The interaction of *Escherichia coli* O157:H7 and *Salmonella Typhimurium* flagella with host cell membranes and cytoskeletal components

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**Abstract**

Bacterial flagella have many established roles beyond swimming motility. Despite clear evidence of flagella-dependent adherence, the specificity of the ligands and mechanisms of binding are still debated. In this study, the molecular basis of *Escherichia coli* O157:H7 and *Salmonella enterica* serovar Typhimurium flagella binding to epithelial cell cultures was investigated. Flagella interactions with host cell surfaces were intimate and crossed cellular boundaries as demarcated by actin and membrane labelling. Scanning electron microscopy revealed flagella disappearing into cellular surfaces and transmission electron microscopy of *S*. Typhimurium indicated host membrane deformation and disruption in proximity to flagella. Motor mutants of *E. coli* O157:H7 and *S*. Typhimurium caused reduced haemolysis compared to wild-type, indicating that membrane disruption was in part due to flagella rotation. Flagella from *E. coli* O157 (H7), EPEC O127 (H6) and *S*. Typhimurium (P1 and P2 flagella) were shown to bind to purified intracellular components of the actin cytoskeleton and directly increase in vitro actin polymerization rates. We propose that flagella interactions with host cell membranes and cytoskeletal components may help prime intimate attachment and invasion for *E. coli* O157:H7 and *S*. Typhimurium, respectively.

**INTRODUCTION**

Bacterial flagella are macromolecular machines predominately associated with movement toward preferred environmental niches [1]. They are composed of a long, hollow, capped filament of polymeric flagellin, connected to a flagella basal body through a hook complex [2]. The basal body houses a motor complex that rotates the basal body rod, hook and semi-rigid helical flagella filament, or flagellum, to move the bacterium [3, 4]. The basal body also contains an adapted type 3 secretion export system that regulates secretion of the different components during flaggellum assembly [5, 6]. Flagellin monomers make up the majority of the flagellum and have a hairpin-like structure with highly conserved associating termini and central variable domains [7]. *Salmonella enterica* serovar Typhimurium can alternately express two flagella types, presumably to help avoid the adaptive immune response; phase 1 (P1)
flagella are composed of FliC flagellin, and phase 2 (P2) flagella are composed of FljB flagellin [8]. In the stacked flagella filament, the flagellin region exposed on the surface shows little conservation between strains and is used for immunological serotyping, resulting in the H-type. The flagellin termini are stacked inside the filament and drive flagellin polymerization [9]. These more conserved and buried regions are important microbe-associated molecular patterns (MAMPs), which activate Toll-like receptor 5 (TLR5) and NOD-like and CARD domain-containing receptor 4 (NLRC4) [10–13].

In peritrichous and monotrichous species, flagella have been shown to sense surfaces and cause a switch from motile to sessile lifestyles, making flagellar adhesion the preliminary step essential for bacterial colonization [14]. More recently, flagellar rotation has been shown to play a role in flagella adherence, surface stiffness sensing and inducing neutrophil extracellular traps [15, 16]. However, mechanistic information beyond near-surface swimming [17] is lacking and direct adherence along the filament shaft to cell surface components is poorly defined.

Important zoonotic enteropathogens, enterohaemorrhagic Escherichia coli O157:H7 and S. Typhimurium both express flagella and type 3 secretion systems (T3SSs), but have different intestinal colonization strategies. T3SSs are structurally related to the flagella basal body and act as molecular needles to inject effector proteins into host cells [18]. E. coli O157:H7 use their T3SSs to inject effector proteins into host intestinal epithelial cells, inducing actin polymerization and the formation of attaching and effacing lesions [19]. This cytoskeletal network modulation allows E. coli O157:H7 to bind very tightly to the surface of host enterocytes while avoiding phagocytosis [20, 21]. In contrast to E. coli O157:H7, S. Typhimurium translocates T3SS effectors into host epithelial cells, leading to changes in the actin cytoskeleton that promote bacterial invasion [22]. Salmonella survival in host cells requires a second T3SS, which manipulates host cell responses and the position and recognition of the Salmonella containing vacuole (SCV) [23]. E. coli O157:H7 is mainly an extracellular bacterium, but is reliant on both a T3SS and its H7 flagella for cell binding and colonization of its reservoir host, cattle [24–26]. Published studies indicate that H7 flagella bind to mucus and more recent work demonstrated binding of different E. coli flagella, including H7, to ionic phospho-and sulpho-lipids [27, 28]. In contrast, S. Typhimurium is invasive in humans and animals, including cattle and pigs, with invasion being dependent on two different T3SSs [29]. However, the significance of flagella expression by S. Typhimurium for initial attachment and the identity of potential protein ligands remain inconclusive [30–32].

This study takes a closer look at the molecular basis of bacterial flagella adherence to host cells in the context of pathogenic colonization. We start by looking at actin associated with bacterial flagella when bacteria are in contact with host cells. We observe deformation and possible disruption of host plasma membranes in proximity to flagella and explore the possible consequences if membranes are breached by flagella. Actin and flagella interactions are confirmed by pull-downs, far-Western blotting and in vitro actin polymerization assays. These data shed some light on the interactions between the flagella of E. coli O157:H7 and S. Typhimurium and host cells, including membrane and cytoskeletal components, that may prime the next steps in the infection process.

**METHODS**

**Bacterial strains and growth conditions**

Bacterial strains (Table 1) were stored as saturated cultures in lysogeny broth (LB) with 25% glycerol at −70°C. Bacteria were grown in LB at 28–30°C (E. coli O157:H7) or 37°C (S. Typhimurium) at 200 r.p.m. The antibiotics, when required, were chloramphenicol or kanamycin at 50 µg ml⁻¹ and ampicillin at 100 µg ml⁻¹. To assess strain motility, fresh colonies were stab-inoculated once into 0.3% (w/v) LB agar. Inoculated plates were incubated at room temperature (RT) for 36 h. Assays were carried out in quadruplicate from separate colonies.

**Mutant strain construction**

The flagellin gene fliC₁₆ from the E. coli O157:H7 strain TUV93-0 was exchanged with fliC₁₆ according to the allelic exchange method published by Blomfield et al. [33]. Briefly, homologous recombination of pEBW6 or pEBW7 (Table S1, available in the online version of this article) with TUVΔfliC (Table 1) and counter-selection of sacB with sucrose was used to generate TUVΔfliC chromosomally complemented with fliC₁₆ (TUVΔfliC₁₆) and fliC₁₆ (TUVΔfliC₁₆), respectively. Strains were validated by PCR with sacB, fliC locus and O157 specific primer pairs (Table S2), Sanger sequencing, motility in 0.3% (w/v) LB agar and wide-field fluorescence microscopy of saturated cultures with 1:1000 dilutions of H6- and H7- specific rabbit IgG (Table S3). The TUV93-0 motA mutant (ZAP1575, Table 1) was constructed by allelic exchange using pTOF25motA (Table S1) and primer sets: No-motA, Ni-motA, Co-motA and Ci-motA (Table S2). The deleted region corresponds to 2653444–2654278 of E. coli O157:H7 strain E. coli (TUV), respectively. This mutant strain construction was made (ZAP1576, Table 1) using the amplicon of primers No-motA/Co-motA (Table S2) and primer sets: No-motA, Ni-motA, Co-motA and Ci-motA (Table S2). The deleted region corresponds to 2653444–2654278 of EDL933 and was verified by sequencing. A chromosomally complemented strain was made (ZAP1576, Table 1) using the amplicon of primers No-motA/Co-motA (Table S2) and cloning and exchange with the resultant pTOF25 construct. This motA complemented version of ZAP1575 has restored motility (Fig. S3). P22 transduction of phase-locked (hin) and motA mutations into the SL1344 background from the LT2 strains (Table 1) was carried out using standard transduction protocols from strains used and verified in previous studies (Table 1).

**Antibodies and purified proteins**

Abbreviations and/or Uniprot accession codes are in parentheses. The sources, details and use of antibodies and stains are described in Table S3. Rabbit skeletal muscle αβ-actin
(ACTB, P29751), human platelet βγ-actin (ACTB, P60709), recombinant human cofilin-1 (CFL1 P23528) and arp2/3 complex (ARPC4 Q148J6) from bovine brain were all purchased from Cytoskeleton, Inc. Recombinant human galectin-4 (GAL4 P56470) and gelsolin (GSN Q3S×14) from bovine plasma were purchased from R&D Systems and Sigma, respectively.

