A Dual Regulatory Role of the PhoU Protein in \textit{Salmonella} Typhimurium

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\textbf{ABSTRACT} Bacteria utilize two-component regulatory systems to sense and respond to their surroundings. Unlike other two-component systems that directly sense through a sensory domain in the histidine kinase (HK), the PhoB/PhoR two-component system requires additional proteins, including the PstSCAB phosphate transporter and the PhoU protein, to sense phosphate levels. Although PhoU is involved in phosphate signaling by connecting the PstSCAB transporter and PhoR histidine kinase, the mechanism by which PhoU controls expression of \textit{pho} regulon genes has not yet been clearly understood. Here, we identified PhoU residues required for interacting with PhoR histidine kinase from the intracellular pathogen \textit{Salmonella} enterica serovar Typhimurium. The PhoU Ala147 residue interacts with the PhoR PAS domain and is involved in repressing \textit{pho} expression in high phosphate. Unexpectedly, the PhoU Arg184 residue interacts with the PhoR histidine kinase domain and is required for activating \textit{pho} expression in low Mg$^{2+}$ by increasing PhoR autophosphorylation, revealing its new function. The substitution of the Arg184 to Gly codon decreased \textit{Salmonella} virulence both in macrophages and in mice, suggesting that PhoU’s role in promoting PhoR autophosphorylation is required during \textit{Salmonella} infection.

\textbf{IMPORTANCE} Bacteria constantly sense and respond to their surroundings through two-component systems. In Gram-negative bacteria, phosphate sensing is mediated by the PhoB/PhoR two-component system with additional components, the PstSCAB phosphate transporter and the PhoU protein. PhoU, a regulatory protein that connects the PstSCAB phosphate transporter to the PhoB/PhoR two-component system, is believed to function as a negative regulator in phosphate signaling because the \textit{phoU} deletion mutant loses the ability to repress \textit{pho} expression in high phosphate. Using single amino acid substitutions in the intracellular pathogen \textit{Salmonella} enterica serovar Typhimurium, PhoU turns out to control PhoR histidine kinase differently, depending on the conditions. The PhoU-PhoR PAS domain interaction is involved in repressing \textit{pho} expression in high phosphate, whereas the PhoU-PhoR HK domain interaction is involved in activating autophosphorylation of PhoR histidine kinase in low Mg$^{2+}$ and thus promotes \textit{Salmonella} virulence. Therefore, PhoU appears to modulate phosphate signaling exquisitely according to external conditions.

\textbf{KEYWORDS} phosphate limitation, low Mg$^{2+}$, \textit{pst} phosphate-transport system, PhoB/PhoR two-component system, phosphate limitation
respond to phosphate levels. In Gram-negative bacteria, phosphate levels are sensed and controlled by the PhoB/PhoR two-component system, which has been well studied in *Escherichia coli* (2). PhoR is a membrane-bound histidine kinase (HK) that consists of a transmembrane (TM) domain with two helices, a Per-Arnt-Sim (PAS) domain, a dimerization and histidine phosphotransfer (DHp) domain, and a catalytic active/ATP-binding (CA) domain (3). In phosphate-limiting conditions, PhoR self-phosphorylates the histidine residue in its own DHp domain and transfers the phosphate group to the aspartate residue of the cognate PhoB response regulator. Phosphorylated PhoB then binds to the promoters of PhoB-dependent genes and increases transcription of the so-called *pho* regulon genes, including the PstSCAB₂ high-affinity phosphate transporter, the PhoU accessory protein, and the PhoB/PhoR two-component system itself (3). Phosphate signaling via PhoR histidine kinase is unique because, unlike other histidine kinases that contain a periplasmic sensory domain involved in sensing a corresponding signal(s), PhoR lacks such a periplasmic domain. Rather, it was suggested that the phosphate-transporting activity through the PstSCAB transporter is linked to the kinase activity of PhoR histidine kinase (4). In *Escherichia coli*, the connection between the PhoB/PhoR two-component system and the PstSCAB₂ phosphate transporter is physically linked by the PhoU protein that is loosely associated with the inner membrane (5, 6). Even though the PhoU protein bridges PhoR histidine kinase and the PstSCAB₂ phosphate transporter, it was reported that the interaction with the former is stronger than that with the latter (5). A previous bacterial two-hybrid analysis determined that, more specifically, PhoU binds to the PAS domain of the PhoR kinase (5). Given that the PAS domain generally functions as a molecular sensor by interacting with other molecules or proteins (7, 8), molecular interactions between PstB, PhoU, and the PhoR PAS domain appear to be critical in phosphate signaling.

The function of the PhoU protein has been suggested as a negative regulator in phosphate signaling based on the fact that, in *E. coli* and other bacteria, deletion of the *phoU* gene constitutively derepressed expression of *pho* regulon genes in both high- and low-phosphate conditions (9–12). The elevated expression of *pho* regulon genes in the *phoU* deletion mutant suggests that the autophosphorylation activity of the PhoR histidine kinase is always active unless PhoU binds to PhoR and represses autophosphorylation activity. Although it is not formally demonstrated, this also led to an assumption that PhoU might be dissociated/released from the PhoR histidine kinase in low-phosphate conditions to promote autophosphorylation activity of PhoR, resulting in activation of *pho* expression. However, it is still unclear how PhoU accommodates its conformation within the PstSCAB-PhoU-PhoR signaling complex in phosphate-limiting or -replete conditions.

Interestingly, a previous direct coupling analysis suggested that the *E. coli* PhoU protein appears to be proximal to the PhoR CA domain in addition to the above-mentioned PAS domain (6). This implies that PhoU could be in close contact with PhoR histidine kinase at multiple sites, although the physiological implications of such interactions are unclear. In addition, several recent studies have reported that PhoU might have an additional role(s) other than a negative regulator in phosphate signaling. When *E. coli* is grown in low-phosphate and low-potassium media, PhoU is likely to connect the phosphate-sensing PhoB/PhoR two-component system and the potassium-sensing KdpE/KdpD two-component system (13). Thus, in the absence of the cognate KdpD histidine kinase, PhoU allows expression of KdpE-dependent potassium transporter genes in response to a phoshorelay via PhoR histidine kinase (13). PhoU also promotes mutagenesis in *E. coli* when a double-strand break becomes mutagenic in a strain activating SOS response and stationary sigma factor (σ^S)-mediated response (14). However, this mutagenic function of PhoU seems to be linked not to phosphate levels but the ArcA histidine kinase-dependent response controlling aerobic respiration (14). Moreover, PhoU from *Caulobacter crescentus* was reported that it is not involved in repressing *pho* signaling in high-phosphate conditions, but it is involved in polyphosphate accumulation (10). Cumulatively, it implies that PhoU might have multiple functions that need to be addressed.
In the intracellular pathogen *Salmonella enterica* serovar Typhimurium, the *pst* and *phoB*/*phoR* genes were highly induced when *Salmonella* was inside macrophages (15). In addition, *Salmonella* virulence protein MgtC binds to the CA domain of PhoR and activates PhoR autophosphorylation, thereby promoting expression of *pho* regulon genes and phosphate uptake (16). Because the *mgtC* gene is also highly expressed inside macrophages (15, 17), high levels of *mgtC* expression support the activation of the PhoB/PhoR two-component system during *Salmonella* infection, the underlying mechanism of which remains unclear (16, 18). Here, we investigated the roles of the PhoU protein from *Salmonella enterica* serovar Typhimurium. Although maintaining phosphate levels appears to be critical for *Salmonella* pathogenesis (16, 19), the role(s) of the PhoU protein in *pho* expression and *Salmonella* pathogenesis has not been investigated yet. Based on protein homology modeling, we identified the PhoU residues that are required for PhoR interaction (Fig. 1). *Salmonella* PhoU Ala147 residue was identified to interact with the PAS domain of PhoR, similar to that previously reported in *E. coli*. Interestingly, we found another residue, PhoU Arg184, which is required for interacting with the PhoR CA domain, and the removal of the interaction by substituting Arg184 to Gly prevents MgtC-mediated PhoR autophosphorylation and subsequent induction of *phoE* mRNA levels. PstB is a part of *pst* high-affinity phosphate-specific transport system together with PstS phosphate-binding protein and two membrane permeases, PstA and PstC proteins.

