**Biochemical Characterization of the O-Linked Glycosylation Pathway in Neisseria gonorrhoeae Responsible for Biosynthesis of Protein Glycans**

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Biochemical characterization of the O-linked glycosylation pathway in *Neisseria gonorrhoeae* responsible for biosynthesis of protein glycans containing N,N′-diacetylbacillosamine

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**Abstract**

The O-linked protein glycosylation pathway in *Neisseria gonorrhoeae* is responsible for the synthesis of a complex oligosaccharide on undecaprenyl diphosphate and subsequent *en bloc* transfer of the glycan to serine residues of select periplasmic proteins. Protein glycosylation (pgl) genes have been annotated on the basis of bioinformatics and top-down mass spectrometry analysis of protein modifications in *pgl*-null strains (1, 2), but relatively little biochemical analysis has been performed to date. In this report, we present the expression, purification and functional characterization of seven Pgl enzymes. Specifically, the enzymes studied are responsible for synthesis of an uncommon uridine diphosphate (UDP)-sugar (PglD, PglI, and PglB-acetyltransferase domain), glycan assembly (PglB-phospho-glycosyltransferase domain, PglA, PglE, and PglH) and final oligosaccharide transfer (PglO). UDP-2,4-diacetamido-2,4,6-trideoxy-α-D-hexose (DATDH), which is the first sugar in glycan biosynthesis, was produced enzymatically and the stereochemistry was assigned as uridine diphosphate N′-diacetylbacillosamine (UDP-diNAcBac) by NMR characterization. In addition, the substrate specificities of the phospho-glycosyltransferase, glycosyltransferases and oligosaccharyltransferase (OTase) were analyzed *in vitro* and in most cases, these enzymes showed strong preferences for the native substrates relative to closely related glycans. In particular, PglO, the O-linked OTase, and PglB(‡), the N-linked OTase from *Campylobacter jejuni*, preferred the native *N. gonorrhoeae* or *C. jejuni* substrates, respectively. This study represents the first comprehensive biochemical characterization of this important O-linked glycosylation pathway and provides the basis for further investigations of these enzymes as antibacterial targets.

In *Neisseria gonorrhoeae*, individual pilin subunits rapidly assemble and disassemble to form the flagellar-like Type IV pili, which mediate essential interactions with host cells and affect many aspects of pathogenicity including surface motility, bacteria-host...
communication, cell signaling, bacterial dissemination and biofilm formation (3–6). Recently, the gonococcal pilin glycosylation system was shown to be a general O-linked system in which many structurally distinct periplasmic proteins undergo glycosylation (2). Glycan-modifications on pili, flagella and other extracellular proteins have been implicated in bacterial pathogenicity, which has led to increased interest in bacterial glycosylation pathways as potential antibacterial targets (4, 7–11). The focus of this study is the protein glycosylation (pgl) locus identified in *N. gonorrhoeae*, which is responsible for glycan addition to distinct serine residues (1, 12).

The protein glycan modifications present in *N. gonorrhoeae* pgl-null strains have been analyzed by top-down mass spectrometry (MS) (1, 2) and the following model of the protein glycosylation pathway has been developed (Figure 1). The core pgl locus contains four genes, three of which (*pglD, pglC*, and *pglB*)2 are required for the synthesis of an undecaprenyl diphosphate 2,4-diacetamido-2,4,6-trideoxy-α-D-hexose (Und-PP-DATDH) (13). The term ‘DATDH’ indicates that the stereochemistry of this sugar has not been previously determined. PglD and PglC perform NADH-dependent dehydratase and aminotransferase reactions, respectively, to convert UDP-HexNAc to UDP-2-acetamido-4-amino-2,4,6-trideoxy-α-D-glucose or (UDP-4-amino), but it is not known if the initial substrate is UDP-GlcNAc or UDP-GalNAc. PglB is a bifunctional enzyme, which catalyzes the amino acetylation of UDP-4-amino to form UDP-DATDH and the transfer of phospho-DATDH to undecaprenyl phosphate (Und-P). The fourth gene, *pglF*, shares homology with ABC transporter-type flippases and is putatively involved in the translocation of the undecaprenyl diphosphate-linked glycan across the periplasmic membrane. Although the function of this gene has not been demonstrated, the *pglF*-null strain does exhibit diminished glycosylation (1, 12).

Interestingly, the other genes involved in pilin glycosylation are not linked to the core pgl locus. The products of the *pglA* and *pglE* genes further elaborate the polyprenyl-linked DATDH with the transfer of two sequential galactose units (Figure 1) (1). The *pglA* and *pglE* genes undergo phase variation in which the genes are alternately turned on and off. Phase-variant *pglA* alleles have been proposed to be associated with more virulent strains of *N. gonorrhoeae*, although these studies have been disputed (1, 8, 9). In addition to PglA and PglE, an alternate glycosyltransferase PglH adds a Glc unit instead of Gal to Und-PP-DATDH (Figure 2) (14). Finally, a gene has been identified, *pglO*, which shares homology with the O-antigen ligase (WaaL) family and was required for formation of the protein-glycan linkage (Figure 1) (2).

Considerable recent studies have focused on the highly homologous pilin glycosylation pathway in the related species *Neisseria meningitidis*. The proposed model of the *N. meningitidis* pathway is similar to the *N. gonorrhoeae* pathway (Figure 2) and was also developed by bioinformatic analysis and experiments with pgl-null strains (15–20). The oligosaccharyltransferase from *N. meningitidis*, PglL(Nm), has been purified to homogeneity and shown to glycosylate pilin using a farnesyl substrate analog (21). In addition, upon

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1Abbreviations: 2-AB, 2-aminobenzamide; AcCoA, acetyl coenzyme A; Cj, *Campylobacter jejuni*; CEF, cell envelope fraction; DDM, n-dodecyl-β-D-maltopyranoside; Gal, galactose; GalNAc, N-acetylgalactosamine; Glc, glucose; GlcNAc, N-acetylglucosamine; NP-HPLC, normal phase-high performance liquid chromatography; MALDI MS, matrix-assisted laser desorption ionization mass spectrometry; Man, mannose; MBP, maltose binding protein; Ng, *Neisseria gonorrhoeae*; Ni-NTA, Ni-nitrilotriacetic acid; Nm, *Neisseria meningitidis*; Pgl, protein glycosylation; PglB-ATD, acetyltransferase domain of PglB; PglB-PGTD, phospho-glycosyltransferase domain of PglB; TMHMM, Tied Mixture Hidden Markov Model (a transmembrane prediction model); UDP, uridine diphosphate; UDP-4-amino, UDP-2-acetamido-4-amino-2,4,6-trideoxy-α-D-glucose or UDP-2-acetamido-4-amino-α-D-quinovose; UDP-DATDH, 2,4-diacetamido-2,4,6-trideoxy-α-D-hexose; UDP-diNAcBac, UDP-N,N’-diacetylbacillosamine or UDP-2,4-diacetamido-2,4,6-trideoxy-α-D-hexose; UDP-4-keto, UDP-2-acetamido-4-keto-2,4,6-trideoxy-α-D-glucose; UDP-2-acetamido-4-keto-α-D-quinovose; Und-P, undecaprenyl phosphate; Und-PP, undecaprenyl diphosphate.

2The accession numbers for the genes are available in the Supplementary Information.

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heterologous expression in *E. coli*, PgL(Nm) was shown to transfer a variety of complex glycan substrates to pilin (21, 22).

