The N-linking glycosylation system from *Actinobacillus pleuropneumoniae* is required for adhesion and has potential use in glycoengineering

Jon Cuccui¹, Vanessa S. Terra¹, Janine T. Bossé², Andreas Naegeli³, Sherif Abouelhadid¹, Yanwen Li², Chia-Wei Lin², Prema Vohra¹, Alexander W. Tucker⁴, Andrew N. Rycroft⁵, Duncan J. Maskell⁴, Markus Aebi³, Paul R. Langford² and Brendan W. Wren¹ on behalf of the BRaDP1T Consortium

¹Faculty of Infectious and Tropical Diseases, London School of Hygiene and Tropical Medicine, Keppel Street, London WC1E 7HT, UK
²Section of Paediatrics, Department of Medicine, Imperial College London, St. Mary’s Campus, London W2 1PG, UK
³Institute of Microbiology, ETH Zurich, Wolfgang-Pauli-Strasse 10, 8093 Zurich, Switzerland
⁴Department of Veterinary Medicine, University of Cambridge, Madingley Road, Cambridge CB3 0ES, UK
⁵The Royal Veterinary College, Hawkshead Campus, Hatfield, Hertfordshire AL9 7TA, UK

The *Actinobacillus pleuropneumoniae* is a mucosal respiratory pathogen causing contagious porcine pleuropneumonia. Pathogenesis studies have demonstrated a major role for the capsule, exotoxins and outer membrane proteins. *Actinobacillus pleuropneumoniae* can also glycosylate proteins, using acytoplasmic N-linked glycosylating enzyme designated NGT, but its transcriptional arrangement and role in virulence remains unknown. We investigated the NGT locus and demonstrated that the putative transcriptional unit consists of *rimO*, *ngt* and a glycosyltransferase termed *agt*. From this information we used the *A. pleuropneumoniae* glycosylation locus to decorate an acceptor protein, within *Escherichia coli*, with a hexose polymer that reacted with an anti-dextran antibody. Mass spectrometry analysis of a truncated protein revealed that this operon could add up to 29 repeat units to the appropriate sequon. We demonstrated the importance of NGT in virulence, by creating deletion mutants and testing them in a novel respiratory cell line adhesion model. This study demonstrates the importance of the NGT glycosylation system for pathogenesis and its potential biotechnological application for glycoengineering.

1. Introduction

*Actinobacillus pleuropneumoniae* is a Gram-negative bacterium and the causative agent of porcine pleuropneumonia, a severe respiratory disease responsible for significant losses to the pig industry worldwide. Economically, this disease has a huge impact on the pig industry, costing an average €6.4 per fattened pig in an affected herd in Europe [1]. *Actinobacillus pleuropneumoniae* enters the lungs and colonizes tissues by binding to mucus proteins and cells of the lower respiratory tract, including ciliated cells of the terminal bronchioli and alveolar epithelial cells [2,3]. There are 15 established serovars that differ in capsular polysaccharide composition [4], with another proposed based on serological results [5]. Several surface structures have been identified as being involved in adhesion, including fimbriae [6] and lipopolysaccharide (LPS) [7].

Advances in DNA sequencing technologies and mass spectrometry techniques reveal that post-translational modification of proteins by glycosylation is not
restricted to a few bacterial species and is often important in pathogenesis [8,9]. Understanding the mechanisms of bacterial glycosylation and its role in pathogenesis can have practical applications such as the design of novel bioglycoconjugate vaccines, antimicrobials and diagnostics [10,11]. Bacterial protein glycosylation systems can be broadly divided into two main categories: glycans that are covalently attached to amide groups of asparagine residues (N-linked) or to hydroxyl groups on serine/threonine residues (O-linked). These categories can be further subdivided depending on the cellular compartment where protein glycosylation takes place. Oligosaccharyltransferases (OTases) function in the periplasmic compartment of a bacterial cell and catalyse the transfer of an oligosaccharide from a lipid donor to an acceptor molecule, usually a protein. The best-characterized OTases are the donor to an acceptor molecule, usually a protein. The best-characterized OTases are the donor to a lipid (OTases) function in the periplasmic compartment of a bacterial protein glycosylation takes place. Oligosaccharyltransferases are subdivided depending on the cellular compartment where threonine residues (en bloc glycosylation operates through an asparagine residues (egon glycosylation) or to hydroxyl groups on serine/threonine residues (O-linked). These categories can be further divided depending on the cellular compartment where protein glycosylation takes place. Oligosaccharyltransferases (OTases) function in the periplasmic compartment of a bacterial cell and catalyse the transfer of an oligosaccharide from a lipid donor to an acceptor molecule, usually a protein. The best-characterized OTases are the donor to an acceptor molecule, usually a protein. The best-characterized OTases are the donor to an acceptor molecule, usually a protein. The best-characterized OTases are the donor to an acceptor molecule, usually a protein. The best-characterized OTases are the donor to a lipid (OTases) function in the periplasmic compartment of a bacterial protein glycosylation takes place. Oligosaccharyltransferases are subdivided depending on the cellular compartment where threonine residues (en bloc glycosylation operates through an asparagine residues (egon glycosylation) or to hydroxyl groups on serine/threonine residues (O-linked). These categories can be further divided depending on the cellular compartment where protein glycosylation takes place. Oligosaccharyltransferases (OTases) function in the periplasmic compartment of a bacterial cell and catalyse the transfer of an oligosaccharide from a lipid donor to an acceptor molecule, usually a protein. The best-characterized OTases are the donor to an acceptor molecule, usually a protein.

Figure 1. Genetic organization of the HMWC enzyme family. In contrast to other bacteria, NTHi has two copies of the HMW locus, and each has a gene encoding an acceptor protein. A. pleuropneumoniae is the only species that has a second glycosyltransferase adjacent to the N-linked enzyme.

