CELLULAR MICROBIOLOGY: PATHOGEN-HOST CELL MOLECULAR INTERACTIONS



The Type III Effector NIeD from Enteropathogenic *Escherichia coli* Differentiates between Host Substrates p38 and JNK

Infection and

MICROBIOLOGY

AMERICAN SOCIETY FOR

Kristina Creuzburg,^a Cristina Giogha,^a Tania Wong Fok Lung,^a Nichollas E. Scott,^a Sabrina Mühlen,^{a,b} Elizabeth L. Hartland,^a Jaclyn S. Pearson^a

Department of Microbiology and Immunology, University of Melbourne at the Peter Doherty Institute for Infection and Immunity, Melbourne, Australia^a; Department of Molecular Infection Biology, Helmholtz Centre for Infection Research, Braunschweig, Germany^b

ABSTRACT Enteropathogenic Escherichia coli (EPEC) is a gastrointestinal pathogen that utilizes a type III secretion system (T3SS) to inject an array of virulence effector proteins into host enterocytes to subvert numerous cellular processes for successful colonization and dissemination. The T3SS effector NIeD is a 26-kDa zinc metalloprotease that is translocated into host enterocytes, where it directly cleaves and inactivates the mitogen-activated protein kinase signaling proteins JNK and p38. Here a library of 91 random transposon-based, in-frame, linker insertion mutants of NleD were tested for their ability to cleave JNK and p38 during transient transfection of cultured epithelial cells. Immunoblot analysis of p38 and JNK cleavage showed that 7 mutant derivatives of NIeD no longer cleaved p38 but maintained the ability to cleave JNK. Site-directed mutation of specific regions surrounding the insertion sites within NIeD revealed that a single amino acid, R203, was essential for cleavage of p38 but not JNK in a direct in vitro cleavage assay, in transiently transfected cells, or in EPEC-infected cells. Mass spectrometry analysis narrowed the cleavage region to within residues 187 and 213 of p38. Mutation of residue R203 within NIeD to a glutamate residue abolished the cleavage of p38 and impaired the ability of NIeD to inhibit AP-1-dependent gene transcription of a luciferase reporter. Furthermore, the R203 mutation abrogated the ability of NIeD to dampen interleukin-6 production in EPEC-infected cells. Overall, this work provides greater insight into substrate recognition and specificity by the type III effector NIeD.

KEYWORDS NIeD, metalloprotease, JNK, p38, EPEC, type III effector proteins, mitogen-activated protein kinases

E nteropathogenic *Escherichia coli* (EPEC) is an extracellular gastrointestinal pathogen that causes significant diarrheal disease in infants, particularly in developing countries. EPEC adheres intimately to the mucosal surface of enterocytes and injects virulence proteins (effectors) directly into host cells via a type III secretion system (T3SS). The genes required to form a functional T3SS, along with genes encoding a subset of effector proteins, are located on a 35-kb genomic pathogenicity island (PAI) called the locus of enterocyte effacement (LEE). Other effector proteins translocated by the EPEC T3SS are located throughout the genome on distinct genomic PAIs and are termed non-LEE-encoded (NIe) effectors. Many NIe effectors from EPEC display novel enzymatic activity and mediate the subversion of host immune signaling processes, such as NF- κ B and mitogen-activated protein kinase (MAPK) activation, apoptosis, and necroptosis. For example, NIeE is an *S*-adenosyl-L-methionine-dependent transferase that specifically modifies the NF- κ B adaptor proteins TAB2/3, eliminating ubiquitin chain binding activity and thus inhibiting downReceived 18 July 2016 Returned for modification 16 August 2016 Accepted 17 November 2016

Accepted manuscript posted online 21 November 2016

Citation Creuzburg K, Giogha C, Wong Fok Lung T, Scott NE, Mühlen S, Hartland EL, Pearson JS. 2017. The type III effector NIeD from enteropathogenic *Escherichia coli* differentiates between host substrates p38 and JNK. Infect Immun 85:e00620-16. https:// doi.org/10.1128/AL.00620-16.

Editor Andreas J. Bäumler, University of California, Davis

Copyright © 2017 American Society for Microbiology. All Rights Reserved. Address correspondence to Jaclyn S. Pearson, jaclynp@unimelb.edu.au. stream NF- κ B activation and interleukin-8 (IL-8) production during EPEC infection (1–3). NIeB is an *N*-acetylglucosamine transferase that inhibits extrinsic apoptosis by adding a single GlcNAc moiety to a conserved arginine residue on the Fas-associated death domain (FADD) protein, as well as other host death domain proteins, including TRADD and RIPK1. This activity predominantly blocks death receptor signaling and subsequent inflammation and apoptosis during infection, thus promoting cell survival and pathogen persistence (4, 5).

EPEC also produces two zinc metalloprotease T3SS effectors, NIeC and NIeD, which were first identified to be effectors secreted by the LEE-encoded T3SS of the EPEC-like mouse pathogen *Citrobacter rodentium* (6). Zinc-containing metalloproteases are a group of proteolytic enzymes found across eukaryotes and prokaryotes and are important in maintaining physiological homeostasis within the organism (7). Collectively known as the "zincins," they contain a consensus motif, HEXXH, where the histidine residues bind a zinc ion and promote nucleophilic attack on peptide bonds using a water molecule at the active site (8). Recent molecular studies have shown that NIeC and NIeD require the HEXXH motif for cleavage of host innate immune-signaling proteins. NIeC specifically cleaves and inactivates NF- κ B Rel proteins, including p65 and p50, and thus contributes to the inhibition of IL-8 production during EPEC infection (9–15). NIeD cleaves and inactivates the mitogen-activated protein (MAP) kinases c-Jun amino-terminal kinase (JNK) and p38 but not extracellular signal-related kinases (ERK) and contributes modestly to the inhibition of IL-8 production and UV-induced apoptosis in EPEC infection (15).

The MAP kinases are members of discrete signaling pathways that mediate responses to multiple extracellular stimuli, including UV light, heat, osmotic shock, inflammatory cytokines (tumor necrosis factor [TNF] and interleukin-1 [IL-1]), and growth factors (colony-stimulating factor 1 [CSF1]) (16). Three distinct members of the MAPK family exist in mammals: ERK, JNK, and p38/stress-activated protein kinases (SAPK). Activation of JNK and/or p38 contributes to a number of important cellular processes, including inflammation, apoptosis, cell differentiation, and cytokine production (17).

Apart from the metalloprotease motif, little is known about other functional regions of NIeD. In this study, we screened a library of transposon and site-directed mutants of NIeD to identify amino acids critical for its proteolytic activity against p38 and JNK. We found that aside from amino acids within the catalytic site (H¹⁴²ELLH¹⁴⁶), R203 was essential for direct p38 cleavage and inhibition of AP-1 activation and IL-6 production by NIeD but dispensable for JNK cleavage. In addition, we demonstrated direct cleavage of p38 by NIeD and propose that cleavage is likely to occur between residues 187 and 213 of p38.

RESULTS

Effect of pentapeptide insertions on proteolytic activity of NleD. To map the contribution of amino acid residues of NIeD to substrate binding and proteolytic cleavage, an extensive pentapeptide scanning mutagenesis screen was performed. A library of 80 mutants was generated using a mutation generation system kit (catalog number F-701; Thermo Scientific) and screened for the insertion of 5 additional amino acids at random positions in NIeD originating from EPEC O127:H6 strain E2348/69. An additional 11 pentapeptide mutants were generated by site-directed mutagenesis (SDM) to fill the gaps where there were more than 5 amino acids between mutants obtained with the transposon-mediated mutagenesis system (Fig. 1; see also Table S1 in the supplemental material). In total, 91 mutants of pGFP-NleD with pentapeptide insertions at 78 different positions were made. The resultant pGFP-NleD derivatives were transfected into HEK293T cells, and the cell lysates were analyzed by immunoblotting for cleavage of JNK and p38 (Fig. 1 and Table S1). pGFP-NleD was selected as the template vector for the mutagenesis screen, as expression can easily be visualized by immunoblotting using anti-green fluorescent protein (anti-GFP) antibodies or directly by immunofluorescence microscopy, if required.



FIG 1 Selected NIeD transposon mutants that display altered proteolytic activity toward JNK and/or p38. (A) Immunoblots of JNK and p38 cleavage in cells transiently transfected with GFP-NIeD or GFP-NIeD mutant derivatives. Cell lysates were probed with anti-JNK and anti-p38 antibodies. Arrows, cleavage products. Anti-GFP antibody was used to show the expression of GFP-NIeD and GFP-NIeD variants. Actin was used as a loading control. The immunoblots are representative of those from at least 3 independent experiments. UT, untransfected. (B) Schematic diagram mapping transposon insertion sites in NIeD that alter JNK and/or p38 cleavage. Dark gray arrowheads, little or no cleavage of JNK or p38; light gray arrowheads, cleavage of JNK but little or no cleavage of p38; white arrowheads, total cleavage of both JNK and p38. aa, amino acids.

NleD from EPEC E2348/69 is 232 amino acids in length, and the HEXXH motif spans amino acids 142 to 146. Mutants harboring insertions of pentapeptides within the 59 amino acids of the N-terminal region of NleD showed proteolytic activity comparable to that of wild-type NleD (Table S1), whereas insertions at 22 different positions downstream of amino acid 59 altered the proteolytic activity of NleD to various extents (Fig. 1A and B and Table S1). Insertion of pentapeptides starting at amino acid positions 73, 196, 202, 203, and 217 in NleD abolished the cleavage of p38 and reduced the level of JNK cleavage compared to that by wild-type NleD (Fig. 1A and Table S1). Insertions starting at amino acid positions 60, 76, 100, 110, 154, 222, and 226 decreased the level of cleavage of p38 to different degrees, whereas cleavage of JNK by these mutants was comparable to that by wild-type NleD (Fig. 1A and Table S1). Insertions at amino acid positions 115 and 179 completely abolished the cleavage of p38 and JNK, as did those within or immediately preceding the H¹⁴²ELLH¹⁴⁶ catalytic motif at positions 144, 146, 147, and 150 (Fig. 1A and Table S1).

Site-directed mutagenesis of NIeD and screening of mutants for loss of proteolytic activity. To further examine potential functional regions of NIeD, 16 sitedirected mutants with mutations consisting of either two or three consecutive alanine substitutions were constructed using a QuikChange II site-directed mutagenesis kit. Mutation positions were selected on the basis of the pentapeptide insertion positions that yielded a significant effect on p38 or JNK cleavage. The site-directed mutants were subsequently tested for their ability to cleave p38 and JNK (Fig. 2A and B and Table S2).



FIG 2 Screen of NIeD multiple site-directed mutants for loss of proteolytic activity against JNK and/or p38. (A) Immunoblots of JNK and p38 cleavage in HEK293T cells transiently transfected with GFP-NIeD or GFP-NIeD multiple site-directed mutants. Cell lysates were probed with anti-JNK and anti-p38 antibodies. Anti-GFP antibody was used to show the expression of GFP-NIeD and GFP-NIeD variants. Arrows, cleavage products. Actin was used as a loading control. The immunoblots are representative of those from at least 3 independent experiments. UT, untransfected. (B) Schematic diagram mapping triple alanine replacement sites in NIeD that alter JNK and/or p38 cleavage. Light gray arrowheads, cleavage of JNK but little or no cleavage of p38; white arrowheads, total cleavage of both JNK and p38. aa, amino acids.

HEK293T cells were transfected with derivatives of pGFP-NleD, and lysates were subjected to SDS-PAGE followed by immunoblot analysis using anti-p38 or anti-JNK antibodies (Fig. 2A). NleD harboring amino acid substitutions across positions 109 to 111, 114 to 116, 146 to 148, 149 to 151, 153 to 155, 178 to 180, 202 to 204, 216 to 218, and 227 to 229 cleaved JNK as efficiently as wild-type NleD. However, substitutions across 109 to 111, 114 to 116, 146 to 148, and 216 to 218 resulted in less efficient cleavage of p38 compared to that by wild-type NleD, and substitutions across positions 149 to 151, 153 to 155, 178 to 180, and 202 to 204 rendered NleD unable to cleave p38 (Fig. 2A and B and Table S2).

Arginine 203 in NIeD is required for p38 cleavage but dispensable for JNK cleavage. Single amino acid changes were made across most of the 3 amino acid substitutions within NIeD that resulted in the loss in p38 cleavage. This included the replacement of glutamate 143 within the H¹⁴²EXXH¹⁴⁶ catalytic motif by alanine (E143A) as a control for the proteolytic activity of NIeD. Of these mutants, pGFP-NIeD with the R203A substitution (pGFP-NIeD_{R203A}) completely lost the ability to cleave p38 but retained the ability to cleave JNK (Fig. 3 and Table S2).



FIG 3 Analysis of NIeD single site-directed mutants for p38 cleavage. Immunoblots of JNK and p38 cleavage in HEK293T cells transiently transfected with GFP-NIeD or GFP-NIeD single site-directed mutants are shown. Cell lysates were probed with anti-JNK and anti-p38 antibodies. Anti-GFP antibody was used to show the expression of GFP-NIeD and GFP-NIeD variants. Cleavage products are indicated by arrows. Actin was used as a loading control. The immunoblots are representative of those from at least 3 independent experiments. UT, untransfected.

