The Bacterial Cytoskeleton Modulates Motility, Type 3 Secretion, and Colonization in Salmonella

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Abstract

Although there have been great advances in our understanding of the bacterial cytoskeleton, major gaps remain in our knowledge of its importance to virulence. In this study we have explored the contribution of the bacterial cytoskeleton to the ability of Salmonella to express and assemble virulence factors and cause disease. The bacterial actin-like protein MreB polymerises into helical filaments and interacts with other cytoskeletal elements including MreC to control cell-shape. As mreB appears to be an essential gene, we have constructed a viable ΔmreC deletion mutant in Salmonella. Using a broad range of independent biochemical, fluorescence and phenotypic screens we provide evidence that the Salmonella pathogenicity island-1 type three secretion system (SPI-1-T3SS) and flagella systems are down-regulated in the absence of MreC. In contrast the SPI-2 T3SS appears to remain functional. The phenotypes have been further validated using a chemical genetic approach to disrupt the functionality of MreB. Although the fitness of ΔmreC is reduced in vivo, we observed that this defect does not completely abrogate the ability of Salmonella to cause disease systemically. By forcing on expression of flagella and SPI-1 T3SS in trans with the master regulators FlhDC and HilA, it is clear that the cytoskeleton is dispensable for the assembly of these structures but essential for their expression. As two-component systems are involved in sensing and adapting to environmental and cell surface signals, we have constructed and screened a panel of such mutants and identified the sensor kinase RcsC as a key phenotypic regulator in ΔmreC. Further genetic analysis revealed the importance of the Rcs two-component system in modulating the expression of these virulence factors. Collectively, these results suggest that expression of virulence genes might be directly coordinated with cytoskeletal integrity, and this regulation is mediated by the two-component system sensor kinase RcsC.

Introduction

Salmonellae remain major global pathogens causing a broad spectrum of disease ranging from gastroenteritis to typhoid fever [1,2]. The emergence of multidrug resistant salmonellae is complicating the management of disease [3,4]. Hence, there is an urgent need to identify novel bacterial targets for the development of new antimicrobial agents or vaccines to combat infection.

The view that bacteria do not possess a cytoskeleton has radically changed in recent years with the discovery of intracellular filamentous protein assemblies with cell-shape defining function [5]. Although there is little primary sequence identity between eukaryotic cytoskeletal proteins and those in prokaryotes, proteins with actin- and tubulin-like structural motifs have been identified in bacteria. Bacterial cytokinesis is dependent on FtsZ which contains a structural fold mirroring tubulin. FtsZ displays similar dynamic properties to tubulin and is able to polymerise unidirectionally in a GTP-dependent manner to produce polymeric filaments [6,7]. Polymers of FtsZ are able to assemble into a transient helical structure and subsequently form a ring-like structure around the circumference of the mid-cell [8]. This Z-ring is required for recruiting proteins for the assembly of the cell division complex [8]. The intermediate filament-like protein crescentin determines the vibroid shape of Caulobacter crescentus cells [9].

The bacterial proteins MreB, Mbl, and ParM display the structural and dynamic properties of eukaryotic actin [10]. Amongst these proteins, MreB is the most homologous to actin in terms of primary sequence, structure, and size [11,12]. The most conserved region of this actin-superfamily is the ATPase domain. MreB can polymerise into helical filamentous structures important for cell morphology. Live cell microscopy in Bacillus subtilis revealed that MreB forms large cables which follow a helical path close to the cytoplasmic membrane [5]. An equivalent MreB protein has been found in Escherichia coli. When MreB is depleted, rod-shaped B. subtilis and E. coli cells become spherical [5,13–15]. In C. crescentus MreB has been implicated to play a role in the...
control of cell polarity [16]. In rod-shaped bacteria the MreB polymeric structures control the localisation of cell wall growth by providing a scaffold for enzymes involved in cell wall assembly [17].

The MreB operon in E. coli and B. subtilis encodes for a number of additional genes, which do not possess any similarity to actin [18]. These include the cellular membrane proteins MreC and MreD, which also have a helical disposition. MreC forms a dimer and interestingly in C. crescentus MreC is localised in spirals in the periplasm [19]. Recent studies by Rothfield and colleagues provide convincing evidence to suggest that in E. coli MreB, MreC and MreD form helical structures independently of each other [20]. Using affinity purification and bacterial two hybrid assays, MreC and MreD appear to interact together [13]. In E. coli there is evidence to suggest that MreB interacts with MreC, but this may not be the case in Rhodobacter sphaeroides or C. crescentus [21]. As well as playing a key role in cell morphogenesis, MreB also has a pivotal function in chromosome segregation [22–24]. Adding the MreB inhibitor A22 [S-(3,4-Dichlorobenzyl) isothiourea] to synchronised cultures of C. crescentus inhibited segregation of GFP-tagged chromosomal origins [22]. However MreB may not function in chromosome segregation in Bacteria [15]. Recently another helically distributed cytoplasmic membrane protein which interacts with MreB named RodZ has been identified [25–27]. Cellular components including the RNA degradosome and lipopolysaccharide have also been identified to be localised in helical structures within the cell [28,29].

In spite of these major advances in our understanding of the structure and organization of the bacterial cytoskeleton, there are major gaps in our knowledge of its role in bacterial pathogenicity. In this study we wished to gain insights into understanding the importance of the integrity of the bacterial cytoskeleton in the ability of Salmonella to cause disease, and thus may provide a novel target for antimicrobial drugs or vaccines.

Author Summary

Salmonella are major global pathogens responsible for causing food-borne disease. In recent years the existence of a cytoskeleton in prokaryotes has received much attention. In this study the Salmonella cytoskeleton has been genetically disrupted, causing changes in morphology, motility and expression of key virulence factors. We provide evidence that the sensory protein RcsC detects changes at the cell surface caused by the disintegration of the bacterial cytoskeleton and modulates expression of key virulence factors. This study provides insights into the ability of Salmonella to cause disease, and thus may provide a novel target for antimicrobial drugs or vaccines.
Table 1. Strains and plasmids.

| Strain | Genotype | Reference |
|--------|----------|-----------|
| SL1344 | Parent Strain | [69] |
| ΔmreC1 | SL1344 mreC::kan | This work |
| ΔmreD1 | SL1344 mreD::kan | This work |
| ΔmreC | SL1344 mreC::kan pTK521 | This work |
| ΔmreD | SL1344 mreD::kan pTK521 | This work |
| ΔSpi-1 | RM69 SPI-1::kan | [70] |
| ΔSpi-2 | 12023 ssaV::kan | [71] |
| ΔflhDC | LT2 flhDC::kan | [72] |
| ΔrcsA | SL1344 rcsA::kan | This work |
| ΔrcsB | SL1344 rcsB::kan | This work |
| ΔrcsC | SL1344 rcsC::cat | This work |
| ΔrcsD | SL1344 rcsD::kan | This work |
| ΔrcsDB | SL1344 rcsDB::kan | This work |
| ΔrcsCBD | SL1344 rcsCBD::kan | This work |
| ΔmreC ΔrcsA | SL1344 mreC::cat rcsA::kan | This work |
| ΔmreC ΔrcsB | SL1344 mreC::cat rcsB::kan | This work |
| ΔmreC ΔrcsC | SL1344 mreC::kan rcsC::cat | This work |
| ΔmreC ΔrcsD | SL1344 mreC::cat rcsD::kan | This work |
| ΔmreC ΔrcsDB | SL1344 mreC::kan rcsDB::cat | This work |
| ΔmreC ΔrcsCBD | SL1344 mreC::cat rcsCBD::kan | This work |
| ΔmreC ΔrcsF | SL1344 mreC::cat rcsF::kan | This work |
| ΔmreC ΔqseF | SL1344 mreC::kan qseF::cat | This work |
| ΔmreC ΔphoBR | SL1344 mreC::kan phoBR::cat | This work |
| ΔmreC ΔyjiGH | SL1344 mreC::kan yjiGH::cat | This work |
| ΔmreC ΔbaeSR | SL1344 mreC::kan baeSR::cat | This work |
| ΔmreC ΔbasSR | SL1344 mreC::kan basSR::cat | This work |
| ΔmreC ΔhydH | SL1344 mreC::kan hydH::cat | This work |
| ΔmreC ΔqseBC | SL1344 mreC::kan qseBC::cat | This work |
| ΔmreC ΔtctDE | SL1344 mreC::kan tctDE::cat | This work |
| ΔmreC ΔcpxAR | SL1344 mreC::kan cpxAR::cat | This work |
| YVM004 | SJW1103 gfp-flIG | [32] |
| YVM004 ΔmreC | SJW1103 gfp-flIG mreC::kan | [32] |
| TH3724 | PfhDC::T-POP (DEL-25) fhCS213::MudJ | [33] |

