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ORIGINAl RESEARCH

Attaching-and-Effacing Pathogens Exploit Junction Regulatory Activities of N-WASP and SNX9 to Disrupt the Intestinal Barrier

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SUMMARY
During enteric infection, the attaching-and-effacing pathogen virulence factor early secreted antigenic target-6 (ESX)-1 secretion-associated protein F (EspF) targets host Neural Wiskott-Aldrich Syndrome protein and sorting nexin 9 to promote junctional disruption and intestinal barrier loss. On this basis, we propose a novel role for the actin cytoskeletal regulatory protein Neural Wiskott-Aldrich Syndrome protein in controlling intestinal epithelial apical junction complex stability.

BACKGROUND & AIMS: Neural Wiskott-Aldrich Syndrome protein (N-WASP) is a key regulator of the actin cytoskeleton in epithelial tissues and is poised to mediate cytoskeletal-dependent aspects of apical junction complex (AJC) homeostasis. Attaching-and-effacing (AE) pathogens disrupt this homeostasis through translocation of the effector molecule early secreted antigenic target-6 (ESX)-1 secretion-associated protein F (EspF). Although the mechanisms underlying AJC disruption by EspF are unknown, EspF contains putative binding sites for N-WASP and the endocytic regulator sorting nexin 9 (SNX9). We hypothesized that N-WASP regulates AJC integrity and AE pathogens use EspF to induce junction disassembly through an N-WASP- and SNX9-dependent pathway.

METHODS: We analyzed mice with intestine-specific N-WASP deletion and generated cell lines with N-WASP and SNX9 depletion for dynamic functional assays. We generated EPEC and Citrobacter rodentium strains complemented with EspF bearing point mutations abolishing N-WASP and SNX9 binding to investigate the requirement for these interactions.

RESULTS: Mice lacking N-WASP in the intestinal epithelium showed spontaneously increased permeability, abnormal AJC morphology, and mislocalization of occludin. N-WASP depletion in epithelial cell lines led to impaired assembly and disassembly of tight junctions in response to changes in extracellular calcium. Cells lacking N-WASP or SNX9 supported actin pedestals and type III secretion, but were resistant to EPEC-induced AJC disassembly and loss of transepithelial resistance. We found that during in vivo infection with AE pathogens, EspF must bind both N-WASP and SNX9 to disrupt AJCs and induce intestinal barrier dysfunction.

CONCLUSIONS: Overall, these studies show that N-WASP critically regulates AJC homeostasis, and the AE pathogen...
effector EspF specifically exploits both N-WASP and SNX9 to disrupt intestinal barrier integrity during infection. (Cell Mol Gastroenterol Hepatol 2018;5:273–288; https://doi.org/10.1016/j.jcmgh.2017.11.015)

Keywords: N-WASP; Cytoskeleton; Junction Regulation; EspF.

See editorial on page 420.

The apical junction complex (AJC), consisting of the tight junction (TJ) and adherens junction (AJs), is a multiprotein organelle that is intimately linked to dynamic networks of apical actin,\textsuperscript{1–3} which, by simultaneously lending the AJC both stability and plasticity, enable it to control intestinal permeability. Neural Wiskott-Aldrich Syndrome protein (N-WASP) is a ubiquitously expressed protein that interacts with and activates the actin-related protein (Arp)2/3 complex, leading to robust polymerization of actin filaments. N-WASP has been shown to stabilize newly formed actin filaments and facilitate their incorporation into perijunctional rings critical for E-cadherin–dependent AJ stability.\textsuperscript{4} There is also indirect evidence of a role for N-WASP in regulating both the delivery and removal of key junction proteins. A role for N-WASP in delivery of AJC proteins was suggested by studies in which chemical inhibition or deletion of N-WASP in epithelial cell lines was associated with defective delivery of occludin,\textsuperscript{5} E-cadherin, and F-actin\textsuperscript{6} to nascent AJCs. Evidence for N-WASP–mediated removal of AJ proteins comes from studies of Drosophila epithelium lacking Cell division cycle (Cdc)42, N-WASP, or Arp3. In these studies, genetic depletion of any member of the Cdc42/N-WASP/Arp3 pathway in fly epithelium led to defective internalization of E-cadherin and destabilization of the AJC.\textsuperscript{7,8}

As the major regulator of intestinal permeability, the AJC is also the target of a number of bacterial pathogens, including enteropathogenic Escherichia coli (EPEC) and enterohemorrhagic E coli (EHEC), and the closely related mouse pathogen Citrobacter rodentium.\textsuperscript{9,10} These pathogens use type III secretion to inject effector proteins that modulate a variety of host cellular processes. The first of these translocated proteins, the translocated intimin receptor (Tir), activates N-WASP, which results in localized actin polymerization and the formation of actin-rich pedestals and destruction of adjacent microvilli, lending this class of bacteria their designation as attaching-and-effacing (AE) pathogens.\textsuperscript{11}

EPEC, EHEC, and C. rodentium also disrupt intercellular junctions and induce intestinal barrier dysfunction.\textsuperscript{12,13} and bacterial effector protein EspF, common to all 3 pathogens, has been shown to be required for AE pathogen-induced junction disruption.\textsuperscript{14} Although the mechanism of EspF-mediated junction disruption remains unknown, EspF contains N-WASP binding sites and is capable of directly stimulating the actin polymerizing activity of N-WASP.\textsuperscript{12} In addition to binding N-WASP, EspF also contains a nonoverlapping binding site for sorting nexin 9 (SNX9).\textsuperscript{12} Sorting nexins are involved in diverse endocytic vesicle sorting activities, and the specific functions of SNX9 within the epithelium are largely unknown.\textsuperscript{13,14} Direct binding of EspF to SNX9 alone appears to be dispensable for mediating EspF-dependent TJ disruption,\textsuperscript{12} although a role for EspF binding to N-WASP in mediating TJ disruption has not been reported.

We sought to evaluate the role that N-WASP, as a key regulator of actin dynamics, plays in maintaining AJC homeostasis, and to test whether AE pathogens, through EspF, exploit N-WASP–dependent pathways to disrupt AJC function. We first show in vivo and in vitro that N-WASP regulates AJC homeostasis, and is required for efficient TJ assembly and disassembly. Moreover, we identify a novel pathway whereby the bacterial effector EspF disrupts tight junctions through exploiting an otherwise homeostatic function of both N-WASP and SNX9 in regulating removal of TJ proteins from the AJC.

Materials and Methods

Mice

Mixed strain (Friend Virus B [FVB]/129) mice bearing a conditional Wasl knockout (KO) allele (Wasl\textsuperscript{fl/fl}fl/fl) were crossed with villin-Cre transgenic mice\textsuperscript{15} to generate Wasl\textsuperscript{fl/fl}fl/fl \textsuperscript{tg}villin-Cre mice. Littermate Wasl\textsuperscript{fl/fl}fl/fl \textsuperscript{tg}villin-Cre mice were used as controls in each experiment. Intestinal N-WASP KO (iNWKO) breeder mice and offspring were housed at either the Center for Comparative Medicine at the Massachusetts General Hospital or the Boston Children’s Hospital animal facility. This study protocol was conducted in accordance with the Animal Research: Reporting of In Vivo Experiments (ARRIVE) guidelines and was approved by the animal ethics committees of both Massachusetts General Hospital and Boston Children’s Hospital.

