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HilE Regulates HilD by Blocking DNA Binding in *Salmonella enterica* Serovar Typhimurium

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ABSTRACT The *Salmonella* type three secretion system (T3SS), encoded in the *Salmonella* pathogenicity island 1 (SPI1) locus, mediates the invasion of the host intestinal epithelium. SPI1 expression is dependent upon three AraC-like regulators: HilD, HilC, and RtsA. These regulators act in a complex feed-forward loop to activate each other and *hilA*, which encodes the activator of the T3SS structural genes. HilD has been shown to be the major integration point of most signals known to activate the expression of the SPI1 T3SS, acting as a switch to control induction of the system. HilE is a negative regulator that acts upon HilD. Here we provide genetic and biochemical data showing that HilE specifically binds to HilD but not to HilC or RtsA. This protein-protein interaction blocks the ability of HilD to bind DNA as shown by both an *in vivo* reporter system and an *in vitro* gel shift assay. HilE does not affect HilD dimerization, nor does it control the stability of the HilD protein. We also investigated the role of HilE during the infection of mice using competition assays. Although deletion of *hilE* does not confer a phenotype, the *hilE* mutation does suppress the invasion defect conferred by loss of *FliZ*, which acts as a positive signal controlling HilD protein activity. Together, these data suggest that HilE functions to restrict low-level HilD activity, preventing premature activation of SPI1 until positive inputs reach a threshold required to fully induce the system.

IMPORTANCE *Salmonella* is a leading cause of gastrointestinal and systemic disease throughout the world. The SPI1 T3SS is required for *Salmonella* to induce inflammatory diarrhea and to gain access to underlying tissue. A complex regulatory network controls expression of SPI1 in response to numerous physiological inputs. Most of these signals impinge primarily on HilD translation or activity. The system is triggered when HilD activity crosses a threshold that allows efficient activation of its own promoter. This threshold is set by HilE, which binds to HilD to prevent the inevitable minor fluctuations in HilD activity from inappropriately activating the system. The circuit also serves as a paradigm for systems that must integrate numerous environmental parameters to control regulatory output.

KEYWORDS *Salmonella*, SPI1, HilD, HilE

Salmonella enterica serovar Typhimurium is a Gram-negative pathogen that, when ingested, is capable of causing a range of diseases from gastroenteritis to potentially lethal systemic infections in a wide range of hosts, including humans. It is a leading cause of bacterial foodborne illness in the United States, with nontyphoid species of *Salmonella* estimated to cause over 1 million infections per year (1). *Salmonella* is defined and differentiated from other *Enterobacteriaceae* by the presence of *Salmonella* pathogenicity island 1 (SPI1). This 40-kb locus carries all of the genes required for a functional type III secretion system (T3SS) and the regulators HilD, HilC, and HilA (2). The additional regulators RtsA (STM14_5188) and HilE (STM14_4514) are encoded elsewhere on the chromosome (3–5). The T3SS forms a needle-like complex that injects bacterial effector proteins into host epithelial cells. These effectors induce

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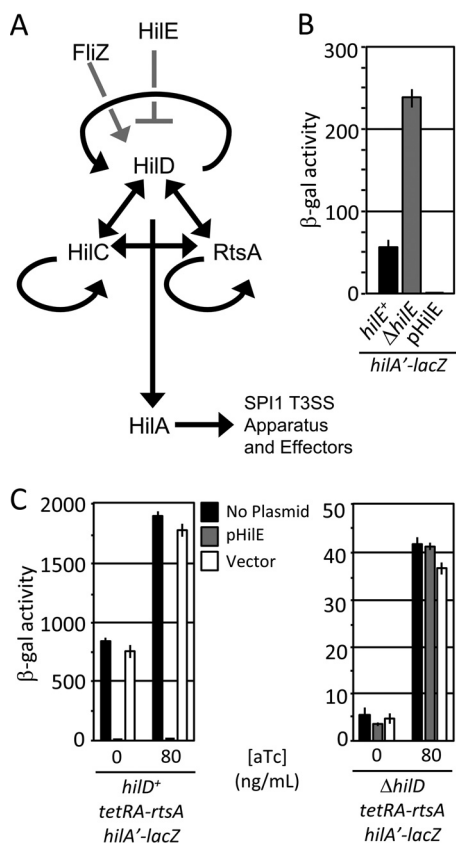


FIG 1 (A) A simplified model for SPI1-T3SS regulation. Black lines indicate transcriptional regulation. Gray lines indicate posttranslational regulation. For clarity, the genes encoding SPI1 regulators are not shown. (B) *HilE* is a negative regulator of SPI1. Strains with *hilE* deleted or overexpressing *hilE* were grown under SPI1-inducing conditions and analyzed via β -galactosidase assay. β -Galactosidase activity units are defined as (micromoles of ONP formed per minute $\times 10^6$)/(OD₆₀₀ \times milliliters of cell suspension) and reported as mean \pm one standard deviation ($n = 6$, 3 sets of duplicate assays on 3 separate days). The strains used were JS749, JS749 with p*hilE*, and JS2103. (C) *HilE* acts via *HilD*. The *rtSA* gene was under the control of a *tetA* promoter, *hilD* was left unaltered or deleted, and *HilE* was overexpressed as indicated. Strains were grown overnight under SPI1-inducing conditions and subcultured 1:100 with the indicated concentration of anhydrotetracycline (aTc) ($n = 4$, 3 sets of duplicate assays on 2 separate days; error bars represent one standard deviation). The strains used were JS953 and JS2104 with p*hilE* or pWKS30.

a number of physiological responses, including inflammatory diarrhea and bacterial engulfment that can lead to a systemic infection (6).

Expression of the SPI1 T3SS is controlled by a complex regulatory network (Fig. 1A). *HilA* functions to activate all of the genes required to synthesize a functional T3SS (7–10). Transcriptional activation of *hilA* is dependent upon three AraC-like proteins, *HilD*, *HilC*, and *RtsA*, which bind to the same sites upstream of activated promoters (11, 12). Each of the three regulators activates, in addition to *hilA*, its own promoter as well as those of the other two regulators, thus forming a complex feed-forward loop (12–15).

