The Type III Effectors NleE and NleB from Enteropathogenic E. coli and OspZ from Shigella Block Nuclear Translocation of NF-κB p65

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Abstract

Many bacterial pathogens utilize a type III secretion system to deliver multiple effector proteins into host cells. Here we found that the type III effectors, NleE from enteropathogenic E. coli (EPEC) and OspZ from Shigella, blocked translocation of the p65 subunit of the transcription factor, NF-κB, to the host cell nucleus. NF-κB inhibition by NleE was associated with decreased IL-8 expression in EPEC-infected intestinal epithelial cells. Ectopically expressed NleE also blocked nuclear translocation of p65 and c-Rel, but not p50 or STAT1/2. NleE homologues from other attaching and effacing pathogens as well as OspZ from Shigella flexneri 6 and Shigella boydii, also inhibited NF-κB activation and p65 nuclear import; however, a truncated form of OspZ from S. flexneri 2a that carries a 36 amino acid deletion at the C-terminus had no inhibitory activity. We determined that the C-termini of NleE and full length OspZ were functionally interchangeable and identified a six amino acid motif, IDSY(M/I)K, that was important for both NleE- and OspZ-mediated inhibition of NF-κB activity. We also established that NleB, encoded directly upstream from NleE, suppressed NF-κB activation. Whereas NleE inhibited both TNFα and IL-1β stimulated p65 nuclear translocation and IκB degradation, NleB inhibited the TNFα pathway only. Neither NleE nor NleB inhibited AP-1 activation, suggesting that the modulatory activity of the effectors was specific for NF-κB signaling. Overall our data show that EPEC and Shigella have evolved similar T3SS-dependent means to manipulate host inflammatory pathways by interfering with the activation of selected host transcriptional regulators.

Introduction

Many bacterial pathogens have the ability to “inject” virulence effector proteins into the host cell using a type III secretion system (T3SS). The effector proteins perform a variety of functions that allow the pathogen to persist in the host and cause disease [1]. Enteropathogenic Escherichia coli (EPEC) and enterohemorrhagic E. coli (EHEC) deliver T3SS effector proteins to the intestinal epithelium that mediate attaching and effacing lesion (A/E) lesion formation. A/E lesions are characterized by intimate bacterial attachment, effacement of the brush border microvilli and actin pedestal formation [2]. T3SS effectors from other pathogens such as Salmonella and Shigella have various roles in invasion, intracellular survival and the inhibition of innate immune responses through targeting host inflammatory signaling pathways [1]. Many of the T3SS effectors belong to conserved protein families that are found in a range of bacterial pathogens of plants and animals. For example, the OspF family of T3SS effectors from Shigella, Salmonella and Pseudomonas exhibit phosphothreonine lyase activity and induce irreversible dephosphorylation of mitogen-activated protein kinases (MAPKs) in the host cell nucleus [3,4,5]. In Shigella, this leads to gene-specific repression of a subset of NF-κB regulated genes, including IL8 [3]. Given the remarkable specificity of their biochemical function, the discovery of the mechanism of action of T3SS effectors remains an important step towards understanding the pathogenesis of many bacterial infections.
Author Summary

Bacterial intestinal pathogens have evolved distinct ways of colonizing the gut and causing disease. Enteropathogenic *E. coli* (EPEC) and its close relative enterohemorrhagic *E. coli* O157:H7 (EHEC) are extracellular pathogens that cause a characteristic lesion on the intestinal mucosa known as an attaching and effacing lesion. In contrast, *Shigella* is an intracellular pathogen that invades the intestinal mucosa and spreads from cell to cell. Both pathogens utilize a bacterial type III secretion system that "injects" virulence effector proteins into the host cell upon contact. We have discovered that an effector shared by EPEC/EHEC and *Shigella*, known as NleE or OspZ, as well as another EPEC/EHEC effector, NleB, inhibit the host cell inflammatory response by preventing translocation of the immune regulator NF-κB to the cell nucleus. Thus, although EPEC/EHEC and *Shigella* have evolved different colonization strategies, they share a common virulence determinant that suppresses the inflammatory response of the host, and both pathogens mediate a multi-effector attack on NF-κB signaling.

The activation of gene expression during inflammation is tightly regulated by transcription factors such as NF-κB. The NF-κB/Rel family comprises five members that share an N-terminal Rel homology domain that mediates DNA binding, dimerization and nuclear translocation [6]. The p65, c-Rel and RelB NF-κB subunits have an additional C-terminal transactivation domain, which strongly activates transcription from NF-κB-binding sites in target genes. The p50 and p52 subunits lack the transactivation domain but still bind to NF-κB consensus sites and act as transcriptional repressors [6]. The most abundant form of NF-κB in mammalian tissues is a p65/p50 dimer that activates the expression of multiple cytokine genes in response to inflammatory signals. In resting cells, NF-κB subunits are held in an inactive form in the cytoplasm by binding IkB proteins. Activation of NF-κB signaling stimulates the phosphorylation and proteosomal degradation of IkB, whereupon the NF-κB dimer is transported into the nucleus through the nuclear pore complex [6]. The canonical NF-κB pathway is stimulated by a range cell surface receptors such as the TNF receptor, IL-1 receptor, Toll-like receptors and T-cell receptor. Although the upstream components of these pathways vary, they converge at the point of IkB kinase (IKK)-mediated phosphorylation of and subsequent degradation of IkB [7].

