Modification of the Campylobacter jejuni N-Linked Glycan by EptC Protein-mediated Addition of Phosphoethanolamine

Received for publication, May 9, 2012, and in revised form, June 20, 2012. Published, JBC Papers in Press, July 2, 2012, DOI 10.1074/jbc.M112.380212

Nicholas E. Scott†§1, Harald Nothaft†, Alistair V. G. Edwards§, Maurizio Labbate‖, Steven P. Djordjevic‖, Martin R. Larsen*, Christine M. Szymanski§, and Stuart J. Cordwell†§

From the †School of Molecular Bioscience, and Discipline of Pathology (School of Medical Sciences), The University of Sydney, Australia 2006, ‡Albetta Glycomics Centre and Department of Biological Sciences, University of Alberta, Edmonton, Alberta T6G 2E9 Canada, §Protein Research Group, Department of Biochemistry and Molecular Biology, University of Southern Denmark, Odense Denmark 5000, and †‖3 Institute, University of Technology Sydney, Sydney, Australia 2007

Background: C. jejuni produces an N-linked heptasaccharide that is attached to multiple proteins and has been linked with full virulence.

Results: A modified N-glycan displaying a phosphoethanolamine (pEtN) moiety linked to the terminal GalNAc was identified attached to 9 proteins.

Conclusion: The addition of pEtN to the N-glycan is mediated by the pEtN transferase EptC.

Significance: Modification of the N-glycan by pEtN confirms that EptC targets multiple substrates in C. jejuni.

Campylobacter jejuni is the major worldwide cause of bacterial gastroenteritis. C. jejuni possesses an extensive repertoire of carbohydrate structures that decorate both protein and non-protein surface-exposed structures. An N-linked glycosylation system encoded by the pgf gene cluster mediates the synthesis of a rigidly conserved heptasaccharide that is attached to protein substrates or released as free oligosaccharide in the periplasm. Removal of N-glycosylation results in reduced virulence and impeded host cell attachment. Since the N-glycan is conserved, the N-glycosylation system is also an attractive option for glycobiology recombination of recombinant vaccines in Escherichia coli. To determine whether non-canonical N-glycans are present in C. jejuni, we utilized high throughput glycoproteomics to characterize C. jejuni JHH1 and identified 93 glycosylation sites, including 34 not previously reported. Interrogation of these data allowed the identification of a phosphoethanolamine (pEtN)-modified variant of the N-glycan that was attached to multiple proteins. The pEtN moiety was attached to the terminal GalNAc of the canonical N-glycan. Deletion of the pEtN transferase eptC removed all evidence of the pEtN-glycan but did not globally influence protein reactivity to patient sera, whereas deletion of the pgIB oligosaccharyltransferase significantly reduced reactivity. Transfer of eptC and the pgl gene cluster to E. coli confirmed the addition of the pEtN-glycan to a target C. jejuni protein. Significantly reduced, yet above background levels of pEtN-glycan were also observed in E. coli not expressing eptC, suggesting that endogenous E. coli pEtN transferases can mediate the addition of pEtN to N-glycans. The addition of pEtN must be considered in the context of glycobiology and may alter C. jejuni glycamediated structure-function interactions.

Campylobacter jejuni is a Gram-negative pathogen responsible for a major proportion of food (typically poultry-derived)- and water-borne diarrheal illness worldwide, with an estimated 1% of the population in the United Kingdom and United States infected annually (1, 2). Prior infection with C. jejuni has also been associated with development of immune-mediated sequelae such as Reiter’s syndrome (3), Guillain-Barré Syndrome (4), and immunoproliferative small intestinal disease (5). Despite studies to identify factors responsible for C. jejuni virulence in humans and colonization of the poultry host, our understanding of these processes remains incomplete (6, 7). Virulence-associated factors include cell-surface structures such as carbohydrates (8–10) and post-translationally modified flagellar and membrane proteins (11–15). These components generate significant variability that manifests in both the derived chemical structures and between C. jejuni isolates. Variations in lipooligosaccharide (LOS), 3 capsule polysaccharide, and O-linked glycosylation of the flagella structural subunit flagellin are thus a quintessential feature of C. jejuni biology (16–21).

C. jejuni contains a conserved N-glycosylation system responsible for the modification of membrane-associated, periplasmic, and secreted proteins (22) and for the generation of a “free” oligosaccharide (fOS) involved in osmotic stability (23). The N-linked glycan is a heptasaccharide (GalNAc–α1,4-GalNAc–α1,4-[GlcB1,3]-GalNAc–α1,4-GalNAc–α1,3-Bac–β1, where Bac is bacillosamine (2,4-diacetamido-2,4,6 trideoxyxylpyranylos)) encoded by the pgf gene cluster (24–26) and is attached to proteins (22, 27) in the periplasm by

* This work was supported by Australian Research Council Discovery Project Grant ARC DP110103573 (to S. J. C.).

† Supported by an Australian Postgraduate Award and a current National Health and Medical Research Council of Australia Overseas (Biomedical) Fellow (APP1037733).

To whom correspondence should be addressed: School of Molecular Biosciences, Bldg. GO8, The University of Sydney, Australia 2006. Tel.: 61-2-9351-6050; E-mail: stuart.cordwell@sydney.edu.au.

3 The abbreviations used are: LOS, lipooligosaccharide; CID, collision-induced dissociation; fOS, free oligosaccharide; HCD, higher energy collisional dissociation; XIC, extracted ion chromatogram; ZIC-HILIC, zwitterionic hydrophilic interaction liquid chromatography; MH, Mueller-Hinton; BHI, brain heart infusion; pEtN, phosphoethanolamine.
the PglB oligosaccharyltransferase (28) at the consensus sequon (D/E)AX(N/S/T) (where X ≠ proline (29)). Removal of components of the N-glycosylation pathway results in reduced adherence to and invasion of gut epithelial cells and lowered colonization of the chicken gastrointestinal tract (11–12). Analysis of fOS (30), N-linked glycan (22, 30), and pathway intermediates (e.g. lipid-bound glycan (31)) suggest that the heptasaccharide is the sole N-glycan formed, although enzymes within the pgl biosynthetic pathway appear to have broader specificity than the substrates used to construct the canonical glycan (25, 32–33). Monoacetylated Bac in the laboratory passaged NCTC 11168 strain (32) remains the only glycan variant identified to date. As passaged C. jejuni isolates differ in the manifestation of phenotypes associated with motility, morphology, and virulence (34, 35), it is unclear if the utilization of monoacetylated Bac is an authentic process common to all C. jejuni.

In contrast to the apparently conserved nature of the N-glycan, other carbohydrate structures such as capsule polysaccharide and LOS are subject to phase variation and can also be modified by phosphate-containing moieties, including phosphoramidate (36–37) and phosphoethanolamine (pEtN (19, 38)). The addition of similar moieties to protein substrates has been documented in other pathogens, most notably the decoration of Neisseria gonorrhoeae pilin with pEtN, phosphocholine (39, 40), and phosphoglycerol (41, 42). C. jejuni also modifies the flagellar rod protein FlgG with pEtN, and this addition is mediated by the pEtN transferase Cj0256 (Ref. 15; recently named EptC (43)). Deletion of eptC resulted in a decrease in motility and increased sensitivity to polymyxin B (15). Modification of FlgG with pEtN is required for full motility (43), whereas pEtN modification of lipid A contributes to polymyxin B resistance (15).

