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Evidence for Differential Glycosylation of Trophoblast Cell Types

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Human placental villi are surfaced by the syncytiotrophoblast (STB), with a layer of cytotrophoblasts (CTB) positioned just beneath the STB. STB in normal term pregnancies is exposed to maternal immune cells in the placental intervillous space. Extravillous cytotrophoblasts (EVT) invade the decidua and spiral arteries, where they act in conjunction with natural killer (NK) cells to convert the spiral arteries into flaccid conduits for maternal blood that support a 3–4 fold increase in the rate of maternal blood flow into the placental intervillous space. The functional roles of these distinct trophoblast subtypes during pregnancy suggested that they could be differentially glycosylated. Glycomic analysis of these trophoblasts has revealed the expression of elevated levels of biantennary N-glycans in STB and CTB, with the majority of them bearing a bisecting GlcNAc. N-glycans terminated with polylactosamine extensions were also detected at low levels. A subset of the N-glycans linked to these trophoblasts were sialylated, primarily with terminal NeuAcα2–3Gal sequences. EVT were decorated with the same N-glycans as STB and CTB, except in different proportions. The level of bisecting type N-glycans was reduced, but the level of N-glycans decorated with polylactosamine sequences were substantially elevated compared with the other types of trophoblasts. The level of triantennary and tetraantennary N-glycans was also elevated in EVT. The sialylated N-glycans derived from EVT were completely susceptible to an α2–3 specific neuraminidase (sialidase S).

The possibility exists that the N-glycans associated with these different trophoblast subpopulations could act as functional groups. These potential relationships will be considered. Molecular & Cellular Proteomics 15: 10.1074/mcp.M115.055798, 1857–1866, 2016.

Human development in utero requires the formation of a functional placenta that mediates the transport of nutrients, the exchange of gases, and the production of hormones that are required to maintain fetal viability. The placenta also serves as the immune interface between the mother and her histoincompatible fetus. Human trophoblasts are specialized placental cells that come into direct physical contact with maternal immune cells at both the villous surface and within the maternal decidua and myometrium. The syncytiotrophoblast (STB) layer is localized to the villous surface, with subsyncytial cytotrophoblasts (CTB) lying just beneath the STB. Regions of the syncytium can become damaged, which occurs at an increased frequency in many complicated pregnancies, and this damage can be repaired by division and fusion of cytotrophoblasts. It is likely that CTB in these damaged regions are at least transiently in direct contact with circulating maternal immune cells (1, 2).

Extravillous cytotrophoblasts (EVT) derived from placental stem cells penetrate through the tips of the anchoring villi and differentiate into invasive trophoblasts that migrate into the decidua, myometrium and spiral arteries (3). Trophoblasts that enter the spiral arteries differentiate into endovascular cytotrophoblasts. These cytotrophoblasts act in concert with maternal NK cells to remodel these arteries into flaccid conduits that support a three- to fourfold increase in the rate of maternal blood flow into the placental intervillous space at a reduced pressure compared with the maternal circulation (4–6).

The abbreviations used are: STB, syncytiotrophoblast; BBSCT, biantennary bisected type; CFG, Consortium for Functional Glycomics; CTB, cytotrophoblasts; DNK, decidual natural killer; EVT, extravillous cytotrophoblasts; HLA, human leukocyte antigen; LacNAc, N-acetyllactosamine; NK, natural killer; PAEP, progestagen-associated endometrial protein; PMAA, partially methylated alditol acetate.
Interactions between villous trophoblasts and maternal immune cells would normally be expected to trigger major histocompatibility responses because of the expression of paternal human leukocyte antigens (HLA). However, STB and CTB do not express the HLA class I molecules that are necessary to evoke such responses (7, 8). This absence of HLA molecules on STB and CTB could make these trophoblasts potential targets for lysis by NK cells (9). However, STB and CTB are highly resistant to lysis by peripheral blood NK cells in vitro (10, 11).

Unlike CTB and STB, EVT express on their cell surface a classical paternal HLA class I molecule designated HLA-C (7, 8). This presentation means that EVT are semiallogeneic during natural pregnancies and are therefore potentially subject to powerful histocompatibility-based immune responses. However, the activation of such responses in a manner that harms the fetus and placenta does not typically occur during normal pregnancies. How EVT evade immune recognition in the gravid human uterus remains a major enigma. In contrast to STB or CTB, EVT are also susceptible to lysis by purified decidual NK (dNK) cells (12–14). Because EVT are the precursors of endovascular trophoblasts that remodel the spiral arteries (15), increased lysis of EVT by activated dNK cells could decrease the number of endovascular trophoblasts, thereby contributing to deficient spiral artery remodeling and inadequate circulation in the intervillous space.