Our data indicate that the majority of regulatory inputs into the SPI1 system are integrated at the level of *HilD* (16). The overall effect is to control the ability of *HilD* to activate its own promoter, the critical step in initiating expression of the SPI1 T3SS (17). Primary examples of factors that influence *HilD* activity are *HilE*, a major negative regulator of SPI1 T3SS expression, and the flagellar gene product *FliZ*, which acts indirectly upon *HilD* to increase activity (10, 16, 18–20). *HilE* and *FliZ* regulate the activity of *HilD* simultaneously and independently (20). Our immediate goal was to understand how *HilE* regulates *HilD* activity at the molecular level to further our understanding of overall integration of signals into the SPI1 regulatory circuit.

Fahlen et al. identified *HilE* as a negative regulator of *hilA* transcription, and Baxter

et al. provided bacterial two-hybrid data indicating that HilE directly interacts with HilD protein (5, 21). We have also provided genetic data demonstrating that HilE operates at the level of the HilD protein (16, 20). HilE controls an ectopically expressed HilD but does not affect expression of *hilD-lacZ* transcriptional or translational fusions in the absence of HilD (20). In this study, we provide further genetic and biochemical data showing that HilE directly binds to HilD and blocks its ability to bind DNA but does not affect the dimerization or stability of HilD. Consistent with our previous genetic data, HilE is specific for HilD and does not bind or affect HilC or RtsA. *In vivo* data suggest that during infection, FlhZ acts to overcome the negative regulation of HilE to switch on SPI1 T3SS expression in the intestine.

RESULTS

HilE acts specifically upon HilD. Previous data suggest that HilE acts directly at the level of the HilD protein (5, 16, 20, 21). We sought to further understand the molecular mechanism by which HilE controls HilD protein activity. Figure 1B shows the effect of HilE on *hilA* expression. In the absence of HilE, *hilA* transcription was increased ~4-fold, whereas overproduction of HilE decreased *hilA* transcription by ~50-fold, consistent with previous data (20, 21). Our previous results showed that HilE does not regulate HilC or RtsA activity in the absence of HilD (16). To confirm this finding, we constructed a strain in which *rtsA* is placed under the control of the *tetRA* cassette. Otherwise isogenic variations of this strain included the deletion of *hilD* and/or overexpression of HilE, which should maximize any effect. Upon induction with anhydrotetracycline under SPI1-inducing conditions, we observed the effect of HilE only in the HilD-producing strain (Fig. 1C). Because these strains express both RtsA and HilC, we can conclude that HilE acts only through HilD, consistent with our previous experiments examining the effects of a *hilE* deletion (16).

HilE binds directly and specifically with HilD. To characterize the interaction between HilE and HilD, we first confirmed the results of Baxter et al. (21) using a bacterial two-hybrid system. In this system, IPTG (isopropyl- β -D-thiogalactopyranoside) induces the expression of a LexA DNA binding domain (LexA_{DBD}) fused to the C terminus of a protein of interest. If the protein of interest dimerizes, the LexA_{DBD} binds to its cognate *sulA* promoter, repressing the expression of a transcriptionally linked *lacZ* gene. Alternatively, a mutant LexA_{DBD} and a half-mutant *sulA* promoter can be used to test for heterodimerization of two proteins (21–23). We created *hilD-lexA_{DBD}* and *hilE-lexA_{DBD}* constructs. The data in Fig. 2A suggest that first, HilD forms homodimers, and second, HilE and HilD indeed interact via protein-protein interaction.

To verify these data and to determine if HilE is capable of interacting with either HilC or RtsA, we performed a coimmunoprecipitation assay using strains containing FLAG-tagged HilE and Myc-tagged HilD, HilC, or RtsA constructs. HilE was immunoprecipitated using anti-FLAG antibodies, and the coprecipitation of HilD, HilC, or RtsA was monitored via Western analysis. The data in Fig. 2B indicate that HilE specifically binds HilD but not HilC or RtsA *in vivo*. Both the two-hybrid and coimmunoprecipitation data are consistent with our genetic model (Fig. 1A) and previous data indicating that HilE controls HilD protein activity but does not affect either HilC or RtsA (16).

HilE specifically disrupts HilD DNA binding activity. One can envision several ways in which HilE could mechanistically regulate the activity of HilD. (i) HilE may simply interact with HilD in such a way that it blocks DNA binding. (ii) Given the results above and our unpublished data indicating that HilD forms homodimers *in vivo*, it is possible that HilE disrupts HilD dimerization and thereby indirectly blocks HilD DNA binding and activation of the *hilA* promoter. (iii) HilD has been reported to be degraded by Lon protease (24, 25). HilE could bind HilD and deliver it to Lon or another protease. (iv) HilE could regulate HilD function after HilD has bound to DNA. However, HilD functions partially by displacing H-NS (11), making the last model more difficult to imagine.

We first examined the ability of HilE to block HilD DNA binding. To provide a direct *in vivo* assay of HilD DNA binding, we created an artificial promoter that includes an overlapping HilD/HilC binding site and which drives the expression of *lacZ*, analogous

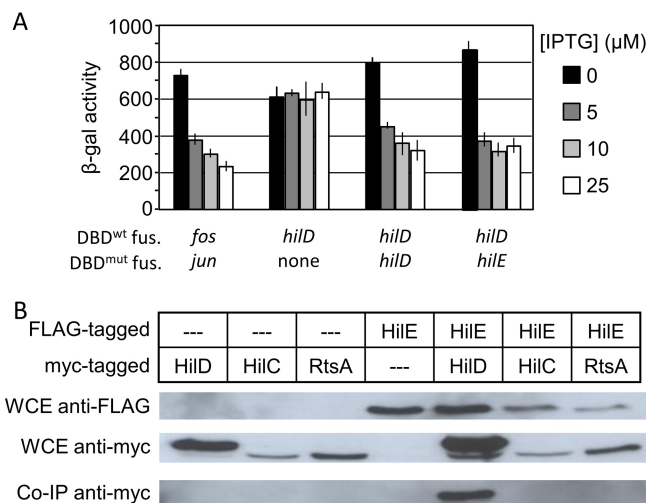


FIG 2 HilE specifically interacts with HilD. (A) A bacterial two-hybrid system with the indicated genes fused to either a wild-type or mutant *lexA* DNA binding domain were grown overnight in LSLB medium at 37°C and then subcultured 1:100 and allowed to incubate at 37°C for 18 h in LSLB medium containing the indicated concentration of IPTG. The genes *fos* and *jun* were employed as a positive control ($n = 6$, 3 sets of duplicate assays on 3 separate days; error bars represent one standard deviation). The strains used were SU202 with either pMS604 and pDP804, pSR659::*hilD*, pSR659::*hilD* and pSR658::*hilE*, or pSR659::*hilD* and pSR658::*hilD*. (B) Strains containing FLAG-tagged HilE and Myc-tagged HilD, HilC, or RtsA were grown overnight and lysed with a French pressure cell. Whole-cell extract (WCE) was then analyzed via Western blotting using either anti-FLAG or anti-Myc antibodies. Extracts were immunoprecipitated using anti-FLAG magnetic beads and subsequently analyzed via Western blotting using anti-Myc. The strains used were 14028 with pLS118, pLS119, or pCE81 and JS2105 with pLS118, pLS119, or pCE81.

