A Salmonella Virulence Factor Activates the NOD1/NOD2 Signaling Pathway

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ABSTRACT The invasion-associated type III secretion system (T3SS-1) of Salmonella enterica serotype Typhimurium (S. Typhimurium) activates the transcription factor NF-κB in tissue culture cells and induces inflammatory responses in animal models through unknown mechanisms. Here we show that bacterial delivery or ectopic expression of SipA, a T3SS-1-translocated protein, led to the activation of the NOD1/NOD2 signaling pathway and consequent RIP2-mediated induction of NF-κB-dependent inflammatory responses. SipA-mediated activation of NOD1/NOD2 signaling was independent of bacterial invasion in vitro but required an intact T3SS-1. In the mouse colitis model, SipA triggered mucosal inflammation in wild-type mice but not in NOD1/NOD2-deficient mice. These findings implicate SipA-driven activation of the NOD1/NOD2 signaling pathway as a mechanism by which the T3SS-1 induces inflammatory responses in vitro and in vivo.

IMPORTANT Salmonella enterica serotype Typhimurium (S. Typhimurium) deploys a type III secretion system (T3SS-1) to induce intestinal inflammation and benefits from the ensuing host response, which enhances growth of the pathogen in the intestinal lumen. However, the mechanisms by which the T3SS-1 triggers inflammatory responses have not been resolved. Here we show that the T3SS-1 effector protein SipA induces NF-κB activation and intestinal inflammation by activating the NOD1/NOD2 signaling pathway. These data suggest that the T3SS-1 escalates innate responses through a SipA-mediated activation of pattern recognition receptors in the host cell cytosol.

The principal pathogenic strategy of Salmonella enterica serotype Typhimurium (S. Typhimurium) is to use its virulence factors to elicit acute intestinal inflammation (1). A by-product of this host response is the generation of a new respiratory electron acceptor, tetrathionate, which enables the pathogen to outgrow other microbes in the intestinal lumen (2), thereby enhancing its transmission (3). S. Typhimurium utilizes a type III secretion system (T3SS-1) encoded by Salmonella pathogenicity island 1 (SPI 1) to elicit intestinal inflammation in animal models (1, 4). The principal function of the T3SS-1 is to deliver proteins, termed effectors, into host cells (5, 6). Five T3SS-1 effectors, termed SipA, SopA, SopB, SopD, and SopE2, act in concert to trigger alterations in the actin cytoskeleton of host cells, thereby promoting epithelial invasion (7–11) and intestinal inflammation (12). S. Typhimurium induces activation of mitogen-activated protein (MAP) kinases, activation of nuclear factor kappa-light-chain-enhancer of activated B cells (NF-κB), and production of proinflammatory cytokines in cultured cell lines in a T3SS-1-dependent fashion (13), suggesting that cytosolic access by the T3SS-1 triggers host cell responses.

Several mechanisms through which cytosolic access by the T3SS-1 can induce inflammatory responses have been proposed. These mechanisms have in common that cytosolic access by the T3SS-1 is detected by host macrophages via cytosolic recognition of translocated bacterial proteins, resulting in caspase 1 activation and consequent proteolytic activation of interleukin-1β (IL-1β) and IL-18, two proinflammatory cytokines (14–16). However, it is also clear that these mechanisms are not sufficient to explain important in vitro and in vivo observations, suggesting that additional pathways by which the T3SS-1 induces inflammatory responses remain to be discovered. Specifically, activation of caspase 1 cannot account for the T3SS-1-dependent activation of NF-κB observed in vitro (13). Furthermore, the invasion-associated T3SS-1 of S. Typhimurium elicits intestinal inflammation in vivo through a myeloid differentiation primary response protein 88 (MyD88)-independent mechanism (17). Caspase 1 activation does not explain the generation of these MyD88-independent proinflammatory responses, because MyD88 is required for signaling through receptors for IL-1β and IL-18 (18–20). Here we identify a new MyD88-independent mechanism by which the T3SS-1 induces host cell responses in vitro and demonstrate that this mechanism contributes to S. Typhimurium-induced intestinal inflammation in vivo.

