

Sensing Glycans as Biochemical Messages by Tissue Lectins: The Sugar Code at Work in Vascular Biology

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Abstract

Although a plethora of players has already been revealed to be engaged in the haemostatic system, a fundamental consideration of the molecular nature of information coding can give further explorations of the mechanisms of blood clotting, platelet functionality and vascular trafficking direction. By any means, looking at ranges of occurrence and of potential for structural versatility, at strategic positioning to influence protein and cell sociology as well as at dynamics of processing and restructuring for phenotypic variability, using sugars as an alphabet of life for generating the glycan part of glycoconjugates is a success story. The handiwork by the complex system for glycan biosynthesis renders biochemical messages of exceptionally high coding capacity available. They are read and translated into cellular effects by receptors termed lectins. The different levels of regulation on both sides, that is, glycan and lectin, establish an intriguingly fine-tuned capacity for functional pairing. The emerging insights into the highly branched routes of glycosylation, into lectin structures up to complete characterization in solution and the shape of lectin networks, first obtained for the three selectins, now extended to considering many other C-type lectins, galectins and siglecs, as well as into intra- and inter-family cross-talk and cooperations are sure to push boundaries in our understanding of the molecular basis of haemostasis.

Keywords

- ▶ C-type lectin
- ▶ galectin
- ▶ mucin
- ▶ selectin
- ▶ siglec
- ▶ von Willebrand factor

Introduction

The cell surface is the platform for a multitude of recognition processes. They can lead to selective cross-linking events within the membrane (lattice formation) and hereby, for example, trigger outside-in signalling. Moreover, bridging between cells or cells and the extracellular matrix can be facilitated. Mechanistically, the complementarity between surface epitopes and their receptors underlies the specificity of these *cis/trans*-interactions. Allegorically spoken, a molecular message is 'read' and 'translated' into a post-binding effect, that is, biochemically coded information is turned into a process and an outcome. Considering the stringent space limitations on the surface and the large size of the pool of signals relevant to cover all aspects of cell sociology, it

immediately becomes apparent that the biochemical information coding is required to reach a high density. To do so, the building blocks (letters of an alphabet of life) of the molecular messages should be able to form many structural isomers (words) with as few constituents as possible. How is this chemically feasible?

In principle, ways of making connections between units to build a biopolymer chain are known from the 5', 3'-phosphodiester of nucleic acids or peptide bonds of proteins. In each case, the same chemistry is applied in a uniform Lego-like manner so that exclusively the sequence matters for information coding. With focus on this aspect, that means that going beyond the spatial order of units linked to each other in exactly the same way will open opportunities to

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increase the informational contents of oligomers. Explicitly, if variations in the positions and topological properties of connecting points (and branching) become regular features of oligomers, then the extent of structural diversity will be greatly enhanced, and, indeed, one class of biomolecules offers to realize this potential: carbohydrates (for common structures, please see ►Fig. 1; for overviews on their diversity and properties, please see Gabius¹). Our review introduces readers to the biochemical basis of the concept of the sugar code and respective recognition systems with relevance for vascular biology.

Known to be present abundantly and ubiquitously in nature, as polysaccharides and as conjugates with lipids and proteins, in these cases positioned strategically as

'sugary coating of cells',² glycans have the widespread occurrence profile and surface presentation expected of a basic coding system.^{3–14} Equally important to note, the positioning of the (glycosidic) bond that brings the anomeric centre (marked for each carbohydrate shown in ►Fig. 1) of an activated sugar donor together with one of the hydroxyl groups of the acceptor during chain elongation has many more options than just one, what is the case in nucleic acids and proteins. Structurally, variability in linkage position and in anomeric status of the glycosidic bond (α or β), the possibility for branching and also the introduction of various site-specific substitutions explain why, 'among all biological molecules, carbohydrates, *in a short sequence*, can potentially display the largest number of ligand structures'.¹⁵