Here we report the identification of multiple proteins modified with an N-linked glycan displaying an additional moiety of +123.01 Da, consistent with the presence of pEtN. High resolution tandem mass spectrometry (MS/MS) confirmed that pEtN was attached to the terminal GalNAc of the canonical N-glycan rather than as a direct modification of the protein substrate. Deletion of eptC (cJ0256) from C. jejuni JHH1 confirmed that EptC generates the pEtN-glycan. Heterologous expression of EptC in combination with the pgl cluster in E. coli led to the production of a pEtN-glycan-modified C. jejuni substrate protein (AcrA (44)). Modification with pEtN may influence C. jejuni and host-pathogen structure-function relationships mediated by the N-linked glycan.

**EXPERIMENTAL PROCEDURES**

**Bacterial Strains and Growth Conditions**—Bacterial strains used in this study are provided in supplemental Table S1. C. jejuni were cultured in parallel on 100 individual Skirrow’s agar plates in a micro-aerophilic environment of 5% O2, 5% CO2, and 90% N2 at 37 °C for 48 h. Collection of cells and generation of protein extracts for glycoproteomics were previously described (27). Growth of C. jejuni wild-type, and deletion/complementation mutants was performed in Mueller-Hinton (MH) or brain heart infusion (BHI) medium supplemented with 1% yeast extract and 7% blood (BHI+). Escherichia coli were grown in Luria-Bertani (LB) broth or on agar at 37 °C under ambient oxygen conditions. Where required for selection, 30 μg/ml kanamycin, 25 μg/ml chloramphenicol, 25 μg/ml trimethoprim, and 100 μg/ml ampicillin were used.

**Protease Digestion for Glycopeptide Enrichment**—Dried proteins were resuspended in 6 μl urea, 2 μl thiourea, 40 mM NH4HCO3, and reduced/alkylated before digestion with Lys-C (1/200 w/w) and then trypsin (1/50 w/w) as previously described (27). For pepsin and thermolysin digestion, proteins were reduced/alkylated and processed according to Chen et al. (45). Briefly, for pepsin digests, samples were diluted 1:4 with 0.1% trifluoroacetic acid (TFA) and adjusted to a pH of ~2.5 with 10% TFA. 1:25 (w/w) of pepsin to protein was added, and digestion was allowed to proceed for 24 h at 25 °C. For thermolysin digestion, samples were adjusted to a dilution of 1:4 with 100 mM NH4HCO3, and 1:25 (w/w) thermolysin to protein added. Samples were incubated for 24 h at 25 °C. All peptide digests were dialyzed against ultrapure water overnight using a Mini Dialysis kit with a molecular mass cutoff of 1000 Da (Amersham Biosciences) and on completion were collected and lyophilized.

**Identification of Glycopeptides Using Zwitterionic–Hydrophilic Interaction Liquid Chromatography (ZIC-HILIC) Enrichment and Reversed Phase LC-MS/MS—ZIC-HILIC enrichment was performed according to Scott et al. (27) with minor modification. Essentially, 100 μg of starting peptide was used to improve detection of the 204 m/z GalNac oxonium ion based on extracted ion chromatograms (XIC) compared with the previously reported 20 μg enrichment (27). ZIC-HILIC fractions were resuspended in 0.1% formic acid and separated using a trapless EASY-nLC system (Proxeon, Odense Denmark) coupled to an LTQ-Orbitrap Velos mass spectrometer (Thermo Scientific, San Jose CA) as in Scott et al. (27). The instrument was operated using Xcalibur v2.2 (Thermo Scientific) with a capillary temperature of 200 °C in a data-dependent mode automatically switching between MS and higher energy collisional dissociation (HCD) MS/MS. The rationale for employing HCD fragmentation alone (rather than collision-induced dissociation (CID) and HCD (27) was that HCD spectra contain both glycan diagnostic ions (e.g. 204.08 m/z; GalNAc) and peptide fragment ions allowing glycopeptide identification. For each MS scan, the three most abundant precursor ions were selected for HCD fragmentation (normalized collision energy 45). MS resolution was set to 60,000 with an AGC of 1e6, maximum fill time of 500 ms, and a mass window of 600–2000 m/z. HCD fragmentation was carried out with an AGC of 2e4, maximum fill time of 500 ms, and mass window 170–2000 m/z. For definition of unusual glycan structures on glycopeptides, CID was employed using an LTQ-Orbitrap Velos mass spectrometer with automatic switching between MS and CID MS/MS. For each MS scan, the 10 most abundant precursor ions were selected for fragmentation with normalized collision energy 35. MS parameters were set as above, whereas CID fragmentation was carried out with an AGC of 2e4, maximum fill time of 100 ms. The mass window for CID was dynamically defined by the selected ion m/z with an upper m/z limit of 2000 and lower m/z of 28% of the selected m/z.

**Data Base Interrogation of Identified Glycopeptides**—Raw files were processed in Proteome Discover v1.0 Build 43
**pEtN Modification of the C. jejuni N-linked Glycan**

(Thermo Scientific) to generate .mgf files and searched using MASCOT against a composite data base composed of the HB93-13, RM1221, 81-176, NCTC11168, and 81116 genomes in FASTA format. Scan events that did not result in peptide identification were exported to GPMAW 8.2 (Lighthouse Data, Odense Denmark) “mgf graph,” which generated .mgf files for MS/MS spectra containing the GalNAc diagnostic oxonium 204.086 m/z ion. These scans were manually annotated based on the presence of the deglycosylated peptide ion (parent mass minus 1405.561 Da and corresponding to the elemental composition C_{32}H_{51}N_{7}O_{34} of the *C. jejuni* glycan) within a tolerance of 20 ppm. To facilitate glycopeptide assignments from HCD scan events, ions below the mass of the predicted deglycosylated peptides were extracted with Xcalibur v2.2 using the Spectrum list function. Ions with a deconvoluted mass above the deglycosylated peptide mass and ions corresponding to known carbohydrate oxonium ions such as 204.08, 366.14, and 407.16 m/z were removed by post-spectral processing. MASCOT v2.2 searches were conducted via the Australasian Proteomics Computational Facility with the *C. jejuni* taxonomy selected. Searches were carried out with a parent ion mass accuracy of 20 ppm and a product ion accuracy of 0.02 Da with no protease specificity and instrument selected as MALDI-QIT-TOF (use of this setting was due to the observation of multiple internal cleavage products and extensive NH3 and H2O loss). Searches were carried out with a parent ion mass accuracy of 20 ppm and a product ion accuracy of 0.02 Da with no protease specificity and instrument selected as MALDI-QIT-TOF (use of this setting was due to the observation of multiple internal cleavage products and extensive NH3 and H2O loss). A MASCOT ion score cut-off of 20 was accepted for positive identifications, and all data were searched with the decoy setting activated to generate a zero false positive rate against a decoy data base. XIC corresponding to ±0.025 m/z of the monoisotopic peak of identified glycopeptides were generated within Xcalibur. Peaks were processed with a 15-point gaussian smooth and the corresponding ions inspected to ensure correct charge state.