Arginine 203 is conserved among metalloprotease effectors from pathogenic bacteria. In order to determine whether R203 was conserved in homologues of NIeD from other pathogenic bacteria, a BLAST analysis was performed and an alignment of proteins was performed using the ESPript (v3) program. R203 was indeed conserved across a number of diverse bacterial pathogens (Fig. 4). To ensure that the phenotype of the NIeD mutant with the R203A substitution (the NIeD_{R203A} mutant) was not only due to the selection of alanine as the substitute amino acid, we replaced R203 with lysine or glutamate, each of which carries a charge similar to or opposite that of arginine, respectively. pGFP-NIeD_{R203A}, pGFP-NIeD_{R203E}, pGFP-NIeD_{R203K}, and GFP-NIeD_{E143A} were transfected into HEK293T cells and the cell lysates were analyzed by immunoblotting using anti-p38 and anti-JNK antibodies. Cleavage of p38 was abrogated upon expression of GFP-NIeD_{R203A}, GFP-NIeD_{R203E}, or GFP-NIeD_{E143A} compared to that obtained upon expression of GFP-NIeD, whereas expression of GFP-NIeD_{R203K} only partially compromised p38 cleavage (Fig. 5A).

Previous studies have suggested that NleD inhibits AP-1 activation (15); therefore, we tested NleD derivatives carrying amino acid substitutions in R203 and the adjacent amino acids, R202 and T204, for their ability to activate AP-1-dependent gene transcription of a luciferase reporter. Upon stimulation with phorbol myristate acetate (PMA) (18), AP-1 activation was evident in cells transfected with the empty vector pEGFP-C2, whereas it was not in unstimulated cells (Fig. 5B). AP-1 activation was

EPEC_E2348/69_NleD Sorangium_cellulosum Morganella_morganii Chelativorans_spJ32 Endozoicomonas_numazuensis Acidovorax_avenae_Acav_3399 Xanthomonas_vasicola Xanthomonas_fuscans_XopG Vibrio_shilonii Pseudomonas_syringae_Hop1 Janthinobacterium_agaricidamnosum Klebsiella_michiganensis Beauveria_bassiana	VVDFHATIVFHELLHVFHNLNGERLKVESSRPESQKYSPLLLEEARTVGLGAFSE IAEPSQVLLGHEMAHALANAEGRRQTGTDPA.PPASQPDIGREEAQAIGTGSHNG EKQFHASVLYHELVHVLHYLQGECIHIIPADSPESIRYPYWEEEARTIGFGPFTS PGISDVLKLFHELVHVVHYLQGECIHIIPADSPESIRYPYWEEEARTVGLGMFRD GLRPAYIGLAHELHHYYYNMRGAVSHSEDSAHFSTVLYEYMCVGLGPWNG NPANAFITLAHELVHAKHHLAGTMMYGGGPVTREASSSRTDAGREELRAVGLGEYAST NPEESFVVLAHELIHAQHLLAGTSKACKGGDRYDDTSEAGKEELRAVGIGKYEYRKT DPESFVVLAHELIHAQHLLAGTSRAYKGGDRYDDTSEAGKEELRAVGIGKYEYRKT DPSLAYVFLGHELHASRTLQGTKM.DGKDELWTTGLGKYSN DFSLAYVFLGHELHASRTLQGTQY.RVSRGPDVDEDSGAEEELRAVGIGVGKYEYRKT DSLAYVFLGHELHASRTLQGTYGGTSDRYDPSTPAAEEEDRAVGIGKYSK DTSLAFVSLAHELVHGYRMMKGTYT.GGTSDRYDTGSPAGQEESRAVGIGKYAG
EPEC_E2348/69_NleD Sorangium_cellulosum Morganella_morganii Chelativorans_spJ32 Endozoicomonas_numazuensis Acidovorax_avenae_Acav_3399 Xanthomonas_vasicola Xanthomonas_fuscans_XopG Vibrio_shilonii Pseudomonas_syringae_Hop1 Janthinobacterium_agaricidamnosum Klebsiella_michiganensis Beauveria_bassiana FIG 4 Alignment of zinc metalloprotease motif a	EVLSENKFREEIGMPRRTSYPHDSALIHDD AYPSENSFRRDLGLPERDNHYWTPDG ETISENTFRAEIGVPLRTH.W EQLCENRLRSEIGVQPRASATSGDWVSYDD AAISENQIRQDAGVPLRTR.YA GEPSENSIRAEHGLPQRTS.YSRSGN RQPSENSIRQEHGLPVRKK.YKPHG NPMSENCFRRQFGLAERTQ.YF EYPSENAIRREHGLAPRTS.YSGNTG EYPSENAIRREHGLPRF.YRAAPD EALSENGIRQEHGLPLRQG.YAAG EALSENGIRQEHGLPLRGQ.YAAG

FIG 4 Alignment of zinc metalloprotease motif and the surrounding area in NIeD proteins from various Gram-negative bacteria. Sequences were identified through BLAST analysis using NIeD from EPEC O127:H6 strain E2348/69 as a reference. A section of the proteins identified by BLAST analysis was aligned using the Clustal Omega program, and presentation of the alignment was performed using the ESPript (v3) program (36). *, HEXXH zinc metalloprotease site; arrow, conserved residue R203.

inhibited in cells transfected with wild-type pGFP-NIeD, pGFP-NIeD_{R202A}, or pGFP-NIeD_{T204A}, whereas ectopic expression of the catalytic mutant GFP-NIeD_{E143A} or GFP-NIeD_{R203E} had no significant impact on AP-1 activation upon PMA stimulation (Fig. 5B). Inhibition of AP-1 activation by GFP-NIeD_{R203K} in response to PMA was similar to that by wild-type NIeD, and we noted that GFP-NIeD_{R203A} still inhibited AP-1 activation in this system, despite being unable to cleave p38 (Fig. 5A and B).

NIeD directly cleaves p38. To test the ability of NIeD to cleave p38 directly, we incubated purified recombinant NIeD with purified p38 and observed cleavage of the substrate by catalytically active NIeD (Fig. 6A and B). Both catalytically inactive NIeD (His-NleD_{E143A}) and His-NleD_{R203E} were unable to cleave p38 directly (Fig. 6A). In order to prove that the R203E mutant was active, we incubated purified His-NIeD_{R203E} with HT-29 cell lysate and observed cleavage of endogenous JNK but not that of p38 (Fig. S1). The in vivo-generated p38 fragments were further assessed by mass spectrometry, which both confirmed their status as p38 cleavage fragments and revealed the putative cleavage site to be between residues 187 and 213 (Fig. 6B and C, S2, and S3). Analysis of the peptide coverage of full-length p38 compared to that of the N- and C-terminal cleavage fragments revealed that peptides spanning the entire protein sequence could be identified, with peptide coverage being restricted to the N- and C-terminal regions within the N- and C-terminal fragments, respectively (Fig. S2). However, there remained 26 residues (187WYRAPEIMLNWMHYNQTVDIWSVGCl²¹²) between the N- and C-terminal cleavage fragments that were inaccessible to analysis due to the composition of the sequence, which lacks sufficient lysine or arginine residues required for tryptic digestion and subsequent mass spectrometry analysis (Fig. S2). The peptide ²¹³MAELLTGR²²⁰ was the most N-terminal peptide identified within the C-terminal cleavage fragment and was not present in the full-length or the N-terminal p38 cleavage fragment tryptic digest, suggesting that M₂₁₃ may be the most likely cleavage site (Fig. S3). Overall, we can safely conclude only that cleavage occurs between residues 187 and 213 of p38.

Arginine 203 in NIeD is essential for p38 cleavage and inhibition of IL-6 secretion during EPEC infection. To examine whether the NIeD R203E variant had the same effect on p38 and JNK cleavage in EPEC-infected cells as in transiently transfected



FIG 5 Analysis of various NleD_{R203} site-directed mutants for p38 cleavage and their effect on AP-1 activation. (A) Immunoblots of JNK and p38 cleavage in HEK293T cells transiently transfected with GFP-NleD or GFP-NleD single site-directed mutants. Cell lysates were probed with anti-JNK and anti-p38 antibodies. Anti-GFP antibody was used to show the expression of GFP-NleD and GFP-NleD variants. Arrows, cleavage products. Actin was used as a loading control. The immunoblots are representative of those from at least 3 independent experiments. UT, untransfected. (B) Fold increase in AP-1-dependent luciferase activity in HEK293T cells expressing GFP or GFP-NleD variants unstimulated or stimulated with PMA for 6 h. Results are the means \pm SEMs from at least three independent experiments carried out in duplicate. *, a result significantly different from the results obtained with GFP-NleD only stimulated with PMA (P < 0.05, by an unpaired, two-tailed t test).

cells, *nleD*, *nleD*_{R203E} and *nleD*_{E143A} were cloned into the bacterial expression vector pTrc99A that carries a C-terminal 2× hemagglutinin (HA) tag to visualize the expression of NleD by immunoblotting. The constructs were expressed in the EPEC E2348/69 Δ PP4/IE6 genetic background. This strain is missing seven effectors (*nleD*, *nleC*, *nleE*, *nleB1*, *nleB2*, *espL*, and *nleG*). Infection of HT-29 cells with wild-type EPEC resulted in the cleavage of p38 and JNK, whereas infection with either the T3SS mutant (Δ *escN*), the Δ PP4/IE6 double island mutant, or the Δ PP4/IE6 mutant complemented with *nleD*_{E143A} had no effect on the cleavage of either substrate (Fig. 7A). Complementation of the Δ PP4/IE6 mutant with wild-type *nleD* resulted in the cleavage of both JNK and p38 in EPEC-infected HT-29 cells, whereas complementation with *nleD*_{R203E} resulted only in the loss of JNK (Fig. 7A).

Studies have shown that activation of p38 MAPK signaling results in the increased secretion of proinflammatory cytokines, including IL-1 β , TNF, and IL-6 (19). Therefore, we examined IL-6 secretion levels in HT-29 cells infected with various EPEC derivatives. Cells infected with wild-type EPEC and stimulated with TNF produced minimal amounts of IL-6, comparable to the findings for uninfected, unstimulated cells (Fig. 6B). This was also the case for cells infected with Δ PP4/IE6 complemented with *nleE*, which is a potent inhibitor of inflammatory cytokine production (2) (Fig. 6B). Expression of wild-type NIeD but not that of NIeD_{E143A} or NIeD_{R203E} in the Δ PP4/IE6 mutant also inhibited IL-6 production, but NIeD was less effective than NIeE (Fig. 7B).

DISCUSSION

Early *in vivo* studies showed no significant virulence phenotype for single *nleC* or *nleD* deletion mutants in *C. rodentium* infection of mice (20) or enterohemorrhagic *E. coli* (EHEC) infection of infant ruminants (21). Subsequently, both NleC and NleD were identified to be metalloprotease effectors. Whereas NleC and its cleavage of NF- κ B are



FIG 6 Direct cleavage of p38 by NIeD *in vitro* and mass spectrometric analysis of putative cleavage site for p38. (A) Immunoblots of GST or GST-p38 incubated with His-NIeD or His-NIeD site-directed mutants. Arrows, p38 cleavage fragment. Reaction mixtures were probed with anti-GST antibodies, and anti-His antibodies were used to detect NIeD and NIeD variants. (B) A Coomassie-stained gel showing the N-terminal (N-term) and C-terminal (C-term) cleavage products of GST-p38 in the presence of His-tagged NIeD after incubation for 2 h or overnight (O/N). White boxes, bands that were excised from the gel and used for further mass spectrometric analysis. (C) Identification of semitryptic peptide ²¹³MAELLTGR²²⁰ only within the C-terminal p38 cleavage fragment supports the suggestion that cleavage of p38 occurs at or before position 213. M_{nv}, oxidized methionine residue; a, b, and y, peptide fragment ions.

very well characterized (9–15, 22), to date only one study has examined the substrate specificity and enzymatic function of NIeD (15). Baruch et al. showed that NIeD cleaved the MAP kinases p38 and JNK and that mutation of glutamate within the metalloprotease motif (H¹⁴²ELLH¹⁴⁶) abolished the cleavage of JNK. In addition, this study demonstrated direct cleavage of JNK by NIeD and identified the conserved TPY motif within the kinase activation loop of JNK to be the specific cleavage site (15). The activation of JNK/p38 influences transcription of the proinflammatory cytokines, such as IL-8 (23, 24). Previous studies showed that EPEC potently inhibits the production of proinflammatory cytokines early in infection (25, 26) and that NIeE and NIeC contribute significantly to this inhibition (2, 3, 12). Cleavage of JNK by NIeD makes a minor contribution to this inhibition, as a modest reduction in the level of IL-8 secretion was observed in cells infected with an EPEC mutant lacking *nIeC*, *nIeD*, *nIeE*, and *nIeB1* and complemented with wild-type *nIeD* (15).