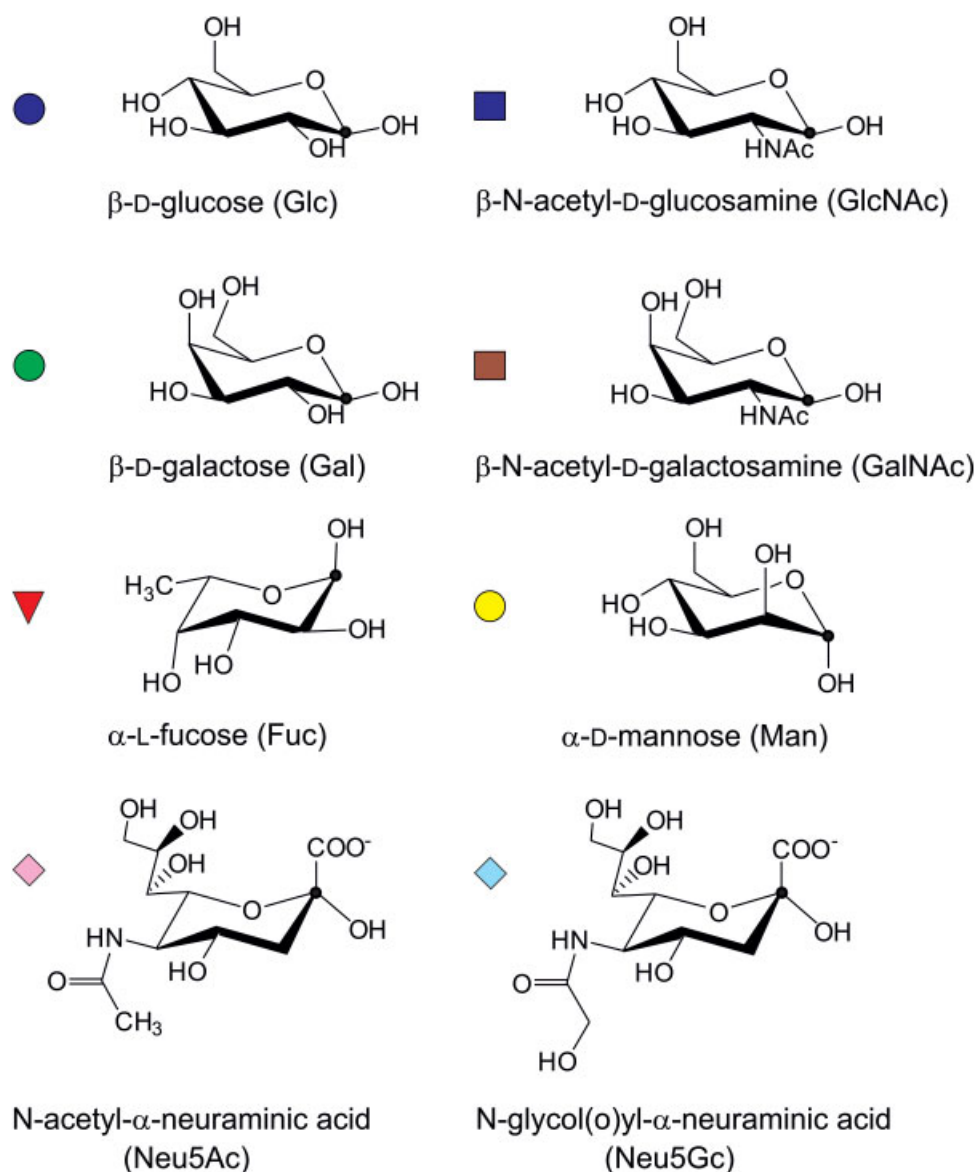


Fig. 1 Main letters of the sugar alphabet present in vertebrate glycans. The anomeric centre, the site of conjugation to an acceptor for glycan chain elongation, is marked by a black dot, and the typical colour symbol is given for each sugar. A change in the position of a single OH group (from equatorial to axial) is sufficient to alter the character of the letter, i.e., by epimer formation (Glc to Gal or to Man), as is its exchange by an N-acetyl group (GlcNAc, GalNAc). Reduction to obtain a deoxy sugar (i.e. for the exocyclic methyl group in l-Fuc) or an introduction of an additional hydroxyl group in the side chain to obtain Neu5Gc from Neu5Ac are further means to increase the pool size of this alphabet of life. The type of anomeric position for each compound is given in the monosaccharide's name.