**RESULTS**

The Ala147 residue of *Salmonella* PhoU is required to interact with PhoR histidine kinase and represses expression of the PhoB-dependent genes in high phosphate. In *E. coli*, a previous study reported that PhoU interacts with the PAS domain of PhoR histidine kinase to repress expression of PhoB-dependent genes in high-phosphate conditions (5, 20). To explore a functional role of PhoU in *Salmonella enterica*, we started to test whether *Salmonella* PhoU could interact with PhoR histidine kinase using a bacterial two-hybrid assay. Indeed, the *E. coli* *cyaA* mutant expressing T18-PhoU and T25-PhoR together exhibited a strong blue color on Luria-Bertani (LB) X-Gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) plates, indicating that N-terminally T18-fused PhoU and N-terminally T25-fused PhoR proteins physically interact with each other to complement...
CyaA adenylate cyclase and produce $\beta$-galactosidase from a cAMP-dependent promoter (Fig. 2A and B).

We then tested whether Ala147 and Arg148 residues of the PhoU protein are required for PhoR interaction based on the following reasons. First, the substitutions of the E. coli PhoU Ala147 or Arg148 residues were reported to weaken the interaction with PhoR, losing the ability to suppress the elevated alkaline phosphatase activity of the chromosomal phoU deletion mutant when expressed heterologously (6). Second, homology-based modeling and amino acid sequence alignment suggested that Ala147 and Arg148 residues in the PhoU proteins from E. coli and S. enterica are well conserved in both species (Fig. 2C; see Fig. S1 in the supplemental material). Similar to those reported in E. coli, T18-PhoU derivatives that harbor the Ala147-to-Glu or Arg148-to-Ala substitutions significantly decreased $\beta$-galactosidase production when coexpressed with T25-PhoR (Fig. 2A and B), indicating that Ala147 and Arg148 residues in the Salmonella PhoU protein are required for PhoR interaction and suppression of phoE mRNA levels in high phosphate. (A) Bacterial two-hybrid assay between PhoR and PhoU or PhoU variant proteins. Escherichia coli BTH101 strains harboring two plasmids, pUT18c and pKT25 derivatives, expressing an N-terminal fusion of the cyaA T25 fragment to the phoR gene and N-terminal fusions of the cyaA T18 fragment to the wild-type phoU, phoU$^{A147E}$, phoU$^{A147K}$, and phoU$^{R148A}$ genes or the pUT18c empty vector (negative) were spotted as indicated. Cells expressing both pUT18-mgtC and pKT25-mgtR are spotted as a positive control (37). Cells were spotted onto LB plates containing 80 $\mu$M X-Gal and 0.1 mM IPTG and incubated at 30°C for 40 h. Blue-colored colonies indicate a positive interaction. (B) $\beta$-Galactosidase assay from strains listed above. The average $\beta$-galactosidase activities (Miller units) are shown as mean ± SD ($n =$ 3, independent measurements). (C) Modeled structure of the Salmonella PhoU monomer predicted from a homology modeling program (Phyre2) based on the crystal structure of Pseudomonas aeruginosa PhoU (PDB ID 4Q25). Arg184 is located at the $\alpha$6 helix and Gln120 is located at the $\alpha$3-$\alpha$4 loop. The Arg184 and Gln120 residues are indicated with gray sticks, and PAS domain-interacting residues (Ala147 and Arg148) are located between $\alpha$4 and $\alpha$5 regions and indicated with pale gray sticks (6). The protein is colored blue to red, going from the amino terminus to the carboxyl terminus. (D and E) Relative mRNA levels of the phoE (D) and phoU (E) genes in Salmonella strains with the wild-type phoU gene (14028s), Ala147 to Glu-substituted phoU gene (SM323), or a strain that deleted the phoU gene. Bacteria were grown for 5 h in N-minimal medium containing 10 mM Pi (high Pi) or 0.01 mM Pi (low Pi). Shown are the means ± SD ($n =$ 3, independent measurements). Relative mRNA levels represent (target RNA/rrsH RNA) × 10,000. 

FIG 2 PhoU Ala147 and Arg148 residues are required for PhoR interaction and suppression of phoE mRNA levels in high phosphate. (A) Bacterial two-hybrid assay between PhoR and PhoU or PhoU variant proteins. Escherichia coli BTH101 strains harboring two plasmids, pUT18c and pKT25 derivatives, expressing an N-terminal fusion of the cyaA T25 fragment to the phoR gene and N-terminal fusions of the cyaA T18 fragment to the wild-type phoU, phoU$^{A147E}$, phoU$^{A147K}$, and phoU$^{R148A}$ genes or the pUT18c empty vector (negative) were spotted as indicated. Cells expressing both pUT18-mgtC and pKT25-mgtR are spotted as a positive control (37). Cells were spotted onto LB plates containing 80 $\mu$M X-Gal and 0.1 mM IPTG and incubated at 30°C for 40 h. Blue-colored colonies indicate a positive interaction. (B) $\beta$-Galactosidase assay from strains listed above. The average $\beta$-galactosidase activities (Miller units) are shown as mean ± SD ($n =$ 3, independent measurements). (C) Modeled structure of the Salmonella PhoU monomer predicted from a homology modeling program (Phyre2) based on the crystal structure of Pseudomonas aeruginosa PhoU (PDB ID 4Q25). Arg184 is located at the $\alpha$6 helix and Gln120 is located at the $\alpha$3-$\alpha$4 loop. The Arg184 and Gln120 residues are indicated with gray sticks, and PAS domain-interacting residues (Ala147 and Arg148) are located between $\alpha$4 and $\alpha$5 regions and indicated with pale gray sticks (6). The protein is colored blue to red, going from the amino terminus to the carboxyl terminus. (D and E) Relative mRNA levels of the phoE (D) and phoU (E) genes in Salmonella strains with the wild-type phoU gene (14028s), Ala147 to Glu-substituted phoU gene (SM323), or a strain that deleted the phoU gene. Bacteria were grown for 5 h in N-minimal medium containing 10 mM Pi (high Pi) or 0.01 mM Pi (low Pi). Shown are the means ± SD ($n =$ 3, independent measurements). Relative mRNA levels represent (target RNA/rrsH RNA) × 10,000. 