Enteropathogenic *Escherichia coli* (EPEC) and enterohemorrhagic *E. coli* (EHEC) utilize a type III secretion system (T3SS) to deliver effector proteins to the intestinal epithelium that induce actin pedestal formation [2]. Multiple additional effectors are transported into the host cell where their targets and effects on host cell biology remain largely uncharacterized [8]. NleE is a highly conserved 27 kDa T3SS effector protein of A/E pathogens encoded in an operon with the 38 kDa effector, NleB. The NleE homologue in the invasive pathogen, *Shigella*, is called OspZ [9,10]. While investigating the effect of EPEC infection on NF-κB activation, we observed that wild type EPEC prevented translocation of the p65 subunit of NF-κB to the host cell nucleus, whereas an nleE mutant was defective for this activity. Here we report that NleE inhibits p65 nuclear translocation, thereby reducing the IL-8 response during bacterial infection, and that OspZ shares this activity. In addition, we show that NleB suppresses NF-κB activation but appears to act in distinct manner to NleE.

Results

EPEC infection blocks NF-κB p65 nuclear translocation, leading to reduced IL-8 production

Recent work has shown that EPEC and EHEC infection inhibits inflammatory cytokine production and NF-κB activation [11,12,13]. Previously, we found that translocated NleE localised to the host cell nucleus and we postulated that NleE had a role in subverting innate immune signaling [10]. Here we investigated the effect of NleE on NF-κB activation during EPEC infection. As actin accumulation beneath adherent EPEC depends on successful translocation of the T3SS effector, Tir [2], we used the fluorescent actin staining (FAS) test as a general marker for the translocation of T3SS effectors. HeLa cells were infected with wild type EPEC E2348/69, a T3SS (escF) mutant, an nleE deletion mutant of EPEC or an nleE mutant complemented with full length nleE. Cell monolayers were infected for 4 h and left unstimulated or infected for 90 min and stimulated with tumour necrosis factor α (TNFα) or interleukin-1β (IL-1β) for 30 min. Nuclear translocation of the p65 NF-κB subunit was visualised by immunofluorescence microscopy of FAS-positive cells for EPEC E2348/69, ΔnleE and ΔnleE [pNleE] (Fig. 1A) and cells with adherent bacteria for ΔeScF. In unstimulated cells, there was no significant difference in p65 nuclear exclusion between wild-type infected cells and the escF mutant (Fig. 1B). In contrast, in cells stimulated with TNFα or IL-1β, wild type EPEC E2348/69 inhibited p65 transport to the nucleus, whereas the escF mutant had little inhibitory effect on p65 nuclear translocation (Fig. 1B). The nleE mutant also showed greatly reduced inhibition of p65 nuclear transport in response to TNFα or IL-1β compared to wild type EPEC E2348/69 which was restored upon complementation of the nleE mutant with a copy of full length nleE. Similar results were obtained in response to IL-1β in Caco-2 intestinal epithelial cells (Fig. S1).

Caco-2 intestinal epithelial cells were then utilised to determine if the inhibition of p65 translocation resulted in the suppression of IL-8 production. Caco-2 cells were incubated with wild type EPEC E2348/69, a T3SS (escF) mutant, the nleE mutant or the nleE mutant complemented with nleE. Cells were then stimulated with IL-1β [14]. Compared to infection with the escF mutant, wild type EPEC inhibited IL-8 production from Caco-2 cells. The nleE mutant showed a diminished capacity to inhibit IL-8 production, which was complemented to wild type levels upon reintroduction of full length nleE (Fig. 2A). A similar trend was observed in CaCo-2 cells left unstimulated, although the differences were not as great as in IL-1β-stimulated cells (Fig. 2B). Real time PCR analysis of IL-8 from Caco-2 cells infected with derivatives of EPEC and stimulated with IL-1β, showed that levels of IL-8 mRNA were suppressed by NleE expression (Fig. 2C).

NleE inhibits nuclear translocation of p65 and c-Rel but not p50 or STAT1/2

To determine if NleE was sufficient for the inhibition of p65 nuclear translocation, we expressed GFP-NleE or GFP transiently in HeLa cells. Upon stimulation with TNFα, the NF-κB p65 subunit was excluded from the nucleus in the presence of ectopically expressed GFP-NleE but not GFP alone (Fig. 3A). To ensure that this effect was not an artefact arising from the over expression of GFP-NleE, we examined a range of transacted cells exhibiting low, moderate and high GFP expression. Even in cells exhibiting low levels of GFP-NleE, p65 was excluded from the nucleus upon stimulation with TNFα (Fig. 3A and data not shown). We also investigated the effect of NleE on nuclear localization of other NF-κB proteins, c-Rel and p50. Similar to p65, NleE blocked nuclear translocation of c-Rel in response to...
TNFα (Fig. 3B), however p50 nuclear localization was unaffected by the presence of NleE (Fig. 3C). A dual-luciferase reporter system measuring the activation of NF-κB-dependent transcription confirmed the absence of NF-κB p65 nuclear activity in GFP-NleE transfected cells stimulated with TNFα (Fig. 3D).

To determine if NleE-mediated inhibition of signaling affected other transcription factors, we tested the ability of NleE to inhibit nuclear translocation and activation of STAT1 and STAT2. Following stimulation with interferon α, both STAT1 and STAT2 were translocated to the nucleus in the presence of GFP or GFP-NleE (Fig. 4A and B). NleE also had no impact on STAT1/2 activation using a ISRE-Luc luciferase reporter (Fig. 4C) [15]. This suggests NleE acts on a subset of signaling pathways that includes NF-κB but not STAT1/2.