This assumption will gain impact when this pool size reaches impressive numbers. Calculated for amino acids and for carbohydrates, a set of 20 types of monomers will theoretically build 6.4×10^7 hexapeptides but to as many as 1.44×10^{15} hexasaccharides.¹⁵ As concisely emphasized by Roseman, this exceptional level of structural diversity has its enormous pros but also imparts a practical con for analytical glycoscience: ‘glycoconjugates are much more complex, variegated and difficult to study than proteins or nucleic acids’,¹⁶ nonetheless giving ample reason to be ‘intrigued as to whether these sugars might be arranged in specific sequences that function as information molecules in biological processes’.¹⁷ Their capacity to store information and to transmit it by inter-molecular recognition is the basis of the sugar code.

Pioneering work to give substance to the concept of the sugar code applied enzymatically editing surface glycans of ³²P-labelled cells and following their in vivo routing. Using an enzyme fraction from *Clostridium perfringens* rich in L-fucosidase activity, ‘alteration of lymphocytes by glycosidases profoundly affects their fate in the body’.¹⁸ Instead of their typical recirculation to lymph nodes, increased quantity of radioactivity was found in the liver.¹⁸ Interestingly, a similar phenomenon of re-programming of routing was discovered when measuring the kinetics of clearance of a ⁶⁴Cu-labelled glycoprotein (i.e. ceruloplasmin) from serum by the liver: enzymatic removal of terminal sialic acids from the glycan chains of ceruloplasmin, what caused presentation of intact galactose (Gal) residues, drastically shortened the period of asialoglycoprotein’s serum presence.¹⁹ Since the status of desialylation can be an indicator of a glycoprotein’s lifetime like a timer, a functional correlation between the level of Gal presence at terminal position and hepatic uptake of the processed glycoprotein makes sense physiologically. Obviously, the hepatocytes (somehow) ‘read’ the message encoded in Gal-terminated glycans, as removal of fucose (Fuc) moieties erases the signal for lymphocyte homing.

In both instances, a distinct glycan signal (on the cell surface or as part of a glycoprotein) appears to serve as molecular equivalent of a postal code (in lymphocyte homing and glycoprotein clearance from circulation). This idea also emerged from studying other systems of cell adhesion such as neuronal or teratoma cells,²⁰ setting the stage to move the limelight to glycans. All that can only happen if the enzymatic assembly of glycans is a non-random process so that a large panel of epitopes with their own physiological meaning can be produced, and this independent of a template. The letters of the sugar alphabet are thus expected to be arranged in a highly ordered manner on proteins (and on sphingolipids), as meaningful words in any language are separated from purely random (non-sense) combinations. In this sense, ‘the significance of the glycosyl residues (of glycoconjugates) is to impart a discrete recognitional role on the protein’.²¹ This fundamental conclusion prompts us to first introduce the ‘letters’, to comment on the chemistry of their conjugation to ‘words’ next and then describe the enzymatic machinery ‘writing’ glycan-encoded messages.

The Sugar Code: Glycans as Signals

The examples of nucleic acids and proteins teach us how an enzymatic process connects letters of the first and second alphabets of life to messages. In general, an activated donor is used for chain elongation, and this at a single site of the acceptor. Taking the place of nucleotide triphosphates and aminoacyl adenylates, known from the other two types of biopolymer synthesis, are glycoside conjugates with nucleotides (uridine diphosphate, guanosine diphosphate or cytidine monophosphate) at the respective sugar’s anomeric centre. These activated sugar derivatives are the substrates for the enzymes to let a sugar chain grow, that is, glycosyltransferases. The most frequently used structural units (letters) for glycan biosynthesis in mammals are shown in ►Fig. 1.

