A Dopamine-Responsive Signal Transduction Controls Transcription of *Salmonella enterica* Serovar Typhimurium Virulence Genes

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**ABSTRACT** We have shown that the ligand-responsive MarR family member SlyA plays an important role in transcription activation of multiple virulence genes in *Salmonella enterica* serovar Typhimurium by responding to guanosine tetraphosphate (ppGpp). Here, we demonstrate that another MarR family member, EmrR, is required for virulence of *S.* Typhimurium and another enteric bacterium, *Yersinia pestis*. EmrR is found to activate transcription of an array of virulence determinants, including *Salmonella* pathogenicity island 2 (SPI-2) genes and several divergent operons, which have been shown to be activated by SlyA and the PhoP/PhoQ two-component system. We studied the regulatory effect of EmrR on one of these genetic loci, i.e., the *pagC-pagD* divergent operon, and characterized a catecholamine neurotransmitter, dopamine, as an EmrR-sensed signal. Dopamine acts on EmrR to reduce its ability to bind to the target promoters, thus functioning as a negative signal to downregulate this EmrR-activated transcription. EmrR can bind to AT-rich sequences, which particularly overlap the SlyA and PhoP binding sites in the *pagC-pagD* divergent promoter. EmrR is a priming transcription regulator that binds its target promoters prior to successive transcription activators, by which it displaces universal silencer H-NS from these promoters and facilitates successive regulators to bind these regions. Regulation of the *Salmonella*-specific gene in *Escherichia coli* and *Y. pestis* reveals that EmrR-dependent regulation is conserved in enteric bacteria. These observations suggest that EmrR is a transcription activator to control the expression of virulence genes, including the SPI-2 genes. Dopamine can act on the EmrR-mediated signal transduction, thus downregulating expression of these virulence factors.

**IMPORTANCE** In this study, MarR family regulator EmrR is identified as a novel virulence factor of enteric bacteria, here exemplified by *Salmonella enterica* serovar Typhimurium and *Yersinia pestis*. EmrR exerts an essential effect as a transcription activator for expression of virulence determinants, including *Salmonella* pathogenicity island 2 genes and a set of horizontally acquired genetic loci that formed divergent operons. EmrR senses the neurotransmitter dopamine and is subsequently released from target promoters, resulting in downregulation of the virulence gene expression. Through this action on EmrR, dopamine can weaken *Salmonella* resistance against host defense mechanisms. This provides an explanation for the previous observation that dopamine inhibits bacterial infection in animal gastrointestinal tracts. Our findings provide evidence that this neurotransmitter can modulate bacterial gene expression through interaction with virulence regulator EmrR.

**KEYWORDS** bacterial signal transduction, catecholamine neurotransmitter dopamine, MarR family virulence regulator EmrR, PhoP/PhoQ and SlyA feedforward loop, *Salmonella* pathogenicity island 2 (SPI-2)

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MarR family regulators play critical roles in multidrug resistance, oxidative stresses, catabolism of environmental aromatic compounds, antimicrobial peptide resistance, and virulence (1–7). Their members display significant diversity, responding to different signaling molecules and recognizing diverse DNA targets. First characterized in *Escherichia coli*, EmrR is a MarR family regulator encoded by the first gene of the *emrRAB* operon. It functions as a transcriptional repressor to control the synthesis of EmrAB, which interacts with the TolC outer membrane porin to construct a tripartite multidrug pump (8–11). Originally referred to as MprA, EmrR regulates the plasmid-borne *mcb* operon for microcin B17 synthesis (12). EmrR downregulates the *emrRAB* operon by binding to an inverted repeat sequence centered near the −10 region of its promoter (13). Accordingly, disruption of the *emrR* gene led to increased expression of EmrAB and elevated resistance to the antibiotic thiolactomycin (9). The presence of structurally unrelated substrates, including carbonyl cyanide *m*-chlorophenylhydrazone, carbonyl cyanide *p*-(trifluoromethoxy)phenylhydrazone, and 2,4-dinitrophenol derepress *emrRAB* expression by binding to EmrR and releasing it from the operon promoter, allowing expression of EmrAB (14). The EmrAB pump can then export these molecules (14). Except for the *emrRAB* operon and plasmid-borne *mcb* loci, it remains largely elusive which genetic loci are regulated by EmrR in *E. coli* or other enteric bacteria.

Although work in the area of MarR family-related regulatory mechanisms has been focused on their functions in transcriptional repression, cumulative evidence suggests that some MarR family regulators play important roles in transcriptional activation. For example, ExpG from the nitrogen-fixing soil bacterium *Sinorhizobium meliloti* activates transcription of three *exp* operons involved in the production of galactoglucon, an exopolysaccharide required for this organism to infect plant roots for nodule formation (15–17). In *Salmonella enterica* serovar Typhimurium, the MarR family SlyA protein has been characterized as a virulence regulator contributing to bacterial infection in part by activating hemolysin and flagellum production (18–20). This characterization was further supported by the observations that a master virulence regulator, i.e., the PhoP/PhoQ two-component system, activated transcription of the *slyA* gene (5, 21). When *Salmonella* experiences Mg²⁺ deprivation conditions, the sensor kinase PhoQ mediates phosphorylation of the response regulator PhoP (for a review, see reference 22), facilitating PhoP binding the target promoters and regulating transcription of these genes, which includes their operon, *phoPQ* and *slyA* (5, 21). Actually, the PhoP protein acts not only as the transcriptional activator of the *slyA* gene but also as one of two transcriptional activators (the other activator being SlyA) for a subset of horizontally acquired gene clusters, including several divergent operons (5, 23). This regulatory circuit, generally designated the feedforward regulatory loop (24), enables bacteria to regulate transcription by integrating different signals, i.e., the environmental low Mg²⁺ sensed by the PhoP/PhoQ system (25) and the cytoplasmic alarmone (ppGpp) sensed by SlyA (23). The regulators PhoP and SlyA recognized the target promoters via the PhoP binding site, TAAAT-N₁₀-TAAAC (a PhoP box in the opposite strand) and the SlyA binding site, AATATT-N₁₀-ATTATT (the SlyA box), respectively, in which they antagonized a global transcriptional silencer H-NS occupying their AT-rich binding sites (5, 23, 26, 27). This working model demonstrated that a MarR family regulator could participate in a regulatory network by coordinating with other signal response regulators to control genetic loci contributing to bacterial pathogenesis.

For the first time, we have elucidated that EmrR, a MarR family regulator sharing sequence similarity with SlyA, governs a signal transduction pathway in response to neurotransmitter dopamine and essential for virulence of enteric bacteria including *S. Typhimurium* and *Yersinia pestis*. We found that EmrR was able to activate transcription of *Salmonella* pathogenicity island 2 (SPI-2) genes and a set of horizontally acquired loci and characterized dopamine as a ligand that could act on EmrR to repress the EmrR-dependent transcription. We revealed that EmrR interacted with the AT-rich regions at its target promoters, by which it displaced a global transcription repressor H-NS from the promoters and enhanced other activators to bind to their binding sites.
EmrR Senses Dopamine To Control Virulence

Therefore, our study illustrates a multifactorial regulatory cascade to control the expression of the S. Typhimurium virulence regulons in response to both positive and negative inputs.

