'Sweetening' Pregnancy: Galectins at the Fetomaternal Interface

Ada G. Blidner¹, Gabriel A. Rabinovich^{2,3}

¹Instituto de Oncología Ángel H. Roffo, Universidad de Buenos Aires, Buenos Aires, Argentina;

²Laboratorio de Inmunopatología, Instituto de Biología y Medicina Experimental (IBYME), Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Buenos Aires, Argentina;

³Laboratorio de Glicómica Funcional, IQUIBICEN-CONICET, Departamento de Química Biológica, Facultad de Ciencias Exactas y Naturales,

Universidad de Buenos Aires, Buenos Aires, Argentina

Keywords

Angiogenesis, fetomaternal tolerance, galectins, immune privilege, implantation, pregnancy

Correspondence

Gabriel A. Rabinovich, Laboratorio de Inmunopatologia, Instituto de Biología y Medicina Experimental (IBYME), Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Vuelta de Obligado 2490, C1428 Buenos Aires, Argentina. E-mail: gabyrabi@gmail.com

Submission January 9, 2013; accepted January 11, 2013.

Citation

Blidner AG, Rabinovich GA. 'Sweetening' pregnancy: galectins at the fetomaternal interface. Am J Reprod Immunol 2013; 69: 369–382

doi:10.1111/aji.12090

AIRI

Successful mammalian pregnancy relies upon acceptance of a semi-allogeneic fetus by the maternal immune system. Lessons learned from studies on protective immunity to microbial infections and tumours, prevention of autoimmunity, and allograft rejection have contributed to delineate the mechanisms leading to T-cell tolerance at the fetomaternal interface. Recent observations highlight the contribution of galectins, a family of endogenous glycan-binding proteins, to critical biological events occurring during mammalian gestation, including immune cell tolerance, inflammation, implantation, and angiogenesis. These multifunctional lectins can hierarchically control a cascade of immunoregulatory events including the expansion, recruitment, and function of regulatory T cells, the promotion of tolerogenic dendritic cells, and the execution of T-cell death programs. In addition, galectins can control cell adhesion and signaling events critical for implantation and are involved in fundamental processes linking tissue hypoxia to angiogenesis. In an attempt to integrate the regulatory roles of galectins to immunological and vascular programs operating during pregnancy. Here we outline the regulated expression and function of individual members of the galectin family within the fetoplacental unit and their biological implications for the development and preservation of successful pregnancies.

Mechanisms leading to fetomaternal tolerance

Homeostatic signals delivered in the form of immunosuppressive cytokines or inhibitory receptors are integrated into a cascade of regulatory circuits that sustain immune tolerance at the fetomaternal interface.^{1,2} Several tolerogenic mechanisms contribute to prevent fetal damage resulting from overexuberant immune responses to fetal antigens.¹ These include a shift toward a T helper (Th) type-2 cytokine profile,^{3,4} differentiation, and/or recruitment of CD4⁺ CD25⁺ FoxP3⁺ T regulatory (T_{reg}) cells^{5–8} and

American Journal of Reproductive Immunology **69** (2013) 369–382 © 2013 John Wiley & Sons A/S expansion of uterine immunomodulatory natural killer (uNK) cells.⁹ Moreover, trophoblast cells express inhibitory signals such as programed deathligand 1 (PD-L1) and indoleamine 2,3-dioxygenase (IDO), synthesize immunosuppressive cytokines such as transforming growth factor- β (TGF- β) and IL-10¹⁰ and promote death of uterine T cells via the Fas ligand (Fas L) and tumour necrosis factor (TNF)-related apoptosis-inducing ligand (TRAIL) path-ways.^{11,12} Furthermore, cells that integrate the fetoplacental unit can elude allorecognition by expression of 'non-classical' HLA molecules¹ and can elaborate a variety of immunomodulatory neuropeptides including vasoactive intestinal peptide.^{13–15} Additionally, a functional cross-talk between immune and endocrine systems, mediated by progesterone, progesterone-induced blocking factor, and human chorionic gonadotropin (hCG) has been reported to contribute to immune privilege during pregnancy.^{16,17} Although there is still no integrated portrait of these regulatory circuits, disruption of single regulatory pathways may lead to substantial and unpredictable inflammatory responses that induce fetal loss.^{18,19} Here, we outline the contribution of galectins to the promotion of fetomaternal tolerance and the development and preservation of a successful pregnancy.

Glycans and glycan-binding proteins in pregnancy

A multiplicity of regulatory mechanisms involving both mother and embryo cells govern embryo implantation, immune tolerance, and establishment of a functional vascularized placenta. These dynamic processes involve the secretion of a variety of cytokines, chemokines, hormones, and growth factors and a network of specialized cells, including trophoblast cells, epithelial cells, decidual stromal cells, endothelial cells and immune cells.²⁰ As glycans and glycan-binding proteins play key roles in immune cell communication and signaling, their involvement in pregnancy-related processes is not surprising. In fact, the enormous variety of glycans that cover the surfaces of mammalian cells have the potential of storing critical biological information that is decoded by a large number of glycan-binding proteins or lectins.²¹ An interesting example illustrating this concept is the complex mucin MUC1. When a mammalian blastocyst penetrates the uterine cavity, the surface epithelium of the uterus is extensively covered by MUC1 which prevents the attachment of the highly adhesive blastocyst to an improper site. Indeed, in the human endometrium, MUC1 is upregulated during the implantation period.²² This suggests that the human endometrial surface epithelium prevents blastocyst adhesion except for the precise place where the embryo attaches. It has been hypothesized that proinflammatory cytokines released in the uterine stroma induce local degradation of MUC1 which enables the blastocyst to attach to a specific area of the uterus.²² Whether lectin–glycan interactions participate in MUC1-driven control of the implantation process remains unknown.

Galectins: key regulators of cellular homeostasis

Emerging evidence indicates that interactions between endogenous glycan-binding proteins and glycosylated receptors are integral for the control of immunological homeostasis.²¹ Although distinct lectin families have been described that contribute to modulate innate and adaptive immune responses, including C-type lectins and siglecs,²³ we will focus here on an ancient family of glycan-binding proteins, termed galectins, which have emerged as central regulators of fetomaternal tolerance.

Galectins specifically recognize complex glycan determinants on cell surface glycoproteins with relatively high affinity in the submicromolar range.²⁴ It has been demonstrated that it is the structure, number and density of glycan epitopes in multivalent glycoproteins, as well as the density of the glycoproteins expressed on the cell surface and the multivalent nature of some galectins, which together determine the avidity of lectin–glycan interactions.²⁵ Multivalent galectin-glycan complexes have been proposed to serve as scaffolds for organizing plasma membrane domains, which in turn modulate the signaling threshold of relevant surface glycoproteins including the T-cell receptor (TCR), B-cell receptor (BCR) and specific cytokine receptors.²⁶

Although galectins do not have the signal sequence required for the classical secretory pathway, most of them are externalized through a nonclassical mechanism and function in the extracellular milieu by interacting with a myriad of glycosylated ligands on the cell surface and extracellular matrix.^{24,27} However, these lectins also play roles inside the cells including modulation of signaling and splicing machineries.²⁸ According to their structure, galectins are classified into 'prototype' galectins (galectin-1, -2, -5, -7, -10, -11, -13, -14 and -15) which have one carbohydrate recognition domain (CRD) that can dimerize, 'tandem-repeat' galectins (galectin-4, -6, -8, -9 and -12) which contain two homologous CRDs in tandem in a single polypeptide chain and galectin-3, which is unique as it contains a CRD connected to a non-lectin N-terminal region that is responsible for oligomerization²⁹ (Fig. 1).

Although galectins were originally defined by their ability to recognize the disaccharide N-acetyllactosamine [Gal β (1–4)-GlcNAc; LacNAc], evidence obtained from glycan microarrays and frontal affinity chromatography analysis revealed substantial differences in the glycan-binding preferences of individual (a) Proto-type galectins (galectin-1, -2, -5, -7, -10, -11, -13, -14, -15)



(b) Chimaera-type galectin-3



(c) Tandem-repeat-type galectins (galectin-4, -6, -8, -9, -12)



Fig. 1 Structure and classification of the galectin family. Galectin are subdivided into three groups: those containing one CRD; galectin-3, which consists of unusual tandem repeats of proline- and glycine-rich short stretches fused onto the CRD; and those containing two distinct CRDs in tandem, connected by a linker peptide. Many galectins are either bivalent or multivalent with regard to their carbohydrate-binding activities: one CRD galectins often exist as dimers; galectin-3 forms pentamers upon binding to multivalent carbohydrates and two CRD galectins have two carbohydrate-binding sites.

members of the galectin family, which might explain differences in biological activity.^{27,30,31} Noteworthy, in addition to their regulated expression in immuneprivileged sites and inflammatory microenvironments, different intrinsic and extrinsic factors may control the biological activity of galectins including (i) their subcellular compartmentalization, (ii) their oligomerization status, (iii) their stability in reducing or oxidative microenvironments and (iv) the active remodeling of N- and O-glycans on target cells.^{27,32}

Interestingly, while some galectins are widely expressed, either constitutively or in an inducible fashion in immune cells and tissues, others have a more limited cellular distribution. For example, galectin-1 is considerably upregulated in activated T and B cells, inflammatory macrophages, tolerogenic DCs, decidual NK cells, and CD4⁺ CD25⁺ T_{Reg} cells, whereas galectin-10 expression appears to be restricted to eosinophils and CD4⁺ CD25⁺ T_{Reg} cells, galectin-5 is selectively expressed in rat reticulocytes, galectin-13 (so called 'placental protein 13' or PP13)

is expressed in human placenta, and galectin-15 appears to be preferentially expressed in ovine placental tissue.²⁷ In the next sections, we will integrate pioneer work and recent findings that contributed to our understanding of the expression and function of galectins during inflammation and pregnancy.