| Plasmid | Description | Reference |
|---------|-------------|-----------|
| pBR322  | Cloning vector | [73,74] |
| pBAD24  | Cloning vector | [75] |
| pZEP08  | GFP+ transcriptional fusion vector | [76] |
| pBR322-gfp | pBR322 with gfp | This work |
| pBR322-mreB-gfp | pBR322gfp with mreB | This work |
| pNDM220 | Low copy cloning vector | [77] |
| pNDM220-mreB-gfp | mreB-gfp subcloned from pBR322-mreB-gfp | This work |
| pKD13   | Lambda Red template | [31] |
| pKD46   | Lambda Red recombinase | [31] |
| pLE7    | gfp-mreB | [36] |
| pTK521  | pNDM220 pA1/O4/O3::mreB::CD | [14] |
| pCS26   | luxCDABE promoter reporter vector | [49] |
| pCS26-hiLA | hiLA | [49] |
| pCS26-hiLC | hiLC | [49] |
| pCS26-hiLD | hiLD | [49] |
Table 1. Cont.

| Plasmid | Description | Reference |
|---------|-------------|-----------|
| pSB401 | luxCDABE promoter reporter vector | [41] |
| pQA09  | sopB promoter reporter | [41] |
| pRG34  | pSB401-flhA | [41] |
| pRG38  | pSB401-flhD | [41] |
| pRG46  | pSB401-flcC | [41] |
| pRG51  | pSB401-flgA | [41] |
| pMKI-lux | luxCDABE promoter reporter vector | [52] |
| pMKI-lux-ssaG | pMKI-lux-ssaG | This work |
| pBAD24-hilA | hilA inducible expression plasmid | This work |
| pBAD24-rscC | rscC complementation plasmid | This work |

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Construction of Complementation Plasmids

The hilC and rscC open reading frames were amplified from SL1344 genomic DNA and cloned into the EcoRI and XbaI or the EcoRI and HindIII sites of pBAD24 to create pBADhlcA and pBADrsc respectively.

Protein Manipulation

Whole cell total protein samples were obtained by pelleting an appropriate volume of bacterial culture, followed by resuspension in SDS-loading buffer and boiling for 10 mins. Culture supernatants were filter sterilized (0.22 μm) and proteins were ammonium sulphate precipitated (4 g 10 ml-1 supernatant) overnight at 4°C. Precipitated secreted proteins were resuspended in H2O and then combined with an equal volume of sample buffer (Biorad). Western blot analysis was performed using Protran nitrocellulose transfer membranes (Schleicher & Schuell) using a wet transfer apparatus (Biorad). Detection was carried out using 4-chloro-1-naphthol (Sigma) with the peroxidase-labelled secondary antibody (Dako Cytomation).

In vivo Inoculation and Growth Curves

Female C57BL/6 mice were purchased from Harlan Olac Ltd., Blackthorn, Bicester, UK). Mice were used when over eight weeks of age. Bacterial suspensions for injection were grown for 16 h as a stationary culture at 37oC in LB broth. Bacteria were diluted in fresh PBS before mounting onto agarose beds.

Construction of Flagella Live Cell Imaging Strains

Wild type Salmonella SJW1103 cells with chromosomal N-terminal GFP fusion to fliG (YVM004) [32] were P22 transduced with the mrvC::kan mutation to create YVM004 ΔmrvC. This strain, along with the WT control, was subsequently transduced with a chromosomally-based inducible fliDC locus derived from TH2919 [33].

Visualisation of Type 3 Secretion Systems and Flagella

Cells were grown to the appropriate growth phase (mid-log for SPI-1 and flagella, or stationary phase for SPI-2) in relevant media (LB or SPI-2 inducing media). Flagella visualisation strains (fliG-gfp), were mounted on 1% agarose beds for imaging. Samples for visualising the type 3 secretion apparatus were fixed in 4% paraformaldehyde diluted in PBS for 1 h before washing for 15 minutes in three changes of PBS. Samples were incubated with either α-SipA, α-SipB, α-SipC, α-SipD (SPI-1) or α-SaeB (SPI-2) antibodies diluted 1:1000 in PBS for 3 h with gentle agitation. Samples were subsequently washed in PBS before incubating in 1:100 Alexa Fluor 488 conjugated goat anti-rabbit antibody (Invitrogen-Molecular Probes, Paisley, U.K.), washed for 30 mins in fresh PBS before mounting onto agarose beds.

Tissue Immunostaining for Fluorescence Microscopy

Half of each organ was fixed overnight in 4% paraformaldehyde diluted in PBS, washed for 90 min in three changes of PBS and then immersed in 20% sucrose (in PBS) for 16 h at 4°C before being embedded in Optimal Cutting Temperature (OCT) (Raymond A Lamb Ltd, Eastbourne, U.K.) in cryomoulds (Park Scientific, Northampton, U.K.). Samples were frozen and stored at -80°C. 30 μm sections were cut, blocked and permeabilised for 10 min in a permeablising solution containing 10% normal goat serum and 0.02% Saponin in PBS (Sigma, Poole, UK). All sections were stained with 1:1000 dilution of rabbit anti-mouse CD18 monoclonal antibody (clone M18/2, BD Pharmingen), together with a 1:500 dilution of rabbit anti-lps CD18 monochonal antibody (clone M18/2, BD Pharmingen), together with a 1:500 dilution of rabbit anti-LPS O4 agglutinating serum (Remel Europe Ltd), for 16 h at 4°C. Subsequently, sections were washed in PBS then incubated with 1:200 Alexa Fluor 488 conjugated goat anti-rabbit antibody (Invitrogen-Molecular Probes, Paisley, U.K.) and a 1:1000 dilution of Alexa Fluor 488 conjugated goat anti-rabbit antibody (Invitrogen-Molecular Probes, Paisley, U.K.). All sections were mounted onto VECTA-BOND-treated glass slides (Vector Laboratories Ltd.) using Vectashield containing DAPI (Vector Laboratories Ltd.).