RESULTS NF-κB activation by the invasion-associated T3SS is invasion independent. The invasion-associated T3SS-1 of S. Typhimurium activates NF-κB in human epithelial cells (see Fig. S1A in the supplemental material) (13, 21). To analyze whether this process is invasion dependent, we used human epithelial cells transfected...
with an NF-κB luciferase reporter construct. A T3SS-1-deficient S. Typhimurium mutant (invA mutant) neither activated NF-κB (Fig. 1A) nor invaded epithelial HeLa cells (Fig. S1B). Introduction of the Yersinia pseudotuberculosis invasin protein increased invasiveness of the T3SS-1-deficient S. Typhimurium mutant (P < 0.05) but did not restore NF-κB activation. Similar results were obtained with human embryonic kidney 293 (HEK293) cells, although inactivation of the invasion-associated T3SS resulted only in a partial inhibition of NF-κB activation in this cell line (Fig. S1C and S1D).

The T3SS-1 effectors SipA, SopA, SopB, SopD, and SopE2 act in concert to promote invasion of epithelial cells (10). An S. Typhimurium mutant lacking these five effector proteins (sipA sopABDE2 mutant) was deficient for invasion (P < 0.05) and NF-κB activation (P < 0.05) (see Fig. S1 in the supplemental material). Introduction of the Y. pseudotuberculosis invasin protein into the sipA sopABDE2 mutant restored invasiveness (P < 0.05) but not NF-κB activation. These data suggested that NF-κB activation required delivery of T3SS-1 effector proteins but not bacterial invasion.

**SipA is a key mediator of NF-κB activation in vitro.** To determine whether host cells can sense delivery of specific effectors, we complemented the S. Typhimurium sipA sopABDE2 mutant with plasmids encoding SipA, SopA, SopB, SopD, or SopE2 (10) and monitored NF-κB activation (Fig. 1). Only a plasmid encoding SipA complemented the sipA sopABDE2 mutant for NF-κB activation in epithelial HeLa cells (P < 0.05) (Fig. 1A). Endogenous TLR5 expression renders HEK293 cells responsive to stimulation with flagellin (23, 24), which might explain why we observed NF-κB activation in response to infection with the sipA sopABDE2 mutant or to stimulation with purified S. Typhimurium flagellin (FliC) (P < 0.05) in this cell line (Fig. 1A and 1B). Consistent with

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**FIG 1** SipA-mediated activation of NF-κB in HeLa and HEK293 cells is MyD88 independent. (A) HeLa cells (gray bars) and HEK293 cells (black bars) transfected with an NF-κB luciferase reporter construct were infected with the S. Typhimurium sipA sopABDE2 mutant complemented with genes encoding the indicated effector proteins. (B) HEK293 cells were transfected with MyD88DN and infected with the S. Typhimurium sipA sopABDE2 mutant and complemented derivatives. Data represent the NF-κB luciferase activity as fold increase over the uninfected controls. Values are the means from at least 3 independent experiments ± standard errors. Brackets indicate the statistical significance of differences. NS, not significant.
this idea, transfection of HEK293 cells with a plasmid encoding a dominant-negative form of the TLR5 adaptor protein MyD88 (MyD88DN) abrogated NF-κB activation induced by the sipA sopABDE2 mutant or purified flagellin (Fig. 1B). Complementation of the sipA sopABDE2 mutant with plasmids encoding SipA or SopE2 induced NF-κB activation in HEK293 cells transfected with MyD88DN (P < 0.05) (Fig. 1B). Collectively, these data suggested that bacterial delivery of SipA activates NF-κB through a MyD88-independent mechanism.

To determine whether SipA is sufficient for NF-κB activation, a green fluorescent protein (GFP)-SipA fusion protein was expressed ectopically in HEK293 cells transfected with an NF-κB luciferase reporter construct (Fig. 2). Ectopic expression of GFP-SipA, but not ectopic expression of GFP, resulted in NF-κB activation (P < 0.05). These data suggested that cytosolic localization of SipA in the absence of other bacterial molecules was sufficient for inducing NF-κB activation in host cells.