Western Blot

For Western blot, intestinal epithelial cells were isolated from 8-week-old littermate control or iNWKO mice by EDTA dissociation as previously described,\textsuperscript{16} and lysed with ice-cold buffer (150 mmol/L NaCl, 50 mmol/L Tris-HCl, pH 7.4, 1% Triton X-100) by 15 minutes at 4°C. Total lysates were centrifuged at 13,000g for 15 minutes at 4°C and the supernatant (Triton-insoluble cytoplasmic fraction) was collected. The pellet was resuspended in lysis buffer containing 1% sodium dodecyl sulfate and centrifuged, and the supernatant (Triton-insoluble...
membrane/cytoskeletal fraction) was collected. To assess short hairpin RNA (shRNA) knockdown in Caco-2 cells, monolayers were grown to approximately 90% confluence then lysed in ice-cold RIPA buffer (50 mmol/L Tris, 150 mmol/L NaCl, 0.1% sodium dodecyl sulfate, 0.5% sodium deoxycholate, 1% NP-40, protease inhibitors). Lysates were separated by sodium dodecyl sulfate gel electrophoresis, and transferred onto nitrocellulose membrane. Immunoblotting was performed with rabbit polyclonal anti-N-WASP (1:2000) (kindly provided by Dr Marc Kirshner), or rabbit polyclonal anti-SNX9 antibody (1:500) (Cell Signaling Technology, Danvers, MA), followed by horseradish-peroxidase-conjugated goat anti-rabbit secondary antibody (1:3000; Cell Signaling), and visualized using enhanced chemiluminescence (GE Healthcare Life Sciences, Piscataway, NJ).

**In Vivo Assessment of Intestinal Permeability**

The fluorescein isothiocyanate (FITC)-dextran assay was performed as previously described.15 In brief, mice were administered 60 mg/100 g body weight 4 kilodaltons FITC-conjugated dextran (Sigma-Aldrich, St. Louis, MO) by oral gavage. Four hours later, whole blood was obtained by cardiac puncture in anesthetized animals and incubated on ice for 1 hour, followed by centrifugation at 13,000 rpm for 5 minutes to collect serum for assessment by direct fluorimetry.

**Polarized Cell Culture**

Caco-2 brush border expressing (BBe) cells were obtained from the epithelial cell biology core at the Harvard Digestive Diseases Center. Cells were maintained in a humidified atmosphere of 5% CO\textsubscript{2} at 37°C, grown in Dulbecco’s modified Eagle medium (Gibco Invitrogen, Paisley, UK) supplemented with 10% fetal bovine serum, 1% nonessential amino acids, 1% penicillin (1000 U/mL), and 1% streptomycin (1000 μg/mL). For permeability and infection studies, the cells were seeded on rat-tail collagen-coated 12-mm polycarbonate permeable Transwell cell culture inserts (0.4-μm pore size; Costar Corning, Inc, Corning, NY) at an initial density of 1×10\textsuperscript{4} cells/well. Culture medium was replaced every 2–3 days as needed, and cells were maintained for up to 31 days to form a differentiated monolayer. Cell confluence was confirmed by microscopic observance and measurement of transepithelial electrical resistance (TER) using a 12-well, 12-mm Endohm voltmeter (World Precision Instruments, Sarasota, FL) as previously described.18–20

**shRNA Knockdown**

N-WASP and SNX9 knockdown were achieved by transducing Caco-2 cells with N-WASP shRNA (antisense sequence: GCTGGAGATACCTTGAAG), SNX9 shRNA (antisense sequence: TACTCGTGATTACATTCCATC), or scrambled control in a plKO.1-based lentiviral vector (Open Biosystems, Pittsburgh, PA). After transduction, cells were selected and maintained in 10 μg/mL puromycin. N-WASP and SNX9 protein levels were determined after selection by Western blot and detection with rabbit polyclonal anti-N-WASP antibody (Cell Signaling Technology) and monoclonal mouse anti-SNX9 antibody (Abcam, Cambridge, MA). Greater than 90% knockdown was confirmed by comparison of the optical density of N-WASP and SNX9 bands relative to tubulin loading controls in ImageJ software (National Institutes of Health, Bethesda, MD).

**Line Scan and Kymographic Analysis**

Raw imaging stacks were analyzed using ImageJ software. A line 60-pixels long was drawn perpendicular to the junctional actin and a kymograph was created from the image sequence using the Multiple Kymograph and surface plot analysis functions of ImageJ software. For the line scan analysis, histogram data were saved as a text file and imported into Excel (Microsoft, Redmond, WA).

**Generation and Passaging of Intestinal Organoids**

Small intestinal crypts were isolated from WT and iNKO mice and subsequently cultured to grow organoids according to the methods described by Miyoshi and Stappenbeck.21 In brief, intestinal tissues were opened longitudinally and villi were removed by scraping with a glass coverslip before mincing and incubation in 2 mmol/L EDTA with gentle rocking. Supernatants obtained from successive incubations were examined to identify crypt-enriched fractions, which subsequently were maintained in basal culture medium consisting of advanced Dulbecco’s modified Eagle medium/F12 supplemented with penicillin (100 U/mL) and streptomycin (100 mg/mL) (Life Technologies, Carlsbad, CA), GlutaMAX (Gibco, Gaithersburg, MD), and 10 mmol/L HEPES. Crypts were resuspended in Matrigel (Corning Life Sciences, Tewksbury, MA) at a density of 2000 crypts/50 μL and allowed to form hemispherical droplets by inversion at 37°C. After solidification of Matrigel droplets, 500 μL complete culture medium consisting of basal culture medium supplemented with B27 supplement (Life Technologies), 1× N2 supplement (Life Technologies), 1 mmol/L N-acetyl-L-cysteine (Sigma-Aldrich), and 50% L-Wnt3A/R-spondin-3/noggin (WRN) conditioned media22 was obtained from the Harvard Digestive Diseases Core. Organoids were passaged after 7–14 days in culture by gently breaking up the Matrigel by pipetting and resuspending organoids in 1 mL basal culture medium followed by gently disruption using a fire-polished Pasteur pipette. Organoids were washed with 10 mL basal culture medium to remove single and dead cells and resuspended in 50 μL Matrigel droplets, which were allowed to solidify before adding complete organoid media.

**Antibodies and Immunofluorescence Microscopy**

Immunostaining was performed as previously described.22 Briefly, for mouse intestinal tissue, 5-μm frozen sections were collected on coated slides, fixed in fresh 4% paraformaldehyde, and washed 3 times with phosphate-buffered saline (PBS) and permeabilized with 0.1% Triton-X 100. After incubation with anti–E-cadherin (Cell Signaling Technology), anti-occludin (Abcam), or anti–zonula occludens-1 (ZO-1) (Abcam) primary antibodies, all at 1:400, slides were washed and incubated with Alexa 594–conjugated goat anti-rabbit or Alexa 488–conjugated goat anti-rabbit secondary antibodies and DAPI (Life Technologies, Grand Island, NY). Standard epifluorescence microscopy was performed using an
AX70 upright microscope (Olympus, Tokyo, Japan). For polarized Caco-2 cells, the Transwell filters were cut out using a scalpel and immediately washed with PBS and fixed in 4% paraformaldehyde. Further staining was performed in 6- or 12-well plates as described for frozen sections. Confocal images were obtained with a Zeiss (Jena, Germany) Axiovert 200M microscope.