Construction of *C. jejuni* JHH1 Chromosomal Mutants—To create the *C. jejuni* JHH1 ΔpglB mutant, primers PglBF and PglBR (supplemental Table S1) were used to amplify pglB::kan from genomic NCTC11168 ΔpglB DNA (22). The resulting PCR product was inserted into the pGEM-T easy vector. Positive insertions were confirmed by PCR, and the resulting plasmid was named pNS1 (supplemental Table S1). To create the ΔeptC mutant, up- and downstream regions of eptC were amplified using 0256-1-PstI-F, 0256-2-HincII-R, 0256-3-HincII-F, and 0256-4-Xhol-R from JHH1 genomic DNA. PCR amplicons digested with HincII and PstI or Xhol, dependent on the amplicon, were introduced into PCR script digested with PstI and Xhol generating PCScript-cj0256up/down. The APH(3′)-III phosphotransferase gene was gel-isolated after digestion of plasmid pWM10 with Smal. The resulting product was inserted into PCScript-cj0256up/down digested with HincII. The candidates were antibiotic-selected, and the resulting plasmid was named pNS2. Plasmids (pNS1 and pNS2) were used to create disruptions in the JHH1 background as described (46). *C. jejuni* JHH1 from overnight confluent plates of MH or BHI+ were resuspended in 15% glycerol, 9% sucrose, and washed 5 times before electroporation using a Micropulser™ (Bio-Rad) at 2.48 kV uniformly achieving ~5–6 ms electroporation events. Cells were diluted with prewarmed BHI and recovered on BHI+ plates for 5 h. Cells were then collected by centrifugation at 8000 × g for 2 min at 25 °C and plated on BHI+ plates containing 30 μg/ml kanamycin. Plates were incubated for up to 4 days at 37 °C, 5% O₂, 5% CO₂, and 90% N₂. The resulting colonies were evaluated for pglB and eptC disruption by PCR.

Western Blotting—Proteins were separated by SDS-PAGE gels and transferred to polyvinylidene difluoride membrane. Membranes were blocked overnight in 5% bovine serum albumin (Sigma) and probed with either α-N-linked glycan (1/10,000, (23)), α-His (1/3000; Rockland, Gilbertsville PA), α-HA (1/4000, Santa Cruz, Santa Cruz, CA), anti-JlpA (1/1000, (47)), or patient sera samples (1/2000, kindly provided by Dr. S. Cawthraw (48)).

Induction of eptC and Isolation of pEtN-glycan-modified AcrA from *E. coli—*E. coli CLM24 cells containing pACYC and pWA2 and either pMLBAD or pNS3 from overnight growth were used to inoculate 100-ml cultures. On achieving an A_{600} of 0.7, cells were induced by the addition of arabinose to 0.2% (w/v). After induction at 37 °C for 5 h, arabinose was added again to ensure expression during overnight cultivation. Cells were harvested by centrifugation at 6000 × g, 4 °C, stored at ~80 °C, and freeze-dried. Cells were resuspended in 10 mM imidazole PBS and sonicated for 4 × 1 min with 1 min on ice between rounds. Cellular debris was removed by centrifugation, and soluble AcrA was purified by nickel-nitritoltriacetic acid as previously described (44). Purified AcrA was separated using 10% SDS-PAGE and subjected to in-gel tryptic digestion (49).

Motility Assay—Motility assays were conducted using semi-solid MH medium supplemented with 0.4% agar. Plates were inoculated using 1 μl of overnight biphasic culture (A_{600} of ~0.5). Plates were incubated for 24 h at 37 °C under microaerophilic conditions, and the diameter of motility was measured.

Determination of Polymyxin B Resistance—Minimum inhibitory concentrations were determined using polymyxin B Etest® strips (Biomérieux, France). Confluent overnight plates of *C. jejuni* were harvested and normalized to A_{600} 0.35, and a 1:10 dilution of culture was then incubated in pre-warmed BHI (with 1% yeast extract) at 37 °C under microaerophilic conditions for 4 h. This diluted culture was then mixed 1:10 with 50°C NZCYM medium (Oxoid, Basingstock UK) supplemented with 0.6% agar. NZCYM top agar was gently poured onto MH plates and allowed to dry. An Etest® strip was placed on the center, and the plates were incubated at 37 °C for 24 h.

**RESULTS**

Identification of a pEtN-modified N-Linked Glycan in *C. jejuni*—To determine whether *C. jejuni* is capable of synthesizing novel N-glycans and attaching them to proteins, whole cell lysates from *C. jejuni* JHH1 were proteolyzed in parallel with thermolysin, pepsin, and trypsin, and the resulting digests were subjected to glycoproteomics analysis (27). A total of 12 ZIC-HILIC enrichments (biological replicates for each digest and protein lysate) were subjected to LC-MS, and 263 unique glycopeptides representing 93 glycosylation sites from 58 gly-
pEtN Modification of the C. jejuni N-linked Glycan

coproteins were identified (supplemental Table S2). These data contained 34 previously uncharacterized glycosylation sites and 15 novel glycoproteins (supplemental Table S3).

We next interrogated the dataset to determine whether modifications to the canonical N-linked glycan could be identified. Manual inspection of HCD MS/MS spectra was undertaken, with an emphasis on high quality spectra that did not generate an acceptable MASCOT score and that contained the diagnostic GalNAc oxonium ion, 204.08 m/z. Such spectra are more likely to contain a modified N-glycan as we relied on high mass accuracy within MS scans and included the predicted mass of the canonical heptasaccharide (1405.561 Da) as a fixed modification. Approximately 5000 MS/MS spectra were examined. We observed no evidence of monoacetylated Bac-containing N-linked glycans. We detected several highly charged (z \geq 3) parent ions that generated fragment ions consistent with the canonical heptasaccharide, but with an overall increased glycan mass of \sim 123.01 Da, as determined by Y0 (deglycosylated peptide) and Y1 (peptide + bacillosamine) fragment ions compared with the parent ion mass. None of these spectra generated MASCOT identifications of the glycopeptide. Manual definition of the y- and b-ion series generated by fragmentation of these parent ions confirmed peptide sequences (Fig. 1A). As the mass of 123.01 Da does not correspond to a known sugar, we searched the small molecule METLIN database to identify possible candidates (mass of 142.03 ± 0.02 Da; corresponding to 123.01 + 19.018 [H2O and H+] Da). METLIN retrieved pEtN (MH+ = 142.0264) as the most likely candidate.

