Title page

Identification and characterisation of O-linked glycans in cervical mucus as biomarkers of sperm transport: a novel sheep model

Key words: Cervical mucus/ fertility/ O-glycans/ UPLC

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Supplementary Data

Supplementary Figures. Fig. S1. Representative FLR-HILIC-UPLC chromatograms of O-glycans released by microwave-assisted non-reductive β-elimination in follicular phase sheep cervical mucin (six European ewe breeds ordered top-down from lowest to highest fertility). Blank in black. Integrated peaks annotated 1-51. To be referenced against Table S2. Fig. S2. Nested dot plots for all significant glycan peak changes across ewe breeds at the natural cycle.
**Fig. S3.** PCA plots comparing ewe breeds at the synchronized cycle to glycan peaks. **Fig. S4.** Nested dot plots for all significant glycan peak differences across ewe breeds at the synchronized cycle. **Fig. S4 (cont).** Nested dot plots for all significant glycan peak changes across ewe breeds at the synchronized cycle. **Fig. S5.** PCA plots comparing ewe breeds at the synchronized cycle to significant features. **Fig. S6.** Mucus viscosity-glycan correlation. Only significant correlations are shown ($P < 0.05$) with correlation coefficient ($\rho = \text{rho value}$). **Fig. S7.** Mucus viscosity-feature correlation. Only significant correlations are shown ($P < 0.05$) with correlation coefficient ($\rho = \text{rho value}$). **Fig. S8.** Workflow for the comprehensive analysis of $O$-glycans from cervical mucus combining FLR-HILIC-UPLC, exoglycosidase digestions and HILIC-UPLC-ESI-MS. (1) Purified cervical mucin samples were analysed per duplicates by HILIC-UPLC for statistical evaluation. (2) Each undigested sample was treated with a panel of exoglycosidase enzymes followed by another round of HILIC-UPLC analysis. The objective was to determine the $O$-glycan sequence and linkage. (3) Undigested samples were analysed by HILIC-UPLC-ESI-MS to confirm exoglycosidase findings thus the presence of these $O$-glycan structures. **Supplementary tables. Table S1.** Reproducibility study of ovine cervical mucin (OCM) sample number 37 outlining coefficient of variation (CV) across seven technical replicate release. Red indicates a CV above 15%. **Table SII.** Glycan list of 2AB labelled HILIC separated $O$-glycans identified in sheep cervical mucin with GU values and peak number. To be referenced against Figure 1 and Figure S1. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SIII.** MS data outlining m/z values as well as relative intensities and charge state; and FLR-HILIC-UPLC data outlining GU values and relative area. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SIV.** Exoglycosidase digestion panel for Suffolk pool. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SV.** Exoglycosidase digestion panel for Belclare pool. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SVI.** Exoglycosidase digestion panel for Ile de France pool. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SVII.** Exoglycosidase digestion panel for Romanov pool. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SVIII.** Exoglycosidase digestion panel for Fur pool. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SIX.** Exoglycosidase digestion panel for NWS pool. – REFER TO SUPPLEMENTARY SPREADSHEET FILE. **Table SX.** $P$-values from feature analysis comparing natural and synchronized cycles of each ewe breed. Green indicates a significant increase of that particular core in the natural cycle.
Abstract

Cervical mucus plays an important role in female fertility, since it allows the entry of motile and morphological normal sperm while preventing the ascent of pathogens from the vagina. The function of cervical mucus is critically linked to its rheological properties that are in turn dictated by \( O \)-glycosylated proteins, called mucins. We aimed to characterize the \( O \)-glycan composition in the cervical mucus of six European ewe breeds with known differences in pregnancy rates following cervical/vaginal artificial insemination with frozen-thawed semen, which are due to reported differences in cervical sperm transport. These were Suffolk (low fertility) and Belclare (medium fertility) in Ireland, Ile de France and Romanov (both with medium fertility) in France and Norwegian White Sheep (NWS) and Fur (both with high fertility) in Norway (n = 28 to 30 ewes/breed). We identified 124 \( O \)-glycans, from which 51 were the major glycans with core 2 and fucosylated glycans as the most common structures. The use of exogenous hormones for synchronization did not affect the \( O \)-glycan composition in both high fertility ewe breeds, but it did in the other four ewe breeds. There was a higher abundance of the sulfated glycan (Galβ1-3[SO3-GlcNAcβ1-6]GalNAc), fucosylated glycan (GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc) and core 4 glycan (GlcNAcβ1-3(GlcNAcβ1-6)GalNAc) in the low fertility Suffolk breed compared with NWS (high fertility). In addition, core 4 glycans were negatively correlated with mucus viscosity. This novel study has identified \( O \)-glycans that are important for cervical sperm transport and could have applications across a range of species including human.
Introduction

In vaginal depositors such as humans and sheep, hundreds of millions to billions of spermatozoa are deposited into the vagina with a small percentage of these entering the cervix and uterus while, only a few hundred spermatozoa are thought to arrive at the site of fertilization in the ampulla. During this long and tortuous journey, spermatozoa encounter a number of physiological barriers, the first of which is the cervix. Cervical goblet epithelial cells secrete large quantities of cervical mucus, which is a viscoelastic fluid that allows the entry of sperm to the upper genital tract while also preventing ascending pathogens from the vagina. In the lead up to ovulation when levels of oestrogen are high, mucus production and hydration increase while mucus viscosity is decreased (Gipson et al. 2001). This is essential to allow motile and morphologically normal sperm reach the site of fertilization (Katz et al. 1980, Ragni et al. 1985). Therefore, sperm migration through the cervix is dependent on the cervical mucus properties, including mucus volume, viscosity and biochemical composition.

The key component of cervical mucus are mucins, high molecular weight glycoproteins that exhibit a complex molecular structure (Lagow et al. 1999). Mucins comprise hydrophobic, partially folded termini and a polyanionic, highly glycosylated, and elongated protein core (Corfield 2015). O-glycans constitute the main component of mucins and confer a hydrophilic characteristic by binding water through hydrogen bonds, which increases mucus hydration (Marczynski et al. 2020). Proteomic analysis has identified differences in O-glycosylation across the phases of the menstrual cycle (Andersch-Bjorkman et al. 2007), where in humans around the time of ovulation there were more neutral fucosylated structures than in the pre- or postovulatory phases. Mucin gene expression has also been examined, showing a peak of mucin production around the time of ovulation, which also promotes mucus hydration (Gipson 2001) and sperm migration (Ma et al. 2016). Mucus structure also changes during the oestrous cycle as pre-ovulatory mucus is arranged in a dense filamentous structure like a net of interconnected fibres while ovulatory mucus consists of dispersed floating globules of aggregated mucin molecules, which allow sperm migration through the mucus mesh around the time of ovulation (Brunelli et al. 2007, Lai et al. 2009). In addition, the sperm glycocalyx is rich in glycans with sialic acids as terminal monosaccharides, which typically interact with
terminal sugars of the cervical mucins. For example, N-acetyl glucosamine on the sperm surface showed high affinity for sialic acid of the cervical mucus (Ma et al. 2016, Tecle and Gagneux 2015). Therefore, O-glycans in the cervical mucus are important modulators of not only mucus viscosity but also mediate biochemical interactions with sperm.