2. Material and methods
2.1. Bacterial strains used and culture conditions
Actinobacillus pleuropneumoniae serovar 15 reference strain, HS143, or derived mutants were grown at 37°C with 5% CO₂ on BHI (Oxoid, UK) agar or broth, supplemented with...
10 μg ml\(^{-1}\) nicotinamide adenine dinucleotide (NAD) and when required with kanamycin (50 μg ml\(^{-1}\)) or chloramphenicol (1 μg ml\(^{-1}\)). *Escherichia coli* TOP10 and Mu Free Donor (MFD) [23] were grown in LB broth or agar (Oxoid) supplemented, when required, with 50 μg ml\(^{-1}\) kanamycin at 37°C. *E. coli* DH10 were grown in LB broth or agar (Oxoid) at 37°C supplemented, when required, with 80 μg ml\(^{-1}\) spectinomycin and/or 100 μg ml\(^{-1}\) trimethoprim.

### 2.2. Genomic DNA extraction

Total genomic DNA was extracted from a 10 ml overnight culture of *A. pleuropneumoniae* HS143, using a proteinase K and phenol: chloroform: isoamyl-alcohol-based procedure as previously described by Cuccui et al. [24].

### 2.3. Construction of Actinobacillus pleuropneumoniae knockout mutants

The *A. pleuropneumoniae* HS143 orthologues of *apl_1634* and *apl_1635* (also known as *agt* and *ngt*, coding for αGalT and NGT, respectively), found in the *A. pleuropneumoniae* L20 genome [25], were deleted using our recently described unmarked mutation system [26]. Primers used to generate the cat-sacB insertion/deletion and the unmarked deletion constructs for each gene are shown in electronic supplementary material, table S2. Briefly, the target genes and approximately 600–900 bp of flanking sequences were amplified using CloneAmp HiFi PCR Premix (Clontech), A-tailed and cloned into pGEMT (Promega), as previously described [26]. Inverse PCR was then used to open up the clones, using the appropriate primers, removing the target sequence and adding 15 bp overhangs to allow insertion of the cat-sacB cassette by In-Fusion cloning (Clontech). Unmarked deletion constructs were generated by amplifying the left and right flanking sequences for each gene, using appropriate primers with added 15 bp overhangs designed to allow direct fusion by overlap-extension PCR. The unmarked deletion mutants were then obtained by two sequential rounds of natural transformation as previously described [27].

### 2.4. Plasmid complementation

The vector pMKExpress [28] was digested with EcoRI and SacI (New England Biolabs, UK) and the resulting digest was gel purified using a Qiagen MinElute gel extraction kit (Qiagen, UK) before being purified using a Qiagen PCR purification kit (Invitrogen). The vectors pMKngt, pMKagt and pMKngt–agt, prior to transformation of the plasmid into One Shot *E. coli* TOP10 cells (Invitrogen) according to manufacturer’s instructions. Transformants were selected on LB agar supplemented with kanamycin (50 μg ml\(^{-1}\)). The complementation vectors were transformed into the mutant recipient strains by natural transformation as previously described [27].

### 2.5. Cell culture

The A549 cell line, adenocarcinoma human alveolar basal epithelial cells, (ATCC, CCL-185, US) was grown at 37°C, 5% CO\(_2\) in F-12 K medium (Gibco) supplemented with 10% fetal calf serum (Sigma).

### 2.6. Adhesion assay using A549 cell line

The A549 cell line (ATCC, CCL-185, USA) was seeded into 12-well tissue culture plates at a concentration of 2.5 × 10\(^4\) cells ml\(^{-1}\) and incubated overnight at 37°C 5% CO\(_2\). Bacterial overnight cultures (HS143 wild-type, isogenic ngt and *agt* mutants and complemented mutants) were used to seed into BHI–NAD medium and grown to an OD\(_{600}\) of 0.6. One millilitre of the suspension was added to the A549 cells at a multiplicity of infection (MOI) of 100 : 1, and the plates incubated at 37°C 5% CO\(_2\). After 3 h, non-adherent bacteria were removed by washing three times with 1 ml DPBS (Gibco), and adherent bacteria were released by adding 100 μl of 0.25% trypsin–EDTA (Sigma) for 5 min at 37°C. Trypsinization was stopped by the addition of 900 μl of DPBS. Serial dilutions were plated onto BHI–NAD plates for quantification of adherent bacteria. In order to determine if any of the recovered bacteria had invaded the A549 cells, controls were treated with gentamycin (Sigma) at a final concentration of 10 μg ml\(^{-1}\) for 1 h to allow for killing of adherent extracellular bacteria. The cells were then lysed by the addition of ice-cold sterile water, and serial dilutions were plated out on BHI–NAD.

### 2.7. Statistical analysis of adhesion assay data

The number of adherent cells was calculated by counting the colony forming units and comparing with the initial inoculum of each individual culture to determine the percentage of adherent cells. The statistical analysis was performed using a one-way analysis of variance followed by a Bonferroni’s multiple comparison test. The significance level was set at 0.05 throughout. Statistical analysis was done using GraphPad Prism v. 4.00 for Windows (GraphPad Software, San Diego, CA, www.graphpad.com).

### 2.8. Reverse transcriptase PCR

An overnight culture of *A. pleuropneumoniae* HS143 was diluted 1 : 20 in BHI–NAD broth. At the time points of 1.5, 3.0, 5 and 24 cycles of 94°C/30 s, 53°C/30 s, 68°C/2 min and a final 68°C/5 min cycle. All ORFs were expressed under the control of the plasmid promoter.