The SipA protein contains an N-terminal domain required for T3SS-1 translocation (NTD), a C-terminal actin-binding domain (ABD), and two central domains mediating SipA focus formation (F1) and SipA-SipA interaction (F2) (25). GFP-SipA fusion proteins lacking one or several of these domains were constructed to investigate which domain(s) is required for NF-κB activation (Fig. 2A). Ectopic expression of these fusion proteins in HEK293 cells suggested that the F2 domain, which is not required for host cell invasion (25), was necessary but not sufficient for NF-κB activation (P < 0.05) (Fig. 2B; see also Fig. S2 in the supplemental material). In contrast, the ABD, which is required to support S. Typhimurium host cell invasion (25), was not necessary for NF-κB activation.

To determine whether the F2 domain was necessary for NF-κB activation upon bacterial delivery of SipA, we complemented the sipA sopABDE2 mutant with plasmids carrying truncated sipA genes encoding the first 425 amino acids (aa) (sipA1-425), the
first 290 amino acids (sipA’1-290), or the first 149 amino acids (sipA’1-149) of SipA. Complementation of the sipA sopABDE2 mutant with genes encoding full-length SipA or a truncated SipA lacking the ABD (sipA’1-425) resulted in NF-κB activation through a MyD88-independent mechanism (Fig. 3). In contrast, NF-κB activation was not observed with proteins lacking the F2 domain, which were encoded by sipA’1-290 and sipA’1-149. In summary, the F2 domain was necessary for NF-κB activation regardless of whether SipA was delivered by bacteria or expressed ectopically.

SipA-mediated NF-κB activation requires RIP2 in vitro. One MyD88-independent mechanism contributing to intestinal inflammation during S. Typhimurium infection is the activation of nucleotide-binding and oligomerization domain 1 (NOD1) and NOD2, two intracellular sensors of bacterial cell wall fragments (26, 27). NOD1 and NOD2 interact with protein kinase receptor-interacting protein 2 (RIP2) to mediate NF-κB activation (28, 29). Stimulation of HEK293 cells with the NOD1 ligand C12-iE-DAP resulted in NF-κB activation (P < 0.05), which was MyD88 independent and could be inhibited with SB203580 (Fig. 3B and 3D), a pyridinyl imidazole inhibitor of RIP2 activity (30). NF-κB activation induced by bacterial delivery of SipA was abrogated when cells were treated with the RIP2 inhibitor SB203580 (Fig. 3A and 3C). These data suggested that SipA-mediated NF-κB activation required RIP2.

SipA-mediated NF-κB activation requires both NOD1 and NOD2 in vitro. We next investigated whether NOD1 or NOD2 was required for SipA-dependent NF-κB activation (Fig. 4).
Transfection of HEK293 cells with plasmids encoding a dominant-negative NOD1 protein (Nod1DN) or a dominant-negative RIP2 protein (Rip2DN) inhibited NF-κB activation elicited by stimulation with the NOD1 ligand C12-iE-DAP (Fig. 4A). Similarly, transfection of HEK293 cells with dominant-negative NOD2 (Nod2DN) or Rip2DN inhibited NF-κB activation elicited...
by stimulation with the NOD2 ligand muramyl dipeptide (MDP) (Fig. 4B). Dominant-negative forms of NOD1 (Nod1DN), NOD2 (Nod2DN), or RIP2 (Rip2DN) did not inhibit NF-κB activation in HEK293 cells stimulated with flagellin (FliC) (Fig. 4C). Importantly, transfection of HEK293 cells with dominant-negative forms of NOD1 (Nod1DN), NOD2 (Nod2DN), or RIP2 (Rip2DN) inhibited NF-κB activation elicited by ectopic expression of GFP-SipA or GFP-SipA425 ($P < 0.05$) (Fig. 4D and 4E). Cotransfection of the dominant-negative forms of NOD1, NOD2, or RIP2 with GFP-SipA425 did not markedly affect protein expression of SipA425 (Fig. 4F). Ectopic expression excluded the possibility that SipA-mediated NF-κB activation was due to contamination with endogenous bacterial ligands, such as MDP. Furthermore, both NOD1 and NOD2 contributed to NF-κB activation in response to ectopic expression of SipA (Fig. 4D), while responses to C12-iE-DAP required only NOD1 (Fig. 4A) and responses to MDP required only NOD2 (Fig. 4B). Collectively, these data suggested that SipA activated NOD1/NOD2 through a pathway that was distinct from that observed with bacterial cell wall fragments.