Here, we screened a library of insertion mutants of NIeD for their ability to cleave the host substrates p38 and JNK during transient transfection of cultured epithelial cells. A total of 78 unique insertion mutants were constructed, and of these, 22 insertions affected p38 and/or JNK cleavage. Further mutagenesis in the form of multiple or single site-directed mutations was performed to assess the contribution of specific amino acids to the cleavage of host substrates. Amino acid substitutions at HNL from positions 149 to 151 [HNL₍₁₄₉₋₁₅₁₎], GER from positions 153 to 155 [GER₍₁₅₃₋₁₅₅₎], TVG from positions 178 to 180 [TVG₍₁₇₈₋₁₈₀₎], and RRT from positions 202 to 204 [RRT₍₂₀₂₋₂₀₄₎]



FIG 7 Effect of the R203E mutation in NIeD on p38 cleavage and IL-6 secretion in EPEC infection. (A) Immunoblots of JNK and p38 cleavage in HT-29 cells infected with various EPEC derivatives. Cell lysates were probed with anti-JNK and anti-p38 antibodies. Anti-HA antibody was used to show the expression of NIeD-2HA and NIeD-2HA variants. Arrows, cleavage products. Actin was used as a loading control. The immunoblots are representative of those from at least 3 independent experiments. (B) HT-29 cells were infected with derivatives of EPEC, as indicated, for 2 h and left unstimulated or stimulated with TNF for 8 h (gray bars). Results are the means \pm SEMs from at least three independent experiments carried out in duplicate (*, *P* < 0.05; **, *P* < 0.005; *P* values were determined by an unpaired, two-tailed *t* test). UI, uninfected; UN, unstimulated.

abolished p38 cleavage, while cleavage of JNK remained unaffected. Analysis of multiple single site-directed mutants revealed that mutation of the R203 residue in NIeD abolished the cleavage of p38 but not that of JNK during both transient transfection and EPEC infection, suggesting that R203 in NIeD discriminates between these host substrates. An alignment of the region surrounding the zinc metalloprotease motif from a subset of effectors from various Gram-negative pathogens revealed that R203 in NIeD was highly conserved. This may reflect the importance of the conserved amino acid in host substrate recognition for other metalloprotease effectors; however, there is currently no structural information available for these effectors. Therefore, the function of R203 in NIeD proteolytic activity remains unknown.

NleD mediated the direct cleavage of p38, which was dependent on the catalytic motif of NleD, as mutation of E143 within the HEXXH motif abolished cleavage of the substrate. In addition, the R203E NleD mutant was also unable to directly cleave p38, further suggesting that R203 is essential for the proteolytic activity of NleD against p38. The purified R203E NleD mutant was still, however, active against JNK. A previous study identified the NleD cleavage site within the activation loop of JNK before residue Y185 of the TPY motif (15). No study has yet, however, identified the cleavage site within p38. Here, we have shown that cleavage does not occur within the TGY motif of p38 (residues 180 to 182), as we observed a clear intact peptide containing this motif within the N-terminal cleavage fragment. We did, however, narrow down the cleavage site to a region just downstream of the TGY motif, between residues 187 and 213, suggesting that the cleavage sites of p38 and JNK differ. Mass spectrometric analysis of full-length, N- and C-terminal fragments of p38 suggests that the most likely cleavage site is M213.

The loss of p38 cleavage attributed to the R203 mutation in NleD significantly

Bacterial strain	Characteristics	Source or reference
EPEC E2348/69	Wild-type EPEC 0127:H6	37
∆nleD	EPEC E2348/69 Δ <i>nleD</i> Km ^r	This study
$\Delta escN$	EPEC E2348/69 ΔescN Km ^r	38
ΔPP4/IE6	EPEC E2348/69 PP4/IE6 double island deletion mutant	2
XL1-Blue	E. coli recA1 endA1 gyrA96 thi-1 hsdR17 supE44 relA1 lac [F' proAB laclqZΔM15 Tn10 (Tetr)]	Stratagene
BL21 C43(DE3)	<i>E. coli</i> strain used for expression of proteins for affinity purification	Novagen

TABLE 1 Bacterial strains used in this study

affected MAPK signaling in host cells. AP-1 is a transcriptional activator located downstream of the MAP kinases p38 and JNK that mediates transcription of proinflammatory cytokines, such as IL-6, in host cells (27, 28). NIeD inhibited transcription of an AP-1 luciferase reporter in response to PMA, which was dependent on the R203 residue in NIeD, as radical mutation of R203 to glutamate most significantly affected NIeD activity.

Since a consequence of AP-1 activation is the production of the proinflammatory cytokine IL-6 (28), we examined the ability of NIeD to suppress IL-6 production. Previous studies have shown that IL-6 is required for protection against colonization, mucosal pathology, and mortality in C. rodentium infection of mice (29). A single nleD deletion mutant of EPEC was unable to cleave JNK or p38; therefore, in order to test the contribution of NIeD to inflammatory cytokine production in EPEC infection, we utilized a double island deletion mutant lacking seven effectors (nleB1, nleB2, nleC, nleD, nleE, nleG, and espL) and complemented this with wild-type nleD or nleD site-directed mutants. The reason that we chose to utilize the seven-effector mutant is because NIeE, NIeB, and NIeC all significantly contribute to the inhibition of inflammatory cytokine production (2, 12, 15). Therefore, we aimed to test the relative contribution of NIeD in this genetic background. Here, wild-type EPEC inhibited IL-6 production in response to TNF stimulation, to which NleD made a significant contribution. Mutation of R203 in NIeD significantly abrogated IL-6 inhibition, suggesting that p38 activation contributed to IL-6 production during EPEC infection and that NIeD specifically targets p38 to dampen the IL-6 response. Interestingly, cleavage of JNK alone by NIeD was insufficient to mediate the suppression of IL-6 during EPEC infection.

Overall, the potent early inhibition of inflammation in EPEC infection is a result of the translocation of multiple type III effectors that specifically target innate immune signaling pathways, including NF- κ B (NIeE, NIeC, NIeB1, NIeH1, NIeH2, Tir) (2–5, 12, 13, 30) and MAPK (NIeC and NIeD) (13, 15). Here we showed that NIeD blocked IL-6 production through cleavage of the MAPK p38 during EPEC infection. Hence, NIeD is yet another example of an effector that targets host immune processes and contributes to the dampening of inflammatory responses during EPEC infection.

MATERIALS AND METHODS

Bacterial strains, plasmids, cell lines, and growth conditions. The bacterial strains and plasmids used in this study are listed in Tables 1 and 2, respectively. All PCR primers are listed in Table 3. Bacteria were grown at 37°C in Luria-Bertani (LB) medium, Dulbecco's modified Eagle's medium (DMEM) with GlutaMAX (Gibco), or RPMI with GlutaMAX (Gibco) where indicated, and the media were supplemented with ampicillin (100 μ g/ml), kanamycin (100 μ g/ml), or chloramphenicol (25 μ g/ml) where necessary. HEK293T cells were grown in DMEM with GlutaMAX (Gibco) supplemented with 10% fetal calf serum (FCS; Sigma) at 37°C with 5% CO₂. HT-29 cells were grown in RPMI with GlutaMAX (Gibco) supplemented with 10% FCS (Sigma) at 37°C with 5% CO₂.

Random-insertion scanning mutagenesis and site-directed mutagenesis. Generation of an NIeD pentapeptide scanning mutagenesis library was carried out in the plasmid pGFP-NIeD using a mutation generation system kit (catalog number F-701; Thermo Scientific) with an entranceposon (M1-Cam') according to the manufacturer's instructions. Three independent mutagenesis reactions were performed. *E. coli* strain XL1-Blue was used as the transformation host for the chemical transformation steps, and agar plates were incubated for 40 h at 37°C. The restriction enzymes EcoRl and BamHI (New England BioLabs) were used to clone *nleD* open reading frames containing the entranceposon into the vector backbone of pEGFP-C2 without entranceposon insertions. The 15-bp insertions using primers pEGFP-C2_F and pEGFP-C2_R in combination with the Notl miniprimer (Table 3). The exact position of the