The resulting ngt only PCR product was digested with EcoRI and SacI, the *agt* only product was digested with PstI and NotI, and the ngt–agt PCR product was digested with EcoRI and NotI before being purified using a Qiagen PCR purification kit. Digested vector and PCR products were ligated using Promega T4 DNA ligase (Promega, UK) to yield the vectors pMKngt, pMKagt and pMKngt–agt, prior to transformation of the plasmid into One Shot *E. coli* TOP10 cells (Invitrogen) according to manufacturer’s instructions. Transformants were selected on LB agar supplemented with kanamycin (50 μg ml\(^{-1}\)). The complementation vectors were transformed into the mutant recipient strains by natural transformation as previously described [27].
24 h, RNA was extracted as previously described [29], with the minor modification that 2 μg of total RNA from each sample was treated with Ambion TURBO DNase (Invitrogen) according to the manufacturer's instructions. cDNA was generated from DNase-treated RNA using the SuperScript II kit (Invitrogen) according to the manufacturer's instructions. For each sample, 2 μl of reverse transcribed cDNA was used as a template in a 25 μl total volume PCR mixture and amplified using MyTaq master mix (Bioline, UK) using the following cycling conditions: 94°C/30 s, followed by 35 cycles of 94°C/10 s, 53°C/10 s, 72°C/10 s and a final single 72°C/30 s cycle using the primers listed in electronic supplementary material, table S2.

2.9. Quantitative real-time PCR validation

Total RNA was extracted from bacteria grown in BHI–NAD broth by using Tri Reagent (Sigma, UK) as previously described. Two micrograms from each sample was treated with Ambion TURBO DNase (Invitrogen) according to the manufacturer's instructions and some modifications. Total RNA was incubated for 1 h at 37°C followed by the addition of another two units of DNase. The sample was then incubated for an extra 1 h before inactivation.

cDNA was generated from DNase-treated RNA using the SuperScript III kit (Invitrogen) using random hexamers (Invitrogen) according to the manufacturer's instructions. One microtitre of this material was used in a QPCR using SYBR Green dye-based PCR amplification and detection system (Applied Biosystems). The A. pleuropneumoniae HS143 WT, HS143∆ngt and HS143∆agt were analysed for absolute quantification of cDNA using an ABI7500 Fast instrument (Applied Biosystems). Amplification was carried out using the following primers at a final concentration of 500 nM. agt fwd: 5'-GAT TGGATA GTG GAA GCC GA-3', agtrev: 5'-CCC TGG CCT TAA GTT ACG GA-3', ngtfwd: 5'-AGT TGG TGA GAG CAA CCG TG-3', ngtrev: 5'-AGT CCG AAT GGT TGG CT-3', rimOfwds2: 5'-CGT CCT CTT GAT GTG CAA GTG TT-3', rimOrevv2: 5'-CAC GTG TCC TGA AAA CCG TT-3'. Samples tested were four biological replicates, each tested as three technical replicates.

For comparative qRT-PCR analysis of A. pleuropneumoniae HS143 WT, HS143∆ngt and HS143∆agt, generation or reduction values were calculated by comparing the normalized values of the wild-type and mutant samples, using the statistical formulation for the threshold cycle (ΔΔCT) method. The threshold value of each gene was first normalized to the value of the constitutively expressed control gene glyA [30] (glyA primers: fwd: 5'-CAA GAC GAT GCA GCA GCT GTT TA-3', glyArev: 5'-CTG TGA TGC CGT AGA GGA CA-3').

2.10. Subcloning and heterologous expression of agt and ngt

The putative NGT operon was PCR amplified using the primers ngt-agtfwd: 5'-TTTTGAATTCGACGAGAAG TGAAGTCG-3' and ngt-agtrev: 5'-TTTTGTACCACCATGAG GCATAGCCGTATTTCT-3' using Accuprime Taq Hifi (Invitrogen) and the following cycling conditions: 94°C/30 s, followed by 24 cycles of 94°C/30 s, 53°C/30 s, 68°C/4 min and a final cycle of 68°C/5 min.

The amplicon was ligated into the vector pEXT20 using T4 DNA ligase (New England Biolabs, UK) following digestion of the plasmid and the PCR product with EcoRI and KpnI. Escherichia coli NEB10β (New England Biolabs, UK) was transformed with the ligation reaction generating the plasmid pJC78. Expression was induced by growing an E. coli colony in LB broth with ampicillin 100 μg ml⁻¹ until an OD₆₀₀ of 0.4 was reached. At that point 1 mM, IPTG was added, and the cultures were incubated at 37°C with shaking for a further 16 h. Expression of NGT and α6GlcT was monitored using SDS–PAGE, Coomassie staining and western blotting.

2.11. Glycosylation of AtaC by NGT and α6GlcT in Escherichia coli cells

Escherichia coli DH10β cells carrying pJC78 were transformed with the construct pMLBADAtaC1666–2428 [17], and cultured in LB broth with ampicillin 100 μg ml⁻¹, trimethoprim 20 μg ml⁻¹ at 37°C with shaking until an OD₆₀₀ of 0.4 was reached, followed by induction with 0.2% L-arabinose and 1 mM IPTG. After 16 h incubation, AtaC was purified. The bacterial cell pellet was isolated by centrifugation at 6000g for 10 min and lysed using a cell homogenizer (Starsted Fluidics Ltd. SPCH-10). Any intact cell debris was thereafter pelleted by centrifugation at 10 000g for 30 min before purification from the supernatant using an Ni–NTA (Qiagen, UK) gravity column (Thermo Scientific, USA).

Glycosylated product was analysed by SDS–PAGE and transferred onto a nitrocellulose membrane before being analysed by immunoblot using a mouse anti-His antibody (AbCam, UK) and an IRDye 680CW goat anti-mouse conjugate secondary antibody. Detection of fluorescent signal was carried out using a LI-COR imaging system.

