gene carried by VP0350.

**OmpU is important for survival under extracellular stress conditions.** ToxRS has been shown previously to be an important stress response regulator in *V. cholerae* (43, 47, 48, 67–69). Additionally, we previously demonstrated that the ToxRS system in *V. parahaemolyticus* plays an important role in the organism’s ability to elicit a successful acid tolerance response (79). Provenzano and Klose demonstrated that ToxRS was important for growth in bile salts and SDS (69); however, a mechanism for how ToxRS protects *V. parahaemolyticus* in the face of these stresses has yet to be determined. Because organic acids as well as bile and other detergents are believed to act on the bacterial cell membrane, we aimed to determine if the regulation of OmpU by ToxRS was the mechanism for stress tolerance. Therefore, we examined the role of ToxRS in the cell stress response to organic acid and anionic detergents, such as bile and SDS. When the ΔtoxRS strain and the wild-type strain were preadapted in 1 mM acetic acid and then subjected to 4 mM acetic acid, there was a significant defect in the survival of the ΔtoxRS strain after 30 min (P < 0.001) (Fig. 3A). After 60 min, there were no detectable viable ΔtoxRS cells, whereas the wild-type strain exhibited nearly 100% survival throughout the 60 min in acetic acid (P < 0.0001) (Fig. 3A). However, when ompU was ectopically expressed in the ΔtoxRS background (designated strain WBW2467), the ability to mount a successful adaptive acid tolerance response was restored, and this strain exhibited 97 and 75% survival after 30 and 60 min, respectively (Fig. 3A). The ΔtoxRS strain also exhibited a defect in the ability to survive in the presence of 15% bile, demonstrating 0.1 and 0.04% survival after 20 and 40 min, respectively (Fig. 3B). This was a significantly lower level of survival than that of the isogenic wild-type strain, which exhibited 31% (P < 0.001) and 24% (P < 0.0001) survival during 20 and 40 min in bile, respectively. WBW2467 exhibited no significant differences in survival compared to that of the wild-type strain for both 20 and 40 min in the presence of 15% bile, indicating that ectopically expressing ompU was enough to restore survival in bile to the ΔtoxRS strain. The ΔtoxRS strain phenotype was similar when subjected to 0.5% SDS stress. The ΔtoxRS strain exhibited 2 and 0.0005% survival in 0.5% SDS, which was significantly lower than that of the wild-type strain for both time points (P < 0.0001) (Fig. 3C). Similarly, WBW2467 showed no significant difference between itself and the wild-type strain, again indicating that the ectopic expression of ompU was enough to rescue the sensitive phenotype of ΔtoxRS. We also complemented our in-frame ΔtoxRS mutation with a functional copy of toxRS (strain WBW0819-0820) to confirm that it has no polar effects, and as expected under all stress conditions examined, the survival of WBW0819-0820 was not significantly different from that of the wild-type strain (Fig. 3). These results indicate that ToxRS is an important stress response regulator in *V. parahaemolyticus* through its regulation of OmpU.

ToxRS is an important mediator of the colonization of the streptomycin-treated adult mouse model. Adult C57BL/6 mice were given an orogastric dose of either PBS or streptomycin (20 mg/animal) 24 h prior to orogastric inoculation with *V. parahaemolyticus* RIMD2210633. Two days postinfection, total intestinal tissues were isolated and assessed for weight and bacterial content. The average intestinal weights in *V. parahaemolyticus*-infected animals pretreated with streptomycin were significantly higher than...
for successful colonization. Having observed that *V. parahaemolyticus* persists in the murine intestine after orogastric challenge only after streptomycin administration, we next assessed whether the organism was distributed equally throughout the gastrointestinal tract or if it preferentially localized to discrete segments of the intestine. Mice given an orogastric dose of either PBS or streptomycin were subsequently infected with *V. parahaemolyticus*. Two days postinfection, the intestinal tissues were removed and fecal contents cleared, with the small intestine, cecae, and large intestine independently homogenized and plated to determine the tissue-associated bacterial burdens (Fig. 4B). As we observed initially, *V. parahaemolyticus* was not detected in tissues from animals pretreated with PBS prior to infection (Fig. 4B), while mice given an orogastric dose of streptomycin were found to have *V. parahaemolyticus* in all of the intestinal segments tested (Fig. 4B). Within an individual animal, the levels of *V. parahaemolyticus* were typically more than 1-log lower in the small intestine compared to those found in either the cecum or large intestine (*P < 0.05*) (Fig. 4B). We also examined both the spleen and mesenteric lymph nodes from both PBS- and streptomycin-treated infected mice and recovered no *V. parahaemolyticus* CFU from either.