to the method used by Luo and Farrand (26). In this construct, HilD or HilC should act as a simple repressor, occluding the binding of RNA polymerase. Thus, HilD and HilC DNA binding could be analyzed independent of displacement of H-NS or activation of RNA polymerase. We placed HilD or HilC under *tetRA* control in strains containing the synthetic promoter construct and in which the chromosomal *hilD*, *hilC*, *rtsA*, and *hilE* loci were deleted. These strains were grown in low-salt LB (LSLB) medium with various concentrations of tetracycline. When induced with tetracycline, there was marked a decrease in β -galactosidase activity in both the HilD- and HilC-producing strains (Fig. 3). These data suggest that both HilD and HilC act as simple repressors when bound to this synthetic promoter.

To determine if HilE prevents HilD from DNA binding, we introduced a plasmid encoding HilE into the above-mentioned strains. As shown in Fig. 3, HilE blocked the reduction in β -galactosidase activity in the HilD-producing strain but had no effect in the HilC-producing background. We interpret these data to indicate that HilE prevents HilD from binding DNA but does not affect HilC. This is consistent with our model (Fig. 1A) and with previous genetic data (14, 16, 20).

We next sought to biochemically verify these results. For these experiments, we used small ubiquitin-related modifier (SUMO)-tagged HilD and HilC. These SUMO-tagged constructs are functional *in vivo* and *in vitro* (see Fig. S1 in the supplemental material) and are more soluble than untagged proteins. We also purified HilE as a SUMO-tagged protein but then proteolytically removed the SUMO tag. We mixed either SUMO-tagged HilD or HilC with a fragment of the *hilA* promoter region and tested whether the addition of HilE would disrupt any shift of the DNA. As shown in Fig. 4, both HilD and HilC shifted a fragment of the *hilA* promoter, consistent with previous findings (11). Addition of HilE blocked DNA binding by HilD but had no effect on the HilC-mediated gel shift. Combined with our findings from Fig. 3, we interpret these data to indicate that HilE prevents HilD from binding DNA but does not affect HilC.

HilE does not affect HilD dimerization. HilE could block HilD DNA binding indirectly by preventing HilD dimerization. To test whether HilD dimerization was

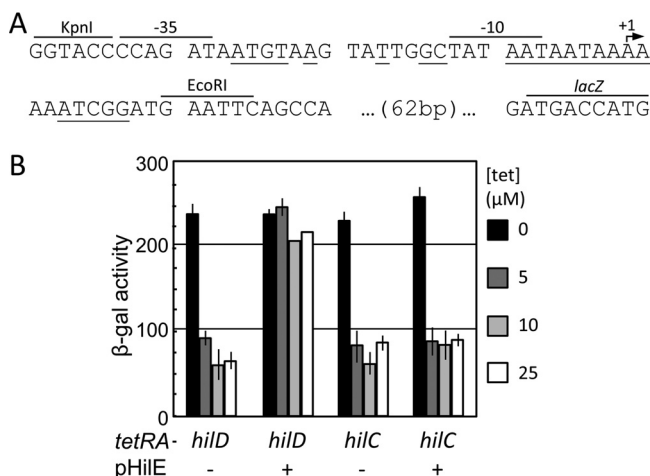


FIG 3 HiE selectively blocks HiD DNA binding. (A) Sequence of the pJG1 HiD/HiC DNA binding reporter plasmid. Bases important for HiD/HiC binding are indicated by underlining (11). The recognition sequences for the restriction enzymes used in the creation of pJG1, the -35 and -10 promoter sequences, the predicted start site of transcription, and the start of the *lacZ* open reading frame are indicated. (B) Strains contained a HiD/HiC-repressible *lac* fusion and *hilD* or *hilC* under the control of a *tetA* promoter. Where indicated, HiE was overexpressed. Strains were incubated overnight at 37°C in LSLB medium with various concentrations of tetracycline ($n = 6$, 3 sets of duplicate assays on 3 separate days; error bars represent one standard deviation). The strains used were JS2106 and JS2107 with or without pHiE.

affected, we employed the bacterial two-hybrid system (22, 23). The data in Fig. 5 suggest that both HiD and HiC homodimerize, as evidenced by an IPTG-dependent decrease in β -galactosidase activity. The addition of HiE to this system did not markedly alter the ability of the HiD- or HiC-LexA_{DBD} proteins to repress *lacZ* expression. Although not conclusive, these data suggested that HiE does not disrupt HiD or HiC homodimerization.

To further test whether HiE affects HiD dimerization, we fluorescently labeled either SUMO-tagged HiD or HiC and mixed the labeled protein in a 5% molar ratio with unlabeled SUMO-tagged HiD or HiC. These samples, with or without the addition of purified HiE, were separated via size exclusion chromatography over a Superose 12 HR 10/30 column, and the fluorescence of each fraction was measured. The use of fluorescently labeled protein allowed us to specifically monitor the elution of HiD or HiC but not HiE, which is unlabeled. The molecular masses of monomers and dimers of SUMO-HiD should be 47.7 kDa and 95.4 kDa, respectively; SUMO-HiC monomers and dimers should be 46.4 kDa and 92.7 kDa, respectively. In Fig. 6, we observe that both HiD and HiC eluted from the column consistent with the predicted mass of SUMO-tagged dimers. Structural prediction suggests that HiE should be a hexamer with a total molecular mass of 100 kDa. The addition of HiE in the HiD sample led to formation of a fluorescent species that eluted with an apparent molecular mass that

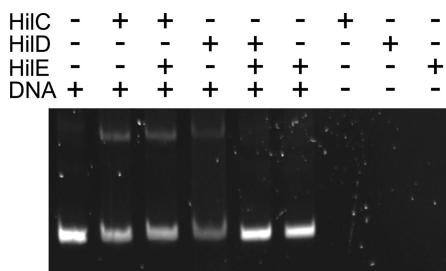


FIG 4 HiE selectively blocks HiD DNA binding. The samples contained 0.7 μM purified HiD or HiC, 0.56 μM DNA, and, where indicated, 2.25 μM HiE. Samples were electrophoresed on a 10% continuous polyacrylamide gel and subsequently stained with SYBR Safe.