Glycomic analyses of glycoproteins, whole cells, tissues, and extracellular matrices have been useful for defining potential carbohydrate functional groups and their diverse roles in many cellular processes (16–21). Glycomic analysis of STB, CTB, and EVT was performed in this study to determine if the N-glycans associated with these cells have potential functional roles.

**EXPERIMENTAL PROCEDURES**

**Isolation of CTB and STB**—The protocol for obtaining placenta used for this portion of this study was approved by the Institutional Review Board of Washington University School of Medicine, Term, singleton placentas from uncomplicated pregnancies were obtained, and CTB were isolated as described previously (22, 27). All reagents were obtained from Sigma-Aldrich (St. Louis, MO) except for DNase, which was from Roche Diagnostics (Indianapolis, IN). In brief, villous tissue was isolated and digested with dispase, trypsin, and Dnase and CTB were isolated on a continuous gradient of Percoll. Over 85% of the cells stained positive for cytokeratin 7, a TB specific marker. CTB were obtained after 24 h of culture in DMEM/10% FBS in 5% CO2/air. To obtain STB, culture was continued for an additional 48 h with daily changes of medium. During this time, spontaneous differentiation and fusion occurs, with over 70% of the nuclei being present in multinucleated syncytiotrophoblasts after 72 h of culture (22). We observed an ~50-fold increase of hCG expression from 24 to 72 h of culture, confirming efficient differentiation.

**Isolation of EVT**—All human tissue collection necessary for this study was approved by the local ethics committee of the Geneva University Hospital. All patients provided their informed written consent prior to their inclusion in the study. Placental tissue was obtained from patients undergoing an elective termination of pregnancy during the first trimester (8–12 weeks of gestation). The procedure for the isolation of EVT was performed as previously described (23). Placental tissue specimens were isolated and washed several times in sterile Hanks balanced salt solution. The tissue samples were subjected to enzymatic digestion five times for 20 min at 37 °C (0.25% trypsin, 0.25 mg/ml Disase I, Roche, Diagnostics GmbH). After this incubation, fetal bovine serum (FBS) was added to neutralize the trypsin mixture. The cells were suspended in Dulbecco’s modified Eagle’s medium (DMEM) (Invitrogen, Switzerland). This cell suspension was filtered through a 50-μm mesh and the resulting supernatant was applied onto tubes containing a Percoll gradient (70–5% Percoll diluted with HBSS). The samples were centrifuged for 25 min at 1200 × g. The 30–45% Percoll layer was collected, and the trophoblast cells were harvested, washed, and resuspended in DMEM containing 10% fetal bovine serum.

The cells were transferred to Petri dishes and incubated for 15 min at 37 °C. Trophoblasts in the culture supernatants were collected by centrifugation and resuspended in culture medium. They were transferred to six-well plates (4 × 10⁵ cells/well) and in 96-well plates (1 × 10⁴ cells/well). After 48 h of culture, 95% of the cells were: (1) negative for vimentin; and (2) positive for cytokeratin 7 and HLA-G.

**Processing of Trophoblasts to Acquire N- and O-glycans**—All cell samples were subjected to a standard protocol (24). Briefly, cells were suspended in lysis buffer (25 mM Tris, 150 mM NaCl, 5 mM EDTA and 1% CHAPS (v/v), pH 7.4) before homogenization and sonication were performed. The homogenates were subsequently dialyzed against a 50 mM ammonia bicarbonate buffer, pH 7.5, after which the samples were lyophilized. Cell/tissue extracts were reduced and carboxymethylated and then treated with trypsin. The treated samples were purified using a C18 cartridge (Oasis HLB Plus, Waters, Milford, MA) prior to the release of N-glycans by peptide N-glycosidase F (recombinant from Escherichia coli, Roche Applied Science, Penzberg, Germany) digestion. Released N-glycans were permethylated and then purified using a Sep-Pak C18 cartridge (Waters) prior to MS analysis.

Purified, undervatized N-glycans were incubated with sialidase S (recombinant from Streptococcus pneumoniae, Prozyme Glyko, Hayward, CA) or β1,4-galactosyltransferase (from bovine milk, Merck, Kenilworth, NJ) separately. Sialidase S digestion was carried out using sialidase S in 50 mM sodium acetate, pH 5.5. The β1,4-galactosyltransferase reaction was performed using purified β1,4-galactosyltransferase in 50 mM 3-(N-morpholino)-propanesulfonic acid (MOPS) containing 45 μM UDP-Gal, pH 7.4. The resulting enzyme-treated samples were lyophilized and permethylated prior to MS analysis.