The persistence of intestinal colonization with *V. parahaemolyticus* was examined by measuring the bacterial loads in fecal samples from infected animals. Animals pretreated with either PBS or streptomycin were infected with *V. parahaemolyticus*, and fresh fecal pellets were collected at daily intervals and plated to determine *V. parahaemolyticus* CFU in the fecal content. High levels of *V. parahaemolyticus* were observed only in those samples collected from mice that were treated with streptomycin prior to infection compared to mice treated only with PBS. The bacterial numbers decreased slowly over time but remained above $10^6$/g in all streptomycin-treated animals for 7 days (Fig. 5). Colonized mice displayed no overt signs of morbidity during the course of infection. We next examined whether or not variation in the inoculum dose affects the establishment of colonization. To accomplish this, we examined mice pretreated with streptomycin in groups (3 to 5 mice/group) given a range of doses orogastrically (1.5 x 10^7, 5 x 10^6, 1.5 x 10^6, and 1.7 x 10^5 CFU). Fecal pellets were collected 24 and 72 h postinfection, and *V. parahaemolyticus* CFU counts were determined. All mice were colonized to the same extent for all doses examined (data not shown).

To directly assess the importance of the ToxRS system in the colonization of the host and to increase the utility of our colonization model, we developed a competition assay by creating a β-galactosidase-positive wild-type strain of *V. parahaemolyticus*, named WBWlacZ. To confirm that WBWlacZ grows similarly to the wild type, we compared growth rates in LB, 3% NaCl at 37°C overnight and found no difference in growth rates between the two strains (data not shown). In addition, we performed *in vitro* competition assays, including growth overnight in LB, 3% NaCl at 37°C of mixed cultures of WBWlacZ and the test strains, and we found no difference in the CFU counts between the strains examined (data not shown). By using the streptomycin mouse model and a mixed culture of the WBWlacZ and ΔtoxRS strains, we can directly measure CFU using a blue/white screen *in vivo*. Equal volumes of WBWlacZ and the test strains were inoculated into streptomycin-treated adult mice. After 24 h postinfection, the entire gastrointestinal tract of each mouse was removed, homogenized, and plated for CFU to assay for the presence of both strains.

**FIG 3** Effect of organic acid, bile, and SDS stress on the survival of the ΔtoxRS deletion mutant. Wild-type (closed circles), ΔtoxRS (closed triangles), WBW2467 (closed squares), or WBW0819-0820 (open triangles) cells were grown in LB with 3% NaCl and subjected to (A) 60 min of organic (4 mM acetic acid, pH 4.5) stress after preadaptation for 30 min in 1 mM acetic acid adjusted to pH 5.5 (HCl), (B) 40 min in 15% sodium cholate, or (C) 40 min in 0.5% SDS stress. Survival was determined by dividing the surviving population at various time points by the initial population. All cultures were grown in triplicate, and each experiment was performed at least twice. Error bars indicate standard deviations. P values were calculated using an unpaired Student’s t test with a 95% confidence interval. Asterisks denote significant differences between groups. ***, P < 0.001; ****, P < 0.0001. WT, wild-type *V. parahaemolyticus* RIMD2210633; WBW2467, a ΔtoxRS strain that ectopically expressed ompU.

those from the control *V. parahaemolyticus*-infected mice pretreated with PBS (*P < 0.05*) (data not shown). Importantly, when these intestinal samples were examined to determine bacterial burdens, tissues from animals pretreated with PBS did not contain detectable levels of *V. parahaemolyticus*, while those pretreated with streptomycin contained significantly higher *V. parahaemolyticus* bacterial loads (*P < 0.0001*) (Fig. 4A). These results demonstrate that the administration of a single antibiotic treatment to mice prior to orogastric challenge with *V. parahaemolyticus* allows...
We first compared WBWlacZ and the wild-type strains in coculture in vivo to confirm that they behave similarly and found no difference in the competitive index between the two strains. We next examined a coculture of WBWlacZ and /H9004 toxRS and determined that after 24 h postinfection the /H9004 toxRS strain is significantly (P < 0.001) outcompeted (4-fold) by the WBWlacZ strain (mean competitive index of 0.25 ± 0.13) (Fig. 6). These results indicate that ToxRS is important for colonization, probably as a positive regulator of the stress response for the in vivo survival of V. parahaemolyticus. We also examined the fitness of /H9004 vscN1 (T3SS-1 mutant) and ΔvscN2 (T3SS-2 mutant) versus WBWlacZ in our colonization model. Both secretion systems are well-characterized virulence factors; however, their role in colonization has thus far not been determined. Using the streptomycin-treated adult mouse model and the WBWlacZ strain, we found no significant difference in the competitive index for either of the T3SS mutants compared to that of the isogenic wild-type strain (Fig. 6).

