Identification and initial characterisation of a protein involved in Campylobacter jejuni cell shape

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Campylobacter jejuni is the leading cause of bacterial food borne illness. While helical cell shape is considered important for C. jejuni pathogenesis, this bacterium is capable of adopting other morphologies. To better understand how helical-shaped C. jejuni maintain their shape and thus any associated colonisation, pathogenicity or other advantage, it is first important to identify the genes and proteins involved. So far, two peptidoglycan modifying enzymes Pgp1 and Pgp2 have been shown to be required for C. jejuni helical cell shape. We performed a visual screen of ~2000 transposon mutants of C. jejuni for cell shape mutants. Whole genome sequence data of the mutants with altered cell shape, directed mutants, wild type stocks and isolated helical and rod-shaped ‘wild type’ C. jejuni, identified a number of different mutations in pgp1 and pgp2, which result in a change in helical to rod bacterial cell shape. We also identified an isolate with a loss of curvature. In this study, we have identified the genomic change in this isolate, and found that targeted deletion of the gene with the change resulted in bacteria with loss of curvature. Helical cell shape was restored by supplying the gene in trans. We examined the effect of loss of the gene on bacterial motility, adhesion and invasion of tissue culture cells and chicken colonisation, as well as the effect on the muropeptide profile of the peptidoglycan sacculus. Our work identifies another factor involved in helical cell shape.

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1. Introduction

Infection by Campylobacter spp, especially Campylobacter jejuni, is considered to be the most prevalent cause of bacterial diarrhoeal disease worldwide [1]. The bacterium is found in the gastrointestinal tract of healthy animals, especially chickens, destined for human consumption. The helical shape of C. jejuni is believed to be important for the bacteria to colonise chickens and during infection, to move through the mucus layer of the gastrointestinal tract and to ‘corkscrew’ into the cells of a human (or other animal) host. There is limited understanding of how C. jejuni adopts a helical morphology. One study identified a mutation in flhB that affected flagella formation and apparently correlated with C. jejuni becoming rod-shaped [2], but mutations at other sites in the same flagellar gene resulted in bacteria that remained helical. A mutant in cj1564 (transducer-like protein 3, Tlp3) has many altered phenotypic characteristics including loss of curvature, but the mechanism for the change in shape is not clear [3]. Occasionally, laboratory strains of C. jejuni lose cell curvature and become rod
C. jejuni can also undergo a transition from helical cells to rod shaped or coccoid forms in older cultures, and under conditions of stress. It is not clear whether C. jejuni can move back and forth between different conformational states during growth. The only genes known to be involved in determination of the helical cell shape of C. jejuni are pgg1 and pgg2 [5–7], and their protein products are peptidoglycan (PG) peptidases that are important for PG modification [5,6].

The bacterial cell wall is important for providing both rigidity and shape to cells and is composed of layers of PG, or murein, which forms the murein sacculus [8]. In Gram-negative bacteria, such as C. jejuni, the murein sacculus is very thin and lies in the periplasm between the inner and outer membranes. PG is a web of glycan polymers joined by peptide side chains, which are either directly crosslinked or joined by short peptide bridges. The peptide side chains are synthesised at the inner membrane as pentapeptides and may be cleaved into shorter fragments by a number of peptidases. Peptidases may be endopeptidases or carboxy-peptidases depending on whether they cleave an internal or C-terminal amino acid, respectively. Peptidases are also classified by whether they hydrolyse the bond between two L-amino acids (DD) or between a D-amino acid and a D-amino acid (LD or DL). The number and length of peptide attached to the glycan backbone provide unique muropeptide profiles for each bacterium. The PG modification pathway in bacteria is known to contain a wide array of carboxy- and endopeptidases responsible for cleaving monomeric, dimeric and trimeric peptides [9].

To date, only two carboxypeptidases involved in cleaving monomeric peptides have been identified in C. jejuni, Pgp1 [5] and Pgg2 [6]. Pgg2 is an L,L-carboxypeptidase, which cleaves disaccharide tetrapeptides into tripeptides [6]. Pgp1 is a D,L-carboxypeptidase, which cleaves disaccharide tripeptides into dipeptides [5]. Pgp1 activity is metal-dependent and requires the activity of Pgg2 to provide the tripeptide substrate [6]. When either of the pgg1 or pgg2 genes are mutated in the laboratory the muropeptide profile radically changes and helical cell shape cannot be maintained [5,6]. Loss of pgg1 causes a decrease in dipeptides and tetrapeptides and an increase in tripeptides [5]. Loss of pgg2 causes a decrease in dipeptides and tripeptides and an increase in tetrapeptides [6]. Furthermore, overexpression of pgg1 in C. jejuni results in a kinked rod morphology, and muropeptide analysis of the pgg1 overexpressing strain demonstrates a decrease in tripeptides and an increase in dipeptides [5]. Combined, these findings suggest that even subtle changes to proportions of peptides in the PG can affect C. jejuni cell shape.

