

# Virulence and stress-related periplasmic protein (VisP) in bacterial/host associations

Cristiano G. Moreira<sup>a,b</sup>, Carmen M. Herrera<sup>c,1</sup>, Brittany D. Needham<sup>c,1</sup>, Christopher T. Parker<sup>a,b</sup>, Stephen J. Libby<sup>d,e</sup>, Ferric C. Fang<sup>d,e</sup>, M. Stephen Trent<sup>c</sup>, and Vanessa Sperandio<sup>a,b,2</sup>

Departments of <sup>a</sup>Microbiology and <sup>b</sup>Biochemistry, University of Texas Southwestern Medical Center, Dallas, TX 75390-9048; <sup>c</sup>Molecular Genetics and Microbiology, University of Texas, Austin, TX 78712-1191; and Departments of <sup>d</sup>Laboratory Medicine and <sup>e</sup>Microbiology, University of Washington, Seattle, WA 98195-7735

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Gram-negative bacteria have an outer membrane containing LPS. LPS is constituted of an oligosaccharide portion and a lipid-A moiety that embeds this molecule within the outer membrane. LPS is a pathogen-associated molecular pattern, and several pathogens modify their lipid-A as a stealth strategy to avoid recognition by the innate immune system and gain resistance to host factors that disrupt the bacterial cell envelope. An essential feature of *Salmonella enterica* Typhimurium pathogenesis is its ability to replicate within vacuoles in professional macrophages. *S. Typhimurium* modifies its lipid-A by hydroxylation by the Fe2+/ $\alpha$ -ketoglutarate-dependent dioxygenase enzyme (LpxO). Here, we show that a periplasmic protein of the bacterial oligonucleotide/oligosaccharide-binding fold family, herein named virulence and stress-related periplasmic protein (VisP), on binding to the sugar moiety of peptidoglycan interacts with LpxO. This interaction inhibits LpxO function, leading to decreased LpxO-dependent lipid-A modifications and increasing resistance to stressors within the vacuole environment during intramacrophage replication promoting systemic disease. Consequently,  $\Delta visP$  is avirulent in systemic murine infections, where VisP acts through LpxO. Several Gram-negative pathogens harbor both VisP and LpxO, suggesting that this VisP-LpxO mechanism of lipid-A modifications has broader implications in bacterial pathogenesis. Bacterial species devoid of LpxO (e.g., *Escherichia coli*) have no lipid-A phenotypes associated with the lack of VisP; however, VisP also controls LpxO-independent phenotypes. VisP and LpxO act independently in the *S. Typhimurium* murine colitis model, with both mutants being attenuated for diverging reasons;  $\Delta visP$  is less resistant to cationic antimicrobial peptides, whereas  $\Delta lpxO$  is deficient for epithelial cell invasion. VisP converges bacterial cell wall homeostasis, stress responses, and pathogenicity.

QseC | chemical signaling

Bacterial–host relationships lead to different outcomes, ranging from beneficial to pathogenic interactions. Bacterial cells search for energy sources and niches for colonization within the host using either mutually beneficial or aggressive strategies that lead to pathogenesis. The host keeps these relationships balanced through the immune system, which is constantly surveying these interactions with microbes. *Salmonella* is a human pathogen responsible for food poisoning and typhoid fever (1). The pathogenesis of *S. Typhimurium* is a complex process. An essential feature of *S. Typhimurium* pathogenesis is its ability to replicate within vacuoles in macrophages (2).

During its evolution, *S. Typhimurium* acquired many pathogenicity islands (3). The main islands involved in *S. Typhimurium* infection are *Salmonella* pathogenicity island 1 (SPI-1) and SPI-2, both of which encode a type three secretion system (T3SS), a syringe-like apparatus used by the bacteria to inject effectors into the host cell. Both of these T3SSs are essential for *S. Typhimurium* virulence (4–8). The SPI-1 T3SS is required for invasion of the intestinal epithelium (4), whereas the SPI-2 T3SS is essential for *S. Typhimurium* replication and survival within macrophages and systemic murine infection (8–10). *S. Typhimurium* is a Gram-negative bacterium, and in addition to the expression of specialized

apparatus such as T3SS, it also uses additional strategies to promote host infection (11). Gram-negative bacteria have LPS in their outer membrane. Although LPS is a pathogen-associated molecular pattern recognized by toll-like receptor 4 (TLR4) (12), several pathogens modify its lipid-A domain to either avoid recognition by the innate immune system and/or gain resistance to host factors that compromise the integrity of the bacterial cell envelope (11). *S. Typhimurium* expresses machinery to modify its lipid-A by addition or removal of acyl chains, addition of phosphoethanolamine or amino-arabinose, and hydroxylation by the Fe2+/ $\alpha$ -ketoglutarate-dependent dioxygenase enzyme (LpxO) (11).