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To assess whether or not the defect in the ΔtoxRS mutant colonization in the streptomycin-treated adult mouse model is due to ompU, we performed a competition assay examining the ΔtoxRS mutant complement with ompU (WWB2467) compared to the wild type (Fig. 6). We found that WWB2467 had a competitive index of 0.72 ± 0.18, which, while significantly higher than that of the ΔtoxRS mutant (P < 0.05), was not significantly different from that of the isogenic wild-type strain. These results indicate that the deficiency in colonization exhibited by the ΔtoxRS strain can be alleviated by ectopically expressing the ompU gene.

DISCUSSION

The consumption of V. parahaemolyticus-contaminated seafood is the leading cause of seafood-associated bacterial gastroenteritis in the United States and Asia (5, 12, 13, 46). To date, relatively little is known about the bacterial mechanisms responsible for colonization following its entry into the human host. In an effort to begin investigating genes necessary for the colonization, survival, and proliferation of V. parahaemolyticus within the gastrointestinal tract, we examined the global regulator ToxRS. Similarly to what has previously been shown for V. cholerae, our ΔtoxRS mutant is sensitive to a number of stress conditions that the bacterium encounters in vivo, such as acid and bile salt stresses. Our data suggest that the defect in stress tolerance is due to the ToxRS regulation of ompU, which encodes a major outer membrane protein found in a wide range of enteric bacteria.

In V. cholerae, the ToxRS regulon is essential for virulence and has previously been shown to control the expression of a cascade of genes involved in bacterial pathogenesis (cholera toxin and toxin coregulated pilus), along with several outer membrane proteins (OmpU and OmpT), which enhance the resistance of the organism to antimicrobial compounds in the intestine (reviewed in references 10, 15, and 63). Our results show that a V. parahaemolyticus mutant of ToxRS does not colonize the mouse intestine as well as the wild type does. The role of ToxRS in the pathogenesis of V. parahaemolyticus has not been defined, even though there is evidence that it controls the expression of outer membrane proteins and bacterial resistance to membrane disruption by intestinal detergents (bile salts) (67). In the V. parahaemolyticus ΔtoxRS mutant, we observed a 500-fold decrease in the expression of ompU (VP2467) relative to that of the wild type by quantitative real-time PCR analysis (Fig. 1). We previously showed that under acidic conditions V. parahaemolyticus expression of toxRS is induced, and in the absence of ToxRS the organism has a significantly reduced ability to survive at low pH (79). In this study, we have demonstrated that the ΔtoxRS strain was also deficient in survival in high concentrations of bile and SDS. These defects, along with the defect in organic acid survival, were shown to be due to this strain’s inability to express the ompU gene, which would implicate OmpU as playing an important role in protection from extracellular stresses. Interestingly, following the coinfection of streptomycin-treated animals with both the V. parahaemolyticus RIMD2210633 ΔtoxRS and WWBWlacZ strains, we observed a significant (P < 0.001) reduction in the ability of the ΔtoxRS mutant to colonize the intestine. Thus, in addition to demonstrating that ToxRS and OmpU are important stress response systems, our results show that both proteins are important for colonization as well.

We have also previously shown that the ΔtoxRS strain exhibits increased cytotoxicity in vitro (79). In an effort to further characterize this phenotype, we have demonstrated in this study that the V. parahaemolyticus ToxRS system is a negative regulator of the type III secretion system found in chromosome I. This regulation is indirect and is mediated by the LeuO transcription factor homologue encoded by VP0350. ToxRS is a positive regulator of leuO expression, and LeuO is a negative regulator of T3SS-1 gene expression. This ToxRS pathway is unique to any Vibrio species thus far described. Previously, Call and colleagues demonstrated that the genes VP1699, VP1698, and VP1701 are homologues of Pseudomonas genes exsA, exsD and exsC, are found at the terminus of the T3SS-1 cluster, and are regulators of T3SS-1 genes (84, 86). They showed that the environmental regulation of T3SS-1 genes is through ExsA, which acts as a positive transcriptional regulator of the T3SS-1 gene cluster. Call et al. found that under T3SS-1 non-inducing conditions (growth in LB, NaCl), ExsD binds to ExsA and prevents the transcriptional activity of this protein. However, under inducing conditions (cell culture media), ExsC is expressed and binds to ExsD. This prevents ExsD from binding ExsA and thus allows for the transcription of the T3SS-1 genes (84). Kodama and colleagues also demonstrated the regulation of T3SS-1 genes by ExsA and proposed that H-NS is a negative regulator of exsA (35). To add to this regulatory pathway, we have demonstrated that ToxRS and LeuO act as upstream regulators of T3SS-1 expression.