The Pgl pathways from *N. gonorrhoeae* and *N. meningitidis* represent the first example of O-linked protein glycans derived from polyprenyl-linked intermediates; all other identified O-linked pathways glycosylate protein substrates by sequential transfer of individual saccharide units from nucleotide or polyprenyl-phosphate activated glycan donors. Another intriguing facet of the *N. gonorrhoeae* O-linked glycosylation pathway is that the first three enzymes (PglD, C, and B) share homology with the first four enzymes in the N-linked protein glycosylation (also designated Pgl) pathway in *Campylobacter jejuni* (23) with the exception that the *C. jejuni* locus encodes separate enzymes for the sequential acetyltransferase and phospho-glycosyltransferase reactions. Both the *N. gonorrhoeae* and *C. jejuni* pathways produce an initial Und-PP-DATDH intermediate, but this intermediate is elaborated in distinct ways (Figure 2). The *N. gonorrhoeae* pathway produces a serine-linked mono-, di- or trisaccharide (13) and the *C. jejuni* pathway generates an asparagine-linked heptasaccharide (24). The *C. jejuni* glycosylation pathway serves as an important model for the *N. gonorrhoeae* system and previous work has resulted in the complete biochemical characterization of the *C. jejuni* Pgl pathway enzymes except for the flippase (PglK) (25–28).

In this study, the biochemical functions of the proteins PglD, C, B, A, E, and O from *N. gonorrhoeae* are characterized for the first time through *in vitro* biochemical analysis. Importantly, the previously undefined stereochemical assignment of the UDP-DATDH produced by PglD, C, and B is unequivocally shown to be UDP-diNACBac, which is also the identity of the first sugar added in the *C. jejuni* N-linked glycosylation pathway. *In vitro* assays demonstrate that the phospho-glycosyltransferase (PglB) and two glycosyltransferases (PglA and PglE) build the glycan on an undecaprenyl-diphosphate linker prior to *en bloc* transfer to protein and that these enzymes display strict specificity for the UDP-saccharide donor. Finally, glycan substrate specificity analyses suggest that the O-linked OTase is highly selective for native *N. gonorrhoeae* glycan substrates *in vitro*.

**Experimental Procedures**

**Common Materials**

All radioactive materials and undecaprenol were obtained from American Radiolabeled Chemicals. UDP-4-amino and UDP-diNACBac were prepared as previously described (28) by enzymes from *C. jejuni* or as described herein by PglD, PglC and PglB. All other chemicals were obtained from Sigma-Aldrich unless stated otherwise. Radioactivity was determined using a LS6500 Beckman Scintillation Counter; organic samples were dried and resuspended in 200 μL Solvable™ (Perkin-Elmer) and 5 mL of scintillation fluid (Opti-Fluor, Perkin-Elmer). Aqueous samples were mixed with 5 mL of Ecolite(+)™ (MP Biomedicals) prior to scintillation counting.

**Preparation of genetic constructs**

The genes *pglD, pglC, pglB, pglA, pglO*, and *pglE* were PCR amplified from the *N. gonorrhoeae* strain MS11 (1, 8, 12), while *pglE* was amplified from *N. gonorrhoeae* strain FA 1090 and *pglH* was amplified from the *N. meningitidis* strain Z2491. The PCR products of *pglD, pglC, pglB, pglA, pglO*, and *pglE* were cloned into BamH I/Xho I in the pET-24at(+) vector (Novagen). The *pglE* and *PglH* genes were cloned into Nde I/Xho I in the pET-24at(+) vector (Novagen). The Xho I site was inserted prior to the stop codon to encode for a His<sub>6</sub> tag at the C-terminal end of each protein.
The acetyltransferase domain of PglB (PglB-ATD) was identified through sequence homology with the related *C. jejuni* protein, PglD. The gene encoding the domain was amplified from the full-length gene using the forward primer 5’-CGCGGATCCATGGCGGGGAATCGCAAACTCG-3’ and the reverse primer 5’-GCAACCCGGCAGCCCTTATTAGCTCGAGCGG-3’ to generate a gene encoding the acetyltransferase domain. The gene was inserted into BamH I/Xho I in a modified pET-30b(+) vector which contains an N-terminal His\(^8\) tag followed by a Tobacco Etch Virus (TEV) protease site. Also, pglH was amplified by PCR and inserted into BamH I/Xho I in the pMAL-c2X vector. This construct encoded for the addition of an N-terminal maltose binding protein (MBP).

**Expression of proteins**

In general, all proteins (PglD, PglC, PglB, PglB-ATD, PglA, PglE, PglO, PglH, PilE (pilin), PglB(Cj) (27) and *S. mutans* undecaprenol kinase (29)) were expressed heterologously in *E. coli* BL21 cells (Agilent). PglD, PglC and PglB-ATD were expressed in the BL21(DE3) pLysS strain; all other proteins were expressed in the BL21-Gold (DE3) strain. A typical expression protocol involved preparation of an overnight culture of cells (5 mL), which was used to inoculate 1 L of LB media with shaking at 37 °C. After the cells reached an optical density of ~0.8 absorbance units, the temperature was lowered to 16 °C and the cells were induced with 0.5 mM iso-β-D-thiogalactosylpyranoside (IPTG). After 16–18 hours of incubation, the cells were harvested and the pellets were stored at −80 °C.

**Protein purification**

In general, all steps of protein purification were carried out at 4 °C. Protein concentrations were determined with the appropriate extinction coefficients at a UV absorbance of 280 nm, with the exception of PglO and pilin, which were quantified with the MicroBCA Assay (Pierce) due to the presence of the highly absorbent detergent Triton X-100.

The cell pellets generated from the expression of the soluble proteins, PglC and PglB-ATD, were resuspended in 50 mL of ice-cold 50 mM HEPES (pH 7.4) and 100 mM NaCl (Buffer A), supplemented with 30 mM imidazole and lysed by sonication. In the case of PglC, 200 μM pyridoxal-5’- phosphate was also added to the buffer. The lysate was cleared by centrifugation (145,000 × g) for 45 min. Cleared lysate was mixed with 2 mL of Ni-nitritriacetic acid (Ni-NTA) resin (Qiagen), tumbled for 4 hours, and then packed into a K 9/15 column (GE Healthcare). Using gravity flow, the resin-bound protein was washed with 10 column volumes of Buffer A containing 30 mM imidazole. The resin was further washed with 20 column volumes of Buffer A supplemented with 40 mM imidazole and then 10 column volumes of Buffer A containing 60 mM imidazole. The protein was eluted in Buffer A supplemented with 250 mM imidazole and 1 mL fractions were collected. Fractions containing purified material were assessed by SDS-PAGE (12%) and Western blot analysis probing for the His\(_8\) tag. Pooled fractions of PglC and PglB-ATD were dialyzed against Buffer A, concentrated, supplemented with a final glycerol concentration of 15% and frozen at −80 °C (Figure S1 of the supplementary information, lanes 2 and 3).

Purification of the glycosyltransferase PglA was similar to PglC and PglB-ATD with a few exceptions. A buffer containing 50 mM Tris (pH 8.0) and 150 mM NaCl (Buffer B) was used instead of Buffer A and the cells were incubated with 1% Triton X-100 for 20 minutes immediately following lysis and prior to centrifugation. In addition, 5% glycerol was added to all buffers. Following elution, the most concentrated 1.5 mL fraction as determined by SDS-PAGE was desalted using a Hi-Trap desalting cartridge (GE Healthcare) with Buffer B and stored at −20°C in 30% glycerol (Figure S1, lane 5).
To purify the membrane-associated proteins (PglD, PglB, PglE, PglO, and pilin), a cell envelope fraction (CEF) was first prepared. The cells were thawed in 40 mL of buffer per L of cell culture and lysed by sonication. PBS supplemented with 200 µM NAD⁺ was used for PglD and Buffer B with 1 mg/mL lysozyme was used for PglB, PglE, PglO and pilin. Cellular debris was cleared by centrifugation at 9000 × g for 45 minutes. The resulting supernatant was transferred to a clean centrifuge tube and subjected to centrifugation at 145,000 × g for 65 min to pellet the CEF. For PglB and PglE, the CEFs were resuspended in half the volume of the unlysed cell pellet weight (i.e. 1.5 mL was used for 3 g cell pellet). The CEF was aliquoted and stored at −80 °C (Figure S1, lanes 4 and 6). The CEFs of PglB and PglE were used in all glycosyltransferase assays.