Pgg2 orthologs are present in a wide range of bacteria that display helical, rod, vibroid (curved rod) or coccoid cell shapes [6]. In contrast, Pgp1 is most highly conserved in helical and vibroid species of the Epsilon- and Delta-proteobacteria [5]. The Pgp1 ortholog in H. pylori, Csd4, has also been characterised as a necessary determinant of cell shape in this helical pathogen. A defined csd4 mutant in H. pylori generates a rod-shaped strain that exhibits a similar muropeptide profile to Δpfg1 in C. jejuni [5,10].

The conserved nature of Pgp1 in particular supports the hypothesis that this protein is fundamental to cell curvature and helical cell shape.

While it is known that peptidases can be redundant [11,12], single and double knockouts of Pgp1 and Pgg2 do not demonstrate any change to levels of peptide crosslinking [5,6], suggesting that there remain unidentified PG peptidases in C. jejuni. Thus, further identification and characterisation of the enzymes involved in PG synthesis and modification systems and how these enzymes are localised and regulated is required before we can fully understand how helical shape is generated in C. jejuni.

We recently performed a visual screen of 1933 transposon (Tn) mutants of C. jejuni for changes in cell morphology [13]. Whole genome sequence (WGS) data of the Tn mutants with altered cell shape, directed mutants, wild type (WT) stocks and isolated helical and rod-shaped ‘WT’ C. jejuni, identified a number of different genetic mutations in ppg1 and pgg2, which result in a change in helical to rod bacterial cell shape [13]. In addition, we identified an isolate with a loss of curvature. In this study, we report the genome change leading to the loss of curvature and initial characterisation of the gene.
rarrangement was confirmed by PCR and sequencing using the primers dare_c1 and ck2, respectively. Primers used in this study are listed in Table 2.

Complementation of CJ81176_1105 and CJM1_1064 in the targeted deletion strains was performed by amplifying CJM1_1064 from DNA isolated from WT helical isolates of strain M1 using primers darec_F and darec_R. The PCR product was ligated into the Campylobacter shuttle vector pCE107/70 (KmR) [18] and transformed into electrocompetent mutants. A novel genetic complementation vector, ApR, KmR [20], was used to complement CJ81176_1105 in the strain 81–176, since the transformation of C. jejuni 81–176 with pCE107/70 was unsuccessful after repeated attempts. DNA isolated from 81 to 176 WT strain served as the template for the amplification of CJ81176_1105 coding sequence using primers darec_F and darec_R. The PCR product was cloned into pSV009 (KmR) using the BamHI and Pst restriction sites. The resulting plasmid, pSV009-pgpc3C (KmR), was confirmed by PCR and sequencing. Following which, the CJ81176_1105 complementation region was amplified by PCR from pSV009-CJ81176_1105C using the primers pSV009_GCampFW1/RV1 and subsequently introduced into C. jejuni 81–176 by electroporation. Primers used in this study are listed in Table 2.

2.5. Muropeptide analysis

PG purification and digestion protocols were adapted from those described in Glauer [25], Li et al. [26] and Frirdich et al. [5]. HPLC of purified and muramidase-digested C. jejuni PG was performed in the same manner and using the same instrumentation as described in Christie et al. [27].

2.6. Motility assay

The motility of C. jejuni was quantified using motility agar made with 0.4%, 0.6%, 0.8% and 1.0% (w/v) select agar (Sigma) in MH broth. Motility agar was used to fill 6-well plates (7 ml of agar per well) 20 min prior to use. C. jejuni isolates were transferred via pipette tip from 12 h lawn growth (on MH agar plates) into each well of the motility agar. For each strain to be tested, three replicate 6-well plates were incubated for each motility agar concentration.