Bacterial pathogens rely on tightly regulated and complex signaling systems to express virulence traits. Virulence gene expression is an expensive process and should only occur within a suitable host. Hence, pathogens evolved to sense diverse environmental cues to promote host colonization (13). Such cues include the hormones epinephrine and norepinephrine that play a central role in stress responses in mammals (14). Stress responses have a profound effect on an organism's physiology and survival, and many pathogens, including *S. Typhimurium*, use the Quorum-sensing *Escherichia coli* (QseC) adrenergic sensor to activate virulence (15, 16). *S. Typhimurium* relies on QseC to cause systemic disease and activates its virulence repertoire when replicating in the spleen and liver (16).

Here, we show that a gene within the QseC regulon in *S. Typhimurium* encodes a periplasmic protein, renamed virulence and stress-related periplasmic protein (VisP), which binds to peptidoglycan and interacts with the LpxO–lipid-A–modifying enzyme inhibiting its function. This inhibition of LpxO-mediated lipid-A modifications is essential for resistance to intravacuolar stressors to promote intramacrophage replication and systemic disease. However, both VisP and LpxO have independent functions in colitis.

## Results

**VisP Is a Member of the QseC Regulon.** QseC plays an important role in *S. Typhimurium* pathogenesis by regulating transcription of SPI-1 and *sifA* (encodes an effector for the SPI-2 T3SS) genes (16). To gain a global view of the QseC regulon in *S. Typhimurium*, we performed transcriptomic studies comparing WT *S. Typhimurium* with a  $\Delta qseC$ . These studies, as well as a previous report (17), identified the gene *ygiW*, encoded upstream of the *qseBC* operon,

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<sup>1</sup>C.H. and B.N. contributed equally to this work.

<sup>2</sup>To whom correspondence should be addressed. E-mail: [vanessa.sperandio@utsouthwestern.edu](mailto:vanessa.sperandio@utsouthwestern.edu).

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as a member of the QseC regulon (Fig. S1). YgiW (protein ygiW precursor *E. coli*) as cryptic gene function encodes a periplasmic protein of the bacterial oligonucleotide/oligosaccharide-binding fold (BOF) family (Figs. S1–S3) (18). BOF proteins (cl01196) have five antiparallel  $\beta$ -strands forming a closed or partly opened barrel, and six conserved pairs of aspartic acid or glutamic/aspartic acid residues are located within the last three  $\beta$ -strands preceding the C terminus (Fig. S2). BOF proteins are predicted to bind oligosaccharides (18), and the crystal structure of the *Escherichia coli* YgiW has been solved (Protein Data Bank ID code 1NNX). YgiW is prevalent in several bacterial species (Table 1 and Fig. S2), and it is involved in antimicrobial resistance in *S. Typhimurium* (19). The  $\Delta ygiW$  ( $\Delta visP$ ) was more sensitive than WT to several stressors, such as cadmium chloride, acidic pH, and hydrogen peroxide (Fig. 1A–D). Additionally, the  $\Delta visP$  grows poorly under low magnesium ( $MgCl_2$ ) conditions (20  $\mu M$ ), but its growth is similar to WT under high  $MgCl_2$  concentrations (200  $\mu M$ ) (Fig. 1E). *S. Typhimurium* is a facultative intracellular pathogen, and its adaptation to acidic pH, low magnesium (20  $\mu M$ ), and reactive oxygen species vacuolar conditions is essential for intramacrophage survival (2). The  $\Delta visP$  presented a four orders of magnitude decrease in intramacrophage survival compared with WT (Fig. 1F), congruent with its increased sensibility to vacuolar stressors. Because YgiW plays a role in stress resistance and virulence in *S. Typhimurium*, we renamed this protein VisP.

**VisP and Lipid-A Modifications.** LPS modification is a common strategy used by many Gram-negative pathogens to survive stress responses (20). Of note, the LpxO enzyme is absent in *E. coli*, which explains the observations that an enterohemorrhagic *E. coli* (EHEC)  $\Delta visP$  is indistinguishable from WT concerning lipid-A modifications (Fig. S4), stress resistance (Fig. S4), and virulence [ability to express virulence genes and form attaching and effacing lesions on epithelial cells, a signature of EHEC gastrointestinal (GI) infection in mammals] (Fig. S4) phenotypes. *S. Typhimurium* expresses machinery to modify its lipid-A by addition or removal of several chemical groups, including hydroxylation by the LpxO enzyme (11). The  $\Delta visP$  presented significant differences in lipid-A modifications (Fig. 1H). Detailed analysis by MS (Fig. S5) identified that the differences in the  $\Delta visP$  were LpxO-mediated (Fig. 1G and H and Fig. S5) (21), specifically an S-2-hydroxymyristate moiety dioxygenase modification, known to be an