The ewe is an excellent model for investigating how spermatozoa are selected in the female reproductive tract as semen is deposited vaginally and cervical artificial insemination (AI) is limited due to poor pregnancy rates when frozen-thawed semen is used. Worldwide, pregnancy rates rarely exceed 30% when frozen-thawed semen is used in conjunction with cervical AI. However, Norway is the exception to this, since they routinely achieve pregnancy rates of 60-70% with vaginal (shot-in-the-dark) AI to a natural oestrous (Paulenz et al. 2005). Previous studies by our group have demonstrated that this is due to the breed of the ewe used in Norway and specifically the inability of frozen-thawed sperm to traverse the cervix of some ewe breeds (Donovan et al. 2004, Fair et al. 2005). These ewe breed differences are not due to the gross anatomy of the cervix, the amount of mucus produced or its viscosity (Abril-Parreno et al. 2020) or differences in the peri-ovulatory endocrine profiles between breeds (Fair et al. 2007, Fair et al. 2019). In a recent study we found fewer sperm in the cervical crypts of Suffolk ewes (low fertility) compared with Belclare ewes (high fertility), which was associated with higher sialic acid content in the cervix of Suffolk ewes compared with Belclare ewes (Richardson et al. 2019). This suggests that large mucin O-glycans may cause the attachment of frozen-thawed sperm to the cervical mucus, thus impeding sperm migration through the cervix in some ewe breeds.

Therefore, the objective of this study was to characterise the O-glycans in cervical mucus around the time of ovulation of both a synchronized and natural cycle in six European ewe breeds known to have differences in pregnancy rates following cervical AI with frozen-thawed semen. Detailed and quantitative comparisons of O-glycan structures between ewe breeds with known differences in sperm transport may help to identify novel biomarkers of sperm transport, which is applicable across a wide range of species, including human.

Results

Most prevalent O-glycans in follicular phase cervical mucin

Representative chromatograms of each ewe breed are shown in Figure 1, outlining the main glycans identified across 51 major integrated peaks (full peak annotation in Supplementary Figure 1). From those, the five glycans with highest average area included: GlcAβ1-3/4galβ1-
3[SO3-Neu5Gcα2-6]GalNAc (peak 36), core 2 glycan Galα1-3Galβ1-3Galα1-3GlcNAcβ1-6]GalNAc (peak 35), core 2 glycan Galβ1-3[Galβ1-4GlcNAcβ1-6]GalNAc (peak 24), core 1 glycan Galβ1-3GalNAc (peak 4), core 2 fucosylated and sulfated glycan GlcAβ1-3/4(Fucα1-2galβ1-3)[SO3-Neu5Gcα2-6]GalNAc (peak 37) (from highest to lowest average area under the curve, respectively). Peeling peaks were not included in this ranking as they are degradation products of the chemical method to release the O-glycans.

Combining FLR-HILIC-UPLC, exoglycosidase digestions and ESI-MS for assignment confirmation, our structural analyses identified 124 O-glycans in total across a range of mucin-type cores (Supplementary Table II). Core 2 glycans were the most common structures with a 45.2 % of the total, followed by core 1 glycans (21.8 %), core 6 (10.5 %), core 4 (6.5 %) and core 3 (5.6 %), graphical representations of each core type are shown in Figure 2. Of these structures, there were 56 (37.8 %) fucosylated glycans, 37 (25 %) sialylated (19 Neu5Ac type and 18 Neu5Gc type), 22 (14.9 %) sulfated, 21 (14.2 %) glycans contained terminal α-galactose epitopes (with an additional three capped by sialic acid). In addition, 73 (49.3 %) O-glycans contained a terminal galactose epitope (both α- and β-linked) and five (3.4 %) structures contained an uncommon glucuronic acid terminal epitope. A total of 21 (14.2 %) peeling products were additionally identified (full list including MS confirmation in Supplementary Table III). Exoglycosidase arrays used to identify these structures are presented in Supplementary Tables IV-IX. Additionally, released N-glycans were analysed using FLR-HILIC-UPLC however, we did not identify N-glycan peaks.

Differences in cervical mucin O-glycan content collected at the follicular phase between a natural and a synchronized oestrous cycle

Individual glycan peaks were compared across the six ewe breeds at both the natural and synchronized oestrous cycles to identify significant cycle-related differences. We identified a ewe breed by type of cycle interaction (p < 0.05; Table I) represented by differences between synchronization and natural cycle for some ewe breeds but not others. In the Suffolk breed (low fertility), type 3 H-antigen (Fucα1-2Galβ1-3)GalNAc (peak 8) and glucuronyl-Tn-antigen GlcAβ1-3GalNAc (peak 14) were higher in the natural cycle compared to the synchronized cycle (p < 0.05). In the medium fertility ewe breeds, Belclare had four glycans with higher expression at the natural than at the synchronized cycle (Neu5Accα2-3GalNAc and GlcNAcβ1-3Galβ1-3GalNAc (both in peak 11) 11; Galα1-3Galβ1-3GalNAc (peak 13) and a fucosylated glycan (Galβ1-3GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc (peak 26)) (p < 0.05). In Ile de France,
core 1 T-antigen Galβ1-3GalNAc (peak 4), and Galβ1-3GlcNAc (peak 5) were increased at the natural cycle ($p < 0.05$). However, a fucosylated glycan GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc (peak 16) was decreased at the natural compared to the synchronized cycle ($p < 0.05$). In Romanov ewes, T-antigen (peak 4) was increased at the natural cycle ($p < 0.05$) while sialyl-Tn antigen (Neu5Aca2-3GalNAc) and GlcNAcβ1-3Galβ1-3GalNAc (both in peak 11) were decreased ($p < 0.05$). There were no differences in the two Norwegian ewe breeds between natural and synchronized cycles ($p > 0.05$).