RESULTS

MarR family member EmrR is a virulence regulator required for S. Typhimurium and Yersinia pestis infection. Since MarR family regulator SlyA had been demonstrated as a virulence factor for Salmonella infection in a mouse model (18), we investigated other members of the MarR family for their possible roles in bacterial virulence. A mouse virulence analysis was carried out by using single in-frame deletion mutations at the genes encoding other MarR family members that shared homology with SlyA. While all the BALB/c mice inoculated orally with 10^3 and 10^4 CFU of wild-type ATCC 14028s (14028) cells died within 11 days, 80% and 40% of the mice that received an isogenic ΔemrR mutant survived 30 days postinoculation, respectively (Fig. 1A). In contrast, all mice died within 10 days after intraperitoneal administration of 10^3 and 10^4 CFU of the wild-type strain and ΔemrR mutant (unpublished results), suggesting that EmrR should be specifically required for the oral route of bacterial infection. A Salmonella survival assay in murine J774.2 macrophages showed that the number of CFU of the ΔemrR mutant was reduced ~13-fold compared with that of the wild-type strain that survived within macrophages (Fig. 1B). The phenotype of the ΔemrR mutant was solely the result of a lack of the EmrR protein because the CFU in macrophages were recovered to a wild-type level in an ΔemrR mutant in which a FLAG-tagged emrR gene was expressed under the control of the P_lac promoter from plasmid pemrR-FLAG (Fig. 1B). A significant feature often associated with Salmonella virulence was the inducible resistance to antimicrobial peptides (AMPs), many of which were regulated through specific virulence regulators (28–33). We investigated whether EmrR contributed to AMP resistance by challenging Salmonella strains with a cationic antimicrobial peptide, polymyxin B, and found that the ΔemrR mutant displayed a susceptible phenotype to polymyxin B in a manner similar to that of the ΔslyA mutant (Fig. 1C). The emrR mutation could be complemented in trans because the survival rates of the ΔemrR mutant harboring pemrR-FLAG was restored to wild-type levels when challenged by polymyxin B at different concentrations (Fig. 1C). Furthermore, we demonstrated that EmrR was a virulence factor not only in Salmonella but also in Yersinia pestis by carrying out a mouse virulence assay described previously (34). When Swiss Webster mice were injected subcutaneously with a wild-type strain KIM6+(pCD1Ap) and its isogenic ΔemrR mutant y10064(pCD1Ap), the ΔemrR mutant was ~1,000 times less virulent than its isogenic wild-type strain (see Fig. S1 in the supplemental material).

EmrR is required for transcription activation of Salmonella pathogenicity island 2 genes and virulence-related divergent operons. As shown previously, the PhoP/PhoQ two-component system built a feedforward loop in association with SlyA to activate transcription of a polymyxin B resistance gene, ugtL, in a PhoP-activating condition (0.01 mM Mg^{2+}; referred to as low Mg^{2+}) (5, 23). Actually, EmrR was able to activate ugtL transcription in a low-Mg^{2+} condition in which the β-galactosidase activity in a strain harboring the chromosomal ugtL-lacZY fusion was 15.5-fold higher than that in its isogenic ΔemrR mutant (Fig. 2A). The deficient ugtL expression in the ΔemrR mutant was merely caused by an absence of this activator and could be rescued by an EmrR protein produced from plasmid pemrR-FLAG. The ugtL activation was specifically dependent on SlyA and EmrR, but not on other known MarR family regulators, since a deletion at the coding region of the marR, STM1100, STM1547, or STM2920 gene did not affect ugtL transcription (Fig. S2A). An RNA-Seq analysis was carried out to profile more Salmonella genetic loci regulated by EmrR under low-Mg^{2+} growth (Table S3). The RNA-Seq data were first validated by increased RNA levels of the emrA and emrB genes in the ΔemrR mutant (5.7- and 5.9-fold, respectively; Table S3), since transcription of these genes was shown to be repressed by EmrR previously (9). Strikingly, almost all genetic loci of Salmonella pathogenicity island 2 (SPI-2) were detected as EmrR-activated genes, since their mRNA levels were significantly higher in
EmrR is an *S. Typhimurium* virulence regulator in a mouse model and contributes to resistance to polymyxin B. (A) Survival of BALB/c mice (five mice in each group) administered orally with *Salmonella* wild-type (WT) strain 14028s (diamonds) and ΔemrR mutant (YS15776) (squares) at doses of $1 \times 10^5$ CFU (solid symbols) and $1 \times 10^6$ CFU (open symbols). WT versus ΔemrR ($1 \times 10^5$), $P < 0.01$; WT versus ΔemrR ($1 \times 10^6$), $P < 0.01$. (B) Intramacrophage growth levels of *Salmonella* wild-type strain 14028s (WT), ΔemrR mutant (YS15776), and ΔemrR mutant harboring complementing plasmid *pemrR-FLAG* (pYS2015) at a ratio of 10 bacteria per J774.2 cell (i.e., multiplicity of infection [MOI] of 10). Macrophages were mixed with bacterial cells, centrifuged, and then incubated for 20 min to permit phagocytosis. Macrophages were lysed after cultured in the medium containing dopamine at the indicated concentrations for 18 h and plated onto LB agar to determine the number of viable bacteria (average numbers of CFU of the intracellular bacteria for 2 h postinfection were $5.7 \times 10^4$ for WT, $5.1 \times 10^4$ for the ΔemrR mutant, and $6.3 \times 10^4$ for ΔemrR mutant plus *pemrR-FLAG*; see details in Materials and Methods). Values that are significantly different from the value for the WT group by *t* test are shown by asterisks as follows: ***, $P < 0.001$; **, $P < 0.01$; *, $P < 0.05$. (C) Survival rates of *Salmonella* wild-type strain 14028s (WT), ΔslyA mutant (YS11068), ΔemrR mutant (YS15776), and ΔemrR mutant harboring *pemrR-FLAG* after the strains were challenged by antimicrobial peptide polymyxin B. Bacterial cultures were supplemented with dopamine at the indicated concentration or not supplemented with dopamine. Values that are significantly different from the value for the strain not treated with polymyxin B by *t* test are indicated by asterisks as follows: ***, $P < 0.001$; **, $P < 0.01$; *, $P < 0.05$. Data in panels B and C are from three independent assays conducted in duplicate, and all values are means ± standard deviations.
EmrR activates transcription of SPI-2 genes and the PhoP- and SlyA-activated divergent operons. (A) β-Galactosidase activity from a chromosomal ugtL-lacZY fusion was determined in the S. Typhimurium wild-type (WT) strain (YS15550), ΔemrR mutant (YS15708), and ΔemrR mutant harboring complementing plasmid pemR-FLAG (pYS2015) grown in 0.01 mM Mg²⁺ for 4 h. The value for the ΔemrR mutant was significantly different (P < 0.01) by t test from the WT value as indicated (**). (B) RNA-Seq analysis shows downregulation of SPI-2 genes and horizontally acquired divergent operons in the ΔemrR mutant (YS15776) and ΔslyA mutant (YS11068) compared to the wild-type strain 14028s (WT) when grown in low Mg²⁺ for 4 h. (C) RT-PCR analysis for the mRNA levels of transcripts from four SPI-2 genes and also the pagC gene in the wild-type strain 14028s (WT), ΔemrR mutant (YS15776), and ΔslyA mutant (YS11068) grown in low Mg²⁺ with or without 1 mM dopamine and from the ΔemrR mutant (YS15776) and ΔslyA mutant (YS11068) grown in low Mg²⁺. The constitutively transcribed rpoD gene was used to indicate similar amounts of total RNAs. The DNA fragment was amplified using the primers listed in Table S2 and separated in an agarose gel. Dopamine was added to the required bacterial culture at a final concentration of 1 mM. (D) β-Galactosidase activity from a chromosomal ssaM-lacZY fusion was determined in the wild-type strain (YS18490), ΔemrR mutant (YS18520), and ΔemrR mutant harboring pemR-FLAG grown in low Mg²⁺ supplemented with 0.2 mM IPTG for 4 h. *, P < 0.05 versus WT value by t test. (E) β-Galactosidase activity from respective chromosomal pagC-lacZY and pagD-lacZY fusions was determined in the WT strains (YS11644 and YS12000), ΔemrR mutants (YS14827 and YS17260), ΔemrR mutants harboring pemR-FLAG, and ΔemrR Δhns double mutants (YS15468 and YS17261) cultured in low Mg²⁺ for 4 h. **, P < 0.01 versus WT blue by t test. Data in panels A, D, and E are from three independent assays conducted in duplicate, and all values are means plus standard deviations.
the wild-type strain than in the ΔemrR mutant (Table S3; also summarized in Fig. 2B). Consistent with the previous observation that SlyA activated transcription of SPI-2 genes (35), RNA-Seq analysis also showed that the mRNA levels of these SPI-2 genes in the wild-type strain were higher than those in the ΔslyA mutant (Table S4; also summarized in Fig. 2B). We further confirmed the RNA-Seq results by a reverse transcription-PCR (RT-PCR) analysis which revealed higher mRNA levels of ssaA, ssaB, sseB, and ssaM genes from different SPI-2 operons in the wild-type strain than in the ΔemrR and ΔslyA mutants, respectively (Fig. 2C) and with a β-galactosidase assay of reporter strains carrying a chromosomal ssaM-lacZ fusion in which the activity in the ΔemrR mutant was 8-fold lower than that in the wild-type strain (Fig. 2D). Consistently, deficient ssaM expression in the ΔemrR mutant could be restored by a heterologous emrR expression from plasmid pemrR-FLAG (Fig. 2D). It was shown that the PhoP/PhoQ and SlyA feedforward loop simultaneously activated transcription of a set of divergent operons, including ugtl-sifB and pagC-pagD (5, 22). Concomitantly, RNA-Seq analysis showed that mRNA levels of ugtl, sifB, pagC, and pagD in the ΔemrR mutant strain were significantly reduced compared to those in the wild-type strain (Table S3; also see Fig. 2B). The EmrR-dependent regulation of this class of genetic loci was further confirmed by a β-galactosidase assay of reporter strains carrying chromosomal pagC-lacZ and pagD-lacZ fusions in which activities in the ΔemrR mutants were 25.2- and 22.5-fold reduced, respectively, compared to those in their isogenic wild-type strains grown in low Mg²⁺ (Fig. 2E). Furthermore, the deficient expression of this divergent operon in the ΔemrR mutant could be restored by heterologous emrR expression from plasmid pemrR-FLAG (Fig. 2E). We found that deletion of the transcriptional silencer hns gene could restore both pagC and pagD transcription in the ΔemrR mutant to wild-type levels (Fig. 2E), suggesting that EmrR should function as an antirepressor to antagonize the H-NS effect at the pagC-pagD divergent promoter in a manner comparable to that of PhoP and SlyA (27). In contrast, EmrR did not exert an effect on transcription merely dependent on the PhoP/PhoQ system because transcription of two PhoP-activated genes, pcgL and STM3595 (36), remained at similar levels in wild-type and ΔemrR strains (Fig. S3A). Therefore, we demonstrated that EmrR could function as a transcription activator to enhance transcription of the SPI-2 genes and those stimulated by both PhoP and SlyA regulators.