NleE from other A/E pathogens and full length OspZ from Shigella inhibit p65 translocation

To determine if the function of NleE and OspZ was conserved across A/E pathogens and Shigella, GFP-NleE and GFP-OspZ fusions generated from enterohemorrhagic E. coli
O157:H7, Citrobacter rodentium, Shigella boydii and Shigella flexneri were expressed by transfection in HeLa cells. NleE from EHEC O157:H7 and C. rodentium, and full length OspZ from S. boydii and S. flexneri serogroup 6 inhibited NF-κB activation and p65 nuclear translocation (Fig. 5A and B). In contrast, OspZ from S. flexneri serogroup 2a which carries a 36 amino acid truncation at the C-terminus (Fig. S2), had no impact on NF-κB activation and did not block p65 nuclear translocation. Further screening of three clinical isolates of S. flexneri 2a revealed that all strains encoded a truncated OspZ protein. Similar to the truncated form of OspZ from S. flexneri 2a, a GFP-NleE truncation lacking the C-terminal 36 amino acid residues, GFP-NleE1-188 was unable to prevent NF-κB activation. However, the C-terminal region was not sufficient for this antagonism, as GFP-NleE188-224 did not inhibit NF-κB activation (Fig. 5C). Interestingly, in contrast to the other full length GFP-NleE/OspZ fusion proteins, GFP-OspZ from S. flexneri 6 and S. flexneri 2a was largely excluded from the nucleus (Fig. 5B). Although the molecular basis of this is unknown, it may indicate that the mechanism of action of NleE/OspZ is in the cytoplasm of the cell since GFP-OspZSF6 inhibited NF-κB activation to the same degree as GFP-NleE (Fig. 5A).

A C-terminal IDSY(M/I)K motif in NleE and OspZ is important for the inhibition of NF-κB activation

To examine further the C-terminal region of NleE and OspZ, we performed a deletion analysis of NleE. Whereas truncated NleE1-214, inhibited NF-κB activation to the same degree as full length NleE, truncated NleE1-208 was inactive (Fig. 6A). This suggested that the region between amino acid residues 208 and 214, with the motif IDSYMK was critical for NleE function. We
then constructed a deleted form of NleE lacking amino acids 1209DSYMK214 which was unable to inhibit NF-κB activation (Fig. 6A) as was a corresponding deleted form of OspZ from S. flexneri lacking amino acids 1209DSYIK214 (Fig. 6B). In fact GFP-OspZΔDSYMK appeared to have a pro-inflammatory effect even in unstimulated cells (Fig. 6B).

Figure 3. Effect of ectopically expressed NleE on NF-κB activation. A. Representative immunofluorescence fields and quantification of p65 nuclear exclusion using anti-p65 (A), anti-c-Rel (B) or anti-p50 antibodies (C) (red) of HeLa cells transfected with pEGFP-C2 (GFP only) or pGFP-NleE (green), stimulated with TNFα for 30 min, and stained for nucleic acid (blue). Results are expressed as the percentage of GFP-positive cells that exclude p65, c-Rel or p50 and are the mean ± SEM of three independent experiments performed in duplicate. At least 100 GFP-positive cells were counted per test. *significantly different to GFP only stimulated with TNFα, P = 0.0005 (unpaired, two-tailed t-test) D. Fold increase in NF-κB dependent luciferase activity in pEGFP-C2 (GFP) or pGFP-NleE transfected cells unstimulated (white bars) or stimulated with TNFα for 30 min (black bars). Results are the mean ± SEM of triplicate wells. *significantly different to GFP only stimulated with TNFα (P<0.05, one way ANOVA).

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The C-terminal 50 amino acids of NleE and OspZ are strongly predicted by Jpred [16] to form an alpha-helical region, therefore it was possible that the IDSY[M/I]KM deletion had disrupted protein secondary structure (Fig. 6C). To account for this possibility and preserve the predicted alpha helix, we changed all six amino acids to alanine to generate GFP-NleE6A. GFP-NleE6A had the same alpha-helical prediction by Jpred as NleE (Fig. 6C). Similar to GFP-NleE6A, GFP-NleE was unable to inhibit NF-κB activation (Fig. 6D), and NleE6A delivered by the T3SS during infection was also unable to inhibit p65 nuclear translocation (Fig. 1B). Further mutation of individual amino acids within the IDSYMK motif of NleE to alanine did not make a significant difference to NF-κB inhibition compared to the fulllength GFP-NleE fusion (Fig. 6D). Expression of all GFP-NleE derivatives was tested by immunoblot using anti-GFP antibodies to ensure that differences in activity were not due to differences in the levels of GFP fusion proteins (Fig. S3).

To determine if variations in amino acid sequence at the C-termini of NleE and full length OspZ had any functional significance (Fig. 6C), we performed a domain swap by exchanging the last 40 amino acids of NleE with the last 46 amino acids of OspZ and vice versa. Both chimeric forms of NleE and OspZ (NleE-OspZ6term and OspZ-NleE6term) were fully functional (Fig. 6E) and they retained their native localization (data not shown). Therefore these regions were functionally interchange-
able, suggesting that NleE and OspZ use the same molecular mechanism to inhibit NF-κB activation.