**Mass Spectrometric Analysis**—MS data were obtained using a Voyager-DE TM STR MALDI-TOF/TOF mass spectrometer (AB SCIEX, Framingham, MA). In the MS/MS experiment, the dissolved sample was dried and then re-dissolved in 10 μl methanol and 1 μl of the sample was mixed with 1 μl of matrix, 20 mg/ml 2,5-dihydroxybenzoic acid (DHB) in 70% (v/v) aqueous methanol and loaded on to a metal target plate. The instrument was run in the reflectron positive ion mode. The accelerating voltage was 20 kV. MS/MS data were acquired using a 4800 MALDI-TOF/TOF mass spectrometer (AB SCIEX, Framingham, MA). In the MS/MS experiment, the dissolved sample was dried and then re-dissolved in 10 μl methanol. 1 μl of the sample was mixed with 1 μl of matrix, 10 mg/ml diaminobenzophenone (DABP) in 70% (v/v) aqueous acetonitrile and loaded on to a metal target plate. The instrument was run in the reflectron positive ion mode. The collision energy was set to 1 kV with argon as the collision gas. The 4700 calibration standard (mass standards kit for the 4700 proteomics analyzer, Applied Biosystems) was used as the external calibrant for the MS and MS/MS modes.

**GC/MS Linkage Analysis**—GC-MS linkage analysis of partially methylated alditol acetates (PMAAs) was carried out using a Perkin-Elmer Life Sciences Clarus 500 instrument (Framingham, MA) fitted with
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RTX-5MS column (30 m × 0.32 mm internal diameter, Restek Corp., Bellefonte, PA). The PMAAs were prepared from permethylated N-glycans as described previously (25). The permethylated glycans were hydrolyzed with 2 ml trifluoroacetic acid at 121 °C for 2 h, and then reduced with 10 mg/ml sodium borohydride in 2 ml aqueous ammonium hydroxide at room temperature for 2 h, and acetylated with acetic anhydride at 100 °C for 1 h. The sample was dissolved in hexanes and injected onto the column after the oven temperature reached 60 °C. The column was maintained at this temperature for 1 min and then heated to 300 °C at a rate of 8 °C/min. Running a blank reached 60 °C. The column was maintained at this temperature for 1

Experimental Design—

CTB N-glycans—(1) All N-glycans were presumed to have a Manα1–6(Manα1–3)Manβ1–4GlcNAcβ1–4GlcNAc core structure based on knowledge of the N-glycan biosynthetic pathway and PNGase F specificity. (2) The composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode were manually interpreted. (3) MALDI-TOF/TOF MS/MS analyses were carried out on the following ions: m/z 2592, 2635, 2837, 2850, 3141, 3212, 3300, and 3416. All MS/MS spectra were annotated manually with the assistance of a glycobiomics tool, GlycoWorkBench (version 1.0.3353). (4) GC-MS linkage analysis was performed on partially methylated alditol acetates (PMAAs). (5) Sialidase S digestion was followed by manual assignment of the composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode. (6) Sialidase S digestion was followed by MALDI-TOF/TOF MS/MS analyses of the following ions: m/z 3143 and 4939. All MS/MS spectra were annotated manually with the assistance of a glycobiomics tool, GlycoWorkBench (version 1.0.3353). (7) The β1,4-galactosyltransferase reaction was followed by manual assignment of the composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode.

STB N-glycans—(1) All N-glycans were presumed to have a Manα1–6(Manα1–3)Manβ1–4GlcNAcβ1–4GlcNAc core structure based on knowledge of the N-glycan biosynthetic pathway and PNGase F specificity. (2) The composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode were manually interpreted. (3) MALDI-TOF/TOF MS/MS analyses were carried out on the following ions: m/z 2592, 2635, 2837, 2850, 3141, 3212, 3300, and 3416. All MS/MS spectra were annotated manually with the assistance of a glycobiomics tool, GlycoWorkBench (version 1.0.3353). (4) GC-MS linkage analysis was performed on partially methylated alditol acetates (PMAAs). (5) Sialidase S digestion was followed by manual assignment of the composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode. (6) Sialidase S digestion was followed by MALDI-TOF/TOF MS/MS analyses of the following ions: m/z 3143 and 4939. All MS/MS spectra were annotated manually with the assistance of a glycobiomics tool, GlycoWorkBench (version 1.0.3353). The β1,4-galactosyltransferase reaction was followed by manual assignment of the composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode.