Microscopy

All phase contrast and fluorescence images were captured using an Andor iXon885 EMCCD camera coupled to a Nikon Ti-E microscope using a 100x/NA 1.4 oil immersion objective. Images were acquired with NIS-ELEMENTS software (Nikon) and processed using ImageJ. Fluorescence images were decon-
| Primer    | Sequence (5’-3’)                                                                 |
|----------|--------------------------------------------------------------------------------|
| mreC-P1  | GGATTGTCTCCTCCTCGAGCGAGAAATACGCATAGCCTGTGATGCTGAGCTGCTTC                      |
| mreC-P4  | CATCAGGCGGCTACATGGCCGAGCGGATGACGGTCTGAGCTGCTTC                              |
| mreC+5   | ATACGGGCAGGATTATCCCT                                                        |
| mreC+3   | GCGCAATAAGAAACGAGAC                                                       |
| mreD-P1  | GGCGCGACCACGGCGGCTCGGCGGAGGGTACGCGTGGACTGCTGCTTC                           |
| mreD-P4  | GGGGAACCGGAAAGGATAGATAGTAGGCTGAGCTGCTTC                                    |
| mreD+5   | ATCAACCGCAAACCTGCTC                                                        |
| mreD+3   | TCAATATCTCGGCGGCA                                                          |
| qseBC-P1 | GTTAACTGAGGCGAGCTCGGAGCTGCTTC                                             |
| qseBC-P4 | AAATGTGCAAGCTCTTTGCTGATAGGCGGATCCGATCCGAGCTGCTTC                           |
| qseBC+5  | ACATCGCCTGCGGAGGACT                                                         |
| qseBC+3  | GCGGTGCGGTGAAATTAGCA                                                       |
| rcsA-P1  | GTAAGGGGAATATTATGCTGAGGCTGAGCTGCTTC                                       |
| rcsA-P4  | AATTGAGCCGGACTGGAGGTAGATGTCGAGCTGCTTC                                     |
| rcsA+5   | GATTATGGTAGGTTACG                                                          |
| rcsA+3   | CGAGAAGCGGAGGAC                                                        |
| rcsB-P1  | GCCTACGTCAAAAGCTTGCTGAGCTGAGCTGCTTC                                       |
| rcsB-P4  | ATAAGGGGAATATTATGCTGAGGCTGAGCTGCTTC                                       |
| rcsB+5   | CGTGAAGAAAGTGTCCAGG                                                      |
| rcsB+3   | TGGTAGCCTGTTAGGCTG                                                           |
| rcsC-P1  | GTCACTACTATTTTACTATTCGAGGCGGAGCTGCTTC                                     |
| rcsC-P4  | TTTTAGGCGAACGGAGGCAGGGGATCCGAGCTGCTTC                                     |
| rcsC+5   | CGTCAATTACCGCTACTTA                                                        |
| rcsC+3   | GGCTACAGGAGCATGCTAC                                                        |
| rcsD-P1  | CTTTCAAGCCCGGCTAGATGCTGAGGCTGAGCTGCTTC                                   |
| rcsD-P4  | ACCGTGCTGCAAGGCTTTTCTAGCGAGGGGATCCGAGCTGCTTC                             |
| rcsD+5   | TTCATTACCCCTTTATGCT                                                        |
| rcsD+3   | CATATTGCTCATGATTGGG                                                        |
| rcsF-P1  | TCTAACATCTACGCTGAGGAAATATTATGCTGAGGCTGCTTC                               |
| rcsF-P4  | GGGGAAGGAAATACCCGCGGATTAGCTAAACATGCTGAGGCTGCTTC                           |
| rcsF+5   | TCTATTAGCGCTCTGCA                                                        |
| rcsF+3   | CGGCGAATTTTTTTTAG                                                          |
| rcsBD-P1 | GTCACTACTTTTACATTCGAGGCGGAGCTGCTTC                                       |
| rcsBD-P4 | CTTTCAAGCCCGGCTAGATGCTGAGGCTGCTTC                                       |
| rcsBD+5  | CGTCAATTACCGCTACTTA                                                        |
| rcsBD+3  | TTCATTACCCCTTTATGCT                                                        |
| rcsDB-P1 | CTTTCAAGCCCGGCTAGATGCTGAGGCTGCTTC                                       |
| rcsDB-P4 | ATAAGGGGAATATTATGCTGAGGCTGAGCTGCTTC                                       |
| rcsDB+5  | TTCATTACCCCTTTATGCT                                                        |
| rcsDB+3  | TGAGTGCTAGTCTAGGGGCTG                                                      |
| phoBR-P1 | ATGGCGCGCCATGATCCCTCAATCAAGCTGAGAACAAATATTATGCTAGGCGTGGAGCTGCTTC           |
| phoBR-P4 | CATCCGCTGCTGGATTAGGAAATATTATGCTGAGGCTGCTTC                                |
| phoBR+5  | TGTACATAATTGACGCA                                                          |
| phoBR+3  | CTGCAAAGAATAAGGCCA                                                        |
| qseF-P1  | GGGCGCGGCGGTCACACAAGTAGGAGGTAACAGGCCATGATAAGAGTGTAGGCTGAGGCTGCTTC          |
| qseF-P4  | TAAACGAATACATTTTCCGCGTACTCTTACCCGAGCATGAATAACATCCCGGGGATCCGCTGACC         |
| qseF+5   | CAAACCGGGAGCTGCTGAG                                                      |
| qseF+3   | GTTCGCTGTTTGTATGCG                                                       |
| cpxAR-P1 | CGCGTCTGAGCTATAATTTCTGCTGAGGAGCTGCTTC                                   |
| cpxAR-P4 | CGAGATAAAATCCGCGCTCCTGATCAGGGCGATGTTTATTCGCGGAGATCCGCTGACC               |
Table 2. Cont.

| Primer | Sequence (5’-3’) |
|--------|-----------------|
| cpxAR 5’ | GAAATTGATGGTACG |
| cpxAR 3’ | CTCCGGTAAATTCGACG |
| tctDE P1 | AATTCCCTTTAATGCGGAGAAACCTTACACAGATTGATGTGATGGCTTGGAGGTGCTTTC |
| tctDE P4 | TTTTGTAACGGTGCTTATTCCGGCTGACAATTTGTCGCAAATTCCGGGGAATCCGGTACG |
| tctDE 5’ | TTTTTGTAACGGTGCTTATTCCGGCTGACAATTTGTCGCAAATTCCGGGGAATCCGGTACG |
| yjIG P1 | TTCCTGCTCAGCTTCCGGCTGAGGACACCCGTTTCTTGTATGAGGCTGAGGCTTTC |
| yjIG P4 | TAAACTCCGCCGGGAGTATCCAGCATGATAACCTCTTAAATTCCGGGGAATCCGGTACG |
| yjIG 5’ | TCTACTTATTCTTCTT |
| yjIG 3’ | GTTCGCCACTGGTAAATAGG |
| HydH P1 | TCTGGTGTCAGTTAGCGAGACAAAACAGGTTGTAGGCTGAGGCTTTC |
| HydH P4 | GTACAGAACATTGCGCTGGGCGCATTGAGCGTGAGAAGAAAATTCCGGGGAATCCGGTACG |
| HydH 5’ | TAAAGGGCTGGGTCTTACT |
| HydH 3’ | GTGGGACGGCTGGTACG |
| BasS P1 | CTCATGGTGTTGGCTGAGGAGAAGCACTGAGGAGGAAATCAGGTGAGGCTGAGGCTTTC |
| BasS P4 | AGTTTTATCTATGTGTGGGTCACGACGTATTAAACGCCTGATTCCGGGGATCCGTCGACC |
| BasS 5’ | CTCGAGGGTGCGCATGAGG |
| BasS 3’ | GTATTGTGGTACGAGG |
| BasSR P1 | TGGTATTACGGCTGAAATGAGGAGGAGGAGGAAATCAGGTGAGGCTGAGGCTTTC |
| BasSR P4 | ATATCGTCTTACGACCTTGTATTTGTTATGCACAAATAATCATTCCGGGGAATCCGGTACG |
| BasSR 5’ | CTCGAGGGTGCGCATGAGG |
| BasSR 3’ | CTCGAGGGTGCGCATGAGG |
| pEGFP5 | GGCGAATTCCTAGTACCCGGCCATGTGCTAGAAGTGAGGGGAGG |
| pEGFP3 | GACAGCTTTACCTTTCAGTACGAGG |
| mreB5 Eco | GGCGAATTCTGAGATGGTTTCAACATAC |
| mreB3 Xba | GCGCTCTAGACCTTTCCGAGAAGGTCGCG |
| ssaG5 Eco | GCGAATTCGCCAGATGAGGAGGAGGAGG |
| ssaG3 Bam | GGCGGATCCACTAATATTGTCGAAATAC |
| BADhilA5 | GGCGAATTCGCCAGATGAGGAGGAGGAGG |
| BADhilA3 | GCGCTCTAGAATTCCATTTAATAC |
| BADresC5 | GGCGAATTCTGGAATACCTTTCGCTT |
| BADresC3 | GGCGAAGCTTTATGCGGCTGTTTACGAG |

Bold indicates restriction enzyme recognition sites.