### Table 1

| Strain or plasmid | Relevant genotype or description | Source and/or reference |
|-------------------|---------------------------------|-------------------------|
| C. jejuni M1      | Chicken and human clinical isolate | Diane Newell [15] |
| M1 Helical        | Helical M1 wild type (M1 isolate, bacteria confirmed to be helical) | This study |
| M1 Rod            | Rod M1 wild type (INDEL in pgp1, 8A–7A, leading to a Stop at amino acid 403) | This study |
| C. jejuni 81-176  | Human clinical isolate, hyperinvasive | This study |
| 81-176_KR         | 81-176 wild type with loss of curvature | This study |
| 81-176 Helical    | Helical 81–176 wild type (81–176 isolate, bacteria confirmed to be helical) | This study |
| 81-176 Rod        | Rod 81–176 wild type (INDEL in pgp1, 8A–7A, leading to a Stop at amino acid 403) | This study |
| CJ81176_1105      | Helical 81–176 background, CJ81176_1105, KmR (loss of curvature) | This study |
| CJ81176_1105comp  | CJ81176_1105 background, CmR KmR (complemented mutant - Helical) | This study |
| CJM1_1064         | Helical M1 background, CJM1_1064, KmR (loss of curvature) | This study |
| CJM1_1064comp     | CJM1_1064 background, pDARE14, CmR KmR (complemented mutant - Helical) | This study |
| E. coli DH5α      | Subcloning Efficiency™ DH5α™ Competent Cells. F− ΔlacZΔM15 Δ(lacZY-aargF) U169 recA1 endA1 hsdR17(rk1-1 mcrB) phoA supe44 thi-1 gyr96 relA1 λ | Thermo Scientific |

### Table 2

| Primer | Target | Sequence (5′–3′) |
|--------|--------|----------------|
| dare008 | cut cassette | gaattggtagctTGGCGCGTTGTTCCCTTTCCAAAGTT |
| dare009 | cut cassette | gattcgtagctGCGCCTTATTCTCTAAAGGTTT |
| dare010 | cut cassette | gcatgctctcgcGGCCGTTGCTATCCTAAAGGTTT |
| dare011 | cut cassette | agtagctcctcggcGGCGTTGCTATCCTAAAGGTTT |
| dare_1001 | CJ81176_1105 upstream | cccggcgattAAAGTCGAGAACGAAAGCTG |
| dare_1002 | CJ81176_1105 upstream | cctgtagcgcAAATGCTCTTGAACCGTTATACCTG |
| dare_1003 | CJ81176_1105 downstream | gtgacacgctCACCATTGCACATAGACGAGG |
| dare_1004 | CJ81176_1105 downstream | gtaagcgcctcCACCATTGCACATAGACGAGG |
| dare_F | CJ81176_1105 | gtcggcgctcCACCATTGCACATAGACGAGG |
| dare_R | CJ81176_1105 | gtcggcgctcCACCATTGCACATAGACGAGG |
| dare_cl1 | CJ81176_1105 | GGCTATGCTTGATAAATTCA |
| dare_cl2 | CJ81176_1105 | AGTTCATTTAGAAAAGCAGC |
| pSV009_GCampFW1 | Genetic complementation region | TAATAGAATTTTTCTGAGCT |
| pSV009_GCampRV1 | Genetic complementation region | CATAAGCATATGTCCTTACAGCT |
| pSV009_seq_FW1 | Sequencing genetic complementation insert | GAGACATTCCCTCCGATTC |
| pSV009_seq_RV1 | Sequencing genetic complementation insert | ACGCAGACAAAAAACACCTAGC |

Upper-case indicates homology to target sequence. Restriction enzyme sites are underlined and preceded by an arbitrary 6-bp sequence.
Motility was measured as the diameter of the halo of motility after 12 h incubation.

2.7. Culture of Caco-2 cells

Caco-2 cell lines were purchased from the ATCC (CC-L244, HTB-37). Cells were grown using DMEM (Gibco) supplemented with 10% FBS and 1% non-essential amino acids. Cells were routinely grown in 75 cm² tissue culture flasks and incubated at 37 °C with 5% CO₂ in a humidified atmosphere.