In MS/MS (CID or HCD), pEtN generates internal fragment ions resulting from the loss of ethanolamine followed by phosphate (Fig. 1B). We manually inspected the +123.01 Da-containing glycopeptide dataset and in each case we observed fragment ions corresponding to those predicted for pEtN as well as internal fragment ions showing a pEtN-GalNAc linkage (Fig. 1C). To define the location of pEtN in the N-glycan, we performed CID MS/MS, which predominantly generates glycan-associated fragment ions (27). This confirmed the location of pEtN attached to the terminal GalNAc in the modified N-glycan (Fig. 2, A–C).

We next refined our spectral processing approach to enhance data base searching by removing pEtN-glycan fragment ions before analysis. MS/MS spectra were then reanalyzed, and a total of 8 pEtN-glycan-modified peptides were confidently identified (minimum MASCOT ion score of 20) from 7 glycoproteins (Table 1 and supplemental Fig. S1). Further support for these pEtN-glycan-modified glycopeptides was provided by the LC elution profiles of pEtN-glycan-modified glycopeptides compared with their canonical N-glycan-modified forms (supplemental Fig. S2). Glycopeptide species were chromatographically related, with pEtN-glycan-modified peptides typically eluting \sim 1–2 min before their canonical glycopeptide forms.

An N-Linked Glycan-modified with pEtN Is Present in C. jejuni NCTC 11168

To examine whether pEtN-glycan modification was unique to C. jejuni JHH1, we subjected peptides from the NCTC 11168 O strain (34) to ZIC-HILIC enrichment and glycopeptide analysis. On this occasion we searched the data to only extract MS/MS spectra containing diagnostic pEtN-glycan fragment ions (pEtN-GalNAc, 327.09; pEtN-GalNAc minus ethanolamine (EtN), 284.05). We observed two glycopeptides with pEtN-glycan modification. These were \sim 123.01/H11001 (the underline highlights the site of glycan attachment) from HisJ (CjAc or Cj0734c; MASCOT score = 20), which was also observed in the C. jejuni JHH1 dataset, and a novel glycopeptide 529QDLNSTLPVVNTNHK454 derived from Cj1013c (Fig. 3A; MASCOT score = 36). Consistent with the behavior of glycopeptides from C. jejuni JHH1, pEtN-glycan-modified glycopeptides from NCTC 11168 eluted \sim 1–2 min before the canonical glycopeptide forms (Fig. 3, B and C).

EptC (Cj0256) Is the C. jejuni N-Glycan pEtN Transferase—EptC (Cj0256) is the sole pEtN transferase predicted in C. jejuni 81-176 (15), and we confirmed this by in silico analysis of the C. jejuni 81116, ICDC07001, BAA-1458, NCTC 11168, IA3902, M1, RM1221, and S3 genomes. Furthermore, no strains are predicted to contain eptC paralogs. Since EptC demonstrates substrate promiscuity (15) by modifying both lipid A and FlgG, we next tested whether EptC is responsible for pEtN modification of the N-glycan. A chromosomal disruption of eptC was created in the C. jejuni JHH1 background. We also deleted the pglB gene to compare the effects of loss of the canonical N-glycan with loss of the pEtN modification alone. Western blotting of protein lysates from C. jejuni JHH1 wild-type and ΔeptC with N-glycan-specific antibody showed no major differences in the N-glycoproteome (Fig. 4A), and anti-JlpA antibody revealed no differences in the pEtN-glycan-modified protein JlpA (Table 1) after eptC deletion (Fig. 4B). This is most likely due to the low mass of pEtN (123.01 Da), which cannot be differentiated from non-pEtN modified forms on SDS-PAGE gels. C. jejuni JHH1 ΔpglB showed a complete loss of glycosylated proteins.

We next compared the C. jejuni JHH1 wild-type and ΔeptC strains by examining XIC from ZIC-HILIC-enriched glycopeptide fractions for all glycopeptides identified in Table 1. In the wild-type, we were readily able to detect both the canonical and pEtN-glycan-modified forms of each glycopeptide, whereas in JHH1 ΔeptC, only the canonical modification could be observed (Fig. 4C). MS/MS confirmed the presence of the pEtN-glycan on glycopeptides from wild-type (Fig. 4, D and E) that corresponded to peaks not observed in the ΔeptC mutant.

Generation of a pEtN-glycan in E. coli—To further demonstrate the involvement of EptC in pEtN transfer to the N-glycan, we employed the E. coli CLM24 expression system containing pACYC (encoding the pgl gene cluster) and pWA2 (containing the glycosylation substrate protein, AcrA) (44, 50). Introduction of EptC with a C-terminal hemagglutinin (HA) tag in the arabinose-inducible system pMLBAD (pNS3; supplemental Table S1) enabled the selective induction of EptC in E. coli (Fig. 5A). Expression of EptC did not lead to gross changes in AcrA levels (Fig. 5B). The glycoforms of His-tagged AcrA were isolated from induced pMLBAD (empty vector control; no eptC) and induced pNS3 (expressing eptC) and subjected to tryptic digestion followed by ZIC-HILIC enrichment. In the presence of EptC, the AcrA peptide 113-133 ATFNASKDFNKR124 was clearly glycosylated both with and without pEtN modification on the canonical C. jejuni glycan (Fig. 5C-D and supplemental Fig. S4)
FIGURE 1. Identification of a pEtN-modified N-linked glycan attached to antigenic protein HisJ (Cj0734c). A, shown is peptide fragmentation (lowercase y and b) by HCD MS/MS supporting the assignment of the peptide sequence $^{27}$ESNASVELK$^{35}$. B, fragmentation of pETN leads to the loss of ethanolamine (43.04 Da) followed by phosphate (79.97 Da). C, shown is pETN and N-glycan fragmentation (uppercase Y and B) supporting the terminal location of pETN attached to GalNAC. Oxonium ions of N-glycan sugars and pETN are also annotated.
as well as on the E. coli-specific glycan variant (Hex6HexNAc, Refs. 31 and 51); Fig. 5, C and D).

Influence of pEtN Modification on the N-Linked Glycoproteome—The observation of pEtN on multiple N-linked glycoproteins, several of which are known targets of the human humoral immune response (47, 49, 52), suggested a possible association with protein immunogenicity. This would also be consistent with studies showing that pEtN is an immunodominant modification of Neisseria meningitidis LOS (53). Coupled with the association of the C. jejuni N-linked glycan itself in reactivity to patient sera (54), we attempted to determine whether pEtN modification influenced immunoreactivity to C. jejuni proteins. Examination of reactivity with eight patient sera from the long term humoral response study of Cawthraw et al. (48) revealed that deletion of pgIB resulted in loss of reactivity; however, no discernible changes in reactivity were observed against proteins from C. jejuni JHH1/etC (Fig. 6A and supplemental Fig. S5). The substrate promiscuity of EptC also lead us to examine the known phenotypes of etC deletion in comparison with a C. jejuni JHH1 ΔetC (Fig. 6A and supplemental Fig. S5). The substrate promiscuity of EptC also lead us to examine the known phenotypes of etC deletion in comparison with a C. jejuni JHH1 ΔetC (Fig. 6A and supplemental Fig. S5). The substrate promiscuity of EptC also lead us to examine the known phenotypes of etC deletion in comparison with a C. jejuni JHH1 ΔetC (Fig. 6A and supplemental Fig. S5). The substrate promiscuity of EptC also lead us to examine the known phenotypes of etC deletion in comparison with a C. jejuni JHH1 ΔetC (Fig. 6A and supplemental Fig. S5). The substrate promiscuity of EptC also lead us to examine the known phenotypes of etC deletion in comparison with a C. jejuni JHH1 ΔetC (Fig. 6A and supplemental Fig. S5). The substrate promiscuity of EptC also lead us to examine the known phenotypes of etC deletion in comparison with a C. jejuni JHH1 ΔetC (Fig. 6A and supplemental Fig. S5).
pEtN Modification of the C. jejuni N-linked Glyceran