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Transepithelial Resistance and Bacterial Effector Translocation Assays

The effect of *Salmonella* infection on transepithelial resistance (TER) was determined for differentiated Caco-2 cells as previously described [34]. Briefly, the Caco-2 cells were grown on transwell inserts (Corning, UK) until differentiated (12–14 days), before the transepithelial resistance was measured for each well. *Salmonella* strains were then added to the cells at a multiplicity of infection (MOI) of 20, and the cells incubated for 4 h. TER measurements were taken every hour and the results given as a ratio of TER (t)/

(t0) to show the percentage change in TER over the course of the experiment. Data were collated and analysed for statistical differences (Student’s t-test) in Minitab.

Samples for the assay of translocated effector proteins were isolated from differentiated Caco-2 cells grown in 6 well plates after infection with an MOI of 20 for 4 h. Excess bacteria were washed off before the cells were solubilised in 0.01% Triton X-100 and centrifuged to remove bacteria and host cell membranes. The host cell cytoplasmic fractions were analysed by western blotting with αSipB antibody.

Results

*In silico* Identification of the *Salmonella* Actin Homologue *mreB*

We wished to identify and characterise putative *Salmonella* cytoskeletal gene homologues. A BLAST search of the *S.*
Typhimurium genome sequence database (www.ncbi.nlm.nih.gov) [35] for the known E. coli actin-homologue MreB identified a putative mre operon of high sequence identity. Comparison of the Salmonella genes to those of E. coli showed 100% (mreB), 90% (mreC) and 94% (mreD) homology at amino acid level, comparisons of these same genes to those in B. subtilis revealed sequence homologies of 57%, 24% and 27% respectively.

MreB Proteins Are Helically Localised

In order to determine the localisation of MreB in Salmonella, vectors expressing N and C terminal fusions of MreB to GFP were used. The N-terminal fusion plasmid has already been described [36], and we constructed a C-terminal fusion vector. Both constructs revealed a helical distribution of MreB along the long axis of the cell. The helices were discerned by assembling a series of z-stack images taken in successive planes by using Metamorph imaging and Huygens deconvolution software (Figure 1A).

Construction of mre Mutants

The mreB gene appears to be essential in bacteria including Salmonella (data not shown), and ΔmreB viable cells often contain compensatory mutations [37]. Each of the components of the cytoskeletal complex, for example MreB, MreC, or MreD, are essential for its function. As an alternative strategy to study the function of the cytoskeleton we therefore generated a mreC deletion strain under conditions designed to minimise selective pressures for undefined secondary compensatory mutations [37]. Using the lambda Red one-step gene disruption method, we successfully constructed a mreCΔkan mutant in the S. Typhimurium wild-type strain SL1344 [31]. This mutation leaves intact the first gene in the operon mreB. Using bacteriophage F22int the mreCΔkan mutation was then transduced into a genetically ‘clean’ SL1344 strain harbouring plc-mre operon (pTK521) [14] and the resulting strain designated ΔmreC. The plc-mre operon is a low copy number plasmid expressing the mre operon from the IPTG-inducible lac promoter. The identity of the mutation was confirmed by PCR and DNA sequencing. Expression of MreC was assessed by western blotting in the mutant strains, revealing no detectable levels MreC unless complementation was induced (Figure S1). In addition to the ΔmreC mutant, the lambda Red method was used to generate ΔmreD.

Morphology and Growth Rates

When the morphology of the ΔmreC mutant was examined microscopically, the cells were no longer rod-shaped but spherical (Figure 1B). Upon the addition of IPTG the morphology of the ΔmreC strain was restored to the wild-type rod shape. Under microscopic examination the ΔmreD mutant displays a similar morphological phenotype to the ΔmreC strain. WT cells were measured to be on 1.61(+/−0.49) μm in length and 0.73(+/−0.17) μm in width, whereas the ΔmreC cells were 2.03(+/−0.60) μm in length and 1.21(+/−0.41) μm in width. Complementation of the ΔmreC mutant with 100 μM IPTG resulted in wild type shaped cells 1.82(+/−0.44) μm in length and 0.78(+/−0.24) μm in width. Measurements were taken from a minimum of 350 cells per strain. Growth rates of the strains were determined in LB media at 37°C revealing a ∼50% increase in the lag phase of the ΔmreC mutants (Figure S2), which subsequently grow at a comparable rate to that of the wild type or complemented mutant strains during log phase.

Motility and Expression of Flagellin Subunits

The motility phenotype of ΔmreC was examined on semi-solid agar plates. In contrast to the isogenic parent, the ΔmreC cells were no longer motile. Surprisingly, this motility defect has not been reported in either E. coli or S. subtilis. Cellular and secreted proteins of the parent SL1344 and ΔmreC were examined by SDS-PAGE and western blotting using antibodies directed against the phase-1 and phase-2 flagellin subunits FlIC and FlIIB. Neither of these subunits were present in either the secreted or cellular proteins, explaining the inability of the cells to swim (data not shown). The non-motile phenotype was fully complementable in trans upon the addition of IPTG to the mutant strain harbouring pTK521 (Figure S3).

Expression of Flagella Genes

We observed that the Salmonella ΔmreC deletion strain was non-motile and failed to express flagella subunits FlIC or FlIIB. The regulation and assembly of flagella in Salmonella is complex. Flagella genes are arranged into 14 operons and their transcription is organised into a regulatory hierarchy of early (class I), middle (class II), and late genes (class III) [38]. The class I flhDC operon is the master regulator, with FlhD and FlhC forming a heterotetramer that is required for transcriptional activation of the class II genes, which encode the hook-basal body complexes and the alternative sigma factor FliA (sigma28). FliA alone or with FlhDC, activates expression of the class III operon genes, which encode the filament protein, hook-associated proteins, motor proteins, and chemotaxis proteins [39,40]. The class III genes are further subdivided into fliC-independent expression class IIIa or class IIIb [41]. In order to systematically investigate the mechanistic basis for the ΔmreC motility phenotype we have taken selected class I, II, and III regulated flagella gene promoter fusions to a luciferase reporter gene, and monitored their expression by luminescence in

Figure 1. Localization and morphological role of the S. Typhimurium Mre proteins. (A) Fluorescence microscopy montage showing z-sections taken of MreB-GFP fusions in WT SL1344 revealing a helical distribution. Slices taken at 0.1 μm intervals on live cells in mid log phase going from left to right followed by maximum intensity projection (boxed). (B) Morphology of WT S. Typhimurium (I) and ΔmreC (II) reveal the mutant has changed from rod to round-shaped, with some heterogeneity in size noted. In all images the bar represents 1 μm.
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wild type and ΔmreC strains. Constructs with flhD (class I), fliA, flgA, (class II), and fliC (class III) promoters fused to the luciferase reporter gene were used. The reporter plasmid pSB401 has a promoterless luxCDABE operon and was used as a control.

The class I flhD promoter displayed a reduction in the level of expression in ΔmreC compared to the wild-type strain suggesting the class I promoter has reduced activity. Notably greater changes in the expression profiles occur in other class II and class III genes. The class II promoters for the operons encoding the transcriptional regulators fliA\_Y and flgAM display significant reductions in expression levels in ΔmreC (Figure 2). As predicted from the western blotting data expression of the flhD class III promoter was significantly reduced. Collectively, the promoter-reporter activity data can account for the motility defect.