Flagella were sheared from the mutants TUVΔfliC H7F (H7, Q7DBI0), TUVΔfliC H6F (H6, B7USU2), TUVΔfliC cis complement fliC, from TUV93-0 This study and TUVΔfliC H6F from E2348/69 This study. TUVΔfliC H7F, TUVΔfliC cis complement fliC from TUV93-0 This study were sheared from the mutants TUV93-0 fliC H7F (H7, Q7DBI0), TUVΔfliC H6F (H6, B7USU2), TUVΔfliC−::pEW5 (H48, P04949), SL1344ΔfljB P2 (P2, FljB E1W A22), using an adapted protocol [34]. TUVΔfliC−::pEW5 was grown in the presence of ampicillin at all times. TUVΔfliC−::pEW5 was further sub-cultured in LB in the same conditions to OD600 0.6, at which point 1 mM IPTG was added and cultures were then grown to OD600 1–1.5. All cultures were centrifuged at 4100 g at 4 °C for 15 min, followed by centrifugation once at 16000 g at 4 °C for 10 min, were undertaken to obtain bacteria-free supernatants. These were centrifuged at 145000 g 4 °C for 1.5 h and resuspended in PBS (or 50 mM Tris/HCl pH 7.5 for actin polymerization assays) at 1:500 of initial culture volumes. Protein concentration and purity was assessed using BCA, and then confirmed by Coomassie staining (Imperial protein stain, Thermo Fisher) and densitometry after SDS-PAGE.

Flagella were monomerized according to the protocol published by Smith et al. [35]. Flagella preparations were incubated at 70 °C for 15 min and then filtered by centrifugation at 5000 g at 4 °C for 30 min using 100 kDa 4 ml filter units (Millipore). Filtrates were confirmed by size-exclusion HPLC to be exclusively monomeric.

**Tissue culture**

All cells were maintained at 37 °C in 5 % CO2 and at 80% humidity. The embryonic bovine lung epithelial

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**Table 1. Strains used or constructed in this study**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Relevant features</th>
<th>Source</th>
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<tbody>
<tr>
<td>TUV93-0</td>
<td><em>E. coli</em> O157:H7; EDL933 (ATCC 700927) stx− [67]</td>
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<tr>
<td>TUVΔfliC</td>
<td>TUV93-0 fliC::sacB::kan r, aflagellate [28]</td>
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<tr>
<td>TUVfliC−</td>
<td>TUV93-0 fliC−, aflagellate [28]</td>
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<tr>
<td>TUVfliC H7F</td>
<td>TUVΔfliC cis complement fliC, from TUV93-0</td>
<td>This study</td>
</tr>
<tr>
<td>TUVfliC H6F</td>
<td>TUVΔfliC cis complement fliC, from E2348/69</td>
<td>This study</td>
</tr>
<tr>
<td>ZAP1574</td>
<td>TUV93-0 ΔmotA</td>
<td>This study</td>
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<tr>
<td>ZAP1575</td>
<td>TUV93-0 motA−</td>
<td>This study</td>
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<tr>
<td>ZAP1576</td>
<td>TUV93-0 motA-complemented</td>
<td>This study</td>
</tr>
<tr>
<td>ZAP734</td>
<td><em>E. coli</em> O157:H7 NCTC 12900, Stx− AHVLA</td>
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<tr>
<td>E2348/69</td>
<td><em>E. coli</em> O127:H6; sequenced [68]</td>
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<tr>
<td>MG1655</td>
<td><em>E. coli</em> K12; sequenced [69]</td>
<td></td>
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<tr>
<td>TUV ΔLEE1-2</td>
<td>8.9 Kb deletion in LEE1-3, ETTA1 [70]</td>
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<td>SL1344</td>
<td><em>S. Typhimurium</em> [71]</td>
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<tr>
<td>SL1344Δβ/β2</td>
<td>SL1344 Δβ/β2 by λred, expresses only P1 [71]</td>
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<tr>
<td>SL1344Δβ/C</td>
<td>SL1344 Δβ/C by λred, expresses only P2 [71]</td>
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<tr>
<td>SL1344Δβ/C Δβ/β2</td>
<td>SL1344 Δβ/C and Δβ/β2 by λred, aflagellate [71]</td>
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<td>ZAP1566</td>
<td><em>S. Typhimurium</em> LT2 fliMS978-GFP motAB::TcR, non-motile P. Aldridge Lab, Newcastle</td>
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<td>SL1344ΔmotA/AB</td>
<td>SL1344 P22 transduced with ZAP1566, non-motile This study</td>
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<tr>
<td>Maskan</td>
<td><em>S. Typhimurium</em> Stock from Lohmann Animal Health</td>
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<tr>
<td>4/74</td>
<td><em>S. Typhimurium</em> [72]</td>
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<tr>
<td>4/74 prgH</td>
<td>prgH::mini-Tn 5Km2 mutant of ST4/74 NalR [73]</td>
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were then sheared for 2 min on ice with at maximum speed with an IKA T-10 homogenizer (Ultra-Turrax). Two to five rounds of centrifugation at 4100 g at 4 °C for 15 min, followed by centrifugation once at 16000 g at 4 °C for 10 min, were undertaken to obtain bacteria-free supernatants. These were centrifuged at 145000 g 4 °C for 1.5 h and resuspended in PBS (or 50 mM Tris/HCl pH 7.5 for actin polymerization assays) at 1:500 of initial culture volumes. Protein concentration and purity was assessed using BCA, and then confirmed by Coomassie staining (Imperial protein stain, Thermo Fisher) and densitometry after SDS-PAGE.

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(EBL; laboratory stocks), murine fibroblast (3T3; laboratory stocks) and HEK293 (laboratory stocks) cell lines were maintained in Dulbecco’s minimal essential medium (DMEM; Sigma) supplemented with 10% (v/v) foetal bovine serum (FBS; Sigma), 1 U of penicillin (Invitrogen), 1 µg ml⁻¹ of streptomycin (Invitrogen) and 2 mM L-glutamine (Invitrogen), unless otherwise stated. The porcine intestinal epithelial cell line IPEC-J2 was maintained as described by Schierack et al. [36] in HAMS/F-12 1:1 with 5% FBS, 1 U of penicillin and 1 µg ml⁻¹ of streptomycin. Prior to bacterial infection, cultured cells were washed in MEM/HEPES and incubated at 37 °C, 5% CO₂, 80% humidity for 1–2 h. All bacterial infections were incubated in these conditions for the time specified unless otherwise stated.

**Primary cell culture**

Bovine rectal and porcine colonic epithelia were isolated from abattoir-derived tissue as described previously [37]. The intestinal tissues were collected from material discarded as part of the normal work of the abattoir; while no licence was required for this, the relevant permissions were obtained from the West Lothian abattoir. The mucosal scrapings from bovine terminal rectal or porcine colonic tissue from a local abattoir were digested in DMEM [1% (v/v) foetal calf serum (FCS), 100 U ml⁻¹ penicillin, 30 µg ml⁻¹ streptomycin, 25 µg ml⁻¹ gentamicin] containing 75 µl ml⁻¹ collagenase and 20 µg ml⁻¹ dispase (Roche) with gentle shaking at 37 °C until isolated crypts could be observed microscopically. Crypt enrichment from undigested contaminating gut microflora and single crypts could be observed microscopically. Crypt enrichment from undigested contaminating gut microflora and single cells including fibroblasts was performed using a series of differential centrifugation steps with DMEM containing 2% (w/v) sorbitol. Approximately 500–700 crypts were seeded per well on to collagen (Nutacon for bovine cultures, Porcogen for porcine cultures) coated plates. The bovine cells were grown for 5 days before use, and the porcine cells were grown to a stage of confluence (approximately 3×10⁵ cells per well, typically at 10–14 days following initial primary epithelial cell culture).