Since hydroxyl groups of acceptor carbohydrates (for hexopyranoses such as glucose [Glc] or Gal shown in ►Fig. 1) are chemically rather equivalent, variability in linkage positions will be possible if the enzymatic assembly line is equipped with respective sets of tools. Indeed, this is the case so that glycosidic linkages with (α or β)1,2- or 1,3- or 1,4- or 1,6-connections can be formed. As a consequence, a disaccharide is not only structurally defined by its sequence but also by these two additional parameters. For example, Gal α 1,4Glc, Gal β 1,4Glc or Gal β 1,6Glc are different compounds (‘words’) that all share the Gal-Glc sequence.

In contrast to homopolysaccharides such as chitin or cellulose with their uniform β 1,4-linkage of Glc(NAc) residues, glycans of cellular glycoconjugates are the proof that the potential of carbohydrates to build oligomers of enormous diversity is actually exploited. The same substrate, that is, the sugar of its nucleotide derivative, can enter an acceptor glycan at different positions depending on enzyme and acceptor presence. This principle that underlies the origin of glycan complexity is graphically illustrated by an example from the maturation of N-glycans of glycoproteins.

The pentasaccharide core of N-glycans shown in ►Fig. 2 is the common substrate for processing towards all forms of complex-type N-glycans. Six sites exist for branch initiation by site- and position-specific addition of N-acetylglucosamine (GlcNAc) in β 1-linkage via distinct N-acetylglucosaminyltransferases (GnTs), as listed in ►Fig. 2. Linkages are in β 1,2/4/6, and

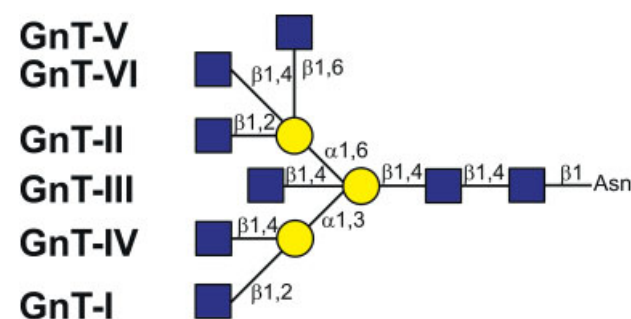


Fig. 2 The six modes of GlcNAc incorporation into the N-glycan core pentasaccharide by N-acetylglucosaminyltransferases (GnTs). Their activity in the order of numbers given to each enzyme initiates production of up to penta-antennary complex-type N-glycans. For explanations of symbols for sugars, please see legend to ►Fig. 1.

each GnT has its specific acceptor, working in the assembly line at a position as indicated by the given numbers. Viewed systematically, large enzyme panels, too, are available for each other type of carbohydrate so that the toolbox of glycosyltransferases is well equipped for making glycan diversity and complexity possible.^{22–37} The connection of impairing certain enzymes of glycan biosynthesis to causing pathological phenotypes of mice deficient in the responsible genes underscores their relevance for cellular physiology.³⁸ In these studies, important clues may even be missed, because compensatory mechanisms have been tracked down that maintain certain glycome features despite the engineered deficiency.^{39,40} The dynamic nature of availability of substrate, acceptor and enzyme accounts for the broad spectrum of the glycome and its spatiotemporal diversity.^{41–47}

Overall, carbohydrates have the chemical features to make the envisioned high-density coding possible, and an enzymatic machinery has been developed to realize this potential. Variations in anomeric position and in linkage points enable the glycome to be composed of a huge number of structures—at the same time posing demanding challenges to structural glycan characterization. Progress in glycan analysis^{48–50} and synthesis⁵¹ has laid the foundation for aiming at delineating structure–function relationships. As important as the presence of the hydroxyl groups (in axial/equatorial positions) is for structural variability, it then is for making directional hydrogen bonding possible, and there is more: in contrast to most peptides, glycans are not highly flexible. Within their conformational space, they often access few low-energy ‘valleys’ (conformers), a boon in the thermodynamic balance sheet when binding to a receptor.⁵² Hereby, the entropic penalty is rather low, favouring binding. In aggregate, the hypothesis for a flow of information from glycans as ligands (counter-receptors) via a recognition process (‘reading’) by receptors and a translation of the glycan-encoded information into biological effects by this functional pairing is attractive and testable.