The catecholamine neurotransmitter dopamine is a signal molecule to abrogate EmrR-activated transcription. In contrast to PhoP-activated slyA transcription (5, 21), transcription of the emrR gene was not regulated by either PhoP/PhoQ system or SlyA, since the levels of emrR transcripts were similar in the wild-type, ΔphoP, and ΔslyA strains (Fig. 3A, top panel). Also, the slyA and phoP mRNA levels were similar in the wild-type and ΔemrR strains (Fig. 3A, middle and bottom panels), which ruled out the possibility that EmrR could stimulate slyA and phoPQ transcription. The result also confirmed that slyA transcription was activated by the PhoP/PhoQ system (5) because the slyA mRNA level in the ΔphoP mutant was significantly reduced than that in the wild-type strain (Fig. 3A, middle panel). We postulate that the ligand-responsive regulator EmrR should also be able to respond to a specific signal molecule since most MarR family regulators bind small-molecule ligands, often phenolic compounds (37, 38). We chose some bioactive phenolic compounds (39) and investigated their effects on transcription of EmrR-regulated pagC gene on the premise that EmrR could sense one of these molecules. Among some 58 chosen phenolic compounds, we found that catecholamine neurotransmitter dopamine caused significant repression of pagC transcription (Fig. S2B). Consistently, β-galactosidase activity in the pagC-lacZ strain grown in low Mg²⁺ was reduced by supplementing dopamine in a concentration-dependent manner (Fig. 3B). Apparently, dopamine exerted its inhibitory effect on transcription of the pagC-pagD divergent operon since it could also reduce the β-galactosidase activity from a pagD-lacZ strain in a concentration-dependent manner (Fig. 3B). On the other hand, dopamine could not act on the PhoP/PhoQ system because the PhoP-activated transcription of the pcgL and STM3595 genes remained unchanged under the conditions with 2 mM dopamine or without dopamine (Fig. S3B). Furthermore, RT-PCR
analysis showed that dopamine could reduce mRNA levels of the tested SPI-2 genes in low Mg²⁺ (Fig. 2C). Concomitantly, dopamine reduced the survival rates of the wild-type strain living within macrophages and challenged by polymyxin B in a concentration-dependent manner, respectively (Fig. 1B and 1C). In this study, our...
results also showed that dopamine exerted a significant negative effect only in the presence of EmrR but could not further lower the survival rate of the ΔemrR mutant within macrophages (Fig. 1B). A sequence alignment analysis indicated that EmrR and SlyA exhibited ~30% amino acid sequence identity (Fig. S2C) and that many of the conserved residues were clustered in the helices α3 and α4 as well as the β wing for DNA recognition (40). As a matter of fact, our previous study revealed that SlyA, at a high level that could not be attained physiologically in a wild-type cell, could stimulate deficient transcription of both pagC and pagD genes caused by an absence of the positive signal, i.e.,alarmine-guanosine pentaphosphate (23). We found that SlyA when overproduced to 4.6-fold higher than the wild-type level in a ΔemrR ΔslyA mutant harboring plasmid psyA-FLAG (36) (Fig. S3C), stimulated pagC transcription even in the absence of EmrR to 61% of the wild-type level (Fig. 3C). Likewise, EmrR at 3.0-fold higher than the wild-type level (Fig. S3D) was able to stimulate the pagC transcription to 29% of the wild-type level in the double mutant harboring pemrR-FLAG in which SlyA was absent (Fig. 3C). Therefore, we took advantage of this partially overlapped function to investigate whether EmrR could be the responsive regulator to dopamine. We found that the EmrR-stimulated pagC transcription in the ΔemrR ΔslyA mutant carrying plasmid pemrR-FLAG was 7.1-fold reduced by 1 mM dopamine, whereas the SlyA-stimulated transcription in this mutant carrying psyA-FLAG was not influenced (Fig. 3D). l-Tyrosine, a precursor for biosynthesis of dopamine (41), did not affect the EmrR-stimulated pagC transcription even at a concentration as high as 40 mM (Fig. 3D). Taken together, we postulate that dopamine can serve as a signal molecule to act on EmrR and subsequently modulate S. Typhimurium gene regulation.