PglD, PglO and pilin were further purified from the CEF. The CEF was homogenized in 10 mL of buffer containing 1% Triton X-100 (PBS with 200 µM NAD⁺ for PglD and Buffer B for PglO and pilin) per liter of cell culture. Each CEF was incubated with detergent for several hours and then centrifuged again (145,000 × g) to remove insoluble material. The resultant supernatants were incubated with 0.5–2 mL of Ni-NTA resin for 1–2 hours; the resins were washed as previously described with the addition of 0.1% Triton X-100 to the wash and elution buffers. The proteins were eluted from the resin in 1 mL fractions. Pooled fractions of PglD were dialyzed against PBS containing 200 µM NAD⁺ and 0.1% Triton X-100, supplemented with a final glycerol concentration of 30%, and frozen at −80°C (Figure S1, lane 1). The most concentrated fractions of PglO and pilin were desalted as described above for PglA and stored at −80 °C (Figure S1, lanes 7 and 9). For the biosynthesis of UDP-diNAcBac, PglB was purified in a manner similar to PglD, except that NAD⁺ was not added to the purification buffers.

PglH was expressed as an MBP-fusion protein and purified as described elsewhere (14) (Figure S1, lane 8). In addition, PglB(Cj) and undecaprenol kinase from S. mutans were expressed and purified as cell envelope fractions as described previously (27, 29).

**Acetyltransferase (PglB-ATD) Activity Assay**

Determination of the kinetic constants for PglB-ATD was carried out using Ellman’s reagent, 5,5’-dithio-bis-(2-nitrobenzoic acid), in a continuous fashion. Ellman’s reagent was utilized to quantify substrate turnover as monitored by measuring conversion of acetyl coenzyme A (AcCoA) to CoASH using the released TNB chromophore (λ_max = 412 nm, ε_max = 14,150 M⁻¹ cm⁻¹). The *in vitro* assay contained 50 mM HEPES (pH 7.4), 1 mM EDTA, 1 mM DTNB, and 25 nM PglB-ATD in a quartz cuvette. The substrate concentrations of AcCoA and UDP-4-amino were varied separately to determine kinetic parameters using initial velocity measurements while keeping the other substrate at saturation. The reaction was initiated with the UDP-4-amino substrate and took place at room temperature over a 200 second time period. The absorbance change at 412 nm was measured. A blank reaction lacking UDP-4-amino was prepared as a background control. Steady-state rate parameters were calculated from equation 1 using the program GraFit 6.0.12 (Erithacus Software).

\[ n = \frac{V_{max}[S]}{K_m + [S]} \]  

(1)

**Aminotransferase (PglC) Activity Assay**

The aminotransferase reaction was assayed by coupling generation of UDP-4-amino from the PglC reaction to the acetyltransferase activity of PglD from *C. jejuni* (Cj) producing CoASH, which was detected by Ellman’s reagent in a similar fashion to the PglB-ATD assay. In a flat bottom 96-well plate (Nunc), 50 mM HEPES (pH 7.4), 1 mM PglD(Cj), 400
mM AcCoA, and 400 nM PglC were added. Since PglC activity was coupled to the turnover of the acetyltransferase PglD(C), addition of excess PglD(C) ensured that the initial velocity measurements were dependent only upon PglC activity. The concentrations of L-glutamate and UDP-4-keto were varied separately to determine kinetic parameters using initial velocity measurements while keeping the other substrate at saturation. Interference of Ellman’s reagent with PglC activity required the implementation of a discontinuous assay in which reactions were initiated with L-glutamate and quenched over a 30 minute time period with 20% n-propanol, 2 mM DTNB, and 1 mM EDTA. The absorbance at 415 nm was followed on an Ultramark EX microplate imaging system (BioRad). A blank reaction without L-glutamate was set up as a background control.

**Biosynthesis and stereochemical assignment of UDP-DATDH**

In order to biosynthesize UDP-DATDH, 0.3 mg of PglD (bound to Ni-NTA resin) was added to 15 mL buffer containing 50 mM HEPES (pH 7.4), 100 mM NaCl, 200 mM NAD$^+$, and 30 mg UDP-GlcNAc. The reaction was carried out at room temperature for 12 hours with gentle rocking. Once conversion to the UDP-4-keto sugar was complete, as verified by capillary electrophoresis as described previously (28), the reaction was filtered and the flow-through collected. The filtrate containing the UDP-4-keto sugar was supplemented with 15 mg PglC (bound to Ni-NTA resin), 20 mM L-glutamate, and 200 µM pyridoxal-5’-phosphate. This reaction was filtered after rocking for 18 hours at room temperature and reaching 80% conversion to the UDP-4-amino sugar. The filtrate was supplemented with 0.2 mg purified, full-length PglB (bound to Ni-NTA resin) and 1.2 mM AcCoA and allowed to react at room temperature with rocking for 12 hours. The slurry was filtered and the flow-through containing the UDP-DATDH sugar was collected. Purification and NMR characterization of the final UDP-DATDH product was completed as previously described (28).

**Preparation of radiolabeled Und-PP-linked substrates**

In general, radiolabeled Und-PP-substrates were prepared at two different specific activity levels. A higher specific activity was used for the OTase assay and analysis by normal phase-high performance liquid chromatography (NP-HPLC) and a lower specific activity was appropriate for the glycosyltransferase assays.

Und-PP-[^3]HdiNAcBac was enzymatically synthesized using *S. mutans* undecaprenol kinase (30) and PglB. An undecaprenol kinase from *N. gonorrhoeae* has not been characterized, and thus the undecaprenol kinase from *S. mutans* (29) was used a tool to affect the undecaprenol phosphorylation *in situ*. A typical reaction contained 3% DMSO, 1% Triton X-100, 50 mM MgCl$_2$, 30 mM Tris-Acetate (pH 8.0), 500 µM undecaprenol, 1 mM ATP, 500 µM UDP-4-amino, 500 µM[^3]HAcCoA (20 mCi/mmol), 15–20 µL of undecaprenol kinase CEF, 15–20 µL PglB CEF and water to a final volume of 100 µL. The reaction was modified to prepare Und-PP-[^3]HdiNAcBac with high specific activity by adjusting the undecaprenol and UDP-4-amino concentrations to 100 µM and the[^3]HAcCoA concentration to 4.5 µM (20 Ci/mmol). After incubation at room temperature for 2 hours, the reactions were quenched into 1 mL 2:1 CHCl$_3$:MeOH and extracted three times with 400 µL of an aqueous extract prepared by dissolving 1.83 g of potassium chloride in 235 mL water, 240 mL chloroform, and 15 mL methanol. The organic layer containing the Und-PP-[^3]HdiNAcBac product was dried down and purified using NP-HPLC as described below.

Und-PP-diNAcBac-[^3]HGal was prepared in a similar manner to Und-PP-diNAcBac. The reaction components are as described above for Und-PP-diNAcBac with the following exceptions; 500 µM UDP-diNAcBac was added instead of UDP-4-amino and AcCoA, and 2 µM PglA and 500 µM UDP-[^3]HGal (20 mCi/mmol) were added to affect the transfer of the
galactosyl unit, which is the second sugar in the glycan. To prepare Und-PP-diNAcBac-[\textsuperscript{3}H]Gal with higher specific activity, undecaprenol and UDP-diNAcBac concentrations were lowered to 100 µM and the UDP-[\textsuperscript{3}H]Gal concentration was adjusted to 4.5 µM (20 Ci/mmol). The reactions were quenched after two hours and extracted as described above.