To investigate interaction of SipA with NOD1, HEK293 cells were transfected with Flag-tagged NOD1 protein (NOD1-Flag) and either Myc-tagged SipA protein (Myc-SipA) or GFP-SipA425. After immunoprecipitation of Flag-NOD1 with anti-Flag antibody, SipA was detected in the immunoprecipitate (Fig. 5). Furthermore, colocalization of SipA and NOD1 was observed microscopically in some HEK293 cells transfected with NOD1-Flag and GFP-SipA (see Fig. S3 in the supplemental material). Collectively, these data suggested that SipA either interacts with NOD1 directly or is present in a protein complex that contains NOD1.

SipA does not trigger mucosal inflammation in NOD1/NOD2-deficient mice. To test the biological significance of our observations, we determined the role of NOD1/NOD2 in SipA-mediated inflammation in vivo using the mouse colitis model of S. Typhimurium infection (4). The T3SS effectors SipA, SopA, SopB, SopD, and SopE2 act in concert to trigger intestinal inflam-

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**FIG 5** Coimmunoprecipitation of NOD1 and SipA. At 48 hours after transfection of HEK293 cells with the indicated constructs, immunoprecipitation was performed using anti-Flag antibodies. Proteins were detected in whole-cell lysates and immunoprecipitates by Western blot analysis with anti-Flag antibody (α-Flag), anti-Myc antibody (α-myc), or anti-GFP antibody (α-GFP).
To restrict our analysis to SipA-mediated mechanisms of inflammation, we compared an *S*. Typhimurium strain producing only SipA (*sopABDE2* mutant) with an isogenic SipA-deficient mutant (*sipA sopABDE2* mutant). Remarkably, the SipA-proficient *S*. Typhimurium strain (*sopABDE2* mutant) triggered mucosal inflammation in wild-type mice (C57BL/6) but not in the NOD1/NOD2-deficient mice 2 days after infection (Fig. 6; see also Fig. S4 in the supplemental material). NOD1/NOD2-deficient mice failed to induce SipA-dependent expression of *Ifng*, the gene encoding gamma interferon (IFN-γ) (Fig. 6A), and *Tnfa*, the gene encoding tumor necrosis factor alpha (TNF-α) (Fig. 6B), in the cecal mucosa. Wild-type (C57BL/6) mice infected with the SipA-proficient *S*. Typhimurium strain (*sopABDE2* mutant) exhibited marked intestinal inflammation, which was absent in wild-type mice infected with the SipA-deficient mutant (*sipA sopABDE2* mutant) or in NOD1/NOD2-deficient mice (Fig. 6C; see also Fig. S4). In summary, a SipA-proficient *S*. Typhimurium...
strain required NOD1/NOD2 for eliciting intestinal inflammation.

**DISCUSSION**

The translocation of multiple proteins by the T3SS-1 can mask the contribution of an individual pathway to inflammation in vivo. For example, previous studies show that S. Typhimurium can induce inflammation in mice deficient for MyD88, RIP2, or caspase 1 (31). However, when mice are infected with an S. Typhimurium strain translocating only the SopE protein (sipA sopB sopE2 sed mutant), it becomes apparent that caspase 1 activation contributes to mucosal inflammation in the mouse colitis model (32). Similarly, here we show that a contribution of NOD1/NOD2 signaling to mucosal inflammation in the mouse colitis model (32). Typhimurium, thereby escalating mucosal inflammatory responses. Consistent with this idea, SipA enhanced mucosal inflammation in the mouse colitis model in a NOD1/NOD2-dependent fashion.