Variation in the glycan profile between ewe breeds under the effect of synchronization were further categorized by feature (total fucosylation, total $N$-Acetylneuraminic sialic acid (Neu5Ac), $N$-Glycolyl sialic acid (Neu5Gc), sulfation, glucuronylation, type of core, and peeling) to determine the significant trends. The only feature differences in this analysis was an overall upregulation of total core 1 structures in Belclare ewe breed at the natural cycle ($p < 0.001$; Supplementary Table X), arising from upregulated Neu5Aca2-3GalNAc and GlcNAcβ1-3Galβ1-3GalNAc ($p < 0.05$; peak 11), Galα1-3Galβ1-3GalNAc ($p < 0.05$; peak 13), and Galβ1-3GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc ($p < 0.05$; peak 26).

**Ewe breed differences in cervical mucin $O$-glycans collected at the follicular phase of a natural oestrous cycle**

Individual glycan peaks and glycan features of six ewe breeds at the natural cycle are presented in Supplementary Tables XI and XII, respectively. Regarding individual structures, glycan peaks 1, 4, 10, 12, 15, 16, 17, 19, 20, 21, 22, 29 and 44 were affected by ewe breed ($p < 0.05$; Supplementary Figure 2). Of these, peak 12 (GlcNAcβ1-3[GlcNAcβ1-6]GalNAc) and peak 15 (Galβ1-3[SO3-GlcNAcβ1-6]GalNAc) had higher abundance in Suffolk (low fertility) compared with NWS (highest fertility; $p < 0.05$; Figure 3). Peak 19 (Neu5Aca2-3Galβ1-3GalNAc) also had higher abundance in Suffolk compared with Fur ewes (high fertility; $p < 0.05$). In addition, peak 29 (Neu5Aca2-3Galβ1-3GlcNAcβ1-3Gal) presented lower abundance in Beclare (medium fertility) compared with the Romanov, Fur and NWS ($p < 0.05$). Out of the features assessed, only core 6 glycans had increased expression in Suffolk ($p < 0.001$), Beclare ($p < 0.05$) and Fur ($p < 0.05$) when compared against Ile de France (medium fertility) which presented the lowest abundance of core 6 structures (Figure 4). This was derived only from the expression of a sulfated and fucosylated glycan (Galβ1-3(Fucα1-6-SO3-GlcNAcβ1-6)GalNAc) peak 20, which showed higher variation within breed in Suffolk and Beclare compared with the other ewe breeds.
Ewe breed differences in cervical mucins O-glycans collected at the follicular phase of a synchronized oestrous cycle

In the synchronized cycle, glycan peaks 1, 6, 7, 8, 9, 10, 11, 12, 14, 16, 17, 18, 19, 25, 26, 28, 30, 35, 36, 39, 42, 43 and 45 were altered by ewe breed (p < 0.05; Supplementary Figure 4 and Supplementary Table XIII). Of these glycan peaks, peak 12 (GlcNAcβ1-3[GlcNAcβ1-6]GalNAc) and peak 16 (GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc) were increased in Suffolk ewes (low fertility) compared with NWS (p < 0.05; Figure 5). Peak 18 (Neu5Gca2-6GalNAc) had lower abundance in Belclare (medium fertility) compared with Ile de France (medium fertility and Fur ewes (high fertility) (p < 0.05). Peak 19 (Neu5Acα2-3Galβ1-3GalNAc) also presented lower abundance in Belclare compared with Ile de France, Romanov and Fur ewes (p < 0.05). In addition, peak 25 (GlcAβ1-3[SO3-Neu5Gca2-6]GalNAc) had lower abundance in Belclare compared with NWS (p < 0.05), as well as Romanov (medium fertility) had lower abundance compare with Fur and NWS (p < 0.05). However, the abundance of this glycan was higher in Suffolk than in Romanov ewes (p < 0.05).

There was a ewe breed by type of cycle interaction in fucosylation, glucuronylation, acetylsialylation, core 1 and core 4 (p < 0.05), represented by more feature differences between ewe breeds at the synchronized compared to the natural oestrous cycle (Supplementary Table XIV). At a synchronized cycle, fucosylation was increased in Belclare compared with Suffolk (p < 0.05). Both Suffolk and NWS had increased glucuronylated structures compared to the Romanov (p < 0.05). Neu5Ac structures were increased in Romanov compared with the Belclare (p < 0.05). Core 1 glycans were increased in Ile de France (p < 0.05) and Romanov (p < 0.05) compared with Belclare. Core 4 glycans were increased in Suffolk compared with Belclare (p < 0.05). Dot plots of all significant feature differences are shown in Figure 6. Principal Component Analysis (PCA) was used to compare each breed with the glycans associated with a particular feature, where there were no significant differences (Supplementary Figure 5). However, clear separation was identified between Suffolk and Belclare for core 4 glycans (peaks 12, 41 and 50; Figure 6). In addition, clear separation on PCA plots was also observed between the low fertility Suffolk and NWS (highest fertility). However, this separation was not significant (Figure 6; p > 0.05).

Relationship between mucus viscosity and O-glycans

Mucus viscosity was correlated against relative feature and glycan abundance without including peeling peaks (Supplementary Figure 6 and 7). Fucosylation had a positive weak
correlation with viscosity ($\rho = 0.27$; $p < 0.05$) while core 4 abundance and glucuronation had a negative weak correlation ($\rho = -0.26$; $p < 0.05$). Glycan peaks 11 (Neu5Acα2-3GalNAc and GlcNAcβ1-3Galβ1-3GalNAc), 24 (Galβ1-3[Galβ1-4GlcNAcβ1-6]GalNAc), 35 (Galα1-3Galβ1-3[Galα1-3GlcNAcβ1-6]GalNAc), 36 (GlcAβ1-3/4galβ1-3[SO3-Neu5Gcα2-6]GalNAc) and 41 (Galβ1-4GlcNAcβ1-3[Galβ1-4(Fucα1-3GlcNAcβ1-6)]GalNAc) were all negatively correlated to mucus viscosity ($\rho = -0.30, -0.31, -0.29, -0.33$ and -0.25, respectively; $p < 0.05$).