Galectins in pregnancy: decision-makers in fetomaternal tolerance, inflammation, and implantation

During the past few years, there has been increasing appreciation for the impact of galectin–glycan interactions in the control of immune cell homeostasis.²¹ Although galectins have been mostly studied in the context of pathologic responses including autoimmune pathology, microbial infection, or tumour growth, their biological roles in physiologic settings, including pregnancy, are just emerging. Here, we will dissect the general roles of individual members of the galectin family with particular emphasis on their function in the context of reproduction and pregnancy.

Galectin-1

Galectin-1, a one CRD galectin, functions as a monomer or a dimer and is highly sensitive to the redox fluctuations in inflammatory microenvironments.³² This endogenous lectin elicits a broad spectrum of immunoregulatory activities in vivo including the attenuation of autoimmune pathology,^{33–36} control of inflammation-induced neurodegeneration,³⁷ resolution of cardiac inflammation during myocardial infarction,³⁸ and promotion of tumour immune escape.^{39–43} Mice lacking galectin-1 show greater Th1 and Th17 responses, higher frequency of immunogenic dendritic cells, aberrant microglia activation and display more severe autoimmune pathology compared with their wild-type counterpart. 35, 37, 44, 45 Mechanistically, galectin-1 selectively kills Th1 and Th17 cell subsets through glycosylation-dependent mechanisms,³⁵ modulates T-cell adhesion and trafficking,^{46–48} promotes the expansion of FoxP3⁺ or FoxP3⁻ regulatory T (T_{reg}) cells,^{34,40,43} induces IL-10 secretion⁴⁹⁻⁵¹ and promotes the differentiation of IL-27⁺ IL-10⁺ tolerogenic dendritic cells.44

Research over the past few years has identified essential roles of galectin-1 during reproduction and pregnancy.^{52,53} Galectin-1 is abundant in the

American Journal of Reproductive Immunology 69 (2013) 369–382 © 2013 John Wiley & Sons A/S

placenta, in reproductive tracts and developing embryo of various species.^{54–61} Moreover, this 14.5kDa β-galactoside-binding lectin is substantially increased in FoxP3⁺ T_{reg} cells,⁶² tolerogenic dendritic cells,⁴⁴ activated T cells,⁶³ inflammatory macrophages,⁶⁴ and uterine NK cells.^{9,65} Within reproductive tracts and the fetoplacental unit, galectin-1 is mainly expressed in the late-secretory phase endometrium,⁵⁹ invasive extravillous trophoblast cells, maternal decidual cells and to some extent also in the villous mesenchyme of human first trimester and term placenta.^{66,67} Moreover, human embryos undergoing pre-implantation development express galectin-1 on the trophectoderm and inner cell mass.⁶¹ Galectin-1 is initially synthesized in the trophectoderm of expanded blastocysts immediately prior to implantation, suggesting that it may be involved in the attachment of the embryo to the uterine epithelium.⁶⁸

Although original studies showed that Gal-1-deficient mice developed normally and did not show any apparent phenotypic abnormality,⁶⁹ recent studies showed that mice lacking this endogenous lectin had higher rates of fetal loss compared with their WT counterpart in allogeneic, but not in syngeneic pregnancies.⁷⁰ Administration of recombinant galectin-1 prevented fetal loss and restored tolerance in vivo through expansion and recruitment of uterine tolerogenic dendritic cells which in turn promote the differentiation of IL-10-producing T_{reg} cells.⁷⁰ Accordingly, these effects were prevented in mice depleted of T_{reg} cells or in mice devoid of IL-10.⁷⁰ Moreover, galectin-1 confers immune privilege to the human trophoblast through modulation of a number of immunosuppressive pathways including induction of $FoxP3^+$ T_{reg} cells and suppression of Th1 cytokines⁶⁷ (Fig. 2). Targeted inhibition of galectin-1 expression through antibody-mediated blockade, disruption of the carbohydrate recognition domain using specific disaccharides or retroviralmediated siRNA strategies prevented these immunoregulatory effects.⁶⁷ Interestingly, secretion of galectin-1 by uNK cells has been shown to contribute to the generation of an immune-privileged environment by inducing apoptosis of decidual T cells.⁷¹ More recently, galectin-1 produced by uterine mast cells (uMC) has been identified as a decisive factor which positively influences spiral artery formation and placentation and prevents fetal abnormalities.⁷² Also, recent studies underscored a role for galectin-1 in promoting immune privilege by regulating expression of HLA-G on human trophoblast cells.⁶¹ Exogenous galectin-1 upregulated synthesis of the membrane-bound HLA-G isoforms (HLA-G1 and G2) in a cytotrophoblast cell line, whereas endogenous galectin-1 induced expression of the soluble isoforms (HLA-G5 and -G6),⁶¹ adding an additional galectin-1-mediated mechanism of immune tolerance.

Interestingly, a progesterone-galectin synergism has been reported in both mouse and human settings.^{67,70} This cross-talk was confirmed by phylogenetic footprinting studies highlighting steroidresponsive elements in the galectin-1 gene (LGALS1) that were gained after the emergence of mammalian placentation.⁷³ Supporting these findings, recent studies revealed that galectin-1 markedly reduces the incidence of resorptions in mice missing the immunophilin FK506-binding protein (FKB)52, a cochaperone that optimizes progesterone receptor (PR) signaling in the uterus.⁷⁴ Notably, galectin-1 was significantly downregulated in both $Fkbp52^{-/-}$ and $Pgr^{-/-}$ uteri compared with WT uteri, suggesting that uterine galectin-1 is an important downstream target progesterone-FKBP52-PR signaling in the of uterus.⁷⁴ In addition, galectin-1 has been shown to act as a regulator of progesterone and hCG synthesis by the chorionic carcinoma cell line BeWo.⁷⁵

In addition to its role in fetomaternal tolerance. galectin-1 has been implicated in non-immunological processes including trophoblast cell invasion and sincytium formation.^{76,77} In the choriocarcinomaderived BeWo cells, galectin-1 inhibits proliferation through binding to the Thomsen-Friedenreich disaccharide on MUC1 and modulation of the receptor tyrosine kinases rearranged during transfection (RET) and Janus kinase (JAK)2 as well as modulation of the vascular endothelial growth factor-3 (VEGFR3).⁷⁷ Moreover, galectin-1–N-glycan interactions have been proposed to play key roles in tumour neovascularization.^{78–81} Tissue expression of this lectin is upregulated during hypoxia through mechanisms involving hypoxia-inducible factor (HIF)-1 α or nuclear factor kappa B (NF- κ B).^{80,82} Although these effects were demonstrated in tumour settings, it is surmised that galectin-induced angiogenesis may also be important in reproductive biology and implantation where hypoxia is a critical regulatory factor. In addition, galectin-1 plays a key role in modulation of innate immunity, host-pathogen interactions, and platelet physiology.^{83–85} These effects may contribute to protective immunity and normal physiology during pregnancy. Thus, in

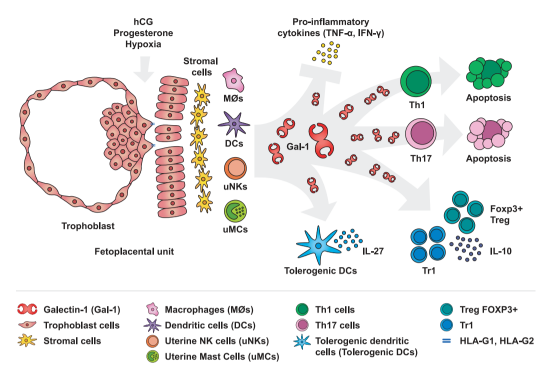


Fig. 2 The multifaceted roles of galectin-1 in pregnancy. Galectin-1 (Gal-1), an endogenous glycan-binding protein, is synthesized by trophoblast cells, stromal cells, uterine NK cells, uterine mast cells, CD4⁺ CD25⁺ regulatory T (T_{reg}) cells, macrophages (M ϕ), and dendritic cells (DC) within the fetoplacental unit. Its expression is regulated by several factors including progesterone, human chorionic gonadotrophin, and hypoxia. Through extracellular mechanisms, this lectin signals apoptosis of Th1 and Th17 cells, suppresses pro-inflammatory cytokine production, drives the differentiation of tolerogenic dendritic cells (tolerogenic DC), promotes the differentiation of FoxP3⁺ IL-10-producing T_{reg} cells and FoxP3⁻ IL-10-producing T_{reg} (Tr1) cells and induces HLA-G1 and G2 isoforms. In addition, this lectin contributes to trophoblast cell invasion, sincytium formation, placentation, and angiogenesis.

addition to the major role of galectin-1 as an essential component of the tolerogenic circuit activated during pregnancy, this endogenous lectin might also regulate other critical processes including placentation, angiogenesis, hemostasis, and control of pathogen invasion.

Galectin-2

Similar to galectin-1, galectin-2 is a 'prototype' galectin that exists as symmetric dimers containing two extended antiparallel β -sheets.⁸⁶ Recombinant galectin-2 induces apoptosis of mucosal T cells and displays anti-inflammatory activity in a model of colitis,⁸⁷ suggesting its major role in mucosal immunity. Human macrophages express significant amounts of galectin-2 which interacts with lymphotoxin α (LTA) and play a key role in the development of innate immunity and inflammation during myocardial infarction. Moreover, galectin-2 binds to epithelial cells at the E-cadherin/ β -catenin complex

and significantly enhances epithelial cell restoration and cellular migration *in vitro*,⁸⁸ suggesting its possible involvement in placentation-related events. However, the precise function of this lectin during pregnancy remains unexplored.

