**Shigella IpaH0722 E3 Ubiquitin Ligase Effector Targets TRAF2 to Inhibit PKC–NF-κB Activity in Invaded Epithelial Cells**

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**Abstract**

NF-κB plays a central role in modulating innate immune responses to bacterial infections. Therefore, many bacterial pathogens deploy multiple mechanisms to counteract NF-κB activation. The invasion of and subsequent replication of *Shigella* within epithelial cells is recognized by various pathogen recognition receptors as pathogen-associated molecular patterns. These receptors trigger innate defense mechanisms via the activation of the NF-κB signaling pathway. Here, we show the inhibition of the NF-κB activation by the delivery of the IpaH E3 ubiquitin ligase family member IpaH0722 using *Shigella*’s type III secretion system. IpaH0722 dampens the acute inflammatory response by preferentially inhibiting the PKC-mediated activation of NF-κB by ubiquitinating TRAF2, a molecule downstream of PKC, and by promoting its proteasome-dependent degradation.

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**Introduction**

The intestinal epithelium deploys multiple sensors and defense systems against microbial intrusion. Microbial components and infection-associated cellular damage are recognized as pathogen-associated molecular patterns (PAMPs) and as danger-associated molecular patterns (DAMPs), respectively. Pattern recognition receptors (PRRs) recognize PAMPs and DAMPs, thus activating the immune system to clear the bacteria and initiate the repair of the injured epithelial lining [1,2]. Nevertheless, many bacterial pathogens such as *Shigella*, *Salmonella*, *Yersinia*, enteropathogenic *Escherichia coli* (EPEC), and enterohemorrhagic *E. coli* (EHEC), are able to efficiently colonize the intestinal epithelium by utilizing highly evolved mechanisms to counteract host innate defense mechanisms [3]. Previous studies have reported that enteric bacterial pathogens possess distinctive mechanisms to attenuate host inflammatory responses, which is a prerequisite for promoting intracellular and extracellular bacterial survival [4,5]. For example, during the invasion of the intestinal epithelium by *Shigella*, the Toll-like receptors (TLRs) and Nod-like receptors (NLRs) are the PRRs that recognize PAMPs and DAMPs [6–9]. These receptors stimulate host inflammatory signaling pathways, including nuclear factor κB (NF-κB) and mitogen activated protein kinases (MAPK), which culminate in the expression of proinflammatory chemokines and cytokines [6–9]. *Shigella* counteract innate immune responses by delivering effector molecules using its type III secretion system (T3SS) [8–10]. The targeting molecules and mechanisms of inhibition of the NF-κB pathway are specific to each of the effectors, and *Shigella* are known to deliver OspG [11], OspI [12], OspZ [15], and IpaH9.8 [14–17], to efficiently attenuate NF-κB activation thus allowing replication within the intestinal epithelium [8,9].

IpaH9.8 belongs to IpaH E3 ubiquitin ligase family [16] and it is also called novel E3 ligase (NEL) [18]. The *ipaH* genes, which are originally identified in *Shigella* large virulence plasmid, are conserved by Gram-negative bacterial pathogens, including *Shigella*, *Salmonella*, *Yersinia*, *Edwardsiella ictluri*, *Bradyrhizobium japonicum*, *Rhizobium sp. strain NGR234*, *Pseudomonas putida*, *P. entomophila*, *P. fluorescens*, and *P. syringae* [16,19,20]. The IpaH family proteins share common structural and functional properties; they contain an N-terminus leucine rich repeat (LRR) and a highly conserved C-terminal region (CTR) [21,22]. Within the conserved CTR there is a Cys residue that is critical for its E3 ubiquitin ligase activity [16]. Each of the IpaH family effectors, including IpaH9.8 (*Shigella*), IpaH7 (*Salmonella*), IpaH10 (*Salmonella*), IpaH2 (*Salmonella*), YopM (*Yersinia*), Y4H (R. jostii), and B1pM (*Torrina*), has been shown to target specific host proteins in a variety of cell types; some of them act as effectors that attenuate host inflammatory responses, while others modulate host defense responses in plants [17,23–25]. The existence of multiple effectors with E3 ligase activity suggests that a divergent array of E3 ligases is required for promoting bacterial infection and antagonizing host innate defense responses.

The *Shigella flexneri* strain, YSH6000, has three *ipaH* genes (*ipaH9.8, ipaH7.8*, and *ipaH4.5*) on its large virulence plasmid and seven *ipaH* genes on its chromosome (*ipaH0722, ipaH0807,*
**Author Summary**

In response to bacterial infection, host cells induce a plethora of innate immune responses to combat the infection. However, many bacterial pathogens have developed sophisticated mechanisms to evade the host’s immune system. Because NF-κB is crucial for innate immune responses against bacterial infection, bacterial pathogens deploy multiple countermeasures to inhibit NF-κB activation. The invasion and replication of *Shigella* within host cells results in cellular damage and the production of bacterial components that trigger NF-κB activation. Here, we show that the *Shigella* type III secretion system (T3SS) effector IpaH0722, a member of the IpaH E3 ubiquitin ligase family, inhibits NF-κB activation during *Shigella* infection. IpaH0722 preferentially targets the PKC–NF-κB pathway, which is activated in response to danger signals caused by disruption of the phagosomal membrane during the dissemination of *Shigella* into the cytoplasm. IpaH0722 inhibits NF-κB activation by targeting TRAF2, which lies downstream of PKC, for ubiquitination and proteasome-dependent degradation.

ipaH1383, ipaH1880, ipaH2022, ipaH2292, and ipaH2610 [26–28]. We previously showed that all of these IpaH effector proteins are secreted via the T3SS [28,29], suggesting they have the potential to act as E3 ubiquitin ligase effectors during infection, although their exact roles and host targets during infection, with the exception of IpaH9.8, remain largely unclear [17]. In the mouse lung infection model, we showed that mice infected with the *ApaH*-chromosome mutant, which lacks all seven chromosomal *ipaH* genes, had a more severe inflammatory response and less colonizing bacteria compared to the WT strain [28], suggesting that modulation of the host inflammatory response by chromosomal IpaH proteins promotes bacterial infection. Therefore, to gain further insight into the strategies employed by *Shigella* to counteract host innate defence systems during infection, we sought to characterize the chromosomally encoded IpaH effectors. We found that IpaH0722 targets TRAF2 and plays an important role in dampening the PKC–NF-κB pathway, in response to membrane damage generated during *Shigella* invasion of epithelial cells.