**NleE/NleB** inhibits NF-κB activation through a distinct mechanism

To compare the activity of NleE with another T3SS effector, NleH, reported to interfere with NF-κB activation [17], we generated GFP-NleH1 and GFP-NleH2 fusions from EPEC E2348/69 and tested these for their ability to inhibit NF-κB activity following stimulation with TNF-α. In this system, NleE showed greater inhibition of NF-κB activation than either NleH1 or NleH2 (Fig. 7A). To test for possible non-specific effects on NF-κB activation of effector over-expression by transfection, we also tested the effect of NleD and NleB on NF-κB activation following stimulation with TNF-α. While NleD had no effect on luciferase

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**Figure 5. Effect of NleE and OspZ homologues on NF-κB activation.**

**A.** Fold increase in NF-κB dependent luciferase activity in HeLa cells transfected with pEGFP-C2 (GFP) or pGFP-NleE/OspZ cloned from C. rodentium (GFP-NleECR), EHEC O157:H7 EDL933 (GFP-NleEO157), EPEC E2348/69 (GFP-NleEPEC), S. flexneri 2a (GFP-OspZSF2a), S. boydii (GFP-OspZSB) and S. flexneri 6 (GFP-OspZSF6) and left unstimulated (white bars) or stimulated with TNF-α for 30 min (black bars). Results are the mean ± SEM of 3 independent experiments performed in triplicate. *significantly different to GFP-NleEPEC (P<0.05, one way ANOVA).

**B.** Representative immunofluorescence fields using anti-p65 antibodies (red) of HeLa cells transfected with derivatives of pGFP-NleE and GFP-OspZ (green) labelled as in Fig. 5A and stimulated with TNF-α for 30 min.

**C.** Fold increase in NF-κB dependent luciferase activity in HeLa cells transfected with pEGFP-C2 (GFP) or pGFP-NleE, pGFP-NleE1-188 or pGFP-NleE188-224 and stimulated with TNF-α for 30 min where indicated. Results are the mean ± SEM of 3 independent experiments performed in triplicate. *significantly different to GFP-NleE stimulated with TNF-α (P<0.05, one way ANOVA).

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Figure 6. Effect of NleE deletions and mutations on NF-κB activation. A. Fold increase in NF-κB dependent luciferase activity in cells transfected with pEGFP-C2 or derivatives of pGFP-NleE as labelled and left unstimulated or stimulated with TNFα for 30 min where indicated. Results are the mean ± SEM of at least 3 independent experiments performed in triplicate. *significantly different to GFP-NleE stimulated with TNFα (P<0.05, one way ANOVA) B. Fold increase in NF-κB dependent luciferase activity in cells transfected with pEGFP-C2, pGFP-OspZ or pGFP-OspZ_{ADSYM} and left unstimulated (white bars) or stimulated with TNFα for 30 min (black bars). Results are the mean ± SEM of at least 3 independent experiments performed in triplicate. *significantly different to GFP-OspZ stimulated with TNFα (P<0.05, one way ANOVA) C. Alignment of the C-terminal region of NleE and OspZ from A/E pathogens and Shigella as well as the NleE_{AD} variant. The IDSY(M/I)K motif is shaded and the Jpred prediction for an amino acid contributing to secondary structure is given as an accuracy score between 0 and 9, where 9 represents the most reliable prediction [16].

D. Fold increase in NF-κB dependent luciferase activity in cells transfected with pEGFP-C2 or derivatives of pGFP-NleE and OspZ as labelled and left unstimulated (white bars) or stimulated with TNFα for 30 min (black bars). Results are the mean ± SEM at least 3 independent experiments performed in triplicate. *significantly different to GFP-NleE stimulated with TNFα (P<0.05, one way ANOVA). E. Fold increase in NF-κB dependent luciferase activity in cells transfected with pEGFP-C2 or derivatives of pGFP-NleE and pGFP-OspZ as labelled and left unstimulated (white bars) or stimulated with TNFα for 30 min (black bars). Results are the mean ± SEM at least 3 independent experiments performed in triplicate. *significantly different to GFP-NleE stimulated with TNFα (P<0.05, one way ANOVA).

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induction in response to TNFα, NleB inhibited NF-κB activation as effectively as NleE (Fig. 7A). NleB is encoded directly upstream from NleE and this organization is highly conserved among A/E pathogens [18]. Fluorescence microscopy of GFP-NleB transfected cells stimulated with TNFα confirmed that GFP-NleB inhibited p65 translocation (Fig. 7B).

NleE inhibited NF-κB activation in response to both TNFα and IL-1β so we tested the ability of NleB to inhibit IL-1β signaling. Whereas, GFP-NleE was effective against both TNFα and IL-1β stimulation, GFP-NleB had no effect on NF-κB activation stimulated by IL-1β (Fig. 7C). This suggested that the two effectors act at different points in the NF-κB signaling cascade. To examine the effect of NleE and NleB on other signaling pathways, we used an AP-1 reporter to monitor JNK/MAPK signaling. Neither NleE nor NleB inhibited AP-1 activation by phorbol 12-myristate 13-acetate (PMA) (Fig. 7D), and NleB also had no effect.
NleE/OspZ/NleB Inhibits p65 Nuclear Translocation

A

NF-κB p65
actin
DAPI
Merge

E2348/69

ΔescN

ΔPP4/E6
(pNleE)

ΔPP4/E6
(pNleB)

ΔPP4/E6
(pNleE)

TNFα

IL-1β

B

% p65 nuclear exclusion in FAS-positive cells

Unstimulated
Stimulated with TNFα

EPEC E2348/69
ΔPP4/E6
ΔPP4/E6 (pNleE)
ΔescN
Uninfected

% p65 nuclear exclusion in FAS-positive cells

Unstimulated
Stimulated with IL-1β

EPEC E2348/69
ΔPP4/E6
ΔPP4/E6 (pNleB)
ΔPP4/E6 (pNleE)

* p < 0.05
** p < 0.01
on STAT1/2 activation (data not shown). This suggests that the effectors target only a subset of signaling pathways involving NF-kB.