2.8. Caco-2 cell infection assays

Caco-2 cells were seeded at 5 × 10⁴ cells on 24 well plates (Greiner) until confluency was observed. Caco-2 cells were infected with different C. jejuni strains at a multiplicity of infection (MOI) of 100. To assay adherence/invasion, infected cells were incubated at 37 °C with 5% CO₂ in a humidified atmosphere for 2 h. At this point, non-adherent bacteria were removed, subjected to 10-fold serial dilutions and plated on BHI blood agar plates with 0.1% Triton X-100 in PBS for 10 min. Serial dilutions of the cell lysates were carried out and plated on BHI blood agar plates with 0.1% Triton X-100 in PBS for 15 min. Lysed cells were subjected to 10-fold serial dilutions and plated on BHI blood agar plates with 5 µg/ml trimethoprim. To determine the number of internalised bacteria, infected Caco-2 cells were incubated at 37 °C with 5% CO₂ in a humidified atmosphere. After 2 h, the media overlaying the infected cells was changed to complete DMEM containing 250 µg/ml gentamycin sulphate and infected cells were incubated at 37 °C with 5% CO₂ in a humidified atmosphere for a further 2 h. Cells were then washed three times with PBS and lysed with 0.1% Triton X-100 in PBS for 10 min. Serial dilutions of the cell lysates were carried out and plated on BHI blood agar plates with 5 µg/ml trimethoprim. Dilutions of mutant C. jejuni strains were plated on BHI blood agar plates containing 10 µg/ml chloramphenicol, whereas genetically complemented strains were recovered on MH blood agar plates supplemented with 10 µg/ml chloramphenicol and 50 µg/ml kanamycin. All plates were incubated for 48 h under microaerophilic conditions at 42 °C before colony counting took place. For both total association and invasion experiments, the percentage of C. jejuni interacting with Caco-2 cells was calculated as a percentage of the non-adherent fraction, to account for various strain survival within DMEM (n = 3).

2.9. Chicken colonisation experiments

All work was conducted in accordance with UK legislation governing experimental animals under project licence 40/3652 and was approved by the University of Liverpool ethical review process prior to the award of the licence.

One-day-old Ross 308 broiler chicks were obtained from a commercial hatchery. Chicks were housed in the University of Liverpool, High Biosecurity Poultry unit. Chicks were maintained in floor pens at UK legislation recommended stocking levels allowing a floor space of 2,000 cm² per bird and were given ad libitum access to water and a pelleted laboratory grade vegetable protein-based diet (SDS, Witham, Essex, UK). Chicks were housed in separate groups at a temperature of 30 °C, which was reduced to 20 °C at 3 weeks of age. Prior to experimental infection, all birds were confirmed as Campylobacter-free by taking cloacal swabs, which were streaked onto selective blood-free agar (mCCDA) supplemented with Campylobacter Enrichment Supplement (SV59; Mast group, Bootle, Merseyside, UK) and grown for 48 h in microaerobic conditions at 41.5 °C. All microbiological media were purchased from Lab M (Heywood, Lancashire, UK).

At 21 days of age birds were infected with 2 × 10⁶ CFU of either C. jejuni M1 or the CJM1_1064 mutant. At 5 days post infection (p.i.), chickens were killed by cervical dislocation. At necropsy the ceca were removed aseptically and the cecal contents plated onto mCCDA Campylobacter selective agar plates for enumeration as previously described [28].

3. Results and discussion

3.1. Identification of a C. jejuni 81–176 isolate with a loss of curvature

When isolating helical and rod bacteria from WT C. jejuni strains based on colony morphology as described in Esson et al. [13], we noticed a colony, which was not quite as grey and flat as the typical rod colony morphology but still distinct from the helical colony morphology, and was composed of bacteria with a loss of curvature (‘kinked rod’ cell morphology, 81176_KR). This cell morphology contrasted with the helical morphology of WT 81–176 and was confirmed by scanning electron microscopy (SEM) (Fig. 1). Based on a qualitative assessment of our SEM analyses, there appears to be a difference in the degree of curvature between shorter (presumably younger cells) and longer (presumably older) cells of the ‘kinked

Fig. 1. Scanning electron micrographs of helical and loss of curvature morphologies of C. jejuni 81–176. (a) 81–176 helical isolate and (b) isolate 81176_KR from the WT C. jejuni 81–176 laboratory frozen stock. Scale bars represent 2.5 µm.
rod’ bacteria.

3.2. Whole genome sequence analysis of the 81–176 isolate with a loss of curvature

The 81–176 isolate with a loss of curvature (81176_KR) was analysed by WGS and was a change in the number of bases in a documented phase variable region (PVR), and two unique point mutations.