2.12. Mutagenesis of the ngt locus

The cloned locus coding for NGT and α6GlcT was mutated using the QuickChange XL II site-directed mutagenesis kit (Agilent Technologies, CA) using the following primers. agt120a: 5'-CAAACAGAAGAAGCTTTTACATTATATTTTCTCAAGAACACCTTAAAGAGCC-3' and agtt120a: 5'-GCCTCTGTTGGTTGAAAATGCTTTAGGTTTATTTCA-3' (underlined nucleotides denotes the change).

The ngt gene was mutated using the following primers: ngt1321g_a132: 5'-GCCTATACGTCTTCAACACCATGCGC CTAAATCCGTATTT CTTAGAA-3' and ngt1321g_a132: 5'-TCTAAAGAATACCGGTTGTTTATGCGTCAT CTATACGCGGTTTCTTGT-3' (underlined nucleotides denotes the change). The following conditions were used: 95°C/60 s followed by 18 cycles of 95°C/50 s, 60°C/50 s, 68°C/8 min and a final 68°C/7 min cycle.

Following amplification, the PCR products were DpnI treated according to the manufacturer’s instructions and used to transform E. coli XL-10 Gold cells (New England Biolabs, UK).

2.13. Western blot analysis of glycosylated AtaC

Purified AtaC from E. coli DH10β was analysed by western blotting. Unglycosylated, fully glycosylated and monoglycosylated AtaC were analysed by dot blot by placing a 5 μl drop of a 1 mg ml⁻¹ solution of protein or dextran (dextran standard
MW 1000 from Leuconostoc, Sigma-Aldrich UK) onto a nitrdocellulose blotting membrane (Amersham Protran, GE Healthcare, Germany) and allowing to air dry before blocking the membrane by incubating with phosphate-buffered saline, 2% milk solution for 1 h at room temperature. The membrane was then probed using a 1 : 1000 dilution of a mouse monoclonal antibody raised specifically against a tetrasaccharide of α1–6 linked glucose (MS α-Dextran Clone DxI, Stem Cell Technologies, Canada). An IRDye 680CW goat anti-mouse antibody at a 1 : 10 000 dilution was used as the secondary antibody. The western blot images were visualized using a LI-COR imaging system.

2.14. Analysis of N-glycans released from AtaC

Glycans were released from 200 µg AtaC using the Ludger Liberate Hydrazinolysis kit, according to the manufacturer’s recommendations. The released glycans were fluorescently labelled with 2-aminobenzamide (2-AB) as described previously [31]. Excess labelling reagent was removed as follows: four discs of filter paper (Whatman) were soaked in 30% acetic acid, inserted into a 1 ml plastic syringe and washed sequentially with 2 × 1 ml acetic acid, 2 × 1 ml water, 2 × 1 ml acetonitrile (ACN) and finally 2 × 1 ml 95% ACN. The labelled glycans were diluted to 500 µl with 95% ACN and loaded onto the column. Of 500 µl 95% ACN was used to rinse the labelling tube and was added onto the column as well. The column was washed with 8 × 1 ml 95% ACN and glycans were finally eluted in 50 µl water twice. Elution fractions were pooled and passed through a 0.45 µm filter (Ultrafree-MC Durapore HV filter unit, Millipore) before analysis by normal-phase HPLC (Supelcosil LC-NH2 column, 80–20% ACN gradient over 90 min, fluorescence detection at 320 nm excitation and 420 nm emission wavelength). Glyko 2-AB glucose homopolymer standard (Prozyme) was used as a reference. The identity of the labelled glycans was confirmed by MALDI mass spectrometry analysis of a self-made tip column (75 µm filter (Ultrafree-MC Durapore HV filter unit, Millipore)) before analysis by normal-phase HPLC (Supelcosil LC-NH2 column, 80–20% ACN gradient over 90 min, fluorescence detection at 320 nm excitation and 420 nm emission wavelength). Glyko 2-AB glucose homopolymer standard (Prozyme) was used as a reference. The identity of the labelled glycans was confirmed by MALDI mass spectrometry. Samples were mixed 1 : 1 with dihydroxybenzonic acid matrix (15 mg ml⁻¹ in 75% ACN in water with 0.1% formic acid (FA)), and spotted onto a matrix-assisted laser desorption/ionization target plate. Data acquisition was performed manually on a Model 4800 Proteomics Analyser (Applied Biosystems, Framingham, MA) with an Nd : YAG laser, and 1000 shots were accumulated in the reflectron positive ion mode.

2.15. Nano-LC – ESI – MS/MS analysis of glycosylated AtaC

For structural analysis, 50 µg of AtaC was reduced, alkylated and digested with trypsin using the filter-aided sample preparation protocol [32]. Samples were analysed on a calibrated LTQ-Orbitrap Velos mass spectrometer (Thermo Fischer Scientific, Bremen, Germany) coupled to an Eksigent-Nano-HPLC system (Eksigent Technologies, Dublin (CA)). Peptides were reuspended in 2.5% ACN and 0.1% FA, and loaded on a self-made tip column (75 µm × 80 mm packed with reverse phase C18 material (AQ, 3 µm 200 A, Bischoff GmbH, Leonberg, Germany) and eluted with a flow rate of 200 nl per min by a gradient from 3% to 30% ACN, 0.1% FA in 22 min, 50% ACN, 0.1% FA in 25 min, 97% ACN, 0.1% FA in 27 min. One scan cycle comprised a full-scan MS survey spectrum, followed by up to 20 sequential collision-induced dissociation (CID) MS/MS on the most intense signals above a threshold of 1500. Full-scan MS spectra (400–2000 m/z) were acquired in the FT-Orbitrap at a resolution of 60 000 at 400 m/z, whereas CID MS/MS spectra were recorded in the linear ion trap. CID was performed with a target value of 1e4 in the linear trap, collision energy at 35 V, Q-value at 0.25 and activation time at 30 min. AGC target values were 5e5 for full FTMS scans and 1e4 for ion trap MSn scans. For all experiments, dynamic exclusion was used with one repeat count, 15 s repeat duration and 60 s exclusion duration.