EVT N-glycans—(1) All N-glycans were presumed to have a Manα1–6(Manα1–3)Manβ1–4GlcNAcβ1–4GlcNAc core structure based on knowledge of the N-glycan biosynthetic pathway and PNGase F specificity. (2) The composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode were manually interpreted. (3) MALDI-TOF/TOF MS/MS analyses were carried out on the following ions: m/z 2519, 2850, 3141, 3212, 3415, 3964, and 4041. All MS/MS spectra were annotated manually with the assistance of a glycobiomics tool, GlycoWorkBench (version 1.0.3353). (4) Sialidase S digestion was followed by manual assignment of the composition (numbers of Hex, Fuc, etc.) of the glycans derived from MALDI-TOF MS in positive ion mode.

RESULTS

Analysis of N-glycans Associated with CTB—A modification of the Kliman procedure was employed to isolate samples of CTB from eight term placentas (22, 27). Each sample was subjected to glycomic profiling exactly as described in a previous study (16). High quality MALDI data were obtained for the N-glycans of all the CTB samples. There were no apparent differences in the spectra between eight individuals. A representative MALDI spectrum of N-glycans derived from CTB was also shown in Fig. 1. The theoretical and observed m/z values of the glycans and the manual interpretation of the composition of these glycans are shown in supplemental Table S1. Signals for high mannose type N-glycans (Manα1–3GlcNAc2) were abundant at m/z 1579.8, 1783.9, 1988.0, 2192.1, and 2396.1. In addition numerous signals for complex-type biantennary, triantennary and tetraantennary N-glycans were observed, and their structures are shown in the cartoon annotations in Fig. 1. To decrease the complexity of all the MALDI spectrum figures, the cartoon annotations for minor signals less than m/z 3250 are shown in supplemental Table S2. Nearly all the complex-type glycans carry core α1–6 linked fucose, consistent with their localization to the plasma membrane, and sialic acid is the major capping sugar on their N-acetyllactosamine antennae. The most abundant complex-type structures give signals at m/z 2431.1 (mono-sialylated biantennary), 2489.2 (non-sialylated, core fucosylated biantennary), 2850.3 (mono-sialylated, core fucosylated biantennary), 2966.3 (disialylated, core fucosylated biantennary), and 3211.4 (disialylated, core fucosylated biantennary). Interestingly three of these glycans have masses consistent with biantennary bisected type (BBSCT) structures (m/z 2489.2, 2850.3, and 3211.4). Minor signals for N-glycans bearing Lewisα (m/z 2592.2, 2837.3, 3024.3) and sialyl-Lewisα type antennae (m/z 3140.3) were also detected. Evidence for N-glycans bearing polylactosamine type sequences was indicated by the signal at m/z 3299.5.

Additional experiments were performed to firmly establish the glycan assignments shown in Fig. 1. First, to confirm assignments of BBSCT glycans, the N-glycan mixture was incubated with a β-galactosyltransferase in the presence of UDP-Gal. This enzymatic modification adds galactose to all antennae bearing terminal GlcNAc except for the bisecting GlcNAc which is sterically inaccessible to the enzyme. After this treatment, there was no detectable addition of galactose.
to putative BBSCT glycans (supplemental Fig. S1). In contrast, the truncated antennae of immature glycans (for example m/z 1835.9) were extended by galactose, confirming that the β-galactosyltransferase was reacting successfully. In addition, GC-MS linkage analysis of the glycan mixture (supplemental Table S3) provided further evidence for the presence of bisecting GlcNAc. The presence of Lewisx and sialyl-Lewisx type antennae on these N-glycans was confirmed by MS/MS analysis (supplemental Figs. S2, S3, S4). The glycan mixture was also treated with sialidase S, a bacterial neuraminidase (sialidase S) that specifically removes terminal α2–3 linked sialic acid. This enzymatic step removed nearly all of the sialic acid from these glycans, confirming that only a minor amount of sialic acid was α2–6 linked on glycoproteins associated with this cell type (Fig. 2). The major N-glycans that were detected after digestion were high mannose type N-glycans and the BBSCT N-glycan. This desialylation step also revealed minor amounts of N-glycans bearing polylactosamine sequences at m/z 3591.6, 3765.6, 3836.6, 4040.6, 4214.8, 4489.6, and 4938.6 (Fig. 2). MS/MS analysis of the peak at m/z 4938.6 (4939) showed that the maximum number of the LacNAc unit observed is five (supplemental Fig. S5).
FIG. 2. Annotated MALDI-TOF MS spectra of permethylated sialidase S treated N-glycans from CTB (upper panel), STB (middle panel) and EVT (lower panel). Profiles were obtained from the 50% acetonitrile fraction from a C18 Sep-Pak column. All ions are \([M+Na]^+\). Putative structures are based on the molecular weight, N-glycan biosynthetic pathway, MS/MS data and enzymatic digestion results. Annotations are simplified to biantennary structures, with additional LacNAc units and capping sugars listed outside the bracket, for mixtures of isobaric multiantennary glycans, some of which have antennae containing extended LacNAc repeats. Minor peaks are only labeled with their m/z values; their putative structures can be found in supplemental Table S2.