The synthesis of Und-PP-diNAcBac-Gal-[\textsuperscript{3}H]Gal utilized unlabeled Und-PP-diNAcBac-Gal, which was prepared as described above with the exception that UDP-Gal was not radioactive. A typical biosynthesis reaction contained 3% DMSO, 0.05% Triton X-100, 50 mM MnCl\textsubscript{2}, 50 mM HEPES (pH 7.5), 20 µM Und-PP-diNAcBac-Gal, 20 µM UDP-[\textsuperscript{3}H]Gal (20 mCi/mmole), 20 µL PgiE CEF and water to a final volume of 100 µL. To prepare the substrate with higher specific activity, the UDP-Gal concentration was lowered to 4.5 µM UDP-[\textsuperscript{3}H]Gal (20 Ci/mmol). The reactions were quenched after two hours and extracted as described above.

The C. jejuni substrates (Und-PP-diNAcBac-[\textsuperscript{3}H]GalNAc and Und-PP-diNAcBac-GalNAc-[\textsuperscript{3}H]GalNAc) for the OTase reactions were prepared as previously described (25–27, 29, 31) with similar specific activities (20 Ci/mmole) to the N. gonorrhoeae OTase substrates.

**Normal phase HPLC purification of Und-PP-linked substrates**

The dried Und-PP-linked substrates were purified via NP-HPLC with a Varian Microsorb column using the previously described gradient (31). The substrates were resuspended in 100 µL of 4:1 CHCl\textsubscript{3}:MeOH for injection onto the column. Fractions of 1 mL were collected and 10 µL of each fraction was solubilized in 200 µL Solvable™ for detection of radioactivity. The fractions containing substrate were combined, aliquoted and stored at −20 °C.

To obtain the NP-HPLC analytical traces, Und-PP-linked glycan fractions were resolubilized in 4:1 CHCl\textsubscript{3}:MeOH and 100 µL of the appropriate sample was injected onto the column. The 1 mL elution fractions were dried completely and resuspended in 200 µL Solvable™ for scintillation counting.

**Preparation and analysis of 2-AB labeled oligosaccharides**

Unlabeled versions of Und-PP-diNAcBac-Gal and Und-PP-diNAcBac-(Gal)\textsubscript{2} were prepared in an identical manner as the radiolabeled substrates, except that unlabeled substrates were used in all reactions. The oligosaccharides were labeled with 2-aminobenzamide as previously described (26, 32) and purified using the GlykoNSep column (Prozyme). The appropriate peaks were collected and matrix-assisted laser desorption ionization mass spectrometry (MALDI MS) was used to determine the mass of the 2-AB labeled glycans.

**Glycosyltransferase substrate specificity assays**

To determine the UDP-sugar specificity of PglB, PglA, and PglE, radioactivity-based were performed on a variety of UDP-linked sugar substrates.

The ability of PglB to transfer UDP-diNAcBac, UDP-Glc, UDP-Gal, UDP-GlcNAc, and UDP-GalNAc was analyzed. The activity of PglB was coupled to the action of S. mutans undecaprenol kinase in order to provide the undecaprenyl phosphate in situ (29). The
specificity assays included 3% DMSO, 1% Triton X-100, 50 mM MgCl₂, 30 mM Tris-Acetate (pH 8.0), 20 µM undecaprenol, 1 mM ATP, 2 µM UDP-[³H]sugar (20 mCi/mmol), 20 µL of undecaprenol kinase CEF, 5 µL PglB CEF and water to a final volume of 100 µL. In the case of UDP-diNAcBac transfer, 2 µM UDP-4-amino and 2 µM [³H]AcCoA (20 mCi/ mmol) were included to assay both activities of the bifunctional PglB, which carries out transfer of the acetyl group to UDP-4-amino and transfer of phospho-diNAcBac to undecaprenyl phosphate. The reactions were initiated with a mixture of ATP and PglB and were monitored by quenching 15 µL aliquots at 20, 40, 60, 80, and 100 seconds. The radioactivity present in the organic and aqueous layers was determined as described above.

The ability of PglA and PglE to transfer UDP-Glc, UDP-Gal, UDP-GlcNAc, and UDP-GalNac was also analyzed. In the case of PglA, the assays contained 3% DMSO, 0.05% Triton X-100, 50 mM MgCl₂, 30 mM Tris-Acetate (pH 8.0), 10 µM Und-PP-diNAcBac, 2 µM UDP-sugar (20 mCi/mmol), 0.1 µM PglA and water to a final volume of 100 µL. The reaction was monitored by quenching 15 µL at 1, 2, 3, 4, and 5 minutes. For PglE, the assays contained 3% DMSO, 0.05% Triton X-100, 50 mM MnCl₂, 50 mM HEPES (pH 7.5), 2 µM Und-PP-diNAcBac-Gal, 5 µM UDP-sugar (20 mCi/mmol), 5 µL PglE CEF and water to a final volume of 100 µL. The reaction was monitored by quenching 15 µL at 2, 4, 6, 8, and 10 minutes. For both assays, the reactions were initiated by addition of enzyme and the radioactivity present in the organic and aqueous layers was determined as described above.

In addition, a coupled reaction was performed to test the ability of PglB and PglA to distinguish between UDP-4-amino and UDP-diNAcBac. The reaction components contained 3% DMSO, 1% Triton X-100, 40 mM MgCl₂, 30 mM Tris-Acetate (pH 8.0), 50 µM undecaprenol, 1 mM ATP, 500 µM UDP-4-amino, 500 µM AcCoA, 90 nM UDP-Gal (20 Ci/mmol), 5 µL undecaprenol kinase CEF, 10 µL PglB CEF and 1 µM PglA with a final volume of 100 µL. The reaction was initiated with a mixture of ATP and PglB. The extent of the reaction was monitored by quenching 15 µL at 1, 3, 5, 7, and 9 minutes. To test if PglB and PglA could recognize UDP-4-amino, a second reaction was prepared in which the AcCoA was omitted. The quenched aliquots were extracted as described above and the radioactivity present in the organic fraction was determined by scintillation counting.

**Oligosaccharyltransferase assays**

The OTase reactions were performed with a variety of Und-PP-linked glycosyl donors. In general, the reactions contained 5% DMSO, 0.7% Triton X-100, 50 mM MnCl₂, 25 mM HEPES (pH 7.5), 70 mM sucrose, 10–20 nM Und-PP-substrate (20 Ci/mmol), 8 µM pilin and 1 µM PglO in 100 µL reaction volume. The reactions were incubated overnight at room temperature with shaking. The glycosylated pilin protein was isolated via Ni-NTA purification. Briefly, the reaction was incubated with 15 µL of Ni-NTA resin for several hours in a 1.5 mL eppendorf tube. The tube was briefly centrifuged and the supernatant was removed. The resin was then washed five times with 500 µL Buffer A containing 30 mM imidazole and 0.1% Triton X-100. For each wash, the buffer was added to the eppendorf tube, the resin was mixed thoroughly with buffer, and the supernatant was removed following a brief centrifugation. The protein was eluted in three fractions of 500 µL of Buffer A containing 300 mM imidazole and 0.1% Triton X-100. Scintillation fluid (Ecolite(+™, MP Biomedicals) was added to all flow-through, wash and elution fractions and the radioactivity of each sample was determined.

Glycosylated protein samples for Western blot analysis were prepared and purified in the same manner, except that unlabeled versions of Und-PP-diNAcBac substrate was used at concentrations of 10–24 µM. Parallel reactions with radioactive substrates were performed at identical concentrations in order to determine reaction yields. The Western blot analysis
was performed following standard protocols. An antibody specific for His$_4$ (Qiagen) was used as a positive control for the purified proteins and a diNacBac-epitope monoclonal antibody termed npg1, which was previously described (33), was used to detect diNacBac-modified protein.

Results

Determination of UDP-DATDH stereochemistry by NMR

The biosynthesis of UDP-DATDH from UDP-GlcNAc was carried out in the presence of purified dehydratase (PglD), aminotransferase (PglC), and acetyltransferase-phospho-glycosyltransferase (PglB). PglC, a soluble protein, was purified to homogeneity (Figure S1, lane 2). TMHMM, a transmembrane prediction program, (34) predicts that PglD and PglB contain four and one transmembrane helices, respectively. SDS-PAGE analysis of purified PglD demonstrated that the desired protein product is the dominant component (Figure S1, lane 1). PglB was purified for this experiment, but the enzyme was used as a partially purified CEF (Figure S1, lane 4) in all other assays to avoid problems with protein stability. The anti-His$_4$ Western Blot analysis revealed that both purified PglD and PglB CEF contained His$_6$-tagged truncation products that formed during protein expression (Figure S1, lanes 1 and 4).