**Transmission Electron Microscopy**

Electron microscopy (EM) was performed in the Microscopy Core in the Program for Membrane Biology at Massachusetts General Hospital. Intestinal tissues were fixed in 2.0% glutaraldehyde in 0.1 mol/L sodium cacodylate buffer (Electron Microscopy Sciences, Hatfield, PA) overnight at 4°C. The following day, samples were postfixed in 1.0% osmium tetroxide for 1 hour at room temperature, rinsed in buffer, and dehydrated through a graded series of ethanol. Samples then were infiltrated with Epon resin (Ted Pella, Redding, CA) in a 1:1 solution of Epon:ethanol overnight. The following day, they were placed in fresh Epon for several hours and then embedded in Epon overnight at 60°C. Thin sections were cut on a Leica (Wetzlar, Germany) EM UC7 ultramicrotome, collected on formvar-coated grids, stained with uranyl acetate and lead citrate, and examined in a JEOL (Peabody, MA) JEM 1011 transmission electron microscope at 80 kV. Images were collected at direct magnification of 30,000× using an AMT digital imaging system (Advanced Microscopy Techniques, Danvers, MA).

**Calcium Switch**

The calcium-switch technique has been described previously. Briefly, for calcium depletion, once monolayers developed maximal TER, normal media was replaced with calcium-free Minimum Eagle Spinner Modification, which had been prewarmed to 37°C. Changes in TER were measured every 10 minutes for 1 hour. At 0-, 15-, and 30-minute time points, wells were harvested, washed in PBS, and fixed in fresh 4% paraformaldehyde for immunostaining as described earlier. For calcium repletion, fully confluent monolayers were cultured overnight in Minimum Essential Medium Eagle Spinner Modification, and the media replaced with complete Dulbecco's modified Eagle medium containing calcium. Serial TER measurements were obtained every 15 minutes for the first hour after calcium repletion, then every 30 minutes for 8 hours. At 0, 3, and 8 hours after calcium repletion, wells were harvested for junction characterization.

**Bacteria and Infections**

For in vitro infections, wild-type EPEC strain E2348/69 (EPEC 0127:H7) or E2348/69 (EspF) complemented with N-WASP– or SNX9-binding–defective EspF constructs (described later) were grown in Luria–Bertani broth with shaking at 37°C for 12 hours, and then maintained without shaking for 8 hours at 37°C to enhance type III secretion. For in vitro infections, bacterial density was assessed by photometric measurement of optical density (optical density of 600), and diluted to a final multiplicity of infection of 100 bacteria per cell in prewarmed Caco-2 medium without antibiotics. Bacteria were allowed to adhere for up to 12 hours, and TER measurements were obtained at the indicated time points in triplicate wells. At 4 hours after infection, Transwell filters were removed and fixed in fresh 4% PFA for actin pedestal assessment and immunostaining. For in vivo infections, wild-type *C. rodentium* strain David B. Schauer (DBS) 100, or DBS100ΔespF (EM282) strains complemented with N-WASP– or SNX9-binding–defective EspF constructs were similarly grown overnight. Mice were infected by a single oral gavage of 1 × 10⁹ bacteria. EspF-deficient EPEC strain UMD874 (E2348/69ΔespF) has been described previously. The EspF-deficient *C. rodentium* strain EM282 (DBS100ΔespF) was generated by using lambda red recombination. Linear polymerase chain reaction (PCR) products containing the Zeocin cassette and its promoter region with flanking tails homologous to *C. rodentium* espF were amplified from pDONR/Zeo. DBS100 containing pKD46, which carries the lambda red recombinase, was induced with arabinose to allow for expression of the lambda red recombinase. The linear PCR fragment then was electroproporated into DBS100. Bacteria were recovered and plated on Zeocin to select for recombinants and deletions were confirmed by PCR.

**Generation of Modular EspF and N-WASP– and SNX9-Binding–Defective EspF**

The modular version of EspF (pTM007) was generated by inverse PCR of pJN61 (a pTrc99A-based vector containing FLAG-tagged espF) using the forward (GCCACGAATTCCCAACCGCTGACACCACTGCAGGCT) and reverse (CCGGATCCGCCAATGCTACGACGCTTACAGTCTCAGACGCTT) primers followed by E. coli RY13 I (EcoRI) digestion and ligation. This yielded a version of EspF that contained the N-terminal 73 amino acids, in addition to the third proline-rich region (PRR) and C-terminus (pTM001). The second PRR was amplified from pJN61 using the forward (GCCACGAAAATGCTACGACGCTTACAGTCTCAGGCT) primers followed by E. coli RY13 I (EcoRI) digestion and ligation. The linear PCR fragment then was electroproporated into DBS100. Bacteria were recovered and plated on Zeocin to select for recombinants and deletions were confirmed by PCR.

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GGCATAAAAAGATCATGCAGCTGCCTATGAACTATCG and reverse (CGATAGTTCATAGGCAGCTGCATGATCTTTTAATG CCTGTGCAATGG) primers for pTM003. The SNX9 binding site was mutated using the forward (CTCGGGGAATTCCAACCGATCAGGCACCACCGCCAC) and reverse (GGTGGCGGTCGGTTGGGAATTCCCCGAG) primers for pTM001, and forward (CCAACTCCGGGCCAACCAGATCGGCCACC) and reverse (GGTGGCGGCGGTGGCGGA TCGTTGGTCCGGAGTTGG) primers for pTM003.

Data
All authors had access to the study data and reviewed and approved the final manuscript.

Results
Mice With Intestine-Specific N-WASP Deletion Have Spontaneous AJC Defects and Increased Intestinal Permeability
Germline deletion of N-WASP in mice results in early embryonic lethality. To investigate the intestinal junction regulatory functions of N-WASP in vivo, conditionally targeted Wasl^flx/flx^ mice were bred with mice expressing Cre recombinase under the intestine-specific promoter villin (tg^vil-Cre). INWKO mice were viable and fertile, although tended to have lower overall average weights compared with littermate controls. H&E staining showed normal crypt-villus architecture with all epithelial cell lineages present and no evidence of spontaneous inflammation.

Electron microscopic analysis showed that iNWKO mice displayed abnormal microvilli and decreased perijunctional actin (Figure 1A). Immunofluorescent staining showed normal junction localization of E-cadherin and ZO-1, but weak junction localization and more diffuse cytoplasmic staining of occludin (Figure 1B) despite comparable levels of total occludin protein in membrane and cytosolic fractions (Figure 1C). Although the pixel intensity by line scan at the tight junction appeared very similar between WT and N-WASP knockdown (NWKD) cells, the intensity of the occludin signal decreased more quickly as one moved away from the junction toward the cytoplasm in the WT cells

Figure 1. Intestine-specific deletion of N-WASP leads to abnormalities in AJC morphology and function in vivo. (A) Electron microscopic examination of small intestine iNWKO mice showing a lack of perijunctional actin (arrow) normally present in WT ileum (asterisk). (B) Immunofluorescence staining of E-cadherin (top), ZO-1 (middle), and occludin (bottom) in WT and iNWKO mice. (C) Occludin Western blot was similar between intestinal lysates from WT and iNWKO mice. (D) Small intestinal organoids generated from WT or iNWKO mice show normal patterns of E-cadherin, ZO-1, Arp2, and myosin light chain staining; organoids lacking N-WASP show decreased junction localization of occludin. (E) iNWKO mice showed significantly increased intestinal permeability by FITC-labeled dextran. Scale bars: (A) 500 nm; (B) 40 μm. I, insoluble; RLU, relative light units; S, soluble.
compared with the NWKD cells, which reflects less discrete junctional staining in the absence of N-WASP. Decreased junction localization of occludin also was observed in N-WASP-deficient small intestinal organoids, whereas localization of other AJC-related proteins, including E-cadherin and ZO-1, as well as Arp2 and myosin light chain (MLC), were similar to WT controls (Figure 1D). To determine if abnormal junction localization of occludin observed in iNWKO mice and organoids correlated with impaired intestinal barrier function, we measured intestinal permeability and found spontaneously increased intestinal permeability to FITC-dextran in iNWKO mice when compared with littermate controls (Figure 1E). There were no significant differences in subcellular localization of Arp family members, including Arp2 and Arp3 (Figure 2A–C), or other AJC-related molecules including myosin light chain (Figure 2A–C) and claudin-1, -2, and -5 (Figure 2D).