**Expression of SPI-1 and SPI-2 Type 3 Secretion System Proteins**

Type 3 secretion systems are essential for the virulence of a range of pathogens including *Salmonella* [42,43]. The secretion apparatus assembles into a supramolecular needle-complex. Secreted effector proteins in the bacterial cytoplasm traverse through the needle-complex and the bacterial multi-membrane envelope, directly into host cells [44–46]. The apparatus anchors to the cell envelope via a multi-ring base. *Salmonella* possess two T3SS’s encoded by pathogenicity islands (SPI’s). The SPI-1 T3SS is important for invasion of intestinal epithelial cells and the SPI-2 T3SS plays a central role in survival within the hostile environment of a macrophage. The SPI-1 T3SS system translocates virulence effector proteins into the cytosol of epithelial cells resulting in rearrangements of the actin cytoskeleton which enable *Salmonella* to invade [47]. To investigate whether the mreC mutation has an impact on SPI-1 T3S, we used western blotting to determine the presence and functionality of the system using antibodies to an apparatus protein PrgH as well as the effector proteins SipA and SipC, in both SL1344 and ΔmreC. In contrast to the wild-type SL1344, the T3S structural and effector proteins were not expressed in the cellular or secreted fractions from the ΔmreC depletion mutant (Figure 3A). This suggests that SPI-1 T3S in the ΔmreC mutant is not fully functional. The expression and secretion phenotypes were fully complementable in trans upon the addition of IPTG (data not shown).

The functional assembly of SPI-1 T3SS was also confirmed using transepithelial resistance (TER) assays in differentiated Caco-2 cells, showing a reduced ability to disrupt epithelial tight junctions in the ΔmreC mutant compared to the wild type strain (Figure 4).

To further assess the disruption of the functionality of the SPI-1 T3S, a translocation assay was performed in Caco-2 cells infected with the strains. Host cell cytoplasmic proteins were probed for the bacterial effector protein SipB using western blotting (Figure S4). This revealed the inability of the ΔmreC mutants to infect host epithelia and disrupt their tight junctions. In addition, ΔmreC was fully complementable in this assay following IPTG induction.

The SPI-2 T3SS is pivotal for the establishment of the *Salmonella* containing vacuole (SCV) inside macrophages and subsequent survival [43]. We next investigated the effect of the ΔmreC mutation on the functionality of the SPI-2 T3SS. The strains were grown under SPI-2 inducing conditions and the T3S of the translocon protein SseB monitored. SseB together with SseC and SseD function as a translocon for other effector proteins and SseB is normally found associated with the outer surface of *Salmonella*. Thus membrane fractions were purified to monitor expression and T3S by western blotting. This revealed that in contrast to the SPI-2 negative control (ssaV), SseB was secreted and associated with the cell surface in the ΔmreC mutant (Figure 3B). This suggested a defect in the translocation machinery of the SPI-2 T3SS. To further investigate this, a translocation assay was performed using host cell cytoplasmic proteins probed for the bacterial effector protein SipB using western blotting. This revealed the inability of the ΔmreC mutants to infect host epithelia and disrupt their tight junctions. In addition, ΔmreC was fully complementable in this assay following IPTG induction.

**Figure 2. Impact of ΔmreC on the transcription of flagellar genes.** Transcriptional expression profiles of flhD, flgA, fliA and fliC promoter reporters in WT SL1344 (blue diamonds) and ΔmreC (red squares) expressing the *Photorhabdus luminescens* LuxCDABE luciferase. Experiments were repeated at least three times and error bars indicate standard deviation.

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Figure 3. Expression of *Salmonella* SPI-1 and SPI-2 effector proteins in Δ*mreC*.  

(A) Expression of SPI-1 proteins in WT SL1344, Δ*mreC*, ΔSPI-1, and ΔSPI-2 mutants during SPI-1 inducing conditions as revealed by western blotting with polyclonal αSipA and αSipC antibodies. Expression of SPI-2 in WT SL1344, Δ*mreC*, ΔSPI-1, and ΔSPI-2 mutants during SPI-2 inducing conditions as revealed by western blotting of membrane fraction samples with polyclonal αSseB antibody. Samples representing total proteins and secreted proteins are shown. Arrows indicate the respective protein bands.  

(B) Transcriptional expression profiles of *hilA*, *hilC*, *hilD*, *sopB* (SPI-1) and *ssaG* (SPI-2) promoter reporters in WT SL1344 (blue diamonds) and Δ*mreC* (red squares). Experiments were repeated at least three times and error bars indicate standard deviation.

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Expression of SPI-1 and SPI-2 Type 3 Secretion System Regulatory Genes

Several environmental signals and transcriptional factors modulate expression of the SPI-1 T3SS. We wished to understand the mechanistic basis by which expression of the SPI1-T3SS is down-regulated. Within SPI-1 there are key transcriptional activators which regulate expression of SPI-1 genes: HilC, HilD, HilA, and InvF. Both HilC and HilD activate expression of SPI-1 genes by binding upstream of the master regulatory gene hilA to induce its expression[46]. HilA binds and activates promoters of SPI-1 operon genes encoding the type 3 secretory apparatus, several secreted effectors, and the transcriptional regulator InvF. InvF activates expression of effector genes inside SPI1 and also effector genes outside SPI-1 such as sopB and sopE[47].

Expression of selected SPI-1 T3SS genes was monitored using transcriptional promoter reporters in ΔmreC, using constructs harbouring the hilA, hilC, hilD, invF and sopB promoters fused to the promoterless luxCDABE operon that produces light in response to gene expression [49–51]. Each construct was introduced into both wild-type SL1344 and ΔmreC depletion mutant, and the level of expression of the promoters in these strains monitored by luminescence assays. WT SL1344 and ΔmreC cells harbouring pCS26 or pSB401 vectors alone were used as controls, and did not produce any luminescence as expected. The reporter assays revealed that the SPI-1 transcription factor gene promoters for hilA, hilC, hilD, and invF were completely inactive in ΔmreC in contrast to the wild-type strain. However the promoter of sopB located in SPI-5 remained active but its activity was marginally lower than in the wild-type strain (Figure 3B). The regulation of many T3SS genes often require multiple signals for maximal expression and clearly other signals remain in the ΔmreC depletion mutant which drive expression of the SopB in SPI-5.

Expression of SPI-2 T3SS genes were monitored using a transcriptional reporter for the SPI-2 gene ssaG, whose promoter was cloned upstream of the luxCDABE luciferase operon in the plasmid pMK1-lux [52]. The construct was transformed into wild-type SL1344 and ΔmreC, and the luminescence and OD600 measured during growth in SPI-2 inducing conditions (Figure 3B). The ssaG promoter remains active in the ΔmreC mutant although expression appears to be delayed, and is marginally less than in WT. This evidence supports the western blot data with αSseB and suggests that in contrast to the SPI-1 T3SS, the SPI-2 T3SS remains functional in the absence of the cytoskeleton.

Function of the RcsC Two-Component System in Regulation of SPI-1 T3S and Motility in ΔmreC

Two-component regulatory systems are vital in sensing environmental and cell surface signals, enabling bacteria to rapidly adapt to ever changing conditions [53,54]. These signals are detected by histidine protein sensor kinases, which subsequently transfer phosphate groups to an aspartate residue in the response regulator proteins, thus modulating their regulatory activities. The environmental signals are thus transmitted by a phosphorelay system to regulate gene expression.

In order to identify putative regulators of the ΔmreC observed phenotypes, we have constructed knockout mutations in a range of two-component systems. As an initial screen, a panel of nine separate two-component system mutant strains were constructed as double mutants with ΔmreC. One two-component system sensor kinase mutation ΔrscC resulted in recovery of SPI-1 effector expression in the ΔmreC background as judged by western blotting using αSipC sera (Figure 5 panels A and B). Interestingly the amount of SipC protein expressed and secreted from the cell was less than the wild-type suggesting there are additional repressors continuing to operate (Figure 5 panels A and B and Figure S5). Furthermore, disruption of rscC also significantly de-repressed motility (Figure 6 and Figure S6) in a ΔmreC mutant similar to SPI-1 expression, again suggesting there are additional repressors involved. Expression of the RscC protein in trans was able to restore the phenotype of ΔmreC ΔrscC back to the equivalent of a ΔmreC

Salmonella Cytoskeleton and Pathogenicity

Figure 4. Percentage change in transepithelial resistance of differentiated Caco-2 cells after 4hr infection. TER of polarised Caco-2 monolayers exposed to Salmonella strains at an MOI of 20. TER change is expressed as a percentage alteration at 4hr compared to the initial value at time zero. Error bars indicate the standard deviations derived from at least three independent experiments. * Indicates statistical difference from WT (p<0.05).