SipA is a protein delivered into the host cell cytosol by the invasion-associated T3SS of S. Typhimurium (6). SipA contributes to invasion of epithelial cell lines by modulating actin assembly through its C-terminal ABD (6, 9–11, 25). The N-terminal 425 amino acids of SipA trigger neutrophil transmigrational migration in vitro (33–36). Finally, SipA contributes to intestinal inflammation in vivo (12, 37), but the underlying mechanism has remained unclear. Here we show that the proinflammatory effects of SipA required NOD1/NOD2 activity. Furthermore, in vitro experiments demonstrated that SipA was necessary and sufficient for NOD1/NOD2-dependent NF-κB activation, a process that did not require its ABD. These data suggested that cytosolic sensing of SipA by the pattern recognition receptors NOD1 and NOD2 served as a signal for cytosolic access by the invasion-associated T3SS-1 of S. Typhimurium, thereby escalating mucosal inflammatory responses. Consistent with this idea, SipA enhanced mucosal inflammation in the mouse colitis model in a NOD1/NOD2-dependent fashion.

**MATERIALS AND METHODS**

**Bacterial strains, tissue culture cells, and culture conditions.** S. Typhimurium strain IR715 (38), a fully virulent, nalidixic acid-resistant derivative of American Type Culture Collection (ATCC) isolate 14028, was used as a wild-type isolate. IR715 derivatives carrying mutations in invA (SPN449) (39); in sopA, sopB, sopD, and sopE2 (ZA20) (12); and in sipA, sopA, sopB, sopD, and sopE2 (ZA21) (12) have been described previously. Plasmids carrying the sopD gene (pMR15), the sopE2 gene (pMR17), the sopB gene (pMR26), the sopA gene (pMR28), or the sipA gene (pMR29) from S. Typhimurium have been described previously (10). The sipA25, sipA290, and sipA149 fragments of the sipA gene were amplified without the sipA promoter from pMR29 and cloned directionally behind the lac promoter of cloning vector pWSK29 (40) with BamHI and XbaI. Plasmid pRI203, encoding the Yersinia pseudotuberculosis invasion protein, has been described previously (22). Bacteria were grown aerobically at 37°C in Luria-Bertani (LB) broth supplemented with antibiotics. The HeLa 57A cell line stably transfected with an NF-κB luciferase reporter construct (41) was generously provided by R. T. Hay (the Wellcome Trust Centre for Gene Regulation and Expression, College of Life Sciences, University of Dundee, United Kingdom). The HEK293 cell line was purchased from ATCC. HeLa 57A cells and HEK293 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal calf serum (FCS) at 37°C in a 5% CO₂ atmosphere.

**Construction of expression plasmids.** The sipA gene was PCR amplified from the chromosome of S. Typhimurium IR715, and the obtained product was cloned into the mammalian expression vector pEGFP-C1 (BD Biosciences Clontech). The mutant forms of sipA were PCR amplified from pEGFP-SipA and cloned in pEGFP-C1. Expression vectors with deletions in the F1 domain (pEGFP-ΔF1AABD) and in the F2 domain (pEGFP-ΔF2) were engineered by overlap extension PCR. The two DNA fragments needed for the construction of these sipA mutant forms were amplified by PCR, gel purified, and fused in one PCR. The full-length sipA and the mutant forms were expressed as a fusion to the C terminus of the enhanced GFP (EGFP). Construction of Myc-SipA was obtained by cloning the full-length sipA gene generated by PCR from pEGFP-SipA into pcMV-myc (Clontech). The full-length human NOD1 and dominant-negative versions of human NOD1 (hNOD1) and hNOD2 were PCR amplified from pUNO-hNOD1 and pUNO-hNOD2 (InvivoGen), respectively. hNOD1DN (amino acids [aa] 127 to 954) lacks the N-terminal caspase recruitment domain (CARD), hNOD2DN (aa 217 to 1041) lacks the two N-terminal CARDs. The obtained PCR products were cloned into a derivative of the mammalian expression vector pT’racer-CMV2 (Invitrogen, Life Technologies). The derivative lacks the GFP gene and contains a 3×Flag tag at the C-terminal cloning site (42). The human RIP2 gene was PCR amplified from cDNA prepared from HEK293 cells. The obtained PCR product was cloned into pT’racer-CMV2, yielding pT’racer-hRIP2 (aa 1 to 540). The dominant-negative form of hRIP2 (aa 1 to 434), lacking the C-terminal CARD, was PCR amplified from pT’racer-hRIP2 and cloned into pcMV-HA (Clontech). The hemagglutinin (HA) tag is fused to the N terminus of hRIP2DN. The plasmid pDeNy-Md88 expressing a dominant-negative human MyD88 gene was purchased from InvivoGen. All constructs were verified by DNA sequencing.