**Confocal microscopy**

Unless otherwise stated, all samples for confocal microscopy outlined below were washed three times in PBS, fixed for >20 min at RT in 4% (w/v) paraformaldehyde and then permeabilized for <1 min with 0.1% (v/v) Triton X-100 prior to immediate washing as above and staining as below. All stains and antibodies were diluted in PBS. Where stained, cell nuclei were stained with DAPI (Table S3), and slides or coverslips were mounted with ProLong Gold (Invitrogen) unless otherwise stated. Image data were acquired with a Zeiss Plan Apochromat 1.4 NA ×63 oil immersion lens and a multi-track (sequential scan) experimental set-up on a Zeiss LSM510, using Axiovision software, unless otherwise stated.

For the examination of H7 flagella interactions with actin, primary bovine rectal epithelial and EBL cells were used. For primary epithelial interactions, bovine crypts isolated from rectal epithelia were seeded onto collagen-coated four-well Thermanox chamber slides as described (see primary cell culture section). Infection was for 1 h with 1×10⁶ E. coli O157:H7 TUV93-0 grown at 28 °C 200 r.p.m, sub-cultured to OD₅₆₀ 0.3–0.4 and resuspended in MEM/HEPES. These were then fixed, permeabilized and stained as described below, except that primary antibodies were labelled with FITC-conjugated α-rabbit IgG (Sigma). F-actin was then stained with Texas red-conjugated phalloidin diluted 1:250, after which cell nuclei were stained with DAPI diluted 1:1000.

Where EBL cells were used, they were seeded onto glass coverslips 24 h prior to infection with exponential-phase E. coli O157:H7 TUV93-0. TUV93-0 was grown in 0.3% (w/v) LB agar, sub-cultured into LB at 30 °C for 16 h 200 r.p.m, and then sub-cultured 1:100 in MEM/HEPES supplemented with 250 nm Fe²⁺ and 0.2% (w/v) glucose at 37 °C to an OD₅₆₀ 0.3. EBL cells were infected with TUV93-0 at a multiplicity of infection (m.o.i.) of 20 for 3 h. Samples were treated with α-H7 rabbit IgG diluted 1:100, then α-O157 mouse IgG diluted 1:1000, followed by a co-incubation of Alexa Fluor₄₈₈-conjugated α-rabbit IgG and Alexa Fluor₅₄₆-conjugated α-mouse IgG, both diluted 1:1000. F-actin was stained with FITC-conjugated phalloidin (Molecular Probes) diluted 1:100 for 2 h. The cell nuclei were stained with DAPI diluted 1:5000.

For the investigation of S. Typhimurium flagella interactions with host epithelia, confluent IPEC-J2 monolayers on glass coverslips were washed three times with MEM/HEPES immediately prior to infection with S. Typhimurium SL1344. Bacterial cultures were incubated in LB with 300 mM NaCl at 37 °C 200 r.p.m for 16 h and then diluted to an OD₅₆₀ of 0.3 in MEM/HEPES. IPEC-J2 cells were infected at an m.o.i. of 20 for 20 min. Samples were fixed with 2% (w/v) PFA for 20 min and then permeabilized with 0.2% (v/v) triton X-100 for 5 min. Samples were treated with P1- or P2-specific rabbit IgG (α-H1 diluted 1:100 and α-H2 diluted 1:500, respectively) and then α-O4 rabbit IgG diluted 1:500. This was then labelled with FITC-conjugated α-rabbit IgG diluted 1:1000. F-actin was stained with Texas red-conjugated phalloidin diluted 1:500 and slides were mounted in VectorShield (Vector Labs). Membrane staining of S. Typhimurium-infected cells was undertaken in the same way, except that cells were infected for 10 min before fixation, and prior to permeabilization, membranes were stained with Texas red-conjugated wheat-germ agglutinin diluted 1:1000 for 2 h [38]. Additionally, instead of Texas red-conjugated phalloidin, F-actin was stained with Alexa Fluor₄₈₈-conjugated phalloidin.

For the investigation of H7 flagella interactions with host cell membranes, HEK293 cells were transfected with a CRD–STREX–GFP construct that leads to fluorescent staining on the inner side of the plasma membrane [39]. Briefly, 5×10⁴ HEK293 cells were seeded onto glass coverslips in DMEM with 10% (v/v) FBS, 100 U ml⁻¹ penicillin and 30 µg ml⁻¹ streptomycin, and cultured to 40–70% confluence. One hundred and fifty nanograms of CRD–STREX–GFP plasmid was diluted in proprietary EC buffer (Qiagen) and enhancer was added to it at 1:125 (µl ng⁻¹ DNA) and incubated for 5 min at RT. Effectene (Qiagen) was added at a 3:1 ratio to the enhancer and incubated for 10 min at RT. HEK293 cells
were washed with PBS and fresh maintenance medium was added. The DNA-Effectene mixture was diluted 1:5 in maintenance medium and added to cells 1:2. Protein expression was determined 24 h post-transfection using GFP expression as a read-out. These cells were infected with E. coli O157:H7 TUV93-0 for 90 min. Fixed then permeabilized samples were labelled with α-H7 rabbit IgG then α-O157 rabbit IgG (Table S3) diluted 1:1000 and 1:100, respectively. Primary antibodies were labelled with Alexa Fluor<sub>488</sub>-conjugated α-rabbit IgG diluted 1:1000. Volumetric 3D reconstructions of deconvolved images were undertaken using Velocity Visualization software version 6.3.

For all of the above confocal microscopy, image data were acquired at optimal z-slice sampling rates as determined by Zeiss software, with a 1024×1024 pixel image size. Image data were deconvolved using Huygens software (Scientific Volume Imaging, Netherlands). Deconvolved models of image data were analysed and reslices of z-stacks, montages and projection views were created using NIH ImageJ software, with final figures assembled in Adobe Photoshop CS5 and CS6.

Co-incidence analysis was undertaken using the 24 October 2004 version of an open-source co-localization plug-in downloadable from http://rsweb.nih.gov/ij/plugins/colocalisation.html. Threshold values were set based on z-stack histograms to determine the Gaussian curve of background readings for each channel, and co-incidence was output at a display value of 150. Display values were designated co-incident, not co-localized, as total co-localization was <50%.

For pre-/post-permeabilization labelling, collagen (rat tail type 1, Sigma)-coated coverslips were coated in 35% (v/v) ethanol for 4 h at 37°C, washed in PBS and then media, and then seeded with IPEC-J2 cells for Salmonella infection and EBL cells for E. coli O157:H7 infection 24 h prior to infection. Cells at 50–80% confluence were washed with PBS and then incubated with MEM/HEPES for 1–2 h prior to infection with 1×10<sup>5</sup> cells of mid-exponential phase S. Typhimurium SL1344::pAJR145 (constitutively expressing eGFP) for 20 min, or E. coli O157:H7 TUV93-0::pAJR145 (constitutively expressing eGFP) for 1 h, respectively. Once infections had taken place, cells were fixed in 4% (w/v) PFA. Cells were labelled 1:500 with α-FliC rabbit polyclonal IgG and then 1:1000 with FITC-conjugated α-rabbit IgG (Table S3). Dishes were not mounted but instead imaged in PBS using a Leica SP5II confocal laser scanning microscope to determine the locations of cell-associated flagellated bacteria on the four reference grids.

Cells were fixed and embedded in preparation for electron tomography as follows. All wash steps and treatments were diluted in ddH<sub>2</sub>O, unless otherwise stated. Fixation was undertaken in 2.5% (w/v) glutaraldehyde (100 mM sodium cacodylate pH 7.2) for 20 min and then washed three times before post-fixing with 1% (w/v) osmium tetroxide for 20 min. After a further three washes, cells were then stained with 3% (w/v) uranyl acetate for 20 min and washed three more times. Cells were then dehydrated with graded ethanol washes for 5 min each at 70, 80, 90, 96 and 100% (v/v) before the addition of 1:1 epon epoxy resin for 1 h 30 min, with rocking, for embedding. Embedded samples were then baked at 200°C for 5 days before removal from glass dishes with snap-freeze/thawing. The reference grids on the remaining resin were then mounted onto stubs, trimmed to areas of interest determined by confocal microscopy (see also Olmos et al. [40]), and cut into 300 nm thick sections using a Leica EM UC7 with an IC80HD camera. Serial sections of areas of interest were placed onto film-coated copper slot grids (Agar Scientific) and further stained with 3% (w/v) uranyl acetate for 20 min to enhance contrast before imaging. Washed grids were labelled with 15 nm gold particles both to the top and bottom of sections for 5 min and then blotted dry.