To ensure that NleE and NleB translocated by the LEE-encoded T3SS conferred the same phenotype as ectopic expression of the effectors by transfection, we infected HeLa cells with wild type EPEC E2348/69 and a double island mutant that lacked the genomic regions, PP4 and IE6 [19]. The double island mutant was used to eliminate genes encoding NleE and NleB in IE6 as well as NleB2, a close homologue of NleB, encoded in PP4 [19]. The ΔPP4/IE6 island mutant was complemented with pNleE or pNleB to examine the contribution of each effector to the inhibition of p65 translocation. In unstimulated cells, there was no difference in p65 nuclear translocation between uninfected cells and those infected with wild type EPEC E2348/69 or the T3SS mutants (escV and escF) (Fig. 8A and B and Fig. 1B). This suggested that, over a 4 h infection, bacterial products such as flagellin and lipopolysaccharide were not sufficient to stimulate signaling. In contrast, the ΔPP4/IE6 island mutant induced substantial p65 nuclear translocation (Fig. 8B), which may indicate that the translocation and/or biochemical function of some effectors is proinflammatory. In infected cells, both NleB and NleE injected by the T3SS had the capacity to inhibit p65 nuclear translocation in response to TNFα but only NleE was effective in response to IL-1β (Fig. 8A and B).

**Effect of NleE, NleB and OspZ on IkB degradation**

Since EPEC infection has been reported to inhibit IkB degradation [12], a critical event in the activation of NF-kB, the effect of NleE and NleB on IkB degradation was examined here in TNFα and IL-1β stimulated cells. Ectopically expressed GFP-NleE and GFP-OspZ inhibited IkB degradation in response to both stimuli whereas GFP-NleB inhibited IkB degradation in response to TNFα only (Fig. 9A). GFP-NleE ΔesK, lacking the ability to inhibit IkB degradation as did GFP-NleEΔAIDSTMK and GFP-OspZΔAIDSTMK (Fig. 9B). In addition, we tested whether NleB and NleE delivered by the T3SS had the same effect as ectopically expressed protein on IkB degradation stimulated by TNFα and IL-1β. Wild type EPEC E2348/69, ΔnleE (pNleE), ΔAPP4/IE6 (pNleE) or ΔAPP4/IE6 (pNleE) inhibited IkB degradation in HeLa cells stimulated with TNFα but IkB degradation was not inhibited in cells infected with ΔAPP4/IE6 (pNleB) and stimulated with IL-1β. This suggests that NleE and NleB act in different ways to interfere with NF-kB signaling (Fig. 9C).

**Discussion**

The activation of NF-kB signaling is a critical host response to infection. In this study, we found that the T3SS effector NleE from EPEC prevented nuclear translocation of the p65 NF-kB subunit, leading to diminished IκB expression and a compromised IκB response. The inhibition of p65 nuclear translocation occurred when NleE was expressed ectopically or when NleE was delivered through the T3SS by infection. We also observed that NleE inhibited nuclear translocation of c-Rel but not nuclear import of activated p50, STAT1 or STAT2. Both p65 and c-Rel are structurally similar and contain transcriptional activation domains that initiate gene expression [6]. In contrast, p50 lacks a transcriptional activation domain, so that p50/p50 homodimers act as transcriptional repressors. Thus, NleE appears to obstruct nuclear translocation of Rel family transcriptional activators while allowing nuclear import of a transcriptional repressor, resulting in the suppression of IκB expression. The selectivity of NleE for p65 and c-Rel is not unprecedented as lack of nuclear translocation of p65 and c-Rel but not p50 was recently reported for oestrogen-induced inhibition of NF-kB activation, although the mechanism is unknown [20].

NleE is one of the conserved core type III effectors of A/E pathogens [19]. We observed that ectopically expressed NleE from EHEC O157:H7 and the murine pathogen, *C. rodentium* also inhibited NF-kB activation and p65 translocation. A close homologue of NleE, OspZ, is found in *Shigella*, however in *S. flexneri* 2a, OspZ is truncated to the length of NleE1,108 [10]. Both the truncated form of OspZ from *S. flexneri* 2a and a C-terminal 36 amino acid deletion mutant of NleE were inactive, suggesting that the C-terminus was critical for the immunosuppressive function of NleE. However, this region was not sufficient for inhibition of p65 nuclear translocation as a region encompassing the last 36 amino acids of NleE alone was unable to prevent NF-kB activation. A domain swap between NleE and OspZ of the last ~40 amino acids showed that these regions were functionally interchangeable and we identified a 6-amino acid motif, IDSYMI/K, that was critical for both NleE and OspZ function.