Phase variation (PV) enables genetic and phenotypic variation in a number of bacteria, including *C. jejuni* [29–31]. Regions of the bacterial genome that are prone to these reversible mutations are called PVR. In *C. jejuni* the PVRs are typically homopolymeric tracts (HTs) that are highly susceptible to slipped-strand mispairings, which alter the length of the tracts and generate frameshift mutations during DNA replication and repair [32,33]. In this way, PVRs are able to randomly switch genes ‘on’ and ‘off’ and stochastically regulate gene expression [32]. The unique PV pattern was in PVR3 and demonstrated a mostly ‘on’ length, which contrasted with the mostly ‘off’ lengths of other 81–176 isolates. PVR3 in strain 81-176 is 118 bp upstream of *CJJ81176_0590*, encoding a putative uncharacterised protein. This PVR correlates to PVR2 in strain M1, which demonstrated ‘on’ lengths in helical M1 Tn mutants and WT isolates (data not shown). For this reason, as well as the absence of a *CJJ81176_0590* ortholog in strain NCTC11168 [29], we hypothesised that the altered polyG tract upstream of *CJJ81176_0590* was not responsible for the loss of curvature of isolate 81176_KR.

One of the unique point mutations in 81176_KR was a non-synonymous SNP (G > A) in *rpiB* at base location 860819 (CP000538.1). This SNP was predicted to cause a single glutamic acid to lysine amino acid change. The protein product of *rpiB*, ribose 5-phosphate isomerase B, is involved in carbohydrate metabolism [34] and our searches did not demonstrate any link between this enzyme and bacterial cell shape. Therefore, we hypothesised that a single amino acid change to RpiB was not responsible for the observed loss of curvature in 81176_KR.

The other point mutation detected in 81176_KR was an INDEL in *CJJ81176_1105*, a predicted LytM peptidase-encoding gene. This single guanine deletion (2G > G) at base location 1022254 (CP000538.1) was predicted to cause a truncation at residue 65 of the 300 amino acid protein product. BLAST analysis of *CJJ81176_1105* revealed that this gene is highly conserved (>50% coverage and >70% identity) in helical Campylobacter spp. (data not shown). We investigated the presence and allelic variances of

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**Fig. 2.** Gene locus and targeted deletion of *CJJ81176_1105* (81–176) and *CJM1_1064* (M1). (a) A targeted deletion of *CJJ81176_1105* was generated in the 81–176 and M1 (CJM1_1064) backgrounds by exchanging the gene with a cat cassette (CmR). The cat cassette along with the flanking regions indicated (CJM1_1064), was cloned into the suicide vector pUC19 (pDARE10, M1 derivative and pDARE12, 81–176 derivative, CmR). A complementing plasmid (pDARE14, M1 derivative, and pSV009- *CJJ81176_1105*, 81–176 derivative, KmR) was generated by cloning M1 CJM1_1064 into pCE107/70, a kanamycin-resistant shuttle vector. (b) CJM1_1064 and *CJJ81176_1105* displayed loss of curvature, complementation with pDARE14 (M1) or pSV009- *CJJ81176_1105* (81–176) (supplying *CJJ81176_1105* in trans), strains CJM1_1064comp and *CJJ81176_1105*comp, rescued the morphology back to helical.


CJJ81176_1105 in 859 genomes of C. jejuni and C. coli. The genomes were from a wide range of isolates: 192 from clinical, agricultural and wild bird sources [35], 319 from multiple stages of poultry processing, including farms, abattoirs and retail chicken meat [36] and 348 from clinical cases [37]. Analysis of these genomes revealed that CJJ81176_1105 is conserved (data not shown), suggesting this gene is core to both Campylobacter species.

### 3.3. Bioinformatic analysis of CJJ81176_1105

A detailed comparison of the translated sequence of CJJ81176_1105 from four laboratory C. jejuni strains (M1, 81116, 81–176 and NCTC11168) demonstrated identical amino acid sequences at all except four residues (Fig. S1). The protein product of CJJ81176_1105 contains prefoldin, coiled-coil and peptidase domains. Prefoldin is a coiled-coil-containing molecular chaperone that assists in the proper folding of polypeptide products [38]. In eukaryotes, prefoldin is responsible for the folding and localisation of the cytoskeleton components actin and tubulin [39]. The peptidase domain is conserved within the Peptidase M23 (LytM) family, which is composed of zinc-dependent endopeptidases often involved in cell division, elongation and shape determination [40].