Samples were mapped to determine the location of flagellated and cell-associated bacteria using a FEI 120kV BioTwin Spirit transmission electron microscope (TEM). The TEM tilt series (from −60° to +60 in 1.5° increments) were acquired using a FEI 200kV twin lens scanning transmission electron microscope and Xplore3D software (FEI). The tilt series were reconstructed into electron tomograms using the standard workflow of IMOD and 3dMOD software packages (Boulder Laboratory for 3D EM of Cells, University of Colorado, USA) and then analysed and presented using Amira 6.0.0, where an anisotropic diffusion filter was applied to the data, and FIJI software.

**Immuno-gold staining and scanning electron microscopy (SEM)**

The ultra-structural details of flagella interaction with specified epithelial cell types were visualized using Hitachi 4700 Field Emission Scanning Electron microscope. The specimens were fixed in 3% gluteraldehyde in 100 mM sodium cacodylate...
buffer (pH 7.4) and processed without permeabilization for SEM as described previously [37]. Immuno-gold labelling was visualized by detecting back-scattered electrons; overlays of back scattering and secondary electrons were false-coloured in Adobe Photoshop CS4 for contrast.

**Haemolysis assays**

All bacterial cultures were inoculated with agar plugs from the leading edge of the motility halo of fresh motility plates of the strains indicated. These cultures were incubated statically in LB at RT for 16 h. Cultures were adjusted to equal OD600 before addition to a V-bottomed 96-well plate at 1:1 (50 µl each) with 50% (v/v) washed sheep red blood cells (RBCs; Oxoid) in PBS. The bacteria were driven into contact with the RBCs by centrifugation at 2000 g for 5 min RT, followed by gentle mixing by pipetting. Plates were then centrifuged 2000 g for 10 min RT to pellet RBCs. The A405 of 100 µl of supernatants mixed by pipetting. Plates were then centrifuged 2000 g for 5 min RT, followed by gentle mixing by pipetting. Plates were then centrifuged 2000 g for 10 min RT to pellet RBCs. The A405 of 100 µl of supernatants was then measured as a read-out of total haem release. Independent replicates were undertaken three–five times, with three–four technical replicates. The mean level of haem release caused by incubation with LB alone was subtracted and data were normalized as a percentage of wild-type haem release. After tests for equal variance were performed on raw data for each strain set in Minitab 17, and homoscedastic two-tailed t-tests were performed on the raw data, between mutant and wild-type strains.

**Preparation of cell lysates**

Bovine primary rectal epithelial cells were harvested by washing twice in PBS followed by incubation with TripLE Express (Gibco) at 37°C, 5% CO₂, 80% humidity for 10 min. An equal volume of PBS was added and then cells were scraped off, collected and centrifuged at 300 g for 2 min at RT. Cells were washed twice by resuspending cell pellets and centrifuging in PBS or Hank's balanced salt solution (HBSS). The cell pellet was resuspended in PBS or HBSS, subjected to five cycles of snap-freezing in ethanol and dry-ice and thawing at RT, and clarified by centrifugation at 18,000 g for 10 min at 4°C.

**Western and far-Western blotting**

Eighty micrograms of primary epithelial cell lysates or 1 µg purified receptor candidates separated in SDS-PAGE gels were Western-blotted onto nitrocellulose (Amersham, GE Healthcare) with Schaeffer–Nielsen buffer [48 mM Tris, 39 mM glycine, 20% (v/v) methanol, 0.04% (v/v) SDS, pH 9.2], at 15V for 30–60 min. Blots were blocked in Carbofree (Vector Labs) or 0.1% (v/v) Tween20 (Sigma) in PBS (PBST) with 5% (w/v) skimmed milk powder (Sigma, Marvel) at RT for 2 hr or 4°C for 16 h. Blots were washed three times between all subsequent steps in PBST alone for 15 min at RT, with rocking. For both Western and far-Western blots, antibodies were diluted in 1% (w/v) skimmed milk powder in PBST and incubated with blots for 1 h, with rocking at RT. Prior to antibody labelling, far-Western blots were first probed with 1 µg ml⁻¹ H7 flagella in PBS for 3 h, with rocking at RT. H7 flagella were then treated with α-H7 rabbit IgG diluted 1:1000, and this was labelled with horseradish peroxidase (HRP)-conjugated α-rabbit IgG diluted 1:1000. Western blots of cell lysates were probed with α-cofilin-1 mouse IgG diluted 1:500 or α-galectin-4 goat IgG diluted 1:1000, and then a 1:1000 dilution of HRP-conjugated α-mouse or goat IgG, respectively. Blots were then developed using Pico-West SuperSignal ECL reagents (Thermo Fisher) in a G-box (Syngene), and captured using GeneSnap (Syngene).

**Pull-down assays**

Cyanogen bromide (CnBr)-activated sepharose 4B lyophilized beads (GE Healthcare) were used according to manufacturer’s instructions. Fifty micrograms of H7 flagella was added 2:1 to CnBr beads for 16 h at 4°C, with rocking, in coupling buffer (100 mM NaHCO₃, pH 8.3 containing 500 mM NaCl). Excess H7 flagella were washed off beads by five cycles of centrifugation at 18,000 g for 30 s followed by resuspension of beads in equal volumes of coupling buffer. Following one additional centrifugation as above, beads were resuspended in blocking buffer (100 mM Tris/HCl, 500 mM NaCl, pH 8.0) and incubated for 2 h statically at RT. CnBr beads were then washed 10 times by centrifugation as above, with alternate cycles of suspension in coupling buffer or wash buffer (100 mM acetic acid, 100 mM sodium acetate, 500 mM NaCl, pH 4.0). On the final centrifugation, beads were suspended in 180 µg BTRE freeze-thawed cell lysate in HBSS and incubated for 16 h at 4°C, with rocking.

CnBr beads were centrifuged at 18,000 g for 30 s and then washed by three cycles of centrifugation as above followed by resuspension in 0.1% (v/v) PBST. Beads were centrifuged as above and eluted by incubation in 2× SDS-PAGE sample buffer (Sigma) for 5 min at 100°C and centrifuged as above. Eluted proteins were then analysed by 12% SDS-PAGE and Imperial protein staining (Thermo Fisher). Bands visible only in the presence of H7 flagella-coated CnBr beads were considered to be likely ligand candidates and equivalent cell lysate bands were excised and identified by mass spectrometry.

**Mass spectrometry**

Mass spectrometry, tandem mass spectrometry and peptide mass fingerprinting were undertaken by Kevin McLean at the Moredun Research Institute Proteomics Facility. UK. Protein bands from Imperial protein-stained SDS-PAGE gels were excised and delivered to the proteomics facility. Here they were trypsinized, enriched and cleaned up, and matrix-assisted laser desorption/ionization time-of-flight tandem mass spectrometry (MALDI-TOF MS/MS) was performed on a Bruker UltraflexII according to the established protocols. Spectra were then input into MASCOT peptide mass fingerprinting software (Matrix Science) as monoisotopic 1+ (m/z) ratios using a±50 p.p.m. tolerance to generate protein identities.
Accurate MW determination of H6, H7, H48, P1 and P2 was undertaken with online HPLC-MS in the proteomics and metabolomics facilities at the Roslin Institute, UK. Samples of sheared flagella were diluted to ~1 pmol µl⁻¹ in 0.1% (v/v) formic acid and ~20 pmol was applied to a microbore HPLC column (Dionex Acclaim C18, 4.6 mm i.d., 150 mm length, 5 µm beads, 120 Å pore size) pre-equilibrated with 0.1% (v/v) formic acid by use of a Ultimate HPLC system (Dionex). Bound components were eluted with a gradient of 0.1% (v/v) formic acid in acetonitrile into the electrospray source of an amaZon ETD ion trap mass spectrometer (Bruker Daltonics, Germany). The mass spectrometer acquired full-scan mass spectra with the final spectra being an average of eight trap fills with a maximum averaging time of 200 ms. Signals corresponding to intact flagellin were summed, the raw data were smoothed and the background was subtracted, and deconvolution was performed using the Bruker proprietary algorithm.