The biosynthesis of UDP-diNacBac by the action of PglD, PglC, and PglB was followed by capillary electrophoresis to ensure complete turnover of the substrates. This method also verified that the HexNAc substrate of PglD is UDP-GlcNAc, and not UDP-GalNAc (data not shown). Purification by reverse phase-HPLC removed unreacted substrates and cofactors leading to a final UDP-DATDH purity of $>$95%. To determine the final stereochemistry of the sugar, $^1$H NMR (Figure S2) was employed to compare the chemical shifts and coupling constants with UDP-diNacBac from the \textit{C. jejuni} pathway (Table 1). The values for the UDP-DATDH sugar from \textit{N. gonorrhoeae} exactly match the values of UDP-diNacBac from \textit{C. jejuni} (28). Further confirmation was provided by the $^{31}$P, $^{13}$C, and $^1$H-$^1$H COSY NMR spectra (Figures S3–S5). Therefore, the stereochemistry of the DATDH sugar in the \textit{N. gonorrhoeae} pathway is confirmed as diNacBac.

Functional characterization of PglB-ATD

The similarity of the \textit{N. gonorrhoeae} protein glycosylation pathway to the pathway in \textit{C. jejuni} suggests that the acetyltransferase domain of PglB acts first on UDP-4-amino to generate UDP-diNacBac, which is then utilized as a substrate by the phospho-glycosyltransferase domain of PglB (PglB-PGTD, Figure 1). The C-terminal acetyltransferase domain of full-length PglB (PglB-ATD, based upon a ClustalW alignment with PglD(\textit{Cj})) was purified (Figure S1, lane 3). This provided a suitable amount of well-behaved, soluble protein in the absence of the N-terminal phospho-glycosyltransferase domain, which is predicted by TMHMM to contain a single transmembrane domain (34). Functional analysis of PglB-ATD described below confirmed definitively that this domain acetylates UDP-4-amino to produce UDP-diNacBac, which is a substrate for PglB-PGTD mediated transfer of P-diNacBac to Und-P.

Kinetic characterization of PglC and PglB-ATD

Both the aminotransferase (PglC) and acetyltransferase (PglB-ATD) reactions exhibited typical Michaelis-Menten kinetics over a wide range of substrate concentrations (Figure S6). Initial velocity data were used to calculate kinetic parameters of L-Glu and UDP-4-keto for PglC and AcCoA and UDP-4-amino for PglB-ATD. Each reaction was run in duplicate and the initial velocities fit to equation 1 to yield the kinetic parameters in Table 2.
Functional characterization of the glycosyltransferases

As mentioned above, TMHMM (34) predicts that PglB has a single N-terminal transmembrane helix. In addition, PglE is predicted to contain two C-terminal transmembrane helices. Purification of these proteins by detergent solubilization and extraction resulted in low yields and loss of activity; to avoid these problems, both PglB and PglE were purified as crude CEFs for the glycosyltransferase assays. SDS-PAGE and Western blot analysis showed that PglB and PglE are the predominant bands present in the respective CEFs (Figure S1, lanes 4 and 6). In all assays involving PglB and PglE, negative controls with CEFs lacking overexpressed PglB or PglE showed no glycosyltransferase activity (data not shown). PglA is predicted to be soluble and was purified to homogeneity (Figure S1, lane 5).

Tritium-labeled products of PglB, PglA, and PglE were analyzed by NP-HPLC. Und-PP-[\(^{3}H\)]diNAcBac, Und-PP-diNAcBac-[\(^{3}H\)]Gal and Und-PP-diNAcBac-Gal-[\(^{3}H\)]Gal were retained on the column with retention times consistent with glycan size (Figure 3). Each product was analyzed separately in order to confirm the identity of the peaks (Figure S7). In addition, the glycosyltransferase products were characterized by a standard 2-AB fluorescence-labeling protocol as previously described (35). The 2-AB labeled disaccharide and trisaccharide were purified and MALDI MS was used to verify the masses of the products (Figure S8). These studies definitively annotate the biochemical functions of PglB, PglA, and PglE as the phospho-glycosyltransferase and the two glycosyltransferases that produce Und-PP-linked mono-, di- and trisaccharides, respectively.

UDP-saccharide specificity of glycosyltransferases

The substrate specificities of PglB, PglA and PglE were explored through the use of radioactivity-based assays. Organic extraction of the hydrophobic undecaprenyl-linked product allowed for quantification of the amount of radiolabeled sugar transferred to the undecaprenyl substrate similar to previously described assays (26, 31). The isoprenyl-linked substrates for the assays (Und-PP-diNAcBac for PglA and Und-PP-diNAcBac-Gal for PglE) were produced enzymatically and purified by NP-HPLC. The undecaprenyl phosphate required for the PglB reaction was generated in situ from undecaprenol and ATP with S. mutans undecaprenol kinase as previously described (29). The activities of the three enzymes were screened with UDP-Glc, UDP-Gal, UDP-GlcNAc, UDP-GalNAc and in the case of PglB, UDP-diNAcBac (Figure 4). In addition, the ability of PglB and PglA to distinguish between UDP-4-amino and UDP-diNAcBac was evaluated through a coupled assay (Figure S9). In all cases, the enzymes were highly specific for the corresponding predicted sugar substrate; PglB exclusively transferred phospho-diNAcBac, while PglA and PglE transferred only Gal (Figure 4).

Undecaprenyl diphosphate disaccharide specificity of PglE

Facile enzymatic synthesis of Und-PP-diNAcBac-Gal and Und-PP-diNAcBac-GalNAc produced by the Neisseria and C. jejuni pathways, respectively, allowed for examination of the substrate specificity of PglE for the acceptor oligosaccharide. Somewhat surprisingly, PglE is able to add a Gal residue to both the native substrate, Und-PP-diNAcBac-Gal, and to the C. jejuni substrate Und-PP-diNAcBac-GalNAc (Figure 5). This confirms in vivo studies in which the C. jejuni PglA was expressed in N. gonorrhoeae and the resultant trisaccharide (diNAcBac-GalNAc-Gal) was observed as a covalent pilin modification (1).

Characterization of PglH, an alternative glycosyltransferase

Recently an alternative glycosyltransferase in N. gonorrhoeae, PglH, was identified and shown to transfer a Glc unit to Und-PP-diNAcBac (14) (Figure 2). PglH was shown to be
responsible for the specific addition of Glc to Und-PP-diNAcBac by a variety of in vivo and in vitro methods. In vitro radiolabeled assays demonstrated that PglH transfers Glc from UDP-Glc to Und-PP-diNAcBac and does not transfer Man, Gal, GlcNAc or GalNAc; MALDI MS of the 2-AB-labeled diNAcBac-Glc confirmed the identity of the PglH product (14).

Herein, we further characterize PglH and compare its function to the other N. gonorrhoeae glycosyltransferases (Figure S1, lane 8). Analysis of the radiolabeled PglH product, Und-PP-diNAcBac-[3H]Glc, by NP-HPLC revealed that the retention time (30 minutes) was very similar to the Und-PP-diNAcBac-[3H]Gal retention time (29–30 minutes) (Figure 6). In addition, in vivo evidence suggested that unlike the C. jejuni disaccharide, the PglH product was not further modified by the third glycosyltransferase PglE (14). This result was validated by the in vitro specificity assay described above for PglE and established that PglE was unable to transfer Gal to Und-PP-diNAcBac-Glc (Figure 5).