**Results**

IpaH0722 dampens NF-κB activation during *Shigella* infection

The infection of mice using the *ApaH*-chromosome mutant, which lacks all of the chromosomal *ipaH* genes, leads to an increased production of the NF-κB-responsive gene MIP-2 [28]. In this current study, we used HeLa cells to determine the effect of the *ApaH*-chromosome mutant on the NF-κB activation by measuring the kinetics of IkBα degradation. HeLa cells were infected with YSH6000 (*S. flexneri* WT) or *ApaH*-chromosome mutant, and then whole cell lysates were harvested at 20, 40, 60, and 80 min post-infection for the detection of IkBα. The degradation rate of IkBα in cells infected with the *ApaH*-chromosome mutant was higher than that of WT, indicating that one or more of the chromosomal IpaH proteins contributed to the dampening of IkBα degradation (Fig. 1A, left). To identify the IpaH proteins that were involved in suppressing NF-κB activation, we generated deletions mutants for each of the seven chromosomal *ipaH* genes. IpaH0722 was critical for the inhibition of NF-κB activation. The degradation rate of IkBα was higher in *ApaH0722* infection compared to WT infection (Fig. 1A, right). IpaH0722, which corresponds to ORF SF0722 of *Shigella flexneri* 2a S501 strain, has a Cys residue in its C-terminal region that is required for E3 ubiquitin ligase activity [16,28,29].

In Cos-7 and HeLa cells, ectopically-expressed IpaH0722 localized to the cell membrane (Fig. 1B and S1A). A mouse lung infection model was used to evaluate the role of IpaH0722 in the pathogenesis of *Shigella*. The survival rates of mice infected with the *ApaH0722* mutant *Shigella* were increased compared to mice infected with WT *Shigella* (Fig. 1C). Next, to investigate whether the Cys379 residue in the IpaH0722 C-terminal region and its E3 ubiquitin ligase activity contributed to the pathogenesis of *Shigella*, we substituted Cys379 residue (IpaH0722CA). When 293T cells expressing IpaH0722 or IpaH0722CA were infected with *Shigella*, the activation of NF-κB was decreased in the presence of IpaH0722 but not IpaH0722CA (Fig. 1D). In addition, to confirm the importance of the E3 ubiquitin ligase activity of IpaH0722 in *Shigella* infection, HeLa cells were infected with *Shigella* harboring *ipaH0722* (*ApaH0722/ipaH0722*) or *ipaH0722CA* (*ApaH0722/ipaH0722CA*). In cells infected with the WT strain or *ApaH0722/ipaH0722* strain, the degradation rate of IkBα was reduced at 60 and 80 min after infection compared to the *ApaH0722* or *ApaH0722/ipaH0722CA* strains (Fig. 1E). It is important to note that IpaH0722 had no effect on Elk-1 and JNK activation (Fig. 1D and E). We speculated that IpaH0722 plays a role in the inhibition of *Shigella*-induced NF-κB activation in an E3 ubiquitin ligase-dependent manner.

IpaH0722 selectively inhibits phorbol myristate acetate (PMA)-induced NF-κB activation

NF-κB activity can be stimulated by multiple signaling pathways that are triggered by various receptors in response to exogenous and endogenous stimuli, including TNF-α-TNFRI, TLRs, and NLRs [30]. To characterize IpaH0722-dependent dampening of NF-κB activation, we measured the levels of NF-κB activation in *ApaH0722*- or IpaH0722CA-expressing 293T cells that were stimulated with PMA, TNF-α, lipopolysacharide (LPS), or IL-1β using NF-κB-luciferase reporter assays. IpaH0722 markedly impaired PMA-dependent, but not TNF-α-, LPS-, and IL-1β-dependent NF-κB activation in an E3 ubiquitin ligase activity-dependent manner (Fig. 2A). To confirm the specificity of IpaH0722-mediated induction of NF-κB activation, we investigated the expression of IL-8 in 293T cells stimulated with PMA using an IL-8 reporter assay. Consistent with the NF-κB findings, IpaH0722 preferentially inhibited the PMA-dependent IL-8 expression (Fig. 2B). Moreover, the preferential targeting of PMA- but not TNF-α-induced NF-κB activation was also demonstrated by ELISA assays (Fig. 2C). Although PMA was also known to induce MAPK activation, IpaH0722 failed to inhibit PMA-dependent AP-1 and Elk-1 activation (Fig. 2D).

Taken together these results clearly indicated that IpaH0722 preferentially inhibited PMA-induced NF-κB activation by targeting factors that modulated the NF-κB signaling pathway.

Ectopically-expressed IpaH0722 localizes to the cell membrane

It has recently reported that *Salmonella* SspH2, an IpaH cognate protein, localizes to the host cell membrane through its modification by cellular palmitoyl transferases. SspH2 has a putative 3-palmitoylation motif in its N-terminal and undergoes modification by host cell palmitoyltransferases, resulting in its localization to the host cell membrane [31]. Since we found a
Figure 1. IpaH0722 inhibits Shigella-induced NF-κB activation. (A) HeLa cells were infected with Shigella WT, ΔipaH-chromosome (left), or ΔipaH0722 (right). The cell lysates were prepared at the indicated time points and subjected to immunoblottings with anti-IκBα. Anti-actin antibody was used as a loading control. The values indicated below the images are the relative intensities of the bands. (B) IpaH0722 localizes to the host cell
plasma membrane. Cos-7 cells were transfected with a Myc-IpaH0722 expression vector and immunostained with anti-Myc, actin, and TO-PRO3. (C) Immunoblotting. As shown in Fig. 3A, Shigella invasion triggered the phosphorylation of PKC by Shigella WT, or the T3SS deficient mutant S325, however the S325 mutant did not (Fig. 3D). In contrast, both WT and ipaB/inv mutant was internalized into the endocytic vacuole and it was unable to disrupt the vacuolar membrane for dissemination into the cytoplasm (Fig. 3D). When HeLa cells were infected with Shigella WT or the T3SS deficient mutant S325, Shigella WT infection triggered the phosphorylation of PKCδ, however the S325 mutant did not (Fig. 3D). We next sought to determine whether vacuolar membrane disruption potentiated DAG–PKC–NF-κB signaling. We previously created an ipaB/inv mutant by introducing Ver5ina into the IpaB mutant [37]. The resulting IpaB/inv mutant was internalized into the endocytic vacuole and it was unable to disrupt the vacuolar membrane for dissemination into the cytoplasm (Fig. 3D). When HeLa cells were infected with Shigella WT or the IpaB/inv mutant, Shigella WT infection triggered the phosphorylation of PKCδ, however the IpaB/inv mutant did not (Fig. 3D). In contrast, both WT and AvrG mutant, which is unable to support intra- and inter-cellular movement, induced PKCδ phosphorylation (Fig. 3D). To further confirm the importance of vacuolar membrane disruption, we investigated PKC phosphorylation during Salmonella infection using a strain that triggers membrane ruffling but remain sequestered in the phagosome. When HeLa cells were infected with Salmonella WT but not Salmonella WT infection triggered the phosphorylation of PKCδ (Fig. 3D). These data supported the notion that membrane rupture by Shigella in infected epithelial cells triggers the activation of the DAG-PKC-NF-κB pathway. 

IpaH0722 inhibits the PKC–NF-κB pathway without targeting the CBM complex.