Enzyme-linked immunosorbent assays (ELISAs)

One microgram per well of 95% pure human platelet βγ-actin, recombinant human galectin-4 and recombinant human cofilin-1 were adsorbed to 96-well Nunc Maxisorb plates in 100 mM sodium bicarbonate pH 9.6 for 16h at 4°C. Wells were blocked using Carbo-free (VectorLabs) for 2h, at RT, before washing. All wash steps involved washing three times in PBST (0.1% Tween20). Polymers and monomers of each flagella type (as verified by size-exclusion chromatography) were diluted in PBS only at 1000, 500, 100 and 0 ng per well, in specific wells, for 3 h, before washing. Antibodies were diluted in PBST and incubated in wells for 1 h, before washing. H6, H7, P1 and P2 flagella-specific rabbit IgG (α-H6 and α-H7 1:1000; α-H1 1:100; α-H2 1:500, respectively, Table S3), followed by HRP-conjugated α-rabbit IgG, were used to label flagella-specific binding.

Flagella binding was detected using Pico-West SuperSignal ECL reagents (Thermo Fisher) in a G:Box using GeneSnap (Syngene) at a fixed distance of 575 mm. Densitometry on Coomassie-stained SDS-PAGE gels was used to adjust flagella values.

Assays were carried out in black opaque 96-well plates (Nunc). First, 1 µm actin was added to wells with 50 mM Tris/HCl pH 7.5. Cofilin-1 and serially diluted flagella preparations were then added, followed by polymerization buffer (at a final concentration of 5 mM KCl, 0.2 mM MgCl2 and 0.1 mM ATP). Samples were excited at 365 nm, and emission data were measured at 407 nm for 1 h, at 30 s intervals. A daily gain value was applied to all wells to normalize actin polymerization capability and this was calculated using pyrene-actin fluorescence after 1 h polymerization at RT as above.

The maximum velocity (Vmax) of actin polymerization was calculated using the formula Vmax=(A−A0)/(T−T0), where A=absorbance at 407 nm and T=time. Differences due to the addition of flagella were assessed for statistical significance using GLMs in Minitab 16.2.4. In these models, Vmax was the response and replicate was a random factor. With flagella titration experiments, GLM analysis was carried out on data from five independent experiments; flagella type and concentration were fixed factors. Post-hoc Tukey pairwise comparisons of the interaction between flagella type and concentration were carried out with flagella±cofilin-1 experiments, GLM analysis was carried out on data from three independent experiments; flagella type and concentration were fixed factors. Post-hoc Tukey pairwise comparisons of the interaction between cofilin-1 concentration and flagella type were carried out as a very conservative estimate of differences based on the set of assumptions in a GLM statistical model. Therefore, if P values are <0.05, they are very likely to be valid and so were taken as statistically significant.

Actin polymerization assays

Actin polymerization assays were carried out as described by Sithidet et al. with minor modifications [41] and informed by Van Troys et al. [42]. G-actin was prepared by resuspending lyophilized pyrene-conjugated rabbit skeletal muscle αβ-actin in G-buffer (10 mM Tris/HCl, 200 µM CaCl2, 200 µM ATP, 1 mM DTT, pH 7.5) for 1 h on ice, in the dark. This was then centrifuged at 100000 g for 2h at 4°C. The top 80% supernatant was kept on ice in the dark. Test proteins were resuspended as purified or dialyzed with U-tube concentrators (Novagen) into 50 mM Tris/HCl pH 7.5. Molar concentrations of all protein components were calculated using a Bradford assay with a BSA standard curve. Densitometry of Coomassie-stained SDS-PAGE gels was used to adjust flagella values.

Statistical analysis was carried out in R on log10 data from four independent experiments. Data residuals were assessed for normality using a normality plot. As it was not biologically appropriate to compare different flagella types or ligand types statistically, separate linear mixed effects models of relative binding were used to assess the statistical significance of each flagella type binding to each ligand candidate (galectin-4, cofilin-1 and actin). Flagella concentration (1000, 500, 100 ng/well) and polymerization status (polymer and monomer) were fixed effects and experiment was a random effect. Pairwise comparisons of the interaction between polymerization statuses for each flagella type in each model were then performed on the data. To correct for the large number of comparisons, an alpha level of P<0.01 was taken as significant.

Size-exclusion chromatography

Samples were applied to an HPLC column (Sigma TSK G4000SWXL, 4.6 mm i.d., 300 mm length, 8 µm beads, 450 Å pore size) pre-equilibrated with 100 mM Tris/HCl, 200 mM NaCl, 1 mM DTT, pH 8.0, by use of an Ultimate HPLC system (Dionex), measuring absorbance at 220 nm. Columns were
calibrated by the maximum peak of elution of individual molecular weight standards: blue dextran (2.5 MDa) at ~5 ml, BSA (66 kDa) at ~10 ml, and ribonuclease A (13.5 kDa) and ATP (0.5 Da) at ~12 ml.

For verification of flagellin polymerization status, polymeric ATP (0.5 Da) at ~12 ml.

RESULTS

**Bacterial flagella labelling can be coincident with host F-actin staining**

In experimental systems that allow bacteria to come into contact with host cells without centrifugation, co-incidence of flagella and F-actin labelling was routinely detected, 60 min post-infection for *E. coli* O157:H7 (Figs 1, S1 and Movie S1) and 20 min post-infection for *S. Typhimurium* (Fig. 1c). For *Salmonella*, co-incidence of flagella with F-actin labelling raised the possibility that flagella were not necessarily confined to the *Salmonella* containing vacuole (SCV) during invasion. It was also possible to observe flagella expressed by extracellular *Salmonella* co-incident with actin at early infection time points. However, not all flagella that were imaged (bacterially associated or not) were found to be coincident with phalloidin-stained actin (Fig. 1d). Where flagella were coincident with F-actin staining, this followed a striated pattern, consistent with the helical wave of flagella filaments.

**Bacterial flagella labelling is intermittently masked by membranes**

Coincidence of bacterial flagella with F-actin staining (Fig. 1) was detected after samples were fixed with 4% (w/v) PFA and then permeabilized with 0.1% (w/v) Triton X-100 (Tx100). Depending on the level of proximity of bacterial flagella to cortical actin, this could involve some contact with the plasma membrane. Biological membranes are sensitive to non-ionic detergents such as Tx100. To observe whether Tx100 treatment altered flagella labelling, confocal microscopy was carried out on GFP-expressing *E. coli* O157:H7 and *S. Typhimurium* SL1344-colonized epithelial cells. Cells were fixed and then flagella were labelled with specific antibodies before (red) and after (green) Tx100 treatment. F-actin was then labelled with Alexa Fluor 647 phalloidin (blue, see the Methods section). Cell-associated flagellated *S. Typhimurium* and *E. coli* O157:H7 were imaged in 3D stacks and representative projections are presented in Fig. 2. Pre-permeabilization staining of flagella was often interrupted or less intense in short portions of the flagella, where post-permeabilization staining of the same flagella was comparatively even (Fig. 2). This indicates that bacterial flagella were intermittently masked by a detergent-sensitive component, such as a membrane, during the initial stages of infection.

**Bacterial flagella are detectable on both sides of host cell membranes**

To investigate how close flagella associations with host cell surfaces might be, flagellated adherent *E. coli* O157:H7 and *S. Typhimurium* were imaged in the context of plasma membrane labelling. For *E. coli* O157:H7 infections, HEK293 cells were transiently transfected with a trans-membrane voltage-gated ion channel fusion protein. This ion channel contains a green fluorescent protein (GFP) in the C-terminal cytoplasmic domain. The fluorescent signal generated reports the position of the internal cytoplasmic face of the plasma membrane only [39]. H7 flagella were imaged passing in and out of the GFP-labelled cytoplasmic face of the plasma membrane (Fig. 3a, c). Volumetric 3D reconstruction of the labelling in these confocal microscopy images allowed the visualization of transverse sections, showing H7 flagella labelling within the cell cytoplasmic boundaries (Fig. 3b, d).