2.16. Database analysis and identification of modified residues

MS and MS/MS data were processed into Mascot generic format files and searched against the Swissprot database (version 201402) through the Mascot engine (v. 2.2) with the consideration of carbamidomethylation at cysteine, oxidation at methionine and N-hexosylation at Asparagine. The monoisotopic masses of 2 + or more charged peptides were searched with a peptide tolerance of 10 ppm and an MS/MS tolerance of 0.6 Da for fragment ions. Only peptides with a maximum of two missed cleavage sites were allowed in database searches. Positive identification of hexosylated peptides was performed by manual inspection of spectra. Peptides modified with extended glycan chains were investigated manually, and their corresponding MS/MS spectra were annotated. Here, XCALIBUR v. 2.2 sp1.48 was used for data processing, and MS deconvolution was performed by XtractRaw file from Thermo Scientific.

2.17. Construction of acceptor protein JC1

Amino acid residues 23–163 of Cj0114 from C. jejuni NCTC 11168 were used as a scaffold to design a novel acceptor protein. The native signal sequence from residues 1 to 23 was removed along with the native tetrasaccharide domain encoded within residues 164–315. Twelve NAT glycosylation sequons were added at the C-terminus of the new protein, each separated by a proline and a glycine. Finally, a hexahistidine tag was added to the C-terminus to enable protein purification. This construct was DNA synthesized (Celtek Genes, USA) and subcloned into BamHI and SpilI digested expression vector pACYC184.

3. Results

3.1. NGT and α6GlcT are required for adhesion of Actinobacillus pleuropneumoniae HS143 to A549 cell lines

Within the genome of A. pleuropneumoniae strain HS143, we identified two ORFs, orthologues of apl_0104 (70% identity, BlastP) and apl_0443 (82% identity, BlastP) in the L20 genome [25], coding for autotransporter adhesins. In silico analysis revealed that these adhesins have 75 and 95 N-X-(S/T) sequons, respectively (PROGLYPROT). Naegeli et al. [17] carried out mass spectrometry analysis of A. pleuropneumoniae’s proteome for strain 4074 serotype 7, and the only
glycopepetides identified belonged to two autotransporter adhesins [17], making these two adhesins the only native substrates for NGT identified so far. In NTHi, deletion of the N-linked glycosylation system results in a significantly reduced adherence phenotype [15]. In order to investigate whether this was the case for *A. pleuropneumoniae*, adhesion of WT HS143, isogenic mutants HS143Δngt and HS143Δagt and its complements to A549 human adenocarcinoma lung epithelial cells was investigated.

*Actinobacillus pleuropneumoniae* strain HS143, the wild-type strain, was found to have a percentage of adherent cells of 8.55 ± 0.84, *n* = 78 to A549 cells after 3 h incubation (figure 2). In order to understand the role of the cytoplasmic NGT in this adherence phenotype, an in-frame deletion mutant of the *ngt* gene was generated in *A. pleuropneumoniae* and found to have a reduced percentage of adherent cells, 2.39 ± 0.25 (*n* = 78, *p* < 0.05), when compared with the wild-type. This phenotype was restored (11.05 ± 1.10, *n* = 30, *p* > 0.05) upon complementation with the *ngt* gene. Furthermore, when an α6GlcT deletion mutant in *A. pleuropneumoniae* was tested for adherence, it was observed that there was a decrease in adherence (2.85 ± 0.60, *p* < 0.05, *n* = 30) to the same level as the NGT mutant. However, complementation with the ORF coding for α6GlcT was unable to rescue this phenotype (figure 2). Gentamycin treatment of cells confirmed that the bacterial counts observed were due to adhering and not invading bacteria.

### 3.2. Agt is part of a conserved putative operon that includes *ngt* and *rimO*

Unlike NTHi, adjacent to the *A. pleuropneumoniae* N-linking transferase gene, *ngt*, the flanking genes do not encode an adhesin or a dedicated adhesin transporter (figure 1). Instead, we identified an ORF coding for a protein with amino acid similarity to 30S ribosomal protein S12 methylthiotransferase, *rimO*, upstream of *ngt*, and a second glycosyltransferase-encoding gene downstream of *ngt*. Analysis of all available *A. pleuropneumoniae* genomes demonstrated that the genetic arrangement of the locus was absolutely conserved in all published *A. pleuropneumoniae* genomes and over 180 sequenced isolates (J.T.B. 2016, personal communication). Reverse transcriptase-PCR (RT-PCR) was used to analyse the expression of this locus in serovar 15 *A. pleuropneumoniae* reference strain HS143. Primers were designed spanning intergenic regions between the three ORFs. Probe 1 tested if an mRNA transcript was generated between *rimO* and *ngt*, and probe 2 tested for the presence of an mRNA transcript between *ngt* and *agt*. The results suggest that all three genes form an operon (figure 3a). Further RT-PCR analysis showed that the promoter driving *rimO* expression was independent of the ORF immediately upstream (figure 3b).

Messenger RNA was extracted at different time points during the growth of *A. pleuropneumoniae* and cDNA was generated by RT-PCR using a probe designed within *ngt*. This showed that *ngt* was transcribed at all-time points tested (electronic supplementary material, figure S1).

### 3.3. Absolute quantification of *rimO*, *ngt* and *agt* by qPCR

In order to further validate the hypothesis that *rimO*, *ngt* and *agt* are co-transcribed, an absolute quantification qPCR was performed. The results were normalized by the absolute number of copies of *rimO* within each sample assuming *rimO* is the first ORF in the operon and therefore the closest to the putative promoter identified by bioinformatics analysis. A trend was observed in all four biological replicates (n = 12; 4 biologicals, 3 technical replicates) indicating a decrease in expression level from *rimO* to *agt* consistent with the genetic organization of the putative operon (*rimO* versus *ngt*, *p* < 0.05; *rimO* versus *agt*, *p* < 0.001; figure 4).