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Glycomic Analysis of Human STB—These trophoblasts were obtained after in vitro differentiation of primary CTB as described previously (22, 27). They were subjected to the same procedure that was employed to analyze the CTB N-glycome (16). The theoretical and observed m/z values of the glycans and the manual interpretations of the composition of these glycans are shown in supplemental Table S4. There were no apparent differences in the spectra between individuals. A representative MALDI spectrum is shown in Fig. 3. There were essentially no differences in glycan expression between STB and CTB. This same overlap was also observed after sialidase S digestion of N-glycans derived from STB (Fig. 2). The major signals were for the high mannose type N-glycans (Man₅₋₉GlcNAc₂) and the BBSCT N-glycan (m/z 2489.1).

FIG. 3. Annotated MALDI-TOF MS spectra of permethylated N-glycans from STB86. The top panel shows the glycans in the mass range from 1499 to 3250 and the bottom panel shows the glycans in the mass range from 3249 to 4700. In the top panel, minor peaks are only labeled with their m/z values; their putative structures can be found in supplemental Table S2. Compared with the top panel, the bottom panel has been magnified ~37 times. Because of the fact that the peak signal in the mass range from 3900 to 4700 in the bottom panel is weaker, this part has been further magnified five times (upper trace). Profiles were obtained from the 50% acetonitrile fraction from a C18 Sep-Pak column. All ions are [M+Na]⁺. Putative structures are based on the molecular weight, N-glycan biosynthetic pathway, MS/MS data and enzymatic digestion results. Glycans at m/z 2966, 3777, and 4587 are clearly annotated, which is because of the fact that their structures are unequivocal because each antenna is capped with a sialic acid and thus they are homogeneous bi-, tri-, and tetraantennary glycans. However, the glycan structure is not always as unequivocal as the glycan at m/z 2966 as biosynthetically non-fully sialylated glycan molecular ion species could be made up of mixtures of structural isomers. Annotations are simplified to biantennary structures, with additional LacNAc units and capping sugars listed outside the bracket, for mixtures of isobaric multiantennary glycans, some of which have antennae containing extended LacNAc repeats.

Glycomic Analysis of Human EVT—This subpopulation of trophoblasts was subjected to glycomic analysis to determine if there were any differences between them and other types of trophoblasts. The theoretical and observed m/z values of the glycans and the manual interpretations of the composition of these glycans are shown in supplemental Table S5. Although the N-glycans linked to EVT were very similar to those associated with CTB and STB, there were major differences in levels of expression (Fig. 4). The larger complex-type N-glycans were far more prevalent in EVT, which can be most readily observed via the comparison of glycans after sialidase S digestion (Fig. 2). The most abundant peak, after the removal of α2–3 linked sialic acid, was m/z 3141.7, which corresponds to a complex type N-glycan with four LacNAc units in its antennae. This peak was accompanied by a prominent
series of N-glycans with extended poly lactosamine antennae containing additional LacNAc moieties (m/z 3590.6, 4039.6, 4488.4, and 4937.3). These glycans were far more abundant in EVT than in either CTB or STB. As a consequence, the relative abundance of the BBSCT N-glycan approximately m/z 2489 became less prominent in the overall spectrum of EVT compared with the other complex N-glycans. Another important difference was the absence of any detectable sialylated glycans after digestion with sialidase S. This result confirmed that all of the sialic acid associated with EVT is attached via α2–3 linkages.