In contrast, wheat germ agglutinin binds to the external sugar-coated surface of epithelial cell membranes [38]. Confocal microscopy of eukaryotic plasma membranes, labelled with fluorescent wheat germ agglutinin after *S. Typhimurium* infection but before fixation, revealed flagella bundles of invasive *S. Typhimurium* passing through regions of plasma membrane staining (Fig. 3e). Flagella were observed passing both from inside-to-outside and outside-to-inside host cell areas. Both *E. coli* O157:H7 and *S. Typhimurium* flagella were observed passing through gross cellular boundaries by confocal microscopy.

**Defining bacterial flagella interactions with host cell membranes by electron microscopy**

The z-plane resolution limit of confocal laser scanning microscopy is ~500 nm with the fluorophores used, but host cell plasma membranes are only ~5 nm thick. O157:H7 immuno-gold SEM was used to take a closer look at H7 flagella interactions with host cell surfaces without Tx100 treatment. As with confocal microscopy of pre-permeabilization labelling, *E. coli* O157:H7 flagella immune-gold staining was interrupted as the filament disappeared at the primary intestinal epithelial cell surface, and then resumed as it curled upwards away from the surface (Fig. 4a, b). SEM of *S. Typhimurium* at early time points of infection (10–30 min, again without Tx100 treatment) also revealed flagella-like filaments interacting intimately with cell surfaces of primary intestinal epithelial cells (Fig. 4c). These filaments could be seen disappearing and reappearing at the cell surface, consistent with the actin coincidence observed by confocal microscopy. *S. Typhimurium* flagella-like filaments were also observed associating with membrane ruffles and protrusions during the invasion process (Fig. 4d, e).

These observations were made with serologically and structurally distinct flagella (Fig. S2), hinting that a biophysical mechanism may be involved. To look for evidence of
Fig. 1. Position of *E. coli* O157:H7 and *S.* Typhimurium flagella relative to host cell F-actin during early colonization, as determined by confocal microscopy. Phalloidin-labelled actin is red, antibody-labelled LPS and flagella are green, and coincidence of the two is yellow. (a, b) Confocal micrographs of *E. coli* O157:H7 (TUV93-0) interacting with primary bovine epithelial cell cultures 1 h post-infection. (a) Interaction of *E. coli* O157:H7 with host actin with its H7 flagellum in XY, XZ and YZ projections (3D rotating image in Movie S1, individual Z-slices of XZ and YZ projections are presented in Fig. S1). (b) Co-incidence of phalloidin-stained actin with an *E. coli* O157-associated coiled H7 flagellum, but not with the bacterium itself, by XY projection, with individual XZ slices shown beneath. (c, d) Confocal micrographs of O4:P2 stained *S.* Typhimurium (SL1344) interacting with IPEC-J2 cells 20 min post-infection. (c) XY and orthogonal projections of a 3D micrograph where a flagellum emanating from a non-invasive bacterium coincident with actin staining is indicated by an arrow. (d) XY projection of a 3D confocal micrograph of P2 flagella (green) adhering to the cell surface. Not all P2 flagella are coincident with actin, but where there is co-incidence, it is periodic. All samples were fixed with 4% (w/v) paraformaldehyde for >20 min prior to permeabilization. Scale bars, 1 µm unless indicated otherwise.
biophysical interactions of flagella with host cell membranes, \textit{S. Typhimurium} flagella closely associated with host cells were located with correlative light and electron microscopy and examined by thick-section transmission electron tomography (Fig. 5). To preserve the details and visibility of the host cell membranes, conventional post-section staining of flagella was not undertaken, but the visibility of the structures in the tomogram slices was enhanced by applying an anisotropic diffusion filter and a blue–orange ICB colour lookup table in Amira software.

Host cell membranes were observed encapsulating bundles of flagella filaments during the \textit{S. Typhimurium} invasion process (Fig. 5a, b, Movies S2 and S3). Consistent with previous microscopy, these flagella were located within gross cellular boundaries at points, but they were separated from the host cytosol by deformed but contiguous host membranes. Visually distinct from these observations was the reduction or interruption of clear membrane staining in proximity to bacterial flagella (Fig. 5c, Movie S4). In this example, the flagella filaments are largely accessible to extracellular antibo body labelling, as shown by the punctae that occur regularly along their length. Fig. 5c2 shows the labelled bundle of flagella filaments drawing close to a clearly defined plasma membrane, but as they come into presumable contact, the membrane boundary is both deformed and less distinct, and in places apparently absent.

**Bacteria with paralysed flagella cause less membrane disruption**

To test whether flagella rotation is even capable of causing physical disruption of host cell membranes, sheep RBCs were used as a model dye-filled eukaryotic plasma membrane, as they contain haem pigment. Short-term incubation of RBCs was undertaken with bacteria and their isogenic flagella and flagella motor (\textit{mot}) mutants; \textit{mot} mutants result in the production of full-length flagella that do not rotate. Bacteria were cultured under conditions known to induce expression of flagella and briefly centrifuged into contact with RBCs to eliminate the contribution of motility and chemotaxis to this process (Fig. 6; see the Methods section).

All flagellate wild-type strains caused membrane disruption upon interaction with RBCs, as determined by haem release. As haem release could be due to the T3SSs of either pathogen, T3SS mutants (\textit{ΔprgH} for \textit{S. Typhimurium}, \textit{ALEE1-2} for \textit{E. coli} O157:H7; Table 1) were tested under these flagella-selective growth conditions (Fig. 6). Mutation of T3SSs resulted in altered mean levels of haem release compared to wild-type for both \textit{S. Typhimurium} (\approx 30\% reduction) and \textit{E. coli} O157:H7 (\approx 10\% increase). Both effects were statistically significant (\textit{P}<0.0001 and 0.0039, respectively), suggesting that the T3SSs were directly involved in haem release.

Mutation of flagella and flagella rotation caused reduced mean levels of haem release for both \textit{S. Typhimurium} and \textit{E. coli} O157:H7. For \textit{S. Typhimurium}, the levels of haem release were approximately half that of the wild-type in both flagellin (\textit{ΔfljB}) and motility (\textit{ΔmotAB}) mutants, with statistically significant values (\textit{P}=0.0001 for both; Fig. 6). For wild-type \textit{E. coli} O157:H7, mean levels of haem release were much less affected by flagellin (\textit{ΔfljC}) and motor (\textit{ΔmotA}) mutation. Reductions of \approx 5\% for \textit{ΔfljC} were not statistically significant (\textit{P}=0.17). In contrast, a reduction of \approx 10\% for \textit{motA} was statistically significant (\textit{P}=0.0046). The subtle effect for \textit{E. coli} O157:H7 may reflect the lower overall flagella expression and the lower rotation speeds and motility of \textit{E. coli} wild-type compared to \textit{Salmonella} (Fig. S3 [43, 44]). We infer from these results that the flagella of \textit{S. Typhimurium} and \textit{E. coli} O157:H7 are physically able to disrupt eukaryotic plasma membranes and flagellar rotation is likely to play a role.