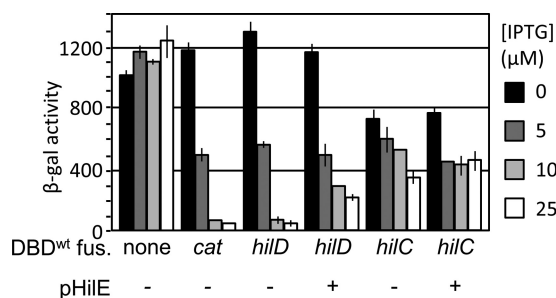


FIG 5 HlIE does not disrupt HlID dimerization. A bacterial monohybrid system with the indicated genes fused to a wild-type *lexA* DNA binding domain were grown under SPI1-inducing conditions with the indicated concentrations of IPTG. Where indicated, *hilE* was overexpressed from pWKS30 ($n = 6$, 3 sets of duplicate assays on 3 separate days; error bars represent one standard deviation). The strains used were SU101 with pSR658, pDD506, pSR658::*hilD*, or pSR658::*hilC* with and without pHiIE.

was significantly greater than 180 kDa. The addition of HlIE to the SUMO-tagged HlC sample had no effect on elution. This is consistent with a predicted protein complex composed of 2 HlID and 12 HlIE monomers (295 kDa). The alternative model is that a hexamer of HlIE would be bound to a HlID monomer, giving a predicted mass of 148 kDa. Combined, these data show that HlIE binds to HlID without disrupting the HlID dimer. Further studies are required to determine the actual stoichiometry of the complex.

HlID remains stable in the presence of HlIE. We next determined whether HlIE regulates HlID activity via modulating its stability. To avoid the complication of HlID autoactivation, we constructed strains in which the expression of a HlID-3×FLAG is under the control of *tetRA*. Furthermore, these strains contained a *hila'*-*lacZ* construct, allowing us to measure HlID activity. We constructed *hilE*⁺, Δ *hilE*, and pHiIE variations of these strains. It has been reported that HlID, HlC, and RtsA are degraded by Lon protease (24, 25). We therefore created otherwise isogenic Δ *lon* Δ *wca* variations of the above-mentioned strains. (The Δ *wca* blocks capsule production, a complicating phenotype of Δ *lon* strains.) All strains were grown overnight and subcultured for 2.5 h with tetracycline at a concentration that leads to activation of *hila* to approximately wild-type levels. After incubation, an aliquot was withdrawn to determine *hila* promoter activity via β -galactosidase assay. The remaining cells were immediately boiled in sodium dodecyl sulfate (SDS) loading buffer. The proteins in each aliquot were then resolved via SDS-PAGE and visualized using anti-FLAG antibody. As shown in Fig. 7, we

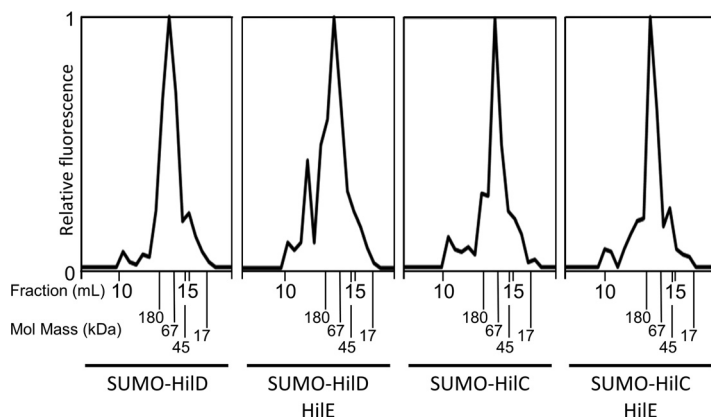


FIG 6 HlIE does not disrupt HlID dimerization. Purified SUMO-HlID or SUMO-HlC was fluorescently labeled and mixed in a 1:20 molar ratio with unlabeled SUMO-HlID or SUMO-HlC. Unlabeled HlIE was mixed in a 12:1 molar ratio with either HlID or HlC where indicated. Proteins were separated by size exclusion chromatography, and the fluorescence of each fraction was assayed. The elution fraction and molecular mass of protein standards are labeled.

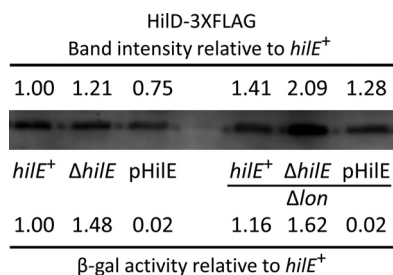


FIG 7 HiE regulates HiD activity independently of stability. The *hiid-3XFLAG* construct is expressed from a *tetA* promoter. Strains with the indicated genotypes were grown to mid-log phase and induced with tetracycline. Western blotting and β -galactosidase assay were performed using aliquots from the same sample. Densitometry of the Western blot was performed using ImageJ. HiD levels and β -galactosidase activity were normalized to those of the *hile*⁺ strain. The strains used were JS2108, JS2109, JS2110, JS2111, JS2112, and JS2113.

observed HiE-dependent *hilA* promoter activity consistent with our genetic model and with previous findings (20). The effect of deleting *hile* is dampened in these backgrounds, likely due to the lack of HiD autoregulation. Importantly, overproduction of HiE completely blocked *hilA* expression. However, the steady-state levels of HiD-3XFLAG were only slightly different in the various backgrounds. Although HiD activity was slightly increased in the Δlon background, overproduction of HiE blocks this activity without having any significant effect on the steady-state levels of the protein. These results suggest that HiE binds and directly blocks HiD function and that degradation of HiD is not a significant mechanism of regulation.