**EmrR binds to the pagC-pagD promoter regions at both PhoP and SlyA recognition sequences.** We conducted an electrophoretic mobility shift assay (EMSA) using a pagC-pagD promoter fragment which possesses both PhoP and SlyA binding sequences (namely, PhoP and SlyA boxes; see reference 23) to investigate EmrR binding to the target promoters. We found that an EmrR-His6 protein shifted this promoter fragment on the gel (Fig. 4A, lane 2). In this in vitro system, dopamine alone was sufficient to reduce the affinity of a pure EmrR protein to the promoter DNA fragment in a concentration-dependent manner (Fig. 4A, lanes 3 to 5). In contrast, dopamine, at a high concentration that actually eliminated EmrR-promoter interaction (Fig. 4A, lane 5), was unable to change the binding of SlyA-His6 protein to the same pagC-pagD promoter fragment (Fig. 5A). This specific in vitro interaction between dopamine and the EmrR protein suggested the possibility that dopamine was a ligand molecule to act on EmrR. A DNase I footprinting analysis confirmed the promoter binding of EmrR and revealed three major EmrR-protected AT-rich regions, marked RI, RII, and RIII (Fig. 4B). We found that the RI, RII, and RIII regions were localized between −67 and −62, −80 and −71, and −109 and −98 nucleotide sequences upstream of the pagC transcription start site, which meanwhile corresponded to the locations between −153 and −148, −144 and −13, and −117 and −106 nucleotide sequences upstream of the pagD transcription start site, respectively (Fig. 4C). Actually, the RI and RII sequences overlapped the PhoP box, TAAAT-(6 nt)-TAAAC, and the RIII sequence partially overlapped the SlyA box, AATATT-(10 nt)-ATTATT (Fig. 4C), thus elucidating that EmrR shared its binding sites with PhoP and SlyA simultaneously. However, there were no consensus sequences that could be found from these binding sites except for their AT-rich feature. It appeared that EmrR might also interact, however weakly, with other AT-rich regions adjacent to RI and RII sites (Fig. 4B). We reasoned that EmrR might recognize AT-rich regions rather than a specific consensus sequence since RI, RII, and RIII were neither homologous to each other (Fig. 4C) nor to an EmrR binding site characterized previously from the E. coli emr promoter (13).

**EmrR acts as a priming regulator to facilitate binding of successive regulators to the target promoters.** In contrast with increased levels of PhoP and SlyA and continuous activation of pagC-pagD transcription in a time-dependent manner, the level of EmrR remained unchanged during the growth of bacterial cells in low Mg²⁺ for 4 h (Fig. 5A and Fig. S5A). We performed chromatin immunoprecipitation analysis (ChIP)
to determine the in vivo interaction of EmrR to the pagC-pagD promoter in the emrR-FLAG strain and observed that the protein level of promoter-bound EmrR gradually decreased during the low-Mg\(^{2+}\) growth (Fig. 5B). Actually, the promoter decreased binding to EmrR and simultaneously increased binding to PhoP and SlyA in a
EmrR interacts with the promoter region to facilitate binding of other regulators in *S. Typhimurium*. (A) Immunoblot analysis for determination of the levels of EmrR-FLAG, PhoP-HA, and SlyA-FLAG proteins from the *phoP-HA* strain (YS15191), *slyA-FLAG* strain (YS10075), and *emrR-FLAG* strain (YS16035) grown in high (10 mM) or low Mg^{2+} for the indicated growth time. The level of CorA-FLAG protein from the *corA-FLAG* strain (YS11477) was used as an Mg^{2+}-independent control. (B) ChIP analysis for in vivo binding of EmrR-FLAG, PhoP-HA, and SlyA-FLAG proteins to the *pagC-pagD* promoter region in the *emrR-FLAG* strain, *phoP-HA* strain, and *slyA-FLAG* strain grown in low Mg^{2+} for 2 to 4 h, respectively. PCRs were conducted using DNA templates in immunoprecipitated samples (IP) and total lysates (Input). (C, left) Immunoblot analysis of the total levels of SlyA-FLAG and PhoP-HA proteins in *slyA-FLAG* and *phoP-HA* strains, WT strains (YS10075 and YS11591) and their isogenic ΔemrR mutants (YS15602 and YS15601) grown in low Mg^{2+} for 4 h. (Right) ChIP analysis for in vivo promoter binding of SlyA-FLAG and PhoP-HA in these bacterial cells. (D) ChIP analysis for the in vivo binding of the EmrR-FLAG protein to the *pagC-pagD* promoter region in the *emrR-FLAG* strain (YS16035) and the isogenic *PhoP-box* mutant (YS17214), and *SlyA-box* mutant (YS17213) grown in low Mg^{2+} for 3 h. (E) ChIP analysis for in vivo binding of EmrR-FLAG, PhoP-HA, and SlyA-FLAG proteins to the *pagC-pagD* promoter region in the *emrR-FLAG* strain (YS16035), *phoP-HA* strain (YS11591), and *slyA-FLAG* strain (YS10075) grown in low Mg^{2+} supplemented with 1 mM dopamine or without dopamine for 3 h. (F, top) Immunoblot analysis of the total levels of EmrR-FLAG protein in the *emrR-FLAG* strain, WT strain (YS16035) and their isogenic Δ*phoP* and Δ*slyA* mutants (YS15538 and YS15537) grown in low Mg^{2+} for 4 h. (Bottom) ChIP analysis for in vivo binding of EmrR-FLAG protein to the *pagC-pagD* promoter region in the Δ*phoP* mutant (YS15538) and Δ*slyA* mutant (YS15537) grown in low Mg^{2+} for 2 to 4 h. (G) ChIP (Continued on next page)
time-dependent manner since the amounts of the promoter fragments pulled down by PhoP-HA and SlyA-FLAG proteins were proportional to their protein levels in the *phaP-pha* _slyA-*FLAG* strain (Fig. 5A and B). Immunoblot analysis showed that the total levels of PhoP-HA and SlyA-FLAG proteins were similar in this *phaP-pha* _slyA-*FLAG* strain and its isogenic Δ*emrR* mutant (Fig. 5C, left panel). However, the levels of PhoP-HA and SlyA-FLAG proteins that bound to the promoter were much lower in the Δ*emrR* mutant than in the wild-type strain (Fig. 5C, right panel), indicating that the role of EmrR was to enhance the binding of PhoP and SlyA to the promoter. In accord with the *in vitro* binding result (Fig. 4B), EmrR interacted specifically with the SlyA and PhoP boxes of the *pagC-pagD* promoter, since nucleotide substitutions at either sequence, which were referred to as *SlyA-box* or *PhoP-box* and shown to cause downregulation of the *pagC-pagD* transcription previously (23), mostly abrogated the promoter-EmrR interaction *in vivo* (Fig. 5D). Consistent with its effect *in vitro* (Fig. 4A), ChIP analysis showed that dopamine added to the culture of *emrR-FLAG* strain caused a reduction in EmrR binding to the *pagC-pagD* promoter *in vivo* (Fig. 5E). Simultaneously, dopamine also caused a reduction in the binding ability of PhoP-HA and SlyA-FLAG proteins to the promoter *in vivo* (Fig. 5E), thus providing further evidence that EmrR enhanced the binding of PhoP and SlyA to the target promoter. It is worth noting that the bacterial cells cultured for 3 h were used to carry out the ChIP analyses in Fig. 5D and E, since more EmrR-FLAG proteins were found to bind to the *pagC-pagD* promoter in the wild-type strain (Fig. 5B). Under this growth condition, a reduced level of the promoter-bound EmrR caused by a mutation of the promoter sequence or supplementing with dopamine could be more accurately compared. Mutation of the _slyA_ and _phaP_ genes did not affect the expression of EmrR, since its total levels were similar in wild-type and mutant cells grown in low Mg$^{2+}$ for 4 h (Fig. 5F, top panel). However, the bound EmrR could not be dissociated from the *pagC-pagD* promoter in the Δ*phaP* mutant, and also more EmrR proteins remained bound to the promoter in the Δ*slyA* mutant than in the wild-type strain (Fig. 5F, bottom panel). We postulate that EmrR should be displaced from the promoter by PhoP and SlyA in a cooperative manner, because the absence of either regulator caused more EmrR bound to the promoter. Importantly, similar to the role in the *pagC-pagD* promoter, EmrR was also displaced by SlyA from the _ssrA_ promoter in SPI-2 because the level of promoter-bound EmrR and SlyA decreased and increased simultaneously in a time-dependent manner during the low-Mg$^{2+}$ growth (Fig. 5G). Subsequently, more EmrR proteins remained bound to this promoter in the Δ*slyA* mutant than in the wild-type strain (Fig. 5G), thus reconfirming that SlyA could displace EmrR from this promoter.