TABLE 2 Plasmids used in this study

PlasmidRelevant characteristicsreferencepEGFP-C2Expression vector carrying enhanced GFP, Kar'ClontechpGFP-NleDFull-length <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G100 mutated to A, Kar'This studypGFP-NleD_113A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G100 mutated to A, Kar'This studypGFP-NleD_113A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid L111 mutated to A, Kar'This studypGFP-NleD_113A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid L111 mutated to A, Kar'This studypGFP-NleD_114A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid L114 mutated to A, Kar'This studypGFP-NleD_114A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid N150 mutated to A, Kar'This studypGFP-NleD_114A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid N150 mutated to A, Kar'This studypGFP-NleD_115A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G160 mutated to A, Kar'This studypGFP-NleD_115A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G160 mutated to A, Kar'This studypGFP-NleD_115A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G160 mutated to A, Kar'This studypGFP-NleD_2020A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G160 mutated to A, Kar'This studypGFP-NleD_2020A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G160 mutated to A, Kar'This studypGFP-NleD_2020A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid C100 mutated to A, Kar'This studypGFP-NleD_2020A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with			Source or
PEGFP-C2 Expression vector carrying enhanced GFP, Kan' Clontech pGFP-NIeD Full-length <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G ₁₁₀ mutated to A, Kan' This study pGFP-NIeD_T11A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G ₁₁₀ mutated to A, Kan' This study pGFP-NIeD_T11A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G ₁₁₀ mutated to A, Kan' This study pGFP-NIeD_T11A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₄₄ mutated to A, Kan' This study pGFP-NIeD_T13A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₄₄ mutated to A, Kan' This study pGFP-NIeD_T13A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₄₄ mutated to A, Kan' This study pGFP-NIeD_T13A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R ₁₅₄ mutated to A, Kan' This study pGFP-NIeD_T13A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R ₁₅₄ mutated to A, Kan' This study pGFP-NIeD_T13A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R ₁₅₄ mutated to A, Kan' This study pGFP-NIeD_T23A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R ₁₅₄ mutated to A, Kan' This study pGFP-NIeD_T23A <i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R ₁₅₆ mutated to A, Kan'	Plasmid	Relevant characteristics	reference
pGPP-NieDFull-length nleD from EPEC E3248/69 in pEGPP-C2, Kan'ZpGP-NieD_0110AnleD from EPEC E3248/69 in pEGP-C2 with amino acid G110 mutated to A, Kan'This studypGP-NieD_111AnleD from EPEC E3248/69 in pEGP-C2 with amino acid G111 mutated to A, Kan'This studypGP-NieD_113AnleD from EPEC E3248/69 in pEGP-C2 with amino acid H111 mutated to A, Kan'This studypGP-NieD_114AnleD from EPEC E3248/69 in pEGP-C2 with amino acid H114 mutated to A, Kan'This studypGP-NieD_114AnleD from EPEC E3248/69 in pEGP-C2 with amino acid H116 mutated to A, Kan'This studypGP-NieD_114AnleD from EPEC E3248/69 in pEGP-C2 with amino acid H116 mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E3248/69 in pEGP-C2 with amino acid H116 mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E3248/69 in pEGP-C2 with amino acid H116 mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E3248/69 in pEGP-C2 with amino acid H116 mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E3248/69 in pEGP-C2 with amino acid R115 mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E3248/69 in pEGP-C2 with amino acid R301 mutated to A, Kan'This studypGP-NieD_202AnleD from EPEC E3248/69 in pEGP-C2 with amino acid R301 mutated to A, Kan'This studypGP-NieD_202AnleD from EPEC E3248/69 in pEGP-C2 with amino acid R301 mutated to A, Kan'This studypGP-NieD_202AnleD from EPEC E3248/69 in pEGP-C2 with amino acid R301 mutated to A, Kan'This studypGP-NieD_202AnleD from EPEC E3248/69 in pEGP-C2 with amino	pEGFP-C2	Expression vector carrying enhanced GFP, Kan ^r	Clontech
pGPP-NieD_110AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G10mutated to A, Kan'This studypGP-NieD_113AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G111mutated to A, Kan'This studypGP-NieD_113AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G111mutated to A, Kan'This studypGP-NieD_113AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G114mutated to A, Kan'This studypGP-NieD_114AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G14mutated to A, Kan'This studypGP-NieD_114AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G14mutated to A, Kan'This studypGP-NieD_114AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G14mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G155mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G100mutated to A, Kan'This studypGP-NieD_115AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200mutated to A, Kan'This studypGP-NieD_2020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200mutated to A, Kan'This studypGP-NieD_2020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200mutated to A, Kan'This studypGP-NieD_2020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200mutated to A, Kan'This studypGP-NieD_2020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200mutated to A, Kan'This studypGP-NieD_2020BnleD from EPEC E23	pGFP-NIeD	Full-length nleD from EPEC E2348/69 in pEGFP-C2, Kan ^r	2
pGPF-NieD_111AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid T111 mutated to A, Kan'This studypGF-NieD_113AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid L141 mutated to A, Kan'This studypGFP-NieD_143AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid L141 mutated to A, Kan'This studypGFP-NieD_143AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid L141 mutated to A, Kan'This studypGFP-NieD_143AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid L141 mutated to A, Kan'This studypGFP-NieD_143AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid L141 mutated to A, Kan'This studypGFP-NieD_153AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid L153 mutated to A, Kan'This studypGFP-NieD_153AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid G135 mutated to A, Kan'This studypGFP-NieD_153AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid G130 mutated to A, Kan'This studypGFP-NieD_203AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid G30 mutated to A, Kan'This studypGFP-NieD_203AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid G30 mutated to A, Kan'This studypGFP-NieD_203AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid G30 mutated to A, Kan'This studypGFP-NieD_203AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid C300 mutated to A, Kan'This studypGFP-NieD_203AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid C300 mutated to A, Kan'This studypGFP-NieD_203AnleD from EPEC E3248/69 in pEGFP-C2 with amino acid C300 mutated to A, Kan'This studypGFP-NieD	pGFP-NIeD _{G110A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid G_{110} mutated to A, Kan ^r	This study
pGPF-NIED_T13AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid F ₁₁₅ mutated to Å, KarrThis studypGFP-NIED_T13AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₄₄ mutated to Å, KarrThis studypGFP-NIED_T14AAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₄₄ mutated to Å, KarrThis studypGFP-NIED_T14AAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₅₄ mutated to Å, KarrThis studypGFP-NIED_T15AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₅₄ mutated to Å, KarrThis studypGFP-NIED_T15AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₅₄ mutated to Å, KarrThis studypGFP-NIED_T15AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₅₄ mutated to Å, KarrThis studypGFP-NIED_T15AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₁₅₄ mutated to Å, KarrThis studypGFP-NIED_R020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid L ₂₀₂ mutated to Å, KarrThis studypGFP-NIED_R020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R ₂₀₃ mutated to Å, KarrThis studypGFP-NIED_R020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R ₂₀₃ mutated to Å, KarrThis studypGFP-NIED_R020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R ₂₀₃ mutated to Å, KarrThis studypGFP-NIED_R020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R ₂₀₃ mutated to Å, KarrThis studypGFP-NIED_R020AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R ₂₀₃ mutated to Å, KarrThis studypGFP-NIED_R020AnleD from EPEC E2348/69 in pEGFP-C2 with the NCR_2720 motif mutated to Å, Karr </td <td>pGFP-NIeD_{T1114}</td> <td><i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid T_{111} mutated to A, Kan^r</td> <td>This study</td>	pGFP-NIeD _{T1114}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid T_{111} mutated to A, Kan ^r	This study
pGFP-NleD_H3AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid E_143 mutated to A, Kar'This studypGFP-NleD_H4AAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_146 mutated to A, Kar'This studypGFP-NleD_H16AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_156 mutated to A, Kar'This studypGFP-NleD_H16AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_156 mutated to A, Kar'This studypGFP-NleD_H15AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_156 mutated to A, Kar'This studypGFP-NleD_H15AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_156 mutated to A, Kar'This studypGFP-NleD_H202AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_H202AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_R02AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_R02AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_R02AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_R02AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_R02AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_R02AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H_200 mutated to A, Kar'This studypGFP-NleD_R02AnleD from EPEC E2348/69 in pEGFP-C2 with the NL(50-91) motif mutated to A, Kar'This study <td>pGFP-NIeD_{F115A}</td> <td><i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid F_{115} mutated to A, Kan^r</td> <td>This study</td>	pGFP-NIeD _{F115A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid F_{115} mutated to A, Kan ^r	This study
pGFP-NIeD_LiaaAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Lia4mutated to A, Kan'This studypGFP-NIeD_Hi46AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Lia4mutated to A, Kan'This studypGFP-NIeD_Hi56AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris5mutated to A, Kan'This studypGFP-NIeD_Hi56AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris5mutated to A, Kan'This studypGFP-NIeD_Hi56AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris5mutated to A, Kan'This studypGFP-NIeD_Hi56AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris5mutated to A, Kan'This studypGFP-NIeD_Hi56AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris6mutated to A, Kan'This studypGFP-NIeD_Ris5AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris6mutated to A, Kan'This studypGFP-NIeD_Ris5AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris6mutated to A, Kan'This studypGFP-NIeD_Ris5AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris6mutated to A, Kan'This studypGFP-NIeD_Ris5AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ris6mutated to A, Kan'This studypGFP-NIeD_NieD_Ris6AnleD from EPEC E2348/69 in pEGFP-C2 with the LN ₇₂₋₂₇₃ motif mutated to AA, Kan'This studypGFP-NIeD_K192-130AnleD from EPEC E2348/69 in pEGFP-C2 with the LN ₇₂₋₂₇₃ motif mutated to AA, Kan'This studypGFP-NIeD_K192-130AnleD from EPEC E2348/69 in pEGFP-C2 with the LN ₇₂₋₂₇₃ motif mutated to AAA, Kan'This studypGFP-N	pGFP-NIeD _{E143A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid E_{143} mutated to A, Kan ^r	This study
pGFP-NIeD_H146A pGFP-NIeD_H146AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H146 mutated to A, Kan'This studypGFP-NIeD_H156AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H146 mutated to A, Kan'This studypGFP-NIeD_H156AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H156 mEPEC E2348/69 in pEGFP-C2 with amino acid H156 mepEGFP-NIeD_H156ANan'This studypGFP-NIeD_H156AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H156 mEPEC E2348/69 in pEGFP-C2 with amino acid H156 mepEGFP-NIeD_H166Nan'This studypGFP-NIeD_H166 mEPEC E2348/69 in pEGFP-C2 with amino acid H250 mepEGFP-NIeD_H2503AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H250 mutated to A, Kan'This studypGFP-NIeD_H2503AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H250 mepEGF NIED_H2503AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H250 mutated to A, Kan'This studypGFP-NIeD_H2503AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H250 mepEGF NIED_H250AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H250 mutated to A, Kan'This studypGFP-NIED_H2513AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid H250 mepEGF NIED_H250AnleD from EPEC E2348/69 in pEGFP-C2 with the LN550-010 mutated to A, Kan'This studypGFP-NIED_H2523AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN550-010 mutated to AA, Kan'This studypGFP-NIED_H2523-023AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN550-010 mutated to AA, Kan'This studypGFP-NIED_H2523-023AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN550-010 mutated to AA, Kan'This studypGFP-NIE	pGFP-NIeD ₁₁₄₄	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid L_{144} mutated to A, Kan ^r	This study
pGFP-NieD Ni50AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ni50 mutated to A, Kan'This studypGFP-NieD FIS5AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid Ni50 mutated to A, Kan'This studypGFP-NieD PINED PINED PGFP-NieD PGFP-NieD PINED PGFP-Nie	pGFP-NIeD	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid H_{146} mutated to A, Kan ^r	This study
pGFP-NIeD pGFP-NieD 	pGFP-NIeD _{N150A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid N_{150} mutated to A, Kan ^r	This study
pGFP-NIeD PGFP-NIeD PGFP-NIeD V179AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R155 mutated to A, Kan'This studypGFP-NIeD PGFP-NIeD C180AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R155 mutated to A, Kan'This studypGFP-NIeD R202AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R202 mutated to A, Kan'This studypGFP-NIeD R202AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R202 mutated to A, Kan'This studypGFP-NIeD R203AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD R203AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD R203AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD R203AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD D218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R211 mutated to A, Kan'This studypGFP-NIeD D15(55-77)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN(72-73) motif mutated to AA, Kan'This studypGFP-NIeD D15(55-77)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LS(72-73) motif mutated to AA, Kan'This studypGFP-NIeD D15(55-77)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LS(72-73) motif mutated to AA, Kan'This studypGFP-NIeD D15(55-77)AAnleD from EPEC E2348/69 in pEGFP-C2 with the VF (147-148) motif mutated to AA, Kan'This studypGFP-NIeD D15(57-77)AAnleD from EPEC E2348/69 in pEGFP-C	pGFP-NIeD _{F154A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid E_{154} mutated to A, Kan ^r	This study
pGFP-NIeD DGFP-NIeD G180AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid V179 mutated to A, Kan'This studypGFP-NIeD G180AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD R203AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD R203AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD R203KnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NIeD R203KnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mutated to A, Kan'This studypGFP-NIeD R203KnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mutated to A, Kan'This studypGFP-NIeD R218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mutated to A, Kan'This studypGFP-NIeD R218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mutated to AA, Kan'This studypGFP-NIeD L0(R2575-77)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the LN(72-73) molf mutated to AA, Kan'This studypGFP-NIeD L0(R27-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the BR(100-101) motif mutated to AA, Kan'This studypGFP-NIeD L0(R10-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the BR(100-101) motif mutated to AA, Kan'This studypGFP-NIeD L0(R114-110)AAnleD from EPEC E2348/69 in pEGFP-C2 with the BR(100-101) motif mutated to AAA, Kan'This studypGFP-NIeD L0(R114-110)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the	pGFP-NIeD _{P155A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R_{155} mutated to A, Kan ^r	This study
pGFP-NIeD GTP-NIeD GTP-NIeD R2003AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid G180 mutated to A, Kan'This studypGFP-NIeD R2003AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to A, Kan'This studypGFP-NIeD R2003BnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to A, Kan'This studypGFP-NIeD R2003BnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to A, Kan'This studypGFP-NIeD R2004nleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mleD from EPEC E2348/69 in pEGFP-C2 with amino acid R201 mutated to A, Kan'This studypGFP-NIeD P204AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R201 mleD from EPEC E2348/69 in pEGFP-C2 with amino acid R201 mutated to A, Kan'This studypGFP-NIeD P218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R201 mleD from EPEC E2348/69 in pEGFP-C2 with the LDK(59-61) motif mutated to AA, Kan'This studypGFP-NIeD PGFP-NIeD PGFP-NIeD PGFP-NIeD PGFP-NIeD PGFP-NIeD PGFP-NIeD PGFP-NIeD PGFP-NIeD PGFP-NIED PGFP-	pGFP-NIeD _{V170A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid V_{170} mutated to A, Kan ^r	This study
pGFP-NleD PGP-NleD PGP-NleD PGPAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to A, Kan'This study This studypGFP-NleD PGP-NleD PGPAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NleD PGP-NleD PGPAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NleD PGP-NleD PGPAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NleD PGP-NleD PCPAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mutated to A, Kan'This studypGFP-NleD PCPAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mutated to A, Kan'This studypGFP-NleD PCPAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R205 motif mutated to AA, Kan'This studypGFP-NleD pGFP-	pGFP-NIeD _{C1804}	nleD from EPEC E2348/69 in pEGFP-C2 with amino acid G ₁₀₀ mutated to A. Kan ^r	This study