Although A/E pathogens stimulate an inflammatory response *in vivo* and proteins such as flagellin are recognised by TLR5 [21,22], previous work has suggested that A/E pathogens modulate that inflammatory response by inhibiting p65 nuclear translocation as well as IkB degradation [11,12,23]. Here, we found that NleE inhibited nuclear translocation of p65 by preventing IkB degradation in response to TNFα and IL-1β. In contrast, we found that NleB inhibited IkB degradation in response to TNFα only. Since TNFα and IL-1β signaling converges at the point of IKK phosphorylation (Fig. 9D) [24], NleE may act on IKK or IkB itself to prevent IkB degradation. TAK1 or other MAPK may also have involvement in IKK phosphorylation leading to JNK activation [24], however, JNK signaling, represented here by the AP-1 reporter, was not affected by NleE and so we predict that NleE interferes with IKK or IkB function directly. Indeed while this work was under review, Nadler *et al* reported that NleE inhibits IKK phosphorylation [25]. The authors also proposed that NleB assists the inhibition of IkB degradation by NleE [25]. Here we hypothesize that NleB acts upstream of IKK in the TNFα pathway since NleB did not inhibit IkB degradation in response to IL-1β (Fig. 9D). We therefore propose a model where NleE and NleB act at different points in the NF-kB signaling pathway and each plays a distinct role in the inhibition of p65 nuclear translocation. In the TNFα pathway, NleE and NleB have overlapping and somewhat redundant inhibitory roles as complementation of the ΔAPP4/IE6 double island mutant with either NleE or NleB was sufficient to block p65 nuclear translocation. In the IL-1β pathway however, NleB was not able to compensate for the lack of NleE. Although we believe that NleB acts independently of NleE, these
results do not exclude the possibility that in IL-1β stimulated cells, NleB acts in concert with NleE [25].

The fact that both NleE and NleB inhibit NF-κB activation raises the possibility that more effectors contribute to the suppression of innate signaling pathways. Although compromised compared to wild type EPEC, the nleE mutant showed significantly greater inhibition of IL-8 secretion than an T3SS mutant, which lacks the ability to translocate all T3SS effectors. While one of the additional effectors inhibiting p65 translocation is clearly NleB, a close homologue, NleB2 may also have anti-inflammatory activity and perhaps other effectors in the genomic islands, PP4 and IE6. In addition, NleH1 and NleH2 were recently reported to interfere with the activation of NF-κB by binding ribosomal protein S3 (RPS3), a co-factor of nuclear NF-κB complexes, and sequestering it in the cytoplasm [17]. We also found that ectopically expressed NleH1 and NleH2 inhibited NF-κB activation, but not to the same degree as NleE and NleB. Together these anti-inflammatory effectors may balance the action of other effectors that through their biochemical activity stimulate inflammatory signaling, as suggested by the ΔΔP4/IE6 double island mutant, which showed increased p65 nuclear translocation in uninfected cells compared to a T3SS mutant. Therefore, despite the fact that EPEC and Shigella infection ultimately induces gut inflammation, we propose that NleE/OspZ and NleB contribute to pathogenesis by inhibiting an initial host inflammatory response to allow the bacteria to persist in the early stages of infection.

A multi-effector attack on NF-κB signaling occurs during Shigella infection, which modulates NF-κB activation through the effectors OspG and OspF [3,26]. Our studies suggest that Shigella strains carrying full length OspG have evolved a different mechanism to modulate NF-κB signaling. This makes the absence of a functional OspG protein in S. flexneri 2a curious and may also explain previous findings that OspZ from Shigella and Salmonella [18] were serotype specific and perhaps other effectors in the genomic islands, PP4 and IE6. Infection of HeLa cells with Shigella is characterized by the absence of a clear NF-κB activation, which is in contrast to the T3SS effectors in Salmonella [10]. To further elucidate this phenomenon, we have characterized the NF-κB signaling machinery during Shigella infection and determined how T3SS effectors influence this process.

In this study, we have ascribed a function to the NleE/OspZ family of T3SS effectors shared by attaching and effacing pathogens and Shigella as well as the EPEC effecter, NleB. Despite the remarkably different infection strategies of these two groups of pathogens, they appear to have a mutual need to inhibit the host inflammatory response during infection. NleE, NleB and OspZ are the latest T3SS effectors to target NF-κB activation and the expression of NF-κB-dependent genes. Neither NleE nor NleB inhibited STAT1/2 or AP-1 signaling, suggesting that the proteins target the NF-κB pathway specifically. The ongoing identification of T3SS effectors that act on this and other inflammatory pathways will continue to provide insight into the molecular mechanisms by which bacterial pathogens inhibit immune signaling and establish infection.

Methods

HeLa and Caco-2 infections, IL-8 secretion and expression

The bacterial strains and plasmids used in this study are listed in Table 1. The construction of vectors and culturing of bacterial strains for infection is described in detail in the supplementary methods (Protocol S1). EPEC strains were used to infect HeLa cells for 4 h without exogenous stimulation or for 90 min after which the media was replaced with DMEM supplemented with 10 ng/ml TNFα or 10 ng/ml IL-1β (eBioscience, San Diego, CA) and the infection was continued for a further 30 min. For Caco-2 cells, EPEC infection continued for 4 h after which the cells were washed, treated with 100 μg/ml gentamicin for 2 h. For mRNA analysis monolayers were incubated for 3 h in media supplemented with 50 μg/ml gentamicin with or without 5 ng/ml IL-1β. For analysis of IL-8 secretion, monolayers were infected for 4 h and incubated for 24 h in media supplemented with 50 μg/ml gentamicin with or without 5 ng/ml IL-1β. IL-8 secretion into cell culture supernatants was measured by ELISA (Peprotech EC). The expression of IL-8 from total RNA was determined using the comparative quantification method included in Rotor-Gene 1.7 software (Qiagen) as described in the supplementary methods using gene specific primers (Protocol S1).