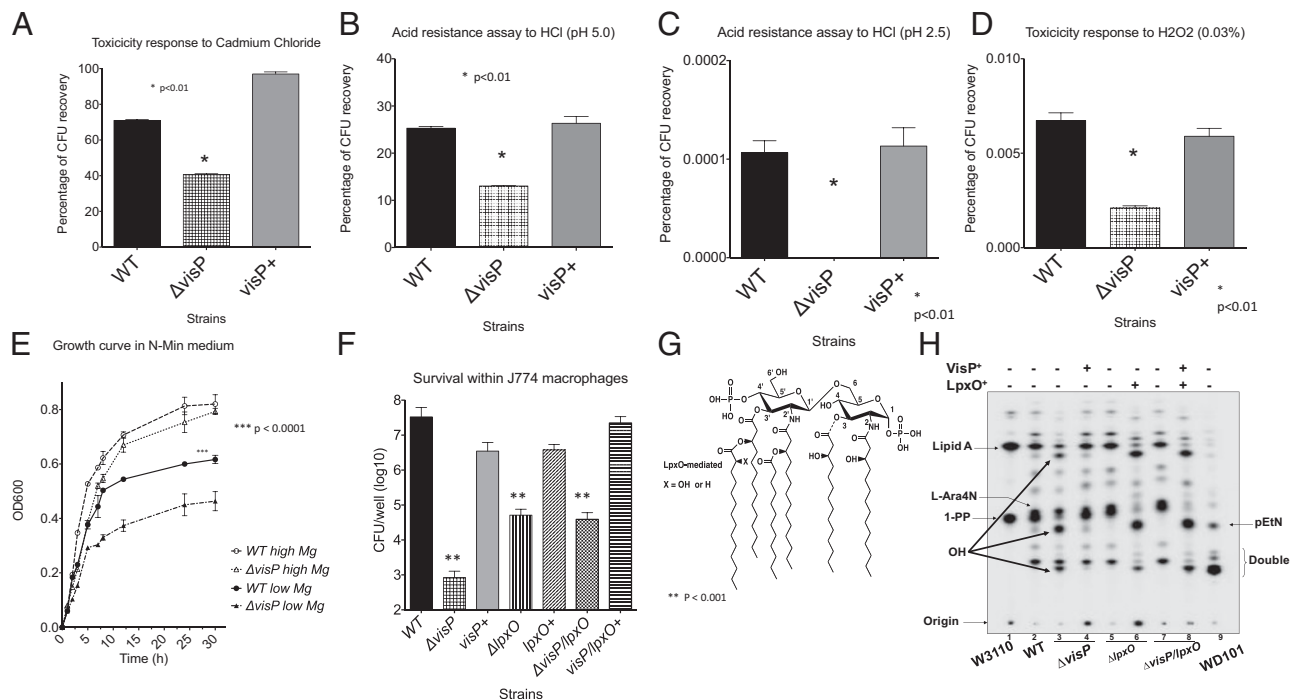
$Fe^{2+}/\alpha$ -ketoglutarate-dependent dioxygenase homolog (21). These data indicate that VisP inhibits LpxO. To better elucidate the VisP/LpxO relationship, we generated  $\Delta lpxO$  and  $\Delta lpxO/visP$  strains. S-2-hydroxymyristate moiety dioxygenase modification to lipid-A was observed in the  $\Delta visP$  by the characteristically lower migration in separation by TLC (Fig. 1H) as well as a mass increase of  $\sim 16$   $m/z$  units to the lipid-A analyzed by MS (Fig. S5). These effects were absent both in  $\Delta lpxO$  and  $\Delta lpxO/visP$  and restored on complementation (Fig. 1H and Fig. S4). Intramacrophage survival was also significantly decreased two orders of magnitude in  $\Delta lpxO$  and  $\Delta lpxO/visP$ , although not as drastically as in  $\Delta visP$  (Fig. 1F). VisP is a periplasmic protein, and LpxO is an inner membrane protein. Far Western blots using VisP-His and LpxO-Flag tagged proteins show that these proteins interact with each other but not with the QseB protein that was used as a negative control (Fig. 2A–C). The interaction between VisP and LpxO was also independently confirmed through coimmunoprecipitation (Fig. S6B). Hence, these results suggest that VisP acts on LpxO by directly interacting with LpxO to modulate its function (Fig. 2A–C and Fig. S6B).