Because Shigella invasion of epithelial cells triggers the DAG–PKC–NF-κB signaling, we sought to determine the host factors in the PKC–NF-κB signaling pathway that were targeted by IpaH0722 E3 ubiquitin ligase. First, we determined which of the steps during the activation of PKC–NF-κB was targeted by
IpaH0722. The NF-κB activity induced by PKC signaling factors was measured in 293T cells with ectopic expression of IpaH0722 or IpaH0722CA. The transiently transfected cells were stimulated with a subset of constitutively active PKC isoforms (PKCζ, -δ, -ε, and 0) and NF-κB activity was determined. IpaH0722, but not IpaH0722CA, inhibited all PKC isoforms (PKCζ, -δ, -ε, and -0) and NF-κB activation, indicating that IpaH0722 targeted PKC itself or factors that lie directly downstream of PKC (Fig. 4A). In HEK293 cells, similar levels of phosphorylated PKCζ were detected in response to Shigella WT and IpaH0722 infection (Fig. 4B). Moreover, no differences were detected in PKC phosphorylation in PMA plus ionomycin-stimulated 293T cells expressing IpaH0722 or IpaH0722CA (Fig. 4C). In addition, IpaH0722 did not bind to PKC, and had no effect on protein stability of PKC, suggesting that IpaH0722 targets downstream of PKC rather than PKC itself (Fig. S2). We therefore focused on the CARMA-Bcl10-MALT1 (CBM) complex because the phosphorylation of CARMA1, 2, and 3 by PKC induces a conformational change that recruits Bcl10-MALT1 to CARMA (CBM signalosome), which is essential for the activation of downstream signaling [38,39]. Therefore, we tested the effect of IpaH0722 on CBM complex formation, and found that IpaH0722 failed to inhibit CARMA1-, CARMA2-, CARMA3-, or Bcl10-induced NF-κB activity (Fig. 4D). Indeed, IpaH0722 did not bind to CARMA1-, CARMA2-, CARMA3-, or Bcl10, and had no effect on their protein stability (Fig. S3). Furthermore, since IpaH0722 did not affect PKC-mediated CARMA phosphorylation (Fig. 4E), we believe that IpaH0722 inhibited the PKC-NF-κB pathway but not via the CBM complex.

IpaH0722 preferentially inhibits TRAF2-mediated NF-κB activation

We subsequently investigated other NF-κB signaling factors, namely TRAF2, TRAF5, TRAF6, NIK, IKKα, IKKβ, or p65 as potential IpaH0722 targets [40]. To this end, 293T cells expressing NF-κB-luciferase reporter gene together with TRAF2, TRAF5, TRAF6, NIK, IKKα, IKKβ, or p65 were transfected with a vector encoding IpaH0722 or IpaH0722CA and luciferase activity was measured (Fig. 5A). IpaH0722, but not IpaH0722CA, inhibited NF-κB activity when the cells that expressed TRAF2 suggesting that TRAF2 is a target for IpaH0722 E3 ubiquitin ligase (Fig. 5A). In previous reports, PKC-mediated phosphorylation of TRAF2 was a prerequisite for NF-κB activation in epithelial cells [41,42]. PKCζ and PKCε phosphorylated TRAF2 at residue Thr117 resulting in IKK recruitment and ultimately NF-κB activation [41,42]. We therefore used siRNA-mediated TRAF2 knockdown cells to measure the effect of TRAF2 on activation of the PKC-NF-κB pathway. The results showed that TRAF2 knockdown decreased NF-κB activation in response to PMA stimulation and Shigella infection (Fig. S4A). To further confirm the effect of TRAF2 knockdown on NF-κB activation we utilized Traf2-knockout mouse embryonic fibroblasts (MEF) [43]. We transduced Traf2 into the Traf2-knockout MEFs by retrovirus infection to generate stably expressing Traf2 cell lines. The degradation of IkBα in Traf2 stably expressing MEFs by retrovirus infection to generate stably expressing Traf2 cell lines. The degradation of IkBα in Traf2 stably expressing MEFs was determined. As shown in Fig. S4B, IpaH0722 precipitated with TRAF2, but not TRAF5, TRAF6, NIK, IKKα, IKKβ, or p65 (Fig. 5B). The interaction between IpaH0722 and TRAF2 was further confirmed by GST-pull down assay. Whole cell lysates of HEK293 and GST-IpaH0722, we were able to pull down endogenous TRAF2, but not Traf2 (Fig. 5C).

In previous reports, structural analysis of IpaH proteins using IpaH1.4 or IpaH3 revealed that the LRR region of the IpaH family is required for substrate recognition, while the CTR region is essential for its E3 ubiquitin ligase activity [44,45]. IpaH proteins share characteristic domains: an N-terminal 60–70 amino acid stretch, a LRR region, and an intervening region flanked by the LRR and the conserved CTR region [46]. The interaction between IpaH0722 and TRAF2 was further confirmed by GST-pull down assay. Using whole cell lysates of HEK293 and GST-IpaH0722, we were able to pull down endogenous Traf2, but not Traf2 (Fig. 5C).

IpaH0722 targets TRAF2 for ubiquitination and proteasomal degradation

Since IpaH0722 inhibited NF-κB activation in an E3 ubiquitin ligase-dependent manner, we tested whether IpaH0722 ubiquitin ligase activity using an in vitro ubiquitination assay. TRAF2, IpaH0722, or IpaH0722CA (purified from E. coli) were combined with E1, ATP, UbcH5b (an E2 ubiquitin conjugating enzyme) in assay medium and subjected to immunoblotting. TRAF2 was ubiquitinated by IpaH0722 but not IpaH0722CA (Fig. 6A). To confirm the fate of ubiquitinated TRAF2, we measured the stability of TRAF2 in 293T cells ectopically expressing IpaH0722 or IpaH0722CA and treated with cycloheximide (CHX; a protein synthesis inhibitor) for 2, 4, and 6 h. TRAF2 degradation was faster in IpaH0722 expressing cells compared to IpaH0722CA expressing cells, which confirmed that IpaH0722 targeted TRAF2 for ubiquitin-mediated protein degradation (Fig. 6B). We also measured TRAF2 degradation in 293T cells that were co-
Shigella Effector Inhibits PKC–NF-κB Activity

A

Time (min) - 10 20 40 60
α-pan-p-PKC
α-p-PKCδ
α-p-PKCδ
α-actin

B

Relative NF-κB activity (fold)

Shigella
PMA+ionomycin

C

Relative NF-κB activity (fold)

Control siRNA
PKCδ siRNA
+ Shigella

D

Time (min) - 10 20 40 60 80
WT S325
α-p-PKCδ
α-PKCδ
α-actin

ΔipaB/inv

ΔvirG

Salmonella

Invasion
Membrane disruption
Escape
Intracellular movement

Shigella WT
S325
ΔipaB/inv
ΔvirG

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Figure 3. *Shigella* triggers PKC–NF-κB activation. (A) HeLa cells were infected with *Shigella*. Cell lysates were prepared at the indicated time points and subjected to immunoblotting with anti-phospho-PKC antibody. *Non-specific bands. (B) NF-κB luciferase assays of 293T cells transiently transfected with an NF-κB reporter plasmid and empty vector or dominant-negative forms of PKC. After 24 h, cells were infected with *Shigella* or treated with PMA for 3 h, and luciferase activity was measured. Results are presented as fold change relative to the activity of uninfected or unstimulated cells. *P* < 0.01. (C) Cells were treated with PKCα siRNA and transiently transfected with an NF-κB reporter plasmid. The cells were infected with *Shigella* and NF-κB reporter activity was measured. (D) *Left*; HeLa cells were infected with *Shigella* WT, invasin expressing *ipaB* mutant, *Avr*G mutant, or *Salmonella*. The cell lysates were harvested at the indicated time points and subjected to immunoblot. *Right*; A model of *Shigella* invasion into the epithelial cells.