**N-WASP Regulates Epithelial Permeability In Vitro**

To investigate the mechanistic role(s) that N-WASP plays in regulating TJ dynamics, we generated N-WASP-deficient Caco-2 cells (Figure 3A). When cultured on permeable Transwell filters, NWKD cells polarized normally and showed apical microvilli marked by phalloidin (Figure 3B, left) and well-defined junction complexes marked by E-cadherin (Figure 3B, right). In contrast to E-cadherin and ZO-1 (Figure 3C, top and bottom), the junctional localization of occludin was less distinct with more cytoplasmic staining in NWKD cells (Figure 3C, middle). By using line scan analysis, we found no quantitative differences in E-cadherin or ZO-1 localization at the AJC between WT and NWKD cells (Figure 3D, top and bottom). In contrast, the occludin signal was diminished at intercellular junctions in NWKD cells when compared with WT cells and was more pronounced in the cytoplasm (Figure 3D, middle).

To next assess whether the aberrant TJ morphology correlated with altered barrier function, we measured the development of TER over the course of 30 days after seeding on permeable Transwell filters and found that the maximal TER achieved by fully polarized NWKD cells was ultimately 25% less than cells expressing normal levels of N-WASP (Figure 3E). Of note, NWKD cells developed TER more rapidly around days 12–13, which may reflect slightly

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**Figure 2. N-WASP depletion does not affect Arp2/3, MLC, or claudin-1, -2, and -5.** (A and B) Immunolocalization showed normal patterns of Arp2, Arp3, and MLC in iNWKO mice and NWKD cells. (C) Western blot for Triton-soluble (S) cytoplasmic fraction and Triton-insoluble (I) membrane/cytoskeletal fractions of primary intestinal epithelial lysates did not show any significant differences in key AJC proteins, including Arp complex proteins, claudin-1 and -2, and MLC in WT and iNWKO mice. (D) Immunolocalization showed similar patterns of claudin-1, -2, and -5 in iNWKO and WT control mice. Scale bars: (A and D) 20 μm; (B) 5 μm.
increased proliferation and more rapid confluency in NWKD monolayers, but TER essentially plateaued and did not significantly increase beyond day 14 in culture.

To determine whether N-WASP influences the ability of the AJC to dynamically alter permeability in response to exogenous signals, we used the calcium-switch technique, in which removal of calcium from the culture medium of mature monolayers rapidly induces junction disassembly and loss of TER, and restoring calcium promotes junction reassembly and recovery of TER. Consistent with a role for N-WASP in internalization of AJC proteins, N-WASP depletion led to a delayed response during calcium withdrawal compared with WT controls: 10 minutes after calcium removal, WT monolayers showed a 27.6% loss of TER (range, 17.7%–36.9%), whereas TER in NWKD monolayers was essentially unchanged from its starting value (99.6%–104%; \( P < .05 \) vs WT), and took approximately twice as long to reach a similar decrement (28% decrease in TER after 19.4 minutes) (Figure 4A). Delayed junction disassembly in NWKD cells also was reflected by the persistence of well-formed tight junctions, with increased persistence of occludin, but not E-cadherin or F-actin, in NWKD cells compared with WT control cells during calcium depletion (Figure 4B).

N-WASP depletion also resulted in delayed junction reassembly. After overnight culture in low-calcium medium, we performed serial TER measurements in WT and NWKD cultures after restoring calcium to the media. Recovery of TER in NWKD cells significantly lagged behind that of WT: although WT cells rapidly recovered monolayer resistance between 2 and 6 hours after calcium repletion, TER in NWKD cultures only gradually began to increase...
after 2 hours, and did not return to steady-state TER until after >8 hours (Figure 4C). The delay in TER recovery induced by N-WASP depletion also correlated with delayed recovery of AJC morphology. Although both WT and NWKD cells displayed a complete disruption of occludin localization after overnight culture in low-calcium media (Figure 4D, left panels), the majority of occludin signal relocated to the cell–cell junctions in WT cells 8 hours after restoring calcium to the culture medium. In contrast, occludin remained primarily localized to the cytoplasm in NWKD cultures at 8 hours (Figure 4D, top panel). In contrast to the effects observed for occludin, junction localization of E-cadherin and F-actin appeared similar in calcium-repleted NWKD and control monolayers (Figure 4D, lower panels).

**Host N-WASP and SNX9 Are Required for EPEC-Induced Barrier Dysfunction and Junction Disruption**

Because EspF, an AE pathogen effector protein essential for TJ disruption, is capable of binding both N-WASP and the endocytic modulator SNX9, we hypothesized that EspF induces intestinal barrier dysfunction by simultaneously activating N-WASP and SNX9 to stimulate the internalization of AJC proteins. We first assessed the ability of WT EPEC, expressing native EspF, to induce TJ disruption and permeability changes. After 4 hours of infection, EPEC induced a 61.4% decrease in TER (range, 51%–72%) in WT monolayers, whereas NWKD monolayers were more resistant to barrier disruption, with TER decreasing by 28.8%
(range, 25%–34% P < .01 for ΔTER in NWKD vs WT) (Figure 5A). As expected,11,34,35 membrane-attached EPEC induced robust actin pedestal formation on WT cells (Figure 5B and C), along with the relocalization of ZO-1 to the tip of actin pedestals (Figure 5B, white arrows). Somewhat unexpectedly, EPEC also was able to induce pedestals on NWKD cells (Figure 5B, second row), but despite the presence of a large number of actin pedestals, did not disrupt ZO-1 or E-cadherin, or induce the relocalization of these AJC proteins to actin pedestals in the setting of reduced N-WASP levels. Apical (XY plane) views showed strong staining of both E-cadherin and ZO-1 at the periphery between neighboring EPEC-infected NWKD cells, and side views (XZ and YZ planes) showed well-defined linear strands of both AJC proteins (Figure 5B and C). In contrast, ZO-1 was not localized at junctions in EPEC-infected WT cells (Figure 5B, top row). Consistent with these findings, surface intensity plots of ZO-1 pixel distribution derived from cell–cell boundaries showed reduced peak intensities in EPEC-infected WT, but not NWKD, cells (data not shown), indicating inefficient TJ disruption in the absence of N-WASP.