A *hile* mutation partially suppresses the invasion defect conferred by loss of FliZ. HiE is a major negative regulator of HiD and therefore of SPI1 T3SS expression. We therefore wished to determine whether HiE-mediated regulation is important during intestinal invasion. To test this, we performed oral and intraperitoneal (i.p.) competition assays, infecting BALB/c mice with a 1:1 mixture of $\Delta hile$ and wild-type strains and, after 2.5 days, determining the ratio of the two strains in the small intestine and the spleen. As shown in Table 1, the $\Delta hile$ strain conferred no significant phenotype in either oral or i.p. competition assays. This is perhaps not surprising, given that the negative regulatory effects of HiE must be normally overcome during intestinal infection.

It has been previously shown that FliZ significantly affects SPI1 T3SS regulation both *in vivo* and *in vitro*, with a $\Delta fliZ$ strain conferring an ~12.5-fold SPI1-dependent invasion phenotype (20). HiE and FliZ act both independently and simultaneously to regulate HiD (16, 20). We therefore asked whether a $\Delta hile$ mutation would suppress a $\Delta fliZ$ phenotype. We performed both oral and i.p. competition assays using a 1:1 mixture of $\Delta hile \Delta fliZ$ and $\Delta fliZ$ strains. After 3 days, we measured the ratio of the strains in the spleen and small intestine. As shown in Table 1, the $\Delta hile \Delta fliZ$ strain was 5-fold more

TABLE 1 Role of HiE during infection

Genotype of strain ^a :		Infection route	Inoculum (CFU)	No. of mice	Spleen		Small intestine	
A	B				Median CI ^b	P ^c	Median CI	P
$\Delta hile$	WT	i.p.	10 ³	5	1.41	NS	0.79	NS
			10 ⁷	5	0.66	NS	0.93	NS
			10 ⁸	15	2.08	NS	1.41	NS
$\Delta hile \Delta fliZ$	$\Delta fliZ$	i.p.	10 ³	9	0.92	NS	0.85	NS
			10 ⁷	9	3.35	0.01	1.88	0.04
			10 ⁸	5	4.97	0.01	4.21	0.06

^aThe strains used were JS135, JS996, JS2115, and JS2116.

^bThe competitive index (CI) was calculated as described in Materials and Methods.

^cThe Student *t* test was used to compare the CIs to those for the inocula. NS, not significant.

invasive than the Δ *fliZ* strain. In all cases, no phenotype was observed after i.p. infection, indicating that the differences in infectivity between the strains were due to differences in invasiveness alone and not survivability within the animal. These data suggest that the deletion of *hilE* is capable of partially rescuing the invasion deficit conferred by Δ *fliZ*.

DISCUSSION

Salmonella enterica has evolved a complex regulatory network to control the expression of the SPI1 T3SS, which is central to the ability of *Salmonella* to cause gastroenteritis and invade the intestinal epithelium. We have developed a model for the SPI1 regulatory network and have begun to understand the mechanisms by which a large number of signals impinge upon this regulatory system (14, 16). In this work, we focused on HilE, a major negative regulator of HilD with a central role in SPI1 regulation.

HilE was previously identified as a negative regulator of the SPI1 T3SS, and two-hybrid data suggested that HilE binds directly to HilD (5, 21). We previously provided genetic evidence that the HilE regulatory effect is mediated through HilD (16, 20). Here we provide genetic and biochemical data showing that HilE directly binds to HilD and blocks its ability to bind DNA. These studies included the use of a synthetic promoter in which a HilD/HilC binding site is integrated into the -10 and -35 sequences, resulting in a promoter that is repressible by HilD and HilC, allowing us to study HilD and HilC DNA binding *in vivo*, independent of any interactions with H-NS or RNA polymerase. HilE does not block dimerization, nor does it significantly affect the stability of HilD. This regulation by direct protein-protein interaction explains the phenotype conferred by both *hilE* null and HilE overproduction, which completely blocks HilD activity.

Accumulated data suggest that HilD acts as the integration point for many of the environmental signals that control SPI1 expression. HilC and RtsA act as amplifiers of these signals. Although the actual mechanism is known in only a few cases, we have shown that regulation is primarily through either translation of the *hilD* message or control of HilD protein activity (14, 16, 19, 27). HilE falls into the latter class. Saini et al. demonstrated that the interaction of HilD with the *hilD* promoter is the trigger for inducing the expression of SPI1 through a stepwise increase of *hilD* promoter activity and autoactivation (17). Thus, the system is switched on when HilD activity reaches some threshold sufficient to efficiently activate its own expression. It is the amount of HilE that sets this threshold, by blocking the activity of low levels of HilD and thus preventing the inevitable minor fluctuations in HilD activity (noise) from inappropriately activating the system.

The N-terminal domains of HilD, HilC, and RtsA share approximately 10% identity, whereas the C-terminal DNA binding domains share 46% identity. The three proteins bind to the same DNA sequences to activate transcription of the *hilD*, *hilC*, *rtsA*, and *hilA* genes (11, 12, 15). Despite this homology, HilE functions solely through HilD. HilE does not interact with HilC or RtsA, nor does it have any regulatory effect in the absence of HilD. Our unpublished data suggest that HilD, HilC, and RtsA form both homodimers and heterodimers. HilE does not apparently affect dimerization. We propose that HilE binds independently to the individual HilD proteins in both homodimers and heterodimers containing HilD. This would allow HilE to efficiently block induction. Otherwise, heterodimers containing HilD would be capable of inappropriately activating expression of the SPI1 system.