The crystal structure of the *E. coli* VisP is solved (Protein Data Bank ID code 1NNX), and the modeled *S. Typhimurium* VisP structure suggests that this protein is conserved in these two organisms and that the six amino acid residues (D88, D90, E110, D112, E119, and D121) of the BOF binding pocket are conserved between them (Fig. 2D). Because these six residues in the binding pocket are involved in ligand binding, which can modulate the function of these proteins (18), we performed site direct mutagenesis of these six residues and showed that E110 is critical for the role of VisP in intramacrophage replication (Fig. 2E). Of note, expression and stability of all six VisP point mutants are indistinguishable from the WT protein (Fig. S6C). The VisP<sub>E110A</sub> mutant is unable to rescue intracellular replication in  $\Delta visP$ , and the VisP<sub>D90A</sub> rescues this phenotype, albeit to a lesser extent than the other four mutants, where complementation restores intramacrophage replication to WT levels (Fig. 2E). BOF proteins bind to oligosaccharides, and a major source of oligosaccharides in the periplasmic space is peptidoglycan. Peptidoglycan is composed of repeating units of *N*-acetyl-glucosamine (NAG) and *N*-acetyl-muramic acid (NAM). In silico analysis of the modeled VisP docking to NAG and NAM (Fig. 3A and Dataset S1) ([www.dockingserver.com](http://www.dockingserver.com)) shows that binding of these sugars, specially for NAM, is near to the predicted binding pocket. The binding of VisP for carbohydrates was also assayed using insoluble peptidoglycan (PGN) and cellulose (Fig. 3B). VisP binding was evaluated by pull-down assays to these insoluble fractions. The VisP<sub>WT</sub> bound better to PGN than VisP<sub>E110A</sub>, whereas VisP<sub>E110A</sub> bound better to cellulose (Fig. 3B). These data show that VisP binds to peptidoglycan in the periplasmic space. Importantly, this binding seems to alter VisP function, given that the VisP<sub>E110A</sub> mutant, which has its ability to bind to PGN decreased, no longer interacts with LpxO (Fig. 2A–C), and is defective for intramacrophage replication (Fig. 2D). Availability of peptidoglycan sugars in the periplasmic space enhances during cell wall remodeling, which occurs during environmental stress conditions, as well as during expression of T3SSs, which use peptidoglycan hydrolases during their assembly (22). Of note, T3SSs are used by *S. Typhimurium* to survive within macrophages (23).

**VisP in *S. Typhimurium* Murine Systemic Disease.** An in vivo systemic infection model (12) in BALB/c mice, (through an i.p. infection route) was used to assess virulence of VisP and LpxO individually and coupled. The  $\Delta visP$  mutant was avirulent in the systemic model, given that all animals survived infection by  $\Delta visP$ , whereas infection with WT led to 100% mortality by day 2 postinfection (Fig. 4A). Additionally,  $\Delta visP$  infected animals presented a decrease of four orders of magnitude in bacterial loads in spleens and livers compared with WT infected animals (Fig. 4B). The  $\Delta lpxO$  and double  $\Delta lpxO/visP$  were also attenuated, albeit to a lesser extent in these animal infections, compared with  $\Delta visP$  (Fig. 4). In the systemic model, VisP is dispensable for virulence if LpxO is absent, which is consistent with a negative regulatory role of VisP on LpxO

**Table 1. Distribution of VisP and LpxO among different bacterial species**

Bacterial species	VisP	LpxO
<i>Erwinia spp</i>	+	+
<i>Pectobacteria spp</i>	+	–
<i>Pantoea spp</i>	+	+
<i>Enterobacter spp</i>	+	+
<i>Serratia spp</i>	+	+
<i>Brucella spp</i>	–	+
<i>Ralstonia spp</i>	–	+
<i>Salmonella spp</i>	+	+
<i>E. coli</i>	+	–
<i>Yersinia spp</i>	+	+
<i>Pseudomonas spp</i>	+	+
<i>Klebsiella pneumoniae</i>	+	+
<i>Burkholderia spp</i>	–	+
<i>Bordetella spp</i>	–	+
<i>Acinetobacter spp</i>	–	+
<i>Xylella spp</i>	–	+
<i>Haemophilus influenzae</i>	+	–
<i>Pasteurella multocida</i>	+	–
<i>Shigella spp</i>	+	–
<i>Citrobacter spp</i>	+	–
<i>Hafnia alvei</i>	+	–



**Fig. 1.** VisP promotes resistance to stressors, virulence, and cell envelope remodeling in bacteria. (A) *S. Typhimurium* WT,  $\Delta visP$ , and complemented strains survival of CdCl<sub>2</sub> (4 mg mL<sup>-1</sup>). (B) Survival of HCl (pH 5.0) (C) Survival of HCl (pH 2.5). (D) Survival of H<sub>2</sub>O<sub>2</sub> (34 nmol L<sup>-1</sup>). (E) Bacterial growth under MgCl<sub>2</sub> low (20  $\mu$ M) and high (200  $\mu$ M) concentrations. (F) Intramacrophage replication of *S. Typhimurium* WT,  $\Delta visP$ ,  $\Delta lpxO$ ,  $\Delta lpxO/visP$ , and complemented strains (+) in J774 macrophage-like cells. (G) LpxO-mediated modifications on the lipid-A chemical structure. (H) TLC of <sup>32</sup>P-labeled lipid-A from *E. coli* K12 W3110 and WD101 (controls), *S. Typhimurium* WT,  $\Delta visP$ ,  $\Delta lpxO$ ,  $\Delta lpxO/visP$ , and complemented strains. Modifications assessed are aminoarabinose (L-Ara4N), 1-di-phosphate (1-PP), phosphoethanolamine (pEtN), double lipid-A modifications (double), and LpxO-mediated dioxigenase (OH). W3110 was the *E. coli* strain used as a control to visualize lipid-A (1,4 bisphosphorylated) and 1-PP species; WD101 was the *E. coli* strain used as phosphoethanolamine and lipid-A double controls. \**P* < 0.01, \*\**P* < 0.001.