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required for PhoR interaction. It is interesting to note that the Ala147-to-Lys substitution did not affect the interaction (Fig. 2A and B), unlike the E. coli counterpart (6). When we created a phoU mutant strain with the Ala147-to-Glu substitution at its chromosomal location, the phoU variant with the Ala147-to-Glu substitution completely lost the ability to repress mRNA levels of the phoE and phoU genes in high phosphate (Fig. 2D and E), supporting the notion that Salmonella PhoU is a negative regulator in phosphate signaling and PhoU Ala147 and Arg148 residues are required for this activity by interacting with PhoR, possibly via the PAS domain.

Given that the PhoU-PhoR interaction was recapitulated with the PhoR PAS domain alone in E. coli (5), one can expect that this is also true for Salmonella. However, the PAS domain of PhoR alone was not sufficient for interacting with PhoU when several T25-PhoR subclones were tested to identify the region(s) required for PhoU interaction (Fig. S2). Only the HK domain exhibited a weak interaction when we incubated for 60 h (Fig. S2D). This suggests that the PhoU-PhoR interaction may require additional residues or regions.

A region that includes the α-helix 6 of PhoU is also required for PhoR interaction. (A) Schematic representation of the Salmonella PhoU protein. Numbers correspond to the positions of each helix in the full-length PhoU sequence (aa 1 to 241). (B) Bacterial two-hybrid assay between PhoR and PhoU or PhoU variant proteins. Escherichia coli BTH101 strains harboring two plasmids (pKT25 and pUT18C derivatives) expressing an N-terminal fusion of the cyaA T25 fragment to the coding region of the phoR gene and N-terminal fusions of the cyaA T18 fragment to the full-length phoU (PhoU), phoU1–44 (α1), phoU1–77 (α2), phoU1–119 (α3), phoU1–147 (α4), phoU1–181 (α5), or phoU1–218 (α6), or the pUT18C empty vector (negative) were spotted as indicated. Cells expressing both pUT18C-mgtC and pKT25-mgtR are spotted as a positive control (37). Cells were spotted onto LB plates containing 80 μM X-Gal and 0.1 mM IPTG and incubated at 30°C for 40 h. Blue-colored colonies indicate a positive interaction. (C) β-Galactosidase assay from strains listed above. The average β-galactosidase activities (Miller units) are shown as mean ± SD (n = 3, independent measurements).
and a T18-PhoU derivative that harbors the coding region up to the end of α-helix 6 (α6, aa 1 to 218), retained the ability to interact with T25-PhoR (Fig. 3B and C). In contrast, all other T18-PhoU derivatives that harbor the coding regions up to α-helix 5 (α5, aa 1 to 181) or less (α1 to α4) lost the ability to interact with PhoR (Fig. 3B and C). These data suggest that a near full length of the PhoU protein is required for PhoR interaction, and the region containing and neighboring α-helix 6 (aa 182 to 218; Fig. S3A) is specifically required for this interaction.

Arg184 in the α-helix 6 of PhoU is required to interact with PhoR histidine kinase. The region corresponding to amino acids 182 to 218 includes the α5-α6 loop and α-helix 6 itself (Fig. S3). We suspected that an additional interaction between PhoU and PhoR might occur near the α5-α6 loop or at the beginning of α-helix 6. This is because Ala147 and Arg148 in the α4-α5 loop are involved in PhoR interaction via the PAS domain (Fig. S3), and, based on structural modeling, it seems unlikely that α-helix 6 is involved in interacting with the same PAS domain, considering that most of α-helix 6 residues are far from the PhoU-PhoR PAS interface (Fig. S4A, top view).

Based on our structural modeling, we selected Glu181, Asp182, and Arg184 residues to test for additional PhoR interactions (Fig. S3 and S4). As control experiments, we also selected Gln120 and Leu124 residues outside α-helix 6 because Gln120 and Leu124 residues are in the α3-α4 loop and are also in close proximity to those selected residues in the α5-α6 loop (Fig. S3). Among them, the Arg184 substitution to Gly completely lost the ability to interact with T25-PhoR (Fig. 4 and Fig. S3), indicating that the Arg184 residue in the α-helix 6 is required for PhoR interaction. We further demonstrated

(PhoU, amino acids [aa] 1 to 231) and a T18-PhoU derivative that harbors the coding region up to the end of α-helix 6 (α6, aa 1 to 218), retained the ability to interact with T25-PhoR (Fig. 3B and C). In contrast, all other T18-PhoU derivatives that harbor the coding regions up to α-helix 5 (α5, aa 1 to 181) or less (α1 to α4) lost the ability to interact with PhoR (Fig. 3B and C). These data suggest that a near full length of the PhoU protein is required for PhoR interaction, and the region containing and neighboring α-helix 6 (aa 182 to 218; Fig. S3A) is specifically required for this interaction.

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FIG 4 The PhoU Arg184 residue is required for PhoR interaction. (A) Bacterial two-hybrid assay between PhoR and PhoU or PhoU variant proteins. Escherichia coli BTH101 strains harboring two plasmids, pUT18c and pKT25 derivatives, expressing an N-terminal fusion of the cyaA T25 fragment to the phoR gene and N-terminal fusions of the cyaA T18 fragment to the wild-type phoU, phoU<sup>R184A</sup>, phoU<sup>R184G</sup> genes, or the pUT18C empty vector (negative) were spotted as indicated. Cells expressing both pUT18-mgt<sub>C</sub> and pKT25-mgt<sub>R</sub> are spotted as a positive control (37). Cells were spotted onto LB plates containing 80 μM X-Gal and 0.1 mM IPTG and incubated at 30°C for 40 h. Blue-colored colonies indicate a positive interaction. (B) β-Galactosidase assay from strains listed above. The average β-galactosidase activities (Miller units) are shown as mean ± SD (n = 3, independent measurements). (C and D) The C-terminally 8×Myc-tagged PhoR protein immunoprecipitates PhoU-His. (C) Crude extracts prepared from Salmonella strains with the phoR-8×Myc gene expressing PhoU-His, PhoU<sup>R184A</sup>-His, PhoU<sup>R184G</sup>-His, or the empty vector (pBAD33) were detected with anti-Myc (top) and anti-His (bottom) antibodies. (D) Eluted fractions prepared from strains listed above were detected with anti-Myc (top) and anti-His (bottom) antibodies after immunoprecipitation with anti-Myc antibody-coated beads.
the requirement of the PhoU Arg184 residue for PhoR interaction by immunoprecipitation. The C-terminally Myc-tagged PhoR protein successfully immunoprecipitated the C-terminally His-tagged PhoU protein (Fig. 4). In contrast, the PhoR-GFP protein failed to immunoprecipitate a PhoU-His derivative with the Arg184-to-Ala or -Gly substitution (Fig. 4), supporting the requirement of Arg184 residue in PhoU-PhoR interaction. We could even recapitulate this interaction with the C-terminally green fluorescent protein (GFP)-tagged HK domain of PhoR even though the HK domain-PhoU interaction was much weaker than the interaction between full-length PhoR and PhoU (Fig. S2F and G). Please note that the Gln120-to-Ala substitution in PhoU also abolished the interaction between PhoU and PhoR (Fig. S3), but we did not follow the substitution further because the Gln120-to-Ala substitution also disrupted the interaction with the PstB ATPase, another PhoU-interacting protein in phosphate signaling (Fig. S5).