How the level of HilE and, thus, the threshold of activation is controlled is only partially understood. Expression of *hilE* is apparently activated primarily by FimZ, but this regulation is modest under normal conditions. Loss of FimZ results in less than a 2-fold increase in *hilA* expression and a slight delay in the timing of SPI1 turning off in an *in vitro* experiment (17, 28). Only when FimZ is artificially overexpressed is there a dramatic effect on SPI1 expression (17, 28). Mlc is a transcriptional repressor considered a global regulator of carbohydrate metabolism (29–31). Lim et al. showed that loss of Mlc leads to a 2-fold increase in *hilE* mRNA levels and a 4-fold HilE-dependent decrease

in invasion (32). They provide evidence that Mlc acts directly in the *hilE* promoter region, but the effects of various carbon sources on SPI1 expression and the physiological relevance are not clear. The transcriptional regulator LeuO is normally repressed by H-NS under laboratory conditions. Ectopic expression of LeuO in *Salmonella* transcriptionally induces *hilE* and leads to decreased expression of all of the SPI1 regulators. However, importantly, production of LeuO decreases *hilD* transcription in the absence of HilD, decreases *hilC* and *rtsA* transcription in the absence of both HilD and HilE, and leads to an invasion defect in the absence of HilE (33). Thus, although there is an apparent effect on *hilE* transcription, the predominant regulatory effects on SPI1 are HilE independent and seem to be global, analogous to the effects of H-NS (16). The PhoPQ two-component system strongly represses SPI1 gene expression (16). Baxter and Jones recently reported that PhoP activates *hilE* transcription in a manner that is somehow dependent on FimZ (34). However, our data show that PhoPQ can repress the system independently of HilD and HilE (reference 6 and unpublished data); thus, this transcriptional induction of *hilE* is apparently superfluous. Baxter and Jones also report that deletion of *pstS*, which induces the PhoBR two-component system, leads to transcriptional induction of *fimZ* and *hilE* and decreased expression of *hilA* (34). Understanding the overall impact of PhoBR in the SPI1 system will require further investigation.

Wang et al. provided evidence that the small RNA *InvS* (STnc470) negatively affects FimZ levels by an undefined mechanism (35). Deletion of *invS* leads to increased levels of FimZ, which should lead to increased expression of *hilE* (not tested). However, deletion of *invS* did not affect transcription of HilD-dependent *invF*, as would be expected if the phenotype were mediated through HilE. Rather, secretion, but not production, of effector proteins was reduced in an *invS* deletion strain, reportedly via control of translation of *prgH*, encoding a component of secretion machinery. The small RNA *IsrM* is reported to directly block translation of *hilE* by base pairing at the initiation codon (36). Deletion of *isrM* decreased SPI1 gene expression, although transcriptional effects were never quantified. However, the *isrM* deletion mutant is significantly attenuated in an animal, apparently during systemic stages of the disease, in which SPI1 is neither expressed nor required. These data suggest that *IsrM* has pleiotropic effects, and further studies are required to determine the relative role of *hilE* translation in these phenotypes. Together, these data suggest that although there is some regulation of *hilE* transcription and translation, this regulation is not dramatic under normal conditions and/or the identified regulators have effects on the system that are independent of HilE. We believe that a low-level regulation is consistent with the primary role of HilE in setting the threshold for activation rather than being a primary mechanism to repress the system in response to particular environmental signals.

Competition assays between Δ *hilE* and wild-type strains indicated no significant phenotype as a result of the loss of HilE (Table 1). This observation is consistent with previously published data indicating that the deletion of *hilE* confers no invasion phenotype in HEp-2 cells (28). It has been previously demonstrated that the deletion of *fliZ*, encoding a strong positive regulator of HilD, causes a defect in the ability of *Salmonella* to invade the host intestinal epithelium (20). However, deletion of *hilE* in a Δ *fliZ* background led to a 3- to 5-fold increase of the competitive index (CI) over a Δ *fliZ* background. These data together suggest that the negative regulatory effects that HilE has upon HilD are overcome by the point at which *Salmonella* reaches the small intestine and the site of invasion, consistent with the overall role of HilE in setting the threshold of activation that is overcome at the appropriate time and place during infection.

It has been suggested that Lon protease regulates SPI1 expression by the degradation of SPI1 regulatory components (24, 25), and HilD stability has been invoked numerous times to help explain regulatory effects (37–41). We previously demonstrated, however, that loss of Lon causes increased SPI1 expression by stabilizing *FliZ* and not the SPI1 regulatory components *per se* (20). We did note a change in the steady-state levels of HilD in response to HilE. The half-life of HilD-FLAG has been

reported to be from 40 to 155 min by various investigators (20, 37, 38, 40, 41). Under our conditions, the HilD half-life changed from ~75 min in a wild-type background to ~40 min when HilE was overproduced (data not shown). However, the effect of HilE upon HilD activity does not correlate with the relative levels of HilD; overexpression of HilE resulted in a 50-fold decrease in *hilA* expression, while there was only 10% less HilD protein at the time that activity was measured (Fig. 7). This shows that the primary effect of HilE is to directly regulate HilD DNA binding activity and that any effect on HilD stability is a secondary effect. We propose that the changes in HilD half-life conferred by changing HilE levels are a result of HilD being slightly more unstable when free in solution than its DNA-bound form. This likely contributes to the effects seen for other regulatory inputs. Proteins that are truly regulated by modulating stability, such as RpoS, RpoH, and SulA, have half-lives on the order of 1 to 2 min under the unstable condition and are stabilized to >30 min in the induced state (42). In those cases where the stability of HilD has been invoked in regulation, the measured half-life under the "unstable" condition remained >15 min. It is difficult to imagine that these purported 2- to 3-fold changes in half-life in what is a relatively stable protein constitute the primary mechanism of regulation. It seems more likely that these various factors have some more direct effect on HilD activity.

Overall, our data suggest that the role of HilE is to simply bind to HilD, disrupting its ability to bind to DNA, thereby putting HilD into an inactive or low-activity state. The ability of HilE to maintain HilD in such a state is then overcome by the positive regulatory effects, including FlhZ, at the time when *Salmonella* reaches the site of invasion in the small intestine. At this point, HilD is able to autoactivate its own promoter, as well as those of HilC and RtsA, leading to robust and rapid induction of the SPI1 T3SS.

MATERIALS AND METHODS

Construction of strains and plasmids. All *Salmonella* strains are isogenic derivatives of *Salmonella enterica* serovar Typhimurium strain 14028 (American Type Culture Collection) and are listed in Table S1 in the supplemental material. Gene deletions and concomitant insertion of an antibiotic resistance cassette were constructed using λ Red-mediated recombination as previously described (43, 44). All constructs were verified by PCR and moved into a clean background via P22HTint105 phage transduction (45). In some cases, the antibiotic resistance cassette was removed by FLP-mediated recombination via the introduction of pCP20 (46).