**EmrR controls the PhoP/PhoQ and SlyA feedforward regulatory loop in many enteric bacteria.** Sequence alignment analysis indicated that *S. Typhimurium* PhoP, SlyA, and EmrR were conserved in many enteric bacteria and specifically that the amino acid sequences revealed 93%, 79%, and 89% identity to their orthologs in *E. coli* and 75%, 93%, and 80% identity to their orthologs in *Yersinia pestis*. However, genetic loci regulated by the feedforward loop via these regulators remain elusive in these organisms. Thus, we investigated whether these orthologs could constitute the same regulatory network by using a reporter plasmid, pYS1031, which harbored a _lacZ_ fusion with a _pagC_ promoter fragment, including the SlyA and PhoP boxes (23). The cloned promoter fragment in this plasmid was sufficient to render not only a PhoP- and SlyA-dependent regulation (23) but also an EmrR-dependent regulation, since β-galactosidase activity in the *Salmonella* Δ*emrR* mutant harboring pYS1031 was 9.9-fold lower than that in the wild-type strain grown in low Mg$^{2+}$ (Fig. 6A). We found

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**FIG 5 Legend (Continued)**

analysis for *in vivo* binding of EmrR-FLAG and SlyA-FLAG proteins to the _ssrA_ promoter region in the _emrR-FLAG_ strain, WT strain (YS16035) and _slyA_ mutant (YS15537), and the _slyA-FLAG_ strain and WT strain (YS10075) grown in low Mg$^{2+}$ for 2 to 4 h. PCRs were conducted using DNA templates in immunoprecipitated samples (IP) and total lysates (Input). Data in panels A to F are from one of three independent assays.
that β-galactosidase activity in an E. coli wild-type strain (BW25113) harboring pYS1031 in low Mg2+ was 13.3-, 11.1-, and 8.4-fold higher than that in the isogenic ΔphoP, ΔslyA, and ΔemrR mutants, respectively (Fig. 6B). Salmonella EmrR protein functioned effectively in E. coli because reduced transcription of the pagC promoter-controlled lacZ fusion in the ΔemrR mutant was fully recovered by emrR overexpression from plasmid pemR-FLAG (Fig. S5B). In a manner similar to the phenotype in the Salmonella ΔslyA mutant, emrR overexpression from pemR-FLAG could also partially stimulate lacZ transcription in the E. coli ΔslyA mutant in which SlyA was absent (Fig. S5B). Furthermore, this EmrR-stimulated transcription in the E. coli ΔslyA mutant was reduced again by adding dopamine (Fig. S5B). Additionally, β-galactosidase activity from a Yersinia pestis wild-type strain (KIM6+) harboring pYS1031 grown in a pH 6.0 medium at 26°C was 7.9-, 8.1-, and 7.7-fold higher than that of the isogenic ΔphoP, ΔrovA (rovA is the slyA homolog in Yersinia; see reference 42), and ΔemrR mutants, respectively (Fig. 6C). These observations suggested that the same regulatory circuit constructed by these
regulators should be conserved in *E. coli* and *Y. pestis* and functional even for regulation of the *Salmonella*-specific *pagC* gene.

**DISCUSSION**

Here, the MarR family regulator EmrR is characterized as a virulence factor of *Salmonella enterica* serovar Typhimurium and *Yersinia pestis* and able to activate transcription by acting on the AT-rich regions at the promoters at genes such as the SPI-2 genes and also many horizontally acquired genetic loci controlled by the PhoP/PhoQ and SlyA-dependent feedforward loop.

A role of EmrR in transcription activation as a priming antirepressor to govern a sequential regulatory module. A well-known function of the MarR family regulators is to repress transcription mainly through exerting steric hindrance of RNA polymerase binding to target promoters by binding at the $-35$ and/or $-10$ promoter elements. Here, we have revealed an EmrR function as a transcriptional antirepressor by acting, in a manner similar to that of SlyA (5, 21, 23), on far upstream of the $-35$ region of the target promoters (Fig. 4B and C). In contrast with a specific direct repeat sequence recognized by SlyA (23), the EmrR binding sites lack a consensus sequence but appear to be AT-rich regions which are exemplified by the SlyA and PhoP box sequences in this study (Fig. 4C). We reason that a less sequence-specific feature will allow EmrR to interact a boarder range of the binding sites of other regulators such as SlyA and PhoP, by which it can counteract H-NS that occupies these AT-rich sequences to repress the transcription of the *ssrA* and *pagC-pagD* genes and many others. Initial binding of EmrR to these promoters is essential for transcription that is activated by other regulators such as SlyA and PhoP but becomes dispensable when H-NS is absent (Fig. 2E). EmrR binding to a promoter is not sufficient for transcription activation; for example, *pagC* transcription is repressed in the $\Delta phoP$ or $\Delta slyA$ mutant despite the fact that there are actually more EmrR proteins associated with the promoter (Fig. 5F). We hypothesize that the priming interaction of EmrR will make local conformation of a specific promoter, such as the *pagC-pagD* promoter, undergo a transition to form "a derepression state" (Fig. 7, left panel), which becomes more favored for subsequent binding of those successive transcription regulators. This is supported by the observation that the successive regulators SlyA and PhoP replace EmrR from the *pagC-pagD* promoter and also SlyA subsequently replaces EmrR from the *ssrA* promoter (Fig. 5B and G), by which the promoter region forms "an activation state" for transcription (Fig. 7, right panel). Therefore, in the $\Delta slyA$ mutant in which SlyA was absent, more EmrR proteins stay on both *pagC-pagD* and *ssrA* promoters (Fig. 5F and G), and in the *Δ phoP* mutant in which both PhoP and SlyA were absent, almost all EmrR remained on the *pagC-pagD* promoter (Fig. 5F). Two EmrR dimers may bind to the adjacent PhoP and SlyA boxes of the *pagC-pagD* promoter in a cooperative manner comparable to the $\lambda Cl$ dimers that bind to two neighboring operators $O_{41}$ and $O_{42}$ (43). This cooperative interaction must be important for EmrR binding to the *pagC-pagD* promoter, since nucleotide substitutions of either the SlyA box or the PhoP box were sufficient to reduce EmrR affinity dramatically to the promoter overall (Fig. 5D). The coordinate interaction between two EmrR dimers is devastated when one successive regulator displaces EmrR first, which should make it easier for the second regulator to replace another EmrR dimer from the promoter. It is worth noting that EmrR can also regulate PhoP- and SlyA-independent genetic loci, including its operon comprising *emrR*, *emrA*, and *emrB* genes encoding a multidrug resistance pump (annotated as *STM2813*, *STM2814*, and *STM2815* loci, respectively, in the *S. Typhimurium* LT2 genome). Our RNA-Seq results showed that transcription of the *emrA* and *emrB* genes in the $\Delta emrR$ mutant were 5.75- and 5.94-fold higher than those in the wild-type strain (see Table S3 in the supplemental material), thus reconfirming that EmrR was a negative regulator of the *emrAB* operon (9). This observation should also rule out the possibility that deletion of the *emrR* gene in our $\Delta emrR$ mutant had a polar effect on transcription of downstream *emrA* and *emrB* genes. Otherwise, this deletion should cause a reduction in transcription of these downstream genes. In support of this conclusion, a complementing plasmid which solely contained
the emrR coding region (pemrR-FLAG) fully recovered EmrR-dependent transcription in the ΔemrR mutant to wild-type levels (Fig. 2A, D, and E).