Functional characterization of oligosaccharyltransferase, PglO

PglO and pilin are integral membrane proteins and were expressed in E. coli, extracted from the CEF with Triton X-100 and purified to homogeneity (Figure S1, lanes 7 and 9). To assay for OTase activity, purified PglO was incubated with pilin and radiolabeled Und-PP-diNAcBac-[3H]Gal glycan donor. After overnight incubation, the reaction mixture was bound to Ni-NTA resin and washed thoroughly to remove most (>99%) of the unreacted Und-PP-diNAcBac-[3H]Gal donor. The pilin protein was then eluted with imidazole and the radioactivity associated with the wash and elution fractions was determined by scintillation counting (Figure S10). Under these assay conditions, in which pilin protein is in excess over Und-PP-diNAcBac-[3H]Gal, PglO transferred ~60% of the sugar substrate to pilin (Figure 7). Interestingly, unlike OTases in N-linked glycosylation pathways (27, 36), PglO does not readily glycosylate a short peptide based on the pilin glycosylation sequence (Figure S11).

These results were further verified via Western blot analysis utilizing a monoclonal antibody recognizing a diNAcBac-associated epitope (Figure 7). For the Western blot analysis, the pilin glycosylation reaction was performed with equimolar amounts of protein substrate and Und-PP-diNAcBac donor and under these conditions ~13% of the pilin protein was associated with glycan. The antibody raised against diNAcBac showed strong staining with the glycosylated pilin and was unreactive with the unmodified pilin (33).

Glycan donor specificity of PglO

To further characterize PglO, a screen of various oligosaccharyl donors was performed. Previous studies by Feldman and coworkers (21, 22) on PglL, the homologous oligosaccharyltransferase found in Neisseria meningitidis, have suggested that these enzymes exhibit relaxed substrate specificity in vivo and can transfer oligosaccharides composed of different sugars, linkages and lengths. Thus, PglO was assayed with the four native substrates (the products of PglB, PglA, PglE and PglH) and with two substrates from the C. jejuni pathway: Und-PP-diNAcBac modified with one or two GalNAc residues. (For structures of the six glycans tested and further discussion of OTase specificity, see Figure S12 of the supporting information.) All glycosyl donors were prepared with high specific activity and purified via NP-HPLC. Surprisingly, and in contrast to the in vivo studies with PglL(Nm) (21, 22), PglO was only able to transfer the four native substrates; the two C. jejuni substrates had <3% turnover (Figure 8).

To verify that the C. jejuni substrates were functional as glycan donors, all six substrates were assayed with PglB(Cj). Pilin was used as the protein substrate in these assays as well, because it contains an N-linked glycosylation sequon (59ENNTS63) adjacent to the site of O-
linked glycosylation (37, 38). As seen in Figure 9, PglB(Cj) transferred the native C. jejuni substrates in addition to the diNAcBac-Gal disaccharide from N. gonorrhoeae; it showed low reactivity with diNAcBac-Glc and diNAcBac-(Gal)_2.

Since pilin is glycosylated by both PglB(Cj) and PglO(Ng), it was important to confirm the identity of the glycosylated residues. Pilin variants were prepared with alanine mutations at the expected sites of glycosylation for PglB(Cj) and PglO. Asn 61 and Ser 63, respectively. PglB(Cj) was unable to glycosylate pilin-N61A, validating this residue as the N-glycan acceptor site. However, PglO showed ~85% of normal activity with the pilin S63A mutant suggesting that another site, potentially Thr 62, is a glycosyl acceptor site in the absence of Ser 63 (Figure S13). Further mutational analysis confirmed this hypothesis; pilin-T62A exhibited normal glycosylation, whereas glycosylation was greatly reduced in the pilin double mutant (T62A/S63A) (Figure S13).

Discussion
Stereochemistry of DATDH defined as diNAcBac

The extreme diversity of bacterial glycans is highly significant since these glycans typically decorate the bacterial cell surface facilitating interactions with host cells (7, 10) and potentially confounding the immune response (8, 39–42). Bacillosamine was originally identified as an unusual 2,4-diamino-2,4,6-trideoxy-α-D-glucose in Bacillus subtilis (43), but it also appears in the mono- and di-aminoacetylated forms in a large variety of bacterial glycoconjugates (44). It has been found frequently in the O-antigen and capsular polysaccharide of Gram-negative bacteria, but has also been identified in the S-layer of Gram-positive bacteria and as the UDP-activated donor in cellular extracts (45). DiNAcBac was initially discovered in N-linked glycans in C. jejuni (24), but more recently, a second route to diNAcBac was biochemically characterized in C. jejuni, in which GDP-diNAcBac is an intermediate in the CMP-legionaminic acid biosynthetic pathway (46). The pilin oligosaccharide in N. gonorrhoeae was thought originally to comprise Gal-α-(1,3)-GlcNAc-β-Ser 63 (47). Mass spectrometry and bioinformatic analysis suggested that the linking sugar unit was DATDH instead of GlcNAc (1). Herein, we confirm the stereochemical assignment of this sugar for the first time showing that the DATDH sugar in N. gonorrhoeae is diNAcBac (Table 1). This adds to the growing number of oligosaccharides identified in bacteria that contain forms of bacillosamine.

PglD, PglC and PglB produce Und-PP-diNAcBac in N. gonorrhoeae; these three enzymes have functional homology to PglF(Cj), PglE(Cj), PglD(Cj), and PglC(Cj) in C. jejuni, which produce the same polyprenyl-linked intermediate (Figure 2). Even though the early enzymes in these two pathways carry out identical functions, the sequence identity is relatively low (25–30%), except for the phospho-glycosyltransferase domain of PglB, which has 52% identity with PglC(Cj) (Table 3). These numbers starkly contrast the sequence identity observed between N. gonorrhoeae and N. meningitidis, which indicate much closer homologies (>84%, Table 3). These numbers imply that C. jejuni and N. gonorrhoeae pathways are only distantly related from an evolutionary standpoint.

Work is ongoing to understand the low level of homology between the C. jejuni and N. gonorrhoeae diNAcBac biosynthetic enzymes. As a first step, this study describes the kinetic parameters of both the aminotransferase (PglC) and the acetyltransferase (PglB-ATD) (Table 2). The apparent K_m of the UDP-4-keto sugar for PglC (233 µM) was comparable to the PglE(Cj) homologue (48 µM (48) and 610 µM (49)) and well within the typical binding efficiency for this type of substrate. Likewise, K_m values of the UDP-4-amino substrate for the acetyltransferases PglB-ATD (122 µM) and PglD(Cj) (410 µM (28)) lead to a similar conclusion. However, the N. gonorrhoeae enzymes presented here are
catalytically much less efficient ($k_{\text{cat}}$ is 10–100 fold less for PglC and 1000-fold less for PglB-ATD) than their *C. jejuni* counterparts with respect to the UDP-sugar. This observation is reflected in the differences between their specificity constants ($k_{\text{cat}}/K_{m}$) (Table 2). The high acetyltransferase activity in the *C. jejuni* pathway is used to drive the biosynthesis of the UDP-diNAcBac sugar (28). A similar phenomenon is observed in the *N. gonorrhoeae* pathway, with a 20-fold enhancement in $k_{\text{cat}}/K_{m}$ of PglB-ATD with respect to aminotransferase activity.

For PglB-ATD, one cannot rule out interplay between the acetyltransferase domain and the missing C-terminal phospho-glycosyltransferase domain. Therefore, care must be taken in interpreting the reduced *N. gonorrhoeae* acetyltransferase efficiency as compared to PglD(*Cj*). Further work will be necessary to clarify how domain interactions affect kinetic parameters. In addition, the low sequence homology between the *C. jejuni* and *N. gonorrhoeae* UDP-diNAcBac pathway enzymes (Table 3) could contribute to the differences in catalytic efficiency observed here.