The PhoU Arg184 residue is required for MgtC-mediated activation of pho genes but not low Pm-mediated activation. Protein modeling suggests that the Arg184 residue is located on the opposite side of the Ala147 and Arg148 residues in the PhoU protein (Fig. 2B and Fig. S4). Given that the Ala147 and Arg148 residues in PhoU interact with PhoR possibly via the PAS domain, Arg184 appears to be in close proximity to the CA domain of the PhoR protein (Fig. S4). This configuration suggests that PhoU Arg184 might have contact with the CA domain in PhoR and also have a regulatory role in the PhoR histidine kinase-mediated signaling. To understand the physiological role of the PhoU-PhoR interaction via the PhoU Arg184 residue, we created a chromosomal mutant strain with the Arg184-to-Gly substitution in PhoU-PhoR interaction via the PhoU Arg184 residue. We then tested the behaviors of the mutant in phosphate signaling. In the wild type, mRNA levels of the phoE gene were repressed in high-phosphate media but highly elevated in low-phosphate media (Fig. 5). Contrary to our expectation, the PhoU Arg184-to-Gly substitution did not affect the expression patterns of phoE mRNA levels in either high- or low-phosphate media (Fig. 5). As a control experiment, phoU deletion derepressed phoE mRNA levels both in high- and low-phosphate media.

Because the phoU Arg184-to-Gly substitution had no effect on PhoB/PhoR-mediated phosphate signaling despite disrupting the PhoU-PhoR interaction, we tried to look for other conditions that could be affected by the PhoU Arg184-to-Gly substitution. In a previous study, the Salmonella MgtC virulence factor also activates the PhoB-dependent genes independently of available phosphate concentration (16). MgtC activates PhoR histidine kinase by interacting with the CA domain (Fig. 1), which coincides with the potential contact site of PhoU Arg184 (Fig. 1 and Fig. S4). This suggests that the PhoU Arg184 residue might affect MgtC-mediated PhoR control. To test this, Salmonella strains with the wild-type phoU gene or the Arg184 to Gly-substituted phoU gene were grown in N-minimal medium with low Mg2+ to induce mgtC expression from the PhoP/PhoQ-dependent promoter (21) (Fig. 5C). As a control, the same strains were grown in N-minimal medium with high Mg2+ to repress mgtC expression. In the wild type, expression of the phoE gene was highly induced in low Mg2+ but repressed in high Mg2+ (Fig. 5), and the low Mg2+-mediated phoE induction is MgtC dependent because the mgtC deletion strain failed to increase in low Mg2+. The phoU Arg184-to-Gly substitution completely lost the ability to induce phoE expression, similar to that observed in the mgtC mutant (Fig. 5), indicating that PhoU is required for MgtC-mediated PhoR activation via the Arg184 residue. As a control, the phoU deletion mutant showed elevated phoE mRNA levels in all tested conditions (Fig. 5).

This effect was further confirmed by introducing the mgtC gene heterologously expressed from an IPTG (isopropyl-β-D-thiogalactopyranoside)-inducible promoter. In the wild type, mgtC expression from an IPTG-inducible promoter increased phoE mRNA levels (Fig. 6). However, the chromosomal phoU mutant strains with Arg184-to-Ala or -Gly substitutions exhibited reduced phoE mRNA levels despite high levels of mgtC expression (Fig. 6), indicating that PhoU Arg184 is required for MgtC-mediated phoE expression.

Given that both PhoU and PhoR histidine kinase belong to PhoB/PhoB-controlled genes, we created strains with the C-terminally FLAG-tagged phoU gene or C-terminally 8×Myc tagged phoR gene at its chromosomal locations and measured PhoU or...
PhoR protein levels in the presence of the **phoU** Arg184-to-Ala or -Gly substitutions. In low Pi, both PhoU and PhoR protein levels increased compared to high Pi and were unaffected by the **phoU** Arg184-to-Ala or -Gly substitutions (Fig. 7). However, similar to what we detected in **phoE** mRNA levels, the **phoU** Arg184-to-Ala or -Gly substitutions exhibited a defect in low Mg\(^{2+}\)-mediated induction of PhoU and PhoR protein levels (Fig. 7), supporting a notion that PhoU Arg184 residue is required for MgtC-mediated PhoR activation, possibly via interacting with the HK domain of PhoR.

Next, we determined epistasis between the **phoU** Arg184-to-Gly and the above-mentioned **phoU** Ala147-to-Glu substitutions. To test this, we created a strain where both Ala147 and Arg184 codons in the **phoU** gene were substituted by Glu and Gly codons, respectively. Then, we investigated the expression behavior of the A147E- and R184G-substituted **phoU** mutant in low phosphate or low Mg\(^{2+}\). The expression patterns of the A147E- and R184G-substituted **phoU** mutant were identical to those of the **phoU** A147E mutant in all tested conditions (Fig. S6), indicating that the **phoU** Ala147-to-Glu substitution is dominant over the **phoU** Arg184-to-Gly substitution.

**FIG 5** The **phoU** Arg184-to-Gly substitution prevents a low Mg\(^{2+}\)-mediated increase in **phoE** mRNA levels but does not affect low Pi-mediated **phoE** induction. (A to C) Relative mRNA levels of the **phoE** (A), **phoU** (B), and **mgtC** (C) genes in *Salmonella* strains with the wild-type **phoU** gene (14028s) or **Arg184 to Gly**-substituted **phoU** (SM235) genes and strains that deleted the **phoU** gene (SM101) or the **mgtC** gene (EL4). Bacteria were grown for 5 h in N-minimum media containing 10 mM Mg\(^{2+}\) and 10 mM Pi (high Mg\(^{2+}\), high Pi), 10 mM Mg\(^{2+}\) and 0.01 mM Pi (high Mg\(^{2+}\), low Pi), or 0.01 mM Mg\(^{2+}\) and 10 mM Pi (low Mg\(^{2+}\), high Pi). Shown are the means \(\pm\) SD (n = 3, independent measurements). Relative mRNA levels represent (target RNA/rrsH RNA) \(\times\) 10,000.
The PhoU Arg184 residue is required for MgtC-mediated PhoR autophosphorylation. MgtC binds to the Leu421 residue of the PhoR CA domain and promotes autophosphorylation, thus increasing expression of PhoB-dependent genes (16). Because the PhoU Arg184 residue is required for the MgtC-mediated increase in phoE mRNA
levels, we wondered whether the PhoU Arg184 residue is also involved in PhoR autophosphorylation when MgtC is present. We grew cells in low Mg\(^{2+}\) to increase MgtC production and isolated membrane fractions to measure PhoR autophosphorylation. When membrane fractions isolated from wild-type cells were incubated with \(\gamma\)-radiolabeled ATP, phosphorylated PhoR proteins were detected from 30 s and strongly increased up to 30 min (Fig. S7). However, the \(\textit{phoU}\) Arg184-to-Gly substitution delayed the appearance of phosphorylated PhoR and also exhibited low levels of phospho-PhoR (Fig. S7), suggesting that the PhoU Arg184 residue is involved in MgtC-mediated PhoR autophosphorylation. As a control, the \(\textit{phoU}\) Arg184-to-Gly substitution did not affect the behavior of PhoR autophosphorylation when grown in low phosphate (Fig. S7), and the levels of phospho-PhoR of both strains were low in a PhoB- and PhoP-repressing (10 mM Pi and 10 mM Mg\(^{2+}\)) condition (Fig. S7).