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transfected with Traf2 plus IpAH0722 or IpAH0722CA, and treated with MG132 (a proteasome inhibitor) or E64D plus pepstatin A (lysosome inhibitors). As shown in Fig. 6C, MG132 treatment prevented Traf2 degradation in the presence of IpAH0722, whereas E64D plus pepstatin A treatment did not prevent Traf2 degradation. These data confirm that IpAH0722 targets Traf2 for ubiquitination and proteasomal degradation.

Previous studies indicated that Traf5 compensates for Traf2 function [43], so we exploited Traf2/Traf5 double knockout MEFs to clarify the requirement of Traf2 on NF-κB activation. We introduced Traf2 into the Traf2/Traf5 double knockout MEFs by retrovirus transduction. Under puromycin selection, we obtained stable Traf2-expressing Traf2/Traf5 double knockout MEFs. Traf2/Traf5 double knockout MEFs and stable Traf2-expressing Traf2/Traf5 double knockout MEFs were infected with *Shigella* WT or *Avr*H0722, and IκBα degradation was measured over time. The degradation rate of IκBα in *Avr*H0722-infected stable Traf2-expressing Traf2/Traf5 double knockout MEFs was higher compared to WT *Shigella* infection (Fig. 6D). In contrast, the degradation rates of IκBα in WT and *Avr*H0722-infected double knockout MEFs were almost equal (Fig. 6D). To evaluate the importance of the in vivo role of E3 ligase activity of IpAH0722 for *Shigella* virulence, we infected mice with a sublethal dose of *Shigella* WT, *Avr*AH0722, *Avr*H0722/0722, or *Avr*H0722/0722CA. Mice infected with *Avr*H0722 or *Avr*H0722/0722CA mutants showed significantly reduced bacterial colonization when compared to that of WT or *Avr*H0722/0722 infection (Fig. S6A). Furthermore, histopathology showed that inflammatory responses, such as suppurative masses, neutrophil infiltration, and macrophage infiltration, were significantly higher in mice infected with *Avr*H0722 or *Avr*H0722/0722CA mutants when compared to that of WT or *Avr*H0722/0722 infections (Fig. S6B). These results supported our notion that the interaction between IpAH0722 and Traf2 is an important mechanism for dampening NF-κB activation during *Shigella* infection of epithelial cells.

**Discussion**

In this study we show that IpAH0722, a chromosomally-encoded IpaH E3 ligase, specifically dampens the PKC-dependent activation of NF-κB in response to *Shigella* invasion of epithelial cells. We find the IpAH0722 exerts its effect by targeting Traf2 for proteasomal-dependent degradation (Fig. 6E). The inhibitory activity possessed by IpAH0722 appears to be an important mechanism for the downregulation of the acute inflammatory response by blocking the activation of PKC-NF-κB pathway. This is a critical step since cell membrane rupture is a part of the process of *Shigella* invasion into epithelial cells, and the recognition of this as a DAMP triggers the recruitment and activation PKC, and ultimately the activation of the innate immune response via PKC-NF-κB pathway. This notion was supported by the experiments using the *Avr*B/mice mutant, which was internalized by the vacuolar membrane but could not rupture the vacuole [36], we showed that *Shigella* WT infection, but not *Avr*B/mice mutant infection, caused in the phosphorylation of PKCα (Fig. 3E). In fact, Dupont et al. recently reported that the vacuolar membrane remnants generated by *Shigella* invasion into epithelial cells undergo ubiquitination and provided a cue to stimulate autophagy and inflammatory responses [46]. Moreover, Shahnazari et al. showed that the DAG generated in *Salmonella* containing vacuole (SCV) membranes serves as a specific signal to induce autophagy through recruitment of PKC into the SCV [47]. Tattoli et al. showed that vacuolar membrane damage triggers amino acid starvation and the downregulation of mTOR activity, thereby inducing autophagy against intracellular *Shigella* and *Salmonella* [48]. Phagosomal membrane disruption by *Listeria monocytogenes* LLO in macrophages, which is known as a cholesterol-dependent cytolysin, triggers the recruitment of PKC to the membrane and its activation [49]. After disruption of the *Listeria* phagosome the membrane damage was recognized by PKC. These data suggest that PKC functions as a part of the repair mechanism of host cells, and PKC recruitment by the membrane rupture is recognized as a DAMP by the innate immune system [49]. These studies also imply that the membrane rupture, which is created by bacterial infection, acts as a double-edged sword for the pathogens; on one hand it benefits the bacteria by promoting further dissemination and multiplication, and on the other it hinders the bacteria by acting as a DAMP that stimulates the host innate immune response. In line with this, we identified IpAH0722 as a *Shigella* effector that plays an important pathogenic role because it counteracts DAMPs-mediated innate immune responses.

Recently we reported that another *Shigella* effector, Ospl targets UBC13 and dampens the DAG–CBM–TRA6–NF-κB pathway, which is triggered by membrane ruffles around the *Shigella* entry sites into epithelial cells [12]. Ospl acts as a glutamine deamidase for UBC13 to convert Q100 to E100, thus inactivating UBC13 E2 ubiquitin conjugating enzymatic activity that is required for Traf6 activation [12]. Currently, although the reason underlying why *Shigella* deliver IpAH0722 in addition to Ospl during the invasion of epithelial cells remains partly unclear, we believe that it is due to the fact that the two different signaling pathways are required to neutralize the acute inflammatory response and are triggered by distinct bacteria-induced cellular events: DAG-CBM–TRA6–NF-κB (membrane ruffles) and DAG–PKC–TRA2–NF-κB (membrane rupture) [12; this study].