**Bacterial flagella can interact with components of the actin cytoskeleton in vitro**

If bacterial flagella can disrupt host cell membranes, perhaps they are also able to interact with cytosolic components just beneath host cell membranes. Initial pull-downs and far-Western blotting with \textit{H7} flagella identified β-actin (ACTB1), cofilin-1 (CFL1) and galectin-4 (GAL4) as potential interactants (Fig. S4). A ‘far’ ELISA was designed to assess relative flagella binding to human β-actin, recombinant human CFL1 and recombinant human GAL4. These purified proteins...
Fig. 3. Position of *E. coli* O157:H7 and *S. Typhimurium* flagella relative to host cell membranes, as determined by confocal microscopy. (a, c) Projections of 3D confocal micrographs of *E. coli* O157:H7 (TUV93-0: LPS and flagella are green) interacting with HEK293 cells transiently transfected with a transmembrane protein–fluorescent protein fusion construct (red, see the Methods section) 1.5 h post-infection. This transfection causes labelling of the cytoplasmic face of the plasma membrane cytosolic leaflet [39]. (b, d) 3D volume-rendered model of images in (a) and (c), respectively, using Volocity Visualization software. Arrows indicate intracellular flagella. (a1) XY and XZ projections of a single flagellum emanating from a bacterium on a transfected HEK293 cell, 3D-rendered in (b1) and enlarged in (a2). A similar view has been 3D-rendered in (b2), where the arrow indicates intracellular flagella. (c) XY projection of the binding of a cluster of bacteria expressing H7 flagella to a transfected HEK293 cell. (d) Transverse cut-through of the HEK293 cell, with flagella visible beneath the layer defined by the cytoplasmic staining of the labelled ion channel protein (arrow). (e) Projections of a 3D confocal micrograph of O4:P2 stained *S. Typhimurium* SL1344 (green) interacting with wheat germ agglutinin-labelled IPEC-J2 cells (red) 10 min post-infection. This labelling stains N-acetylglucosamine and sialic acid moieties on the external face of cell membranes [38]. (e1) YZ projection of the *XY* inset labelled in (e2), with a long flagella bundle that passes out then back into the cell (arrow). Coincidence of flagella and membrane staining is shown in yellow. (e2) XY projection of whole image shown in (e1). All samples were fixed in 4% (w/v) para-formaldehyde for >20 min prior to permeabilization. Scale bars, 5 µm unless indicated otherwise.
Fig. 4. SEM of *E. coli* O157:H7 and *S. Typhimurium* flagella-like structures disappearing and reappearing at host cell surfaces. (a, b) *E. coli* O157:H7 (ZAP734) interactions with bovine primary epithelial cell cultures 3 h post-infection. (a) Secondary electron (SE) image of a snapped flagella-like filament, which appears to penetrate the cell surface (arrows). (b) A false-coloured SE image (cyan) superimposed with a false-coloured back-scattered image (red). H7 flagella and O157 LPS are both immuno-gold labelled. Filament staining occurs adjacent to the bacterium and is then absent (arrow 1). The filament is then broken and curls back on itself (arrow 2), with the remnant embedded filament (arrow 3). (c1, c2) SE images of flagella-like filaments disappearing into and coming out of the surface of IPEC-J2 epithelial cells (arrows) within the first 30 min of infection with *S. Typhimurium* (Maskan). (d1) SE image of *S. Typhimurium* (Maskan) micro-colonies on IPEC-J2 cells 40 min post-infection. (d1) Actin ruffling proximal to invading *Salmonella*. (d2) Enlarged from the inset indicated in (d1). Wavy flagella-like filaments are interacting with ruffled and unruflled cell surfaces (arrows). (e1) Long filaments disappearing into the cell surface (arrows). (e2) Higher resolution image of the area indicated in (b1), which shows long filaments interacting with a large macropinocytic protrusion (arrow). Samples were fixed in 3% (w/v) glutaraldehyde and were not permeabilized prior to sample processing for SEM. Imaging was undertaken on a Hitachi 4700 field emission scanning electron microscope. Scale bars, 1 µm unless indicated otherwise.

Bacterial flagella can increase actin polymerization rates *in vitro*

To further confirm the *in vitro* binding interactions of H6, H7, P1 and P2 polymeric flagellin with actin, their effect on actin polymerization was assessed (Fig. 8a). We used 1 mM pyrene-conjugated rabbit αβ-actin in 10 mM Tris/HCl, 200 µM CaCl₂, 200 µM ATP, 1 mM DTT, pH 7.5 as a baseline (actin control). Actin polymerization was initiated by the addition of final concentrations of 5 mM KCl, 0.2 mM MgCl₂, and 0.1 mM ATP (actin). The actin-severing activity of CFL1 at 2:1 ratio with actin generates more rapidly polymerizing ‘plus’ ends of actin [46], so 500 nM CFL1 was added as a positive control for enhanced rates of actin polymerization (cofilin-1). The bacterial flagella filaments tested showed all enhanced *in vitro* rabbit αβ-actin polymerization to variable extents, implying a direct interaction with physiologically active actin (Fig. 8a). Higher median actin polymerization Vₘₐₓ rates were determined for all flagella tested when compared to actin alone, with P2 flagella showing a statistically significant dose-dependent effect (*P* = 0.003, Fig 8a, b).

To investigate whether H6 and H7 flagella interactions with CFL1 enhanced the effect they had on actin polymerization, αβ-actin was co-incubated with the four flagella types and were coated to 96-well plates and probed with 1000–100 ng flagellins, and then binding was detected using appropriate antibodies. Binding data were normalized to negative (no flagellin added) and positive (wells coated with 1 µg flagellin) detection controls to give a measure of relative binding (see the Methods section). Binding of polymeric and monomeric flagellin from *E. coli* O127:H6, *E. coli* O157:H7, commensal *E. coli* K12:O175:H48 and *S. Typhimurium* O4:H1/H2 expressing either P1 or P2 flagella was assessed (Figs 7 and S2).

For all interactions tested, except for H48-CFL1 (*P* = 0.55), monomeric flagellins bound at significantly lower levels than polymeric flagellins (Fig. 7, *P* = 0.0023–0.0001 or less, depending on the flagellin tested), serving as a suitable baseline for non-binding in this assay. This was particularly the case for the actin interactions, where binding by flagellin monomers was virtually undetectable in this assay for each flagellin type tested.

There were different patterns of binding between *E. coli* and *S. Typhimurium* polymeric flagellins; *E. coli* flagellins showed higher relative binding to CFL1 compared to βγ-actin under these conditions, and conversely the *S. Typhimurium* flagellins tested showed a stronger association with βγ-actin, compared to CFL1. All polymeric flagellins tested bound to GAL4, although there was no particular distinction between *S. Typhimurium* and *E. coli* relative levels. Binding to GAL4 was not a consequence of flagella post-translational glycosylation; determination of molecular weights by mass spectrometry of H6 and H7 sheared preparations indicated that these flagellins were unmodified, whilst modified forms of *S. Typhimurium* flagellins were consistent with methylation (data not shown [45]).
Fig. 5. Electron tomography of host cell membranes in proximity to *S. Typhimurium* flagella-like structures. *S. Typhimurium* SL1344::pAJR145 interacting with IPEC-J2 cells 20 min post-infection. Fixation with 4% (w/v) paraformaldehyde was undertaken for >20 min prior to flagella labelling and samples were not treated with any detergents. Confocal microscopy was used to locate bacteria with flagella in contact with host cells and correlative electron tomography was undertaken (see the Methods section). The contrast of insets was enhanced using the blue_orange_ICB look-up table in Fiji software. (a1) Flagellated *S. Typhimurium* within an epithelial cell. (a2) Inset in (a1); the flagella cross the gross vacuole boundary but are present inside an intact membrane channel (arrow). (b1) Flagellated *S. Typhimurium* inside a membrane ruffle. (b2) Inset in (b1); the flagella bundle passes through the gross boundaries of the ruffle but is adjacent to a distinct membrane boundary (arrow). (c1) Flagellated *S. Typhimurium* at the point of induced uptake into an epithelial cell. (c2) Inset in (c1); flagella in proximity to the disrupted epithelial plasma membrane (arrow). (c3) Structures in (c2) are labelled for clarity. Scale bars, 500 nm unless indicated otherwise.
two different concentrations of CFL1 (Fig. 8c). Without CFL1, 250 nM of each flagella type was sufficient to cause a statistically significant increase in the $V_{\text{max}}$ of actin polymerization, with *S. Typhimurium* P2 flagella being the most effective. This low concentration of flagella was chosen to allow detection of increases in actin polymerization rates that may occur as a result of synergy between flagella and CFL1 interactions. Adding an equimolar concentration of CFL1 (250 nM) caused a trend towards further enhanced rates of actin polymerization, but this was not statistically different to addition of CFL1 alone for any of the flagella types tested ($P=0.124–0.856$). At a 2:1 molar concentration of CFL1:flagella, at which the CFL1 concentration was optimal, further increases in the $V_{\text{max}}$ of actin polymerization were apparent for all flagella tested. These increases were only statistically significant with P1 and P2 flagella, compared to the effect of CFL1 alone ($P=0.035$ and 0.004 respectively). Additionally, there was no statistically significant interaction between CFL1 concentration and flagella type ($P=0.6$ see the Methods section), indicating that the effects on $V_{\text{max}}$ were additive and not due to any synergistic effects of flagella and CFL1 on actin polymerization.