### 3.4. Reconstruction of the NGT glycosylation operon and its functional transfer and expression in Escherichia coli

Following on from the RT-PCR studies indicating that both *agt* and *ngt* were co-transcribed, we amplified by PCR the two ORFs as a single amplicon and cloned them into the IPTG-inducible expression vector pEXT20 [33], to generate the plasmid pJC78. When the ORFs encoding α6GlcT and *ngt* were co-expressed with a fragment of an autotransporter adhesin from *A. pleuropneumoniae* (AtAC), which is a natural acceptor [17], a reduction in protein migration on SDS–PAGE was observed, indicating an increase in molecular weight consistent with the addition of an oligosaccharide (figure 5). To further understand the *in vivo* glycosylation operon, individual mutations in *ngt* or *agt* were constructed within the plasmid pJC78. NGT activity was abolished by substituting the conserved lysine residue at position 441 by alanine (K441A) [18,20], whereas α6GlcT activity was abolished by the replacement of the leucine codon at amino acid position 7 with a stop codon (L7*). Schwarz et al. [19] indicated that *in vitro*, NGT and α6GlcT could assemble a glucose polymer between two and six residues on an acceptor peptide. We reasoned therefore that a commercially available antibody...
specific for α1–6 linked glucose tetrasaccharide (isomaltotetraose) may be able to detect and verify the nature of the polysaccharide generated by the cloned agt–ngt operon and of the knockouts. Ni–NTA purified proteins from the three construct combinations were tested for expression by dot blot analysis using an anti-dextran monoclonal antibody (mAb). This showed that a recognizable epitope could only be generated when NGT and α6GlcT were both functional (figure 5, top panel). SDS–PAGE and western blot analysis using an anti-HIS monoclonal antibody showed a smear visible above the point at which AtaC should migrate, but only when NGT and α6GlcT are both functional. This smearing was also detected using the anti-dextran monoclonal antibody (figure 5f, lane 1). AtaC glycosylated appears to migrate less than when detected by anti-HIS antibody, because the anti-dextran antibody will only recognize AtaC modified with four or more glucoses per site. Removing the function of NGT yielded an AtaC fragment that migrated to its unglycosylated location losing the epitope recognized by the anti-dextran mAb (figure 5e, lane 2). Finally, knocking out the function of α6GlcT reduced protein migration to a slightly higher level than that observed with NGT mutation alone (figure 5e, lane 3). This can be explained by glycosylation with a single hexose at multiple sites within the acceptor protein. Furthermore, this material was not recognized by anti-dextran mAb, suggesting that glycosylation

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**Figure 3.** (a) Transcriptional analysis of the *A. pleuropneumoniae* NGT locus. Lane 1: cDNA as template; lane 2: RNA as template; lane 3: *A. pleuropneumoniae* HS143 genomic DNA positive control; lane 4: negative PCR control (no template). (b) Transcriptional analysis of the region upstream of ngt. Lane 1: cDNA as template; lane 2: genomic DNA positive control; lane 3: PCR control (RNA as template).

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**Figure 4.** Absolute quantification of rimO, ngt and agt in *A. pleuropneumoniae* strain HS143 expressed in fold change. Horizontal bars indicate pairs of columns that are significantly different when compared with each other. Asterisk indicates $p < 0.05$ ($n = 12$).

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**Figure 5.** Glycosylation analysis of AtaC1866–2428 alongside NGT and α6GlcT by anti-dextran and anti-HIS western blots. Top panel: anti-dextran dot blot; (a) dextran; (b) AtaC with NGT K441A and functional α6GlcT; (c) AtaC with functional NGT and α6GlcT; (d) AtaC with functional NGT only. Bottom panel: anti-HIS and anti-dextran Western blots (e, f, respectively); lane 1, AtaC with functional NGT and α6GlcT; lane 2, AtaC with NGT K441A and functional α6GlcT; lane 3, AtaC with functional NGT only.
with an asterisk. (A) A glucose homopolymer (dextran) ladder serves as a reference for retention time (red). The peak originating from excess 2-AB label (rt = 4.1 min) is marked with an asterisk. (b,c) MALDI mass spectrometry analysis of the same glycan sample confirms the identity of the observed glycan chains as a hexose polymer.

Figure 6  (a) NP-HPLC analysis of 2-AB labelled glycans released from purified AtaC which was co-expressed with NGT and αGlcT in E. coli DH10B (black). A glucose homopolymer (dextran) ladder serves as a reference for retention time (red). The peak originating from excess 2-AB label (rt = 4.1 min) is marked with an asterisk. (b,c) MALDI mass spectrometry analysis of the same glycan sample confirms the identity of the observed glycan chains as a hexose polymer.

had occurred, but that no polymer had been generated (figure 5f, lane 3).

3.5. Confirmation of hexose build-up on AtaC

In order to confirm the identity of the observed post-translational modification of AtaC, the glycans from purified protein were released by hydrazinolysis, fluorescently labelled with 2-aminobenzamide (2-AB) and analysed by normal-phase HPLC (figure 6a).

With a labelled dextran ladder used as a reference, this analysis revealed the presence of glycan chains of varying lengths (1–27 monosaccharide units). This was further confirmed by MALDI-MS analysis. Peaks differing in mass by 162 Da suggested potential for glucose or galactose attachment (figure 6b,c). Therefore, the results from both methods were in agreement showing a hexose polymer ranging up to at least 20 units.