Traf2 and Traf5 have been extensively studied in TNFR–NF-κB signaling in immune cells, and it has also been reported that Traf2 is essential in Nod1−, PKC−, and non-canonical NF-κB signaling in non-myeloid cells [42,50,51]. In previous studies, PKC-mediated phosphorylation of Traf2 at Ser11, Ser55, and Thr117 was required for prolonged NF-κB activation, but not JNK activation [42,52–54]. Consistent with these observations, our data showed that treatment of siRNA-mediated knockdown of Traf2 decreased PMA-induced NF-κB activation, suggesting that Traf2 is involved in PKC–NF-κB signaling (Fig. S4). According to Hasegawa et al., Traf2/Traf5 double knockout MEFs but not Traf6 knockout MEFs had decreased NF-κB activation upon
Figure 4. IpaH0722 targets PKC–NF-κB activation but not via the CBM complex. (A) IpaH0722 inhibits PKC-induced NF-κB activation. Luciferase assays of 293T cells that were transiently transfected with dominant-active forms of PKC and empty vector, IpaH0722, or IpaH0722CA expressing plasmids. Results are presented as fold change relative to the activity of uninfected or unstimulated cells. *P < 0.01. (B) HeLa cells were infected with Shigella WT or the ipaH0722 mutant. The cell lysates were harvested at the indicated time points and subjected to immunoblotting. (C) Cells that transiently expressed empty vector, IpaH0722, or IpaH0722CA were treated with PMA for 20 min, followed by subjecting the lysates to immunoblotting. (D) Luciferase reporter assays of 293T cells were transiently transfected with NF-κB and empty vector, IpaH0722, or IpaH0722CA expressing plasmids. Results are presented as fold change relative to the activity of uninfected or unstimulated cells. (E) 293T cells were transfected with FLAG-CARMAM1 with or without PKCα, Myc-IpaH0722, or Myc-IpaH0722CA. After 24 h, cells were harvested and subjected to immunoprecipitation (IP).

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Nod1-RIP2 stimulation, suggesting that TRAF2 was also important in the Nod1–NF-κB pathway [51]. Since Nod1 recognizes the peptidoglycan that is released from Shigella at the entry site and in the cytoplasm, and plays an important role in Shigella-induced NF-κB activation, IpaH0722 likely inhibits both the PKC-NF-κB pathway and the Nod1–RIP2–TRAF2–NF-κB pathway during Shigella infection [55–57]. In fact, we see that IpaH0722 also inhibited Nod1-stimulated NF-κB activation in our reporter assay (data not shown). We have provided evidence that IpaH0722 interacted with TRAF2 via its substrate recognition site, LRR, and modified its ubiquitination (Fig. 6 and Fig. S5). The ubiquitination of TRAF2 by IpaH0722 led to its proteasomal-degradation and inhibited NF-κB activation (Fig. 6). Though not yet fully understood, IpaH0722 does not inhibit NF-κB in response to TNF-α; unidentified conformational changes or signaling pathways involving TRAF2 likely occur during Shigella infection or PMA stimulation.

Herein, we provide the first evidence that Shigella invasion induced membrane rupture of epithelial cells, which stimulates PKC-NF-κB signaling. Moreover, Shigella used the E3 ubiquitin ligase effector, IpaH0722, to target TRAF2 for proteasomal-degradation dependent degradation (Fig. 6E). This is the first report IpaH0722 as a non-plasmid-encoded T3SS effector protein in Shigella.

Materials and Methods

Materials

Shigella flexneri strain YSH6000 was used as the wild type and S325 (mexE::Tn5) was used as a T3SS-deficient negative control. Salmonella typhimurium SB300 strain was used as the wild-type strain [50]. Construction of the non-polar mutant of ipaH chromosomal, ipaH0722, ipaB, and virG of S. flexneri YSH6000 were carried out as described previously [29,37]. The ipaH0722-FLAG or ipaH0722CA-FLAG were cloned into pWKS130 to yield p-IPA0722-FLAG and p-ipaH0722CA-FLAG. The resultant plasmids were introduced into the AIPA0722 strain. The ipaH0722 or ipaH0722CA coding sequences were amplified by PCR and cloned into pCMV-FLAG, pEGFP, pGEX-4T-1, pcDNA-Myc (6× Myc), and pcDL-SRα-Myc vectors, cDNAs for human Traf2, Traf5, Traf6, Carm1, Carm2, Carm3, Bcl10, IKKα, IKKβ, b65, and a series of P6 mutants were cloned into pCMV-FLAG, pEGFP, pcDNA-Myc, pcDL-SRα-Myc, and pcDNA-FLAG vectors. cDNA for mouse Traf2 was cloned into pGEX 4T-1. Site direct mutageneis of ipaH0722 or P6c was performed using the QuickChange site directed mutagenesis kit (Stratagene).

Materials

The anti-M2 FLAG monoclonal FLAG antibody (Sigma-Aldrich), anti-Myc 9B11, phospho-PKC (pan) (BII Ser660), phospho-PKD/PKCα (Ser744/748), phospho-(Ser) PKC substrate, phospho-JNK antibody (Cell signaling), anti-actin (MILLI-PORE), anti-PKCδ (C-20), PKCα (C-20), Traf2, Traf6 antibody (Santa Cruz Biotechnology), and anti-1xβ (BD transduction) were obtained commercially. The anti-IpaH antibody was described previously [29]. PMA, LPS, ionomycin, E64D, and pepstatin A were obtained from Sigma. IL-1β and TNF-α were obtained from Peprotech. MG132 was obtained from Peptide Inst.

Cell culture

HeLa cells were cultured in Eagle’s minimal essential medium (Sigma) supplemented with 10% fetal calf serum. Cos-7 and 293T cells were cultured in Dulbecco’s modified Eagle medium (Sigma) supplemented with 10% fetal calf serum. The Traf2-deficient or Traf2/Traf5-deficient MEFs were maintained as described previously [43]. To construct Traf2−/− or Traf2/Traf5−/− cells stably expressing Traf2, cDNAs encoding these genes were subcloned into pMX-puro retroviral expression vectors. Retroviral supernatants were produced in Plat-E cells. Target cells were transduced with supernatants in the presence of DO-TAP (Roche) and then cloned under puromycin selection.

Bacterial infection

HeLa cells were infected with various strains of Shigella at a multiplicity of infection (moi) of 100. In the case of the Shigella strains expressing afimbrial adhesin (Afa), these cells were infected at moi of 10. To adjust the intracellular bacterial number, AipaB/inv were set to 3 times moi of WT. Infection was initiated by centrifuging the plate at 700 × g for 10 min. After incubation for 20 min at 37°C, the plates were washed three times with PBS, transferred into fresh medium containing gentamicin (100 μg/ml) and kanamycin (60 μg/ml) to kill extracellular bacteria. Salmonella infection was conducted as described previously [58]. After incubation for indicated time, the cells were washed with PBS and harvested into 2 × Laemmli’s sample buffer for immunoblotting. The density of each band was quantified by measuring the mean intensity using NIH image software version 1.63 and the expression levels were normalized to the levels of β-actin. All immunoblotting experiments were repeated multiple times and representative data of similar results are presented.