function. These *in vivo* results are also congruent with the ability of these strains to replicate within macrophages, where  $\Delta visP$  replication is strikingly decreased, whereas  $\Delta lpxO$  and the double mutant, although also decreased for intracellular survival compared with WT, are not as attenuated as  $\Delta visP$  (Fig. 1F). The  $\Delta lpxO$  and the double mutant intermediary attenuation, compared with  $\Delta visP$ , suggests that absence of LpxO–lipid-A modifications hampers pathogenesis, but the complete deregulation of this phenotype (increased modification of lipid-A) in the absence of VisP has even more profound effects in pathogenesis. It has been proposed that LpxO–lipid-A modifications enhance *S. Typhimurium* membrane permeability, although a  $\Delta lpxO$  presented permeability similar to WT (24), and it was also as resistant to detergent treatments as WT (Fig. S64), suggesting that loss of LpxO does not have a profound effect in membrane permeability. Our data suggest that, although 2-hydroxymyristilation of lipid-A during intramacrophage replication leads to optimal resistance to intramacrophage stressors, enhancement of this modification in the absence of VisP inhibition of LpxO activity is deleterious to survival within these cells. These data suggest that VisP/LpxO-dependent remodeling of the bacterial cell wall within macrophages is an important feature of *S. Typhimurium* systemic infection (Fig. 4).

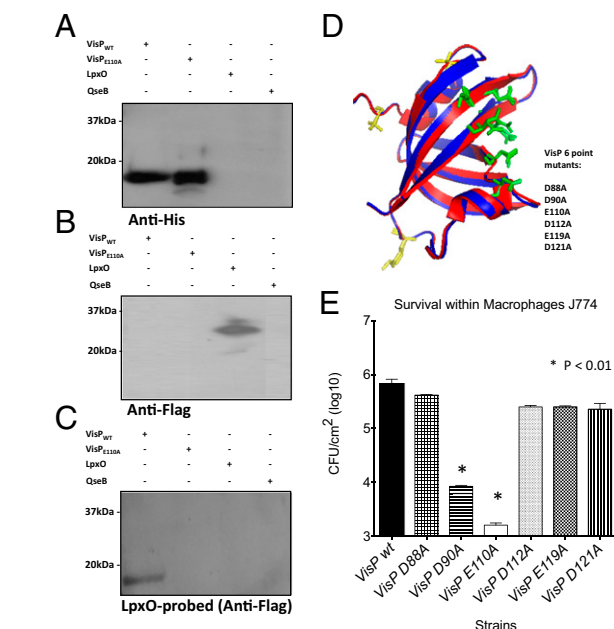
**LpxO-Independent VisP Phenotypes.** Because *S. Typhimurium* also causes gastroenteritis, we assessed whether VisP and/or LpxO played any role in the *S. Typhimurium* colitis model (oral inoculation of BALB/c mice pretreated with streptomycin) (25). In contrast to the systemic murine infections, in the colitis model, all three mutants ( $\Delta visP$ ,  $\Delta lpxO$ , and  $\Delta lpxO/visP$ ) were attenuated three to four orders of magnitude compared with WT (Fig. 5A). These data indicate that, although both VisP and LpxO have important roles in *S. Typhimurium* infection of the GI tract, they may play independent roles in this host compartment. During GI

infection, *S. Typhimurium* encounters cationic antimicrobial peptides, which are an important host defense in the GI tract (2). The *visP* mutant is more susceptible to cationic peptides (e.g., polymyxin B) (Fig. 5B) (19), whereas  $\Delta lpxO$  presented similar susceptibility to WT (Fig. 5B). The double  $\Delta lpxO/visP$  mutant also presented reduced resistance to polymyxin B, further supporting the hypothesis that VisP also performs LpxO-independent functions. The  $\Delta visP$ -increased susceptibility to polymyxin B could be rescued on complementation with WT VisP but not with the VisP<sub>E110A</sub> mutant, suggesting that this mutation in the VisP ligand binding pocket also affects LpxO-independent phenotypes (Fig. 5B). Although the increased susceptibility to cationic antimicrobial peptides could substantiate the attenuation of  $\Delta visP$  in the colitis model, it cannot explain the attenuation of  $\Delta lpxO$ . However, another important facet of *S. Typhimurium* GI infection relies in the ability of this pathogen to invade intestinal epithelial cells (2). The  $\Delta visP$  mutant invades epithelial cells at similar levels to WT, whereas both  $\Delta lpxO$  and  $\Delta lpxO/visP$  have decreased ability (four to five orders of magnitude) to invade epithelial cells compared with WT (Fig. 5C). These data suggest that VisP is important for antimicrobial peptide resistance, whereas LpxO is required for efficient epithelial cell invasion by *S. Typhimurium* in the GI compartment.