**Neurotransmitter dopamine acts on EmrR to downregulate transcription of virulence genes.** Some gut bacteria affect the host gastrointestinal and psychological health by producing and responding to particular neuroactive compounds such as dopamine, serotonin, and norepinephrine (44). Dopamine has been considered a catecholamine signaling molecule for host-pathogen communication in the gut lumen (41, 45). Here for the first time, characterization of dopamine as a signal to interact with EmrR provides evidence that enteric bacteria are able to respond to dopamine through this MarR family regulator and probably also other specific sensor proteins. It is reasonable to say that dopamine should influence the virulence of enteric bacteria through EmrR as it is a virulence factor (Fig. 1A and Fig. S1). Dopamine is a precursor molecule of norepinephrine and epinephrine, both of which are required for full virulence of *Citrobacter rodentium* during mammalian infection. It was shown that host neurotransmitters epinephrine and/or norepinephrine in the gut act on two bacterial adrenergic sensors QseC and QseE to increase the virulence gene expression in enterohemorrhagic *E. coli* and *Citrobacter rodentium* (46). However, dopamine does not act on QseC or QseE, and transcription of the *pagC-pagD* operon is not regulated by these sensor kinases under all tested growth conditions (unpublished data). Therefore, our findings reveal the first bacterial signal transduction pathway in response to dopamine. Our study also illustrates that dopamine acts as a negative signal through sensor EmrR to inhibit transcription of specific *S. Typhimurium* genetic loci, including the SPI-2 genes and a set of divergent operons that are essential for bacterial pathogenesis. In support of its negative effect in these genetic loci, we found that dopamine reduced the ability of *Salmonella* to survive within macrophages and resistance to antimicrobial peptide (Fig. 1B and C), both of which were required for *Salmonella* virulence (28, 47). Also, several previous studies have shown that dopamine was able to

![Diagram of EmrR-dependent gene regulation in *S. Typhimurium*.](https://example.com/diagram.png)

**FIG 7** Model illustrating an EmrR-dependent gene regulation in *S. Typhimurium*. The EmrR proteins bind to the SlyA box and the PhoP box sequences through coordinate interaction to replace the transcription repressor H-NS from the pagC promoter region. (Left) The pagC promoter forms a “derepression state” after binding to EmrR. When *S. Typhimurium* is grown in low-Mg\(^{2+}\) conditions (and perhaps acidic pH or specific cationic antimicrobial peptides), the PhoP protein is phosphorylated (P\(^{i}\)). Transcription of the slyA gene is activated by the phosphorylated PhoP protein. Then, EmrR dimers in the pagC promoter region enhance the phosphorylated PhoP to bind the PhoP box, and also the ppGpp-bound SlyA dimer to bind the SlyA box. (Right) The PhoP- and SlyA-bound promoter further forms an “activation state” for transcription initiation. On the other hand, dopamine interacts with the EmrR protein and reduces its ability to bind the promoter region, thereby downregulating pagC transcription.
decrease early S. Typhimurium uptake into Peyer’s patches where invasive bacteria could penetrate the specialized epithelial M cells to initiate their murine infection (48, 49). We reason that in such a host environment experienced by Salmonella, dopamine will act on specific sensor protein(s) to downregulate expression of those genes required for bacterial invasion. Thus, the establishment of a link between dopamine and EmrR should provide insights into molecular mechanisms to highlight how EmrR regulates Salmonella virulence genes and how dopamine modulates the action of EmrR in specific niches. Particularly in this study, we also provide a singular example of signal transduction requiring the action of EmrR, in which this regulator senses the level of negative signal dopamine to modulate a regulatory circuit stimulated by two positive signals, environmental low Mg\(^{2+}\) through the PhoP/PhoQ system and cytoplasmic ppGpp level through SlyA in Enterobacteriaceae.

MATERIALS AND METHODS

**Bacterial strains, growth conditions, and oligonucleotides.** Bacterial strains used in this study are listed in Table S1 in the supplemental material. All Salmonella enterica serovar Typhimurium strains were derived from the wild-type strain ATCC 14028s (14028). All Escherichia coli strains were derived from the wild-type strain BW25113. Salmonella and E. coli cells were grown at 37°C in Luria-Bertani (LB) broth (Difco) or N minimal medium (50) (pH 7.4) supplemented with 0.1% Casein amino acids and 36 mM glycerol. MgCl\(_2\), and 4832 was added to the broth or medium. All Yersinia pestis strains were derived from the wild-type strain KIM6+ (Pgm\(^{+}\)). Bacterial cells were grown routinely at 26°C on Congo red agar from glycerol stocks to confirm the pigmentation (Pgm) phenotype of Y. pestis strains (51) and then grown on tryptose blood agar (TBA) and heart infusion broth (HIB) (Difco) (52). When necessary, antibiotics were added to bacterial cultures at final concentrations of 50 μg/ml for ampicillin (Ap), 20 μg/ml for chloramphenicol (Cm), or 50 μg/ml for kanamycin (Km). E. coli DH5α and BL21-Gold (DE3) were used as hosts for the preparation of plasmid DNA and protein production, respectively. Oligonucleotides used in this study are given in Table S2.