Galectin-3

Galectin-3 is the only 'chimera-type' member of the galectin family. It shows an extended N-terminal region composed of tandem repeats of short amino acid segments (a total of approximately 120 amino acids) connected to a C-terminal CRD. Unlike galec-tin-1 and -2, this CRD does not exist as homodimers in the crystal. By nuclear magnetic resonance spectroscopy, galectin-3 was found to exist as monomers in solution, and the N-terminal fragment has an unfolded, extended structure. However, galectin-3 can oligomerize in the presence of multivalent carbohydrate ligands and is capable of cross-linking glycans on the cell surface, thereby initiating

transmembrane signaling events and affecting various cellular functions.²⁸ This self-association property is dependent on the N-terminal region of the protein. As this region is sensitive to proteases, such collagenase and matrix metalloproteinases as (MMPs), the in vivo biological activities of galectin-3 are likely to be modulated by these enzymes.^{89,90} Compared with other galectins, intracellular functions of galectin-3 are more extensively documented. Intracellularly, galectin-3 functions as an antiapoptotic protein and induces pre-mRNA splicing.28 Extracellularly, galectin-3 binds in a carbohydratedependent manner to extracellular matrix proteins, including integrins, thereby influencing cell-cell adhesion, migration, cytokine production, and signaling.²⁸ In addition, galectin-3 can form lattices with selected cell surface glycans and modulate signaling, mobility, and endocytosis of different glycosylated receptors including the TCR, epidermal growth factor receptor and transforming growth factor.²⁶ Galectin-3 affects the differentiation and growth of various immune cells; it induces apoptosis of T cells and neutrophils and activates several lymphoid and myeloid cells, including mast cells, neutrophils, and T cells, resulting in production of reactive oxygen species, degranulation, and cytokine production.⁹¹ In vivo studies using galectin-3-deficient (Lgals $3^{-/-}$) mice supported the notion that galectin-3 amplifies inflammatory responses, suggesting that this endogenous lectin might play a role during early implantation, where proinflammatory cytokines, such as TNF and IFN- γ play decisive roles. In addition, a number of studies supported a role of galectin-3 in the modulation of Th1-Th2 cvtokine balance.²⁹ Galectin-3, probably the best studied member of the family, has also been proposed to play key roles in several physiologic and pathologic processes including angiogenesis,⁹² tumour development and progression,⁹³ and wound healing.⁹⁴ Collectively, these observations support a major role for galectin-3, acting either intracellularly or extracelllularly, at different stages of pregnancy development.

Galectin-3 mRNA and protein were higher in pregnant compared to non-pregnant mouse endometrium, and their maximum levels were reached on days 2 and 4, being localized at the luminal and glandular epithelia. Interestingly, the numbers of embryos implanted decreased substantially when galectin-3 was knocked down selectively in mouse endometrium.⁹⁵ Northern blot analysis of total RNA prepared from separated fetal and maternal compo-

nents of utero-placental complexes demonstrated different patterns of galectin-3 expression. Relative levels of galectin-3 mRNA peaked at midgestation in the implantation site and during the second half of gestation remain elevated in the placenta but declined in the uterus.96 Moreover, in bovine placentomas, galectin-3 was confined to uterine epithelial cells.⁹⁷ In human endometrium, galectin-3 expression increased significantly during the secretory phase of the menstrual cycle.⁵⁹ This lectin was observed in endometrial cells of the primary decidual zone immediately after implantation and at later stages of pregnancy in the decidua basalis and metrial gland and all trophoblastic elements of the placenta.⁵⁴ In fact, both galectin-1 and galectin-3 expression correlated with the differentiation pathways of trophoblasts. Galectin-3 was detected in all trophoblastic lineages including villous cytotrophoblasts and extravillous trophoblasts (trophoblastic cell columns, infiltrating trophoblasts, endovascular trophoblasts, and placental bed giant cells).⁵⁶ However, despite considerable evidence on the expression pattern of galectin-3 at the fetoplacental unit of several species, its physiologic roles during pregnancy remain obscure. Crider-Pirkle and colleagues identified cubilin, as an endogenous binding partner for galectin-3. Cubilin occurred in yolk sac epithelium throughout pregnancy but was endocytosed by uterine NK cells, apparently via interaction with galectin-3.98 Whether cubilin-galectin-3 interactions play any role in uNK cell physiology remains to be determined. In addition, in ovine placental tissue, galectin-1 and -3 have demonstrated antagonic immunomodulatory effects in vitro. While galectin-3 favored T-cell proliferation and activation, galectin-1 displayed T-cell inhibitory activity.^{57,99}

Galectin-4

Galectin-4, a tandem-repeat two CRD galectin, is predominantly expressed in mucosal tissues including the intestines.¹⁰⁰ The role of galectin-4 in lipid raft formation has been studied in different settings. Sulphatides with long-chain hydroxylated fatty acids, which are enriched in lipid rafts, were identified as high-affinity ligands for galectin-4. By interacting with sulphatides, galectin-4 fosters the clustering of lipid rafts and contributes to apical delivery of proteins.¹⁰¹ Galectin-4 regulates the development of inflammatory bowel disease by stimulating CD4⁺ T cells to produce IL-6.¹⁰² Recently, Nishida et al. identified an inducible colitis-associated glycome that can be identified through the binding of galectin-4 in local, but not systemic, memory CD4⁺ T cells exposed to intestinal inflammatory conditions. The inflammation-associated glycome represents an immature core 1-expressing O-glycan structure selectively expressed in inflamed intestinal tissue.¹⁰³ In addition, galectin-4 contributes to wound healing and regulates apoptosis of mucosal T cells.⁸⁸ These observations suggest a major role of galectin-4 in immune cell activation, inflammation, and expansion of memory T cells in mucosal tissue. Although scarce information is available regarding the function of galectin-4 in feto-placental tissue, its expression has been found to be restricted to uterine epithelial cells and blood vessel walls in the bovine endometrium.⁹⁷ Furthermore, during rat placentation, galectin-4 was downregulated in differentiated trophoblasts. Immunohistochemical analysis showed that galectin-4 was preferentially located in the maternal-fetal junctional zone, suggesting a possible role for this lectin in the promotion of trophoblast cell differentiation.104

Galectin-7

Galectin-7, a one CRD 'prototype' galectin, is found mainly in stratified squamous epithelium and its expression correlates with keratinocyte differentiation. The crystal structure of human galectin-7 revealed formation of complexes with galactose, lactose, and N-acetyllactosamine.¹⁰⁵ Galectin-7 is an early transcriptional target of the tumour suppressor protein p53, its expression is upregulated rapidly after ultraviolet B irradiation of epidermal keratinocytes and promotes apoptosis of these cells.¹⁰⁶ This lectin is also implicated in wound healing and epithelial cell migration.⁹⁴ However, the regulated expression and function of this lectin during pregnancy has not yet been explored.

Galectin-8

Three mRNAs encode six different galectin-8 isoforms; three with two CRDs in tandem and three with only one CRD.¹⁰⁷ Recombinant galectin-8 inhibits epithelial cell adhesion through binding to $\alpha 1\beta 3$ integrin. Binding of galectin-8 results in strong activation of integrin-mediated signaling which regulates cell adhesion and survival.¹⁰⁸ By cross-linking specific glycoconjugates on αM integrin, galectin-8

stimulates innate immunity through induction of superoxide production by neutrophils and control of neutrophil adhesion to extracellular matrix.¹⁰⁹ Moreover, both galectin-8 and galectin-4, which are expressed in mucosal tissue, recognize and kill human blood group antigen-expressing invading bacteria replacing recognition by specific antibodies.¹¹⁰ Recent findings demonstrated that galectin-8 is required for T-cell proliferation¹¹¹, platelet activation, ¹¹² and angiogenesis.¹¹³ Collectively, these findings suggest a possible role for galectin-8 in cellular migration, host-pathogen interactions, immunity, and neovascularization during pregnancy. Although galectin-8 is selectively expressed by villous and extravillous trophoblast in human placenta,¹¹⁴ the functional relevance of this lectin during pregnancy has not yet been ascertained.

Galectin-9

Three isoforms of galectin-9 differing in the lengths of their linker sequences have been identified.¹¹⁵ Similar to other two CRD galectins, galectin-9 has also been found in lipid rafts. Galectin-9 can either initiate or terminate adaptive immunity through modulation of T-cell survival or activation of antigen-presenting cells. In fact, recombinant galectin-9 induces selective apoptosis of Th1 cells and terminates autoimmune inflammation but, paradoxically, promotes dendritic cell maturation and macrophage activation through binding to the cell surface molecule TIM-3.^{116,117} However, TIM-3-independent mechanisms of immune cell activation, Th1/Th17 cytokine production, and apoptosis have also been described.¹¹⁸ Moreover, galectin-9 regulates glucose homeostasis by facilitating retention of GLUT-2 on the surface of pancreatic β cells.¹¹⁹ These observations suggest potential roles for galectin-9 in immunological and metabolic pathways during pregnancy.

Notably, galectin-9 is found in human endometrium, specifically in endometrial epithelial cells, but not in stromal cells or immune cells. It is expressed at very low concentrations during the proliferative phase and the early secretary phase and showed a sharp increase in the mid and late-secretory phase, the window of implantation, as well as in the deciduas.¹²⁰ Moreover, a recent study profiled expression of galectin-9 (*LGALS9*) isoforms at the fetomaternal interface in mouse and human pregnancy. Decidual galectin-9 was considerably altered in a mouse model of spontaneous abortion. The *LGALS9* D5 isoform selectively dampened IFN- γ production by decidual NK cells. In human patients, six *LGALS9* splice variants were detected, and a decrease in *LGALS9* D5/10 was associated with recurrent fetal loss.¹²¹ Given the inhibitory role of TIM-3 during microbial and tumour immunity, the role of galec-tin-9–TIM-3 interactions during gestation remains to be investigated.