Each PRR from EspF has nonoverlapping binding sites for both N-WASP and SNX9, suggesting that the ability to simultaneously interact with both of these host proteins may be important for the maximal activity of EspF.12 To investigate whether SNX9, similar to N-WASP, was required for EPEC-induced junction disruption, we

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**Figure 5.** N-WASP and SNX9 are required for EspF-mediated junction disruption during infection with EPEC. (A) Change in TER after 4 hours of EPEC infection in NWKD or SNX9 cells. (B and C) Confocal immunofluorescence of WT, NWKD, and SNX9KD monolayers infected with EPEC. Both NWKD (middle row) and SNX9KD (bottom row) cells supported robust actin-pedestal formation during EPEC infection (actin pedestals on higher-magnification insets), comparable with WT cells, but although EPEC disrupted the normal localization of ZO-1 with re-localization to the pedestal (upper row, white arrows) in WT cells, junctional localization of E-cadherin and ZO-1 was essentially preserved in both NWKD and SNX9KD cells. Orthogonal views (to the right and below each main panel) also showed preserved linear strands of E-cadherin and ZO-1 in NWKD and SNX9KD cells whereas linear strands of ZO-1 were lost in EPEC-infected WT cells. In NWKD and SNX9KD, there were numerous cytoplasmic perijunctional puncta enriched for E-cadherin (yellow arrows), which were rarely observed in infected WT cells. (D) Western blot confirming efficient knockdown of SNX9 protein using shRNA. Scale bars: 10 μm.
generated Caco-2 cell lines with stable SNX9 knockdown (SNX9KD) (Figure 5D). Similar to the NWKD cells, SNX9KD cells were partially resistant to EPEC-induced permeability changes, with TER decreasing by 43.7% (range, 22%–58%; P < .05 for ΔTER in SNX9KD vs WT) after 4 hours in EPEC-infected SNX9KD cells, compared with a 61.4% decrease in TER (range, 51%–72%) in WT cells (Figure 5A). After 4 hours, EPEC-infected SNX9KD cells showed a large number of actin pedestals and loss of the brush border (Figure 3B, bottom row); however, similar to NWKD cells, E-cadherin and ZO-1 retained their discrete junctional localization patterns without relocalization of ZO-1 to actin pedestals (Figure 5B and C, bottom rows). Of note, both NWKD and SNX9KD cells showed numerous cytoplasmic perijunctional puncta (mean, 4.08/cell ± 0.41 SEM for NWKD; 4.26/cell ± 0.47 SEM for SNX9KD), which were enriched for E-cadherin but not ZO-1 (Figure 5C, yellow arrows), which may represent vesicles in an E-cadherin–specific recycling pathway that have accumulated owing to impaired efficiency of endocytosis in the absence of N-WASP or SNX9.

EspF Requires Both N-WASP and SNX9 Binding to Induce Efficient Relocalization of TJ Proteins

We next sought to determine whether EPEC strains expressing EspF variants that are unable to bind either N-WASP or SNX9 are able to induce epithelial barrier dysfunction. We introduced point mutations in each of the 3 C-terminal PRR repeats in either the N-WASP (L31A) or SNX9 (R3D) binding domains of espF (Figure 6A), previously shown to be important for binding to N-WASP and SNX9.12 We then complemented the EspF-deficient EPEC strain UMD874 (E2348/69ΔespF) with N-WASP-binding–defective EspF (pEspFNWBD) or SNX9-binding–defective EspF (pEspFSNX9BD), generating EPECΔespF/pEspFNWBD and EPECΔespF/pEspFSNX9BD.

Figure 6. Mutations in N-WASP– and SNX9-binding regions of EspF attenuate EPEC-induced junction disruption in vitro. (A) Schematic of point mutations made in each PRR of EspF, rendering it unable to bind to N-WASP or SNX9. (B) WT EPEC induced robust actin pedestal formation, disruption of ZO-1 localization, and recruitment of ZO-1 to the tips of actin pedestals; in contrast, EPECΔespF/pEspFNWBD and EPECΔespF/pEspFSNX9BD were both attenuated in their ability to induce TJ disruption, with persistent junction localization of ZO-1 (arrowheads) in infected cells. (C) WT cells infected with EPEC expressing WT EspF showed a 61.4% reduction in TER at 4 hours after infection; in contrast, EPECΔespF/pEspFNWBD was only able to induce a 28.4% reduction in TER, and there was minimal change in TER at 4 hours in monolayers infected with EPECΔespF/pEspFSNX9BD (P < .01 for WT EPEC vs EPECΔespF/pEspFNWBD; P < .01 for WT EPEC vs EPECΔespF/pEspFSNX9BD). Scale bars: 10 μm.
pEspFSNX9BD, respectively. WT cells infected with EPEC expressing WT EspF showed a 61.4% reduction in TER at 4 hours after infection; in contrast, EPECΔespF/pEspFSNX9BD was only able to induce a 28.4% reduction in TER, and there was minimal change in TER at 4 hours in monolayers infected with EPECΔespF/pEspFSNX9BD (P < .01 for WT EPEC vs EPECΔespF/pEspFSNX9BD, P < .01 for WT EPEC vs EPECΔespF/pEspFSNX9BD) (Figure 6C). Both EPECΔespF/pEspFSNX9BD and EPECΔespF/pEspFSNX9BD were impaired in their ability to induce tight junction disruption, and ZO-1 remained discretely localized in subapical linear strands at the junction without relocating to actin pedestals (Figure 6B). Together, these results show that EspF specifically targets and disrupts tight junctions, and binding to both host N-WASP and SNX9 are required for its full activity.

C. rodentium EspF Must Bind Both N-WASP and SNX9 to Induce Intestinal Pathology In Vivo

To test whether N-WASP and SNX9 binding are important for the pathogenicity of AE bacteria in vivo, we generated a C. rodentium (CR) strain lacking EspF, and complemented this with a N-WASP- or SNX9-binding-defective EspF constructs. WT mice were infected with a single dose of 5 x 10⁸ colony-forming units WT CR, CRΔespF, CRΔespF/pEspFSNX9BD or CRΔespF/pEspFSNX9BD and followed up for 7 days. We first confirmed that WT CR could efficiently colonize WT mice and noted numerous actin pedestals and AE lesions in WT, but not iNWKO, mice by EM (Figure 7A, arrows). As previously described,36,37 mice infected with WT CR lost weight (Figure 7B) and shed high titers of CR in the stool throughout the first 7 days of infection (Figure 5C). In contrast, disease was attenuated in mice infected with isogenic CRΔespF strains expressing variant EspF unable to bind either N-WASP or SNX9. Mice infected with CRΔespF/pEspFNWBD or CRΔespF/pEspFSNX9BD did not lose weight (Figure 7B). Both the WT and mutant CR strains were able to efficiently colonize mice (Figure 7C). Consistent with previous reports,9 CR expressing WT EspF induced marked disruption of junction localization of ZO-1 (Figure 7D, upper right). In contrast, and despite the presence of at least 10⁷ colony-forming units 2018 N-WASP Regulates TJs and EspF Activity 283

Figure 7. Mutations in N-WASP- and SNX9-binding regions of EspF attenuate CR-induced junction disruption in vivo. (A) WT CR induced robust actin-dense pedestal formation in WT, but not iNWKO mice. (B) WT mice infected with WT CR (black line), but not CR complemented with N-WASP- or SNX9-binding–defective EspF (red and blue lines, respectively), showed weight loss during acute infection. (C) Bacterial colonization by WT CR, CRΔespF/pEspFNWBD and CRΔespF/pEspFSNX9BD. (D) CR expressing wild-type EspF induced marked disruption of normal ZO-1 patterns by 7 days after infection (upper right); in contrast, CRΔespF strains complemented with N-WASP or SNX9-binding–defective EspF failed to disrupt localization of ZO-1 at intercellular junctions. (E) Loss of EspF binding to host N-WASP (CRΔespF/pEspFNWBD) attenuates increases in intestinal permeability to FITC-dextran induced by CR during in vivo infection. Scale bars: 20 μm. RLU, relative light unit.
units/g of stool CR, the strains complemented with N-WASP- and SNX9-binding–defective EspF constructs were unable to induce AJC disruption, and ZO-1 appeared essentially unaffected (Figure 7D, bottom panels). Attenuation in the ability of CR expressing N-WASP–binding–defective EspF also correlated with protection from pathogen-induced increases in intestinal permeability to FITC-dextran compared with mice infected with WT CR (Figure 7E).