### TABLE 1

pEtN-glycan modified glycopeptides identified in *C. jejuni* JHH1 and NCTC11168 O. 9 modified peptides from 8 proteins were identified.

| Cj # | Protein Name                          | Precursor mass (charge) | Precursor MH | Peptide mass (MH+) | Peptide sequence | MASCOT score |
|------|---------------------------------------|-------------------------|--------------|-------------------|------------------|--------------|
| Cj0131 | Putative peptidase M23 family protein | 979.74768 (+3)         | 2937.22715   | 1408.6261         | 73DDNTSAMVIADEK   | 34           |
| Cj0131 | Putative peptidase M23 family protein | 962.76045 (+3)         | 2886.26548   | 1357.6780         | 6ILHKDNTSAM    | 48           |
| Cj0168c | Putative periplasmic protein          | 938.38813 (+3)         | 2813.14853   | 1284.3815         | 4ANTPSDVNQHTF    | 34           |
| Cj0289c | Major antigenic peptide Peb3          | 746.64479 (+3)         | 2237.91849   | 709.3515          | 2DFNVSRE        | 20           |
| Cj0399 | Colicin V production protein homolog  | 1006.11781 (+3)        | 3016.33756   | 1487.7700         | 1LQDIVSDLNNTQK  | 45           |
| Cj0734c | Histidine-binding protein Hsl/CjaC    | 835.69240 (+3)         | 2505.06131   | 976.4946          | 2ESNASVELK       | 31           |
| Cj0982c | Putative amino acid transporter periplasmic soluble-binding protein | 917.06178 (+3) | 2749.16947 | 1220.6005 | 3DSNITSVDELK | 47 |
| Cj0983c | Surface-exposed lipoprotein JlpA      | 840.03658 (+3)         | 2518.09385   | 989.5262          | 4GEANASISIK     | 21           |
| Cj1013c | Putative cytochrome c biogenesis protein | 962.76045 (+3) | 3279.48667  | 1750.9082 | 15QDLNSTLPVNTNHAK | 36 |

* MASCOT score given for *C. jejuni* JHH1 identification. MASCOT score = 20 for identical peptide in *C. jejuni* NCTC 11168 O.

### DISCUSSION

*N*-linked glycosylation has been associated with virulence in *C. jejuni* (11, 12), and protein targets have now been identified across several strains (Refs. 22, 26–27, 29, and 32 and this study). Unlike the complex branch, trimming, and additional sugar subunits associated with glycan diversity in eukaryotic *N*-glycans, all evidence until now has suggested that the *C. jejuni* *N*-glycan is rigidly conserved as a heptasaccharide consisting of an Asn-linked bacillosamine, five HexNAc (GalNAc), and a Hex branch (glucose) from the fourth position (3rd GalNAc). Our glycopeptide ZIC-HILIC enrichment approach (27) combined with the use of multiple, parallel proteolytic digests provided enhanced glycoproteome coverage that enabled the identification of 34 novel glycosylation sites from 15 previously uncharacterized *C. jejuni* glycoproteins. The size of the *C. jejuni* glycoproteome is considered to be as low as 150 protein targets (55) and no more than 260 proteins (predicted periplasmic, secreted, and membrane-associated proteins containing an *N*-linked sequon (27)). Our work combined with that of others in different *C. jejuni* strains has brought the number of verified *C. jejuni* glycoproteins to 70, making this the most comprehensive *N*-glycoproteome characterized for a free-living organism.

Our ability to mine deeply into the *C. jejuni* glycoproteome allowed us to search for evidence of non-canonical *N*-linked glycans. Manual interrogation of MS/MS spectra concentrated on high quality scans, with clear glycan fragment ions but no positive assignment by MASCOT. These spectra contained no evidence for the monoacetylated Bac linked to Asn previously observed in the laboratory passaged NCTC 11168 strain (32). Furthermore, we found no examples of truncated or elongated *N*-glycans modified by the addition or subtraction of GalNAc residues. Multiple glycopeptides containing a modified *N*-linked glycan consistent with the addition of a terminal pEtN were identified. The pEtN-glycan was not restricted to *C. jejuni*.
JHH1 as pEtN-modified glycopeptides were also identified in C. jejuni NCTC 11168 O. Extensive internal fragmentation of the pEtN-glycan and a predominant charge state that is non-ideal for HCD fragmentation (z ≈ +3 (27)) meant that pEtN-glycan-modified peptide sequences were difficult to identify even within this optimized workflow. Several MS/MS spectra containing clear pEtN-glycan fragment ions did not generate discernible peptide-related fragment ions, suggesting that additional sites of modification are likely to exist.

C. jejuni contains only a single protein, EptC (Cj0256), that is predicted to possess pEtN transferase activity (15), in contrast to other bacteria (e.g. E. coli) that contain multiple pEtN transferases (43). Neisseria sp. also encode multiple pEtN transferases that target specific substrates, including LptA, which modifies lipid A (56), Lpt3 and Lpt6, which target heptose within the lipopolysaccharide inner core (57), and phosphoform transferase A that can modify protein substrates (58, 59). In C. jejuni, EptC modifies both lipid A and FlgG and thus demonstrates broader substrate specificity than pEtN transferases from other organisms. It is currently unknown if protein targets in addition to FlgG are modified directly by EptC, although considering the broad range of substrates, this appears probable. Disruption of eptC resulted in the loss of the pEtN-glycan modification from all glycopeptides identified in C. jejuni JHH1 wild type and ΔpglB and ΔeptC deletion mutants. Western blotting with anti-N-linked glycan antibodies (A) and anti-JlpA (0G, 1G, and 2G refer to the number of occupied N-glycosites (47) (B) is shown. C, shown is XIC of pEtN-glycan and canonical N-glycan attached to peptide 27ESNASVELK35. CID MS/MS spectra of pEtN-glycan-modified (D) and canonical N-glycan-modified (E) forms of peptide 27ESNASVELK35, taken from wild-type and ΔeptC C. jejuni JHH1, respectively.