Galectin-10

Galectin-10 has a single CRD, sharing only six of eight residues directly involved in lactose binding. Surprisingly, it has affinity for mannose but not for β -galactosides.¹²² In spite of its modest sequence homology, the overall structural fold of galectin-10 is very similar to other galectins, especially galectin-7.¹²² This lectin is constitutively expressed in eosinophils, basophils, and CD4⁺ CD25⁺ T_{reg} cells. Inhibition of endogenous galectin-10 by RNA interference abrogated the suppressive activity of T_{reg} cells.¹²³ These effects appear to be mediated via an intracellular mechanism that is not completely understood. Although unexplored in pregnancy settings, the selective expression of galectin-10 in T_{reg} cells suggests its potential role in fetomaternal tolerance in human pregnancies.

Galectin-12

While the N-terminal domain of galectin-12, a 'tandem-repeat' galectin, contains all sequence elements predicted to form the two β -sheets found in other galectins as well as conserved carbohydrate-interacting residues, its C-terminal domain shows considerable divergence from the consensus sequence.¹²⁴ Galectin-12 is over-represented in adipocytes compared to other tissues and has been proposed to play key roles in cell cycle regulation, adipocyte differentiation, and insulin sensitivity.^{125,126} The role of this 'tandem-repeat' lectin during pregnancy remains to be investigated.

Galectin-13

Placental protein 13 (PP13) was cloned from human term placenta. As sequence analyses, alignments, and computational modeling showed its conserved structural and functional homology to members of the galectin family, the protein was designated as galectin-13. Similar to human eosinophil galectin-10 but not other galectins, its weak lysophospholipase activity was confirmed.¹²⁷ It has been proposed that PP13 serves as a placental alarmin and undergoes lipid raft-associated subcellular redistribution in the syncytiotrophoblast.¹²⁸ Sugar-binding assays revealed that N-acetyl-lactosamine, mannose, and N-acetylglucosamine residues widely expressed in human placenta had the strongest binding affinity to both the purified and the recombinant PP13/galectin-13. PP13 was found to be localized to syncytiotrophoblasts in the chorionic villi and to occasional multinucleated luminal trophoblasts within converted decidual spiral arterioles.¹²⁷ Extracellular PP13 aggregates were found around decidual veins associated with immune cell-containing decidual zones of necrosis,¹²⁹ suggesting their role in trophoblast invasion, conversion of maternal spiral arterioles, and modulation of immunity. Exposure of cultured trophoblasts to PP13 resulted in depolarization of calcium ions, followed by liberation of linoleic and arachidonic acids from the trophoblast membrane, and subsequent elevation of prostacyclin and tromboxanes,¹³⁰ supporting its potential role in the control of trophoblast physiology.

Galectin-14

Ovine galectin-14 is selectively expressed in eosinophils and is released into lung and gastrointestinal tissues following allergen or parasite challenge. Galectin-14 is active in carbohydrate-binding assays and is involved in promoting cellular adhesion and changing mucus properties during parasite infection and allergies.¹³¹

The possibility that eosinophil-derived galectin-14 could modulate mucus properties during implantation or could influence host–pathogen interactions during pregnancy remains to be explored.

Galectin-15

Galectin-15 (also known as OVGAL11) is expressed specifically in the endometrial luminal and superficial glandular epithelia of the uterus in concert with blastocyst elongation during the pre-implantation period in the sheep, but it is also expressed in the uterus of goats and pigs.¹³² Carbohydate-binding assays revealed its ability to bind lactose and mannose sugars. Progesterone upregulates galectin-15 mRNA in the endometrial epithelium of pre-implantation conceptus.¹³³ In pregnant sheep, expression

appeared in the epithelia between days 10 and 12 and increased between days 12 and 16. Interestingly, galectin-15 forms crystals in the trophectoderm and regulates implantation and placentation by functioning as a heterophilic cell adhesion molecule between the conceptus trophectoderm and endometrium luminal epithelium.¹³⁴

Galectins in pathologic pregnancies

In human pregnancy, the implantation period is characterized by an initial inflammatory response that later shifts toward a tolerogenic state.¹³⁵ An inappropriate dysregulation of proinflammatory/ anti-inflammatory mediators may represent a major cause of fetal loss. Interestingly, a significant reduction in circulating galectin-1 and a higher prevalence of antigalectin-1 autoantibodies were found in sera from patients with recurrent spontaneous abortions (RSA) compared with fertile women, suggesting a possible mechanism to explain pregnancy loss in RSA patients.⁶⁷ Decreased levels of galectin-1 were also found in RSA patients who suffered subsequent miscarriages compared with normal pregnant women.⁶¹ Supporting these findings, proteomic analysis of placental villous showed the lower expression of galectin-1 in placental tissue from patients with early pregnancy loss compared with placental tissue from normal pregnant women.¹³⁶ However, the levels of serum galectin-1 and antigalectin-1 autoantibodies did not differ significantly between healthy pregnant women and patients with pre-eclampsia¹³⁷; yet, galectin-1 expression was markedly upregulated in placental tissue from patients with pre-eclampsia.¹³⁸ These results underscore possible differences in the expression of local versus systemic galectin-1 in different pathophysiologic settings. Moreover, increased galectin-1 mRNA expression was found in chorioamnionitis when analyzed in patients with pre-term pre-labor rupture of the membranes (PPROM). Here, its expression was mainly localized in amniotic epithelium, chorioamniotic mesodermal cells, and apoptotic bodies.¹³⁹ Importantly, both galectin-1 and galectin-3 were upregulated on placental tissue from patients with intrauterine growth retardation (IUGR), whereas only galectin-1 was upregulated in decidual tissue from pre-eclamptic placentas.¹⁴⁰ Moreover, immunoreactivity for both galectin-1 and galectin-3 was found to be increased in gestational trophoblastic disease (GTD), suggesting possible roles of these

lectins in the invasiveness of the transformed trophoblastic cells.58 On the other hand, PP13 (galectin-13) has been proposed to be a practical, individual predictor of women at risk of developing pre-eclampsia.141 Levels of PP13 in first-trimester pregnancies increased with gestational age in controls and pre-eclampsia cases, but at different rates. PP13 levels were decreased in women with preeclampsia compared to controls.¹⁴² Thus, changes in the expression and function of different galectins may contribute to the transition from normal to pathologic pregnancies. Alternatively, selective variations in galectin expression levels could serve as possible biomarkers of pregnancy complications including RSA, pre-eclampsia, IUGR, and GTD.

Conclusions and implications

The present review aims at integrating scattered information on the role of galectins, a family of endogenous glycan-binding proteins, in the development of normal and pathologic pregnancies. Galectins play diverse roles during the initiation, amplification and resolution of inflammatory response and can also regulate non-immunological processes including wound healing, cellular adhesion, migration, and angiogenesis. Although many of these functions have not yet been explored at the fetomaternal interface, galectins have been shown to play key roles in implantation and fetomaternal tolerance. Particularly, galectin-1 produced by trophoblast cells, uMC and uNK cells hierarchically regulates immunosuppression during pregnancy including differentiation of tolerogenic dendritic cells, expansion of T_{reg} cells, and apoptosis of decidual T cells, but also contributes to placentation through non-immunological mechanisms.

Future studies should examine the *in vivo* role of individual members of the galectin family (in addition to galectin-1) during pregnancy. Given the diverse range of galectin-deficient mice that are now available, it will be feasible to determine the impact of galectins and their glycosylated receptors in feto-maternal tolerance, implantation, and placentation. These data will set the basis to improve clinical outcomes in pathologic pregnancies and/or to provide novel biomarkers of successful gestation or early fetal loss. However, before galectin-based therapeutic agents can be extrapolated to clinical settings, a more thorough understanding of the mechanisms involved in galectin functions is essential. In this

regard, it will be critical to evaluate the results of side-by-side studies of the immunoregulatory activities of different members of the galectin family and to study the cross-talk between galectins and other established regulatory pathways including PD-L1, STAT-3, and IDO. Finally, given the complexity of galectin–glycan boundaries and the multiple parameters influencing these molecular interactions, further work is required, involving multidisciplinary approaches, to achieve a global comprehensive view of the role of endogenous galectins and their specific carbohydrate ligands during pregnancy.

Acknowledgements

Work in G.A.R's laboratory is supported by grants from the National Agency for Promotion of Science and Technology (Argentina; PICT 2010-870), Sales Foundation for Cancer (Argentina), University of Buenos Aires (Argentina), National Council for Scientific and Technical Investigation (CONICET, Argentina), Prostate Action (UK) and National Multiple Sclerosis Society (USA). A.G.B. is a postgraduate fellow of CONICET.

References

- 1 Terness P, Kallikourdis M, Betz AG, Rabinovich GA, Saito S, Clark DA: Tolerance signaling molecules and pregnancy: IDO, galectins, and the renaissance of regulatory T cells. *Am J Reprod Immunol* 2007; 58:238–254.
- 2 Chaouat G, Petitbarat M, Dubanchet S, Rahmati M, Ledée N: Tolerance to the foetal allograft? *Am J Reprod Immunol* 2010; 63:624–636.
- 3 Blois SM, Joachim R, Kandil J, Margni R, Tometten M, Klapp BF, Arck PC: Depletion of CD8+ cells abolishes the pregnancy protective effect of progesterone substitution with dydrogesterone in mice by altering the Th1/Th2 cytokine profile. *J Immunol* 2004; 172:5893–5899.
- 4 Chaouat G: The Th1/Th2 paradigm: still important in pregnancy? Semin Immunopathol 2007; 29:95–113.
- 5 Aluvihare VR, Kallikourdis M, Betz AG: Regulatory T cells mediate maternal tolerance to the fetus. *Nat Immunol* 2004; 5:266–271.
- 6 Zenclussen AC, Gerlof K, Zenclussen ML, Sollwedel A, Bertoja AZ, Ritter T, Kotsch K, Leber J, Volk HD: Abnormal T-cell reactivity against paternal antigens in spontaneous abortion: adoptive transfer of pregnancy-induced CD4+ CD25+ T regulatory cells prevents fetal rejection in a murine abortion model. *Am J Pathol* 2005; 166:811–822.
- 7 Kallikourdis M, Andersen KG, Welch KA, Betz AG: Alloantigenenhanced accumulation of CCR5+ 'effector' regulatory T cells in the gravid uterus. *Proc Natl Acad Sci U S A*. 2007; 104:594–599.
- 8 Fainboim L, Arruvito L: Mechanisms involved in the expansion of Tregs during pregnancy: role of IL-2/STAT5 signalling. J Reprod Immunol 2011; 88:93–98.