**Discussion**

In the intestinal epithelium, the AJC is intimately linked to dynamic networks of actin, and the ability of N-WASP to rapidly catalyze actin polymerization makes it an ideal candidate for controlling junction protein mobilization and turnover, which may ultimately influence AJC permeability. We found that mice lacking N-WASP in the intestinal epithelium have abnormal junction morphology and increased intestinal permeability. The N-WASP levels of ZO-1, claudin-1, and E-cadherin, and increased filament treadmilling, in which actin polymerization is counterbalanced with depolymerization, and the relative contributions of these closely linked processes enables actin filaments to generate propulsive forces and serve as dynamic structural scaffolds. The N-WASP–Arp2/3 complex primarily promotes polymerization of actin filaments and the formation of branched-actin–filament networks, whereas actin depolymerization factor (ADF) family proteins (ADF, coflin-1, coflin-2) severs and disassembles existing filaments, an activity that is significantly enhanced by the cofactor actin-interacting protein 1. Notably, several recent studies have suggested a key role for ADF/cofilin in regulating the remodeling and permeability of epithelial junctions because loss of either coflin-1, ADF, or actin-interacting protein 1 was associated with increased epithelial permeability, delayed AJC assembly, and increased susceptibility to dextran sodium sulfate–induced colitis. Although ADF/cofilin proteins primarily promote depolymerization during actin treadmilling, the activity of ADF/cofilin on existing filaments produces free barbed ends for subsequent Arp2/3 filament nucleation, and the overall result may be to promote N-WASP–Arp2/3–dependent actin polymerization, in which newly polymerized actin filaments are more stable than on older filaments. Therefore, in addition to defects in AJC protein endocytosis, loss of N-WASP in the intestinal epithelium also may disrupt the critical balance of actin polymerization and depolymerization by failing to nucleate new actin filaments at ADF/cofilin-produced free barbed ends. The overall result may be defective perijunctional force-generating polymer motors and protein scaffolds leading to increased permeability and impaired dynamic AJC assembly and disassembly.

The significance of cytoskeletal regulation and its relationship to intestinal barrier maintenance also is highlighted by enteric pathogens that specifically target host proteins, including N-WASP, as part of their pathogenic strategy. EPEC, EHEC, and C. rodentium induce diarrheal illness in part by injecting the highly conserved virulence factor EspF into enterocytes via type III secretion. Although EspF has been shown to be a key mediator of AE pathogen-induced disruption of the intestinal barrier, the mechanism of EspF-dependent effects on the AJC is largely unknown. Because we found that N-WASP plays a role in the homeostatic maintenance of AJCs, and EspF has been shown to directly bind and activate N-WASP with high affinity, we reasoned that EspF might act on the intercellular junction through an N-WASP–dependent pathway. Indeed, we found that N-WASP–depleted cells were resistant to EPEC-induced loss of barrier integrity in vitro. Because N-WASP also has been shown to play a role in facilitating the translocation of bacterial effector proteins by EHEC, we considered whether the lack of junction disruption might be owing to ineffective type III secretion and impaired delivery of EspF. Several pieces of data indicate that translocation abnormalities in NWKD cells are unlikely to account for the failure of EspF-mediated barrier disruption. First, NWKD cells supported EPEC-induced actin pedestal formation, which is
dependent on Tir translocation.\textsuperscript{52} We acknowledge that the finding of actin pedestals on EPEC-infected NWKD Caco-2 cells, but not during \textit{C. rodentium} infection in mice, was somewhat unexpected. We suspect that given the central importance of Tir translocation and formation of the actin pedestal formation, AE pathogens have optimized mechanisms for recruiting and using even low levels of N-WASP protein, which may be present in the knockdown cells, but not in the epithelium of genetically depleted mice. In addition, ectopic expression of a WT EspF:green fluorescent protein (GFP) reporter construct induced strong cytoplasmic relocalization of TJ proteins in transfected WT, but not NWKD, cells (data not shown). Mutating N-WASP (or SNX9) binding sites in EspF, which is not predicted to affect EspF translocation, impaired the ability of EspF to induce TJ disruption in WT cells, which resembled the impairment seen in NWKD cells infected with EPEC expressing WT EspF. Taken together, these data show that EspF-induced tight junction disruption is mediated through N-WASP.

In addition to EspF/N-WASP interactions, we found that EspF also must bind SNX9 to disrupt the AJC. Previous studies have shown that an EPEC strain expressing SNX9-binding–deficient mutant EspF could disrupt TER in polarized T84 cells,\textsuperscript{12} suggesting that the ability to interact with SNX9 may be dispensable for EPEC-induced junction disruption. However, more recent work by Tapia et al.\textsuperscript{13} showed that EPEC EspF disrupts epithelial polarity, induces endocytosis, redistributes Na\textsuperscript{+}/K\textsuperscript{+} adenosine triphosphatase, and disrupts TJs by displacing Crumbs (Crb)3 from the Crb polarity complex of Crb3/Pals1/Patj. In these studies, the ability of EspF to induce mislocalization of Crb3 was abolished by either blocking the interaction of EspF/SNX9 or inhibiting the guanosine triphosphatase activity of dynamin. We found that SNX9-depleted Caco-2 cells were resistant to EPEC-induced loss of TER and that EPEC\textit{ΔespF} transcomplemented with SNX9-binding–defective EspF failed to disrupt TER and AJC morphology. These discrepant results may be owing to differences in the expression of SNX9 and other closely related sorting nexins in different colon cancer cell lines.\textsuperscript{54–56} Collectively, our findings show that EspF is similarly dependent on SNX9 and N-WASP to mediate EPEC-induced TJ alterations.

In vivo, EspF binding to either N-WASP or SNX9 appeared to play an important role in pathogen-induced weight loss (Figure 5B). Other investigators\textsuperscript{57,58} have reported a modest degree of weight loss during CR infection, and we speculate that this may be owing to several factors. First, CR expressing WT EspF induced significant AJC disruption, which is associated with increased enteric fluid and electrolyte loss. In addition, although CR complemented native EspF. Differences in bacterial burden, reflecting enhanced pathogenicity of CR expressing nonvariant EspF, also may contribute to significant weight loss through changes in the underlying microbiota.\textsuperscript{59,60}

The current work highlights the dual role that the master cytoskeletal regular N-WASP plays in dynamic AJC assembly and disassembly. Further work will be required to determine how N-WASP facilitates the delivery and removal of specific TJ components to maintain intestinal homeostasis. A common theme in host–pathogen interactions is the utilization of otherwise homeostatic cell processes by invading microbes to enhance their pathogenesis. Consistent with the requirement of N-WASP for optimal junction plasticity and stability, we also found that EspF, an SNX9 interactor and direct inducer of N-WASP–dependent actin polymerization,\textsuperscript{12} requires both N-WASP and SNX9 for efficient junction disruption. Although EspF has been shown previously to bind to both N-WASP and SNX9, we show, both in vitro and in vivo, that this binding is required for AE pathogens to induce intestinal barrier dysfunction. In addition to EPEC, EHEC, and \textit{C. rodentium}, numerous microbes manipulate the host cytoskeletal machinery including the clinically important species \textit{Helicobacter pylori} (associated with gastric ulcers and cancer),\textsuperscript{61,62} adherent-invasive \textit{E. coli} (associated with Crohn’s disease),\textsuperscript{63} and segmented filamentous bacteria (a potent inducer of Th17 immune responses).\textsuperscript{64} By understanding the functions of N-WASP, and cytoskeletal regulation more generally, additional insight will be gained into host cell processes that are critical for cellular homeostasis as well as targets for many important human pathogens.