To confirm that particular sites on the protein were modified with these elongated glycans, LC–ESI–MS/MS analysis of glycosylated AtaC was performed (figure 7 and table 1). This showed that previously identified glycosylation sites were occupied [17]. In total, 15 asparagine (Asn) residues were identified as being modified with glycan chains of variable length. For example, glycopeptide GNLSTAADVTDK could be detected modified with an N-linked glycan consisting of 1–29 hexose units (table 1). On other sites, only short glycan chains could be detected, whereas one peptide (NISTVVK) could only be detected as being modified with glycan chains of more than 14 hexoses. These results are summarized in table 1 and confirm western blot evidence that co-expression of NGT and αGlcT leads to the formation of Asn-linked, linear hexose chains of up to 29 units in length.

3.6. The ngt/agt operon can be used to modify alternative substrates with dextran

Following assembly of plasmid pJC78, we began testing if the ngt/agt operon could be used to make N-linked glucose polymers on non-native substrate proteins in a similar manner to NGT alone [17]. We selected Cj0114 from the α-proteobacterium Campylobacter jejuni as a scaffold for designing a new acceptor protein. The native Cj0114 tetra-ticopeptide domain was removed to reduce protein toxicity and simplify purification. At the C-terminus of the protein, the 12 added NAT glycosylation sequons were followed by a hexa-histidine tag to enable protein purification. The new protein, named JC1, was constitutively expressed from the plasmid pJC1. Combining the plasmids pJC1 and pJC78 generated an epitope that could be recognized by the anti-dextran mouse mAb; this disappeared upon knocking out the function of ngt or agt (figure 8). The marginally different sizes in the anti-His and anti-dextran western blot are due to the recognition epitope for the anti-dextran antibody, where only highly polymerized proteins are detected (acceptors modified with four or more glucose residues). These findings indicated that NGT and αGlcT can be made to target any protein.
4. Discussion

Novel bacterial glycosylation systems are regularly being discovered as glycan analyses methodologies improve [34–37]. The functions of these glycosylation systems are yet to be fully appreciated, but it is now apparent that glycosylation is a feature common to most bacteria.

In this study, we report the investigation of a cytoplasmic glycosylation system in a member of the Pasteurellaceae family, *A. pleuropneumoniae*. Our results demonstrate that despite similarities between NGT and its orthologue, HMW1C, in NTHi, the system described here is unique. The *A. pleuropneumoniae* N-linking locus consists of two co-transcribed glycosyltransferases (*ngt* and *agt*) with no associated adhesin or

Figure 7. Deconvoluted MS spectra show that peptide GNLSTAADVTDK from AtaC is modified with a hexose polymer ranging from 1 to 28 units. (b) MS/MS spectrum of *m/z* 758.3497 (+2), corresponding to GNLSTAADVTDK modified with two hexoses, showed continuous fragmentation ions, which confirm the peptide identity. The hash tag marks doubly charged ion with neutral loss of hexose from precursor ion. y' indicates the y ion without hexoses.
transporter. Another significant difference between the *A. pleuropneumoniae* system and that of NTHi is that the promoter for *rimO*, upstream of *ngt*, appears to be responsible for driving transcription of *ngt* and *agt* (figure 3). Recent studies have shown that in *Aggregabacter aphrophilus* and *Haemophilus ducreyi* [16,21] *hmwC* is also located downstream of *rimO*, although a transcriptional link has yet to be proven [21]. In *E. coli*, RimO is an enzyme that catalyses the methylation of ribosomal subunit S12 at the universally conserved D88 residue. Furthermore, it has been shown that knocking out *rimO* in *E. coli* leads to a growth defect [38,39]. The significance of RimO has also been reported in *Thermus thermophilus*, where residue D88 cannot be mutated [40], leading to the conclusion that although methylation is not essential in every organism, RimO clearly plays an important role in maintaining bacterial fitness [39]. Our tests indicate that the *A. pleuropneumoniae rimO* promoter is active at every time point tested, suggesting that *ngt* and *agt* are constitutively expressed (electronic supplementary material, figure S1), and therefore the cytoplasmic N-linking glycosylation system is always available to modify substrate proteins. Furthermore, qPCR analysis of the locus indicated transcriptional levels consistent with an operonic structure, where the highest level of transcription detected was of *rimO*, followed by *ngt* and *agt*, respectively (figure 4). This is in agreement with the findings reported by Lim *et al.* [41] whereby the expression level of the genes proximal to the promoter was greater than the ones farthest from it. Bioinformatic analysis of the DNA sequence surrounding the putative *rimO*/*ngt*/*agt* operon identifies a transcriptional promoter just upstream of *rimO* and a Rho-independent terminator downstream of *agt* (electronic supplementary material, figure S2) [42]. Nevertheless, to gain further insights into the regulation of the locus, other approaches such as RNAseq could be carried out. Furthermore, *in silico* analysis of all publically available genomes and over 180 others (J.T.B. 2016, personal communication) indicates that the gene order is absolutely conserved (data not shown).

In this work, we also demonstrate that NGT plays an important biological role in the ability of *A. pleuropneumoniae* to adhere to A549 human adenocarcinoma lung epithelial cells, which, although from human origin, are from biologically relevant tissue. The rationale for using A549 cells, instead of St Jude Porcine lung (SJPL) cells, which have been widely used to assess *A. pleuropneumoniae* adhesion [4,43,44], was that the SJPL cell line was found to be misclassified, and is simian in origin [45]. In order to draw absolute conclusions regarding the role of this N-linked glycosylation system in aiding *A. pleuropneumoniae* pathogenesis in the pig, the adhesion assay data that obtained in this study could be extended to investigate other tissues such as *ex vivo* organ cultures [46] possibly primary cell cultures from pig lung epithelial cells.