Because the \(\textit{phoU}\) Arg184-to-Gly substitution decreased PhoU and PhoR protein levels in low Mg\(^{2+}\) due to its autoregulation (Fig. 7), the autophosphorylation defect of the \(\textit{phoU}\) Arg184-to-Gly mutant in low Mg\(^{2+}\) might be due to low levels of PhoU and PhoR proteins. To avoid this, we also measured autophosphorylation of \textit{Salmonella} strains that were grown in high Mg\(^{2+}\) to produce similar levels of PhoR and PhoU proteins (Fig. 7) and expressing MgtC protein from a plasmid. Consistently to what we detected in \(\textit{phoE}\) and \(\textit{phoU}\) mRNA levels, wild-type \textit{Salmonella} expressing MgtC protein from the plasmid increased the rate of PhoR autophosphorylation compared to those expressing the empty vector (Fig. 6D and E). However, MgtC expression from the \(\textit{phoU}\)
Arg184-to-Ala or -Gly substitution mutants did not affect the rate of PhoR autophosphorylation (Fig. 6G and I), indicating that the PhoU Arg184 residue is indeed required for MgtC-mediated PhoR autophosphorylation.

The PhoU Arg184 residue is required for Salmonella virulence. Given that the phoU Arg184-to-Gly substitution affects MgtC-mediated phoE expression (Fig. 5) and that the mgtC gene is required for intramacrophage survival (22, 23) (Fig. 8A), we wondered whether PhoU Arg184-mediated PhoR autophosphorylation is required for Salmonella survival inside macrophages. When we infected Salmonella into the macrophage-like cell line J774A.1, the phoU variant with the Arg184-to-Gly substitution decreased replication within macrophages, reaching only 20% of that of the wild type (Fig. 8A). This is almost as defective as a mutant strain with the phoU deletion, suggesting that the Arg184 residue largely contributes to PhoU’s role in Salmonella pathogenesis. Similarly, when we injected ~3,000 CFU of Salmonella strains intraperitoneally into C3H/HeN mice, the Salmonella strain with the phoU Arg184-to-Gly substitution attenuated mouse virulence compared to the wild type (Fig. 8B). As controls, strains that deleted either the phoU or mgtC genes were completely defective in mouse virulence. Collectively, these data indicate that the PhoU Arg184 residue is involved in Salmonella virulence, possibly by controlling the expression of pho regulon genes via PhoR autophosphorylation.

DISCUSSION

Using on model-based substitutions, we determined that the PhoU regulator binds to PhoR histidine kinase via the HK domain. Because the interaction between PhoU and the PAS domain of PhoR histidine kinase was already established in E. coli (5), this suggests that PhoU could interact with PhoR histidine kinase in at least two different sites, the PAS domain and the CA domain (Fig. 1). Substitution of Ala147 to Glu in the Salmonella PhoU protein, which likely prevents PhoU-PhoR PAS domain interaction, derepressed phoE mRNA levels in high phosphate, in agreement with a previous finding in E. coli that PhoU functions as a negative regulator via interaction with the PhoR PAS domain (5). Constitutively elevated phoE expression in the phoU Ala147-to-Glu

![FIG 8](image-url) PhoU Arg184-mediated PhoR autophosphorylation is required for Salmonella virulence. (A) Arg184-to-Gly substitution in PhoU decreases survival inside macrophages. Survival inside J774A.1 macrophage cells of the wild-type Salmonella (14028s), the phoU chromosomal mutant with Arg184 replaced by the Gly codon (SM235), the phoU deletion mutant (SM101), and the mgtC deletion mutant (EL4) at 21 h postinfection (T21). Fold replication represents the number of bacteria at T21/number of bacteria at T1. Shown are the means and SD from three independent infections. (B) Arg184-to-Gly substitution in PhoU decreases mouse virulence. Survival of C3H/HeN mice inoculated intraperitoneally with ~10^4 CFU of the Salmonella strains listed above.
substitution suggests that PhoU’s binding to the PhoR PAS domain suppresses autophosphorylation activity, probably by inducing a conformation change. However, repression of phoE expression could also be achieved by either preventing phosphate transfer from PhoR to PhoB or stabilizing phosphatase activity toward phosphorylated PhoB protein. Given that the phosphatase activity and the transition between kinase and phosphatase activities of Salmonella PhoR histidine kinase have not been carefully examined yet, it needs to be investigated to understand a mechanism underlying PhoU-mediated inhibition of PhoR activity.

Interestingly, the introduction of the phoU Arg184-to-Gly substitution that disrupts the interaction between PhoU and PhoR HK domain did not affect phoE mRNA levels in high- or low-phosphate conditions (Fig. 5), suggesting that the interaction between the PhoU-PhoR HK domain is not involved in repressing expression of pho regulon in high phosphate. Instead, PhoU’s interaction with the PhoR HK domain seems to be required for inducing expression of the pho regulon when Salmonella is within macrophages because the phoU Arg184-to-Gly substitution failed to induce phoE mRNA levels under the conditions when the MgtC virulence factor was highly expressed (Fig. 5 and 6). Given that the PhoU protein exerts its opposing effects on expression of the pho regulon by interacting with PhoR histidine kinase at two different sites, PhoU should be considered a modulator/molecular adaptor of pho expression rather than simply a negative regulator. By creating amino acid substitutions instead of the phoU deletion, we could dissect PhoU’s regulatory roles in repressing pho expression in high phosphate and in activating pho expression in the presence of the MgtC virulence factor that is highly expressed in low Mg2+ or during Salmonella infection (24).

PhoU’s function as a molecular adaptor was also described in a previous report (13). In E. coli, when both phosphate and potassium are limiting, the phosphate-sensing PhoB/PhoR two-component system also controls a counterion signaling pathway via the potassium-sensing KdpE/KdpD two-component system (13). In this case, PhoU seems to connect these two regulatory pathways given that PhoU interacts with both KdpD and PhoR histidine kinases in the bacterial two-hybrid assay and phoU deletion derepresses KdpE-dependent expression of KdpFABC K+ transporter independent of K+ concentration in the kdpD deletion background. However, its physiological relevance remains unclear, given that the effect of PhoU on potassium signaling can be clearly seen in a strain lacking the KdpD sensor kinase (13). Similar to PhoU, nonphosphorylated EI*AIII, a nitrogen PTS component, is also involved in K+ homeostasis in E. coli via protein-protein interaction. However, instead of connecting two regulatory pathways, monophosphorylated EI*AIII controls K+ homeostasis differently depending on available K+ concentration. In low K+, EI*AIII binds to KdpD histidine kinase and activates KdpE-dependent expression of KdpFABC high-affinity K+ transporter (25). In high K+, EI*AIII directly binds to TrkA, a peripheral membrane protein of Trk low-affinity K+ transporter, and inhibits its activity (26), which is a major K+ transporter at high K+ conditions.