- 9 Koopman LA, Kopcow HD, Rybalov B, Boyson JE, Orange JS, Schatz F, Masch R, Lockwood CJ, Schachter AD, Park PJ, Strominger JL: Human decidual natural killer cells are a unique NK cell subset with immunomodulatory potential. *J Exp Med* 2003; 198:1201–1212.
- 10 Guleria I, Khosroshahi A, Ansari MJ, Habicht A, Azuma M, Yagita H, Noelle RJ, Coyle A, Mellor AL, Khoury SJ, Sayegh MH: A critical role for the programmed death ligand 1 in fetomaternal tolerance. *J Exp Med* 2005; 202:231–237.
- 11 Kayisli UA, Selam B, Guzeloglu-Kayisli O, Demir R, Arici A: Human chorionic gonadotropin contributes to maternal immunotolerance and endometrial apoptosis by regulating Fas-Fas ligand system. J Immunol 2003; 171:2305–2313.
- 12 Phillips TA, Ni J, Pan G, Ruben SM, Wei YF, Pace JL, Hunt JS: TRAIL (Apo-2L) and TRAIL receptors in human placentas: implications for immune privilege. *J Immunol* 1999; 162:6053– 6059.
- 13 Tariverdian N, Theoharides TC, Siedentopf F, Gutiérrez G, Jeschke U, Rabinovich GA, Blois SM, Arck PC: Neuroendocrine-immune disequilibrium and endometriosis: an interdisciplinary approach. *Semin Immunopathol* 2007; 29:193–210.
- 14 Arck PC, Gilhar A, Bienenstock J, Paus R: The alchemy of immune privilege explored from a neuroimmunological perspective. *Curr Opin Pharmacol* 2008; 8:480–489.
- 15 Fraccaroli L, Alfieri J, Larocca L, Calafat M, Roca V, Lombardi E, Ramhorst R, Leirós CP: VIP modulates the pro-inflammatory maternal response, inducing tolerance to trophoblast cells. *Br J Pharmacol* 2009; 156:116–126.
- 16 Arck P, Hansen PJ, Mulac Jericevic B, Piccinni MP, Szekeres-Bartho J: Progesterone during pregnancy: endocrine-immune cross talk in mammalian species and the role of stress. *Am J Reprod Immunol* 2007; 58:268–279.
- 17 Arruvito L, Giulianelli S, Flores AC, Paladino N, Barboza M, Lanari C, Fainboim L: NK cells expressing a progesterone receptor are susceptible to progesterone-induced apoptosis. *J Immunol* 2008; 180:5746–5753.
- 18 Zenclussen AC, Schumacher A, Zenclussen ML, Wafula P, Volk HD: Immunology of pregnancy: cellular mechanisms allowing fetal survival within the maternal uterus. *Expert Rev Mol Med* 2007; 9:1– 14.
- 19 Nakamura K, Sheps S, Arck PC: Stress and reproductive failure: past notions, present insights and future directions. *J Assist Reprod Genet* 2008; 25:47–62.
- 20 Oreshkova T, Dimitrov R, Mourdjeva M: A cross-talk of decidual stromal cells, trophoblast, and immune cells: a prerequisite for the success of pregnancy. *Am J Reprod Immunol* 2012; 68:366–373.
- 21 Rabinovich GA, Croci DO: Regulatory circuits mediated by lectinglycan interactions in autoimmunity and cancer. *Immunity* 2012; 36:322–335.
- 22 Meseguer M, Pellicer A, Simón C: MUC1 and endometrial receptivity. *Mol Hum Reprod* 1998; 4:1089–1098.
- 23 Toscano MA, Ilarregui JM, Bianco GA, Campagna L, Croci DO, Salatino M, Rabinovich GA: Dissecting the pathophysiologic role of endogenous lectins: glycan-binding proteins with cytokine-like activity? *Cytokine Growth Factor Rev* 2007; 18:57–71.
- 24 Rabinovich GA, Toscano MA, Jackson SS, Vasta GR: Functions of cell surface galectin-glycoprotein lattices. *Curr Opin Struct Biol* 2007; 17:513–520.
- 25 Dam TK, Brewer CF: Lectins as pattern recognition molecules: the effects of epitope density in innate immunity. *Glycobiology* 2010; 20:270–279.

- 26 Dennis JW, Nabi IR, Demetriou M: Metabolism, cell surface organization, and disease. *Cell* 2009; 139:1229–1241.
- 27 Rabinovich GA, Toscano MA: Turning 'sweet' on immunity: galectin-glycan interactions in immune tolerance and inflammation. *Nat Rev Immunol* 2009; 9:338–352.
- 28 Liu FT, Rabinovich GA: Galectins: regulators of acute and chronic inflammation. *Ann N Y Acad Sci* 2010; 1183:158–182.
- 29 Yang RY, Rabinovich GA, Liu FT: Galectins: structure, function and therapeutic potential. *Expert Rev Mol Med* 2008; 10:e17.
- 30 Hirabayashi J, Hashidate T, Arata Y, Nishi N, Nakamura T, Hirashima M, Urashima T, Oka T, Futai M, Muller WE, Yagi F, Kasai K: Oligosaccharide specificity of galectins: a search by frontal affinity chromatography. *Biochim Biophys Acta* 2002; 1572:232–254.
- 31 Stowell SR, Arthur CM, Mehta P, Slanina KA, Blixt O, Leffler H, Smith DF, Cummings RD: Galectin-1, -2, and -3 exhibit differential recognition of sialylated glycans and blood group antigens. *J Biol Chem* 2008; 283:10109–10123.
- 32 Di Lella S, Sundblad V, Cerliani JP, Guardia CM, Estrin DA, Vasta GR, Rabinovich GA: When galectins recognize glycans: from biochemistry to physiology and back again. *Biochemistry* 2011; 50:7842–7857.
- 33 Rabinovich GA, Daly G, Dreja H, Tailor H, Riera CM, Hirabayashi J, Chernajovsky Y: Recombinant galectin-1 and its genetic delivery suppress collagen-induced arthritis via T cell apoptosis. *J Exp Med* 1999; 190:385–398.
- 34 Toscano MA, Commodaro AG, Ilarregui JM, Bianco GA, Liberman A, Serra HM, Hirabayashi J, Rizzo LV, Rabinovich GA: Galectin-1 suppresses autoimmune retinal disease by promoting concomitant Th2- and T regulatory-mediated anti-inflammatory responses. *J Immunol* 2006; 176:6323–6332.
- 35 Toscano MA, Bianco GA, Ilarregui JM, Croci DO, Correale J, Hernandez JD, Zwirner NW, Poirier F, Riley EM, Baum LG, Rabinovich GA: Differential glycosylation of TH1, TH2 and TH-17 effector cells selectively regulates susceptibility to cell death. *Nat Immunol* 2007; 8:825–834.
- 36 Perone MJ, Bertera S, Shufesky WJ, Divito SJ, Montecalvo A, Mathers AR, Larregina AT, Pang M, Seth N, Wucherpfennig KW, Trucco M, Baum LG, Morelli AE: Suppression of autoimmune diabetes by soluble galectin-1. *J Immunol* 2009; 182:2641–2653.
- 37 Starossom SC, Mascanfroni ID, Imitola J, Cao L, Raddassi K, Hernandez SF, Bassil R, Croci DO, Cerliani JP, Delacour D, Wang Y, Elyaman W, Khoury SJ, Rabinovich GA: Galectin-1 deactivates classically activated microglia and protects from inflammationinduced neurodegeneration. *Immunity* 2012; 37:249–263.
- 38 Seropian IM, Cerliani JP, Toldo S, Van Tassell BW, Ilarregui JM, González GE, Matoso M, Salloum FN, Melchior R, Gelpi RJ, Stupirski JC, Benatar A, Gómez KA, Morales C, Abbate A, Rabinovich GA: Galectin-1 controls cardiac inflammation and ventricular remodeling during acute myocardial infarction. *Am J Pathol* 2013; 182:29–40.
- 39 Rubinstein N, Alvarez M, Zwirner NW, Toscano MA, Ilarregui JM, Bravo A, Mordoh J, Fainboim L, Podhajcer OL, Rabinovich GA: Targeted inhibition of galectin-1 gene expression in tumor cells results in heightened T cell-mediated rejection; a potential mechanism of tumor-immune privilege. *Cancer Cell* 2004; 5:241– 251.
- 40 Juszczynski P, Ouyang J, Monti S, Rodig SJ, Takeyama K, Abramson J, Chen W, Kutok JL, Rabinovich GA, Shipp MA: The AP1-dependent secretion of galectin-1 by Reed Sternberg cells fosters immune privilege in classical Hodgkin lymphoma. *Proc Natl Acad Sci U S A*. 2007; 104:13134–13139.