Immunofluorescence, fluorescence activation staining test and confocal microscopy

Plasmids were transfected into HeLa cells for ectopic expression of GFP fusion proteins using Lipofectamine 2000 in accordance with the manufacturer’s recommendation (Invitrogen, Carlsbad CA, USA). Transfected HeLa cells were treated with 20 ng/ml TNFα, 10 ng/ml IL-1β or IFNα (500 U/ml; Calbiochem, La Jolla, CA, USA) for 30 min at 37°C and 5% CO2. Transfected or infected cells were fixed in 3.7% (wt/vol) formaldehyde (Sigma) in PBS for 10 min and permeabilized with acetic acid-methanol (1:1, vol/vol) at −20°C for 15 min. Following a 30 min blocking in PBS with 3% (wt/vol) bovine serum albumin (Amresco, Ohio, USA) samples were exposed to rabbit polyclonal anti-p65 (SC-109, Santa Cruz, Santa Cruz CA, USA), anti-c-Rel (#4727, Cell Signaling, Beverly MA, USA), anti-STAT1 (SC-345, Santa Cruz), anti-STAT2 (SC-476, Santa Cruz) or mouse monoclonal anti-p30 (2E6, Novus Biologicals, Littleton CO, USA). Antibodies were used at a 1:100, or 1:30 for anti-c-Rel, (vol/vol) diluted in blocking solution for 1 h at 20°C. Alexa Fluor 488 or Alexa Fluor 568 (Invitrogen) conjugated anti-mouse or anti-rabbit immunoglobulin G were used at 1:2000. Coverslips were mounted onto microscope slides with Prolong Gold containing 4,6-diamidino-2-phenylindole (DAPI, Invitrogen). For the fluorescence activation staining (FAS) test, HeLa and Caco-2 cells were infected with bacterial strains, fixed and permeabilized as described above and cells were incubated with 0.5 ng/ml phalloidin conjugated to rhodamine for 30 min. Images were acquired using a confocal laser scanning microscope (Leica LCS SP2 confocal imaging system) with a
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**Table 1.** Bacterial strains and plasmids used in this study.

| Strains                  | Characteristics                          | Source/reference |
|--------------------------|------------------------------------------|-----------------|
| EPEC E2348/69            | Wild type EPEC O127:H6                   | [27]            |
| ΔnleE                   | EPEC E2348/69 ΔnleE Cm<sup>R</sup>       | [10]            |
| ΔPP4/E6                 | EPEC E2348/69 PP4/E6 double island deletion | O. Marchés   |
| EHEC EDL933             | Wild type EHEC O157:H7                   | [28]            |
| C. rodentium ICC169     | Spontaneous nalidixic acid resistant derivative of wild-type C. rodentium biotype 4280 (Na<sup>R</sup>) | [29] |
| S. flexneri 2104        | Wild type S. flexneri 2a                 | Roy Robins-Browne |
| S. flexneri 0106164     | Wild type S. flexneri serotype 6         | Roy Robins-Browne |
| S. boydii SBA1384       | Wild type S. boydii serotype 4           | Ben Adler       |

**Plasmids**

| pTrc99A | Cloning vector for expression of proteins from Ptrec | Pharmacia Biotech |
| pNleE | nleE from EPEC E2348/69 in pTrc99A | This study |
| pNleE<sub>6A</sub> | nleE from EPEC E2348/69 in pTrc99A carrying alanine substitutions for each amino acid in the IDSYMK motif | This study |
| pNleB | nleB from EPEC E2348/69 in pTrc99A | This study |
| pEGFP-C2 | Green fluorescent protein (GFP) expression vector | Clontech |
| pGFP-NleE | Full length nleE from EPEC E2348/69 in pEGFP-C2 | [10] |
| pGFP-NleE<sub>157</sub> | Full length nleE from EHEC EDL933 in pEGFP-C2 | This study |
| pGFP-OspZ<sub>20</sub> | Full length ospZ from S. flexneri 2104 in pEGFP-C2 | This study |
| pGFP-OspZ<sub>6A</sub> | Full length ospZ from S. flexneri 0106164 in pEGFP-C2 | This study |
| pGFP-OspZ<sub>6B</sub> | Full length ospZ from S. boydii SBA1384 in pEGFP-C2 | This study |
| pGFP-NleH1 | Full length nleH1 from EPEC E2348/69 in pEGFP-C2 | This study |
| pGFP-NleH2 | Full length nleH2 from EPEC E2348/69 in pEGFP-C2 | This study |
| pGFP-NleD | Full length nleD from EPEC E2348/69 in pEGFP-C2 | This study |
| pGFP-NleB | Full length nleB from EPEC E2348/69 in pEGFP-C2 | This study |
| pGFP-NleE<sub>188</sub> | Truncation of nleE from EPEC E2348/69 encoding amino acids 1-188 in pEGFP-C2 | This study |
| pGFP-NleE<sub>200</sub> | Truncation of nleE from EPEC E2348/69 encoding amino acids 1-200 in pEGFP-C2 | This study |
| pGFP-NleE<sub>208</sub> | Truncation of nleE from EPEC E2348/69 encoding amino acids 1-208 in pEGFP-C2 | This study |
| pGFP-NleE<sub>214</sub> | Truncation of nleE from EPEC E2348/69 encoding amino acids 1-214 in pEGFP-C2 | This study |
| pGFP-NleE<sub>IDSYMK</sub> | Full length nleE from EPEC E2348/69 carrying the IDSYMK deletion in pEGFP-C2 | This study |
| pGFP-NleE<sub>Id</sub> | Full length nleE from EPEC E2348/69 in pEGFP-C2 carrying alanine substitutions for each amino acid in the IDSYMK motif | This study |
| pGFP-NleE<sub>209A</sub> | Full length nleE from EPEC E2348/69 carrying the mutation I209A in pEGFP-C2 | This study |
| pGFP-NleE<sub>210A</sub> | Full length nleE from EPEC E2348/69 carrying the mutation D210A in pEGFP-C2 | This study |
| pGFP-NleE<sub>211A</sub> | Full length nleE from EPEC E2348/69 carrying the mutation S211A in pEGFP-C2 | This study |
| pGFP-NleE<sub>212A</sub> | Full length nleE from EPEC E2348/69 carrying the mutation Y212A in pEGFP-C2 | This study |
| pGFP-NleE<sub>213A</sub> | Full length nleE from EPEC E2348/69 carrying the mutation M213A in pEGFP-C2 | This study |
| pGFP-NleE<sub>214A</sub> | Full length nleE from EPEC E2348/69 carrying the mutation K214A in pEGFP-C2 | This study |
| pGFP-OspZ<sub>IDSYIK</sub> | Full length OspZ from S. flexneri carrying the IDSYIK deletion in pEGFP-C2 | This study |
| pGFP-NleE-OspZ<sub>term</sub> | NleE<sub>1-183</sub> fused to the last 46 amino acids of OspZ | This study |
| pGFP-OspZ-NleE<sub>term</sub> | OspZ<sub>1-183</sub> fused to the last 40 amino acids of NleE | This study |
| pRL-TK | Renilla luciferase vector | Promega |
| pNF-xB-Luc | Vector for measuring NF-xB dependent luciferase expression | Clontech |
| p(9-27)4thLucter | Vector for measuring STAT1/2 dependent luciferase expression | [15] |
| pAP-1-Luc | Vector for measuring AP-1 dependent luciferase expression | Clontech |