Further analysis revealed CJJ81176_1105 to be orthologous to csd1 (cell shape determinant 1) in H. pylori [41]. In the helical pathogen H. pylori, a targeted deletion of csd1 results in a curved rod morphology, which is fully complemented when csd1 is supplied elsewhere on the chromosome [41]. Sycuro et al. [41] compared the Csd1 protein product from H. pylori to the crystallised LytM endopeptidase from Staphylococcus aureus, which demonstrated conserved LytM active site residues in Csd1.

Although the csd1 ortholog in C. jejuni was identified by Sycuro et al., the gene was unable to complement the Δcsd1 phenotype in H. pylori [41]. Another group also investigated the role of the csd1 ortholog in C. jejuni morphology but results from these preliminary studies were reported as inconclusive (unpublished work mentioned in Frirdich et al. [5]).

Due to the sequence similarity between csd1 and CJJ81176_1105 and the similar ‘intermediate’ morphologies of the H. pylori Δcsd1 strain and 81176_KR, we hypothesised that the frameshift mutation in CJJ81176_1105 was responsible for loss of curvature morphology of 81176_KR. Moreover, due to its predicted endopeptidase function [41,42], we hypothesised that the CJJ81176_1105 protein product might be involved in the same PG modification cascade as Pgp1 and Pgp2.

### 3.4. Defined gene deletion mutants of CJJ81176_1105 alter C. jejuni motility and interaction with Caco-2 cells, but not chicken colonisation

To test whether the mutation in CJJ81176_1105 was responsible for the loss of curvature of 81176_KR, we constructed targeted deletions and complemented strains, on different C. jejuni WT backgrounds (CJJ81176_1105 for strain 81–176 and CJM1_1064 for strain M1). The defined mutants displayed loss of curvature morphologies, which were restored to helical morphologies by complementation (Fig. 2). From these data, we conclude that CJJ81176_1105, and its homolog in other strains, is necessary for a fully helical morphology in C. jejuni. We next compared physiological characteristics of the CJJ81176_1105 and CJM1_1064 mutants and complemented strains with helical-shaped and rod-shaped WT isolates.

We tested the motility of WT helical and rod isolates, against the CJJ81176_1105 and CJM1_1064 mutants and complemented strains, across a range of motility agar concentrations (Fig. 3). The results showed that migration through increasing agar concentrations (decreasing porosity) was significantly reduced in the WT-rod (INDEL in pgp1) and CJJ81176_1105 and CJM1_1064 mutants compared to the helical isolates (WT-helical and complemented strains). Our work demonstrates that at 0.4 and 0.6% (w/v) agar, the motility of the CJJ81176_1105 and CJM1_1064 mutants is slightly greater (although not statistically significant) from the WT-rod isolates. At 0.8% (w/v) agar the motility of the WT-rod (INDEL in pgp1) and CJJ81176_1105 and CJM1_1064 mutants were comparable, and significantly reduced from the helical isolates. All the isolates were effectively non-motile through 1.0% (w/v) agar, i.e. all isolates measured 1 mm in diameter, roughly equivalent to the original pipette stab.

Next, the ability of the CJJ81176_1105 and CJM1_1064 mutants to adhere to, and invade, Caco-2 cells was measured (Fig. 4). The WT-rod and CJJ81176_1105 and CJM1_1064 mutants displayed statistically significant reductions in adhesion and invasion compared to the WT-helical and complemented strains. For both C. jejuni backgrounds, M1 and 81–176, the adherence and invasion of the mutant was slightly greater (although not statistically significant) from the WT-rod isolate (INDEL in pgp1).

Chicken colonisation experiments were performed using C. jejuni strain M1 WT-helical isolate, a natural poultry isolate which is an efficient coloniser of chickens [15], and the CJM1_1064 mutant. Chickens were inoculated with 2 × 10⁹ CFU of either strain. At 5 days post infection (p.i.), chickens were killed and the cecal contents plated onto mCCDA Campylobacter selective agar plates as previously described [28]. The viable counts per gram of cecal contents revealed that there was no difference in the colonisation of the WT or the CJM1_1064 mutant (Fig. 5).
Fig. 4. Adhesion (a) to, and invasion (b) of Caco-2 cells by C. jejuni M1 and 81–176 rod and helical isolates, CJ81176_1105 and CJM1_1064 mutants and complemented strains. Data is represented as percentage of wild-type (n ≥ 3) and plotted as means and SEM. Statistical significance was calculated using a Mann-Whitney test where * P < 0.05 and **P < 0.005.
3.5. Muropeptide analysis of helical WT and CJ81176_1105 (and CJM1_1064) mutant C. jejuni