DISCUSSION

The N-glycans associated with different types of human trophoblasts were analyzed in this study. A specific caveat to this analysis is that the STB were differentiated from CTB in vitro (22, 27). However, the microvillous membrane of human STB derived from the placenta stain intensely with lectins that bind to: (1) high mannose type N-glycans (Concanavalin A); (2) biantennary bisecting type N-glycans (erythromedagglutinating phytohemagglutinin); (3) N-glycans bearing a core α1–6 linked fucose (Pisum sativum agglutinin); and (4) terminal α2–3 linked sialic acid (Maackia amurensis agglutinin) (28). By contrast, these microvillous membranes stain very poorly with a lectin that specifically binds to α2–6 linked sialic acid (Sambucus nigra agglutinin) (28). These lectin binding results are consistent with the glycomic analysis of STB presented in this report.

The human placenta nourishes the fetus during its development. This organ also negotiates the peaceful co-existence of the mother and her histoincompatible fetus during pregnancy. Specific adaptations have been made that enable 85–90% of women to fully accommodate their fetus until development in utero is complete. Trophoblasts are placental...
cells that encounter maternal immune cells at two distinct interfaces. STB and CTB interact with maternal immune cells at the villous interface. EVT and their derivatives come into direct physical contact with maternal immune cells during their invasion of the decidua, myometrium and spiral arteries. The current study was undertaken to determine if functional glycosylation of trophoblasts could contribute to accommodation of the histoincompatible human fetus.

**Glycosylation of Villous Trophoblasts**—Our glycomic analyses of CTB and STB reveal that these cells express both high mannose and complex-type glycans. The majority of the latter are core fucosylated, biantennary structures with LacNAc or oligo-LacNAc antennae that are partially capped with 2–3 linked (major) or 2–6 linked (minor) sialic acid. A tiny portion of the glycome carries fucosylated antennae in Lewisx/y and sialyl-Lewisx-type sequences. Significantly, many of the biantennary glycans have a bisecting GlcNAc. These N-glycans have previously been implicated in the suppression of NK cell cytotoxicity in highly specialized assay systems (29, 30). STB and CTB are highly resistant to NK cell-mediated cytotoxicity in vitro (10, 11). Whether or not this elevated expression of biantennary bisecting type N-glycans on STB and CTB plays a role in protecting these HLA class I negative cell types from NK cell mediated cytotoxicity is a topic for further investigation.

**Expression of Glycans on EVT**—Notably, the profile of N-glycans in EVT was different from those observed in villous trophoblasts, with EVT expressing relatively lower levels of biantennary bisecting type N-glycans and much higher levels of triantennary and tetraantennary N-glycans. N-glycans decorated with polylactosamine sequences were also considerably elevated in EVT compared with either CTB or STB. Unlike these villous trophoblasts, the sialylated N-glycans associated with EVT were also exclusively capped with α2–3 linked NeuAc. Our finding of the expression of polylactosamine-decorated N-glycans in EVT support previous findings by Fisher and coworkers. These investigators demonstrated that HLA-G produced by EVT and secreted into amniotic fluid migrates as a polydisperse band between 35–50 kDa that is collapsed into a single 35–36 kDa band after digestion with endo-β-galactosidase, an enzyme that specifically depolymerizes polylactosamine sequences (31, 32). Together, these findings suggest that a substantial proportion of native HLA-G molecules in EVT are decorated with polylactosamine sequences.

The expression of bisecting type N-glycans, though lower than in STB or CTB, could nonetheless provide EVT with some measure of protection from dNK cell responses. However, the exact level of protection remains a matter of conjecture with the available data. Results from one study suggest that dNK cells are incapable of forming mature activating synapses with EVT and thus are not cytotoxic (33). Other groups have reported that EVT are susceptible to lysis by dNK cells (12–14). Fisher and coworkers have provided evidence that decidual macrophages induce tolerance to EVT via their secretion of transforming growth factor-β1 (12). These observations are complicated by the presence of glycodelin-A and CA125, two soluble factors that are produced in abundance in the endometrium and decidua from the initiation of implantation up until the 20th week of gestation. Glycodelin-A and CA125 also suppress NK cell cytotoxicity in vitro (34–38). NK cell cytotoxicity is also directly suppressed by galectin-3 that is also present at elevated levels in the uterus during pregnancy (39, 40). These findings suggest that there are likely redundant pathways for the induction of NK cell tolerance in the endometrium and decidua during the first two trimesters of human pregnancy.

**Expression of Galectins (LGALS) and Galectin Ligands in the Pregnant Uterus**—Galectins are a family of β-galactoside binding proteins that express one or two conserved carbohy-
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