Previously, the MgtC virulence protein activated autophosphorylation of PhoR histidine kinase by directly binding to the PhoR CA domain and promoted expression of pho genes (16). The Leu421 residue in the PhoR CA domain was determined as a key residue required for this interaction. Because MgtC stimulates PhoR autophosphorylation independently of available phosphate concentration, it was unclear whether PhoU still binds to PhoR or participates in MgtC-mediated activation of PhoR histidine kinase. The fact that the PhoU Arg184 residue is also required for interacting with the PhoR HK domain (possibly via the CA domain) and stimulating MgtC-mediated PhoR autophosphorylation led us to speculate that the Salmonella MgtC virulence protein might activate pho expression via three molecular interactions between MgtC, PhoU, and PhoR histidine kinase. However, curiously enough, MgtC did not directly interact with PhoU in a previous two-hybrid study (16), despite both PhoR and PhoU proteins being required for MgtC-mediated activation of PhoR autophosphorylation. Thus, it seems that PhoR histidine kinase makes contact with both MgtC and PhoU proteins via its CA
domain but accommodates these proteins at different surface areas, which needs further attention to understand how these molecular interactions occur. Finally, by creating substitution mutants that enable us to address a detailed function, we uncovered an unexpected role of PhoU, which was previously hidden by an effect of gene deletion. This finding will allow us to understand a detailed molecular mechanism of phosphate signaling in bacteria.

MATERIALS AND METHODS

Bacterial strains, plasmids, oligodeoxynucleotides, and growth conditions. Bacterial strains and plasmids used in this study are listed in Table S1 in the supplemental material. All S. enterica serovar Typhimurium strains were derived from the wild-type strain 14028s (27) and were constructed by the one-step gene inactivation method (28) and/or P22-mediated transduction as previously described (29). DNA oligonucleotides are listed in Table S1. Bacteria were grown at 37°C in LB-Bertani (LB) broth and N-minimal media (30) supplemented with 0.1% Casamino Acids, 38 mM glycerol, and the indicated concentrations of MgCl₂. For low-phosphate N-minimal media, 10 mM KH₂PO₄ in the N-minimal medium was replaced by 0.01 mM KH₂PO₄. E. coli DH₅₅C was used as the host for preparing plasmid DNA, and BTH101 lacking the cya gene was used as the host for the bacterial two-hybrid system (31). Ampicillin was used at 50 μg mL⁻¹, chloramphenicol at 20 μg mL⁻¹, kanamycin at 20 μg mL⁻¹, tetracycline at 10 μg mL⁻¹, and fusaric acid (32) at 12 μg mL⁻¹. IPTG (isopropyl β-D-1-thiogalactopyranoside) was used at 0.25 mM, L-arabinose at 0.2% (wt/vol), and X-Gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) at 80 μM.

Bacterial two-hybrid assay. To assess protein-protein interactions in vivo, a bacterial two-hybrid (BACTH) assay was conducted as described (31). The Escherichia coli strain BTH101 lacking the cya adenylate cyclase gene was cotransformed with derivatives of the pUT18, pUT18c, and pK725 plasmids. The strains were grown overnight at 37°C in LB supplemented with ampicillin (50 μg mL⁻¹) and kanamycin (50 μg mL⁻¹). Then, 4 μL of cells were spotted on solid LB medium with 500 μM IPTG, 100 μM ampicillin, 100 μM kanamycin, and 80 μM X-Gal, followed by incubation for 40 h or 60 h at 30°C. (For quantitaive analysis, β-galactosidase assays were performed (33).)

Protein structure modeling and protein docking modeling. We used Protein Homology/analogy Recognition Engine v2.0 (Phyre2) (34) to model the structures of the cytoplasmic portion of the PhoR protein (corresponding to amino acids 52 to 431) and the full-length of the PhoU protein from Salmonella enterica serovar Typhimurium 14028s. The sequence of the PhoR protein was modeled based on the structure of Vich from Streptococcus mutans (PDB ID 4155), and the structure of the PhoU protein was modeled based on the structure of the PhoU homolog 2 from Thermotoga maritima (PDB ID 1SU4) or PhoU from Pseudomonas aeruginosa (PDB ID 4Q2S). Then, we used the ClusPro web server to investigate a potential interaction between Salmonella PhoR and PhoU proteins (35).

Immunoprecipitation assay. The interaction between the PhoU and PhoR proteins was investigated in wild-type Salmonella expressing the phoU gene or its derivatives from an arabinose-inducible promoter (pBAD33-phoU-His, pBAD33-phoU[Arg184Gly]-His, or pBAD33-phoU[Arg184Ala]-His) in the phoR-8×Myc background (phoR-8×Myc) (36). Cells were grown overnight in N-minimal medium containing 10 mM MgCl₂. A 1:100 dilution of the overnight grown bacterial culture was inoculated in 15 mL of N-minimal medium containing 10 mM MgCl₂ and grown for 3 h. Cells were then washed and transferred to 15 mL of N-minimal medium containing 0.5 mM MgCl₂ and 1 mM L-arabinose and grown for 1 h. Cells were normalized by measuring the optical density at 600 nm (OD₆₀₀). Crude extracts were prepared in Tris-buffered saline (TBS) buffer by sonication. For a pulldown assay with anti-GFP antibodies, 50 μL of the crude extracts were kept for input, and 400 μL of the protein extracts were mixed with 2 μL of anti-Myc (MBL Life Science) and 40 μL of protein G Sepharose 4 Fast Flow (Cytiva; catalog no. 17-0618-01) for 2 h at 4°C on a nutator (Benchmark Scientific). Beads were washed with TBS washing buffer three times, and then the bound proteins were eluted in SDS sample buffer. The eluates were resolved on 12% SDS-polyacrylamide gels, transferred to nitrocellulose membrane, and analyzed by Western blotting using anti-His (1:20,000 dilution; Rockland; catalog no. 600-401-382) and anti-Myc (1:3,000 dilution; MBL Life Science; catalog no. M192-3) antibodies overnight. The blots were washed and hybridized with anti-mouse IgG horseradish peroxidase-linked whole antibody (1:10,000 dilution; Amersham; catalog no. NA931) for 1 h and detected using SuperSignal West Femto maximum sensitivity substrate (Thermo Fisher). To investigate the interaction between PhoU and the HK domain of the PhoR protein, Salmonella strains with the C-terminally FLAG-tagged phoU gene and its derivatives expressing the C-terminally gfp-tagged HK domain of the phoR gene (pTGFP-phoR₆₉) or the empty vector (pTGFP) were used, and an immunoprecipitation assay was performed. The pulled-down PhoR₆₉-GFP proteins and the immunoprecipitated PhoU-FLAG proteins were detected with anti-GFP (1:1,000 dilution; Roche; catalog no. 1181446001) and anti-FLAG (1:3,000 dilution; Millipore; catalog no. F7425) antibodies, respectively. For secondary antibody, anti-mouse IgG horseradish peroxidase (HRP)-linked whole antibody (1:10,000 dilution; Amersham; catalog no. NA931) and anti-rabbit IgG horseradish peroxidase-linked antibody (1:10,000 dilution; Thermo Fisher; catalog no. 31460) were used to detect PhoR₆₉-GFP and PhoU-FLAG proteins, respectively.

Membrane vesicle preparation. Cells were for 5 h in N-minimal media containing combinations of 10 mM (high) or 0.01 mM (low) MgCl₂ and 10 mM (high) or 0.01 mM (low) Pi. Cells were normalized by measuring OD₆₀₀. Crude extracts were prepared in TBS buffer by sonication and centrifuged supernatants for 2 h at 40,000 × g (Optima TLX ultracentrifuge, type 90 Ti rotor; Beckman Coulter). The pellets
were resuspended in 50 μL TBS buffer. The protein concentration in the prepared membrane fractions was determined using a NanoDrop machine (Thermo Fisher).