Media, reagents, and enzymatic assays. Strains were routinely grown using high-salt lysogeny broth (HSLB) (0.5% yeast extract, 1% tryptone, 1% NaCl) medium (47) at 37°C, except for strains containing the temperature-sensitive plasmid pKD46, p λ Int, or pCP20, which were grown at 30°C. Where indicated, strains grown under *Salmonella* pathogenicity island 1 (SPI1)-inducing conditions were cultured overnight in 3 ml low-salt LB (LSLB) medium (0.5% NaCl) in 13-mm culture tubes at 37°C on a rotary drum and then subcultured 1:100 and grown in 3 ml HSLB medium in standing cultures (low oxygen) in 13-mm tubes at 37°C for 12 to 16 h. For non-SPI1-inducing conditions, cells were grown in LSB medium at 37°C in a rotary drum for 12 to 16 h. Antibiotics were used at the following concentrations: ampicillin, 25 or 50 μ g/ml; chloramphenicol, 20 μ g/ml; kanamycin, 50 μ g/ml; apramycin, 50 μ g/ml; and tetracycline, 13 μ g/ml. Antibodies were purchased from Sigma (monoclonal anti-FLAG M2 and monoclonal anti-c-Myc) or R&D Systems (rabbit anti-mouse IgG conjugated to horseradish peroxidase). Enzymes were purchased from Invitrogen, New England BioLabs, or Stratagene and used according to the manufacturers' recommendations. Primers were obtained from IDT.

β -Galactosidase assays were performed using a 96-well microtiter plate assay as previously described (48). Unless otherwise indicated, cells were grown under SPI1-inducing conditions for these assays. The indicated amount of anhydrotetracycline (aTc) or isopropyl- β -D-thiogalactopyranoside (IPTG) was added for specific gene induction. β -Galactosidase activity units are defined as (micromoles of *ortho*-nitrophenol [ONP] formed per minute \times 10⁶)/(optical density at 600 nm [OD₆₀₀] \times milliliters of cell suspension) and are reported as mean \pm one standard deviation.

Coimmunoprecipitation. We constructed strains containing Myc-tagged HilD, HilC, or RtsA on a plasmid and a FLAG (SPA)-tagged HilE inserted into the chromosome. These strains were grown overnight and subsequently harvested by centrifugation and then lysed in a French pressure cell. The protein concentration was determined by bicinchoninic acid (BCA) assay (Pierce). Anti-Myc magnetic beads (Pierce/Thermo Fisher) were washed twice in Tris-buffered saline (TBS) with Tween (TBST) (25 mM Tris, 150 mM NaCl, 0.05% Tween 20) after being collected in a magnetic stand. Whole-cell extracts (WCE) containing 100 μ g of protein were mixed with a 4-fold excess of TBST and magnetic beads and then allowed to incubate for 30 min while mixing via rotation. The beads were then collected using a magnetic stand and washed twice in 5 \times TBST. Bound protein was then eluted from the magnetic beads using SDS-PAGE sample buffer (50 mM Tris-HCl, pH 6.8, 2% SDS, 10% glycerol, 1% β -mercaptoethanol, 12.5 mM EDTA, 0.02% bromophenol blue) and heating the samples to 95°C for 15 min before analysis via

Western blotting. The samples were separated on a 10% discontinuous SDS-polyacrylamide gel and transferred to Hybond ECL membranes (Amersham). The membranes were blocked three times for 10 min each using 5% (wt/vol) nonfat dry milk in Tris-buffered saline (TBS) (5% milk, 50 mM Tris, 150 mM NaCl, pH 8.0). The blots were then exposed to anti-FLAG M2 or anti-c-Myc antibody diluted 1:1,000 in TBS–5% milk overnight at 4°C. The blots were washed three times for 10 min each time in TBS–5% milk and then exposed to rabbit polyclonal anti-mouse IgG conjugated to horseradish peroxidase (R&D Systems) diluted 1:1,000 in TBS–5% milk for 1 h. The blots were then washed in Tris-buffered saline (TBS) (50 mM Tris, 150 mM NaCl, pH 8.0) three times for 5 min each repetition. The membranes were then exposed to Amersham ECL Western blot detection reagents (GE Healthcare) for 1 min and exposed to film (Kodak).

HiID/HiIC DNA binding reporter. To build a construct in which HiID or HiIC acts as a simple transcriptional repressor, we synthesized four pairs of complementary oligonucleotides. Each pair consisted of a consensus promoter into which a HiID/HiIC binding sequence had been integrated at various positions relative to the –10 and –35 promoter sequences. These fragments were cloned into pDX1 upstream of the promoterless *lacZ* gene between the unique KpnI and EcoRI sites (10). After passage through the *Salmonella* strain JS198 ($r^- m^+$), the plasmids were transformed and stably integrated in single copy at *attλ* by Int-mediated recombination (49), thus creating single-copy transcriptional fusions of the synthetic promoter and *lacZ*. The resulting constructs were verified to have a single integrated copy of the plasmid by PCR. These *lacZ* fusions were then moved via P22 transduction into Δ *hilE* Δ *rtxA* Δ *SP11* strains containing either tetracycline-inducible *hiID* or *hiIC*. The resulting colonies were then screened by β -galactosidase assay to determine the extent to which HiID or HiIC was able to repress expression from the synthetic promoter. The strain with the greatest repression upon tetracycline induction was selected for further studies.

To further increase the responsiveness of the strain to the induction of HiID or HiIC, we mutagenized the –35 region of the synthetic promoter via PCR mutagenesis and reconstructed the fusion strains as described above. The resulting mutants were screened on medium containing tetracycline and X-Gal (5-bromo-4-chloro-3-indolyl- β -D-galactopyranoside). Colonies demonstrating a robust reduction of β -galactosidase activity on X-Gal were selected and analyzed further. The colony with the most robust decrease of β -galactosidase activity upon induction with tetracycline was then selected, and the DNA sequence was determined. Final isogenic strains were then grown overnight in HSLB medium with 0, 0.78, 1.56, or 3.13 mg/ml tetracycline and analyzed via β -galactosidase assay.