References

1. Ivanov AI, McCall IC, Parkos CA, Nusrat A. Role for actin filament turnover and a myosin II motor in cytoketons-driven disassembly of the epithelial apical junctional complex. Mol Biol Cell 2004;15:2639–2651.
2. Mitic LL, Anderson JM. Molecular architecture of tight junctions. Annu Rev Physiol 1998;60:121–142.
3. Shen L, Turner JR. Actin depolymerization disrupts tight junctions via caveolae-mediated endocytosis. Mol Biol Cell 2005;16:3919–3936.
4. Kovacs EM, Verma S, Ali RG, Ratheesh A, Hamilton NA, Akhmanova A, Yap AS. N-WASP regulates the epithelial junctional actin cytoskeleton through a non-canonical post-nucleation pathway. Nat Cell Biol 2011;13:934–943.
5. Ivanov AI, Hunt D, Utech M, Nusrat A, Parkos CA. Differential roles for actin polymerization and a myosin II motor in assembly of the epithelial apical junctional complex. Mol Biol Cell 2005;16:2636–2650.
6. Otani T, Ichii T, Aono S, Takeichi M. Cdc42 GEF Tuba regulates the junctional configuration of simple epithelial cells. J Cell Biol 2006;175:135–146.
7. Georgiou M, Marinari E, Burden J, Baum B. Cdc42, Par6, and aPKC regulate Arp2/3-mediated endocytosis to control local adherens junction stability. Curr Biol 2008;18:1631–1638.
8. Leibfried A, Fricke R, Morgan MJ, Bogdan S, Bellaiche Y. Drosophila Cip4 and WASP define a branch of the Cdc42-Par6-aPKC pathway regulating E-cadherin endocytosis. Curr Biol 2008;18:1639–1648.
9. Guttman JA, Li Y, Wickham ME, Deng W, Vogl AW, Finlay BB. Attaching and effacing pathogen-induced
tight junction disruption in vivo. Cell Microbiol 2006; 8:634–645.

10. Ma C, Wickham ME, Guttmann JA, Deng W, Walker J, Madsen KL, Jacobson K, Vogl WA, Finlay BB, Vannallce BA. Citrobacter rodentium infection causes both mitochondrial dysfunction and intestinal epithelial barrier disruption in vivo: role of mitochondrial associated protein (Map). Cell Microbiol 2006; 8:1669–1886.

11. Lommel S, Benesch S, Rohde M, Wehland J, Rottner K. Enterohaemorrhagic and enteropathogenic Escherichia coli use different mechanisms for actin pedestal formation that converge on N-WASP. Cell Microbiol 2004; 6:249–254.

12. Alto NM, Weflen AW, Rardin MJ, Yarar D, Lazar CS, Tonikian R, Koller A, Taylor SS, Boone C, Sidhu SS, Schmid SL, Hecht GA, Dixon JE. The type III effector EspF coordinates membrane trafficking by the spatiotemporal activation of two eukaryotic signaling pathways. J Cell Biol 2007; 178:1265–1278.

13. van Weering JR, Sessions RB, Traer CJ, Kloer DP, Bhatia VK, Stamou D, Carlsson SR, Hurley JH, Cullen PJ. Molecular basis for SNX-BAR-mediated assembly of distinct endosomal sorting tubules. EMBO J 2012; 31:4466–4480.

14. Worby CA, Dixon JE. Sorting out the cellular functions of sorting nexins. Nat Rev Mol Cell Biol 2002;3:919–931.

15. el Marjou F, Janssen KP, Chang BH, Li M, Hindie V, Chan L, Louvard D, Chambon P, Metzger D, Robine S. Tissue-specific and inducible Cre-mediated recombination in the gut epithelium. Genesis 2004;39:186–193.

16. Whitehead RH, Robinson PS. Establishment of conditionally immortalized epithelial cell lines from the intestinal tissue of adult normal and transgenic mice. Am J Physiol Gastrointest Liver Physiol 2009;296:G455–G460.

17. Napolitano LM, Koruda MJ, Meyer AA, Baker CC. The impact of femur fracture with associated soft tissue injury on immune function and intestinal permeability. Shock 1996;5:202–207.

18. Cereijido M, Gonzalez-Mariscal L, Contreras RG, Gallardo JM, Garcia-Villegas R, Valdes J. The making of a tight junction. J Cell Sci Suppl 1993;17:127–132.

19. Diamond JM. Twenty-first Bowditch lecture. The epithelial junction: bridge, gate, and fence. Physiologist 1977;20:10–18.

20. Wegener J, Sieber M, Galla HJ. Impedance analysis of epithelial and endothelial cell monolayers cultured on gold surfaces. J Biochem Biophys Methods 1996; 32:151–170.

21. Miyoshi H, Stappenbeck TS. In vitro expansion and genetic modification of gastrointestinal stem cells in spheroid culture. Nat Protoc 2013;8:2471–2482.

22. Clayburgh DR, Barrett TA, Tang Y, Meddings JB, Van Eldik LJ, Watterson DM, Clarke LL, Msrny RJ, Turner JR. Epithelial myosin light chain kinase-dependent barrier dysfunction mediates T cell activation-induced diarrhea in vivo. J Clin Invest 2005;115:2702–2715.

23. Gumbiner B, Simons K. A functional assay for proteins involved in establishing an epithelial occluding barrier: identification of a uvomorulin-like polypeptide. J Cell Biol 1986;102:457–468.

24. Levine MM, Nataro JP, Karch H, Baldini MM, Kaper JB, Black RE, Clements ML, O'Brien AD. The diarrheal response of humans to some classic serotypes of enteropathogenic Escherichia coli is dependent on a plasmid encoding an enteroadhesiveness factor. J Infect Dis 1985;152:550–559.

25. McNamara BP, Donnenberg MS. A novel proline-rich protein, EspF, is secreted from enteropathogenic Escherichia coli via the type III export pathway. FEMS Microbiol Lett 1998;166:71–78.

26. Datsenko KA, Wanner BL. One-step inactivation of chromosomal genes in Escherichia coli K-12 using PCR products. Proc Natl Acad Sci U S A 2000; 97:6640–6645.

27. Nougayrede JP, Donnenberg MS. Enteropathogenic Escherichia coli EspF is targeted to mitochondria and is required to initiate the mitochondrial death pathway. Cell Microbiol 2004;6:1097–1111.

28. Cheng HC, Skehan BM, Campellone KG, Leong JM, Rosen MK. Structural mechanism of WASP activation by the enterohaemorrhagic E. coli effector EspF(U). Nature 2008;454:1009–1013.

29. Snapper CM, Shen Y, Khan AQ, Colino J, Zelazowski P, Mond JJ, Gause WC, Wu QZ. Distinct types of T-cell help for the induction of a humoral immune response to Streptococcus pneumoniae. Trends Immunol 2001; 22:308–311.

30. Cotta-de-Almeida V, Westerberg L, Maillard MH, Onaldi D, Wachtel H, Melu P, Chung UI, Xavier R, Alt FW, Snapper SB. Wiskott Aldrich syndrome protein (WASP) and N-WASP are critical for T cell development. Proc Natl Acad Sci U S A 2007;104:15424–15429.