**Measuring autophosphorylation of PhoR histidine kinase.** As previously described (16), 50 μL of membrane vesicles expressing wild-type PhoU or PhoU Arg184Gly were incubated in 100 μL of TBS containing 1 mM MgCl2 at room temperature. The reaction was started with the addition of [γ-32P]ATP (10 μCi, PerkinElmer) to the mixture. A 10-μL aliquot was taken at different time points and mixed with 10 μL of 5× SDS loading buffer (Biosesang) to stop the reaction. The samples were kept on ice until they were loaded onto a 12% SDS-polyacrylamide gel. After electrophoresis, the gel was dried on Whatman filter paper using model 583 gel dryer (Bio-Rad), and the phosphorylated PhoR proteins were visualized by a phosphoimaging on a Typhoon scanner (GE Healthcare). The phosphorylated PhoR proteins were identified using samples prepared from wild-type and the phoR mutant Salmonella grown for 5 h in N-minimal medium containing 0.01 mM Pi, a PhoB/PhoR-inducing condition.

**Quantitative real-time PCR.** Total RNA was isolated using RNeasy kit (Qiagen) according to the manufacturer’s instructions. The purified RNA was quantified using a NanoDrop machine (Thermo Fisher). cDNA was synthesized using PrimeScript RT reagent kit (TaKaRa). The mRNA levels of each target gene were calculated using a standard curve of 14028s genomic DNA with known concentration, and data were normalized to the levels of 16S rRNA ampli-

**Western blot analysis.** Cells were grown for 5 h in 15 mL of N-minimal medium containing combinations of 10 mM or 0.01 mM MgCl2 and 10 mM or 0.01 μM P. Cells were normalized by measuring the OD600. Crude extracts were prepared in TBS by sonicating. Whole-cell lysates were resolved on 12% SDS-polyacrylamide gels and were transferred onto nitrocellulose membranes. The blots were incubated with monoclonal anti-FLAG antibodies (1:3,000 dilution; Millipore; catalog no. F7425) and anti-Myc antibodies (1:3,000 dilution; Millipore; catalog no. M192-3) overnight to detect PhoU-FLAG and PhoR-8×Myc proteins, respectively. The blots were developed by incubation with anti-rabbit IgG horseradish peroxidase-linked antibody (Ab) (1:10,000 dilution; Thermo Fisher; catalog no. 31460) and anti-mouse IgG HRP-linked whole Ab (1:10,000 dilution; Amersham; catalog no. NA931) for 1 h and were visualized using SuperSignal West Femto maximum-sensitivity substrate (Thermo Fisher).

**Macrophage survival assay.** Intramacrophage survival assays were conducted using the macrophage-like cell line J774 A.1. Briefly, 5 × 106 macrophages in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS) were seeded in 24-well plates and cultured at 37°C. Overnight grown bacteria were added to the macrophages at a multiplicity of infection (MOI) of 10. The plates were centrifuged at 1,000 rpm for 5 min at room temperature and incubated for an additional 20 min. Then, the extracellular bacteria were washed three times with phosphate-buffered saline (PBS) and killed by incubation with DMEM supplemented with 10% FBS and 120 μg mL−1 gentamicin for 1 h. For measuring the number of bacteria at 1 h, the DMEM was replaced after 1 h with fresh DMEM containing 0.1% Triton X-100 and plated on Luria-Bertani (LB) plates with appropriate dilutions. For measuring the number of bacteria at 21 h, the DMEM was replaced after 1 h with fresh DMEM containing 12 μg mL−1 gentamicin, and the incubation was continued at 37°C. After 21 h, cells were lysed with PBS containing 0.1% Triton X-100 and plated on LB plates. The percentage survival was obtained by dividing the number of bacteria recovered after 21 h by the number of bacteria recovered at 1 h. All experiments were performed in triplicate, and the results are representative of at least three independent experiments.

**Mouse virulence assay.** Six- to 8-week-old female C3H/HeN mice were inoculated intraperitoneally with ~105 CFU of Salmonella strains. Mouse survival was followed for 21 days. Virulence assays were conducted three times with similar outcomes, and the data correspond to groups of five mice. All animals were housed in a temperature- and humidity-controlled room, in which a 12-h light/12-h dark cycle was maintained. All procedures were performed according to the protocols (KW-181010-1) approved by the institutional Animal Care and Use Committee of the Kangwon National University.

**Data availability.** All other relevant data are available from the corresponding author upon reasonable request.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**TEXT S1**, DOCX file, 0.04 MB.

**FIG S1**, TIF file, 1.8 MB.

**FIG S2**, TIF file, 1.6 MB.

**FIG S3**, TIF file, 1.2 MB.

**FIG S4**, TIF file, 2 MB.

**FIG S5**, TIF file, 0.9 MB.

**FIG S6**, TIF file, 0.9 MB.

**FIG S7**, TIF file, 0.7 MB.

**TABLE S1**, DOCX file, 0.1 MB.
REFERENCES

1. Nelson DL, Cox MM, Lehninger AL. 2017. Lehninger principles of biochemistry, 7th ed. W.H. Freeman and Company, Hingham, Basingstoke: Macmillan Higher Education, New York, NY.

2. Hsieh YJ, Wanner BL. 2010. Global regulation by the seven-component Pi signaling system. Curr Opin Microbiol 13:198–203. https://doi.org/10.1016/j.mib.2010.01.014.

3. Lamarche MG, Wanner BL, Crepin S, Harel J. 2008. The phosphate regulon. A regulatory network connecting phosphate homeostasis and pathogenesis. FEMS Microbiol Rev 32:461–473. https://doi.org/10.1111/j.1574-6976.2008.00301.x.

4. Piepenbreier H, Fritz G, Behrard S. 2017. Transports as information processors in bacterial signalling pathways. Mol Microbiol 104:1–15. https://doi.org/10.1111/mmi.13633.

5. Gardner SG, Johns KD, Tanner R, McCleary WR. 2014. The PhoU protein from Escherichia coli interacts with PhoR, PstB, and metals to form a phos- phase-signaling complex at the membrane. J Bacteriol 196:1741–1752. https://doi.org/10.1128/JB.00029-14.

6. Gardner SG, Miller JB, Dean T, Robinson T, Erickson M, Ridge PG, McCleary WR. 2015. Genetic analysis, structural modeling, and direct coupling analysis suggest a mechanism for phosphate signaling in Escherichia coli. BMC Genet 16(Suppl 2):S2. https://doi.org/10.1186/1471-2156-16-S2-S2.

7. Hefti MH, Francois KJ, de Vries SC, Dixon R, Vervoort J. 2004. The PAS fold. A redefinition of the PAS domain based upon structural prediction. Eur J Biochem 271:1198–1208. https://doi.org/10.1111/j.1432-1033.2004.04023.x.

8. Ponting CP, Aravind L. 1997. PAS: a multifunctional domain family comes to light. Curr Biol 7:R674 –R687. https://doi.org/10.1016/s0960-9822(06)00352-6.