Similarly to our study, a significant reduction in adhesion was reported for *E. coli* expressing the cloned *hmw1* locus from NTHi when the function of HMW1C (the NGT orthologue) was removed [15]. The *hmw1*/hmw2 loci and the *ngt* operon differ in that the NTHI loci encode an adhesin and an adhesin transporter alongside *hmwC*, whereas the *A. pleuropneumoniae* locus encodes an α6GlcT polymerizing glucosyltransferase (figure 1). Surprisingly, knocking out the function of the
α6GlcT transferase also resulted in a significant reduction in adherence comparable to that detected when NGT activity was abolished. Plasmid-based complementation with agt only (figure 2) and ngt/agt (data not shown) proved insufficient to rescue the adhesion phenotype seen with the wild-type. Our failed attempts to rescue the phenotype suggest a need for fine transcriptional control of agt levels in the bacterium. It is possible that, when ngt is being expressed in the chromosome and agt is on a plasmid, over-glycosylation of target sites occurs, resulting in incorrect adhesin structural conformation. Potentially, this incorrect folding could prevent surface presentation or adhesin function.

Some of the best-studied examples of glycosylated autotransporter adhesins are O-linked, and found in E. coli. O-linked glycosyltransferases can glycosylate TibA, Ag43 and AIDA-I in their passenger domains [47–49]. Bioinformatic analysis of the autotransporter adhesin used in our study, AtaC from A. pleuropneumoniae AP76 (GenBank accession number ACE61172.1), indicated the presence of 72 NX-(S/T) sequons. The majority of these (59/72) are localized in the passenger domain of the adhesin, further demonstrating the similarities with the O-linked counterparts. Whether glycosylation is required, so that the adhesin assumes the correct conformation and is not degraded as observed in AIDA-I, or if it is so that the adhesin can adopt a conformation suited for adhesion as described in TibA [50], remains to be determined even in these well-studied proteins.

Our demonstration of glucose polymer assembly within E. coli cells, when the ngt/agt locus is overexpressed alongside an acceptor protein (figure 5), led us to investigate if this glycan could be detected on the surface of A. pleuropneumoniae. Immunofluorescence studies using an anti-dextran antibody that recognizes isomaltooltetrose as a minimum epitope failed to detect any signal, even in permeabilized cells (data not shown). This suggests that although the capability exists to form α-1,6 glucose chains greater than four subunits heterologously, in A. pleuropneumoniae the necessary epitope for detection with anti-dextran antibody does not appear to be formed. Analysis of the glycosylated peptides generated within E. coli, as determined by peak quantification of the HPLC chromatogram (figure 6), revealed a steady decrease in abundance of the oligosaccharide with increasing chain length. It was however noteworthy that the peak corresponding to Glc2-2AB (retention time: 8.8 min) was considerably smaller than the peaks corresponding to Glc2-2AB and Glc2-2AB, indicating that the addition of the first α-1,6-linked glucose might be considerably slower than the subsequent transfer reactions. This suggests that the first α6GlcT-catalysed reaction is, in fact, the rate-limiting step in the biosynthesis of these extended N-glycan chains.

In a review of the HMWC literature, we found instances where adhesin glycopeptides with dihexose modifications have been reported, in the absence of a co-localized ORF-like agt [15,51]. Rempe et al. [21] report dihexose modifications on four glycopeptides belonging to the autotransporter adhesin of K. kingae. These raise several possibilities; the first is that the reported glycopeptides actually contain two individual hexose attachments and not two hexoses together. Second, it may be possible that the HmwC from K. kingae is able to catalyse N-linked attachment and subsequent polymerization. Third, one cannot rule out there may be another glycosyltransferase in the genome that is enabling dihexose assembly.

A review of the NGT-specific literature reveals an interesting disparity in the function of this enzyme when tested in vitro and in vivo. Choi et al. [51] reported that in vitro, NGT is capable of forming dihexoses. However, this study indicates that α6GlcT is essential for glycosidic bond formation and extension of the glucose polymer in vivo. This is in agreement with a previous study by Naegeli et al., which failed to detect any polymerization when NGT alone was expressed in E. coli to glycosylate an acceptor protein [17].

Our finding that α6GlcT function is necessary to maintain adhesion in A. pleuropneumoniae indicates that this enzyme must be extending glucose residues at some sites within the autotransporter adhesins. However, by transferring the N-linking glycosylation locus into E. coli, we showed that α6GlcT and NGT are unable to fully complement each other’s functions. Our study also provides further evidence that ‘orphan’ HMWC family of enzymes that have not evolved to be co-localized with their target substrate continue to modify proteins involved in adhesion. It is noteworthy that every bacterial species reported thus far with this genetic arrangement uses the glycosylation system to target autotransporter adhesins [16,21]. Glycosylation has been linked to protection from proteolytic degradation, correct protein folding and correct transport to the surface, all of which would have an effect on cell adhesion. Further studies are ongoing to ascertain the level of interaction between α6GlcT/NGT and the target protein(s).

By demonstrating how to harness the ngt/agt operon, we have shown potential for glycoengineering applications, including the generation of N-linked glucose-based conjugate vaccines against A. pleuropneumoniae. The genetic conservation of the ngt operon in A. pleuropneumoniae would favour the development of a glycoconjugate vaccine against multiple A. pleuropneumoniae serovars. Other potential applications include the development of dextran-based conjugates that may be useful against bacteria such as Helicobacter pylori [52]. Recently, such conjugates have been shown to be immunogenic, and post-immune sera from rabbits vaccinated with dextran-based conjugates exhibited activity against strains of H. pylori that contain α(1–6) glucose as part of their LPS [52].

The field of bacterial glycobiology is burgeoning and investigations into various glycosylation systems, such as the NGT/α6GlcT system reported here, help to understand their functional roles. Our results demonstrate the importance of genetic and phenotypic screens for investigating glycosylation systems, and that this data can directly benefit bacterial glycoengineering.

Data accessibility. The datasets supporting this article have been uploaded as part of the electronic supplementary material.

Authors’ contribution. V.S.T. and J.C. wrote the manuscript, carried out most investigations into various glycosylation systems, and that this data can directly benefit bacterial glycoengineering.

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