## Discussion

BOF proteins are present in a number of Gram-negative pathogenic bacterial strains and have unknown function. The association of genes encoding BOF proteins with prophage inserts and virulence plasmids suggests that these genes were inherited horizontally through mobile genetic elements, providing a mechanism for rapid evolution and adaptation to different niches. The BOF family lacks highly conserved amino acid residues involved in nucleotide binding, suggesting that these proteins can bind oligosaccharides but not nucleotides (18). All BOF proteins also harbor





**Fig. 2.** VisP-LpxO interactions. (A) Control far Western blot of immobilized LpxO-Flag, VisP<sub>WT</sub>-His, VisP<sub>E110A</sub>-His, and QseB (untagged) probed with  $\Delta visP\Delta lpxO$  whole-cell lysate and then probed with anti-His antiserum. (B) Control far Western blot of immobilized LpxO-Flag, VisP<sub>WT</sub>-His, VisP<sub>E110A</sub>-His, and QseB (untagged) probed with  $\Delta visP\Delta lpxO$  whole-cell lysate and then probed with anti-Flag antiserum. (C) Far Western blot of immobilized LpxO-Flag, VisP<sub>WT</sub>-His, VisP<sub>E110A</sub>-His, and QseB (untagged) probed with  $\Delta visP\Delta lpxO$  whole-cell lysate and then probed with anti-Flag antiserum. (D) Superposition of VisP modeled *S. Typhimurium* structure to the *E. coli* VisP structure (1NNX). *E. coli* structure is represented in red, *S. Typhimurium* is blue, three modified residues/structures are shown in yellow, and six site-directed point mutants are in green: VisP<sub>D88A</sub>, VisP<sub>D90A</sub>, VisP<sub>E110A</sub>, VisP<sub>D112A</sub>, VisP<sub>E119A</sub>, and VisP<sub>D121A</sub>. (E) Intramacrophage replication of  $\Delta visP$  complemented with WT VisP or the VisP point mutants in J774 macrophage-like cells. \**P* < 0.01.

sec-dependent secretion signal sequences and are predicted to be periplasmic proteins (18). An important source of free saccharides within the periplasmic space, especially within stress environments or expression of T3SSs that induces remodeling of the bacterial cell wall, is peptidoglycan. Here, we have described that a previously uncharacterized BOF protein YgiW, herein renamed VisP, binds to the sugar moiety of peptidoglycan (Fig. 3). All BOF proteins have very conserved amino acid residues within their ligand binding pockets (18), and disruption of one of these residues (E110), although not altering protein expression/stability, alters its binding to peptidoglycan and prevents its function (Figs. 2, 3, and 5).

VisP binds to the inner membrane protein LpxO (Fig. 2), whose function is to promote an S-2-hydroxymyristate moiety dioxygenase modification on lipid-A (21). The GI pathogen *S. Typhimurium* harbors LpxO, and in addition to LpxO lipid-A modifications, it is also known to use a variety of other modifications to promote adaptations to different environments (11). Although LpxO-mediated lipid-A modifications have been exquisitely defined biochemically (21, 26), the biological function of this modification has remained a mystery. Here, we show that controlling the LpxO-lipid-A modifications is key for *S. Typhimurium* pathogenesis. VisP and LpxO mediate resistance to stressors and virulence in *S. Typhimurium* (Figs. 1, 2, 3, 4, and 5). VisP binds to PG. Decreased peptidoglycan (PG) binding alters the ability of VisP to interact with LpxO and LpxO function (Figs. 2 and 3), suggesting that VisP PG binding plays a role in decreasing LpxO-dependent lipid-A modifications to increase resistance to stressors within the vacuole environment during *S. Typhimurium* intramacrophage replication promoting systemic disease (Figs. 1, 2, 3, and 4).

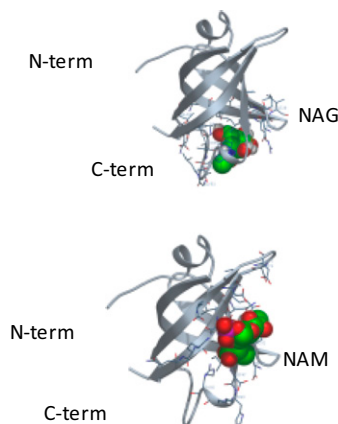
However, VisP also has LpxO-independent functions (Fig. 5), suggesting that it could either have self-sufficient activities or engage with other protein partners. The distribution of VisP and/or LpxO varies among different Gram-negative species, with them either occurring together or by themselves (Table 1). Hence, these two proteins could perform synergistic or individual functions in niche/host adaptations, suggesting that they may have broader implications in bacterial pathogenesis. VisP and LpxO relationships converge bacterial cell wall homeostasis, stress responses, and pathogenicity. Importantly, expression of *visP* itself is regulated by host stress molecules (epinephrine and norepinephrine) sensed by the QseC bacterial adrenergic receptor (Fig. S1), further linking bacterial/host stress in modulation of pathogenesis. This study highlights the coevolution and the fundamental relationship between mammals and microbes, and it highlights that these signaling systems fully integrate bacterial and mammalian cell behavior.