The streptomycin treatment of animals prior to their orogastric infection resulted in the recovery of substantial numbers of V. parahaemolyticus from gastrointestinal tissues 2 days postinfection and high levels of fecal shedding starting day 1 postinfection, whereas no V. parahaemolyticus was detectable in the tissues or stools from similarly infected PBS-treated animals. Vibrio parahaemolyticus was typically found throughout the gastrointestinal tracts of treated mice; however, higher bacterial burdens were observed in the cecum and large intestines. Following the establishment of colonization, the organism was maintained within intestinal tissues, as fecal pellets shed by infected animals contained high levels of V. parahaemolyticus for greater than 1 week postinfection. Interestingly, our histological assessment of tissues of the large intestine 7 days postinfection indicated that there were no overt signs of pathology, such as the disruption of colonic crypt structure or epithelial or brush border degradation (data not shown).

The use of murine models to assess V. parahaemolyticus pathogenesis has been frequently reported in the literature (3, 21, 22, 26, 51, 78, 81–83). Several of these studies utilized intraperitoneal inoculation for bacterial delivery, an infectious route that mimics septicemic infection, to determine and compare the 50% lethal dose (LD₅₀) of strains. In some cases, the LD₅₀ of strains following the orogastric inoculation of untreated animals was determined (3, 22, 78). However, our initial attempts to reproduce these orogastric challenge experiments for the characterization of mutant strains of V. parahaemolyticus RIMD2210633 failed to elicit mortality or observable morbidity in animals at any time postinfection. The examination of several parameters, including the use of either inbred (C57BL/6 and BALB/c) or outbred (CD-1) animals as well as infant mice, the type of bacterial growth media (LB versus brain heart infusion broth), or the growth phase of the inoculum (log-phase versus stationary-phase cultures), still did not result in the productive infection of animals (data not shown). Interestingly, a recent study by Call’s group suggests that previously reported orogastric rabbit and mouse models that produced...
septicemia are inadvertent pulmonary models (65). This study also described two novel animal models, an orogastric piglet model and an intrapulmonary inoculated mouse model, to examine the role of the two T3SSs in *V. parahaemolyticus* pathogenesis (65), and although useful models, neither addresses the most common route of infection. To our knowledge, the colonization model system we report here is the first allowing for the carriage of *V. parahaemolyticus* in the gastrointestinal tract of adult mice after oral inoculation.

Conventionally reared mice are resistant to gastrointestinal infection following oral challenge with a number of enteric pathogens, and therefore the pretreatment of animals with antibiotics to initiate the development of a productive infection has frequently been utilized to overcome the inability to infect adult animals (18, 70, 72, 77). The orogastric administration of streptomycin to mice has been shown to alter many of the normal physiological parameters of the intestine, rapidly resulting in shifts in the microbial flora, luminal pH, and the types of short-chain volatile fatty acids present, which, in combination, presumably play a role in removing barriers that might prohibit pathogen colonization (18, 70, 72, 77). For example, the antibiotic pretreatment of mice has been shown to result in a rapid increase inecal weights owing to fluid accumulation, with an obvious enlargement of tissue upon gross morphological examination (4, 71). Two separate studies have recently described models in which adult mice were orogastrically infected by *V. cholerae*, resulting in productive infections with intestinal colonization lasting for several days (55, 57–59). In these studies, the method by which mice were infected differs substantially from the protocol described here, with one requiring the use of ketamine/xylazine and the other isoflurane (55, 57–59). In the system described here, anesthesia was not utilized.

In summary, our aim was to characterize the ToxRS system in *V. parahaemolyticus* and to develop a murine model allowing for the colonization of adult mice following orogastric infection with *V. parahaemolyticus*, which is the typical route for bacterial entry into the host. We demonstrated that ToxRS is an important regulator of the stress response protein OmpU, and that this regulation is important for survival in a number of stress conditions commonly encountered in the human host. We have demonstrated that the ToxRS system also plays a role in the regulation of the T3SS-1 through its regulation of VP0350, a transcription regulator homologous to LeuO. This branch of the ToxRS regulon is thus far unique to *V. parahaemolyticus*. To our knowledge, the orogastric murine model presented in this study is the first for this particular species of *Vibrio*, and it provides a valuable new tool for studies to begin dissecting both the bacterial and host parameters that are crucial for the bacterial colonization of the gastrointestinal tract. Additionally, we have generated a β-galactosidase-expressing strain of *V. parahaemolyticus*, which for the first time will allow the coinfection of animal models to directly compare the effects of gene deletion in vivo. Through the use of this model, we have demonstrated that *V. parahaemolyticus* ToxRS and OmpU are important mediators of colonization. In summary, our data show a practical application for this new streptomycin-treated murine model for orogastric colonization by *V. parahaemolyticus*.

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