Historically, an antibody-to-antigen-like association has been discussed approximately 70 years ago.^{53,54} Worded prophetically, ‘rather than trying to force all biological specificity into the immunological compartment, we might have to consider the latter as merely a special case of the more universal biological principle, namely, *molecular key-lock configuration as a mechanism of selectivity*’,⁵⁴ here drawing on Fischer’s famous lock-and-key analogy derived from his work with the glycosidases ‘Invertin und Emulsin’.⁵⁵ Within this concept, the two key questions that we must answer are as to whether (antibody-like) receptors for glycans exist and, if positive, whether their presence shows the necessary degree of diversification for accomplishing the assumed task of decoding of a large array of glycan-encoded signals.

The Sugar Code: Lectins as Readers/Translators

A simple and robust assay was crucial to detect lectin activity, that is, haemagglutination. The bridging of erythrocytes by at least bivalent proteins that leads to visible cell aggregation was

the experimental read-out for the discovery of (haem) agglutinins, first in rattlesnake venom (1860), and then in plant extracts (1888). Mitchell put one drop of venom on a slide, ‘and a drop of blood from pigeon’s wounded wing (was) allowed to fall upon it. They were instantly mixed. Within three minutes the mass had coagulated firmly’,⁵⁶ the landmark discovery of receptors bridging cells by ligand binding (an involvement of plasma compounds in the ‘coagulation’ was rigorously excluded ~40 years later,⁵⁷ and the active protein biochemically characterized more than 100 years later).⁵⁸

Giving these agglutinins further medical relevance, members of this family of effectors were described that could even equal the ability of serum antibodies in distinguishing the blood group status of human donors (for historical survey, please see Refs. ^{46,47} and ^{59–61}). That constituents of plant extracts could thus ‘read’ surface determinants of erythrocytes as the blood group-specific antibodies do, which Landsteiner described as cause for fatal incompatibilities in blood transfusion,^{62,63} which inspired Boyd to shape a generic name for these activities: ‘it would appear to be a matter of semantics as to whether a substance not produced in response to an antigen should be called an antibody even though it is a protein and combines specifically with a certain antigen only. It might be better to have a different word for the substances and the present writer would like to propose the word *lectin* from Latin *lectus*, the past principle of *legere* meaning to pick, choose or select’.⁶⁴

Instead of using a term like antibody-like substance with implicit structural comparison, the emphasis was wisely placed on a feature of ligand binding, ‘intending thus to call attention to their specificity without begging the question as to their nature’, while certainly being intrigued by the observations that ‘lectins imitate the behaviour of immune antibodies’.⁶⁵ When asked (in 1973) how he would define a lectin, Boyd replied: ‘that although he had invented the word he had no right to dictate what it should mean. He said that once a word goes into general circulation it becomes independent of its originator and eventually becomes whatever people think it ought to mean! He added that he would like to use the word to mean a protein that had a more or less specific action and that there is no reason to think it is an antibody’.⁶⁶ Over time, lectin became the most common name for agglutinating activities, although the synonym (phyto)haemagglutinin is still in use, for example, for two bean (*Phaseolus vulgaris*) lectins (PHA-E/L; their mixture holds a special place in lectinology, because the induction of mitogenesis in human leukocytes by PHA proved elicitor capacity of lectin binding for the first time⁶⁷) and for a lectin of the influenza viral surface. Thus, lectins qualify as receptors, and investigating their specificity to cellular targets unveiled glycans as counter-receptors.

That their specificity is directed to glycans can historically be traced to reporting a ‘de-agglutination’ of erythrocyte aggregates that had formed in the presence of ricin or abrin preparations by using hog gastric mucin, a potent lectin binder.^{68,69} Blocking haemagglutination in the presence of blood group-specific lectins by a sugar, first Fuc inhibiting the activity of the lectin of eel serum to cross-link type O(H)

erythrocytes,⁷⁰ proved carbohydrate binding beyond doubt, and this approach was systematically followed in transfusion medicine and in lectinology in general.⁷¹ Terminologically, carbohydrate binding is the first criterion for a lectin, a feature yet shared by other classes of proteins such as immunoglobulins (Igs), as the mentioned antibodies attest, and glycosyltransferases. Thus, besides carbohydrate-binding antibodies, lectins need to be strictly separated from any enzyme using a carbohydrate as substrate, from sensors/transporters for free mono- and oligosaccharides and carbohydrate-binding modules of bacterial and fungal glycoside hydrolases.^{72,73}