American Journal of Reproductive Immunology **69** (2013) 369–382 © 2013 John Wiley & Sons A/S

- 41 Banh A, Zhang J, Cao H, Bouley DM, Kwok S, Kong C, Giaccia AJ, Koong AC, Le QT: Tumor galectin-1 mediates tumor growth and metastasis through regulation of T-cell apoptosis. *Cancer Res* 2011; 71:4423–4431.
- 42 Soldati R, Berger E, Zenclussen AC, Jorch G, Lode HN, Salatino M, Rabinovich GA, Fest S: Neuroblastoma triggers an immunoevasive program involving galectin-1-dependent modulation of T cell and dendritic cell compartments. *Int J Cancer* 2012; 131:1131–1141.
- 43 Dalotto-Moreno T, Croci DO, Cerliani JP, Martinez-Allo VC, Dergan-Dylon S, Mendez Huergo SP, Stupirski JC, Mazal D, Osinaga E, Toscano MA, Sundblad V, Rabinovich GA, Salatino M: Targeting galectin-1 overcomes breast cancer associated immunosuppression and prevents metastatic disease. *Cancer Res* 2013; 73:1107–1117.
- 44 Ilarregui JM, Croci DO, Bianco GA, Toscano MA, Salatino M, Vermeulen ME, Geffner JR, Rabinovich GA: Tolerogenic signals delivered by dendritic cells to T cells through a galectin-1-driven immunoregulatory circuit involving interleukin 27 and interleukin 10. *Nat Immunol* 2009; 10:981–991.
- 45 Liu SD, Lee S, La Cava A, Motran CC, Hahn BH, Miceli MC: Galectin-1-induced down-regulation of T lymphocyte activation protects (NZB × NZW) F1 mice fromlupus-like disease. *Lupus* 2011; 20:473–484.
- 46 Rabinovich GA, Ariel A, Hershkoviz R, Hirabayashi J, Kasai KI, Lider O: Specific inhibition of T-cell adhesion to extracellular matrix and proinflammatory cytokine secretion by human recombinant galectin-1. *Immunology* 1999; 97:100–106.
- 47 Norling LV, Sampaio AL, Cooper D, Perretti M: Inhibitory control of endothelial galectin-1 on in vitro and in vivo lymphocyte trafficking. *FASEB J* 2008; 22:682–690.
- 48 He J, Baum LG: Galectin interactions with extracellular matrix and effects on cellular function. *Methods Enzymol* 2006; 417:247–256.
- 49 van der Leij J, van den Berg A, Harms G, Eschbach H, Vos H, Zwiers P, van Weeghel R, Groen H, Poppema S, Visser L: Strongly enhanced IL-10 production using stable galectin-1 homodimers. *Mol Immunol* 2007; 44:506–513.
- 50 Stowell SR, Qian Y, Karmakar S, Koyama NS, Dias-Baruffi M, Leffler H, McEver RP, Cummings RD: Differential roles of galectin-1 and galectin-3 in regulating leukocyte viability and cytokine secretion. *J Immunol* 2008; 180:3091–3102.
- 51 Cedeno-Laurent F, Opperman M, Barthel SR, Kuchroo VK, Dimitroff CJ: Galectin-1 triggers an immunoregulatory signature in Th cells functionally defined by IL-10 expression. *J Immunol* 2012; 188:3127–3137.
- 52 Rabinovich GA, Ilarregui JM: Conveying glycan information into T-cell homeostatic programs: a challenging role for galectin-1 in inflammatory and tumor microenvironments. *Immunol Rev* 2009; 230:144–159.
- 53 Than NG, Romero R, Kim CJ, McGowen MR, Papp Z, Wildman DE: Galectins: guardians of eutherian pregnancy at the maternal-fetal interface. *Trends Endocrinol Metab* 2012; 23:23–31.
- 54 Phillips B, Knisley K, Weitlauf KD, Dorsett J, Lee V, Weitlauf H: Differential expression of two beta-galactoside-binding lectins in the reproductive tracts of pregnant mice. *Biol Reprod* 1996; 55:548– 558.
- 55 Dettin L, Rubinstein N, Aoki A, Rabinovich GA, Maldonado CA: Regulated expression and ultrastructural localization of galectin-1, a proapoptotic beta-galactoside-binding lectin, during spermatogenesis in rat testis. *Biol Reprod* 2003; 68:51–59.
- 56 Maquoi E, van den Brûle FA, Castronovo V, Foidart JM: Changes in the distribution pattern of galectin-1 and galectin-3 in human

placenta correlates with the differentiation pathways of trophoblasts. *Placenta* 1997; 18:433–439.

- 57 Iglesias MM, Rabinovich GA, Ivanovic V, Sotomayor C, Wolfenstein-Todel C: Galectin-1 from ovine placenta–amino-acid sequence, physicochemical properties and implications in T-cell death. *Eur J Biochem* 1998; 252:400–407.
- 58 Bozić M, Petronijević M, Milenković S, Atanacković J, Lazić J, Vićovac LJ: Galectin-1 and galectin-3 in the trophoblast of the gestational trophoblastic disease. *Placenta* 2004; 25:797–802.
- 59 von Wolff M, Wang X, Gabius HJ, Strowitzki T: Galectin fingerprinting in human endometrium and decidua during the menstrual cycle and in early gestation. *Mol Hum Reprod* 2005; 11:189–194.
- 60 Dong M, Ding G, Zhou J, Wang H, Zhao Y, Huang H: The effect of trophoblasts on T lymphocytes: possible regulatory effector molecules–a proteomic analysis. *Cell Physiol Biochem* 2008; 21:463– 472.
- 61 Tirado-González I, Freitag N, Barrientos G, Shaikly V, Nagaeva O, Strand M, Kjellberg L, Klapp BF, Mincheva-Nilsson L, Cohen M, Blois SM: Galectin-1 influences trophoblast immune evasion and emerges as a predictive factor for the outcome of pregnancy. *Mol Hum Reprod* 2013; 19:43–53.
- 62 Garín MI, Chu CC, Golshayan D, Cernuda-Morollón E, Wait R, Lechler RI: Galectin-1: a key effector of regulation mediated by CD4+ CD25+ T cells. *Blood* 2007; 109:2058–2065.
- 63 Fuertes MB, Molinero LL, Toscano MA, Ilarregui JM, Rubinstein N, Fainboim L, Zwirner NW, Rabinovich GA: Regulated expression of galectin-1 during T-cell activation involves Lck and Fyn kinases and signaling through MEK1/ERK, p38 MAP kinase and p70S6 kinase. *Mol Cell Biochem* 2004; 267:177–185.
- 64 Rabinovich GA, Castagna L, Landa C, Riera CM, Sotomayor C: Regulated expression of a 16-kd galectin-like protein in activated rat macrophages. *J Leukoc Biol* 1996; 59:363–370.
- 65 Karimi K, Arck PC: Natural Killer cells: keepers of pregnancy in the turnstile of the environment. *Brain Behav Immun* 2010; 24:339 –347.
- 66 Bevan BH, Kilpatrick DC, Liston WA, Hirabayashi J, Kasai K: Immunohistochemical localization of a beta-D-galactoside-binding lectin at the human maternofetal interface. *Histochem J* 1994; 26:582–586.
- 67 Ramhorst RE, Giribaldi L, Fraccaroli L, Toscano MA, Stupirski JC, Romero MD, Durand ES, Rubinstein N, Blaschitz A, Sedlmayr P, Genti-Raimondi S, Fainboim L, Rabinovich GA: Galectin-1 confers immune privilege to human trophoblast: implications in recurrent fetal loss. *Glycobiology* 2012; 22:1374–1386.
- 68 Poirier F, Timmons PM, Chan CT, Guénet JL, Rigby PW: Expression of the L14 lectin during mouse embryogenesis suggests multiple roles during pre- and post-implantation development. *Development* 1992; 115:143–155.
- 69 Poirier F, Robertson EJ: Normal development of mice carrying a null mutation in the gene encoding the L14 S-type lectin. *Development* 1993; 119:1229–1236.
- 70 Blois SM, Ilarregui JM, Tometten M, Garcia M, Orsal AS, Cordo-Russo R, Toscano MA, Bianco GA, Kobelt P, Handjiski B, Tirado I, Markert UR, Klapp BF, Poirier F, Szekeres-Bartho J, Rabinovich GA, Arck PC: A pivotal role for galectin-1 in fetomaternal tolerance. *Nat Med* 2007; 13:1450–1457.
- 71 Kopcow HD, Rosetti F, Leung Y, Allan DS, Kutok JL, Strominger JL: T cell apoptosis at the maternal-fetal interface in early human pregnancy, involvement of galectin-1. *Proc Natl Acad Sci USA* 2008; 105:18472–18477.