Luciferase reporter assays

293T cells were seeded into 24-well plates. After 24 h, the cells were transfected with reporter plasmids (pNF-κB-luc, pLIL-κB, pEIK-1-luc, pAP-1-luc), Renilla luciferase constructs (phRL-TK, Promega) using FuGENE6 transfection reagent (Roche). The IL-8 reporter plasmid was constructed by amplifying the 181 bp human IL-8 promoter sequence from the genomic DNA of HeLa cells, which was then inserted into the pmET-luc reporter plasmid (Clontech). Equal amounts of empty vector were used to control for the transfection process. After 24 h, cells were infected with Shigella WT (moi = 30) or treated with PMA (25 nM), TNF-α.
Shigella Effector Inhibits PKC–NF-κB Activity

A

Relative NF-κB activity (fold)

vector lpaH0722 lpaH0722CA

+ TRAF2

+ TRAF5

+ TRAF6

+ NIK

vector lpaH0722 lpaH0722CA

+ IKKα

+ IKKβ

+ p65

B

Mycg-vector

Mycg-lpa0722CA

FLAG-TRA2

Mycg-vector

Mycg-lpa0722CA

GFP-TRA2

Mycg-vector

Mycg-lpa0722CA

FLAG-TRA6

Mycg-vector

Mycg-lpa0722CA

FLAG-NIK

C

Pull down

GST

GST fusion protein

Input : α-FLAG

IP : α-Myc

IB : α-FLAG
Expression and purification of recombinant proteins

For GST fusion proteins, *E. coli* BL21 (DE3) strain harboring pGEX4T-1 derivatives were cultivated in L-broth supplemented with ampicillin (50 μg/ml) 3 h at 30°C. Expression was induced by the addition of 1 mM IPTG and incubation for 3 h at 30°C. Bacteria were disrupted by sonication and lysozyme treatment. Purification of the GST fusion proteins with glutathione-Sepharose 4B (GE Healthcare) was performed according to the manufacturer’s protocol.

Pull-down assay

GST-IpaH0722 bound to glutathione sepharose 4B beads was mixed with HeLa cell lysates for 2 h at 4°C. After centrifugation, the beads were washed five times with 0.5% Triton X-100-PBS and subjected to immunoblotting. The immunoblotting experiments were repeated multiple times and representative data of similar results are presented.

Immunoprecipitation

293T cells were transiently transfected using FuGENE 6 (Roche). Cells were washed with PBS and lysed for 30 min at 4°C in lysis buffer containing 150 mM NaCl, 50 mM HEPES pH 7.5, 1 mM EDTA, 0.5% NP-40, and Complete protease inhibitor cocktail (Roche). Lysates were cleared by centrifugation and proteins were immunoprecipitated 2 h with anti-Myc (9B11) or anti-M2 FLAG and Protein G beads (Sigma) at 4°C. Immunoprecipitates were washed five times with lysis buffer and subjected to immunoblotting. The immunoblotting experiments were repeated multiple times and representative data of similar results are presented.

*In vitro* ubiquitination assays

*In vitro* TRAF2 ubiquitination assays were performed in 40 μl reaction mixture containing reaction buffer (25 mM Tris-HCl [pH 7.5], 50 mM NaCl, 5 mM ATP, 10 mM MgCl2, and 0.1 mM DTT), 1 μg TRAF2, 0.5 μg E1, 2 μg UbcH5b, and 2 μg ubiquitin purified from *E. coli* in the presence and absence of GST-IpaH0722 or GST-IpaH0722CA. Reactions were incubated at 37°C for 1 h and stopped by the addition of 5× Laemmli sample buffer. All immunoblotting experiments were repeated multiple times and representative data of similar results are presented.

Cycloheximide chase assay

At 24 h after transfection, 293T cells were treated with cycloheximide (50 μg/ml) (Wako) for the indicated times and cell lysates were harvested for immunoblotting. The density of each band was quantified by measuring the mean intensity using NIH image software version 1.63. The expression levels were normalized to the levels of β-actin. All immunoblotting experiments were repeated multiple times and representative data of similar results are presented.

RNAi

Human PKCδ- or human TRAF2-specific siRNA were prepared by Sigma as follows: 5'-CGUGUGGACACGCGCA-CAUUAU-3’ and 5’-AAUGUGGCCGUGUCCACACGGA-3’ (PKCδ) or 5’-GGCCAGUCAAAGCAUGAACA-3’ and 5’-UCUAUGCGUUGACGUCCUCG-3’ (TRAF2). Cells were transfected using RNAiMax (Invitrogen). siRNA treated cells were utilized after 72 h for further analyses.

Murine pneumonia model

The pulmonary infection model in the mice was described previously [20]. In brief, five-week-old female Balb/c mice (CREA Japan) were housed in the animal facility of the Institute of Medical Science, University of Tokyo, in accordance with University guidelines. *S. flexneri* were suspended in sterile saline, and 20 μl of the bacterial suspension was administered intranasally at 1×10⁷ cfu (survival assay) or 1×10⁸ cfu (colonized bacterial number assay). For histological analysis, mice were sacrificed at 24 h after infection, and their lungs were harvested, fixed in 4% paraformaldehyde in PBS, and frozen in liquid nitrogen for sectioning. The sections were stained with hematoxylin and eosin and examined under a microscope. Viable bacteria in the lung tissue were counted by culturing homogenized tissue for 18 h on LB agar plates. Each data point is the mean of the values for 6 infected mice in each group.

Ethics statement

All animal experiments were carried out in strict accordance with the University of Tokyo’s regulations for Animal Care and Use protocol, which was approved by the Animal Experiment Committee of the Institute of Medical Science, the University of Tokyo (approval number; PA11-92). The committee acknowledged and accepted both the legal and ethical responsibility for the animals, as specified in the Fundamental Guidelines for Proper Conduct of Animal Experiment and Related Activities in Academic Research Institutions under the jurisdiction of the Ministry of Education, Culture, Sports, Science and Technology, 2006 (Japan). All surgery was performed under carbon dioxide euthanasia, and all efforts were made to minimize suffering.

Statistical analysis

Values are reported as means ± standard deviation (SD) or means ± standard error (SEM) of data obtained for independent experiments. Statistical analysis was performed Student’s t-test or a one-way ANOVA. Kaplan-Meier survivor curves were generated. p-values<0.05 were considered significant.
Shigella Effector Inhibits PKC–NF-κB Activity

A

B

C

D

E

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Figure 6. IpaH0722 targets TRAF2 for ubiquitination. (A) In vitro ubiquitination assay using TRAF2 and a mixture of E1, UbCH5b, ATP, and ubiquitin in the presence or absence of IpaH0722 or IpaH0722CA. (B) 293T cells were transfected with FLAG-TRAF2 and with and without Myc-IpaH0722 or Myc-IpaH0722CA. After 24 h, the cells were treated with CHX (50 mg/ml) and cell lysates were prepared at the indicated time points. (C) 293T cells were transfected with FLAG-TRAF2 with and without FLAG-IpaH0722 or FLAG-IpaH0722CA. After 24 h, the cells were treated with DMSO or CHX (50 mg/ml) plus MG132 or E64D+pepsatin A. Cell lysates were prepared after 3 h and samples were subjected to immunoblotting. (D) TRAF2/TRAP53−/− MEFs and TRAF2/TRAP53−/− MEFs stably expressing TRAF2 were infected with Shigella WT or ΔipaH0722. Cell lysates were harvested at the indicated time point and subjected to immunoblotting with anti-IkBα Anti-actin antibody was used as a loading control. (E) A model of this study. Shigella invasion induced phagosome membrane rupture of epithelial cells, which stimulates DAG–PKC–NF-κB activation. However, Shigella deliver IpaH0722 to inhibit PKC–NF-κB activation by ubiquitinating TRAF2 and promoting its proteasome degradation.