To have the implied broad impact on many aspects of cellular physiology, the range of glycan-lectin recognition must not be narrow. Implicitly, this means that lectins must not be a rare commodity in the proteome. The implied functional pairing is assumed to account for a co-evolution towards diversity. Indeed, matching diversity on the level of glycans, more than a dozen protein folds have developed capacity to interact with glycans (for three examples, please see ►Fig. 3; complete illustration of the folds of the families of animal/human lectins is given in Fujimoto et al⁷⁴ and in the Gallery of Lectins in Solís et al,⁷⁵ along with a detailed listing and description of methods to analyse complex formation of a lectin with its ligand in that paper's table 1).

This is similarly seen on the level of glycozymes. They code for glycosyltransferases, glycosidases, glycan-modifying enzymes such as sulfotransferases (please see below) and enzymes for carbohydrate biosynthesis such as for the two sialic acids shown in the bottom part of ►Fig. 1 and for the ensuing generation of the activated carbohydrates as well as for transporters of activated sugars. Diversification of the gene for each ancestral lectin (or glycozyme product) has led to families of genes. In the case of lectins, they code for homologous proteins with differences in fine specificity (due to the sequence alterations within the carbohydrate recognition domain [CRD]) and in the architecture in terms of modular display. Intriguingly, a CRD can be embedded into a complex structural context, with often not yet resolved physiological relevance.

In fact, the CRD can be part of a multi-modular protein, and individual regions can functionally cooperate, for example, to form aggregates (such as galectin-3; please see below) or to guide a lectin to the extracellular space, where it then mediates ordered cell migration.⁷⁶ When thoroughly examining occurrence of proteins with any of the three folds illustrated in ►Fig. 3 as examples, it turned out to be a common theme within phylogenesis that lectin diversity is truly attained commonly on the level of families. This apparent co-evolution with glycans resulting in large lectin numbers strongly argues in favour of the assumed role of lectins as translators for the large panels of glycan-encoded messages.⁷⁷ This intra-family divergence is described in further detail for C-type lectins (please see reviews^{78–80}), for sialic acid-binding Ig superfamily lectins (siglecs) (please see^{81,82}) and for ga (lactose-binding)lectins^{83–88}. Notably, in all three cases, species differ in qualitative aspects of gene display and also with respect to gene number and organization, for example, for mammalian galectins^{84,89} so that extrapolations from animal models to the clinical situation should always be performed with adequate caution.

Having identified this wide range of carbohydrate-binding motifs and detected a large number of endogenous lectins, the two fundamental questions posed at the end of the previous chapter are convincingly answered with a 'Yes'. The current challenge therefore is to proceed from completing the phase of lectin discovery to entering the era of building a functionally meaningful puzzle from the individual proteins.⁹⁰ First, a lectin must be assigned to its binding partners and pairing is very selective. Next, respective studies are revealing that members of lectin families appear to be expressed in networks. This raises the possibility for functional crosstalk (antagonism or cooperation) in *cis/trans*-interactions and in sensing non-self-signals. Obviously, the concept of lectins expressed in networks deserves to become a paradigm. The multiplicity of lectin representation is, for example, in certain cell types or tissues, attested by discovering a large group of myeloid C-type lectins⁹¹ and by revealing overlapping roles of C-type lectins in anti-microbial and anti-fungal immunity.^{92,93}