**Discussion**

This is the first published study that has characterised the $O$-glycome of sheep cervical mucus. We did this using a novel European model composed of six ewe breeds with known differences in pregnancy rates and cervical sperm transport following cervical AI with frozen-thawed semen. The overall aim was to identify $O$-glycans as biomarkers of cervical sperm transport. No $N$-glycans were detected but we identified significant variations in the profile of $O$-glycans between high and low fertility ewe breeds and between a naturally occurring and hormonally synchronized oestrous in some ewe breeds. Results indicate that a core 4 glycan (GlcNAcβ1-3[GlcNAcβ1-6]GalNAc) was related with impaired sperm transport since there was higher abundance in the low fertility Suffolk breed compared to NWS (highest fertility) at both a natural and a synchronized cycle. In addition, a core 2 sulfated glycan (Galβ1-3[SO3-GlcNAcβ1-6]GalNAc) and a fucosylated glycan (GlcNAcβ1-3[Fucα1-2Galβ1-3]GalNAc) were also increased in Suffolk compared to NWS at the natural and synchronized cycle, respectively.

The first major challenge of this study was to optimize the method developed by Wilkinson et al. (2021) as it is the first time that this method is used for the release of $O$-glycans from cervical mucin samples. Here we used a new microwave assisted $\beta$-elimination method, which reduces reaction time in comparison with other chemical methods (reductive $\beta$-elimination and nonreductive $\beta$-elimination) and also provides comparable yields of released $O$-glycans minimising the peeling products as reviewed by Wilkinson and Saldova (2020).

By combining FLR-HILIC-UPLC, exoglycosidase digestions and ESI-MS we identified 124 $O$-glycans across a range of mucin-type cores from which 51 were the major $O$-glycans of the cervical mucins. The majority of glycans were core 2 (45%) followed by core 1 (22%). Other cores were also identified, for example, core 6 (11%), core 4 (6%) and core 3 (5.6%). It is well known, that core 1 glycans are ubiquitously expressed. In contrast, core 2 glycans are more cell-type specific and they act as a scaffold for the production of selectin ligands that act in
regulating inflammation. Relatively few tissues show high levels of core 3 and core 4, except for the gastrointestinal, respiratory tracts and salivary glands, where mucin production is normally high. Thus, core 1 to 4 glycans represent the most common O-glycan structures produced \textit{in vivo}, with core 2 subtype having the highest abundance as we identified in the present study. Other core structures, such as core 6 glycan also detected in our study, has been previously found on mucins from human embryonic gut and from ovarian cysts (Varki and Schauer 2009).

Our results are in broad agreement with a study of peri-ovulatory cervical mucus of women which showed T-antigen (Core 1 glycan type) having the highest levels around the time of ovulation (Argueso et al. 2002). In our study, T-antigen was one of the 51 major detected O-glycans. In addition, that study demonstrated that those T-antigen glycans were carried by the \textit{MUC5B} which was increased around the time of ovulation. This suggests that changes in mucin O-glycosylation modulates the mucus rheological properties such as mucus viscosity and water retention capacity and allow sperm to traverse the cervix around the time of ovulation.

In a proteomic study using human cervical mucus from different phases across the oestrous cycle, core 2 glycans were found in higher levels around the time of ovulation (Andersch-Bjorkman et al. 2007). In the current study, the levels of neutral glycans such as fucosylated glycans were the most abundant with 56 structures across all six ewe breeds. Analysis of mucus in mouse gastrointestinal tract showed lower sialylation in the stomach compared to the colon (Holmén Larsson et al. 2013) as well as it has been demonstrated that increase sialylation contributes to defense against pathogens in the respiratory tract (Denneny et al. 2020). Cervical mucus rheology and hydration seems to be related to the presence of negatively charged terminal groups on glycoproteins, modifying the distension and flexibility of the mucin protein and increasing mucus viscosity (Varki and Schauer 2009). Taken together, these results indicate that cervical mucus collected from all six ewe breeds at the follicular phase had high abundance of neutral mucins and low proportion of acidic glycans, resulting in a more hydrated and less viscous mucus at the follicular phase. This is supported by previous studies using histochemical and complementary biochemical methods that have also suggested cyclic glycosylation variation over the menstrual cycle due to the variation of hormonal levels, mainly oestrogen (Argueso et al. 2002, Iacobelli et al. 1971, Wakefield and Wells 1985).

The use of exogenous hormones to synchronize the oestrous cycle had an effect on the \textit{O}-glycome in some ewe breeds but not others. While the low fertility Suffolk breed had higher
abundance of the type 3 H-antigen (Fuca1-2Galβ1-3GalNAc, peak 8) and glucuronyl-Tn-antigen (GlcAβ1-3GalNAc, peak 14) at a natural cycle compared with the synchronized cycle, there was no difference between natural and synchronized oestrous in both ewe breeds with the highest fertility (Fur and NWS). The results of this study would suggest that the ewe breed has a significant role in the cervical response to exogenous hormones and could help explain the contradictory results regarding the effect of exogenous hormones in gross mucus properties and sperm transport that have been previously reported. In our previous study (Abril-Parreño et al. 2021), we identified increased mucus production at the synchronized cycle compared to the natural cycle at the follicular phase in all six ewe breeds, however mucus viscosity did not follow the same trend in all six ewe breeds. Further studies on mucus viscosity have not been performed in this study due to the limited amount of cervical mucus although the analysis of the addition of sialidases and/or fucosidases to the cervical mucus and subsequently analysis of mucus viscosity are warranted. This could be due to the insensitivity of the chamber filling method used to assess viscosity in the field. Rexroad and Barb (1977) also reported an increase of mucus production while (Smith and Allison 1971) found decreased levels of mucus produced at the synchronized cycle compared to the natural cycle. Maddison et al. (2016) reported similar mucus volumes when Merino ewes were synchronized or not. There have also been reports of changes in spinnbarkeit (Adams and Tang 1979), protein content (Rexroad and Barb 1977) and reduced fertility in sheep (Crocker et al. 2007) due to the use of exogenous hormones.