31. Lyubimova A, Garber JG, Upadhyay G, Sharov A, Anastasoiaie F, Yajnik V, Cotsarelis G, Dotto GP, Botchkarev V, Snapper SB. Neural Wiskott-Aldrich syndrome protein modulates Wnt signaling and is required for hair follicle cycling in mice. J Clin Invest 2010; 120:446–456.

32. Mallick EM, Garber JG, Vanguri VK, Balasubramanian S, Blood T, Clark S, Vingadassalom D, Louissaint C, McCormick B, Snapper SB, Leong JM. The ability of an attaching and effacing pathogen to trigger localized actin assembly contributes to virulence by promoting mucosal attachment. Cell Microbiol 2014;16: 1405–1424.

33. Smutny M, Cox HL, Leerreberg JM, Kovacs EM, Conti MA, Ferguson C, Hamilton NA, Parton RG, Adelstein RS, Yap AS. Myosin II isoforms identify distinct functional modules that support integrity of the epithelial zonula adherens. Nat Cell Biol 2010;12:696–702.

34. Kalman D, Weiner OD, Goosney DL, Sedat JW, Finlay BB, Abo A, Bishop JM. Enteropathogenic E. coli acts through WASP and Arp2/3 complex to form actin pedestals. Nat Cell Biol 1999;1:389–391.

35. Crepin VF, Girard F, Schuller S, Phillips AD, Mousnier A, Frankel G. Dissecting the role of the Tir:Nck and Tir:ITRTKS/IRSp53 signalling pathways in vivo. Mol Microbiol 2010;75:308–323.

36. Lupcherio SA, Newman JV, Danger CA, Schrenzel MD, Brenner DJ, Steigerwalt AG, Schauer DB. Citrobacter.
rodentium, the causative agent of transmissible murine colonic hyperplasia, exhibits clonality: synonymy of C. rodentium and mouse-pathogenic Escherichia coli. J Clin Microbiol 2000;38:4343–4350.

37. Schauer DB, Falkow S. The eae gene of Citrobacter freundii biotype 4280 is necessary for colonization in transmissible murine colonic hyperplasia. Infect Immun 1993;61:4654–4661.

38. Zhou K, Sumigray KD, Lechler T. The Arp2/3 complex has essential roles in vesicle trafficking and transcytosis in the mammalian small intestine. Mol Biol Cell 2015;26:1995–2004.

39. Attar N, Cullen PJ. The retromer complex. Adv Enzyme Regul 2010;50:216–236.

40. Bonifacino JS, Hurley JH. Retromer. Curr Opin Cell Biol 2008;20:427–436.

41. Collins BM. The structure and function of the retromer complex. Traffic 2008;9:1811–1822.

42. Citalan-Madrid AF, Vargas-Robles H, Garcia-Ponce A, Shibayama M, Betanzos A, Nava P, Salinas-Lara C, Rottner K, Mennigen R, Schoo M. Cortactin deficiency causes increased RhoA/ROCK1-dependent actomyosin contractility, intestinal epithelial barrier dysfunction, and disproportionately severe DSS-induced colitis. Mucosal Immunol 2017;10:1237–1247.

43. Goley ED, Welch MD. The ARP2/3 complex: an actin nucleator comes of age. Nat Rev Mol Cell Biol 2006;7:713–726.

44. Mohri K, Ravi H, Smith EM, Goode BL. Aip1 and cofilin-1 have unique and overlapping functions in the pathogenesis of Citrobacter rodentium colitis. Infect Immun 2008;76:436–443.

45. Nadkarni AV, Brieher WM. Aip1 destabilizes cofilin-saturated actin filaments by severing and accelerating monomer dissociation from ends. Curr Biol 2014;24:2749–2757.

46. Okada K, Ravi H, Smith EM, Goode BL. Aip1 and cofilin promote rapid turnover of yeast actin patches and cables: a coordinated mechanism for severing and capping filaments. Mol Biol Cell 2006;17:2190–2199.

47. Nadkarni AV, Brieher WM. Aip1 destabilizes cofilin-saturated actin filaments by severing and accelerating monomer dissociation from ends. Curr Biol 2014;24:2749–2757.

48. Okada K, Ravi H, Smith EM, Goode BL. Aip1 and cofilin promote rapid turnover of yeast actin patches and cables: a coordinated mechanism for severing and capping filaments. Mol Biol Cell 2006;17:2190–2199.

49. Lechuga S, Baranwal S, Ivanov AI. Actin-interacting protein 1 controls assembly and permeability of intestinal epithelial apical junctions. Am J Physiol Gastrointest Liver Physiol 2015;308:G745–G756.

50. Wang D, Naydenov NG, Feygin A, Baranwal S, Kuenmerle JF, Ivanov AI. Actin-depolymerizing factor and cofilin-1 have unique and overlapping functions in regulating intestinal epithelial junctions and mucosal inflammation. Am J Pathol 2016;186:844–858.

51. Andrianantoandro E, Pollard TD. Mechanism of actin filament turnover by severing and nucleation at different concentrations of ADF/cofilin. Mol Cell 2006;24:13–23.

52. Campellone KG, Rankin S, Pawson T, Kirschner MW, Tipper DJ, Leong JM. Clustering of Nck by a 12-residue Tir phosphopeptide is sufficient to trigger localized actin assembly. J Cell Biol 2004;164:407–416.

53. Tapia R, Kralicek SE, Hecht GA. EPEC effector EspF promotes Crombr3 endocytosis and disrupts epithelial cell polarity. Cell Microbiol 2017;19(11). https://doi.org/10.1111/cmi.12757, Epub 2017 Jul 27.

54. Zhang J, Zhang X, Guo Y, Xu L, Pei D. Sorting nexin 33 induces mammalian cell micronucleated phenotype and actin polymerization by interacting with Wiskott-Aldrich syndrome protein. J Biol Chem 2009;284:21659–21669.

55. Andrianantoandro E, Pollard TD. Mechanism of actin filament turnover by severing and nucleation at different concentrations of ADF/cofilin. Mol Cell 2006;24:13–23.

56. Wickham ME, Brown NF, Boyle EC, Coombes BK, Darfeuille-Michaud A, Boudeau J, Bulois P, Neut C, Karaoz U, Wei D, Goldfarb KC, Santee CA, Lynch SV, Tanoue T, Imaoka A, Itoh K, Takeda K, Umesaki Y, Karaoz U, Wei D, Goldfarb KC, Santee CA, Lynch SV, Tanoue T, Imaoka A, Itoh K, Takeda K, Umesaki Y, Honda K, Littman DR. Induction of intestinal Th17 cells by segmented filamentous bacteria. Cell 2009;139:485–498.
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Author contributions
John Garber was responsible for the study concept and design, acquisition, analysis, and interpretation of data, and drafting the manuscript; Emily Mallick and Karen Scanlon were responsible for the acquisition and analysis of data; Jerrold Turner was responsible for the analysis and interpretation of data, and critical revision of the manuscript for important intellectual content; Michael Donnenberg was responsible for the analysis and interpretation of data, and critical revision of the manuscript for important intellectual content; John Leong was responsible for the study concept and design, analysis and interpretation of data, and critical revision of the manuscript for important intellectual content; and Scott Snapper was responsible for the study concept and design, interpretation of data, drafting the manuscript, and study supervision.

Conflicts of interest
The authors disclose no conflicts.

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