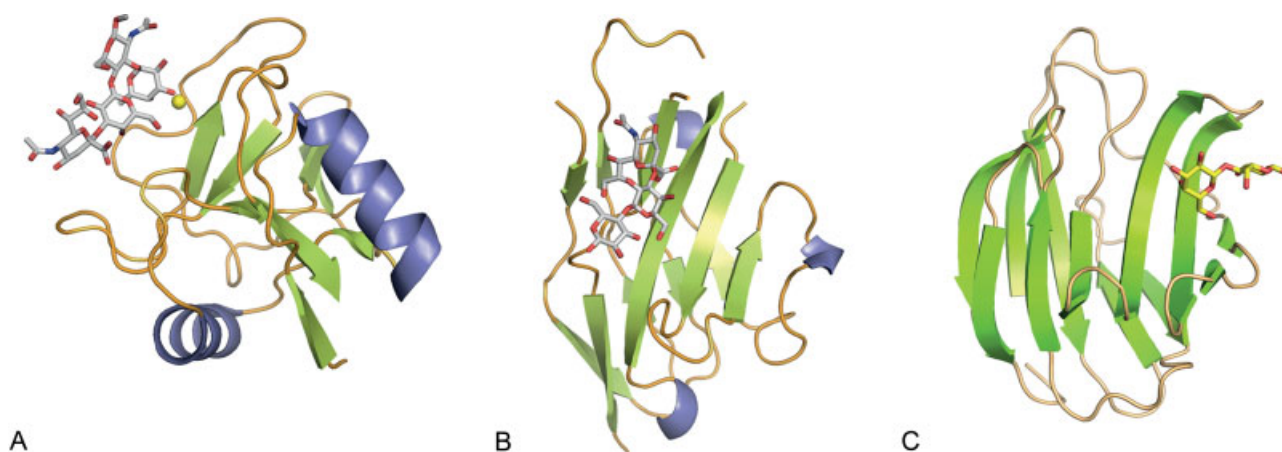


Fig. 3 Illustration of three examples of basic folds of mammalian lectins with bound ligand, that is, C-type lectin E-selectin (A; Protein Data Bank [PDB] code: 1G1T), siglec-1 (sialoadhesin) (B; PDB code: 1QFO) and galectin-1 (C; PDB code: 1GZW).

First detected by immunohistochemistry with non-cross-reactive antibody preparations, galectins analysed up to the level of full-scale monitoring of the complete set of family members are expressed in distribution profiles with individual features and overlaps.^{94–98} These results, of course, have immediate relevance for the strategy how to analyse lectin activities, that is, in mixtures. Initial comparative study of galectin activities alone and in mixtures has been performed by testing surface-tailored glycodendrimersomes in bridging assays^{99,100} and by using cell-based functional assays.^{101–103} The obtained evidence for functional interplay encourages to study in more detail what will happen when lectins are tested alone and in combinations simulating *in vivo* conditions, and this by using members of the same family and of different families.

In summary, presence of sophisticated machineries for lipid and protein glycosylation, producing the ligand side, and for glycan recognition, the complementary side, clearly supports the concept of an information transfer by glycan–protein interactions (sugar code) and gives cracking the sugar code direction.¹⁰⁴ The aim is to define the meaning of a glycan in its cellular context. Following this concept, distinct carbohydrate epitopes of glycan chains of cellular glycoconjugates can be expected to engage in functional pairing with certain tissue (endogenous) lectins. Any structural processing (for example, stepwise chain elongation) should then change the glycan's ligand properties, that is, its biochemical meaning. By looking exemplarily at main products of N- and mucin-type O-glycan biosynthesis, the hypothesis that a series of functional pairings occurs along the biosynthetic pathways is next put to the test, first for branch ends of complex-type N-glycans. At the same time, the following description of routes of glycan processing itself documents that many branch points exist to let glycan diversity reach an amazing level.

Functional Pairing: N-Glycans

N-glycosylation is a co-translational process that is started at the signal-peptide-dependent entry of a nascent protein into the endoplasmic reticulum (ER). The $\text{Glc}_3\text{Man}_9\text{GlcNAc}_2$ oligosaccharide is transferred from its dolichol pyrophosphate donor to the N-atom of an asparagine (Asn) moiety within a protein's N-glycosylation (sequon) motif [i.e. Asn-Xaa (no Pro)-Ser/Thr-Xaa (no Pro)].²⁹ During the early stage of the pathway of N-glycan processing and maturation, initial removal of two to three Glc moieties makes a routing signal with this glycan accessible. It is a molecular signal for vectorial transport of the respective glycoprotein from the ER to the Golgi. Delivery is performed by a cargo transporter (M_r 53,000) present in the ER-Golgi intermediate compartment (ERGIC), called ERGIC-53.^{105,106} Impairment of its binding to partially or fully deglycosylated N-glycans of target glycoproteins by mutations in the ERGIC-53 gene or the gene of its luminal interaction partner, that is, the multiple coagulation factor deficiency protein 2, is cause of an autosomal recessive bleeding disorder characterized by combined reduction of blood levels of coagulation factors V and VIII.^{107–109} This case study of combined factors V and VIII deficiency underlines the emerging relevance of glycan/lectin recognition for haemostasis.