pGFP-NleP PGFP-NleP R203AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This study This studypGFP-NleP R203BnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to K, Kan'This studypGFP-NleP R203RnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R203 mutated to A, Kan'This studypGFP-NleP R203RnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mleD from EPEC E2348/69 in pEGFP-C2 with amino acid R204 mleD from EPEC E2348/69 in pEGFP-C2 with the LN(r2-73) motif mutated to AA, Kan'This studypGFP-NleD L0K(59-61)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN(r2-73) motif mutated to AA, Kan'This studypGFP-NleD L0K(59-61)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN(r2-73) motif mutated to AA, Kan'This studypGFP-NleD L0K(59-61)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN(r2-73) motif mutated to AA, Kan'This studypGFP-NleD L0K(109-11)AAnleD from EPEC E2348/69 in pEGFP-C2 with the NF(r14-r148) motif mutated to AA, Kan'This studypGFP-NleD L0F(114-116)AAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(r14-r148) motif mutated to AAA, Kan'This studypGFP-NleD L0F(114-116)AAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(r14-r148) motif mutated to AAA, Kan'This studypGFP-NleD L0F(114-116)AAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(r14-r148) mleD from EPEC E2348/69 in pEGFP-C2 with the HNF(r14-r148) mleD from EPEC E2	pGFP-NIeD Bases	nleD from EPEC E2348/69 in pEGFP-C2 with amino acid R ₂₀₀ mutated to A. Kan ^r	This study
pGFP-NleD PGFP.NleD PGP-NleD P204AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to E, Kan'This studypGFP-NleD P204AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to E, Kan'This studypGFP-NleD P204AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to A, Kan'This studypGFP-NleD P218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to A, Kan'This studypGFP-NleD P218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid R200 mutated to A, Kan'This studypGFP-NleD LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN(50-61) motif mutated to AA, Kan'This studypGFP-NleD LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LN(50-61) motif mutated to AA, Kan'This studypGFP-NleD LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the HC(100-101) motif mutated to AA, Kan'This studypGFP-NleD LN(100-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the HC(100-101) motif mutated to AA, Kan'This studypGFP-NleD P(1114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HC(140-148) motif mutated to AA, Kan'This studypGFP-NleD PGFP-NleD LN(149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HV(140-148) motif mutated to AAA, Kan'This studypGFP-NleD LN(149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HV(140-148) motif mutated to AAA, Kan'This studypGFP-NleD LN(149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HV(140-148) motif mutated to AAA, Kan'This studypGFP-NleD LN(149-151)AAAnleD from EPEC	pGFP-NIeD _{R202A}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R_{202} mutated to A, Kan ^r	This study
pGFP-NIeD PGFP-NIeD PCFP-NIeD PCFP-NIeD PCFP-NIeD PCFP-NIeD PCFP-NIeD PCFP-NIeD PCFP-NIED 	pGFP-NIeD _{P2025}	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with amino acid R_{203} mutated to E. Kan ^r	This study
pGFP-NIeD TOPALnleD from EPEC E2348/69 in pEGFP-C2 with amino acid T 204 mutated to A, Kan'This studypGFP-NIeD T218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid T 204 mutated to A, Kan'This studypGFP-NIeD LDK(59-61)AAAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid T 204 mutated to A, Kan'This studypGFP-NIeD LDK(59-61)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the LDK(29-61) motif mutated to AA, Kan'This studypGFP-NIeD LE(575-77)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LDK(272-73) motif mutated to AA, Kan'This studypGFP-NIeD LE(575-77)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HS(100-101) motif mutated to AAA, Kan'This studypGFP-NIeD LE(575-77)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HS(100-101) motif mutated to AAA, Kan'This studypGFP-NIeD LF(147-148)AAnleD from EPEC E2348/69 in pEGFP-C2 with the HS(100-101) motif mutated to AAA, Kan'This studypGFP-NIeD DFH114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(147-148) motif mutated to AAA, Kan'This studypGFP-NIeD DFFNIED LF(146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(146-148) motif mutated to AAA, Kan'This studypGFP-NIED PNIED LFVC1146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(146-148) motif mutated to AAA, Kan'This studypGFP-NIED LFVC1VG(178-180)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(146-148) motif mutated to AAA, Kan'This studypGFP-NIED LFVL146-148nleD from EPEC E2348/69 in pEGFP-C2 with the EMC(149-151) motif mutated to AAA, Kan'This study <td< td=""><td>pGFP-NIeD_{R203E}</td><td>nleD from EPEC E2348/69 in pEGFP-C2 with amino acid R₂₀₃ mutated to L, Kan^r</td><td>This study</td></td<>	pGFP-NIeD _{R203E}	nleD from EPEC E2348/69 in pEGFP-C2 with amino acid R ₂₀₃ mutated to L, Kan ^r	This study
pGFP-NIED_DIALIntel from EPEC E2348/69 in pEGFP-C2 with amino acid N217 mutated to A, Kan'This studypGFP-NIED_T218AnleD from EPEC E2348/69 in pEGFP-C2 with amino acid N217 mutated to A, Kan'This studypGFP-NIED_LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with amino acid N217 mutated to AA, Kan'This studypGFP-NIED_LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LDN(59-61) motif mutated to AAA, Kan'This studypGFP-NIED_LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LDN(59-61) motif mutated to AAA, Kan'This studypGFP-NIED_HR(100-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the HS(100-101) motif mutated to AAA, Kan'This studypGFP-NIED_RGT(109-111)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RGT(109-111) motif mutated to AAA, Kan'This studypGFP-NIED_DFH(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RGT(109-111) motif mutated to AAA, Kan'This studypGFP-NIED_DFH(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RGT(109-111) motif mutated to AAA, Kan'This studypGFP-NIED_DFH(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVF(146-148) motif mutated to AAA, Kan'This studypGFP-NIED_HNIC149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVF(146-148) motif mutated to AAA, Kan'This studypGFP-NIED_HNIC149-180/AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVG(178-180) motif mutated to AAA, Kan'This studypGFP-NIED_HNIC149-180/AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RT(129-197) motif mutated to AAA, Kan'This studypGFP-NIED_RT(202-200)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RT(pGFP-NIeD _{Table}	nleD from EPEC E2348/69 in pEGEP-C2 with amino acid T ₂₀₃ mutated to A, Kan ^r	This study
pGFP-NIED_T18AInto Trutt PECpECpECpCwith the min or cid T218mutated to A, Kan'This studypGFP-NIED_LDK(59-61)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the LDK(59-61) motif mutated to AA, Kan'This studypGFP-NIED_LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LDK(59-61) motif mutated to AA, Kan'This studypGFP-NIED_LN(72-73)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LDK(59-77) motif mutated to AA, Kan'This studypGFP-NIED_RGT(109-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the EGT(109-111) motif mutated to AA, Kan'This studypGFP-NIED_RGT(109-111)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RGT(109-111) motif mutated to AA, Kan'This studypGFP-NIED_VF(147-148)AAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(147-148) motif mutated to AA, Kan'This studypGFP-NIED_HVIC149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(147-148) motif mutated to AAA, Kan'This studypGFP-NIED_HVIC149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVF(146-148) motif mutated to AAA, Kan'This studypGFP-NIED_HVIC149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVF(146-148) motif mutated to AAA, Kan'This studypGFP-NIED_HVIC149-151)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVC(178-180) motif mutated to AAA, Kan'This studypGFP-NIED_RVIC178-180)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RT(202-204) motif mutated to AAA, Kan'This studypGFP-NIED_RRE(202-204)AAnleD from EPEC E2348/69 in pEGFP-C2 with the RR(202-204) motif mutated to AAA, Kan'This studypGFP-NIED_RRE(202-204)AAA<	nGEP-NIeD	nleD from EPEC E2348/69 in pEGEP-C2 with amino acid N_{204} instanced to A Kan ^r	This study
pGFP-NIED_LDK(59-GI)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the LDK(59-GI) motif mutated to AAA, Kan'This studypGFP-NIED_LDK(29-GI)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the LDK(59-GI) motif mutated to AAA, Kan'This studypGFP-NIED_HR(100-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the LB(75-77) motif mutated to AAA, Kan'This studypGFP-NIED_HR(100-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the HR(100-101) motif mutated to AAA, Kan'This studypGFP-NIED_FR(110-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the HR(100-101) motif mutated to AAA, Kan'This studypGFP-NIED_FR(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(147-148) motif mutated to AAA, Kan'This studypGFP-NIED_FH(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the DFH(114-116) motif mutated to AAA, Kan'This studypGFP-NIED_FH(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the DFH(114-116) motif mutated to AAA, Kan'This studypGFP-NIED_HNE(146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HNL(149-151) motif mutated to AAA, Kan'This studypGFP-NIED_HNE(146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HNL(149-151) motif mutated to AAA, Kan'This studypGFP-NIED_HNE(146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the GER(153-155) motif mutated to AAA, Kan'This studypGFP-NIED_TNC(178-180)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RT(120-204) motif mutated to AAA, Kan'This studypGFP-NIED_RE(152-197)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RT(202-204) motif mutated to AAA, Kan'This studypGFP-NIED_RE(152-180)AAAnleD fro	pGFP-NIeD	nleD from EPEC E2348/69 in pEGEP-C2 with amino acid T _{exp} mutated to A. Kan ^r	This study
pGFP-NIED_EUX[22-73]AAnleD from EPEC E2348/69 in pEGFP-C2 with the $LN(22-73)$ motif mutated to AA, Kar'This studypGFP-NIED_HR(100-101)AAnleD from EPEC E2348/69 in pEGFP-C2 with the $LS(25-77)$ motif mutated to AA, Kar'This studypGFP-NIED_RGT(109-111)AAnleD from EPEC E2348/69 in pEGFP-C2 with the RGT(109-111) motif mutated to AA, Kar'This studypGFP-NIED_RGT(109-111)AAnleD from EPEC E2348/69 in pEGFP-C2 with the RGT(109-111) motif mutated to AA, Kar'This studypGFP-NIED_FH(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the VF(147-148) motif mutated to AA, Kar'This studypGFP-NIED_DFH(114-116)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the DFH(114-116) motif mutated to AAA, Kar'This studypGFP-NIED_HXC146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the DFH(114-116) motif mutated to AAA, Kar'This studypGFP-NIED_HXC146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVF(146-148) motif mutated to AAA, Kar'This studypGFP-NIED_HXC146-148)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVF(146-148) motif mutated to AAA, Kar'This studypGFP-NIED_GER(153-155)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the HVF(146-148) motif mutated to AAA, Kar'This studypGFP-NIED_RTVG(178-180)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RE(195-197) motif mutated to AAA, Kar'This studypGFP-NIED_RET(152-159)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RE(195-197) motif mutated to AAA, Kar'This studypGFP-NIED_RTTC20-204)AAAnleD from EPEC E2348/69 in pEGFP-C2 with the RT(202-204) motif mutated to AAA, Kar'This studypGFP-NIED_DNT(216-218)AAAnl	pGFP-NIeD, pyrso control	<i>nleD</i> from EPEC E2348/69 in pEGEP-C2 with the LDK _{resc} motif mutated to AAA. Kan ^r	This study
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	nGEP-NIeD W/20 2010	nleD from EPEC E2348/69 in pEGEP-C2 with the $IN_{(39-61)}$ motif mutated to AA Kan ^r	This study
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	nGEP-NIeD	<i>nleD</i> from EPEC E2348/69 in pEGEP-C2 with the IES_{res} motif mutated to AAA Kan ^r	This study
pGFP-NIeD _{RGT(199-111)AAA} pGFP-NIeD _{RGT(199-111)AAA} pGFP-NIeD _{DFH(14-148)AA} pGFP-NIeD _{DFH(14-148)AA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(14-148)AAA} pGFP-NIeD _{DFH(146-148)AAA} pGFP-NIeD _{DFH(146-148)AAA} pGFP-NIeD _{DFH(146-151)AAA} pGFP-NIeD _{DFH(149-151)AAA} pGFP-NIeD _{DFNIeD} pGFP-NIeD _{DFH(149-151)AAA} pGFP-NIeD _{DF} (149-151)AAA pGFP-NIeD _{DF} (1	nGEP-NIeD	nleD from EPEC E2348/69 in pEGEP-C2 with the HB motif mutated to AA Kan ^r	This study
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	nGEP-NIeD	nleD from EPEC F2348/69 in pEGEP-C2 with the RGT and motif mutated to AAA Kan ^r	This study
$ \begin{array}{llllllllllllllllllllllllllllllllllll$	nGEP-NIeD.	nleD from EPEC E2348/69 in pEGEP-C2 with the VE motif mutated to AA Kan ^r	This study
pGFP-NleD pCF-NleD pCF-NleD pCF-NleD pCFO	pGFP-NIeD	nleD from EPEC F2348/69 in pEGEP-C2 with the DEH model motif mutated to AAA. Kan'	This study
pGFP-NleD pGFP-NleD GFP-NleD GFP-NleD DFF-NleD GFP-NleD D HeD from EPEC E2348/69 in pEGFP-C2 with the RLH (227-229)AAA A N-terminal 6× His Tag cloning/expression vector Nevagen Nevagen MleD from EPEC E2348/69 in pET28a, Kan' This study This study This study MleD from EPEC E2348/69 in pET28a with amino acid E143 Mutated to A, Kan' This study This study This study This studyGFT-NleD D<	pGFP-NIeDuvrs(14c-140)AAA	nleD from EPEC F2348/69 in pEGEP-C2 with the HVE (114-116) motif mutated to AAA. Kan'	This study
pGFP-NIED pCT-NIED pET-NIED pET-NIED pET-NIED pET-NIED pET-NIED pFT-NIED pET-NIED pFT-NIED pET-NIED pFT-NIED pET-NIED<	pGFP-NIeD	nleD from EPEC E2348/69 in pEGFP-C2 with the HNL/140-143) motif mutated to AAA. Kan'	This study
pGFP-NIeD pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-NIED pGFP-CI pCT284 pGFP-NIED pGFP-CI pCT-NIED pGFP-CI pCT-NIED pGFP-CI pCT-NIED pGFP-CI pCT-NIED pGFP-CI pCT-NIED pGFP-CI pCT-NIED pGFP-CI pCT2848/69 in pECC pCT2848/69 in pECC pCT2848/69 in pET28a with amino acid E143 pGFT-NIED pCT-	pGFP-NIeD CER(152, 155)AAA	nleD from EPEC E2348/69 in pEGFP-C2 with the GER (163 157) motif mutated to AAA. Kan'	This study
pGFP-NleD pCFT-NleD pET-NleD <b< td=""><td>pGFP-NIeDTyc(178-180)AAA</td><td><i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with the $TVG_{(178, 180)}$ motif mutated to AAA. Kan'</td><td>This study</td></b<>	pGFP-NIeDTyc(178-180)AAA	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with the $TVG_{(178, 180)}$ motif mutated to AAA. Kan'	This study
pGFP-NIeD pET-NIED pET-NIeD pET-NIeD pET-NIeD pFT-NIeD pET-NIeD pFT-NIeD pFT-NIeD pFT-NIeD pFT-NIeD pFT-NIeD pFT-NIeD pFT-NIeD pET-NIeD pFT-NIeD pFT-NIeD pFT-NIeD pFT-NIeD pFT-NIED pHD from FPEC E2348/69 in pFT28a with amino acid E143 mutated to A, Kan' pFT-NIED pFT-NIED pFT-NIED pFT-NIED pFT-NIED pHD from FPEC E2348/69 in pFT28a with amino acid E143 mutated to A, Kan' pFT-NIED prom FPEC FT28a with amino acid E143 mutated to A, Kan' pHD from FPEC FT28a with amino acid E143 mutated to A, Kan' pHD from FPEC FT28a with amino acid E143 mutated to A, Kan' pHD from FPEC FT28a with amino acid E143 mutated to A, Kan' pHD from FPEC FT28a with amino	pGFP-NIeD	nleD from EPEC E2348/69 in pEGFP-C2 with the REE (107, 107) motif mutated to AAA, Kan'	This study
pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pGFP-NleD pET-NleDnleD from EPEC E2348/69 in pEGFP-C2 with the DNT (216-218) motif mutated to AAA, Kan' mutated to AAA, Kan' This studynleD from EPEC E2348/69 in pEGFP-C2 with the QVR (225-227) AAA pET-NleD pET-NleDnleD from EPEC E2348/69 in pEGFP-C2 with the QVR (225-227) motif mutated to AAA, Kan' This studynleD from EPEC E2348/69 in pEGFP-C2 with the RLH (227-229) MAA pET-NleD <td>pGFP-NIeD REE(195-197)AAA</td> <td>nleD from EPEC E2348/69 in pEGFP-C2 with the RRT(202, 204) motif mutated to AAA, Kan^r</td> <td>This study</td>	pGFP-NIeD REE(195-197)AAA	nleD from EPEC E2348/69 in pEGFP-C2 with the RRT(202, 204) motif mutated to AAA, Kan ^r	This study
pGFP-NIeD pGFP-NIeD pT-NIeDnleDfromEPECE2348/69inpEGFP-C2with theQVR(225-227)mutated toAAA, Kan'This studynleDfromEPECE2348/69inpEGFP-C2with theQVR(225-227)motifmutated toAAA, Kan'This studypET28anleDfromEPECE2348/69inpEGFP-C2with theRLH(227-229)motifmutated toAAA, Kan'This studypET-NIeDnleDfromEPECE2348/69inpET28a, Kan'NovagenNovagenpET-NIeDnleDfromEPECE2348/69inpET28a, with amino acidE143mutated toA, Kan'This studypET-NIeDnleDfromEPECE2348/69inpET28a, with amino acidE143mutated toA, Kan'This studypET-NIeDnleDfromEPECE2348/69inpET28a, with amino acidE143mutated toA, Kan'This studypET-NIeDnleDfromEPECE2348/69inpET28a, with amino acidE143mutated toA, Kan'This study	pGFP-NIeD RRT(202-204)AAA	nleD from EPEC E2348/69 in pEGEP-C2 with the DNT ₍₂₁₂₋₂₀₄₎ motif mutated to AAA. Kan'	This study
pGFP-NleD pET-NleD pET-NleD pET-NleD pET-NleD pET-NleD pET-NleD pET-NleDnleD from EPEC E2348/69 in pEGFP-C2 with the RLH (227-229) motif mutated to AAA, Kan'This studynleD from EPEC E2348/69 in pEGFP-C2 with the RLH (227-229) motif mutated to AAA, Kan'Novagen NovagenpET-NleD pET-NleD	pGFP-NIeDovr(216-218)AAA	<i>nleD</i> from EPEC E2348/69 in pEGFP-C2 with the OVR $_{(216-218)}$ motif mutated to AAA. Kan ^r	This study
pET28a N-terminal $6 \times$ His Tag cloning/expression vector Novagen pET-NleD $nleD$ from EPEC E2348/69 in pET28a, Kar' This study pET-NleD _{E143A} $nleD$ from EPEC E2348/69 in pET28a with amino acid E ₁₄₃ mutated to A, Kar' This study pET-NleD _{e143A} $nleD$ from EPEC E2348/69 in pET28a with amino acid E ₁₄₃ mutated to A, Kar' This study	pGFP-NIeD UV227 220) AAA	nleD from EPEC E2348/69 in pEGFP-C2 with the RLH(225-227) motif mutated to AAA, Kan ^r	This study
pET-NIeD nleD from EPEC E2348/69 in pET28a, Kan' This study pET-NIeD nleD from EPEC E2348/69 in pET28a with amino acid E143 mutated to A, Kan' This study pET-NIeD nleD from EPEC E2348/69 in pET28a with amino acid E143 mutated to A, Kan' This study pET-NIeD nleD from EPEC E2348/69 in pET28a with amino acid E143 mutated to A, Kan' This study	pET28a	N-terminal $6 \times$ His Tag cloning/expression vector	Novagen
pET-NIeD _{E143A} nleD from EPEC E2348/69 in pET28a with amino acid E ₁₄₃ mutated to A, Kan' This study pET-NIeD _{E143A} nleD from EPEC E2348/69 in pET28a with amino acid E ₁₄₃ mutated to F. Kan' This study	pET-NIeD	nleD from EPEC E2348/69 in pET28a. Kan ^r	This study
pFT-NIE Datase nleD from FPEC F2348/69 in pFT28a with amino acid Baco mutated to F. Kan' This study	pET-NIeD	<i>nleD</i> from EPEC E2348/69 in pET28a with amino acid E_{142} mutated to A. Kan ^r	This study
	pFT-NIeD	nleD from EPEC E2348/69 in pET28a with amino acid R ₂₀₂ mutated to F. Kan ^r	This study
DGEX-47-1 N-terminal GST cloning/expression vector. Amp ^r GE Healthcare	pGEX-4T-1	N-terminal GST cloning/expression vector. Amp ^r	GE Healthcare
pGEX-p38 Human p38 (MAPK14 transcript variant 2) in pGEX-4T-1, Amp ^r This study	pGEX-p38	Human $p38$ (MAPK14 transcript variant 2) in pGEX-4T-1. Amp ^r	This study
pTre99A Bacterial expression vector containing a <i>lact</i> promoter Amp ^r Pharmacia Biotech	pTrc99A	Bacterial expression vector containing a <i>lacl</i> promoter. Amp ^r	Pharmacia Biotech
nNIeD nIeD trom FPEC 2348/69 in pTrc99A Amp ^r 12	pNIeD	nleD from EPEC 2348/69 in pTrc99A. Amp	12
nleD from EPEC F3348/69 in pTrc99A with amino acid File mutated to A. Amp ^r This study	pNIeD	nleD from EPEC E2348/69 in pTrc99A with amino acid E ₄₄₀ mutated to A. Amp ^r	This study
p NIeD area need to be a need t	pNIeD _{P2025}	nleD from EPEC E2348/69 in pTrc99A with amino acid R ₂₀₂ mutated to F. Amp ^r	This study
pNIeD-ZUSA	pNIeD-2HA	nleD from EPEC 2348/69 in pTrc99A with a 2× C-terminal HA tag. Amp ^r	This study
pNIeD-ran-2HA $n eD$ from EPEC E2348/69 in pTrc99A with amino acid Examitated to A and a 2× This study	pNIeD	<i>nleD</i> from EPEC E2348/69 in pTrc99A with amino acid F mutated to A and a $2\times$	This study
C-terminal HA tag. Amp ^r	E143A	C-terminal HA tag. Amp ^r	
pNIeD ₀₂₀₂ -2HA $nleD$ from EPEC E3348/69 in pTrc99A with amino acid R ₂₀₂ mutated to E and a 2× This study	pNIeD _{B2025} -2HA	<i>nleD</i> from EPEC E2348/69 in pTrc99A with amino acid R_{202} mutated to E and a 2×	This study
C-terminal HA tag, Amp ^r	• N2UJL	C-terminal HA tag, Amp ^r	