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Supporting Information

Figure S1 IpaH0722 localizes to host cell membrane. (A) A series of GFP-tagged IpaH0722 mutants were transiently expressed in HeLa cells and their cellular localization was examined by immunofluorescence analysis. (B) Luciferase reporter assays of 293T cells transiently transfected with an NFκB reporter plasmid and an empty vector or a series of IpaH0722 mutants. After 24 h, the cells were treated with PMA for 3 h and luciferase activity was measured. Results are presented as change relative to the activity of uninfected or unstimulated cells. *P<0.01.

Figure S2 IpaH0722 targets PKC–NF-κB activation but not PKC itself. (A) 293T cells were transfected with series of MycC-terminal PKC with FLAG-IpaH0722CA. After 24 h, cells were harvested and subjected to immunoprecipitation (IP) and immunoblotting. (B) 293T cells were transfected with series of PKC with and without IpaH0722 or IpaH0722CA. After 24 h, the cells were treated with CHX (50 mg/ml) and cell lysates were prepared at the indicated time points. In the case of PKC8, endogenous PKCα levels were detected.

Figure S3 IpaH0722 did not target the CBM complex. (A) 293T cells were transfected with FLAG-tagged CARMA1−, CARMA2−, CARMA3−, or Bcl10 with or without MycC-tagged IpaH0722CA. After 24 h, cells were harvested and subjected to immunoprecipitation (IP) and immunoblotting. (B) 293T cells were transfected with FLAG-tagged CARMA1−, CARMA2−, CARMA3−, or Bcl10 with and without FLAG-IpaH0722 or FLAG-IpaH0722CA. After 24 h, the cells were treated with CHX (50 mg/ml) and cell lysates were prepared at the indicated time points.

Figure S4 TRAF2 is involved in PMA and Shigella-induced NF-κB activation. (A) Cells with siRNA-mediated knockdown of TRAF2 were transiently transfected with an NF-κB reporter plasmid. These cells were stimulated PMA or infected with Shigella for 3 h and NF-κB reporter activity was measured. (B) TRAF2−/− MEFs and TRAF2−/− MEFs stably expressing TRAF2 were infected with Shigella. Cell lysates were harvested at the indicated time points and subjected to immunoblotting with anti-IkBα. Anti-actin antibody was used as a loading control.

Figure S5 The LRR region of IpaH0722 is required for TRAF2 binding. (A) Schematic representation of full length and truncated IpaH0722. (B) GST-tagged IpaH0722 or IpaH0722 truncated forms were prepared from E. coli and used to GST pull down assays with HeLa whole cell lysates.

Figure S6 IpaH0722 modulates the host inflammatory response in a murine model of pneumonia infection. (A) The number of viable bacteria in the lungs of mice infected with the S. flexneri WT strain, ΔipaH0722 mutant, ΔipaH0722/ ipaH0722 complementation strain, or ΔipaH0722/ ipaH0722CA complementation strain at 24 h after inoculation. Data are the mean values of 6 mice per group (±SD). *P<0.05 (B) Histological analysis of mouse lungs at 24 h after inoculation. Sections of infected lungs were stained with hematoxylin and eosin.

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Author Contributions

Conceived and designed the experiments: HA. Performed the experiments: HA. Analyzed the data: HA CS. Contributed reagents/materials/analysis tools: HA. Wrote the paper: HA CS.

References

1. Chen GY and Nunez G. (2010) Sterile inflammation: sensing and reacting to damage. Nat Rev Immunol 10: 826–837.
2. Davis BK, Wen H, and Ting JP. (2011) The inflammasome NLRs in immunity, inflammation, and associated diseases. Annu Rev Immunol 29: 707–735.
3. Ashida H, Ogawa M, Kim M, Mimuro H, Sakasawa C. (2012) Bacteria and host interactions in the gut epithelial barrier. Nat Chem Biol 8: 36–45.
4. Galan MM. (2009) Common themes in the design and function of bacterial effectors. Cell Host Microbe 5: 571–578.
5. Rahman MM and McFadden G. (2011) The inflammasome: a danger sensor that triggers pyroptosis. Science 334: 496–501.
6. Takeuchi O, and Akira S. (2010) Pattern recognition receptors and inflammation. Cell 140: 805–820.
7. Schroder K, and Tschopp J. (2010) The inflammasomes. Cell 140: 821–832.
8. Ashida H, Ogawa M, Kim M, Suzuki S, Sanada T, et al. (2011a) Shigella deploy multiple countermeasures against host innate immune responses. Curr Opin Microbiol 14: 16–23.
9. Ashida H, Ogawa M, Mimuro H, Kobayashi T, Sanada T, et al. (2011b) Shigella are versatile mucosal pathogens that circumvent the host innate immune system. Curr Opin Immunol 23: 448–455.
10. Parsot C. (2009) Shigella type III secretion effectors: how, where, when, for what purposes?Curr OpinMicrobiol 12:110–116.
11. Kim DW, Lenzen G, Page AL, legrain P, Sansonetti PJ, et al. (2005) The Shigella flexneri effector OspG interferes with innate immune responses by targeting ubiquitin-conjugating enzymes. Proc Natl Acad Sci USA 102: 14046–14051.
12. Sanada T, Kim M, Mimuro H, Suzuki M, Ogawa M, et al. (2012) The Shigella flexneri effector OspF deamidates UBC13 to dampen the inflammatory response. Nature 483: 623–626.
13. Newton HJ, Pearson JS, Badea L, Kelly M, Lucas M, et al. (2010) The type III effectors NicE and NicB from enteropathogenic E. coli and OspF from Shigella block nuclear translocation of NF-κB p65. PLoS Pathog 6: e1000891.
14. Haraga A, Miller SL. (2003) A Salmonella enterica serovar Typhimurium translocated leucine-rich repeat effector protein inhibits NF-κB-dependent gene expression. Infect Immun 71:4052–4053.
15. Okuda K, Toyotome T, Katohaka N, Ohno M, Abe H, et al. (2005) Shigella effector IpaH9.8 binds to a splicing factor U2AF35 to modulate host immune responses. Biochem Biophys Res Commun 333: 351–359.
16. Rohde JR, Breitkreutz A, Chenal A, Sansonetti PJ, Parsot C. (2007) Type III secretion effectors of the IpaH family are E3 ubiquitin ligases. Cell Host Microbe 1: 77–83.
17. Ashida H, Kim M, Schmidt-Supprian M, Ma A, Ogawa M, et al. (2010) A bacterial E3 ubiquitin ligase IpaH9.8 targets NEMO/IKKγ to dampen the host NF-κB-mediated inflammatory response. Nat Cell Biol 12: 66–73.

18. Quezada CM, Hicks SW, Galán JE, Steblins CE. (2009) A family of Salmonella virulence factors functions as a distinct class of autoregulated E3 ubiquitin ligases. Proc Natl Acad Sci U S A 106: 4864–4869.