- 72 Woidacki K, Popovic M, Metz M, Schumacher A, Linzke N, Teles A, Poirier F, Fest S, Jensen F, Rabinovich GA, Maurer M, Zenclussen AC: Mast cells rescue implantation defects caused by ckit deficiency. *Cell Death Dis* 2013; 4:e462.
- 73 Than NG, Romero R, Erez O, Weckle A, Tarca AL, Hotra J, Abbas A, Han YM, Kim SS, Kusanovic JP, Gotsch F, Hou Z, Santolaya-Forgas J, Benirschke K, Papp Z, Grossman LI, Goodman M, Wildman DE: Emergence of hormonal and redox regulation of galectin-1 in placental mammals: implication in maternal-fetal immune tolerance. *Proc Natl Acad Sci USA* 2008; 105:15819–15824.
- 74 Hirota Y, Burnum KE, Acar N, Rabinovich GA, Daikoku T, Dey SK: Galectin-1 markedly reduces the incidence of resorptions in mice missing immunophilin FKBP52. *Endocrinology* 2012; 153:2486 –2493.
- 75 Jeschke U, Reimer T, Bergemann C, Wiest I, Schulze S, Friese K, Walzel H: Binding of galectin-1 (gal-1) on trophoblast cells and inhibition of hormone production of trophoblast tumor cells in vitro by gal-1. *Histochem Cell Biol* 2004; 121:501–508.
- 76 Kolundžić N, Bojić-Trbojević Ž, Kovačević T, Stefanoska I, Kadoya T, Vićovac L: Galectin-1 is part of human trophoblast invasion machinery–a functional study in vitro. *PLoS ONE* 2011; 6:e28514.
- 77 Fischer I, Weber M, Kuhn C, Fitzgerald JS, Schulze S, Friese K, Walzel H, Markert UR, Jeschke U: Is galectin-1 a trigger for trophoblast cell fusion?: the MAP-kinase pathway and syncytium formation in trophoblast tumour cells BeWo. *Mol Hum Reprod* 2011; 17:747–757.
- 78 Thijssen VL, Barkan B, Shoji H, Aries IM, Mathieu V, Deltour L, Hackeng TM, Kiss R, Kloog Y, Poirier F, Griffioen AW: Tumor cells secrete galectin-1 to enhance endothelial cell activity. *Cancer Res* 2010; 70:6216–6224.
- 79 Mathieu V, de Lassalle EM, Toelen J, Mohr T, Bellahcène A, Van Goietsenoven G, Verschuere T, Bouzin C, Debyser Z, De Vleeschouwer S, Van Gool S, Poirier F, Castronovo V, Kiss R, Feron O: Galectin-1 in melanoma biology and related neoangiogenesis processes. J Invest Dermatol 2012; 132:2245–2254.
- 80 Croci DO, Salatino M, Rubinstein N, Cerliani JP, Cavallin LE, Leung HJ, Ouyang J, Ilarregui JM, Toscano MA, Domaica CI, Croci MC, Shipp MA, Mesri EA, Albini A, Rabinovich GA: Disrupting galectin-1 interactions with N-glycans suppresses hypoxia-driven angiogenesis and tumorigenesis in Kaposi's sarcoma. J Exp Med 2012; 209:1985–2000.
- 81 Laderach DJ, Gentilini LD, Giribaldi L, Delgado VC, Nugnes L, Croci DO, Al Nakouzi N, Sacca P, Casas G, Mazza O, Shipp MA, Vazquez E, Chauchereau A, Kutok JL, Rodig SJ, Elola MT, Compagno D, Rabinovich GA: A unique galectin signature in human prostate cancer progression suggests galectin-1 as a key target for treatment of advanced disease. *Cancer Res* 2013; 73:86– 96.
- 82 Zhao XY, Zhao KW, Jiang Y, Zhao M, Chen GQ: Synergistic induction of galectin-1 by CCAAT/enhancer binding protein alpha and hypoxia-inducible factor 1alpha and its role in differentiation of acute myeloid leukemic cells. *J Biol Chem* 2011; 286:36808– 36819.
- 83 Cooper D, Norling LV, Perretti M: Novel insights into the inhibitory effects of Galectin-1 on neutrophil recruitment under flow. J Leukoc Biol 2008; 83:1459–1466.
- 84 Vasta GR, Ahmed H, Nita-Lazar M, Banerjee A, Pasek M, Shridhar S, Guha P, Fernández-Robledo JA: Galectins as self/non-self recognition receptors in innate and adaptive immunity: an unresolved paradox. *Front Immunol* 2012; 3:199.

- 85 Pacienza N, Pozner RG, Bianco GA, D'Atri LP, Croci DO, Negrotto S, Malaver E, Gómez RM, Rabinovich GA, Schattner M: The immunoregulatory glycan-binding protein galectin-1 triggers human platelet activation. *FASEB J* 2008; 22:1113–1123.
- 86 Lobsanov YD, Gitt MA, Leffler H, Barondes SH, Rini JM: X-ray crystal structure of the human dimeric S-Lac lectin, L-14-II, in complex with lactose at 2.9-A resolution. *J Biol Chem* 1993; 268:27034–27038.
- 87 Paclik D, Berndt U, Guzy C, Dankof A, Danese S, Holzloehner P, Rosewicz S, Wiedenmann B, Wittig BM, Dignass AU, Sturm A: Galectin-2 induces apoptosis of lamina propria T lymphocytes and ameliorates acute and chronic experimental colitis in mice. *J Mol Med (Berl)* 2008; 86:1395–1406.
- 88 Paclik D, Lohse K, Wiedenmann B, Dignass AU, Sturm A: Galectin-2 and -4, but not galectin-1, promote intestinal epithelial wound healing in vitro through a TGF-beta-independent mechanism. *Inflamm Bowel Dis* 2008; 14:1366–1372.
- 89 Ochieng J, Green B, Evans S, James O, Warfield P: Modulation of the biological functions of galectin-3 by matrix metalloproteinases. *Biochim Biophys Acta* 1998; 1379:97–106.
- 90 Pasquini LA, Millet V, Hoyos HC, Giannoni JP, Croci DO, Marder M, Liu FT, Rabinovich GA, Pasquini JM: Galectin-3 drives oligodendrocyte differentiation to control myelin integrity and function. *Cell Death Differ* 2011; 18:1746–1756.
- 91 Sundblad V, Croci DO, Rabinovich GA: Regulated expression of galectin-3, a multifunctional glycan-binding protein, in haematopoietic and non-haematopoietic tissues. *Histol Histopathol* 2011; 26:247–265.
- 92 Markowska AI, Liu FT, Panjwani N: Galectin-3 is an important mediator of VEGF- and bFGF-mediated angiogenic response. J Exp Med 2010; 207:1981–1993.
- 93 Liu FT, Rabinovich GA: Galectins as modulators of tumour progression. Nat Rev Cancer 2005; 5:29–41.
- 94 Cao Z, Said N, Amin S, Wu HK, Bruce A, Garate M, Hsu DK, Kuwabara I, Liu FT, Panjwani N: Galectins-3 and -7, but not galectin-1, play a role in re-epithelialization of wounds. *J Biol Chem* 2002; 277:42299–42305.
- 95 Yang H, Lei C, Zhang W: Expression of galectin-3 in mouse endometrium and its effect during embryo implantation. *Reprod Biomed Online* 2012; 24:116–122.
- 96 Lee VH, Lee AB, Phillips EB, Roberts JK, Weitlauf HM: Spatiotemporal pattern for expression of galectin-3 in the murine uteroplacental complex: evidence for differential regulation. *Biol Reprod* 1998; 58:1277–1282.
- 97 Froehlich R, Hambruch N, Haeger JD, Dilly M, Kaltner H, Gabius HJ, Pfarrer C: Galectin fingerprinting detects differences in expression profiles between bovine endometrium and placentomes as well as early and late gestational stages. *Placenta* 2012; 33:195–201.
- 98 Crider-Pirkle S, Billingsley P, Faust C, Hardy DM, Lee V, Weitlauf H: Cubilin, a binding partner for galectin-3 in the murine uteroplacental complex. J Biol Chem 2002; 277:15904–15912.
- 99 Iglesias MM, Rabinovich GA, Ambrosio AL, Castagna LF, Sotomayor CE, Wolfenstein-Todel C: Purification of galectin-3 from ovine placenta: developmentally regulated expression and immunological relevance. *Glycobiology* 1998; 8:59–65.
- 100 Troncoso MF, Elola MT, Croci DO, Rabinovich GA: Integrating structure and function of 'tandem-repeat' galectins. *Front Biosci* (Schol Ed) 2012; 4:864–887.
- 101 Delacour D, Gouyer V, Zanetta JP, Drobecq H, Leteurtre E, Grard G, Moreau-Hannedouche O, Maes E, Pons A, André S, Le Bivic A,

American Journal of Reproductive Immunology **69** (2013) 369–382 © 2013 John Wiley & Sons A/S Gabius HJ, Manninen A, Simons K, Huet G: Galectin-4 and sulfatides in apical membrane trafficking in enterocyte-like cells. *J Cell Biol* 2005; 169:491–501.

- 102 Hokama A, Mizoguchi E, Sugimoto K, Shimomura Y, Tanaka Y, Yoshida M, Rietdijk ST, de Jong YP, Snapper SB, Terhorst C, Blumberg RS, Mizoguchi A: Induced reactivity of intestinal CD4 (+) T cells with an epithelial cell lectin, galectin-4, contributes to exacerbation of intestinal inflammation. *Immunity* 2004; 20:681– 693.
- 103 Nishida A, Nagahama K, Imaeda H, Ogawa A, Lau CW, Kobayashi T, Hisamatsu T, Preffer FI, Mizoguchi E, Ikeuchi H, Hibi T, Fukuda M, Andoh A, Blumberg RS, Mizoguchi A: Inducible colitis-associated glycome capable of stimulating the proliferation of memory CD4+ T cells. J Exp Med 2012; 209:2383–2394.
- 104 Arikawa T, Simamura E, Shimada H, Nishi N, Tatsuno T, Ishigaki Y, Tomosugi N, Yamashiro C, Hata T, Takegami T, Mogami H, Yamaguchi K, Nakamura T, Otani H, Hatta T, Shoji H: Expression pattern of Galectin 4 in rat placentation. *Placenta* 2012; 33:885– 887.
- 105 Leonidas DD, Vatzaki EH, Vorum H, Celis JE, Madsen P, Acharya KR: Structural basis for the recognition of carbohydrates by human galectin-7. *Biochemistry* 1998; 37:13930–13940.
- 106 Kuwabara I, Kuwabara Y, Yang RY, Schuler M, Green DR, Zuraw BL, Hsu DK, Liu FT: Galectin-7 (PIG1) exhibits pro-apoptotic function through JNK activation and mitochondrial cytochrome c release. *J Biol Chem* 2002; 277:3487–3497.
- 107 Bidon-Wagner N, Le Pennec JP: Human galectin-8 isoforms and cancer. *Glycoconj J* 2004; 19:557–563.
- 108 Zick Y, Eisenstein M, Goren RA, Hadari YR, Levy Y, Ronen D: Role of galectin-8 as a modulator of cell adhesion and cell growth. *Glycoconj J* 2004; 19:517–526.
- 109 Nishi N, Shoji H, Seki M, Itoh A, Miyanaka H, Yuube K, Hirashima M, Nakamura T: Galectin-8 modulates neutrophil function via interaction with integrin alphaM. *Glycobiology* 2003; 13:755–763.
- 110 Stowell SR, Arthur CM, Dias-Baruffi M, Rodrigues LC, Gourdine JP, Heimburg-Molinaro J, Ju T, Molinaro RJ, Rivera-Marrero C, Xia B, Smith DF, Cummings RD: Innate immune lectins kill bacteria expressing blood group antigen. *Nat Med* 2010; 16:295–301.
- 111 Cattaneo V, Tribulatti MV, Campetella O: Galectin-8 tandemrepeat structure is essential for T-cell proliferation but not for costimulation. *Biochem J* 2011; 434:153–160.
- 112 Romaniuk MA, Tribulatti MV, Cattaneo V, Lapponi MJ, Molinas FC, Campetella O, Schattner M: Human platelets express and are activated by galectin-8. *Biochem J* 2010; 432:535–547.
- 113 Delgado VM, Nugnes LG, Colombo LL, Troncoso MF, Fernández MM, Malchiodi EL, Frahm I, Croci DO, Compagno D, Rabinovich GA, Wolfenstein-Todel C, Elola MT: Modulation of endothelial cell migration and angiogenesis: a novel function for the "tandem-repeat" lectin galectin-8. *FASEB J* 2011; 25:242– 254.
- 114 Kolundžić N, Bojić-Trbojević Z, Radojčić Lj, Petronijević M, Vićovac LJ: Galectin-8 is expressed by villous and extravillous trophoblast of the human placenta. *Placenta* 2011; 32:909–911.
- 115 Chabot S, Kashio Y, Seki M, Shirato Y, Nakamura K, Nishi N, Nakamura T, Matsumoto R, Hirashima M: Regulation of galectin-9 expression and release in Jurkat T cell line cells. *Glycobiology* 2002; 12:111–118.
- 116 Zhu C, Anderson AC, Schubart A, Xiong H, Imitola J, Khoury SJ, Zheng XX, Strom TB, Kuchroo VK: The Tim-3 ligand galectin-9