Changes in sialylation are widely known to participate in sperm regulation, as sialic acids on highly sialylated beta-defensin 126 is required for human sperm to migrate within the mucin mesh of the cervical mucus (Lyons et al. 2018) and has been shown in the bull to affect sperm motility, ability to penetrate mucus in vitro and interaction with the oocyte vestments (Fernandez-Fuertes et al. 2018). However, the role of specific O-glycans from cervical mucin in the selection of sperm is unknown. In the present study, we identified lower abundance of the Sialyl-T-antigen (Neu5Acα2-3Galβ1-3GalNAc; peak 19) in the low fertility Suffolk breed compared with Fur ewes (high fertility) at the natural cycle. At the synchronized cycle, Belclare (medium fertility) had lower abundance of this glycan compared with Fur ewes. The structure (2,3)-sialyl-T-antigen is formed by adding sialic acid to the T-antigen structure, which is antigenic and could represent recognition sites for lectins. This glycan was also identified in the cervical mucus of women (Andersch-Bjorkman et al. 2007) although its role in fertility is unknown.
There are contradictory results regarding the role of sialic acid in the regulation of sperm transport in vivo. Lütjen et al. (1985) found decreased terminal sialic acid in infertile patients compared with cervical samples from fertile women. While other have reported the opposite, a decrease in sialic acid content is related to an increase in mucus receptivity to spermatozoa (Moghissi et al. 1976). In the current study, Sialyl-Tn antigen (Neu5Gcα2-6GalNAc, peak 18) was related to fertility, having lower abundance in Belclare ewes (medium fertility) than in Fur ewes (high fertility). However, Richardson et al. (2019) demonstrated increased amounts of Neu5Ac/Neu5Gc α2-6Gal- GalNAc in Suffolk ewes (low fertility) using a lectin assay. This result was associated with low numbers of sperm in the cervical channels of the Suffolk ewes. This suggests that higher levels of sialylated O-glycans in the low fertility breed are binding sperm and thus inhibiting the sperm progression through the cervix. Specifically, α2,6 linked sialic acid terminal may be involved in the interaction with sialic acid receptors (Siglecs) on the sperm surface, thus causing sperm to become immobilized in the cervical mucins. Sialic acid also plays an important role in the survival of sperm within the female reproductive tract (Crocker et al. 2007), as when sperm are detected an immune response is initiated, which consists of a leukocytic reaction that destroy the majority of the sperm. Specifically, neutrophils are the most abundant immune cell in the human leukocytic reaction following insemination (Thompson et al. 1992). For instance, sialylated cervical mucins seem to silence neutrophils in cows by a sialic acid dependent mechanism (Bornhöft et al. 2019). Further in vitro and in vivo functional studies are required to understand the precise role of specific sialic acid terminals and Siglecs in cervical sperm transport.

It has also been reported that sulfated mucins can inhibit adhesion of bacteria to target gastric cells (Kamisago et al. 1996). In the present study, we identified higher abundance of the sulfated core 2 glycan (Galβ1-3[SO3-GlcNAcβ1-6]GalNAc, peak 15) in the low fertility Suffolk compared with NWS (highest fertility) at the natural cycle. This suggests that these glycosylation changes may be considered a “side effect” of the presence of bacteria and pathogens. In addition, fucosylated core 1 glycan (GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc, peak 16) also had higher abundance in Suffolk compared with NWS ewes (highest fertility) at the synchronized cycle. Fucosylated O-glycans mediate ligand adhesion, pathogen–host interactions and cellular processes via signaling mechanisms (Li et al. 2018). It has been reported that cervical mucus with high fucose protect the vaginal epithelial cells against the binding of Candida albicans (Domino et al. 2009). It is possible that cervical O-glycans bind Candida albicans in order to avoid binding to epithelial cells. This could explain the role of
this fucosylated glycan that was present in higher proportions in the low fertility ewe breeds as an indicator of an active immune response that could have a negative effect on sperm transport. Core 4 glycan (GlcNAcβ1-3[GlcNAcβ1-6]GalNAc, peak 12), the most abundant peak contributing to the total core 4 glycans, was found in higher abundance in the Suffolk than in the highest fertility ewe breed (NWS) at both the natural and synchronized cycle. Core 4 O-GalNAc glycans (as peak 12) have been identified in secreted mucins of some tissues, such as bronchi, colon, and salivary glands (Varki and Schauer 2009) but there are no published studies on its role in sperm transport. Although, a transcriptomic analysis of the bovine cervix (Pluta et al. 2012) identified lowest gene expression of the Glucosaminyl (N-Acetyl) Transferase 2 (GCNT2), which synthetizes the core 4 structures, besides it was correlated with maximal mucin gene expression around the time of ovulation. Therefore, sperm transport can be negatively affected by the high presence of core 4 O-glycans. However, we identified a negative correlation with mucus viscosity and core 4 O-glycans (between other glycans). Our result supports the essential role of O-glycans holding water due to their hydrophilic character and negative charge, which contributes to the viscosity and adhesiveness of mucus (Varki and Schauer 2009). Yet, whether core 4 O-glycans contribute to mucin barrier function and how the rheological properties modulate sperm transport through the cervical mucus remain unclear.

In conclusion, this is the first published study of the O-glycome of sheep cervical mucus. We characterised 124 O-glycans, from which 51 were the major glycans with core 2 and fucosylated glycans as the most common structures (Figure 7). Therefore, higher fucosylation and lower sialylation seems to be the optimal environment for the sperm survival in the cervical mucus. We found no differences in the O-glycan composition between the synchronized and natural cycle in both high fertility ewe breeds (NWS and Fur) but there was an effect in the other ewe breeds. Despite no differences in gross mucus properties previously reported by our group (Abril-Parreno et al. 2021), we identified O-glycan changes and potential biomarkers that could be targeted for the improvement of assisted reproduction techniques such as cervical insemination. For example, sulfated glycan (Galβ1-3[SO3-GlcNAcβ1-6]GalNAc), fucosylated glycan (GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc) and core 4 glycan (GlcNAcβ1-3[GlcNAcβ1-6]GalNAc), which had higher abundance in the low fertility Suffolk breed compared with NWS (high fertility). In addition, core 4 glycans were negatively correlated with mucus viscosity. Future in vitro and in vivo studies using the specific O-glycans found in this study are warranted.

Material and methods
Ethical approval

Protocols were developed in accordance with the Cruelty to Animals Act (Ireland 1876, as amended by European Communities regulations 2002 and 2005) and the European Community Directive 86/609/EC. In Ireland, the study was approved by the Teagasc Animal Ethics Committee and all animal procedures performed were conducted under experimental license from the Health Products Regulatory Authority. In Norway, the study was approved by Norwegian Food safety Authority (FOTS ID 13168). In France, the study was approved by the ethics committee and the Ministry of Research.