FIGURE 4. Comparison of glycoproteomes from C. jejuni JHH1 wild type and ΔpglB and ΔeptC deletion mutants. Western blotting with anti-N-linked glycan antibodies (A) and anti-JlpA (0G, 1G, and 2G refer to the number of occupied N-glycosites (47) (B) is shown. C, shown is XIC of pEtN-glycan and canonical N-glycan attached to peptide 27ESNASVELK35. CID MS/MS spectra of pEtN-glycan-modified (D) and canonical N-glycan-modified (E) forms of peptide 27ESNASVELK35, taken from wild-type and ΔeptC C. jejuni JHH1, respectively.
pEtN Modification of the C. jejuni N-linked Glycan

A) 

B) 

m/z = 968.39-968.41
RT: 28.38
MA: 17656919

(i)

m/z = 927.39-927.41
RT: 28.64
MA: 118179827

(ii)

m/z = 976.74-976.76
RT: 29.40
MA: 480183

(iii)

m/z = 935.74-935.76
RT: 29.79
MA: 253141875

(iv)

C) 

(i)

(ii)

(iii)

(iv)
tified in this study, confirming that EptC also acts as an N-glycan pEtN transferase.

Evidence from previous studies suggests that the EptC-mediated addition of pEtN to the N-glycan occurs after the PglK flippase translocation of the complete canonical N-glycan into the periplasm. EptC has already been shown to modify lipid A and FlgG, both of which are periplasmic substrates (15), and EptC itself is N-glycosylated at Asn-215 (27), confirming that at least this region of the protein is exposed to the periplasm. We have thus far been unable to identify a pEtN-glycan FOS (data not shown), suggesting modification occurs after attachment of the canonical glycan to protein substrates; however, we cannot state unequivocally that such a FOS does not exist. Finally, the exact position of pEtN on the terminal GalNAC remains unresolved, as it is not yet possible to produce sufficient amounts of modified N-glycan for structural analyses.

Expression of EptC in E. coli containing the pg1 gene cluster resulted in the addition of the pEtN-glycan to the target protein, AcrA. Intriguingly, we also found evidence of low level, yet above background, addition of pEtN to the N-linked glycan (20% of that seen in E. coli expressing the pg1 gene cluster and eptC, based on comparison of area under the curve for XIC from E. coli expressing eptC or empty vector control) in the absence of eptC (supplemental Fig. S6). Non-EptC pEtN modification of the C. jejuni N-linked glycan in E. coli at even these low levels demonstrates that at least one of the five predicted E. coli pEtN transferases also possesses this function. When expressed in E. coli, we were also able to identify glycopeptides from AcrA that were modified with an extended C. jejuni N-glycan containing an additional hexose (supplemental Fig. S6) as well as an N-glycan containing an unusual Asn-linked carbohydrate of mass 244.12 Da (supplemental Fig. S6). These

FIGURE 5. Expression of the C. jejuni N-linked glycosylation system, EptC-HA, and AcrA-His in E. coli. A, upper, induction (I) of EptC with arabinose from pNS3 compared with non-induced controls (NI) and empty vector control pMLBAD is shown. Lower, comparison of AcrA glycosylation (0G, 1G, and 2G refer to the number of occupied N-glycosites). B, XIC of pEtN-glycan-modified (i) and iii), E. coli heptasaccharide-modified (i and ii) and canonical C. jejuni N-glycan-modified (iii and iv) glycopeptides from AcrA117ATFENASKDFNR124 are shown. C, confirmation of glycan structures by CID MS/MS (i–iv).

FIGURE 6. The effect of Δpg1B and ΔeptC on C. jejuni phenotypes. A, shown is a Western blot using patient serum. B, shown are motility assays of C. jejuni JHH1 wild type, ΔeptC, and Δpg1B. C, polymyxin B sensitivity of C. jejuni JHH1 wild-type, Δpg1B, and ΔeptC. D, shown is restoration of polymyxin B resistance in C. jejuni NCTC 11168 Δpg1B:pg1B.
findings confirm that further N-glycan variations exist within the *E. coli* expression system than previously reported (31, 51) and that endogenous processes within *E. coli* can generate unexpected glycoforms.

Detection of even very low levels of endogenous pEtN modification on N-linked glycans in *E. coli* CLM24, the favored strain used for glycoprotein expression studies (60), may be a concern with respect to the homogeneity of glyco-conjugated vaccines expressed in this heterologous system. The addition of pEtN to *C. jejuni* substrates by *E. coli* pEtN transferases does have precedent, as *E. coli* EptA is capable of modifying *C. jejuni* lipid A (15). It is unclear whether the addition of pEtN will have an influence on vaccine reactivity or efficiency; however, pEtN has been noted as an immunodominant epitope (53), and therefore, care must be taken to ensure glycoprotein vaccines derived from this system either do not display the pEtN moiety or that the pEtN-glycan is harmless to recipients, and immunological reactivity is minimal compared with the intended epitope. Alternatively, these concerns may be overcome by the generation of an *E. coli* CLM24 derivative, in which the genes encoding endogenous pEtN transferases have been removed to limit background addition of pEtN to glyco-conjugates. Determination of the relative quantities of peptides modified with the canonical and pEtN-modified N-glycans is not feasible with currently available technology. It is, therefore, impossible to state unequivocally the ratio in which the two forms co-exist in *C. jejuni*; however, the enrichment, separation, and MS resolution and sensitivity required to detect the pEtN-glycan suggest it is present at relatively low levels, although factors beyond abundance and including the higher charge state, lack of peptide fragment ions, and change in LC retention time may account for reduced detection of pEtN-glycan-modified peptides.

Direct attachment of pEtN to FlgG by EptC (15) and to the pilin (40) of *Neisseria* sp. by phosho-form transferase A (39) have previously been observed, and pEtN modification of LOS has also been documented in a number of *C. jejuni* strains (37, 38). In *N. gonorrhoeae* there is recognition of the interplay between glycosylation and pEtN modification (39, 40), with overlapping site occupation observed between the O-linked glycosylation system and phospho-form transferase A-mediated phospho-form modifications (59). Extensive heterogeneity of pEtN and glycan modifications have been recognized in at least three proteins: Pile (39, 40) NGO1043, and NGO1237 (59), with two of these, Pile and NGO1043, recognized as major immunogens (40, 59). The link between pEtN and immunogenicity in *Neisseria* proteins and our identification of pEtN-glycan modification on previously identified *C. jejuni* immunogens, such as JlpA (47), Peb3 (61), and HisJ (62), provided further circumstantial evidence regarding a possible role in immunogenicity. We were, however, unable to detect a global effect of *eptC* deletion on reactivity of proteins with patient sera. In contrast, deletion of *pglB*, which results in loss of N-glycan attachment to proteins, significantly reduced protein reactivity. Disruption of *eptC* in the JHH1 background resulted in phenotypes that were reported previously (15); however, similar effects were also observed for *pglB* deletion mutants. Others have reported that deletion of genes within the *pgl* cluster, such as *pglE*, can result in motility defects (63); however, alterations in other genetic backgrounds have not displayed this phenotype (11). Increased sensitivity of Δ*pglB* mutants to polymyxin B was unexpected and suggests modification with the *N*-linked glycan enhances resistance; however, whether this is mediated globally via modification of a specific protein or at the level of the fOS is currently unknown.