19. Busey JM, Stover CK, Oakes EV, Venkatesan M, Kopecko DJ. (1987) Molecular cloning of invasion plasmid antigen (ipa) genes from Shigella flexneri: analysis of the gene products and genetic mapping. J Bacteriol 169: 2561–2569.

20. Venkatesan MM, Busey JM, Kopecko DJ. (1989) Use of IpaH gene sequences for the general identification of Shigella spp. and enteroinvasive Escherichia coli. J Clin Microbiol 27: 2667–2681.

21. Hartman AB, Venkatesan M, Oaks EV, Buysse JM. (1991) Sequence and molecular characterization of a multicopy invasion plasmid antigen gene, *ipaH*, of *Shigella flexneri*. J Bacteriol 172: 1905–1915.

22. Busey JM, Venkatesan M, Oakes EV, Busey JM. (1990) Sequence variation in two *ipaH* genes of *Shigella flexneri* 5 and homology to the LRG-like family of proteins. Mol Microbiol 5: 2435–2445.

23. Hicks SW, Galán JE. (2010) Hijacking the host ubiquitin pathway: structural strategies of bacterial E3 ubiquitin ligases. Curr Opin Microbiol 13: 41–46.

24. Bernal-Bayard J, Ramos-Morales F. (2009) *Salmonella* type III secretion effector SipP is an E3 ubiquitin ligase for mammalian thioreredoxin. J Biol Chem 284: 27587–27595.

25. Bernal-Bayard J, Cardenal-Muñoz E, Ramos-Morales F. (2010) The *Salmonella* type III secretion effectors SipB, SipT, and SipP, target the human chaperone ERP60. J Biol Chem 285: 16360–16368.

26. Buchrieser C, Glauser P, Rusniok C, Nedjari H, D’Hauteville H, et al. (2000) The virulence plasmid pWR100 and the repertoire of proteins secreted by the type III secretion system of *Shigella flexneri*. Mol Microbiol 36: 760–771.

27. Jin Q, Yuan Z, Xu J, Wang Y, Shen Y, et al. (2002) Genome sequence of *Shigella flexneri* 2a: insights into pathogenicity through comparison with genomes of *Escherichia coli* K12 and O157. Nucleic Acids Res 30: 4432–4441.

28. Ashida H, Toyotome T, Nagai T, Sasakawa C. (2007) *Shigella* chromosomal *IpaH* proteins are secreted via the type III secretion system and act as effectors. Mol Microbiol 63: 680–693.

29. Toyotome T, Suzuki T, Konoe A, Nonaka T, Fukuda H, et al. (2001) *Shigella* protein IpaH8 is secreted from bacteria within mammalian cells and transported to the nucleus. J Biol Chem 276: 32071–32079.

30. Bhoj VG, Chen ZJ. (2009) Ubiquitylation in innate and adaptive immunity. Nature 458: 430–457.

31. Hicks S.W., Charron G., Hang H.C., Galañ J.E. (2011) Subcellular targeting of *Salmonella* effector proteins to the nucleus. J Biol Chem 286: 32071–32079.

32. Ashida H, Toyotome T, Nakai T, Sasakawa C. (2007) *Shigella* chromosomal *IpaH* proteins are secreted via the type III secretion system and act as effectors. Mol Microbiol 63: 680–693.

33. Rosse C, Linch M, Kermorgant S, Cameron AJ, Boccellari K, et al. (2010) PKC and the control of localized signal dynamics. Nat Rev Mol Cell Biol 11: 103–112.

34. Cossart P, Sansonetti PJ. (2004) Bacterial invasion: the paradigms of *Salmonella* and *Shigella* effector proteins. Nat Rev Microbiol 2: 112–121.

35. Cossart P, Sansonetti PJ. (2006) Bacterial invasion: the paradigms of *Salmonella* and *Shigella* effector proteins. Nat Rev Microbiol 6: 11–16.

36. Carayol N, Tran Van Nhieu G. (2013) Tips and tricks about *Salmonella* infection. Cell Microbiol 15: 1302–1308.

37. Dupont N, Lucas-Gervais S, Bertout J, Paz I, Freche B, et al. (2009) *Shigella* phagocytic vacuolar membrane remnants participate in the cellular response to pathogen invasion and are regulated by autophagy. Cell Host Microbe 6: 137–149.

38. Shahnazari S, Yen WL, Birmingham CL, Shi J, Namolovan A, et al. (2011) A diacylglycerol-dependent signaling pathway contributes to regulation of antiphagocytic autophagy. Cell Host Microbe 8: 137–146.

39. Tattoli I, Sorbara MT, Vuckovic D, Lang A, Soares F, et al. (2012) Amino acid starvation induced by invasive bacterial pathogens triggers an innate host defense program. Cell Host Microbe 11: 563–573.

40. Shaugnessy LM, Lipp P, Lee KD, Scwasson JA. (2007) Localization of protein kinase Cε to macropage vacuoles perforated by *Listeria monocyctogenes* cytolysin. Cell Microbiol 9: 1695–1704.

41. Hayden MS, Ghosh S. (2008) Shared principles in NF-κB signaling. Cell 132: 344–362.

42. Hasegawa M, Fujimoto Y, Lucas PC, Nakano H, Fukake K, et al. (2008) A critical role of RICK/RIP2 phosphorylation in Nod1-induced NF-κB activation. EMBO J 27: 373–383.

43. Blackwell K, Zhang L, Thomas GS, Sun S, Nakano H, et al. (2009) TRAF2 phosphorylation modulates tumor necrosis factor alpha-induced gene expression and cell resistance to apoptosis. Mol Cell Biol 29: 303–314.

44. Thomas GS, Zhang L, Blackwell K, Habibollah H. (2009) Phosphorylation of TRAF2 within its RING domain inhibits stress-induced cell death by promoting IKK and suppressing JNK activation. Cancer Res 69: 3665–36672.

45. Zhang L, Blackwell K, Ahuva A, Shi Z, Habibollah H. (2011) TRAF2 phosphorylation promotes NF-κB-dependent gene expression and inhibits oxidative stress-induced cell death. Mol Biol Cell 22: 128–140.

46. Girardin SE, Boneca IG, Carneiro LA, Antignac A, Jehanno M, et al. (2003) Nod1 detects a unique muropeptide from gram-negative bacterial peptidoglycan. Science 300: 1584–1587.

47. Kufner TA, Kremmer E, Adam AC, Phillipp DJ, Sansonetti PJ. (2007) The pattern-recognition molecule *Nod1* is localized to the plasma membrane at sites of bacterial interaction. Cell Microbiol 9: 477–486.

48. Travassos LH, Carneiro LA, Ramjeet M, Hussey S, Kim YG, et al. (2010) Nod1 and Nod2 direct autophagy by recruiting ATG16L1 to the plasma membrane at the site of bacterial entry. Nat Immunol 11: 55–62.

49. Watarai M, Kamata Y, Konishi S, Sasakawa C. (1997) rho, a small GTP-binding protein, is essential for *Shigella* invasion of epithelial cells. J Exp Med 185: 281–292.