**Construction of strains with chromosomal mutations, harboring lac gene fusions, or epitope-tagged proteins.** S. Typhimurium and Yersinia strains harboring deletions were generated as described previously (53). If needed, the antibiotic resistance cassette was removed using plasmid pcP20. In Salmonella, deletions of the emrR, STM2920, STM1100, marR, STM1547, ugtL, and ssaM genes were carried out using primer pairs 1857 and 1858, 2535 and 2536, 2537 and 2538, 1880 and 1881, 4478 and 4479, 2624 and 2625, and 3475 and 3476, respectively, to amplify the chloramphenicol resistance (Cmr) gene or kanamycin resistance (Kmr) cassette from plasmid pKD3 or pKD4 (53). The PCR products were electroporated into wild-type cells harboring pKD46, and Cmr or Kmr colonies were selected. Integration of the drug resistance cassette into the chromosomes in these mutants was confirmed by colony PCR. To construct the Salmonella emrR-FLAG strain (YS16035), a PCR fragment was synthesized with primers 1907 and 1858 from pKD3 and then electroporated into the wild type harboring pKD46, and Cmr colonies were selected. The FLAG fusion was confirmed using colony PCR and DNA sequencing. Similarly, in Y. pestis, deletions of the emrR and rovA genes were carried out using primer pair 4480 and 4481 and primer pair 4482 and 4483, respectively, to amplify the chloramphenicol resistance (Cmr) or kanamycin resistance (Kmr) cassette from pKD46. Cmr or Kmr colonies were verified by colony PCR. The Cmr cassette was removed by using plasmid pCP20.

**Plasmid construction.** All plasmids used in this study are listed in Table S1. Plasmid pemrR-FLAG was constructed using PCR fragments containing the emrR coding region generated with primers 1910 and 1858 from pKD46 and chromosomal mutations in these mutants were confirmed by colony PCR. To construct the Salmonella emrR-FLAG strain (YS16035), a PCR fragment was synthesized with primers 1907 and 1858 from pKD3 and then electroporated into the wild type harboring pKD46, and Cmr colonies were selected. The FLAG fusion was confirmed using colony PCR and DNA sequencing. Similarly, in Y. pestis, deletions of the emrR and rovA genes were carried out using primer pair 4480 and 4481 and primer pair 4482 and 4483, respectively, to amplify the chloramphenicol resistance (Cmr) or kanamycin resistance (Kmr) cassette from pKD46. Cmr or Kmr colonies were verified by colony PCR. The Cmr cassette was removed by using plasmid pCP20.

**Stranded RNA-Seq analysis to collect transcriptomic data.** Total RNAs were isolated from wild-type and ΔemrR strains using SV Total RNA Isolation System (Promega) according to the manufacturer’s instructions from bacterial culture. Total RNAs were ribo-depleted using a Ribo-Zero rRNA removal kit (Illumina). The ribo-depleted RNAs were then enzymatically sheared to roughly 350 bp in size using KAPA’s stranded RNA-Seq kit (Roche). KAPA’s stranded RNA-Seq kit along with Illumina-compatible adapters (IDT) was used for the remaining library construction. The adapter-ligated molecules were cleaned using AMPure beads (Agencourt Bioscience/Beckman Coulter) and amplified with KAPA’s HiFi enzyme. Each library was then analyzed for fragment size on an Agilent Tapestation and quantified by qPCR (KAPA) on Quantsstudio 5 (Thermo Fisher Scientific) before multiplex pooling and sequencing on a 2x75 flow cell on the NextSeq500 platform (Illumina) at the Arizona State University Genomics Core facility. RNA-Seq reads for each sample were quality checked using FastQC v0.10.1 and aligned to the S. Typhimurium LT2 genome from NCBI (https://www.ncbi.nlm.nih.gov/nuccore/NC_003197.2) by TopHat v2.0.9. RNA-Seq data analysis was performed by the bacterially specific tool Rockhopper (https://cs.wellesley.edu/~btjaden/Rockhopper/index.html), including alignment, upper quartile normalization, reference-based and novel transcript identification, abundance quantification, differential gene expression detection, and operon prediction.
Reverse transcription-PCR (RT-PCR). Bacterial cells were grown for 4 h in N medium supplemented with 0.01 mM MgCl₂. DOPamine (1 mM) was added when needed. Total RNAs were isolated from the bacterial culture using the Total RNA Isolation System (Promega) according to the manufacturer’s instructions. The concentration of RNA was determined by spectrophotometry at 260 nm. The quality of RNA was confirmed by agarose gel electrophoresis. cDNA was synthesized using murine leukemia virus reverse transcriptase and random primers (New England Biolabs). DNA was amplified with primers 2031 and 2628 for emrR, primers 2529 and 2530 for rpoD, primers 2035 and 2036 for phoP, 2037 and 2038 for slyA, 3959 and 3960 for ssdB, 2732 and 2738 for sseB, 3477 and 3485 for fadA, and 3447 and 3448 for ssaM (Table S2) using Taq polymerase (New England Biolabs) and performed in a thermocycler (Bio-Rad).

5. Typhimurium survival assays inside J774.2 cells. Macrophages were plated at 5 × 10⁵ cells per well in a 24-well plate and incubated overnight at 37°C in 5% CO₂. They were then infected with stationary-phase bacteria at 5 × 10⁴ CFU in DMEM supplemented with 10% FCS and incubated for 30 min. The wells were washed three times with PBS and incubated with DMEM, 10% FCS, and 100 μg/ml gentamicin for 2 h (2 h postinfection). Bacterial invasion was determined at this time point by counting the number of CFU of the intracellular S. Typhimurium cells recovered from lysed macrophages in three wells infected by each bacterial strain. Then, the medium was replaced with DMEM, 10% FCS, and 10 μg/ml gentamicin, and the plate was incubated for another 16 h for bacterial cell counting. Dopamine was added to the DMEM medium at concentrations of 0.2 and 1.0 mM, or dopamine was not added. Heterologous expression of emrR from complementing plasmid pemrR-FLAG was induced by adding 0.2 mM IPTG. After washing the wells three times with PBS, macrophages in the wells were lysed with PBS containing 1% Triton X-100. Recovered bacterial cells were counted after grown on LB plates overnight at 37°C.

β-Galactosidase assay. The β-Galactosidase assay was carried out in triplicate; overnight cultures were diluted with N medium and grown for 4 h, and the activity was determined as described previously (55). M₉₀⁺ was added to 0.01 mM (namely, low M₉₀⁺), and 10 mM (high M₉₀⁺). Complementation was carried out by using complementing plasmid pemrR-FLAG, and heterologous expression of emrR from this plasmid was induced by adding 0.2 mM IPTG or indicated specifically. When needed, salicylate (Sigma-Aldrich) or dopamine (Alfa Aesar, Thermo Fisher Scientific) was added to the required concentrations. Data correspond to three independent assays conducted in duplicate, and all values were means ± standard deviations.

Expression of the pagC gene in the presence of phenolic compounds. The β-galactosidase activity of the pagC-lacZY strain was determined after bacterial cells were grown for 4 h in N medium in low M₀⁺ (0.01 mM) supplemented with or without a natural phenolic compound selected from a published list (39) and picked up from our storage. These phenolic compounds were dissolved in ethanol, DMSO, methanol, or water by following the instructions from their manufacturers and suppliers (Sigma-Aldrich, Thermo Fisher Scientific, and VWR). We chose a final concentration of these compounds for this test that did not cause a loss of bacterial viability during growth. Data correspond to three independent cultures, and all values were means ± standard deviations.