100x/1.4 NA HCX PL APO CS oil immersion objective. Nuclear exclusion of NF-κB, STAT1 and STAT2 was quantified from at least 3 independent experiments for both transfection and infection studies.

NF-κB, STAT and AP-1 reporter assay

To examine the activity of NF-κB, a dual luciferase reporter system was employed. HeLa cells were seeded into 24-well trays and co-transfected with derivatives of pEGFP-C2 (1.0 μg) together with...
with 0.2 μg of pNF-xB-Luc (Clontech, Palo Alto CA, USA) and 0.04 μg of pRL-TK (Promega, Madison WI, USA). Approximately 24 h after transfection, cells were left untreated or stimulated with 20 ng/ml TNF-α or 10 ng/ml IL-1β for 16 h. Firstly and Renilla luciferase levels were measured using the Dual-luciferase reporter assay system (Promega) in the Topcount NXT instrument. For each sample, the expression of firefly luciferase was normalized for Renilla luciferase measurements and NF-κB activity was expressed relative to unstimulated pEGFP-C2 transfected cells.

To measure the induction of STAT1 and 2, the IFN-α/β-responsive luciferase reporter plasmid p9(-27)/4(-39)LucIFR (ISRE-Luc) [15] was used in combination with the Renilla luciferase plasmid pRL-TK. HeLa cells were transfected with both plasmids as described above and stimulated with IFNα (500 U/ml; Calbiochem) for 30 min. Luciferase activity was measured as described above. To measure the induction of AP-1, the cAMP response element (CRE)-dependent luciferase vector pAP-1-Luc was used in combination with the Renilla luciferase plasmid pRL-TK. HeLa cells were transfected with both plasmids as described above and stimulated with 25 ng/ml phorbol 12-myristate 13-acetate (PMA) for 30 min. Luciferase activity was measured as described above.

Detection of IkB degradation by immunoblot
To test the effect of ectopically expressed NleE, NleB and OspZ and on IkB degradation, HeLa cells were mock transfected or transfected with pEGFP-C2 or pEGFP-NleE, pEGFP-NleB, pEGFP-OspZ and derivatives and incubated for 16 h before being left untreated or treated with TNF-α or IL-1β for 10, 20 or 30 min. Cell lysis was performed by incubating cells in cold lysis buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 5 mM EDTA, 1% NP-40) on ice for 5 min before collecting lysate and incubating on ice for a further 10 min. Cell debris was pelleted and equal volumes of supernatant were collected for SDS-PAGE. Proteins transferred to nitrocellulose membranes were probed with mouse monoclonal anti-IκB α (Cell Signaling) diluting 1:1000 or rabbit polyclonal anti-p65 (Santa Cruz) diluted 1:1000. For infection studies, HeLa cells were infected with derivatives of EPEC E2348/69 for 90 min before stimulation with TNF-α or IL-1β for 30 min as described above.

Supporting Information
Figure S1 CaCo-2 cells infected with derivatives of EPEC E2348/69, stimulated with IL-1β where indicated and stained for actin (green), nucleic acid (blue) and p65 (red). Found at: doi:10.1371/journal.ppat.1000898.s001 (2.10 MB TIF)

Figure S2 Alignment of NleE and OspZ from A/E pathogens and Shigella species showing amino acid conservation. Found at: doi:10.1371/journal.ppat.1000898.s002 (0.92 MB TIF)

Figure S3 Ectopic expression of GFP-NleE and GFP-OspZ derivatives in HeLa cells. Found at: doi:10.1371/journal.ppat.1000898.s003 (0.73 MB TIF)

Protocol S1 Supplementary methods. Found at: doi:10.1371/journal.ppat.1000898.s004 (0.08 MB DOC)

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Author Contributions
Conceived and designed the experiments: HJN JS LP MK ML GH KMW DAJ. Wrote the paper: GF ELH.

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