Muropeptide analysis via high-performance liquid chromatography (HPLC) and mass spectrometry (MS) was used to compare the PG sacculi of WT, CJ81176_1105 and CJM1_1064 mutant and complemented C. jejuni strains. Muropeptide profiles of mutanolysin-digested PG sacculi isolated from CJ81176_1105 and CJM1_1064 deletion strains appeared virtually identical to muropeptide profiles derived from the parental strains (Fig. 6). Hence in contrast to Pgp1 and Pgp2, CJ81176_1105 (and CJM1_1064) activity in modelling the PG sacculus appears to be below the threshold for detection via the muropeptide analysis technique, and definitive identification of the hydrolytic bond specificity against PG will require further attention.

4. Conclusion

There is no effective vaccine against C. jejuni and preventative measures aimed at reducing environmental contamination have so far proved ineffective. There is a need for alternative strategies to reduce campylobacteriosis. Most of the Campylobacteraceae are helical and it appears that the helical shape of C. jejuni is important for its ability to colonise its hosts and cause disease. To address this hypothesis, it is essential to know how helical shape is determined in C. jejuni, both genetically and biochemically, but we currently have limited understanding of this. Loss of helical cell shape through interference may hold therapeutic potential by reducing this pathogen’s virulence or ability to colonise animals.

This work identifies CJ81176_1105 as a novel cell shape
determinant in C. jejuni. CJ81176_1105 was identified by the isolation and WGS analysis of a bacterium with loss of curvature within our laboratory WT C. jejuni 81-176 stock. This isolate was found to contain a nonsense mutation in CJ81176_1105. Targeted deletions of the gene in both the C. jejuni 81-176 and M1 backgrounds reproduced the loss of curvature morphology of the original isolate. This intermediate cell shape was rescued by supplying the gene in trans, which confirmed that it is necessary for a fully-helical C. jejuni morphology. The homology of CJ81176_1105 to the endopeptidase Csd1 in H. pylori suggests that it may also be involved in this muropeptide cascade.

The intermediate nature of the loss of curvature morphology implies that there exists a hierarchy to helical cell shape maintenance within C. jejuni. Based on the evidence demonstrating the importance of endo- and carboxypeptidases in helical cell shape [5,6,10,41], this hierarchy is likely a product of PG peptide lengths and the degree of crosslinking that promotes the cell wall to twist.

As a possible explanation for the differences between the helical, loss of curvature and rod forms of C. jejuni, we hypothesise that shorter peptides within the cell wall are localised to the inside of the helix. As such, while the loss of di- and tripeptides in pgp1 and pgp2 mutants prevents the maintenance of any curvature [5,6] the predicted reduction of tetra- and pentapeptide substrates in the CJ81176_1105 mutant may merely ratios of the distribution of di- and tripeptide products throughout the PG, lessening the tension of the helix. To address this hypothesis, future work will require an investigation into the distribution of PG peptides throughout the cell wall in situ.

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Glossary

| Abbreviation | Description |
|--------------|-------------|
| BHI          | brain heart infusion |
| Cat          | chloramphenicol acetyl transferase |
| CFU          | colony forming units |
| Cm           | chloramphenicol |
| DMEM         | Dulbecco's Modified Eagle's Medium |
| FBS          | fetal bovine serum |
| HPLC         | high performance liquid chromatography |
| INDELS       | insertions and deletions |
| Km           | kanamycin |
| mCCDA        | modified charcoal-cefooperazone deoxycholate agar |
| MH           | Mueller Hinton |
| MOI          | multiplicity of infection |
| MS           | mass spectrometry |
| PBS          | phosphate buffered saline |
| PG           | peptidoglycan |
| p.i.         | post infection |
| PV           | phase variable |
| PVR          | phase variable region |
| RE           | restriction enzyme |

SEM scanning electron microscopy
SNPs single nucleotide polymorphisms
Tn transposon
Tp trimethoprim
WGS whole genome sequence
WT wild type
w/v weight per volume

Appendix A. Supplementary data
Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.micpath.2017.01.042.

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