9. Steed PM, Wanner BL. 1993. Use of the rep allele for allele replacement to construct mutants with deletions of the pstSCAB-phoU operon: evidence of a new role for the PhoU protein in the phosphate regulon. J Bacteriol 175:6797–6809. https://doi.org/10.1128/JB.175.21.6797-6809.1993.

10. Lubin EA, Henry JT, Fiebig A, Crosson S, Laub MT. 2016. Identification of the Pho8 regulon and role of PhoU in the phosphate starvation response of Caulobacter crescentus. J Bacteriol 198:187–200. https://doi.org/10.1128/JB.00658-15.

11. de Almeida LG, Ortiz JH, Schneider RP, Spira B. 2015. PhoU inactivation in Pseudomonas aeruginosa enhances accumulation of ppGpp and poly- phospho. Appl Environ Microbiol 81:3006–3015. https://doi.org/10.1128/AEM.01648-15.

12. Wang X, Mao Y, Yu J, Zhu L, Li M, Wang D, Dong D, Liu J, Gao Q. 2013. PhoY2 of mycobacteria is required for metabolic homeostasis and stress response. J Bacteriol 195:243–252. https://doi.org/10.1128/JB.01556-12.

13. Schramek H, Laermann V, Tegetmeyer HE, Brachmann A, Jung K, Atendorf K. 2017. Revisiting regulation of potassium homeostasis in Escherichia coli: the connection to phosphate limitation. Microbiolgenoply 6:00438. https://doi.org/10.1002/mibp.438.

14. Gibson JL, Lombardo MU, Aponyi I, Vera Cruz D, Ray MP, Rosenberg SM. 2015. Appy, a role for PhoU in mutagenic break repair under stress in Escherichia coli. PLoS One 10:e0123315. https://doi.org/10.1371/journal.pone.0123315.

15. Eriksson S, Lucchini S, Thompson A, Rhen M, Hinton JC. 2003. Unravelling the biology of macrophage infection by gene expression profiling of intracellular Salmonella enterica. Mol Microbiol 47:103–118. https://doi.org/10.1046/j.1365-2958.2003.03313.x.

16. Choi S, Choi E, Cho YJ, Nam D, Lee J, Lee EJ. 2019. The Salmonella virulence protein MgtC promotes phosphate uptake inside macro- phages. Nat Commun 10:3326. https://doi.org/10.1038/s41467-019-11318-2.

17. Lee EJ, Groisman EA. 2012. Control of a Salmonella virulence locus by an ATP-sensing leader messenger RNA. Nature 469:271–275. https://doi.org/10.1038/nature10199.

18. Bruna RE, Kendra CG, Groisman EA, Pontes MH. 2021. Limitation of phos- phate assimilation maintains cytoplasmic magnesium homeostasis. Proc Natl Acad Sci U S A 118:e2021370118. https://doi.org/10.1073/pnas.2137011118.

19. Röder J, Felgner P, Hensel M. 2021. Single-cell analyses reveal phosphate availability as critical factor for nutrition of Salmonella enterica within mammalian host cells. Cell Microbiol 23:e13374. https://doi.org/10.1111/cmi.13374.

20. Muda M, Rao NN, Torriani A. 1992. Role of PhoU in phosphate transport and alkaline phosphate regulation. J Bacteriol 174:8057–8064. https://doi.org/10.1128/jb.174.24.8057-8064.1992.

21. Soncini FC, Garcia VE, Solomon F, Groisman EA. 1996. Molecular basis of the magnesium deprivation response in Salmonella typhimurium: identifi- cation of Pho-regulated genes. J Bacteriol 178:5092–5099. https://doi.org/10.1128/jb.178.17.5092-5099.1996.

22. Blanc-Potard AB, Groisman EA. 1997. The Salmonella secL locus contains a pathogenicity island mediating intramacrophage survival. EMBO J 16:5376–5385. https://doi.org/10.1093/emboj/16.17.5376.

23. Lee EJ, Pontes MH, Groisman EA. 2013. A bacterial virulence protein promotes pathogenicity by inhibiting the bacterium’s own F1Fo ATP synthase. Cell 154:146–156. https://doi.org/10.1016/j.cell.2013.06.004.

24. Smith RL, Kaczmarek MT, Kucharски LM, Maguire ME. 1998. Magnesium transport in Salmonella typhimurium: regulation of mgtA and mgtB during invasion of epithelial and macrophage cells. Microbiology 144:1835–1843. https://doi.org/10.1099/00221287-144-7-1835.

25. Luttmann D, Heermann R, Zimmer B, Hillmann A, Rampf IS, Jung K, Gorke B. 2009. Stimulation of the potassium sensor KdpD kinase activity by interaction with the phosphotransferase protein IIA(Ntr) in Escherichia coli. Mol Microbiol 72:978–994. https://doi.org/10.1111/j.1365-2958.2009.06704.x.

26. Lee CR, Cho SH, Yoon MJ, Peterkofsky A, Seok YJ. 2007. Escherichia coli enzyme IIA(Ntr) regulates the K+ transporter TrkA. Proc Natl Acad Sci U S A 104:4124–4129. https://doi.org/10.1073/pnas.0609897104.

27. Fields PI, Swanson RV, Haidar C, Jeffrey T, Hefron F. 1986. Mutants of Salmonella typhimurium that cannot survive within the macrophage are avirulent. Proc Natl Acad Sci U S A 83:5189–5193. https://doi.org/10.1073/pnas.83.14.5189.

28. Datseiko KA, Wanner BL. 2000. One-step inactivation of chromosomal genes in Escherichia coli K-12 using PCR products. Proc Natl Acad Sci U S A 97:6640–6645. https://doi.org/10.1073/pnas.062163297.

29. Davis RW, Bolstein D, Roth JR. 1980. Advanced bacterial genetics. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.

30. Snarey MD, Miller CG, Maguire ME. 1991. The mgtB Mg2+- transport locus of Salmonella typhimurium encodes a P-type ATPase. J Biol Chem 266:815–823. https://doi.org/10.1016/S0021-9258(17)35246-8.

31. Karimova G, Pidoux J, Ullmann A, Ladant D. 1998. A bacterial two- component system based on a reconstituted signal transduction pathway.
32. Maloy SR, Nunn WD. 1981. Selection for loss of tetracycline resistance by *Escherichia coli*. J Bacteriol 145:1110–1111. https://doi.org/10.1128/jb.145.2.1110-1111.1981.

33. Miller JH. 1972. Experiments in molecular genetics. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.

34. Kelley LA, Sternberg MJ. 2009. Protein structure prediction on the web: a case study using the Phyre server. Nat Protoc 4:363–371. https://doi.org/10.1038/nprot.2009.2.

35. Comeau SR, Gatchell DW, Vajda S, Camacho CJ. 2004. ClusPro: a fully automated algorithm for protein-protein docking. Nucleic Acids Res 32:W96–W99. https://doi.org/10.1093/nar/gkh354.

36. Lee EJ, Groisman EA. 2012. Tandem attenuators control expression of the Salmonella mgtCVR virulence operon. Mol Microbiol 86:212–224. https://doi.org/10.1111/j.1365-2958.2012.08188.x.

37. Alix E, Blanc-Potard AB. 2008. Peptide-assisted degradation of the Salmonella MgtC virulence factor. EMBO J 27:546–557. https://doi.org/10.1038/sj.emboj.7601983.