**Significant glycan diversity in *N. gonorrhoeae* protein glycans**

The *C. jejuni* and *N. gonorrhoeae* pathways diverge after the synthesis of Und-PP-diNAcBac. The *C. jejuni* pathway continues to N-linked glycan assembly with the successive addition of five α-(1,4)-linked GalNAc units and a branching Glc unit. However, while the *C. jejuni* N-linked heptasaccharide is highly conserved, *N. gonorrhoeae* strains display high O-linked glycan diversity. Strains have been identified which contain not only O-linked disaccharide (Gal-α-(1,3)-diNAcBac) and trisaccharide (Gal-β-(1,4)-Gal-α-(1,3)-diNAcBac) produced by PglA and PglE, respectively, but also an alternate disaccharide (Glc-α-(1,3)-diNAcBac) produced by PglH (14). Further glycan modification occurs from the addition of O-acetyl groups by PglI (1). In addition, an alternate allele (*pglB2*) has been identified in *N. meningitidis* that contains a domain proposed to transfer a glyceroyl moiety instead of an acetyl group to produce 4-glyceramido-2-acetamido-2,4,6-trideoxy-α-D-hexose (GATDH) (20). This combination of biosynthetic enzymes allows neisserial strains to display a glycan repertoire with at least 13 identified glycan permutations (14). Additional glycan variation can occur within a single strain as phase variation of the genes encoding for PglA, PglE and PglH acts as another mode of glycan regulation (1).

In light of the significant amount of protein glycan variation present within strains of *N. gonorrhoeae*, it is surprising that the glycosyltransferases display such strict specificity (Figure 4, Figure S9). We have demonstrated that PglB, PglA and PglE are specific for the native substrate (UDP-diNAcBac or UDP-Gal) and will not accept any other form of the nucleotide-activated sugars commonly found in vivo, even though one of the alternate substrates contains only a single stereochemical change (UDP-Glc vs. UDP-Gal) and another contains only an additional acetamido group (UDP-GalNAc vs. UDP-Gal). Parallel work has shown similar strict specificity of the fourth glycosyltransferase, PglH (14). These results suggest that glycan identity is regulated at the level of biosynthesis and that these enzymes have evolved to selectively catalyze reactions in the milieu of intracellular NDP-sugars.

Along the same lines, PglE transferred a Gal unit onto native Und-PP-diNAcBac-Gal, but it showed little activity with Und-PP-diNAcBac-Glc, the alternate PglH disaccharide (Figure 5). PglE has evolved to detect the stereochemical difference between Glc and Gal, which is consistent with the model that PglA and PglE have evolved in tandem to produce a trisaccharide that is structurally distinct from the disaccharide produced by PglH (14). In contrast, it is surprising that PglE would recognize the *C. jejuni* substrate Und-PP-diNAcBac-GalNAc (Figure 5), but it is consistent with the hypothesis that the glycosyltransferases exhibit specificity relative to other substrates present in the organism.
PglE may not have developed selectivity against the additional acetamido group in the *C. jejuni* disaccharide, because it is not found in the native *N. gonorrhoeae* glycome.

The diverse O-linked glycans produced by *N. gonorrhoeae* decorate a number of extracellular proteins, but were first discovered as covalent modifications of Ser 63 in pilin. This residue is found within the α-β loop, which is surface exposed in the assembled pilin structure (50) and highly modified by other post-translational modifications including addition of phosphoethanolamine and phosphocholine (12). This modified loop also contains a bacterial N-linked sequon that overlaps the O-linked glycosylation site, 59ENNTS63, which can act as a protein acceptor substrate for the N-linked OTase PglB(Cj) (38, 51). The presence of the N-linked sequon in the pilin structure is interesting from an evolutionary standpoint suggesting that post-translational modification of this conserved loop may be important for pilin structure and function and that in other bacteria, this loop could contain N-linked glycans. A survey of sequenced bacterial genomes that contain a PglB(Cj) OTase homolog revealed at least one bacterium, *Nitrosococcus halophilus*, with a predicted pilin homolog containing the requisite Asn consensus sequence.

**En bloc glycan transfer from polyprenyl-linked intermediate to Ser residue in pilin**

The *C. jejuni* and *N. gonorrhoeae* pathways culminate in transfer of the oligosaccharide to protein; in the bacterial N-linked glycosylation pathway, PglB(Cj) transfers a heptasaccharide en bloc to the amide side chain of asparagine residues. Herein we demonstrate that PglO acts in a similar en bloc manner to transfer mono-, di- and trisaccharides to hydroxyl side chains of serine or threonine residues (Figures 7, 8).

PglO was originally identified as the enzyme responsible for glycosylation of *N. gonorrhoeae* Type IV pili, but has since been shown to glycosylate a wide variety of periplasmic and extracellular lipoproteins. In all examples of non-pilin glycosylation substrates, the acceptor serine or threonine residues are present in loop regions predicted to have undefined structures rich in Ala, Ser, and Pro residues (2). Interestingly, PglO was not active in the presence of a peptide substrate modeled on the pilin sequence (Figure S10). The inactivity of PglO contrasts the behavior of the bacterial N-linked OTase, PglB(Cj), which under comparable conditions is capable of glycosylating a peptide substrate (Figure S10). PglO glycosylates a wide range of periplasmic proteins containing serine and threonine residues in vivo, but it is unclear what binding determinants affect this reaction. Further biochemical and structural analyses are needed to understand how PglO recognizes pilin and non-pilin protein substrates.

**PglO in vitro assays demonstrate specificity for native glycans**

Recent in vivo analyses revealed that both PglL(Nm), which has 95% sequence identity with PglO, and the N-linked OTase PglB(Cj) were promiscuous enzymes that transferred a variety of Und-PP-linked O-antigen substrates to serine or asparagine residues in proteins, respectively (21, 22). These experiments were performed by co-expressing PglL(Nm) or PglB(Cj) heterologously in *E. coli* with the glycan-acceptor protein and a locus encoding for the biosynthesis of an Und-PP-linked substrate. In addition, previous studies showed that *N. gonorrhoeae* strains with heterologously expressed PglA(Cj) contained proteins modified by the *C. jejuni* disaccharide, diNAcBac-GalNAc, implying that PglO can recognize this glycan in vivo (1). However, in this study, we have found that the OTases do not show a comparable substrate promiscuity in vitro; in fact, it appears that PglO and PglB(Cj) are both specific for their native substrates in vitro (Figures 8, 9, S12).

The previous in vivo studies on PglO, PglL(Nm) and PglB(Cj) were performed in the absence of native substrate and under conditions in which the non-native undecaprenyl-
linked glycan accumulated in the membrane (21, 22). Thus, the local concentration of the substrate within the two-dimensional plane of the membrane was likely to be much higher than in the \textit{in vitro} assay, which would promote reaction with PglO. Additionally, in the context of a lipid bilayer, the role of the membrane-bound undecaprenyl moiety may play a greater role in enzyme recognition of the substrate. In our assay, the concentration of undecaprenyl substrate (10–20 nM) was well below 2.7 µM, the apparent \(K_m\) of PglB(Cj) for Und-PP-disaccharide (37), and thus specificity differences between native and non-native substrates were easily distinguished. In addition, it should be noted that the OTases exhibit substrate specificity in native cellular contexts. In the native bacteria, PglO, PgL(Nm), and PglB(Cj) selectively transfer the correct oligosaccharide to pilin in the presence of other undecaprenyl-linked substrates, including those involved in capsular polysaccharide biosynthesis in \textit{N. meningitidis} (52) and \textit{C. jejuni} (53) and the peptidoglycan subunits in the cellular membranes of all three species (54).