15-bp insertion within *nleD* was determined by DNA sequencing of selected transformants using primer pEGFP-C2_R (Table 3).

Site-directed mutagenesis of *nleD* was performed in plasmid pGFP-NleD, pNleD, or pNleD-2HA as the template. The oligonucleotides used to generate the mutants are listed in Table 3. The mutant strand synthesis reaction was performed with *Pfu* DNA polymerase (Promega) under the following conditions: 95°C for 30 s; 18 cycles of 95°C for 30 s, 55°C for 1 min, and 68°C for 5.5 min; and a final extension at 72°C for 7 min. The amplification products were digested with Dpnl (New England BioLabs) and directly transformed into chemically competent cells of *E. coli* strain XL1-Blue. Mutations were confirmed by DNA sequencing using either primer pair pEGFP-C2_F/pEGFP-C2_R or primer pair pTrc_F/pTrc_R (Table 3).

Construction of plasmids to express NIeD and p38. pNIeD-2HA was constructed by amplifying *nIeD* from EPEC E2348/69 genomic DNA by PCR using the primer pair NIeD-2HA_F/NIeD-2HA_R. PCR amplification consisted of an initial denaturation step at 95°C for 10 min, followed by 30 cycles of 94°C

Primer	Sequence ^a
Pos. 18 _F	GTCATCAACGTCGAGCT TGCGGCCGCAGAGCT CAATGTCAGATACAGATATCG
Pos. 18 _R	CGATATCTGTATCTGACATTG AGCTCTGCGGCCGCA AGCTCGACGTTGATGAC
Pos. 50 _F	CGTTTCCAGGGGACTGATCA TGCGGCCGCAGATCA TAATATATATCAGCAGATTG
Pos. 50 _R	CAATCTGCTGATATATATTA TGATCTGCGGCCGCA TGATCAGTCCCCTGGAAACG
Pos. 60 _F	GCAGATTGAAGCAGCACTC TGCGGCCGCACACTC GATAAGATTGGCTCTACAG
Pos. 60 _R	CTGTAGAGCCAATCTTATC GAGTGTGCGGCCGCA GAGTGCTGCTTCAATCTGC
Pos. 91 _F	CAGTGGTAATACACCT TGCGGCCGCACACCT CAACTCTTCCAGACTAGG
Pos. 91 _R	CCTAGTCTGGAAGAGTTG AGGTGTGCGGCCGCA AGGTGTATTACCACTG
Pos. 121 _F	CGATTTTCACTGTAATCTGAAT TGCGGCCGCATGAAT GCAGTTGAATATCCCTGTGG
Pos. 121 _R	CCACAGGGATATTCAACTGC ATTCATGCGGCCGCA ATTCAGATTACAGTGAAAATCG
Pos. 128 _F	CAGTTGAATATCCCTGTGG TGCGGCCGCATGTGG GGAGGGGATTAGCGTGGTG
Pos. 128 _R	CACCACGCTAATCCCCTCC CCACATGCGGCCGCA CCACAGGGATATTCAACTG
Pos. 133 _F	GTGGGGAGGGATTAGCGT TGCGGCCGCAAGCGT GGTGGACTTTCATGCGAC
Pos. 133 _R	GTCGCATGAAAGTCCACC ACGCTTGCGGCCGCA ACGCTAATCCCCTCCCCAC
Pos. 144 _F	CGACTATIGITITICATGAG TGCGGCCGCAATGAG TTGCTCCATGTTTTCCAC
Pos. 144 _R	GIGGAAAACAIGGAGCAA CTCATTGCGGCCGCA CICAIGAAAAACAAIAGICG
Pos. 167	
Pos. 167 _R	
Pos. 191 _F	
Pos. 191 _R	
Pos. 207 _F	GATGULUGTAGAAUTULTATGUGGULGUATULTATUL
POS. 207 _R	
LN72-73AA	
ΙΕ375-77ΔΔΔ	
$HB100-101\Delta\Delta$	
HR100-101AA_	
DEH114-116AAA_	
DFH114-116AAA	GGGATATTCAACTGCATTCAGATTACACGCCGCCGCGGAACCAGTCCCCCGATGG
HVF146-148AAA _F	CGACTATTGTTTTCATGAGTTGCTC GCGGCGCG CACAATTTAAATGGGGAGCGTTTG
HVF146-148AAA _p	CAAACGCTCCCCATTTAAATTGTG CGCCGCCGC GAGCAACTCATGAAAAACAATAGTCG
VF147-148AA	GACTATTGTTTTCATGAGTTGCTCCAT GCGGCG CACAATTTAAATGGGGAGC
VF147-148AA _B	GCTCCCCATTTAAATTGTG CGCCGC ATGGAGCAACTCATGAAAAACAATAGTC
HNL149-151AAA _F	CATGAGTTGCTCCATGTTTTC GCGGCGGCG AATGGGGAGCGTTTGAAAGTTGAGAGTTC
HNL149-151AAA _R	GAACTCTCAACTTTCAAACGCTCCCCATT CGCCGCCGC GAAAACATGGAGCAACTCATG
GER153-155AAA _F	CCATGTTTTCCACAATTTAAAT GCGGCGGCG TTGAAAGTTGAGAGTTCCCGAC
GER153-155AAA _R	GTCGGGAACTCTCAACTTTCAA CGCCGCCGC ATTTAAATTGTGGAAAACATGG
TVG178-180AAA _F	CTCGAAGAAGCCAGG GCGGCGGCG TTGGGGGGCTTTTTCAGAG
TVG178-180AAA _R	CTCTGAAAAAGCCCCCAA CGCCGCCCC CCTGGCTTCTTCGAG
REE195-197AAA _F	GGTGCTTTCAGAAAATAAATTC GCGGCGGCG ATTGGGATGCCCCGTAGAACC
REE195-197AAA _R	GGTTCTACGGGGCATCCCAAT CGCCGCCGC GAATTTATTTTCTGAAAGCACC
RRT202-204AAA _F	CGAAGAGATTGGGATGCCC GCGGCGGCG TCCTACCCGCACGACTCAGC
RRT202-204AAA _R	GCTGAGTCGTGCGGGTAGGA CGCCGCCGC GGGCATCCCAATCTCTTCG
DNT216-218AAA _F	GACTCAGCTCTTATTCATGAT GCGGCGGCG GTGAGTCTGGGATTCCAACAGG
DNT216-218AAA _R	CCTGTTGGAATCCCAGACTCAC CGCCGCCGC ATCATGAATAAGAGCTGAGTC
QVR225-227AAA _F	CAGTGAGTCTGGGATTCCAA GCGGCGCGCG CTGCATCCATTGCTTTAG
QVR225-22/AAA _R	
RLH227-229AAA _F	GACAATACAGTGAGTCTGGGATTCCAACAGGTA GCGGCGCG CCATTGCTTAG
RLH227-229AAA _R	
E143A _F	
C110A	
GTIOA _F	
T111A	
Τ111Δ	
F115Δ	
F115A_	
1144A_	GACTATIGTITTCATGAG GCG CTCCATGTTTCCACAATTTAAATGG
1144A	CCATTTAAATTGTGGAAAACATGGAG CGC CTCATGAAAAACAATAGTC
H146A-	
H146A _p	CCATTTAAATTGTGGAAAACCGGGAGCAACTCATGAAAAACAATAGTCG
N150A	GTTGCTCCATGTTTTCCAC GCG TTAAATGGGGAGCGTTTG
N150A _R	CAAACGCTCCCCATTTAA CGC GTGGAAAACATGGAGCAAC

(Continued on next page)

TABLE 3 (Continued)