Experimental design

The experimental design has previously been described (Abril-Parreño et al. 2021). Briefly, this study was performed as a part of larger project which aimed to interrogate the ewe breed differences in cervical mucus properties and anatomical characteristics across the oestrous cycle (at both follicular and luteal phases) of both a synchronized (using progestogen pessaries combined with equine chorionic gonadotropin) and a natural cycle. In this study, we characterized the O-linked glycan composition in the cervical mucus of six ewe breeds (n = 28 to 30 ewes per breed) across three countries: Ireland (Suffolk and Belclare; low and medium fertility, respectively), France (Ile de France and Romanov; both with medium fertility) and Norway (Norwegian White Sheep (NWS) and Fur; both with high fertility). All the ewes used in this study were multiparous in a range of 4 to 5 years old. We used these ewe breeds due to their known differences in pregnancy rates following cervical/vaginal AI with frozen-thawed semen. Cervical mucus samples were collected from each ewe at the follicular (around the time of ovulation) of a synchronized (56 h post pessary removal) and a natural oestrus cycle (following oestrus detection with a teaser ram), which was then replicated three times with the same ewes over a period of approximately 6 months. After each mucus collection, samples were transported to the laboratory (within 1 h) and an aliquot assessed for cervical mucus properties (weight, viscosity and colour; reported in Abril-Parreño et al. (2021). The remaining sample was stored at -20°C until mucin purification.

Mucin purification from cervical mucus

Mucus samples were ranked according to viscosity at collection (as assessed by the time to fill a 20 µm deep chamber slide) and following thawing were pooled according to viscosity post collection to ensure a sufficient yield of mucins for analysis. Each pool consisted of mucus from five ewes (five most viscous pooled together, then the next five and so on) of the same

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breed collected over three replicates (total of 15 samples pooled across the five ewes). The volume of mucus in each pool/group was ~ 5 mL. Each pooled sample was mixed with an equal volume of 8 M guanidine hydrochloride (GdnCl; Sigma Aldrich, Arklow, Co Wicklow, Ireland) in order to solubilize the mucus and homogenise the samples (the final volume was made up to 30 mL). Dithiothreitol (DTT) (Sigma Aldrich) was added to a final concentration of 10 mM and incubated at 37 °C for 5 h. Iodacetamide (Sigma Aldrich) was added to a final concentration of 25 mM and the samples were incubated at room temperature overnight. The isopycnic density gradient centrifugation was carried out in CsCl/GdnCl. The density of the samples was adjusted to 1.40 g/mL with CsCl in Beckman Ultra-clear tubes which were centrifuged at 65,000 rpm for 18 h at 10 °C in an Optima L-100 XP (Beckman Coulter Inc, Brea, USA) ultracentrifuge using the 70 Ti rotor without break. The density gradient created was unpacked in 1 mL fractions from the top to the bottom of the tube. Following this duplicate 5 μL aliquots from each fraction were blotted onto a polyvinylidene difluoride membrane using a Whatmann manifold Slot blot apparatus (Schleicher & Schnell, Inc, Keene, NH) and stained with periodic acid–Schiff stain (PAS staining (VWR, Radnor, Pennsylvania, USA) in order to assess the relative intensity of carbohydrate in the sample. The carbohydrate rich fractions observed from the slot blot which also had a density profile between 1.35 to 1.45 g/mL were pooled and then separated by size exclusion chromatography. The samples were loaded on a Sepharose CL-4B column (Sigma Aldrich) and eluted with 50 mM Tris/100 mM KCl, pH 7.5, as mobile phase. The eluate was collected in fractions of 4 mL using a fraction collector and then a sample (~20 μL) of all fractions was slot blotted and stained with PAS as described before. Mucin rich fractions were pooled and freeze-dried, then resuspended in H2O and desalted on a Bio-gel P6 column (Bio-Rad Laboratories, Hercules, CA). Fractions collected with the fraction collector were analysed by slot blotting and PAS staining (40 μL), then carbohydrate rich fractions were pooled and freeze-dried. The freeze dried material was used as the final pool of purified mucins from each sample and was weighed and stored at -20 °C until analysis.

**N-glycan release**

Purified cervical mucin samples (starting material: 1 mg) were prepared in tubes using the gel method as previously reported (Royle et al. 2008) and N-glycans were released from the glycoproteins by adding PNGase F as previously described (Kuster et al. 1997), following steps were performed using the fractions that only contain O-glycans. N-glycans released were kept for further analysis.
**O-glycan release using microwave assisted non-reductive β-elimination**

Our group has recently developed a protocol for the extraction and characterisation of O-linked glycans (Wilkinson et al. 2021) with the following modifications. The remaining gels after N-glycan release were crushed by passing them through a small hole in the bottom of a 0.5 mL tube into a 1.5 mL tube at 14,000 rpm and dried down. Then, the crushed gels were resuspended in 250 μL of 40% dimethylamine in water containing 0.1 g/mL ammonium carbonate and transferred to G4 vials (Anton Paar, Graz, Austria). The vials were inserted into a Monowave 450 microwave reactor (Anton Paar), in which the vials were subjected to 20 min microwave radiation at 70 °C and 600 W. After the reaction, the remaining solution including the pieces of gel were transferred to 2 mL tubes and neutralised with 1 M HCl (all samples were at pH 7). Samples were desalted using a Hypersep Hypercarb SPE cartridge (Thermo Fisher Scientific, Waltham, USA) and dried.

**Reproducibility analysis**

Firstly, we determined the viability of the method previously described by Wilkinson et al. (2021), using 1 mg of fetuin glycoprotein (Sigma Aldrich) where the four main O-glycan structures expected in O-glycan release of fetuin were analyzed and presenting a coefficient of variation (CV) below 15%. Then, we validated it for ovine cervical mucin (OCM) by using seven technical replicate release of a randomly selected sample (OCM 37) with all major peaks presenting a CV below 15% (Supplementary Table I). A total of 72 (in duplicate) cervical mucin samples were analyzed in this study. This comprised of six samples per ewe breed (each was a pool of five ewes), two types of oestrous (synchronized and natural) and six ewe breeds.

**Glycan labelling with 2AB**

The N- and O-glycans released were then fluorescently labelled through reductive amination where the aldehyde group of the reducing glycans undergo a condensation reaction with an aromatic amine in excess to drive the formation of a Schiff base. The aromatic amine label used in this study was 2-aminobenzamide (2AB) (Bigge et al. 1995). After the incubation with 1% formic acid for 40 minutes at room temperature, 2AB labelling mixture was added and incubated at 65 °C for 2 hours. Excess label was removed using a Hypersep Diol SPE cartridge (Thermo Fisher Scientific) (Adamczyk et al. 2017).