## Materials and Methods

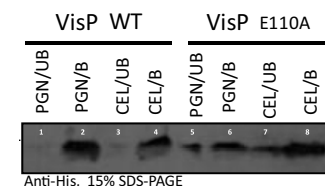
**Strains and Plasmids.** All strains and plasmids used in this study are listed on Table S1. Recombinant DNA and molecular biology techniques were performed as previously described (27). All oligonucleotides used are listed on Table S2.

**Construction of the Isogenic Mutants.** Construction of isogenic nonpolar *S. Typhimurium* SL1344 *visP* and *lpxO* single mutants, *visP/lpxO* double KO, and

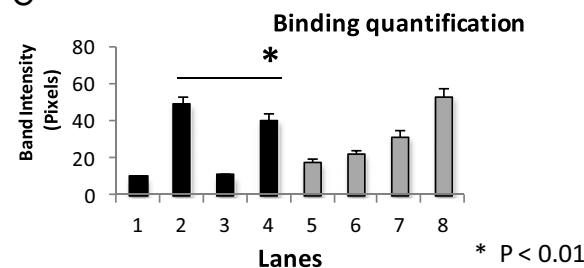
## A VisP docking



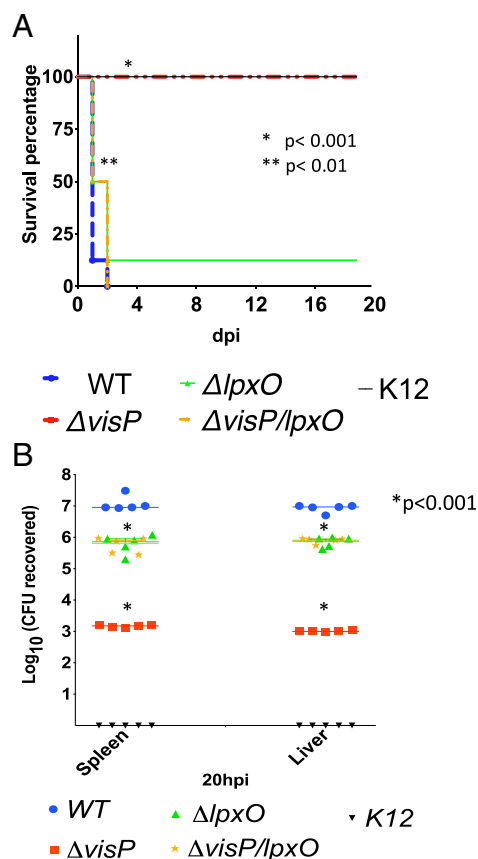
## B



## C



**Fig. 3.** VisP PG binding. (A) Docking modeling of *S. Typhimurium* VisP molecule against NAG and NAM. (B) Pull-down binding assays of purified VisP<sub>WT</sub> and VisP<sub>E110A</sub> with insoluble PGN or cellulose (CEL). B, bound fraction; UB, unbound fraction. (C) Quantification of pull-down binding assays (performed in triplicates). \**P* < 0.01.



**Fig. 4.** ViSP and LpxO in *S. Typhimurium* systemic murine infection. (A) Survival curves of mice infected i.p. with *S. Typhimurium* [days postinfection (dpi)]. (B) Organ loads of mice infected i.p. with *S. Typhimurium* 20 h post-infection (hpi). \* $P < 0.001$ , \*\* $P < 0.01$ .

EHEC  $\Delta visP$  (CP171) was achieved using  $\lambda$ -red mutagenesis (28). The *visP* mutant (CGM300) was complemented with the *visP* gene cloned into pBAD-MycHisA (Invitrogen) (20 copies per cell) vector, generating strain CGM301. The *lpxO* mutant (CGM302) was complemented with the *lpxO* gene cloned into pBAD33 vector (29) (15 copies per cell), generating strain CGM303. The *visP/lpxO* mutant (CGM304) was double complemented with pBADMycHisA and pBAD33c constructions, generating strain CGM305.

**Microarray Analysis.** The microarray study compared the *qseC* mutant with WT in LB broth. The *Salmonella* arrays were performed as previously described (30). The Gene Expression Omnibus database accession number for the microarray results is GSE38353.

**Quantitative Real-Time RT-PCR.** Quantitative RT-PCRs were performed as previously described (31).

**Macrophage Infection.** Intramacrophage replication of *S. Typhimurium* in J774 murine macrophage-like cells was performed as previously reported (32–34).