Isolation of the EmrR-His₆ and SlyA-His₆ proteins. E. coli BL21-Gold(DE3) harboring plasmids pET28a-EmrR-his₆ (pYS2017) and pET28a-slyA-his₆ (pYS1277) was grown at 37°C with shaking to an OD₆₀₀ of 0.5 in 500 ml of LB medium; then IPTG (final concentration, 1 mM) was added, and bacteria were incubated for 2 h. Cells were harvested, washed with PBS once, resuspended in 10 ml of PBS, and lysed by sonication. The whole-cell lysates were used for EmrR-His₆ and SlyA-His₆ purification by mixing with His-Select Nickel Affinity Gel (Qiagen) following the instructions from the manufacturer. Pure EmrR-His₆ and SlyA-His₆ samples were tested using silver staining (Pierce) following the instructions from the manufacturer.

Electrophoretic mobility shift assay (EMSA). Primers were labeled using T4 polynucleotide kinase (New England Biolabs) and [γ⁻³²P]ATP (PerkinElmer Life Sciences). ³²P-labeled DNA fragments (10 nmol) containing pagC-pagD promoter regions, amplified by PCR from S. Typhimurium chromosomes with primer 522 and ³²P-labeled primer 523 (23), were incubated at room temperature for 30 min with 0 or 50 pmol of EmrR-His₆ and SlyA-His₆, regulator proteins in 20 μl of an EMSA buffer (56), respectively. When required, dopamine (final concentrations of 1, 10, and 100 μM) and cold PCR products (20 μM) were added to reduce regulator protein-promoter binding and compete for the binding of regulator protein, respectively. After the addition of the DNA dye solution, the mixture was directly subjected to 4% polyacrylamide electrophoresis. Signals were detected by autoradiography.

DNase I protection assay. DNase I protection assays were performed using DNA fragments amplified by PCR using chromosomal DNA as the template. Before PCR, primer 523 was labeled with T4 polynucleotide kinase and [γ⁻³²P]ATP. The pagC promoter region was synthesized with primers 522 and ³²P-labeled primer 523. Approximately 25 pmol of labeled DNA and 0, 50, 100, or 200 pmol of the EmrR-His₆ protein were used in a 100-μl reaction mixture. DNase I digestion was conducted as described previously (57) using 0.05 U of DNase I (Invitrogen) per reaction mixture. Samples were analyzed by 6% denaturing polyacrylamide electrophoresis by comparison with a DNA sequence ladder generated with the appropriate primer by Maxam and Gilbert A⁻G reaction. The positions of radioactive DNA fragments in the gels were detected by autoradiography.

Immunoblot assay. Bacteria producing epitope-tagged proteins were inoculated into N minimal medium (pH 7.4) with 10 mM MgCl₂ (high Mg²⁺) for overnight growth at 37°C with aeration, and the cultures were shifted to N minimal medium (pH 7.4) with 10 μM MgCl₂ (low Mg²⁺) and grown for 4 h. Bacterial cells were harvested and lysed by sonication. The protein concentration of whole-cell lysates was determined by using the BCA Protein Assay Kit (Pierce). Protein samples were separated by 15% SDS-PAGE and transferred to nitrocellulose membranes (Amersham), and the PhoP-HA, SlyA-FLAG, and EmrR-FLAG proteins were monitored with ECL Western blotting substrate (Amersham) after incubation with monoclonal anti-HA or anti-FLAG M2 antibody (Sigma-Aldrich).
Chromatin immunoprecipitation assay. S. Typhimurium strains harboring chromosomally encoded PhoP, SlyA, and EmrR proteins with a C-terminal HA or FLAG epitope were grown in 25 ml N medium as described above. Bacterial cultures were incubated for 4 h or an indicated time. After culturing, bacterial cells were washed once with PBS and suspended in 25 ml PBS. Proteins were cross-linked to promoter DNA by adding formaldehyde to a 1% final concentration. Chromatin immunoprecipitation (ChIP) assays were performed as described previously (58). Enriched DNA fragments containing the pagC promoter region were detected by PCR using primer pair 522 and 523.

Bacterial virulence analysis in mice. All animal procedures were approved by the Arizona State University Animal Care and Use Committee. Single colonies of each S. Typhimurium strain from LB agar plates were used to inoculate LB broth and grown overnight at 37°C. Bacteria were diluted (1:20) with 50 ml of fresh LB and grown to an OD600 of 0.85. The cells were then harvested and suspended in 1 ml PBS with gelatin. Female 7-week-old BALB/c mice from The Jackson Laboratory were inoculated by oral administration or intraperitoneal injection with bacterial suspension. Actual numbers of CFU inoculated were determined by plating serial dilutions onto LB. Mice were observed over a 30-day period to record survival. To determine the virulence of Y. pestis, single colonies of each strain from Congo red agar plates were used to inoculate HIB broth and grown overnight at 26°C. To maintain plasmid pCD1Ap, ampicillin was added to the medium at a concentration of 25 μg/ml. Bacteria were diluted with 10 ml of fresh HIB enriched with 0.2% xylose and 2.5 mM CaCl2 to obtain an OD600 of 0.1 and incubated at 26°C for subcutaneous (s.c.) infection (bubonic plague) or at 37°C for intranasal (i.n.) infection (pneumonic plague). Both cultures were grown to an OD600 of 0.6. The cells were then harvested and suspended in 1 ml PBS. Female 7-week-old Swiss Webster mice from Charles River Laboratories (Wilmington, MA) were inoculated by s.c. injection under the skin on the back of the neck with 100 μl of bacterial suspension or by the i.n. route with 20 μl of bacterial suspension (around 107 CFU/mouse). Actual numbers of CFU inoculated were determined by plating serial dilutions onto TBA. Mice were observed over a 21-day period to record survival. Data are expressed as means ± SDs. The log rank test was used for analysis of the survival curves. A P value of <0.05 was considered significant.

Antimicrobial peptide killing assay. The antimicrobial peptide susceptibility assay was described previously (59). Briefly, bacterial cultures were grown in N medium (pH 7.4) with 10 μM MgCl2 for 4 h. Dopamine was added to the N medium at concentrations of 0, 1.5, and 2.0 mM. Heterologous expression of emrR from the complementing plasmid pemrR-FLAG was induced by adding 0.2 mM IPTG. Then, bacterial cells were diluted to 1 × 105 to 2 × 106 ml−1 in fresh N medium. Five microliters of serially diluted polymyxin B (Sigma-Aldrich) was mixed with 45 μl of the bacterial suspension in each well of a 96-well plate (Cell Culture Cluster; Costar) and incubated at 37°C for 1 h. Bacterial mixtures were diluted and transferred to LB agar plates, and the plates were incubated overnight at 37°C. The number of CFU on the plates was counted. The percentage of survival was calculated as follows: survival (%) = (CFU of peptide-treated culture/CFU of no-peptide culture) × 100.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.02772-18.

FIG S1, TIF file, 0.2 MB.
FIG S2, TIF file, 0.8 MB.
FIG S3, TIF file, 0.3 MB.
FIG S4, TIF file, 0.1 MB.
FIG S5, TIF file, 0.3 MB.
TABLE S1, PDF file, 0.1 MB.
TABLE S2, PDF file, 0.03 MB.
TABLE S3, XLSX file, 0.1 MB.
TABLE S4, XLSX file, 0.1 MB.

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W.K. and Y.S. designed experiments and wrote the main manuscript text. D.Y., Y.K., W.S., W.K., and Y.S. carried out experiments, collected data, and analyzed results. All authors reviewed the manuscript.

We declare that we have no conflicts of interest.

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