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strain, with respect to repressing SPI-1 type 3 secretion and motility. These complementation studies provide further evidence supporting the regulatory role of RcsC in the ΔmreC phenotypes (Figure S7).

Rcs is a highly complex multi-component phosphorelay system and was originally identified in regulating genes involved in capsule synthesis in *Escherichia coli* [55,56]. The RcsC sensor kinase phosphorylates RcsD, which subsequently phosphorlylates the DNA binding response regulator RcsB. The unstable RcsA protein and additional auxiliary proteins can also interact and regulate RcsB. The Rcs system is involved in down-regulating the expression of flagella, SPI1-T3S and increasing biofilm formation [57].

We therefore also constructed ΔmreC ΔrcsA, ΔmreC ΔrcsB, ΔmreC ΔrcsC, ΔmreC ΔrcsD, ΔmreC ΔrcsF, ΔmreC ΔrcsDB, and ΔmreC ΔrcsCBD double mutants, which however did not restore either SPI-T3S or motility (Figures 5, 6, and S6). We propose that in the absence of RcsC signalling, phosphorylated levels of RcsB are depleted enabling de-repression of FlhDC and motility. The presence of RcsDB appears essential for restoring motility in the absence of RcsC [55]. The functionality of SPI-1 T3SS in the ΔmreC ΔrcsC and ΔmreC ΔrcsDB mutants were assessed in a TER assay, which revealed partial restoration of tight junction disruption in the ΔmreC ΔrcsC mutant, but not in the ΔmreC ΔrcsDB (Figure S8).

It has been suggested that the outer membrane protein RcsF may perceive some of the environmental signals necessary to activate the Rcs phosphorelay system. To investigate this we constructed a ΔmreC ΔrcsF mutant which failed to restore motility or SPI-1 T3S and appeared phenotypically identical to ΔmreC (Figure 5, S6). This would suggest that RcsF is not involved in the observed ΔmreC phenotypes. Furthermore as the auxiliary protein RcsA can interact and regulate RcsB, we therefore disrupted the rcsA gene in ΔmreC and which also resulted in no impact on the observed phenotypes (Figure 5, S6).

In summary, we propose that RscC is sensing cell surface perturbations [58] in ΔmreC, resulting from a disrupted cytoskeleton, and subsequently down-regulating the expression of SPI-1 T3S and motility. This signalling appears to be independent of both RcsF and RcsA.

**Chemical Genetic Inactivation of the Essential MreB Protein**

A cell permeable compound named A22 [3-(3,4-Dichlorobenzyl)isothiourea] has been demonstrated to perturb MreB function [59]. As an alternative approach to genetically disrupting the essential gene *mreB*, we exposed wild-type *Salmonella* cultures to A22 and observed a morphological change from rod to spherical-shaped cells. In addition we phenotypically screened and tested A22-treated cells for motility and T3S. The A22-treated cells were phenotypically...
ically identical to ΔmreC with respect to cell shape, motility, SPI-1 T3S, and also SPI-2 T3S (data not shown). The effects of A22 were completely reversible following its removal (data not shown). Thus the chemical genetic inactivation of MreB, independently corroborates the phenotypic observations made with ΔmreC.

The Salmonella mre Operon Plays an Important Role in Colonization during in vivo Infection

The ΔmreC defect clearly has an impact on the expression of important virulence determinants of Salmonella in vitro. We therefore wished to investigate the contribution of the bacterial cytoskeleton on the virulence of Salmonella in vivo using the mouse model. We observed that the SPI-1 T3SS in ΔmreC is completely down-regulated, and as this virulence system is important for infection through the oral route of inoculation the strain would be attenuated.

We therefore explored the colonization of ΔmreC using the intravenous route allowing us to examine the impact of the host on the further down-stream stages of infection. Groups of 5 female C57/BL6 mice were inoculated intravenously with circa 10^3 colony forming units of either control SL1344 or ΔmreC. The times taken for clinical symptoms to appear were determined. Viable bacterial numbers in the spleen and liver for SL1344 were determined at days 1 and 4, and ΔmreC at days 1, 4, 7, and 10. The in vivo bacterial net growth curves vividly demonstrate two clear phenotypic effects upon the growth of ΔmreC compared to the wild-type. Firstly, there is a greater initial kill of ΔmreC, and this is secondly followed by a slower net growth rate. However, in spite of the reduced growth rate of ΔmreC, the bacterial numbers steadily increase over 6 days. This eventually causes the onset of clinical symptoms necessitating termination of the experiment at day 10 (Figure 7). During these stages Salmonella infect and multiply within macrophages and the

Figure 6. Motility of Salmonella mutant cells. Representative images showing the motility of SL1344 WT, ΔflhDC ΔmreC1, ΔmreC, ΔmreC plus IPTG, ΔrcsC, ΔmreC ΔrcsC, ΔmreC ΔrcsDB, and SL1344 WT plus A22 cells grown on motility agar at 37°C. White circles highlight the limits of motility on the agar plates.
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SPI-2 T3SS is essential for survival. Thus providing further evidence to support the presence of a functional SPI-2 T3SS in \( \text{DmreC} \). Collectively, these observations imply the \( \text{mreC} \) defect reduces the virulence of the strain, but does not completely abrogate its ability to multiply and cause disease systemically \( \text{in vivo} \).

**Morphology \( \text{in vivo} \)**

Strains recovered from \( \text{in vivo} \) passage were tested for changes in morphology, motility and T3S, and were found to be identical to the input strain. Furthermore the \( \text{in vivo} \) morphology of the strain within livers and spleens was determined by immunofluorescence microscopy. Sections of livers and spleens were taken and stained as described in the materials and methods. Figure 8 demonstrates the \( \text{Salmonella DmreC} \) mutant strain retains the round morphology \( \text{in vivo} \) compared to the rod shaped wild-type control. Collectively these data suggests that the mutation has remained stable during the \( \text{in vivo} \) passage for the virulence phenotypes tested.

**Role of the Cytoskeleton in the Assembly, Regulation and Function of SPI-1 T3SS and Flagella**

The regulation and assembly of SPI-1 T3SS and flagella are complex. When the bacterial cytoskeleton is disrupted both the

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**Figure 7. Contribution of \( \Delta \text{mreC} \) to \( \text{in vivo} \) colonization.** In \( \text{vivo} \) growth kinetics of WT SL1344 and \( \Delta \text{mreC} \) in livers and spleens of C57BL/6 mice inoculated intravenously with \( 10^7 \) colony forming units. Viable bacterial counts in the spleen and liver were performed at days 1, 4, 7 and 10, and expressed as mean log\(^6\) viable count +/- standard deviation. 
doi:10.1371/journal.ppat.1002500.g007