Since EptC is multifunctional and targets several different substrates, it is difficult to assess the functional implications of the pEtN-glycan. Although no clear differences in reactivity to patient sera were observed, this does not preclude a role for the pEtN-glycan in innate immunity or in modifying host cell interactions. For example, previous studies have shown that *C. jejuni* N-linked glycoproteins and LOS displaying terminal GalNAc can bind the human macrophage galactose-type lectin (MGL) (64). Although speculative, the addition of pEtN to the *N*-glycan may block interactions between the previously terminal GalNAc of the canonical glycan and MGL, thus disrupting adherence between *C. jejuni* and MGL-expressing dendritic and macrophage cells. To assess this possibility, large-scale production of the pEtN-glycan will be required. Additionally, deletion of *eptC* results in loss of motility and is, therefore, likely to have a significant impact on the ability of *C. jejuni* to colonize both human and chicken hosts. Furthermore, a motility-reduced transposon mutagenesis library identified *eptC* (referred to as cf0256) leading to a 35% reduction in motility that was also associated with reduced invasion of INT-407 cells (65). Again, however, the multifunctional nature of EptC makes interpretation of such results problematic, as the individual contributions of pEtN modification of lipid A, FlgG, and the N-glycan cannot be determined. The motility defect observed in strain JHH1 Δ*eptC* is consistent with strain 81-176 (15); however, the magnitude of this reduction (25%) is less than observed for 81-176 (45%) and more consistent with published reports for strain 480 (35%) (65). The strain variations observed in these studies may be due to a multitude of currently unknown factors that influence motility in *C. jejuni*, including expression in clinical environments and unique genetic composition.

In conclusion, the depth of our *C. jejuni* glycoproteome data set enabled us to identify a novel pEtN modification of the canonical N-linked glycan, which is attached to nine glycosylation sites in eight glycoproteins derived from two different *C. jejuni* strains. The addition of pEtN to the N-linked glycan is mediated by EptC and was transferable into *E. coli*. Although the biological role of the N-glycan and pEtN-glycan are currently unknown, the loss of the pEtN modification does not appear to have the same global effect on patient sera reactivity as seen for the N-linked glycan alone. The addition of pEtN, however, is likely to influence any glycan-mediated charge-based interactions and thus alter glycan-associated structure/function relationships. Further studies are required to fully understand N-glycan diversity and to discern the role of these glycans in *C. jejuni* virulence.

REFERENCES
1. Butzler, J. P. (2004) *Campylobacter*, from obscurity to celebrity. *Clin. Microbiol. Infect.* 10, 868 – 876
2. Snelling, W. J., Matsuda, M., Moore, J. E., and Dooley, J. S. (2005) *Campylobacter jejuni*. *Lett. Appl. Microbiol.* 41, 297 – 302
pEtN Modification of the C. jejuni N-linked Glycan

Hanna, T., Kauppi, M., Tuomala, M., Laaksonen, I., Klemets, P., and Kuusi, M. (2004) Reactive arthritis following an outbreak of Campylobacter jejuni infection. J. Rheumatol. 31, 528–530

Hannifry, O. M., Shashkov, A. S., Moran, A. P., Senchenkova, S. N., and Savage, A. V. (2001) Chemical structure of the core oligosaccharide of aerotolerant Campylobacter jejuni O:2 lipopolysaccharide. Carbohydr. Res. 330, 223–229

Hannifry, O. M., Shashkov, A. S., Moran, A. P., Prendergast, M. M., Senchenkova, S. N., Knirel, Y. A., and Savage, A. V. (1999) Chemical structure of a polysaccharide from Campylobacter jejuni 176.83 (serotype O:41) containing only furanose sugars. Carbohydr. Res. 319, 124–132

Hannifry, O. M., Shashkov, A. S., Moran, A. P., and Szymanski, C. M. (2008) in Campylobacter jejuni O:2 lipopolysaccharide. Proc. Natl. Acad. Sci. U.S.A. 105, 15019–15024

Hannifry, O. M., Shashkov, A. S., Moran, A. P., Prendergast, M. M., Senchenkova, S. N., and Savage, A. V. (2001) Chemical structure of the core oligosaccharide of aerotolerant Campylobacter jejuni O:2 lipopolysaccharide. Carbohydr. Res. 330, 223–229

Hannifry, O. M., Shashkov, A. S., Moran, A. P., Prendergast, M. M., Senchenkova, S. N., Knirel, Y. A., and Savage, A. V. (1999) Chemical structure of a polysaccharide from Campylobacter jejuni 176.83 (serotype O:41) containing only furanose sugars. Carbohydr. Res. 319, 124–132

Hannifry, O. M., Shashkov, A. S., Moran, A. P., and Szymanski, C. M. (2008) in Campylobacter jejuni O:2 lipopolysaccharide. Proc. Natl. Acad. Sci. U.S.A. 105, 15019–15024
pEtN Modification of the C. jejuni N-linked Glycan

S. Vinogradov, E., and Brisson, J. R. (2003) Detection of conserved N- linked glycans and phase-variable lipooligosaccharides and capsules from Campylobacter cells by mass spectrometry and high resolution magic angle spinning NMR spectroscopy. J. Biol. Chem. 278, 24509–24520

38. St Michael, F., Szymanski, C. M., Li, J., Chan, K. H., Khieu, N. H., Laroque, S., Wakarchuk, W. W., Brisson, J. R., and Monteiro, M. A. (2002) The structures of the lipooligosaccharide and capsule polysaccharide of Campylobacter jejuni genome sequenced strain NCTC 11668. Eur. J. Biochem. 269, 5119–5136

39. Aas, F. E., Egge-Jacobsen, W., Winther-Larsen, H. C., Løvold, C., Hitchen, P. G., Dell, A., and Koomey, M. (2006) Neisseria gonorrhoeae type IV pili undergo multisite, hierarchical modifications with phosphoethanolamine and phosphocholine requiring an enzyme structurally related to lipooligosaccharide phosphoethanolamine transferases. J. Biol. Chem. 281, 27712–27723

40. Hegge, F. T., Hitchen, P. G., Aas, F. E., Kristiansen, H., Løvold, C., Egge-Jacobsen, W., Panico, M., Leong, W. Y., Bull, V., Virji, M., Morris, H. R., Dell, A., and Koomey, M. (2004) Unique modifications with phosphocho-line and phosphoethanolamine define alternate antigenic forms of Neisseria gonorrhoeae type IV pili. Proc. Natl. Acad. Sci. U.S.A. 101, 10798–10803

41. Stimson, E., Virji, M., Barker, S., Panico, M., Blench, I., Saunders, J., Payne, G., Moxon, E. R., Dell, A., and Morris, H. R. (1996) Discovery of a novel protein modification. α-Glyceroephosphatase is a substitute of meningococcal pilin. Biochem. J. 316, 29–33