Primer	Sequence ^a
E154A _F	GTTTTCCACAATTTAAATGGG GCG CGTTTGAAAGTTGAGAG
E154A _B	CTCTCAACTTTCAAACG CGC CCCATTTAAATTGTGGAAAAC
R155A _F	CCACAATTTAAATGGGGAG GCG TTGAAAGTTGAGAGTTCCCGAC
R155A _R	GTCGGGAACTCTCAACTTTCAA CGC CTCCCCATTTAAATTGTGG
V179A _F	CGAAGAAGCCAGGACT GCG GGGTTGGGGGCTTTTTC
V179A _R	GAAAAAGCCCCCAACCC CGC AGTCCTGGCTTCTTCG
G180A _F	CGAAGAAGCCAGGACTGTT GCG TTGGGGGGCTTTTTCAGAGG
G180A _R	CCTCTGAAAAAGCCCCCAA CGC AACAGTCCTGGCTTCTTCG
R202A _F	CGAAGAGATTGGGATGCCC GCG AGAACCTCCTACCCGCACG
R202A _R	CGTGCGGGTAGGAGGTTCT CGC GGGCATCCCAATCTCTTCG
R203A _F	GATTGGGATGCCCCGT GCG ACCTCCTACCCGCAC
R203A _R	GTGCGGGTAGGAGGT CGC ACGGGGCATCCCAATC
R203E _F	GATTGGGATGCCCCGT GAG ACCTCCTACCCGCAC
R203E _R	GTGCGGGTAGGAGGT CTC ACGGGGCATCCCAATC
R203K _F	GATTGGGATGCCCCGT AAG ACCTCCTACCCGCAC
R203K _R	GTGCGGGTAGGAGGT CTT ACGGGGGCATCCCAATC
T204A _F	GAGATTGGGATGCCCCGTAGA GCG TCCTACCCGCACGACTCAG
T204A _R	CTGAGTCGTGCGGGTAGGA CGC TCTACGGGGCATCCCAATCTC
N217A _F	GCTCTTATTCATGATGAC GCG ACAGTGAGTCTGGGATTCC
N217A _R	GGAATCCCAGACTCACTGT CGC GTCATCATGAATAAGAGC
T218A _F	CTCTTATTCATGATGACAAT GCG GTGAGTCTGGGATTCCAACAG
T218A _R	CTGTTGGAATCCCAGACTCAC GC ATTGTCATCATGAATAAGAG
NIeD _F	AAGAATTCATGCGCCCTACGTCCCTC
NIeD _R	GGAGTCGACGCTAAAGCAATGGATGCAGTCTTAC
p38 _F	GGGGATCCATGTCTCAGGAGAGGCCCACG
p38 _R	CGCTCGAGTCAGGACTCCATCTTCTTGGTC
NIeD-2HA _F	AACCATGGATGCGCCCTACGTCCCTC
NIeD-2HA _R	CCGGATCCCTACGCATAATCCGGCACATCATACGGATACGCATAATCCGGCAC
	ATCATACGGATAAAGCAATGGATGCAGTCTTACC
pTrc _F	AACACCCCCATCGGCG
pTrc _B	GTAACCATTATAAGCTGC
pEGFP-C2 _F	AACACCCCCATCGGCG
pEGFP-C2 _R	GTAACCATTATAAGCTGC
Notl	TGCGGCCGCA

^aBoldface nucleotides indicate mutated codons.

for 45 s, 50°C for 40 s, and 70°C for 1 min and then a final elongation step of 70°C for 10 min. The PCR product was digested with Ncol and BamHI and ligated into pTrc99A.

pET-NleD, pET-NleD_{E143A'} and pET-NleD_{R203E} were constructed by amplifying *nleD*, *nleD*_{E143A'} and *nleD*_{R203E} from pGFP-NleD, pGFP-NleD_{E143A'} and pGFP-NleD_{R203E} template DNA, respectively, by PCR using the primer pair NleD_F/NleD_R. PCR amplification consisted of an initial denaturation step at 95°C for 10 min, followed by 30 cycles of 94°C for 45 s, 50°C for 40 s, and 70°C for 1 min and then a final elongation step of 70°C for 10 min. The PCR products were digested with EcoRI and Sall and ligated into pET-28a to produce N-terminal 6×His tag fusions to NleD.

pGEX-p38 was constructed by amplifying human *p38* (MAPK14 transcript variant 2) from HeLa cell cDNA using the primer pair $p38_{\rm F}/p38_{\rm R}$. PCR amplification consisted of an initial denaturation step at 98°C for 30 s, followed by 35 cycles of 98°C for 10 s, 63.4°C for 30 s, and 72°C for 30 s and then a final elongation step of 72°C for 5 min. The PCR product was digested with BamHI and Xhol and ligated into pGEX-4T-1 to produce an N-terminal glutathione S-transferase (GST) tag fusion to p38.

EPEC infection. HT-29 cell monolayers were seeded into 24-well tissue culture trays (Corning) at 24 h prior to infection. EPEC derivatives were inoculated into LB broth and grown with shaking at 37°C overnight. On the day of infection, the overnight EPEC cultures were subcultured 1:75 in DMEM with GlutaMAX (Gibco) or RPMI with GlutaMAX (Gibco) and grown statically for 3 h at 37°C with 5% CO₂. Where necessary, cells were induced with 1 mM isopropyl- β -D-thiogalactopyranoside (IPTG) for 30 min prior to infection. Cells were washed twice with phosphate-buffered saline (PBS) and infected with EPEC cultures at an optical density at 600 nm of 0.03 for 2 h.

Transfection. Transfections were carried out in HEK293T cells using the Fugene 6 (Promega) transfection reagent according to the manufacturer's instructions. Cells were seeded into 24-well tissue culture trays (Corning) and transfected with 0.4 μ g of pEGFP-C2 derivatives 24 h later for a period of 18 h.

Dual-luciferase reporter assay. For the AP-1 dual-luciferase assay, HEK293T cells were seeded into 24-well trays (Corning) and cotransfected with derivatives of pEGFP-C2 (0.4 μ g), 0.05 μ g of pRL-TK (Promega), and 0.2 μ g of pAP-1-Luc (Clontech). At approximately 18 h posttransfection, cells were left untreated or stimulated with 25 ng/ml phorbol myristate acetate (PMA; Calbiochem) for 6 h. Firefly and *Renilla* luciferase levels were measured using a dual-luciferase reporter assay system (Promega) in a CLARIOstar microplate reader (BMG Labtech). The expression of firefly luciferase was normalized to *Renilla* luciferase measurements, and luciferase activity was expressed relative to that of unstimulated pEGFP-C2-transfected cells.

Immunoblot analysis. For immunoblot analysis following EPEC infection or transfection, cells were collected and lysed in cold lysis buffer (1% Triton X-100, 50 mM Tris-HCl, pH 7.4, 1 mM EDTA, 150 mM NaCl) with cOmplete protease inhibitor (Roche), 2 mM Na₃VO₄, 10 mM NaF, 1 mM phenylmethylsulfonyl fluoride and incubated on ice for 10 min to complete lysis. The samples were pelleted at 4°C, and the supernatants were added to 4×Bolt lithium dodecyl sulfate sample buffer containing 50 mM dithiothreitol (DTT) (Thermo Fisher), heated to 70°C for 10 min, and resolved on Bolt 4 to 12% bis-Tris Plus gels (Thermo Fisher) by PAGE. Proteins were transferred to nitrocellulose membranes using an iBlot2 gel transfer device (Thermo Fisher) and probed with one of the following primary antibodies: rabbit polyclonal anti-SAPK/JNK (Cell Signaling), rabbit polyclonal anti-p38 MAPK (Cell Signaling), mouse monoclonal anti-HA (Covance), mouse monoclonal anti-GFP (clones 7.1 and 13.1; Roche), and mouse monoclonal anti- β -actin (clone AC-15; Sigma). The antibodies were diluted in Tris-buffered saline (TBS) with 5% bovine serum albumin (BSA) (Sigma) and 0.1% Tween (Sigma). Proteins were detected using anti-rabbit or anti-mouse IgG secondary antibodies conjugated to horseradish peroxidase (PerkinElmer) diluted in TBS with 5% BSA and 0.1% Tween and developed with an enhanced chemiluminescence (ECL) immunoblotting reagent (Amersham). Images were visualized using an MFChemiBis imaging station (DNR Bio-Imaging Systems). At least three biological replicates were performed for all experiments.

In vitro cleavage assay. Plasmids for the expression of 6×His-tagged proteins (pET-NleD, pET-NleD_{E143A}, and pET-NleD_{R203E}) or GST-tagged proteins (pGEX-4T-1 and pGEX-p38) were transformed into *E. coli* BL21 C43(DE3). Overnight LB cultures of BL21 containing the appropriate plasmid were used to inoculate 200 ml LB broth 1:100, in which the cultures were grown at 37°C with shaking to an optical density (A_{600}) of 0.6. The cultures were then induced with 1 mM IPTG and grown for a further 2.5 h at 37°C (for His-NleD constructs and GST) or 18°C (for GST-p38) before being pelleted by centrifugation. Purification of proteins was performed using Novagen His · Bind and GST · Bind kits according to the manufacturer's protocols. Protein concentrations were determined using a bicinchoninic acid (BCA) kit (Thermo Scientific).

In vitro cleavage assays were performed by incubating 2 μ g of purified GST or GST-p38 either alone or with 2 μ g of His-NleD, His-NleD_{E143A}, or His-NleD_{R203E} in buffer [TBS supplemented with 1 mM Tris(2-carboxyethyl)phosphine hydrochloride] for 4 h at 37°C. Sample buffer was then added to the incubation mixtures before they were heated at 70°C for 10 min, resolved by SDS-PAGE, and subsequently transferred to nitrocellulose membranes. The membranes were probed with mouse monoclonal anti-His (clone AD1.1.10; AbD Serotec) or rabbit polyclonal anti-GST (Cell Signaling) primary antibodies and horseradish peroxidase-conjugated anti-mouse or anti-rabbit immunoglobulin secondary antibodies (PerkinElmer) and developed as described above.

For detection of the activity of purified NIeD proteins within HT-29 cell lysates, HT-29 cells were grown in 10-cm dishes for 48 h before being washed with PBS and then collected and resuspended in 800 μ l PBS. Cells were then lysed by passage through a 26-gauge needle 50 times and pelleted, and the supernatant was used for incubations. Forty microliters of the HT-29 cell lysate was incubated alone or with 0.5 μ g of His-NIeD, His-NIeD_{E143AV} or His-NIeD_{R203E} for 4 h at 37°C. Sample buffer was added to the incubation mixtures before they were heated at 70°C for 10 min and immunoblotted as described above.

Detection of IL-6 by ELISA. For analysis of IL-6 secretion, HT-29 cell monolayers were infected for 2 h before 8 to 12 h of incubation with 50 μ g/ml gentamicin with or without 20 ng/ml TNF (Calbiochem, EMD4Biosciences). The HT-29 cell supernatant was collected for subsequent analysis of IL-6 secretion. Analysis of IL-6 secretion was performed using a human IL-6 enzyme-linked immunosorbent assay (ELISA) set (Max Deluxe; BioLegend) as per the manufacturer's instruction. All experiments were performed at least three independent times with samples from three independent infections performed in duplicate.

Mass spectrometry analysis of p38 cleavage fragments. For mass spectrometric analysis, 15 μ g of GST-p38 and 1 μ g of His-NIeD were incubated together or individually at 37°C for 2 h or overnight. Sample buffer was then added to the incubation mixtures before they were heated at 70°C for 10 min and subjected to SDS-PAGE. Polyacrylamide gels were fixed and visualized with Coomassie G-250 according to the protocol of Kang et al. (31).

Tryptic digestion of full-length p38 and cleavage products. Bands of interest were excised from the Coomassie-stained gels and destained in a 50:50 solution of 50 mM NH₄HCO₃-100% ethanol for 20 min at room temperature with shaking at 750 rpm. Destained samples were then washed with 100% ethanol, vacuum dried for 20 min, and rehydrated in 10 mM DTT in 50 mM NH₄HCO₃. Reduction was carried out for 60 min at 56°C with shaking. The reducing buffer was then removed, and the gel bands were washed twice in 100% ethanol for 10 min to remove the residual DTT. Reduced ethanol-washed samples were sequentially alkylated with 55 mM iodoacetamide in 50 mM NH₄HCO₃ in the dark for 45 min at room temperature (RT). Alkylated samples were then washed for 2 rounds with 100% ethanol and vacuum dried. Alkylated samples were then rehydrated with 12 ng/µl trypsin (Promega, Madison, WI) in 40 mM NH₄HCO₃ at 4°C for 1 h. Excess trypsin was removed, and gel pieces were covered in 40 mM NH₄HCO₃ and incubated overnight at 37°C. Peptides were concentrated and desalted using C₁₈ stage tips (32, 33) before analysis by liquid chromatography-mass spectrometry (LC-MS).

Identification of p38 sequence coverage using reversed-phase LC-MS. Purified peptides were resuspended in buffer A* (0.1% trifluoroacetic acid, 2% acetonitrile [MeCN]) and separated using a two-column chromatography setup composed of a PepMap100 C₁₈ trap column (20 mm by 75 μ m) and PepMap C₁₈ analytical columns (500 mm by 75 μ m) (Thermo Scientific). Samples were concentrated onto the trap column at 5 μ l/min for 5 min and infused into an Orbitrap Fusion Lumos Tribrid mass spectrometer (Thermo Scientific) at 300 nl/min via the analytical column using a Dionex Ultimate 3000 ultraperformance liquid chromatograph (Thermo Scientific). A 90-min gradient from 2% buffer B to 28% buffer B to 40% buffer B in the next 5 min was run; it was then

increased to 100% buffer B over a 2-min period, held at 100% buffer B for 2.5 min, and then dropped to 0% buffer B for another 20 min. The Lumos mass spectrometer was operated in a data-dependent mode automatically switching between the acquisition of a single Orbitrap MS scan (120,000 resolution) every 3 s and ion-trap collision-induced dissociation MS-MS, Orbitrap higher-energy collision-induced dissociation MS-MS, and Orbitrap electron-transfer/higher-energy collision dissociation for each selected precursor (maximum fill time, 100 ms; automatic gain control, 5×10^4 with a resolution of 30,000 for Orbitrap MS-MS scans).