**Figure 8. Morphology of \( \Delta \text{mreC} \) in host tissues.** Representative fluorescence micrograph of \( \text{Salmonella} \) SL1344 WT and \( \Delta \text{mreC} \) within a phagocyte in infected livers of C57BL/6 mice at 72 h p.i. CD18+ expressing cells (red), \( \text{Salmonella} \ \Delta \text{mreC} \) (green), nucleic acid is indicated by DAPI (blue). Scale bar, 5 \( \mu \)m. 
doi:10.1371/journal.ppat.1002500.g008
SPI-1 T3SS and flagella expression are down-regulated. A hypothesis is that the integrity of the cytoskeleton is essential for the correct assembly of these complex macromolecular structures and in its absence the SPI-1 and flagella gene expression are down-regulated to conserve resources. Alternatively, in the absence of a functional cytoskeleton the bacterial cell is stressed and shuts down the expression of energetically expensive “non-essential” machinery. To test these ideas we wished to force on the expression of SPI-1 T3S and flagella genes, and examine whether these systems are correctly assembled and functional. We therefore expressed in trans from heterologous inducible promoters either the flagella master regulator FliC or the SPI-1 T3S regulator HilA in a panel of strains including ΔmreC. Strikingly, expression of FliDC restored both the expression and assembly of flagella on the cell surface as determined by fluorescence microscopy (Figure 9A) and motility assays (data not shown) in ΔmreC. Furthermore, expression of HilA in trans up-regulated expression of the SPI-T3SS and its assembly on the cell surface as determined by immunofluorescence microscopy (Figure 9B) western blotting with SipB antibody (Figure S9) or functionally by TER measurements (Figure 4). In contrast to SPI-1 T3SS and flagella, the expression of the SPI-2 T3SS was not turned off in the ΔmreC mutant as shown in (Figure 9C). Interestingly, in WT cells the SPI-1 T3S apparatus and flagella appear to be present in around six to eight copies mainly along the long axis of the cell. In marked contrast the SPI-2 apparatus is typically present in one or two copies located at the poles of the bacterial cell [42], whereas their localisation appears less clear in the ΔmreC mutant, possibly due to perturbations in the cell envelope and the indistinct cell polarity in these cells caused by disruption of the cytoskeleton. The complementation of the functional assembly of SPI-1 T3SS was also confirmed using TER assays, where the levels of decrease in resistance after infection with ΔmreC strain reverted to that of the parent strain upon induction of the transcriptional regulator hilA (Figure 9B and S9), or complementation of the ΔmreC mutation (Figure 4). Taken together the data support the notion that the cytoskeleton is not required for the correct assembly of these virulence factors but essential for their expression.

Discussion

Bacterial cells possess dynamic cytoskeletons composed of diverse classes of self-assembling polymeric proteins. Some of these proteins resemble eukaryotic actin, tubulin, and intermediate filaments both structurally and functionally [5,7,11,12]. The bacterial tubulin FtsZ plays a key role in cell division. Bacterial actins provide vital functions in maintaining cell morphology, segregating DNA, and positioning bacterial organelles. It has recently been demonstrated in Helicobacter pylori, that MreB is essential not for cell shape but for maintenance of the full enzymatic activity of urease, an essential virulence factor [60]. Furthermore the MreB cytoskeleton is also essential for the polar localisation of pili in Pseudomonas aeruginosa [61].

Using a variety of approaches we have demonstrated the importance of the bacterial cytoskeleton in the pathogenicity of Salmonella. MreC and MreD form a complex in the cytoplasmic membrane, which subsequently interacts with MreB. The mreB gene appears to be essential in many organisms including as we discovered in Salmonella. Viable mreB mutants often contain compensatory changes in other genes e.g. ftsZ which compensate for the lethality of the mreB lesion [37]. As an alternative strategy to investigate the function of the bacterial cytoskeleton and avoid these deleterious effects, we carefully constructed deletion mutants of mreC in strains harbouring a single-copy plasmid expressing the MreB operon from the lac promoter. In addition we confirmed the phenotypic effects of the mreC genetic lesion by disrupting the functions of MreB using a chemical genetics approach and inactivating MreB with A22.

Removal of the gratuitous inducer IPTG from the growth medium of the ΔmreC deletion mutant resulted in cells changing from rod to a spherical shaped morphology. Using fluorescence microscopy MreB was observed to be no longer distributed in a helical fashion throughout the cell but rather diffusely throughout the cytoplasm (data not shown). Presumably MreB polymers are no longer able to contact the cytoplasmic membrane via MreD attachment sites resulting in mis-assembly of the entire cytoskeleton. In growing cells, this disruption of the cytoskeleton leads to loss of the rod-shape.

We next examined the motility of ΔmreC deletion strain to assess the functionality of flagella. The strains were non-motile and western blotting revealed absence of the flagellin filament subunit proteins FliC and FliB in both secreted and also cytoplasmic protein fractions, suggesting expression of these alternative subunits had been switched off. Flagella gene expression is complex and involves a regulatory hierarchy of Class I, Class II, and Class III genes [38]. The class I flhDC operon is the master regulator, and FlhDC complex is required for transcriptional activation of the class II genes including the specialized flagellar sigma factor FlsA. FlsA alone or with FlhDC complex, activates expression of the class III operon genes encoding motor proteins, hook-associated proteins, the filament protein, and chemotaxis proteins [39,40]. Expression of the FlhDC complex was reduced but still appeared comparable between the wild-type and the ΔmreC suggesting changes in the promoter activity of flhDC alone are not responsible for the observed phenotype. Class II gene expression was significantly reduced. Expression of the Class III gene flsC was completely down-regulated confirming the western blot observations. Hence these independent observations are in accordance with the ΔmreC motility data. Thus in the absence of the cytoskeleton expression of class II and class III flagella genes appears to be down-regulated.

Expression of the SPI-1 T3S system is essential for invasion of intestinal epithelial cells and the SPI-2 T3SS plays a central role in survival within the hostile environment of a macrophage [43]. Western blotting revealed the SPI-1 T3S structural protein PrgH in ΔmreC was no longer expressed or secreted in the ΔmreC deletion mutant. The phenotype was fully complementable by the addition of IPTG. Several environmental signals and transcriptional factors modulate expression of the SPI-1 and SPI-2 T3SS [43,45,62]. We wished to understand the mechanistic basis by which expression of the SPI-1 T3SS is down-regulated. Within SPI-1 there are key transcriptional activators which regulate expression of SPI-1 genes: HilC, HilD, HilA, and InvF. Using promoter-luciferase transcriptional reporter assays it
was revealed that the SPI-1 transcription factor gene promoters for \( hilA, hilC, hilD \), and \( incF \) were completely inactive in \( \Delta mreC \), in marked contrast to the control wild-type strain. Surprisingly, the promoter of \( sopB \) located outside of SPI-1 in \( \Delta mreC \) remained active but its activity was marginally lower than in the wild-type strain.

The regulation of many T3SS genes often require the input of multiple signals for maximal expression and clearly other signals remain in the \( \Delta mreC \) deletion mutant which drive expression of the SopB in SPI-1. It therefore appears that the SPI-1 T3SS is completely down-regulated in the absence of an cytoskeleton by an unidentified regulatory factor. In contrast, the SPI-2 T3SS remains functional as evidenced by western blotting with SseB antibody and promoter-reporter assays. This is further corroborated with the \( in \_vivo \) evidence that following systemic inoculation, \( \Delta mreC \) is able to survive and multiply within the host. This takes place within the hostile environment of the macrophage where SPI-2 T3S is essential for biogenesis of the \( Salmonella \) containing vacuole and survival [43,63,64].

We wished to gain further insights into the mechanistic basis of the down-regulation of both SPI-1 T3SS and motility in \( \Delta mreC \). Two-component systems play an essential role in sensing and responding to environmental and cell surface signals [54]. To investigate if two-component systems contribute to the regulation of the \( \Delta mreC \) phenotypes, we constructed a panel of separate two-component system mutant strains in an \( \Delta mreC \) background. The double mutants were screened for recovery of motility and expression of the SPI-1 T3SS. A mutation in the \( rcsC \) sensor kinase gene resulted in significant but not complete recovery of both motility and expression of the SPI-1 T3SS.

The \( Rcs \) phosphorelay system regulates a broad range of genes from capsule synthesis in \( E. \ coli \) to increasing biofilm formation [58]. \( RcsC \) also plays an important role in repressing expression of flagella and SPI-1 T3SS in \( Salmonella \) Typhi [57]. The \( RcsC \) sensor kinase normally phosphorylates \( RcsD \), which subsequently phosphorylates the DNA binding response regulator \( RscB \). However, in \( \Delta mreC \) \( \Delta rcsDB \) and \( \Delta mreC \) \( \Delta rcsCBD \) there was no restoration of either motility or expression of the SPI-1 T3SS suggesting that \( RcsC \) signals repression and requires the presence of \( rcsDB \) to mediate this effect. We propose that in \( \Delta mreC \), the sensor kinase \( RcsC \) detects cell surface perturbations and down-regulates expression of flagella and the SPI-1 T3SS apparatus [58]. This signalling is independent of both the outer membrane lipoprotein \( RcsA \) sensor and the auxiliary regulatory protein \( RcsA \).