Protein purification and gel filtration. Translational fusions of *hiID*, *hiIC*, or *hiIE* to the small ubiquitin-related modifier (SUMO) were constructed using the pET-SUMO plasmid (Invitrogen; the SUMO-HiID construct was a gift from Marc Erhardt). These constructs were then transformed into *Escherichia coli* BL21(DE3) (50). These expression strains were grown overnight and subsequently subcultured 1:100 and incubated for 4 h at 37°C. At this point, 1 mM IPTG was added for induction, and the cells were incubated overnight at room temperature. The cells were then pelleted by centrifugation at $7,000 \times g$ for 10 min and resuspended in NPI buffer (50 mM NaH_2PO_4 , 300 mM NaCl, 10 mM imidazole) with a protease inhibitor cocktail (Roche). The cells were lysed with a French press and the lysate clarified by centrifugation at $22,000 \times g$ for 1 h. The proteins were isolated by passing the clarified cell lysate over Ni-nitrilotriacetic acid (Ni-NTA)-agarose columns (Qiagen), washed with NPI buffer containing 90 mM imidazole, and eluted from the columns using NPI buffer with 250 mM imidazole. The proteins were concentrated by the use of centrifugal concentrators (Amicon). The protein was further purified via size exclusion chromatography over a Superose 12 HR 10/30 column and reconcentrated via centrifugal concentrators (Amicon) before use. The protein concentration was determined by the BCA assay (Pierce).

HiE was cleaved from the SUMO tag using SUMO protease (Invitrogen) and the protease and cleaved SUMO tag removed from the solution by passage over an Ni-NTA column (Qiagen). The HiE was subsequently reconcentrated with a centrifugal concentrator and further purified by size exclusion chromatography. The protein elutes as a series of overlapping peaks, consistent with structural predictions of multimerization to a hexamer (see Fig. S2 in the supplemental material). The Superose 12 HR 10/30 column used for both purification and size exclusion chromatography of the HiID-HiE complex has a resolution 1 to 300 kDa and an exclusion limit of 2,000 kDa.

Electromobility shift assay. HiID, HiIC, and HiE were purified as described above. The *hilA* promoter region is a known target of both HiID and HiIC binding (11). Part of the *hilA* promoter (corresponding to base pairs 3033211 to 3033421, inclusive; NCBI genome accession number CP001363.1) was amplified via PCR and purified using a DNA spin column (Qiagen). Purified HiID or HiIC was mixed with DNA in protein binding buffer (20 mM HEPES, 20 mM KCl, 1% glycerol, 50 μ M EDTA) to establish a solution of 1 μ M HiID or HiIC and 15 nM DNA. HiE was added as indicated to a final concentration of 3 μ M. A continuous 10% polyacrylamide gel was prerun for 30 min before loading the samples. The gel was then stained with SYBR Safe DNA gel stain (Invitrogen) and the image obtained on a transilluminator (Bio-Rad).

Bacterial mono- and two-hybrid methods. Translational fusions of *hiID*, *hiIC*, or *hiIE* to the LexA DNA binding domain were constructed in pSR659 and pSR659 (23). These constructs were then transformed into either the SU101 (homodimer) or SU202 (heterodimer) *E. coli* strain containing a LexA-repressible *sulA* promoter that drives the expression of *lacZ*. These strains were grown overnight in 3 ml HSLB medium at 37°C and then subcultured 1:100 and grown for 18 h at 37°C in 3 ml HSLB medium containing 0, 5, 10, or 25 μ M IPTG. A plasmid producing only the LexA DNA binding domain (pSR658) or a Cat-LexA_{DBD} fusion served as a negative or positive control for the bacterial monohybrid system (SU101), respectively (22). Plasmids pSR658 and pSR659 encoding only the DNA binding domains or encoding *fos* and *jun* were used as negative and positive controls for the two-hybrid (SU202) system, respectively. The strains were then analyzed via β -galactosidase assay as described above.

Stability assay methods. To measure HilD protein stability, we cultured a *tetRA-hilD-3×FLAG attλ::pDX1::hilA'-lacZ⁺* strain in 3 ml LSLB medium in 13-mm culture tubes overnight at 37°C on a rotary drum. The cells were subcultured 1:100 in 10 ml HSLB medium containing 0.8 μg/ml tetracycline in 18-mm culture tubes and grown in a rotary drum for 2.5 h at 37°C. After 2.5 h, the OD₆₀₀ of each sample was measured and the volume adjusted with HSLB medium to equalize the concentration of cells. One milliliter of each sample was then withdrawn for β-galactosidase assays as described above. Additional one-milliliter aliquots were removed from each culture and immediately centrifuged for 1 min at 14,000 × g. The pellets were resuspended in 120 μl sodium dodecyl sulfate (SDS) loading buffer (50 mM Tris-HCl [pH 6.8], 2% SDS, 10% glycerol, 1% β-mercaptoethanol, 12.5 mM EDTA, 0.02% bromophenol blue) and boiled for 10 min before being analyzed via Western blotting as described above using anti-FLAG antibodies. Densitometry of the resulting blots was performed using NIH ImageJ, and protein levels were determined by measuring the area under the curve for the resulting protein bands.

Virulence assays. We inoculated 6- to 8-week-old BALB/c mice (Envigo) via oral gavage or intraperitoneal (i.p.) injection with 0.2 ml of bacterial cell suspension. In all the competition assays, the inoculum consisted of a 1:1 mixture of two bacterial strains. The actual CFU and relative percentage of each strain were determined by direct plating of the inocula. Cells were cultured overnight in HSLB medium in 13-mm tubes at 37°C in a rotary drum. For oral infections, the cells were washed in sterile 0.1 M phosphate-buffered saline (PBS) (pH 8.0) and suspended at 5 × 10⁷ or 5 × 10⁸ cells/ml. For i.p. infections, the cells were washed in sterile saline and diluted to 5 × 10³ cells/ml in 0.15 M NaCl. Food was withdrawn 4 h prior to oral infection. Food and water were provided *ad libitum* after infection. The mice were sacrificed via CO₂ asphyxiation 2.5 days after infection, and the small intestines and spleens were harvested. The organs were homogenized, and aliquots of serial dilutions were plated to appropriate selective medium to determine the number of CFU per organ. The ratio of the two strains was determined by replica plating to appropriate antibiotic-containing medium. The competitive index (CI) was calculated as (CFU of strain A recovered/CFU of strain B recovered)/(CFU of strain A inoculated/CFU of strain B inoculated). Each mutant strain was reconstructed at least once to verify that the observed phenotypes were a result of the designated mutation. The Student *t* test was used to determine whether the output ratio was significantly different from the input ratio. All animal work was reviewed and approved by the University of Illinois Institutional Animal Care and Use Committee and performed under protocols 10050 and 13030.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/JB.00750-17>.

SUPPLEMENTAL FILE 1, PDF file, 0.2 MB.

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