**Gentamicin protection assay.** HEK293 and HeLa 57A cells were seeded in a 24-well tissue culture plate at approximately 50% confluence. At the time of infection, the cells had a confluence of approximately 100%. Overnight cultures of S. Typhimurium strains were diluted 1 in 50 and grown for 3 h at 37°C. HEK293 and HeLa 57A cells were infected with S. Typhimurium strains at approximately 10⁵ CFU/ml. The bacteria were incubated for 1 h at 37°C to allow invasion. The cells were washed three times with Dulbecco’s phosphate-buffered saline (DPBS; pH 7.4) to remove extracellular bacteria. DMEM containing 10% FCS and gentamicin (0.1 mg/ml) was added, and the cells were incubated for 90 min at 37°C. The cells were washed in DPBS and lysed with 0.5 ml of 1% Triton X-100, and the lysates were transferred to a sterile Eppendorf tube. The wells were washed with 0.5 ml PBS and transferred to the Eppendorf tubes. Serial dilutions were plated on LB plates with the appropriate antibiotics to count the intracellular bacteria.

**Transient transfection.** HEK293 cells were grown in 24- or 48-well tissue culture plates in DMEM containing 10% FCS until 40% confluence was reached (~24 h). HEK293 cells were transiently transfected with a total of 250 ng of plasmid DNA, consisting of 25 ng of the reporter plasmid pNF-κB-luc, 25 ng of normalization vector pTK-LacZ, and 200 ng of either empty control vector, pUNO-hMyD88DN (InvivoGen), or pEGFP-SipA constructs. For the cotransfection experiments with pEGFP-SipA constructs and hNOD1DN-3×Flag, hNOD1DN-3×Flag, or HA-hRIP2DN, 100 ng of DNA for each plasmid was used. FuGene HD (Roche) was used as transfection reagent at a lipid-to-DNA ratio of 1 to 5. After 48 h of incubation, the cells were infected with the appropriate bacterial strains or the NOD1 ligand C12-IE-DAP, the NOD2 ligand MDP, or the TLR5 ligand flagellin (InvivoGen). The HEK293 cells were stably transduced with an NF-κB-luciferase reporter system. HeLa 57A cells stably transduced with an NF-κB-luciferase reporter construct and transiently transfected HEK293 cells were infected with the indicated S. Typhimurium strains (10⁵ CFU/ml) for 3 h, after which the cells were washed with DPBS and incubated at 37°C for an additional 2 h in the presence of DMEM containing 10% FCS. The cells were washed in DPBS and lysed in 0.1 ml of reporter lysis buffer (Promega). Firefly luciferase activity was measured with the luciferase assay system (Promega) using a plate reader. For normalization of the efficiency of transfection, luciferase values were adjusted to β-galactosidase values as determined with the β-galactosidase assay (Promo-
Protein expression analysis. HEK293 cells transfected with fusion proteins were lysed in lysis buffer (50 mM Tris–HCl, pH 7.6, 150 mM NaCl, 0.2% [vol/vol] NP-40, and 1 mM EDTA). An 0.01-mg amount of total protein was separated by SDS-polyacrylamide gel electrophoresis. Proteins were transferred to a polyvinylidene fluoride membrane (Millipore) in a semidry transfer process (Bio-Rad). The membranes were incubated in PBS containing 3% nonfat dry milk and 0.05% Tween 20 to block nonspecific binding. The membranes were incubated with polyclonal primary antibodies raised against tubulin (Cell Signaling Technology), mouse anti-Flag (Sigma), or mouse anti-HA (Covance). For GFP detection, a horseradish peroxidase (HRP)-conjugated anti-mouse IgG antibody (Sigma) and an HRP-conjugated anti-rabbit IgG (Promega) were used as the secondary antibodies for tubulin, Flag, and HA detection. Horseradish peroxidase activity was visualized by adding Immobilon Western chemiluminescent substrate (Millipore) to the membrane. Images were recorded and processed by a BioSpectrum imaging system (UVProbe).