negatively regulates T helper type 1 immunity. *Nat Immunol* 2005; 6:1245–1252.

- 117 Anderson AC, Anderson DE, Bregoli L, Hastings WD, Kassam N, Lei C, Chandwaskar R, Karman J, Su EW, Hirashima M, Bruce JN, Kane LP, Kuchroo VK, Hafler DA: Promotion of tissue inflammation by the immune receptor Tim-3 expressed on innate immune cells. *Science* 2007; 318:1141–1143.
- 118 Oomizu S, Arikawa T, Niki T, Kadowaki T, Ueno M, Nishi N, Yamauchi A, Hirashima M: Galectin-9 suppresses Th17 cell development in an IL-2-dependent but Tim-3-independent manner. *Clin Immunol* 2012; 143:51–58.
- 119 Ohtsubo K, Takamatsu S, Minowa MT, Yoshida A, Takeuchi M, Marth JD: Dietary and genetic control of glucose transporter 2 glycosylation promotes insulin secretion in suppressing diabetes. *Cell* 2005; 123:1307–1321.
- 120 Popovici RM, Krause MS, Germeyer A, Strowitzki T, von Wolff M: Galectin-9: a new endometrial epithelial marker for the mid- and late-secretory and decidual phases in humans. J Clin Endocrinol Metab 2005; 90:6170–6176.
- 121 Heusschen R, Freitag N, Tirado-González I, Barrientos G, Moschansky P, Muñoz-Fernández R, Leno-Durán E, Klapp BF, Thijssen VL, Blois SM: Profiling Lgals9 splice variant expression at the fetal-maternal interface: implications in normal and pathological pregnancy. *Biol Reprod* 2013; 88:22.
- 122 Swaminathan GJ, Leonidas DD, Savage MP, Ackerman SJ, Acharya KR: Selective recognition of mannose by the human eosinophil Charcot-Leyden crystal protein (galectin-10): a crystallographic study at 1.8 A resolution. *Biochemistry* 1999; 38:13837–13843.
- 123 Kubach J, Lutter P, Bopp T, Stoll S, Becker C, Huter E, Richter C, Weingarten P, Warger T, Knop J, Müllner S, Wijdenes J, Schild H, Schmitt E, Jonuleit H: Human CD4+ CD25+ regulatory T cells: proteome analysis identifies galectin-10 as a novel marker essential for their anergy and suppressive function. *Blood* 2007; 110:1550–1558.
- 124 Guardia CM, Gauto DF, Di Lella S, Rabinovich GA, Martí MA, Estrin DA: An integrated computational analysis of the structure, dynamics, and ligand binding interactions of the human galectin network. J Chem Inf Model 2011; 51:1918–1930.
- 125 Yang RY, Hsu DK, Yu L, Chen HY, Liu FT: Galectin-12 is required for adipogenic signaling and adipocyte differentiation. *J Biol Chem* 2004; 279:29761–29766.
- 126 Yang RY, Yu L, Graham JL, Hsu DK, Lloyd KC, Havel PJ, Liu FT: Ablation of a galectin preferentially expressed in adipocytes increases lipolysis, reduces adiposity, and improves insulin sensitivity in mice. *Proc Natl Acad Sci USA* 2011; 108:18696–18701.
- 127 Than NG, Pick E, Bellyei S, Szigeti A, Burger O, Berente Z, Janaky T, Boronkai A, Kliman H, Meiri H, Bohn H, Than GN, Sumegi B: Functional analyses of placental protein 13/galectin-13. *Eur J Biochem* 2004; 271:1065–1078.
- 128 Balogh A, Pozsgay J, Matkó J, Dong Z, Kim CJ, Várkonyi T, Sammar M, Rigó J Jr, Meiri H, Romero R, Papp Z, Than NG: Placental protein 13 (PP13/galectin-13) undergoes lipid raftassociated subcellular redistribution in the syncytiotrophoblast in preterm preeclampsia and HELLP syndrome. *Am J Obstet Gynecol* 2011; 205:156.e1–156.e14.

- 129 Kliman HJ, Sammar M, Grimpel YI, Lynch SK, Milano KM, Pick E, Bejar J, Arad A, Lee JJ, Meiri H, Gonen R: Placental protein 13 and decidual zones of necrosis: an immunologic diversion that may be linked to preeclampsia. *Reprod Sci* 2012; 19:16–30.
- 130 Burger O, Pick E, Zwickel J, Klayman M, Meiri H, Slotky R, Mandel S, Rabinovitch L, Paltieli Y, Admon A, Gonen R: Placental protein 13 (PP-13): effects on cultured trophoblasts, and its detection in human body fluids in normal and pathological pregnancies. *Placenta* 2004; 25:608–622.
- 131 Young AR, Barcham GJ, Kemp JM, Dunphy JL, Nash A, Meeusen EN: Functional characterization of an eosinophil-specific galectin, ovine galectin-14. *Glycoconj J* 2009; 26:423–432.
- 132 Lewis SK, Farmer JL, Burghardt RC, Newton GR, Johnson GA, Adelson DL, Bazer FW, Spencer TE: Galectin 15 (LGALS15): a gene uniquely expressed in the uteri of sheep and goats that functions in trophoblast attachment. *Biol Reprod* 2007; 77:1027– 1036.
- 133 Satterfield MC, Bazer FW, Spencer TE: Progesterone regulation of preimplantation conceptus growth and galectin 15 (LGALS15) in the ovine uterus. *Biol Reprod* 2006; 75:289–296.
- 134 Gray CA, Adelson DL, Bazer FW, Burghardt RC, Meeusen EN, Spencer TE: Discovery and characterization of an epithelial-specific galectin in the endometrium that forms crystals in the trophectoderm. *Proc Natl Acad Sci USA* 2004; 101:7982–7987.
- 135 Mor G, Cardenas I: The immune system in pregnancy: a unique complexity. Am J Reprod Immunol 2010; 63:425–433.
- 136 Liu AX, Jin F, Zhang WW, Zhou TH, Zhou CY, Yao WM, Qian YL, Huang HF: Proteomic analysis on the alteration of protein expression in the placental villous tissue of early pregnancy loss. *Biol Reprod* 2006; 75:414–420.
- 137 Molvarec A, Blois SM, Stenczer B, Toldi G, Tirado-Gonzalez I, Ito M, Shima T, Yoneda S, Vásárhelyi B, Rigó J Jr, Saito S: Peripheral blood galectin-1-expressing T and natural killer cells in normal pregnancy and preeclampsia. *Clin Immunol* 2011; 139:48–56.
- 138 Than NG, Erez O, Wildman DE, Tarca AL, Edwin SS, Abbas A, Hotra J, Kusanovic JP, Gotsch F, Hassan SS, Espinoza J, Papp Z, Romero R: Severe preeclampsia is characterized by increased placental expression of galectin-1. *J Matern Fetal Neonatal Med* 2008; 21:429–442.
- 139 Than NG, Kim SS, Abbas A, Han YM, Hotra J, Tarca AL, Erez O, Wildman DE, Kusanovic JP, Pineles B, Montenegro D, Edwin SS, Mazaki-Tovi S, Gotsch F, Espinoza J, Hassan SS, Papp Z, Romero R: Chorioamnionitis and increased galectin-1 expression in PPROM –an anti-inflammatory response in the fetal membranes? *Am J Reprod Immunol* 2008; 60:298–311.
- 140 Jeschke U, Mayr D, Schiessl B, Mylonas I, Schulze S, Kuhn C, Friese K, Walzel H: Expression of galectin-1, -3 (gal-1, gal-3) and the Thomsen-Friedenreich (TF) antigen in normal, IUGR, preeclamptic and HELLP placentas. *Placenta* 2007; 28:1165–1173.
- 141 Odibo AO, Zhong Y, Goetzinger KR, Odibo L, Bick JL, Bower CR, Nelson DM: First-trimester placental protein 13, PAPP-A, uterine artery Doppler and maternal characteristics in the prediction of pre-eclampsia. *Placenta* 2011; 32:598–602.
- 142 Cowans NJ, Stamatopoulou A, Khalil A, Spencer K: PP13 as a marker of pre-eclampsia: a two platform comparison study. *Placenta* 2011; 32:S37–S41.