**HeLa Invasion and Adhesion Assays.** Invasion of HeLa cells was performed as previously reported (9, 24, 33–35).

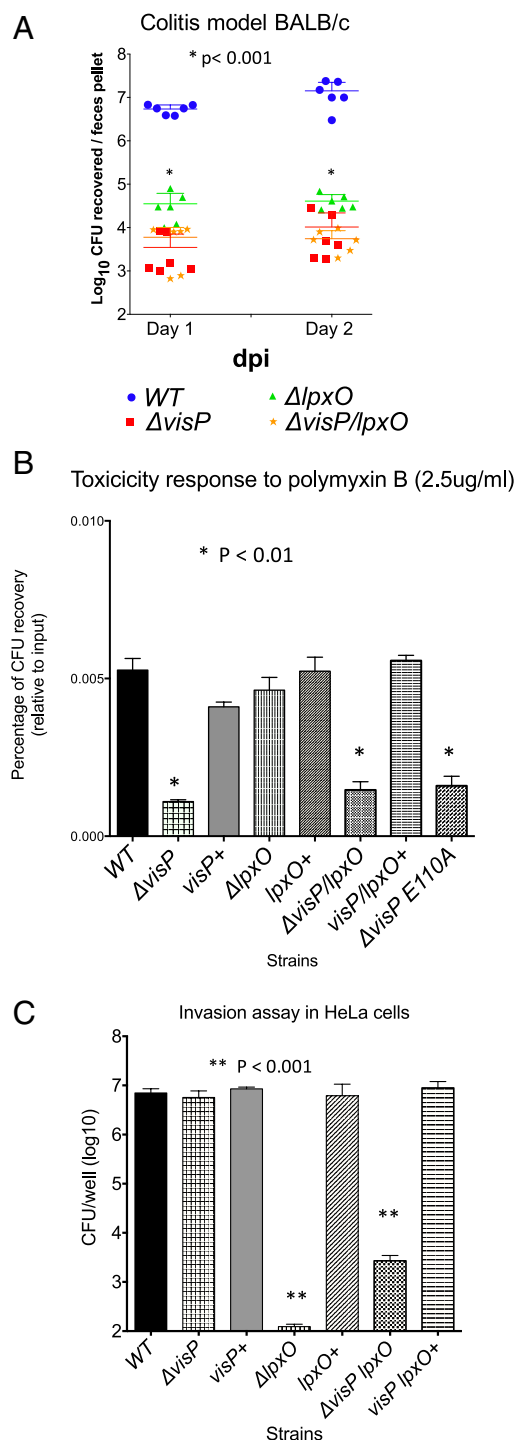
**Docking Assays.** Docking modeling was performed according to [www.dockingserver.com](http://www.dockingserver.com) instructions (36–39).

**Structure Prediction of *Salmonella* ViSP.** A structure for the BOF-containing fragment of *Salmonella* ViSP was predicted using MODELER v9.10 (40, 41).

**Protein Purification.** Performed according to PeriPreps kit manufacturer's instructions (EPICENTRE Biotechnologies). The His-tagged and Flag-tagged protein fractions were isolated as previously described (27), whereas the

ANTI-FLAG M2 affinity gel (Sigma) was employed according to manufacturer's directions.

**Far Western.** Same equimolar amounts of purified His-tagged protein were separated on an SDS gel and transferred to membranes. Replicate purified fractions were then probed with whole-cell lysates of the  $\Delta visP/lpxO$  double mutant or the double mutant overexpressing either Flag-tagged LpxO or His-tagged ViSP. As another (negative) control, a replicate membrane was left unprobed by the whole-cell lysate. All



**Fig. 5.** ViSP and LpxO in *S. Typhimurium* murine colitis. (A) CFUs of *S. Typhimurium* in feces days 1 and 2 postinfection (dpi). \* $P < 0.001$ . (B) Resistance to polymyxin B. \* $P < 0.01$ . (C) Invasion of HeLa cells. \*\* $P < 0.001$ .

membranes were then probed with either anti-His or anti-Flag primary antibodies and then incubated with a secondary antibody. ECL reagent (GE) was added, and membranes were exposed to film to detect interacting proteins.

**PGN Binding Assays.** These binding assays were performed as previously described (42).

**Stress Response Assays.** The resistance assays were performed in the presence of hydrogen peroxide and cadmium chloride (survival test; adapted from previous studies) (43). The acid resistance assay was performed as previously reported (44).

**Polymyxin B Sensitivity Assay.** The sensitivity assays were performed as previously reported (32–34).

**Site Direct Mutagenesis.** VisP site-directed mutagenesis was performed using the QuikChange II Site Directed Mutagenesis kit (Stratagene) using the manufacturer's recommendations.

**Isolation of Labeled Lipid-A.** Lipid-A isolations were performed as previously reported (45).

**MS.** Lipid-A was isolated for analysis by MA as previously described (46). Lipid samples were analyzed using an MALDI-TOF/TOF (ABI 4700 Proteomics Analyzer) mass spectrometer in the negative ion linear mode as previously described (47).

**Fluorescent Actin Staining Assay.** Fluorescent actin staining was performed as previously described (48).

**Colitis Model and Systemic Infections with *S. Typhimurium*.** Mice (BALB/c, 7- to 9-wk old, female) were infected orally for the colitis model as previously described (49). The systemic infections were also performed with mice (BALB/c, 7- to 9-wk old, female) infected using an i.p. route as previously described (32).

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