In conclusion, this work represents the first complete biochemical characterization of the unusual O-linked glycosylation pathway in \textit{N. gonorrhoeae}. The stereochemistry of the DATDH sugar has been identified as diNAcBac. In addition, the substrate preferences of the glycosyltransferases have been characterized and in general these enzymes are shown to be specific for their native substrates. Finally, \textit{in vitro} characterization of the OTases from \textit{N. gonorrhoeae} and \textit{C. jejuni} has suggested that these enzymes prefer their respective native glycans to closely related oligosaccharides. The O-linked pathways found in \textit{N. gonorrhoeae} and \textit{N. meningitidis} are interesting hybrids of O-linked and N-linked glycosylation pathways. While the role of the O-linked glycans in \textit{Neisseria} pathogenicity is not yet understood, the Pgl enzymes may represent unique virulence targets and this initial study provides the foundation for further investigations into the biochemistry of the enzymes.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

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Figure 1.
Biosynthetic pathway of the pilin glycan in *Neisseria gonorrhoeae*.
Figure 2.
Schematic representations of bacterial protein glycosylation pathways: (top) O-linked pathway in *Neisseria gonorrhoeae* and *Neisseria meningitidis*; (middle) alternative O-linked pathway in *Neisseria* species; (bottom) N-linked pathway in *Campylobacter jejuni*. The proteins predicted to contain transmembrane domains by TMHMM are indicated by a black box.
Figure 3. Normal phase HPLC separation of radiolabeled Und-PP-[\(^3\)H]diNAcBac (27 minutes), Und-PP-diNAcBac-[\(^3\)H]Gal (30 minutes), and Und-PP-diNAcBac-Gal-[\(^3\)H]Gal (45 minutes). Fractions were eluted at 1 mL/min and radioactivity (DPM) was determined by scintillation counting.
Figure 4. Specificity analyses of PglB (A), the phospho-glycosyltransferase, and PglA (B) and PglE (C), the galactosyltransferases, in the presence of a panel of UDP-sugar substrates. The reactions were carried out in a volume of 100 µL. The assays were performed in triplicate and the error bars indicate standard deviation.
Figure 5.
Determination of the isoprenyl-linked substrate preferences of PglE in the presence of Und-PP-diNAcBac-Glc (dashed line), Und-PP-diNAcBac-Gal (solid line), or Und-PP-diNAcBac-GalNAc (dotted line). The reactions were carried out in a volume of 100 µL. The assays were performed in triplicate and the error bars indicate standard deviation.
Figure 6.
Normal phase HPLC separation of radiolabeled Und-PP-diNAcBac-[\(^3\)H]Glc, the product of PglH (30 minutes). Fractions were eluted at 1 mL/min and radioactivity (DPM) was determined by scintillation counting.
Figure 7.
Left, PglO reaction turnover after overnight incubation with Und-PP-diNacBac-[³H]Gal in the presence of pilin or a negative control with no protein substrate. Right, Western blot analysis of unmodified pilin and PglO-glycosylated pilin using His₄ antibody (left blot, positive control) and a diNacBac-epitope recognizing monoclonal antibody termed npg1 (right blot, specific for glycan).
Figure 8.
PglO reaction turnover after overnight incubation with pilin protein in the presence of a panel of Und-PP-linked substrates from \( N.\ gonorrhoeae \) (diNAcBac-Gal, diNAcBac-(Gal)\(_2\), diNAcBac-Glc), \( C.\ jejuni \) (diNAcBac-GalNAc, diNAcBac-(GalNAc)\(_2\)) or both (diNAcBac). A negative control is shown in which the assay is performed in the presence of Und-PP-diNAcBac-Gal and the absence of PglO. The assays were performed in triplicate and the error bars indicate standard deviation.
Figure 9.
PglB reaction turnover after overnight incubation with pilin protein in the presence of a panel of Und-PP-linked substrates from *N. gonorrhoeae* (diNAcBac-Gal, diNAcBac-(Gal)$_2$, diNAcBac-Glc), *C. jejuni* (diNAcBac-GalNAc, diNAcBac-(GalNAc)$_2$) or both (diNAcBac). A negative control is shown in which the assay is performed in the presence of Und-PP-diNAcBac-GalNAc and the absence of PglB. The assays were performed in triplicate and the error bars indicate standard deviation.
Table 1

Comparison of *C. jejuni* and *N. gonorrhoeae* UDP-diNAcBac $^1$H chemical shift and coupling constant assignments.

<table>
<thead>
<tr>
<th>Moiety</th>
<th>C. jejuni (28)</th>
<th>N. gonorrhoeae</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>d$_H$ (ppm)</td>
<td>J</td>
</tr>
<tr>
<td>H1</td>
<td>5.48 (dd)</td>
<td>J$<em>{1,2}$ = 3.2 Hz, J$</em>{1,P}$ = 6.9 Hz</td>
</tr>
<tr>
<td>H2</td>
<td>4.02 (m)</td>
<td></td>
</tr>
<tr>
<td>H3</td>
<td>3.79 (at)</td>
<td>J$<em>{2,3}$ = 10.2 Hz, J$</em>{3,4}$ = 10.2 Hz</td>
</tr>
<tr>
<td>H4</td>
<td>3.69 (at)</td>
<td>J$_{4,5}$ = 10.2 Hz</td>
</tr>
<tr>
<td>H5</td>
<td>4.05 (m)</td>
<td></td>
</tr>
<tr>
<td>H6</td>
<td>1.19 (d)</td>
<td>J$_{5,6}$ = 6.2 Hz</td>
</tr>
</tbody>
</table>
Table 2
Steady-state parameters for PglC and PglB-ATD.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Substrate</th>
<th>$k_{cat}$ (s$^{-1}$)</th>
<th>$K_m$ (µM)</th>
<th>$k_{cat}/K_m$ (s$^{-1}$M$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PglC</td>
<td>L-Glutamate</td>
<td>0.025 ± 0.001</td>
<td>4900 ± 900</td>
<td>5.1</td>
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<tr>
<td>PglC</td>
<td>UDP-4-keto</td>
<td>0.039 ± 0.002</td>
<td>233 ± 35</td>
<td>167</td>
</tr>
<tr>
<td>PglB-ATD</td>
<td>AcCoA</td>
<td>0.928 ± 0.032</td>
<td>456 ± 34</td>
<td>2035</td>
</tr>
<tr>
<td>PglB-ATD</td>
<td>UDP-4-amino</td>
<td>0.416 ± 0.016</td>
<td>122 ± 17</td>
<td>3410</td>
</tr>
</tbody>
</table>
Table 3

Percent sequence identity (%) between *N. gonorrhoeae* (*Ng*) and *C. jejuni* (*Cj*) and between *N. gonorrhoeae* (*Ng*) and *N. meningitidis* (*Nm*) proteins.

<table>
<thead>
<tr>
<th>Function</th>
<th><em>N. gonorrhoeae</em></th>
<th><em>C. jejuni</em></th>
<th>% <em>Ng</em> and <em>Cj</em></th>
<th><em>N. meningitidis</em></th>
<th>% <em>Ng</em> and <em>Nm</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>dehydratase</td>
<td>PglD</td>
<td>PglF</td>
<td>29.8</td>
<td>PglD</td>
<td>92.5</td>
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<tr>
<td>aminotransferase</td>
<td>PglC</td>
<td>PglE</td>
<td>21.2</td>
<td>PglC</td>
<td>92.8</td>
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<tr>
<td>acetyltransferase</td>
<td>PglB-ATD</td>
<td>PglD</td>
<td>29.7</td>
<td>PglB-ATD</td>
<td>84.9</td>
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<tr>
<td>P-glycosyltransferase</td>
<td>PglB-GTD</td>
<td>PglC</td>
<td>52.3</td>
<td>PglB-GTD</td>
<td>90.3</td>
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<tr>
<td>glycosyltransferase</td>
<td>PglA</td>
<td>PglA</td>
<td>21.3</td>
<td>PglA</td>
<td>95.5</td>
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<tr>
<td>glycosyltransferase</td>
<td>PglE</td>
<td>PglJ</td>
<td>12.0</td>
<td>PglE</td>
<td>93.0</td>
</tr>
<tr>
<td>OTase</td>
<td>PglO</td>
<td>PglI</td>
<td>11.8</td>
<td>PglL</td>
<td>95.0</td>
</tr>
</tbody>
</table>