42. Chamot-Rooke, J., Mikaty, G., Malosse, C., Soyer, M., Dumont, A., Gault, J., Linhares, A. F., Martin, P., Trellet, M., Clary, G., Chafey, P., Camoin, L., Nilges, M., Nassif, X., and Doménil, G. (2011) Posttranslational modification of pili upon cell contact triggers N. meningitidis dissemination. Science 331, 778–782

43. Cullen, T. W., Madsen, J. A., Ivanov, P. L., Brodbelt, J. S., and Trent, M. S. (2012) Characterization of a unique modification of the flagellar rod protein FlgG by the Campylobacter jejuni lipid A phosphoethanolamine transferase, linking bacterial locomotion and antimicrobial peptide resistance. J. Biol. Chem. 287, 3326–3330

44. Feldman, M. F., Wacker, M., Hernandez, M., Hitchens, P. G., Marolda, C. L., Kowarik, M., Morris, H. R., Dell, A., Valvano, M. A., and Aebl, M. (2005) Engineering N-linked protein glycosylation with diverse O antigen lipooligosaccharide structures in Escherichia coli. Proc. Natl. Acad. Sci. U.S.A. 102, 3016–3021

45. Chen, R., Jiang, X., Sun, D., Han, G., Wang, F., Ye, M., Wang, L., and Zou, H. (2009) Glycoproteomics analysis of human liver tissue by combination of multiple enzyme digestion and hydrazide chemistry. J. Proteome Res. 8, 651–661

46. Davis, L., Young, K., and DiRita, V. (2008) Genetic manipulation of Campylobacter jejuni. Curr. Protoc. Microbiol. Chapter 8, Unit 8.2 1.8A 2.17

47. Scott, N. E., Bogema, D. R., Connolly, A. M., Falconer, L., Djordjevic, S. P., and Cordwell, S. J. (2009) Mass spectrometric characterization of the surface-associated 42 kDa lipoprotein IlpA as a glycosylated antigen in strains of Campylobacter jejuni. J. Proteome Res. 8, 4654–4664

48. Cawthra, S. A., Feldman, R. A., Sayers, A. R., and Newell, D. G. (2002) Long term antibody responses following human infection with Campylo- bacter jejuni. Clin. Exp. Immunol. 130, 101–106

49. Cordwell, S. J., Len, A. C., Touma, R. G., Scott, N. E., Falconer, L., Jones, D., Connolly, A., Crossett, B., and Djordjevic, S. P. (2008) Identification of membrane-associated proteins from Campylobacter jejuni strains using complementary proteomics technologies. Proteomics 8, 122–139

50. Wacker, M., Linton, D., Hitchen, P. G., Nita-Lazar, M., Haslam, S. M., North, S. I., Panico, M., Morris, H. R., Dell, A., Wren, B. W., and Aebl, M. (2002) N-Linked glycosylation in Campylobacter jejuni and its functional transfer into E. coli. Science 298, 1790–1793

51. Linton, D., Dorrell, N., Hitchen, P. G., Amber, S., Karlyshev, A. V., Morris, H. R., Dell, A., Valvano, M. A., Aebl, M., and Wren, B. W. (2005) Functional analysis of the Campylobacter jejuni N-linked protein glycosylation pathway. Mol. Microbiol. 55, 1695–1703

52. Pawelec, D., Jakubowska-Mróz, J., and Jagusztyn-Krynicka, E. K. (1998) Campylobacter jejuni 72Dzx/92 cjaC gene coding 28-kDa immunopositive protein, a homologue of the solute-binding components of the ABC transport system. Lett. Appl. Microbiol. 26, 69–76

53. Szymanski, C. M., Yao, R., Ewing, C. P., Trust, T. J., and Guerry, P. (1999) Evidence for a system of general protein glycosylation in Campylobacter jejuni. Mol. Microbiol. 32, 1022–1030

54. Nothhaft, H., Amber, S., Aebl, M., and Szymanski, C. M. (2008) in Campylo- bacter (Nachamkin, I., Szymanski, C. M., and Blaser, M. J., eds) 3rd Ed., pp. 447–469, American Society for Microbiology, Washington, D.C.

55. Cox, A. D., Wright, J. C., Liu, J., Hood, D. W., Moxon, E. R., and Richards, J. C. (2003) Phosphorylation of the lipid A region of meningococcal lipopolysaccharide: identification of a family of transferases that add phosphoethanolamine to lipopolysaccharide. J. Bacteriol. 185, 3270–3277

56. Wenzel, C. Q., St Michael, F., Stupak, J., Li, J., Cox, A. D., and Richards, J. C. (2010) Functional characterization of Lpt3 and Lpt6, the inner-core lipoo- ligosaccharide phosphoethanolamine transferases from Neisseria meningitidis. J. Bacteriol. 192, 208–216

57. Naessan, C. L., Egge-Jacobsen, W., Heiniger, R. W., Wolfgang, M. C., Aas, F. E., Rohr, A., Winther-Larsen, H. C., and Koomey, M. (2008) Genetic and functional analyses of PptA, a phospho-form transferase targeting type IV pili in Neisseria gonorrhoeae. J. Bacteriol. 190, 387–400

58. Anonsen, J. H., Egge-Jacobsen, W., Aas, F. E., Berud, B., Koomey, M., and Vik, A. (2012) Novel protein substrates of the phospho-form modification system in Neisseria meningitidis and their connection to O-linked protein glycosylation. Infect. Immun. 80, 22–30

59. Ibbsen, J., Kowarik, M., Diplitowso, S., Tanner, C., Wacker, M., Thöny- Meyer, L. (2010) Production of glycoprotein vaccines in Escherichia coli. Microb. Cell Fact. 9, 61–74

60. Pei, Z. H., Ellison, R. T., 3rd, and Blaser, M. J. (1991) Identification, purifica-tion, and characterization of major antigenic proteins of Campylobac- ter jejuni. J. Biol. Chem. 266, 16363–16369

61. Wyszyńska, A., Tomczyk, K., and Jagusztyn-Krynicka, E. K. (2007) Com-parison of the localization and post-translational modification of Campylo- bacter coli CjaC and its homolog from Campylobacter jejuni, Cj0734c/ HisA. Acta Biochim. Pol. 54, 143–150

62. Vijayakumar, S., Merkx-Jacques, A., Ratnayake, D. B., Gryski, I., Obhi, R. K., Houle, S., Dozois, C. M., and Creuzenet, C. (2006) Cj1121c, a novel UDP-4-keto-6-deoxy-GlcNAc C-4aminotransferase essential for protein glycosylation and virulence in Campylobacter jejuni. J. Biol. Chem. 281, 27733–27743

63. van Sorge, N. M., Bleumink, N. M., van Vliet, S. J., Saeland, E., van der Pol, W. L., van Kooyk, Y., and van Putten, J. P. (2009) N-glycosylated proteins and distinct lipooligosaccharide glycoforms of Campylobacter jejuni tar-get the human C-type lectin receptor MGL. Cell Microbiol. 11, 1768–1781

64. Golden, N. J., and Acheson, D. W. (2002) Identification of motility and autoagglutination Campylobacter jejuni mutants by random transposon mutagenesis. Infect. Immun. 70, 1761–1771