Data analysis. MaxQuant (v1.5.3.1) software (34) was used for identification of p38 cleavage products. A database search of the sequences against a GST-p38 protein sequence supplemented with data from an *E. coli* K-12 database (from UniProt, 16 March 2015) was carried out with the following search parameters: carbamidomethylation of cysteine as a fixed modification; oxidation of methionine, deamidation (NQ), and acetylation of protein N termini as variable modifications; and semitrypsin cleavage specificity. The precursor mass tolerance was set to 20 ppm for the first search and 10 ppm for the main search, and a maximum false discovery rate of 1.0% was set for protein and peptide identifications. To enhance the identification of peptides between full-length GST-p38 and cleavage products, the Match between Runs option was enabled with a precursor match window set to 2 min and an alignment window of 10 min. The resulting protein group output was processed within the Perseus (v1.4.0.6) (35) analysis environment to remove reverse matches and common protein contaminants prior to analysis.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/ IAI.00620-16.

TEXT S1, PDF file, 1.8 MB.

ACKNOWLEDGMENTS

This work was supported by grants to E.L.H. from the Australian National Health and Medical Research Council (NHMRC) (APP1100609). K.C. is supported by a Deutsche Forschungsgemeinschaft (DFG) research fellowship. J.S.P. is the recipient of an NHMRC early career research fellowship. C.G. is the recipient of an Australian Postgraduate Award (APA). T.W.F.L. is a recipient of a University of Melbourne International Research Scholarship (MIRS). N.E.S. is the recipient of an NHMRC early career research fellowship.

REFERENCES

- Nadler C, Baruch K, Kobi S, Mills E, Haviv G, Farago M, Alkalay I, Bartfeld S, Meyer T, Ben-Neriah Y, Rosenshine I. 2010. The type III secretion effector NIeE inhibits NF-κB activation. PLoS Pathog 6:e1000743. https:// doi.org/10.1371/journal.ppat.1000743.
- Newton HJ, Pearson JS, Badea L, Kelly M, Lucas M, Holloway G, Wagstaff KM, Dunstone MA, Sloan J, Whisstock JC, Kaper JB, Robins-Browne RM, Jans DA, Frankel G, Phillips AD, Coulson BS, Hartland EL. 2010. The type III effectors NIeE and NIeB from enteropathogenic *E. coli* and OspZ from *Shigella* block nuclear translocation of NF-κB p65. PLoS Pathog 6:e1000898. https://doi.org/10.1371/journal.ppat.1000898.
- Zhang L, Ding X, Cui J, Xu H, Chen J, Gong Y-N, Hu L, Zhou Y, Ge J, Lu Q, Liu L, Chen S, Shao F. 2011. Cysteine methylation disrupts ubiquitinchain sensing in NF-κB activation. Nature 481:204–208. https://doi.org/ 10.1038/nature10690.
- Li S, Zhang L, Yao Q, Li L, Dong N, Rong J, Gao W, Ding X, Sun L, Chen X, Chen S, Shao F. 2013. Pathogen blocks host death receptor signalling by arginine GlcNAcylation of death domains. Nature 501:242–246. https://doi.org/10.1038/nature12436.
- Pearson JS, Giogha C, Ong SY, Kennedy CL, Kelly M, Robinson KS, Lung TWF, Mansell A, Riedmaier P, Oates CVL, Zaid A, Mühlen S, Crepin VF, Marches O, Ang C-S, Williamson NA, O'Reilly LA, Bankovacki A, Nachbur U, Infusini G, Webb AI, Silke J, Strasser A, Frankel G, Hartland EL. 2013. A type III effector antagonizes death receptor signalling during bacterial gut infection. Nature 501:247–251. https://doi.org/10.1038/nature12524.
- Deng W, Puente JL, Gruenheid S, Li Y, Vallance BA, Vazquez A, Barba J, Ibarra JA, O'Donnell P, Metalnikov P, Ashman K, Lee S, Goode D, Pawson T, Finlay BB. 2004. Dissecting virulence: systematic and functional analyses of a pathogenicity island. Proc Natl Acad Sci U S A 101:3597–3602. https://doi.org/10.1073/pnas.0400326101.
- Miyoshi S-I, Shinoda S. 2000. Microbial metalloproteases and pathogenesis. Microbes Infect 2:91–98. https://doi.org/10.1016/S1286 -4579(00)00280-X.

- 8. Potempa J, Pike R. 2005. Bacterial peptidases. Contrib Microbiol 12: 132–180.
- Giogha C, Wong Fok Lung T, Mühlen S, Pearson JS, Hartland EL. 2015. Substrate recognition by the zinc metalloprotease effector NleC from enteropathogenic *Escherichia coli*. Cell Microbiol 17:1766–1768. https:// doi.org/10.1111/cmi.12469.
- Hodgson A, Wier E, Fu K, Sun X, Yu H, Zheng W, Sham H, Johnson K, Bailey S, Vallance B, Wan F. 2015. Metalloprotease NleC suppresses host NF-kappaB/inflammatory responses by cleaving p65 and interfering with the p65/RPS3 interaction. PLoS Pathog 10:e1004705. https://doi.org/ 10.1371/journal.ppat.1004705.
- Mühlen S, Ruchaud-Sparagano M-H, Kenny B. 2011. Proteasomeindependent degradation of canonical NFκB complex components by the NIeC protein of pathogenic *Escherichia coli*. J Biol Chem 286: 5100–5107. https://doi.org/10.1074/jbc.M110.172254.
- Pearson JS, Riedmaier P, Marches O, Frankel G, Hartland EL. 2011. A type III effector protease NIeC from enteropathogenic *Escherichia coli* targets NF-κB for degradation. Mol Microbiol 80:219–230. https://doi.org/ 10.1111/j.1365-2958.2011.07568.x.
- Sham HP, Shames SR, Croxen MA, Ma C, Chan JM, Khan MA, Wickham M, Deng W, Finlay BB, Vallance BA. 2011. Attaching and effacing bacterial effector NIeC suppresses epithelial inflammatory responses by inhibiting NF-κB and p38 mitogen-activated protein kinase activation. Infect Immun 79:3552–3562. https://doi.org/10.1128/IAI.05033-11.
- Yen H, Ooka T, Iguchi A, Hayashi T, Sugimoto N, Tobe T. 2010. NIeC, a type III secretion protease, compromises NF-κB activation by targeting p65/RelA. PLoS Pathog 6:e1001231. https://doi.org/10.1371/journal .ppat.1001231.
- Baruch K, Gur-Arie L, Nadler C, Koby S, Yerushalmi G, Ben-Neriah Y, Yogev O, Shaulian E, Guttman C, Zarivach R, Rosenshine I. 2011. Metalloprotease type III effectors that specifically cleave JNK and NF-κB. EMBO J 30:221–231. https://doi.org/10.1038/emboj.2010.297.

- Zarubin T, Han J. 2005. Activation and signaling of the p38 MAP kinase pathway. Cell Res 15:11–18. https://doi.org/10.1038/sj.cr.7290257.
- 17. Morrison DK. 2012. MAP kinase pathways. Cold Spring Harbor Perspect Biol 4:a011254. https://doi.org/10.1101/cshperspect.a011254.
- Karin M. 1995. The regulation of AP-1 activity by mitogen-activated protein kinases. J Biol Chem 270:16483–16486. https://doi.org/10.1074/ jbc.270.28.16483.
- Guan Z, Buckman SY, Pentland AP, Templeton DJ, Morrison AR. 1998. Induction of cyclooxygenase-2 by the activated MEKK1 → SEK1/MKK4 → p38 mitogen-activated protein kinase pathway. J Biol Chem 273: 12901–12908. https://doi.org/10.1074/jbc.273.21.12901.
- Kelly M, Hart E, Mundy R, Marchès O, Wiles S, Badea L, Luck S, Tauschek M, Frankel G, Robins-Browne RM, Hartland EL. 2006. Essential role of the type III secretion system effector NIeB in colonization of mice by *Citrobacter rodentium*. Infect Immun 74:2328–2337. https://doi.org/10.1128/ IAI.74.4.2328-2337.2006.
- Marchès O, Wiles S, Dziva F, La Ragione RM, Schüller S, Best A, Phillips AD, Hartland EL, Woodward MJ, Stevens MP, Frankel G. 2005. Characterization of two non-locus of enterocyte effacement-encoded type III-translocated effectors, NIeC and NIeD, in attaching and effacing pathogens. Infect Immun 73:8411–8417. https://doi.org/10.1128/IAI .73.12.8411-8417.2005.
- 22. Li W, Liu Y, Sheng X, Yin P, Hu F, Liu Y, Chen C, Li Q, Yan C, Wang J. 2014. Structure and mechanism of a type III secretion protease, NIeC. Acta Crystallogr D Biol Crystallogr 70:40–47. https://doi.org/10.1107/S13 99004713024619.
- Kang S-S, Woo SS, Im J, Yang JS, Yun C-H, Ju HR, Son CG, Moon E-Y, Han SH. 2007. Human placenta promotes IL-8 expression through activation of JNK/SAPK and transcription factors NF-κB and AP-1 in PMAdifferentiated THP-1 cells. Int Immunopharmacol 7:1488–1495. https:// doi.org/10.1016/j.intimp.2007.07.011.
- Roger T, Out AT, Mukaida N, Matsushima K, Jansen MH, Lutter R. 1998. Enhanced AP-1 and NF-κB activities and stability of interleukin 8 (IL-8) transcripts are implicated in IL-8 mRNA superinduction in lung epithelial H292 cells. Biochem J 330:429–435. https://doi.org/10.1042/bj3300429.
- Hauf N, Charkraborty T. 2003. Suppression of NF-kappaB activation and proinflammatory cytokine expression by Shiga toxin-producing *Escherichia coli*. J Immunol 170:2074–2082. https://doi.org/10.4049/jimmunol .170.4.2074.
- Ruchaud-Sparagano M, Maresca M, Kenny B. 2007. Enteropathogenic *Escherichia coli* (EPEC) inactivate innate immune responses prior to compromising epithelial barrier function. Cell Microbiol 9:1909–1921. https://doi.org/10.1111/j.1462-5822.2007.00923.x.
- 27. Angel P, Karin M. 1991. The role of Jun, Fos and the AP-1 complex in cell-proliferation and transformation. Biochim Biophys Acta 1072: 129–157. https://doi.org/10.1016/0304-419X(91)90011-9.
- 28. Hershko DD, Robb BW, Luo G, Hasselgren P-O. 2002. Multiple transcrip-

tion factors regulating the IL-6 gene are activated by cAMP in cultured Caco-2 cells. Am J Physiol Regul Integr Comp Physiol 283:R1140–R1148. https://doi.org/10.1152/ajpregu.00161.2002.

- Dann SM, Spehlmann ME, Hammond DC, limura M, Hase K, Choi LJ, Hanson E, Eckmann L. 2008. IL-6-dependent mucosal protection prevents establishment of a microbial niche for attaching/effacing lesion forming enteric bacterial pathogens. J Immunol 180:6816–6826. https:// doi.org/10.4049/jimmunol.180.10.6816.
- Royan SV, Jones RM, Koutsouris A, Roxas JL, Falzari K, Weflen AW, Kim A, Bellmeyer A, Turner JR, Neish AS, Rhee K-J, Viswanathan VK, Hecht GA. 2010. Enteropathogenic *E. coli* non-LEE encoded effectors NleH1 and NleH2 attenuate NF-κB activation. Mol Microbiol 78:1232–1245. https:// doi.org/10.1111/j.1365-2958.2010.07400.x.
- Kang D, Gho Y, Suh M, Kang C. 2002. Highly sensitive and fast protein detection with Coomassie brilliant blue in sodium dodecyl sulfatepolyacrylamide gel electrophoresis. Bull Korean Chem Soc 23: 1511–1513. https://doi.org/10.5012/bkcs.2002.23.11.1511.
- Ishihama Y, Rappsilber J, Mann M. 2006. Modular stop and go extraction tips with stacked disks for parallel and multidimensional peptide fractionation in proteomics. J Proteome Res 5:988–994. https://doi.org/ 10.1021/pr050385g.
- Rappsilber J, Mann M, Ishihama Y. 2007. Protocol for micro-purification, enrichment, pre-fractionation and storage of peptides for proteomics using StageTips. Nat Protoc 2:1896–1906. https://doi.org/10.1038/nprot .2007.261.
- Cox J, Mann M. 2008. MaxQuant enables high peptide identification rates, individualized p.p.b.-range mass accuracies and proteome-wide protein quantification. Nat Biotechnol 26:1367–1372. https://doi.org/ 10.1038/nbt.1511.
- Tyanova S, Temu T, Sinitcyn P, Carlson A, Hein MY, Geiger T, Mann M, Cox J. 2016. The Perseus computational platform for comprehensive analysis of (prote)omics data. Nat Methods 13:731–740. https://doi.org/ 10.1038/nmeth.3901.
- Gouet P, Courcelle E, Stuart DI, Metoz F. 1999. ESPript: analysis of multiple sequence alignments in PostScript. Bioinformatics 15:305–308. https://doi.org/10.1093/bioinformatics/15.4.305.
- Levine MM, Nalin DR, Hornick RB, Bergquist EJ, Waterman DH, Young CR, Sotman S, Rowe B. 1978. *Escherichia coli* strains that cause diarrhoea but do not produce heat-labile or heat-stable enterotoxins and are noninvasive. Lancet 311:1119–1122. https://doi.org/10.1016/S0140-6736(78) 90299-4.
- Zurawski DV, Mumy KL, Badea L, Prentice JA, Hartland EL, McCormick BA, Maurelli AT. 2008. The NIeE/OspZ family of effector proteins is required for polymorphonuclear transepithelial migration, a characteristic shared by enteropathogenic *Escherichia coli* and *Shigella flexneri* infections. Infect Immun 76:369–379. https://doi.org/10.1128/IAI.00684-07.