**DISCUSSION**

We have known for some time now that bacterial flagella are more than just motility organelles. The idea of flagella as long adherence multi-protein complexes for early host surface attachment by commensal and pathogenic bacteria alike is now widely accepted. This raises the question of why there are so few published examples of specific cell-surface protein receptors for flagella.

This paper presents the masking of *Salmonella* and *E. coli* flagella antigens by a detergent-sensitive component, which indicates very intimate interactions of these flagella with host cell lipids. Such a close interaction between flagella and membranes was borne out by confocal microscopy of membrane labelled samples. Additionally, flagella of *S. Typhimurium* and *E. coli* O157:H7 have been shown to bind to cholesterol and ionic sulpho- and phospho-lipids *in vitro*,
Fig. 8. Impact of flagella type on actin polymerization kinetics. (a) Effects of H6, H7, P1 and P2 flagella on αβ-actin polymerization. Pyrene-conjugated globular (g)-actin polymerization was triggered by the addition of 5 mM KCl, 0.2 mM MgCl₂ and 0.1 mM ATP in the presence of different concentrations of flagella, and arbitrary fluorescence (AFU) was measured every 30 s for 1 h at RT. 500 nM cofilin-1 (CFL) was used as a positive control for increased actin polymerization rates in sub-optimal conditions [42]. Data shown are a representative experiment. (b) Maximum velocity ($V_{max}$) of actin polymerization in the presence of titrating concentrations (125 nM–2 µm) of H6 (light blue), H7 (dark blue), P1 (yellow) and P2 (red) flagella. Data were analysed from five independent experiments and post-hoc Tukey pairwise comparisons of differences between specific flagella concentrations and actin alone are presented; $P < 0.01$ (**). (c) $V_{max}$ of actin polymerization in the presence of 250 nM H6, H7, P1 and P2 flagella, and sub-optimal (250 nM) and optimal (500 nM) concentrations of cofilin-1 (CFL). Data represent the results from three independent experiments and post-hoc Tukey pairwise comparisons of the difference between CFL with and without different types of flagella are presented; $P < 0.05$ (*), $P < 0.01$ (**), $P < 0.001$ (***)
membrane-labelled samples. Such overlaps could be routinely detected by confocal microscopy of actin- and adherence may go beyond surface adhesion. This was this process could be a consequence of pathogen-induced membranes, is not known. Alternatively, on epithelial cells proximities for favourable charged flagella colliding with high-resolution of moving samples, labelled in a microscopy approaches may provide more definitive answers about the position of flagella in relation to host membranes. However, at the time of writing, the technology to obtain 3D sub-micrometre resolution of moving samples, labelled in a way that will not interfere with interacting surfaces, is not readily available. Nevertheless, the observations are supported by the additional phenotypic data that E. coli O157:H7 and S. Typhimurium cause 10 and 50%, respectively, less disruption of biological membranes without flagella rotation, compared to wild-type. It is appreciated that these RBC lysis experiments are under conditions where there is high ratio of flagellated bacteria to RBCs and such ‘overload’ activity is unlikely to be reproduced in vivo.

Where potential disruption of host cell membranes in proximity to bacterial flagella was observed by electron tomography, the mechanism by which they do this remains speculative. The haemolysis data support a role for flagella in membrane disruption in vivo. There are under conditions where there is high ratio of flagellated bacteria to RBCs and such ‘overload’ activity is unlikely to be reproduced in vivo.

large-scale membrane rearrangements seen during S. Typhimurium and E. coli O157:H7 colonization, trapping flagella at random, or directed by receptor-mediated endocytosis, triggered by flagella binding to relevant surface molecules. Pseudomonas aeruginosa flagella bind to MUC1 and heparin-sulfated proteoglycans and E. coli flagella bind to ganglioside receptor GM1 and charged glycerolipids, resulting in immuno-modulation and invasion [28, 52–54]. It is also possible that membrane penetration could be caused by a combination of these factors.

Interaction with actin cytoskeletal proteins was not restricted to flagella from pathogenic bacteria. As established, the predicted surface epitopes of the different polymerized flagellins studied are markedly distinct (Fig. S2), complicating analyses of key surface regions required for CFL1, galectin-4 and β-actin binding. Friedlander et al. [51] proposed that flagella binding to artificial surfaces involves low-affinity but high-avidity co-operative binding to surface substrates. Multiple low-affinity interactions of the different polymeric flagellins may also account for the direct binding to cytoskeletal proteins observed in the present study. This supports the concept that this co-operative binding of flagellin subunits within a filament is a generic property of flagella. In a pathogenic context this could lend itself to binding repetitive substrates such as actin filaments. Flagella in a non-pathogenic context may be capable of binding to actin and actin-binding proteins, but for this to be relevant, the flagella would have to gain access through the mucus layer to the epithelial surface. Conversely, motility- and flagella-deficient strains of E. coli and S. Typhimurium can still be pathogenic [55, 56]. However, where expressed, the importance of flagella in the colonization of enteropathogenic and enterohaemorrhagic E. coli, and S. Typhimurium has been established [17, 24, 30, 57], and this study highlights another potential reason for this.

Furthermore, a recent paper has identified a proteolytic site in the hypervariable region of flagella, present in at least 74 bacterial species, including several species of Clostridium. This metalloprotease domain, named flagellinolysin, was not found in either E. coli or S. enterica [58]. However, this raises the possibility that these flagella are capable of other enzymatic activities against actin or membrane lipids. Additionally, by using flagellinolysin-negative species, this study could serve as a baseline in assessing the relative contribution of flagellinolysin to flagella-dependent colonization processes in the future.

Flagella interactions with actin were undertaken with isolated components in physiological conditions, confirming direct binding. Many pathogens reorganize host actin architecture during host colonization. The first bacteria described to use actin rearrangement were the intracellular bacteria Listeria monocytogenes and Shigella flexneri [59, 60]. They both generate filamentous actin ‘comet tails’ that propel the bacteria through the cytosol, allowing colonization of neighbouring cells. Modulation of actin and actin binding proteins, such as Arp2/3 and CFL1, has also been described for certain pathogenic E. coli and Salmonella spp. secreted effector proteins.
However, the findings in this current study provide the first evidence of potential actin manipulation by bacterial flagella.

That flagella may be able to plug into this complex system by themselves in standard conditions is intriguing, given the lifestyles of their associated pathogens. For *E. coli* O157:H7, it has been recently shown that the T3SS can be activated by physical forces acting on the bacterium through uncoupling of GrlA/R binding [63] and it is interesting to speculate that flagella surface interactions with membranes could lead to such activation. Cofilin binding by flagella could stabilize F-actin at the site of colonization, setting up the process of intimate attachment that requires T3SS translocation of specific effector proteins [64]. We note that there was no evidence in this study that the *in vitro* binding interactions of H7 flagella with CFL1 affected actin polymerization synergistically. It is possible that additional factors are required for this; if flagella can gain access to the host cytosol directly, the presence of other actin regulators and different micro-environments may inhibit or enhance these interactions to affect host cell binding. For *S. Typhimurium*, recent work has shown how methylation of flagella enhances membrane interactions that promote invasion [65] and subsequent interactions of flagella with actin could contribute to the dynamic regulation of actin by T3SS effectors that enable bacterial invasion [22]. Counter to these pro-infection concepts, intracellular flagella are activators of innate responses via the NLRC4 inflammasome [11, 12] and *S. Typhimurium* flagella activation of NLRC4 is known to induce actin-dependent host cell surface stiffening to prevent further bacterial invasion in macrophages [66].

In summary, the results of this study demonstrate that *S. Typhimurium* and *E. coli* O157:H7 flagella become intimately associated with host cell membranes during initial adherence. This association either results in or is a consequence of host membrane deformation that may cause disruption, with any disruption likely to be a consequence of flagella rotation. *S. Typhimurium* and *E. coli* O157:H7 flagella can bind to actin and actin-binding proteins. These interactions can influence cytoskeletal dynamics *in vitro* and this may be relevant during bacterial colonization. An understanding of how connected these two phenomena are will require further work to elucidate the molecular mechanisms involved.

References


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