**Ultra-performance liquid chromatography (UPLC) analysis of 2AB labelled glycans**
The 2AB labelled O-glycans were separated by ultra-performance liquid chromatography (UPLC) analysis using a Glycan BEH Amide, 130Å, 1.7 μm column (Waters Corporation, Milford, MA) on an Aquity H-Class hydrophilic-interaction liquid chromatography-ultra-performance liquid chromatography (HILIC-UPLC) system (Waters Corporation) coupled with a fluorescence detector (Waters Corporation, Milford, MA). Solvent A was 50 mM ammonium formate, prepared with 50 mM formic acid, adjusted to pH 4.4 with ammonium hydroxide solution; and solvent B was acetonitrile. The column temperature was set to 40 °C.

Non-reductive β-eliminated 2AB labelled O-glycans were resuspended in 20 μL 88% ACN and an injection volume of 19 μL was used in a flow rate of 561 μL/min over a 30 min run using the following gradient: 0.00 to 1.47 min – 12% A, 1.47 to 25.00 min – 12% → 47.6% A, 25.00 to 25.60 min – 47.6% → 70% A (flow rate 300 μL/min), 25.60 to 26.80 minutes – 70% A (flow rate 300 μL/min), 26.80 to 28.00 min – 70% → 12% A (flow rate 300 μL/min), 28.00 to 30.00 min – 12% A (flow rate 561 μL/min). The separation temperature was set at 40 °C whilst the sample temperature was 5°C. Excitation and emission wavelengths for fluorescence detection were λ = 330 nm and λ = 420 nm, respectively. External calibration was performed using 2AB labelled glucose oligomers, creating a dextran ladder with retention times of all identified glycan peaks expressed in glucose units (GUs). The 2AB labelled N-glycans were also analysed by UPLC using the method described by Saldova et al. (2014). No N-glycans were found in the cervical mucus samples and the further analysis were only performed using O-glycans.

Mass-spectrometry

Prior to analysis, the 2AB labelled O-glycans were desalted using 10 μL normal phase Phytip® Columns (Phynexus Inc, San Jose, CA). The 2AB labelled O-glycans were resuspended in 10 μL 88% ACN and 9 μL was injected into an Aquity H-Class hydrophilic-interaction liquid chromatography-ultra-performance liquid chromatography (HILIC-UPLC) system (Waters Corporation) with a BEH Glycan column (1.0 x 150 mm, 1.7 μm particle size; Waters Corporation) and an AQUITY fluorescence detector coupled in line with a Waters Xevo G2 QTof system (Waters Corporation). The flow rate was 150 μL/min and the column temperature was set to 60 °C. Solvent A was 50 mM ammonium formate, prepared with 50 mM formic acid, adjusted to pH 4.4 with ammonium hydroxide solution; and solvent B was acetonitrile. The oligosaccharides were eluted over a 30 min run using the following gradient: 0.00 to 1.00 min – 12% A, 1.00 to 25.00 min – 12% → 47% A, 25.00 to 25.50 min – 47% → 70% A, 25.50 to 25.55 min – 70% A (flow rate 100 μL/min), 25.55 to 26.50 min – 70% A (flow rate 100 μL/min), 26.50 to 27.00 min – 70% → 12% A (flow rate 100 μL/min), 27.00 to 30.00 minutes
– 12% A (flow rate 150 μL/min). Excitation and emission wavelengths for fluorescence detection were \( \lambda = 330 \text{ nm} \) and \( \lambda = 420 \text{ nm} \), respectively. The instrument was operated in negative ion sensitivity mode with a capillary voltage of 1.80 kV. The ion source block and nitrogen desolvation gas temperatures were set at 120 °C and 400 °C, respectively. The desolvation gas was set to a flow rate of 600 L/h. The cone voltage was maintained at 50V. Fullscan data for glycans were acquired over \( m/z \) range of 450 to 2500. The data were collected and processed using MassLynx 4.1 software (Waters Corporation).

**Exoglycosidase digestions**

Exoglycosidase digestions were undertaken to determine the \( O \)-glycan sequence and linkage as has been previously described (Saldova et al. 2014). All enzymes were from New England Biolabs (Hitchin, Herts, U.K.) except for NAN1 which was purchased from Prozyme (San Leandro, CA). The 2AB-labelled \( O \)-glycans were incubated in a volume of 10 μL for 18 h at 37 °C in 50 mM sodium acetate buffer, pH 5.5. An array of the following enzymes was used: sialidase cloned from *Streptococcus pneumoniae* and expressed in *E. coli* (NAN1, EC 3.2.1.18), 5 U/mL; sialidase cloned from *Arthrobacter ureafaciens* and expressed in *E. coli* (ABS, EC 3.2.1.18), 1000 U/mL; \( \beta \)-galactosidase cloned from bovine testis and expressed in *Pichia pastoris* (BTG, EC 3.2.1.23), 200 U/mL; \( \beta \)-galactosidase cloned from *Streptococcus pneumoniae* and expressed in *E. coli* (SPG, EC 3.2.1.23), 80 U/mL; \( \alpha \)-fucosidase cloned from bovine kidney and expressed in *E. coli* (BKF, EC 3.2.1.51), 800 U/mL; \( \beta \)-N-acetylglucosaminidase cloned from *Streptococcus pneumonia* and expressed in *E. coli* (GUH, EC 3.2.1.30), 400 U/mL; \( \alpha \)-mannosidase cloned from *Canavalia ensiformis* (Jack Bean) and expressed in *Pichia pastoris* (JBM, EC 3.2.1.24), 400 U/mL; \( \alpha \)-fucosidase cloned from the sweet almond tree (*Prunus dulcis*) and expressed in *Pichia pastoris* (AMF, EC 3.2.1.111), 400 U/mL; \( \beta \)-N-acetylhexosaminidase cloned from *Streptomycyes plicatus* and overexpressed in *E. coli* (JBH, EC 3.2.1.52), 800 U/mL; and \( \alpha \)-galactosidase cloned from green coffee bean and expressed in *E. coli* (CBG, EC 3.2.1.22), 800 U/mL.

**Glycan identification and data validation**

Firstly, chromatograms from the HILIC-UPLC analysis were separated into peaks. Each peak is composed of one or more \( O \)-glycans, of which some could have the same retention time or mass. To separate these glycans, combination of exoglycosidase digestions was used to elucidate the different structures within each peak, since the structural analysis of \( O \)-glycans involves not only monosaccharide composition, but also the type of linkages. HILIC-UPLC
and exoglycosidase digestions analysis was complemented by mass spectrometry data to fully assign and confirm the presence of these glycan structures (Supplementary Figure 8).