Immunoblotting. The whole-cell lysate was incubated for 2 h at 4°C with Dyna protein G beads (Invitrogen) coated with anti-Flag M2 antibody (Sigma). After 10 washes with the lysis buffer and 2 washes with 50 mM ammonium bicarbonate, samples of the immunoprecipitate were collected and subjected to SDS-PAGE and immunoblotting. The proteins of interest were detected with anti-Flag M2 antibody (Sigma), anti-Myc antibody (Cell Signaling), and anti-GFP antibody (Sigma).

Fluorescence microscopy. HEK293 cells were transfected with GFP-SipA and NOD1-Flag. After 48 h, the cells were washed and fixed with 3% paraformaldehyde. The cells were permeabilized with 0.1% saponin–10% goat serum (blocking reagent) in PBS for 30 minutes at room temperature. After blocking, cells were incubated with anti-Flag M2 antibody (1:1,000 dilution; Sigma) in 0.1% saponin and 10% goat serum in PBS for 1 h at room temperature. The cells were rinsed twice in 0.1% saponin in PBS and then twice in PBS and incubated (1 h) with Alexa Fluor 647 goat anti-mouse IgG (1:1,000 solution; Invitrogen). Flow cytometry was used. After the cells were washed twice in 0.1% saponin in PBS, twice in PBS, and once in H2O, they were embedded in FluorSave (Calbiochem) and analyzed with an LSM700 confocal microscope (Zeiss).

Animal experiments. Streptomycin (20 mg/mouse) (Sigma)-pretreated C57BL/6 mice (Jackson Laboratory) and NOD1/NOD2-deficient mice (generously provided by D. Portnoy) were mock infected with 0.1 ml of sterile LB broth or infected orally with 1 × 10⁶ CFU (in 0.1 ml of LB broth) of an S. Typhimurium sopABDE2 mutant (ZA21) or a sipA sopABDE2 mutant (ZA21) carrying the plasmid pHP45 (Haire et al.). After infection, mice were sacrificed, and samples of the cecum were snap-frozen in liquid nitrogen for isolation of mRNA.

Histopathology. Formalin-fixed, hematoxylin- and eosin-stained ce cal tissue sections were blind for evaluation by a veterinary pathologist. The following pathological changes were scored: (i) neutrophil infiltration, (ii) infiltration by mononuclear cells, (iii) submucosal edema, (iv) epithelial damage, and (v) inflammatory exudate. The pathological changes were scored on a scale from 0 to 4 as follows: 0, no changes; 1, detectable; 2, mild; 3, moderate; 4, severe. Images were taken using an Olympus BX41 microscope.

Real-time PCR. Samples of the cecum were collected, immediately snap-frozen in liquid nitrogen, and stored at −80°C. RNA was extracted using TrizReagent (Molecular Research) according to the instructions of the manufacturer. Reverse transcription was performed on 1 μg of DNase-treated RNA with TaqMan reverse transcription reagent (Applied Biosystems). For each real-time reaction, 4 μl of cDNA was used. Real-time transcription–PCR (RT–PCR) was performed using Sybr green (Applied Biosystems) and an ABI 7900 RT–PCR machine (Applied Biosystems). The fold change in mRNA levels was determined using the comparative threshold cycle (ΔCt) method (Applied Biosystems). Target gene transcription was normalized to the levels of glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNA.

Statistical analysis. A paired Student t test was used to confirm statistical significance in the tissue culture experiments. To determine statistical significance of the relative mRNA transcription between treatment groups in the animal experiments, an unpaired Student t test was used. To determine the statistically significant differences in the total histopathology scores, a Mann-Whitney U test was used. A two-tailed P value of <0.05 was taken to be significant.

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SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at http://mbio.asm.orglookup/suppl/doi:10.1128/mBio.00266-11/-/DCSupplemental.

Figure S1, PDF file, 0.1 MB.
Figure S2, PDF file, 0.2 MB.
Figure S3, PDF file, 0.7 MB.
Figure S4, PDF file, 2.8 MB.

REFERENCES