There are a number of explanations to provide a bacterial rational for this shutdown in expression. In the absence of a functional cytoskeleton the flagella and SPI-1 T3SS are either not being correctly assembled, triggering a feedback loop to repress expression, or alternatively are down-regulated to prevent the cell from wasting valuable resources under these conditions. To test the assembly idea, we forced on the expression of flagella and SPI-1 T3SS genes by expressing the regulators \( flhDC \) or \( hilA \) in trans in \( \Delta mreC \). Using independent methods we observed the correct assembly and function of these macromolecular machines suggesting the cytoskeleton is not essential for functionality. The cytoskeleton could also have a role in sensing cellular stress, as has recently been suggested by Chiu and colleagues [65]. They propose that the integrity of the cytoskeleton may be exploited by the cell to monitor oxidative stress and physiological status. If the cytoskeleton disintegrates in the absence of \( MreC \), this may be sensed by the cell leading to a shut-down of the SPI-1 T3S apparatus and down-regulation of flagella protein expression. We have provided mechanistic insights into the regulation of motility and SPI-1 T3S in \( \Delta mreC \). We have identified the two-component system sensor \( RcsC \) as an important regulator controlling expression of these systems, presumably as a consequence of sensing membrane perturbations brought about by the disruption of the cytoskeleton [58].

With a non-functional SPI-1 T3SS, we would expect the \( \Delta mreC \) would be attenuated in mice when administered by the oral route as it is unable to invade intestinal epithelial cells by the SPI-1 T3SS. We therefore explored the colonization of \( \Delta mreC \) \( \Delta sopB \) using the intravenous route of inoculation [66]. This provides an opportunity to examine the impact of \( RcsC \) on the downstream stages of infection, \( Salmonella \) infect and multiply within macrophages during the systemic stages of infection. Survival within the hostile environment of the macrophage would require a functional SPI-2 T3SS in the \( Salmonella\)-containing vacuole to remodel the host cell environment and survive attack from reactive oxygen free radicals [64,67,68]. By examining the \( in \_vivo \) net bacterial growth curves within livers and spleens two clear phenotypic effects were revealed with \( \Delta mreC \) compared to the wild-type. Greater initial killing of \( \Delta mreC \) is followed by a slower net growth rate with the bacterial numbers steadily increasing over six days. Clinical symptoms begin to appear and by day ten these symptoms necessitate termination of the experiment. The phenotypic data clearly imply the \( \Delta mreC \) defect reduces the colonization of \( Salmonella \), but does not completely abrogate its ability to multiply and cause disease systemically \( in \_vivo \). This would suggest that the second T3S in \( Salmonella \) encoded on SPI-2 remains sufficiently functional to permit growth in the absence of the cytoskeleton.

In the absence of an intact cytoskeleton in \( \Delta mreC \) the expression of the SPI-1 T3SS and flagella are clearly down-regulated. Strikingly however, the SPI-2 T3SS appears to remain functional contributing to the virulence of the \( \Delta mreC \) strain observed \( in \_vivo \). A possible explanation could be that the regulation of the SPI-2 T3SS is co-ordinated independently of the integrity of the cytoskeleton in contrast to flagella and SPI-1 T3SS. Collectively these data highlight the importance of the bacterial cytoskeleton in the ability of \( Salmonella \) to cause disease, and may provide opportunities for the development of new antimicrobials to target the cytoskeleton.

### Supporting Information

**Figure S1 Expression of MreC in complemented \( \Delta mreC \) cells.** Western blot of total protein samples from SL1344 WT, \( \Delta mreC \), \( \Delta mreC \), and \( \Delta mreC \) plus 100 \( \mu \)M IPTG cells using \( \alpha \_MreC \) antibody. MreC is indicated at approximately 38kDa and is distinguishable from background bands. (TIF)

**Figure S2 Growth curve of \( Salmonella \) mutant cells.** Log phase growth of SL1344 WT, \( \Delta flhDC \), \( \Delta mreC \), and \( \Delta mreC \) plus 100 \( \mu \)M IPTG, and A22 treated SL1344 WT cells. Strains were grown in LB media at 37°C. (TIF)

**Figure S3 Motility of \( Salmonella \) \( \Delta mre \) mutant cells.** Motility of SL1344 WT, \( \Delta flhDC \), \( \Delta mreC \), \( \Delta mreC \), \( \Delta mreC \) plus 100 \( \mu \)M IPTG, and A22 treated SL1344 WT shown as a percentage of the wild type. Strains were grown on motility agar at 37°C. Experiments were repeated at least three times and error bars indicate SD. * Indicates statistical difference from WT \( p<0.05 \). (TIF)

**Figure S4 Translocation of SipB SPI-1 effector protein into Caco-2 cells.** Western blot of host cytosol fractions with \( \alpha \_SipB \) antibody following infection of cells with \( Salmonella \).
SL1344 WT, ΔSPI-1, ΔmreC1, ΔmreC (+/− IPTG) mutants. SipB is indicated at approximately 63kDa. (TIF)

**Figure S5** Secretion of SPI-1 effector protein SipC in ΔrcsC mutant cells. Western blot of secreted protein samples from SL1344 WT, ΔmreC, ΔSPI-1, ΔSPI-2, ΔrcsC, and ΔmreC ΔrcsC cells using αSipC antibody. SipC is indicated at approximately 43kDa. (TIF)

**Figure S6** Motility of Salmonella Δrcs mutant cells. Motility of SL1344 WT, ΔmreC, ΔfliDCG, ΔrcsA, ΔrcsB, ΔrcsC, ΔrcsD, ΔrcsE, ΔrcsDB, ΔmreC ΔrcsA, ΔmreC ΔrcsD, ΔmreC ΔrcsE, ΔmreC ΔrcsDB, and ΔmreC ΔrcsCBD cells shown as a percentage of the wild type. Experiments were repeated at least three times and error bars indicate SD. Strains were grown on motility agar at 37°C. (TIF)

**Figure S7** Effect of rcsC expression on SipC production and motility. Panels A and B show western blots from SL1344 WT, ΔmreC, ΔrcsC, and SPI-1 control strains, and SL1344 WT pBADΔrcsC, mreC pBADrcsC, ΔrcsC pBADΔrcsC, and mreC ΔrcsC pBADΔrcsC strains (+/− arabinose) with αSipC antibody. SipC is indicated at approximately 43kDa. Panel C shows motility of SL1344 WT, mreC, SL1344 WT pBADΔrcsC, mreC pBADrcsC, ΔrcsC pBADΔrcsC, and mreC ΔrcsC pBADΔrcsC strains (+/− arabinose) shown as a percentage of the wild type. Experiments were repeated at least three times and error bars indicate standard deviation. (TIF)

**Figure S8** Percentage change in transepithelial resistance of differentiated Caco-2 cells after 4hr infection with Δrcs mutant strains. TER of polarised Caco-2 monolayers exposed to *Salmonella* strains at an MOI of 20. TER change is expressed as a percentage alteration at 4hr compared to the initial value at time zero. Error bars indicate the standard deviations derived from at least three independent experiments. *Indicates statistical difference from WT (p<0.05). (TIF)

**Figure S9** Complementation of *Salmonella* Pathogenicity Island SPI-1 in ΔmreC mutant. Expression of SPI-1 proteins in WT SL1344, ΔSPI-1, and ΔmreC mutants, and complemented ΔmreC pBADΔrbc strain during SPI-1 inducing conditions as revealed by western blotting with polyclonal αSipB antibody. SipB is indicated at approximately 63kDa, and a breakdown product is evident. (TIF)

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