**Statistical analysis**

**Glycan analysis.** The UPLC profiles were separated into peaks and the area under each glycan peak was expressed as a relative percentage area derived from the UPLC total profile. Thus, the logit transformed value was used to map the data onto a more Gaussian distribution as previously described by Saldova et al. (2014). Peak data were checked for normal distribution using a Kolmogorov-Smirnov non-parametric test. Multivariate analysis of variance (MANOVA) with post-hoc analysis using Tukey’s Honest Significant difference test was then performed using SPSS software (IBM Corp., Armonk, N.Y., USA) to identify significant peak differences across groups. The pool of mucus was the experimental unit and the model included fixed effects for ewe breed, type of cycle (synchronized and natural) and their interaction.

**Feature analysis.** Glycan peaks were pooled based on similar structural features (core 1 to 4, core 6, fucosylated, sialylated, glucuronylated and sulfated). Features pertaining to a peak were determined based on the major glycan members of that peak. See supplementary table III, where all glycans in all peaks are described, including their amount and peaks included in a particular feature are highlighted in red. We performed statistical analysis on features, as described above.

**Viscosity analysis.** A Spearman correlation was performed to assess the relationship between O-glycan profile and mucus viscosity. The comparisons were considered statistically significant when \( p < 0.05 \).

**Declaration of Competing Interest**
The authors declared no competing interests.

**Author contributions**
L. Abril-Parreño, X. Druart and A. Krogenaes collected the samples from the ewes. L. Abril-Parreño performed the mucin purifications, the O-glycan characterisation and drafted the manuscript. S. Fair, A. Krogenaes, X. Druart and conceived and designed the experimental model and secured funding. R. Saldova designed the glycomic analysis. M. E Gallagher and Colm Reid facilitated the mucin purification. L. Abril-Parreño, H. Wilkinson, J. Morgan, and R. Saldova performed the glycomic data analysis. All authors proof read the manuscript.

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Abbreviations
2AB, 2-aminobenzamide; ACN, acetonitrile; AI, artificial insemination; CV, coefficient of variation; ESI-MS, Electrospray Ionization Mass Spectrometry; FLR-HILIC-UPLC, Fluorescence Hydrophilic Interaction Ultrahigh Performance Liquid Chromatography; Fuc, Fucose; Gal, Galactose; GalNAc, N-acetylglalactosamine; GCNT2, Glucosaminyl (N-Acetyl) Transferase 2; GdnCl, Guanidine hydrochloride; GlcA, Glucuronic acid; GlcNAc, N-acetylglucosamine; GU, glucose unit; HILIC-UPLC, Hydrophilic-interaction liquid chromatography-ultra-performance liquid chromatography; NWS, Norwegian White Sheep; OCM, ovine cervical mucin; PCA, principal component analysis;

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### Tables

**Table I** Cervical mucus O-glycans that were significantly different at the follicular phase of a natural and synchronized cycle in six European ewe breeds. Green indicates a significant increase of that particular glycan in the natural cycle and red indicates a significant decrease of that particular glycan in the natural cycle. Only O-glycans with significant differences are shown ($p < 0.05$).

| Major component of Peak | Peak Suffolk | Peak Belclare | Peak Ile de France | Peak Romanov | Peak Fur | Peak NWS |
|-------------------------|--------------|---------------|--------------------|--------------|---------|----------|
| Galβ1-3GalNAc           | 4            | 0.052         | 0.103              | 0.001        | 0.001   | 1.000    | 0.147    |

Natural vs. Synchronized (P value)
Galβ1-3GlcNAc 5 0.276 1.000 0.020 0.116 0.976 0.587  
(Fucα1-2Galβ1-3)GalNAc 8 0.012 0.521 0.659 1.000 1.000 0.159  
Neu5Aca2-3GalNAc and GlcNAcβ1-3Galβ1-3GalNAc 11 1.000 0.903 1.000 0.006 0.245 1.000  
Galα1-3Galβ1-3GalNAc 13 0.270 0.024 1.000 1.000 0.973 0.649  
GlcAβ1-3GalNAc 14 0.002 0.287 0.810 1.000 0.922 0.867  
GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc 16 1.000 1.000 0.049 0.999 1.000 1.000  
Galβ1-3GlcNAcβ1-3(Fucα1-2Galβ1-3)GalNAc 26 0.996 0.020 1.000 0.977 0.136 0.985  

NWS = Norwegian White Sheep

Legends to Figures

**Figure 1.** (A) Representative FLR-HILIC-UPLC chromatograms of O-glycans released by microwave-assisted non-reductive β-elimination in sheep follicular phase cervical mucin (six
European ewe breeds ordered top-down from lowest to highest fertility). Blank in black.

Retention times of all identified glycan peaks expressed in glucose units (GUs). Major components of each peak shown. (B) Structural symbols for the O-glycans and common core structures found in this study.

**Figure 2.** Pie charts showing the relative abundance of (A) of the core glycan types and (B) the main types of glycan modifications in sheep cervical mucins at the follicular phase, including pictograms of representative structures of each type of core and modification, respectively.
Figure 3. Nested plots of significantly different O-glycans in cervical mucus of six ewe European ewe breeds collected at the follicular phase of a natural cycle. * ($p < 0.05$), ** ($p < 0.01$), *** ($p < 0.001$). NWS = Norwegian white sheep.

Figure 4. Nested plot of core 6 glycosylation of cervical mucin from six European ewe breeds at the follicular phase of a natural cycle. * ($p < 0.05$), ** ($p < 0.01$), *** ($p < 0.001$). NWS = Norwegian white sheep.
Figure 5. Nested plots of significantly differentially expressed O-glycans in cervical mucus of six European ewe breed collected at the follicular phase of a synchronized cycle. * (p < 0.05), ** (p < 0.01), *** (p < 0.001). NWS = Norwegian white sheep.
Figure 6. (A) Nested plots of significant feature changes (fucosylation, glucuronylation, Neu5Ac, core 1 and core 4) in cervical mucus of six European ewe breeds at the follicular phase of a synchronized cycle. * ($p < 0.05$), ** ($p < 0.01$), *** ($p < 0.001$). (B) PCA plot shows the three core 4 associated peaks (12, 41, 50) which are driving the significant difference between Suffolk and Belclare (samples within each breed are indicated by the same colours).

Figure 7. Overview of the main findings of the characterization of $O$-glycans in cervical mucus across six ewe breeds with divergent fertility at the follicular phase (around the time of ovulation) of the oestrous cycle.