Since the ER and the route from the ER to the Golgi are equipped with quality controls by lectins to sort out misfolded glycoproteins, glycans have more than one meaning already at this stage.^{110–112} Following the glycoprotein further along the Golgi route, each N-glycan can be trimmed to the pentasaccharide core as shown in ►Fig. 2 so that, for example, complex-type structures arise from the ensuing maturation. In the Golgi, as shown in ►Fig. 2, six GnTs are capable to add a GlcNAc residue at specific sites of the core pentasaccharide to initiate their synthesis. Incomplete maturation or restructuring by stepwise degradation of mature N-glycan branches are the means to let GlcNAc become presented for protein binding at branch ends.

Relevant for a route of platelet clearance, GlcNAc-terminating N-glycans of the GPIIb α sub-unit of the von Willebrand factor (VWF) complex associate with the α_M -chain of the $\alpha_M\beta_2$ -integrin of phagocytic cells (Mac-1, CD11b/CD18, CR3; Mac-1 was defined as granulocyte- and monocyte-specific antigen by a monoclonal antibody [clones M1/70] raised against mouse spleen cells¹¹³; for details on the Mac-2 antigen [clones M3/31 and M3/38], please see below).^{114,115} Terminal β GlcNAc in N-glycans (α -anomeric linkage is known from mucins¹¹⁶) is not only a ligand for an integrin but also for the C-type lectins langerin (CD207),^{117,118} the liver and lymph node sinusoidal endothelial cell lectin (LSEctin)¹¹⁹ and its close relative human dendritic cell-specific ICAM-3-grabbing nonintegrin-related protein (DC-SIGNR, CD299)¹²⁰ or its murine homologue SIGN-R2.¹²¹ The first epitope along the illustrated route of N-glycan processing therefore fulfils the expectation for functional pairing.

Elongation of the N-glycan chain by adding a Gal moiety (in β 1,4-linkage, as shown in the centre of ►Fig. 4) drastically changes the ligand characteristics, underscoring the required specificity. As noted above, accessible Gal units are docking sites for hepatic clearance, and this also applies to coagulation factor VIII and to VWF¹²² or to tissue plasminogen activator.¹²³ This type of interaction between desialylated glycoproteins on the surface of (senile) platelets and the hepatic asialoglycoprotein receptor is more than a clearance process. In fact, it is 'the long-elusive physiological ligand–receptor pair regulating hepatic thrombopoietin messenger ribonucleic acid production'.¹²⁴ Since the circulatory lifespan of platelets is shortened during systemic infection due to the activity of bacterial sialidases on cell surface glycans, the lectin-mediated clearance is a means of haemostatic adaptation limiting the severity of disseminated intravascular coagulation during sepsis.¹²⁵ Physiologically, one means to mask this determinant (postal code) for hepatic uptake is by α 2,3-sialylation (►Fig. 4, top, left). Fittingly, engineered deficiency in a respective glycosyltransferase, that is, α 2,3-sialyltransferase-IV, accelerated clearance of these glycoproteins and hereby (likely) accounts for prolonged bleeding and coagulation times in the knockout (KO) mice.¹²⁶ In contrast, the alternative to α 2,3-sialylation, that is, α 2,6-sialylation of N-acetyllactosamine (LacNAc) shown in ►Fig. 4, can be tolerated by the hepatic C-type lectin, as shown with neoglycoproteins and natural glycoproteins.^{127–129} Considering the frequent occurrence of LacNAc at the terminal position of

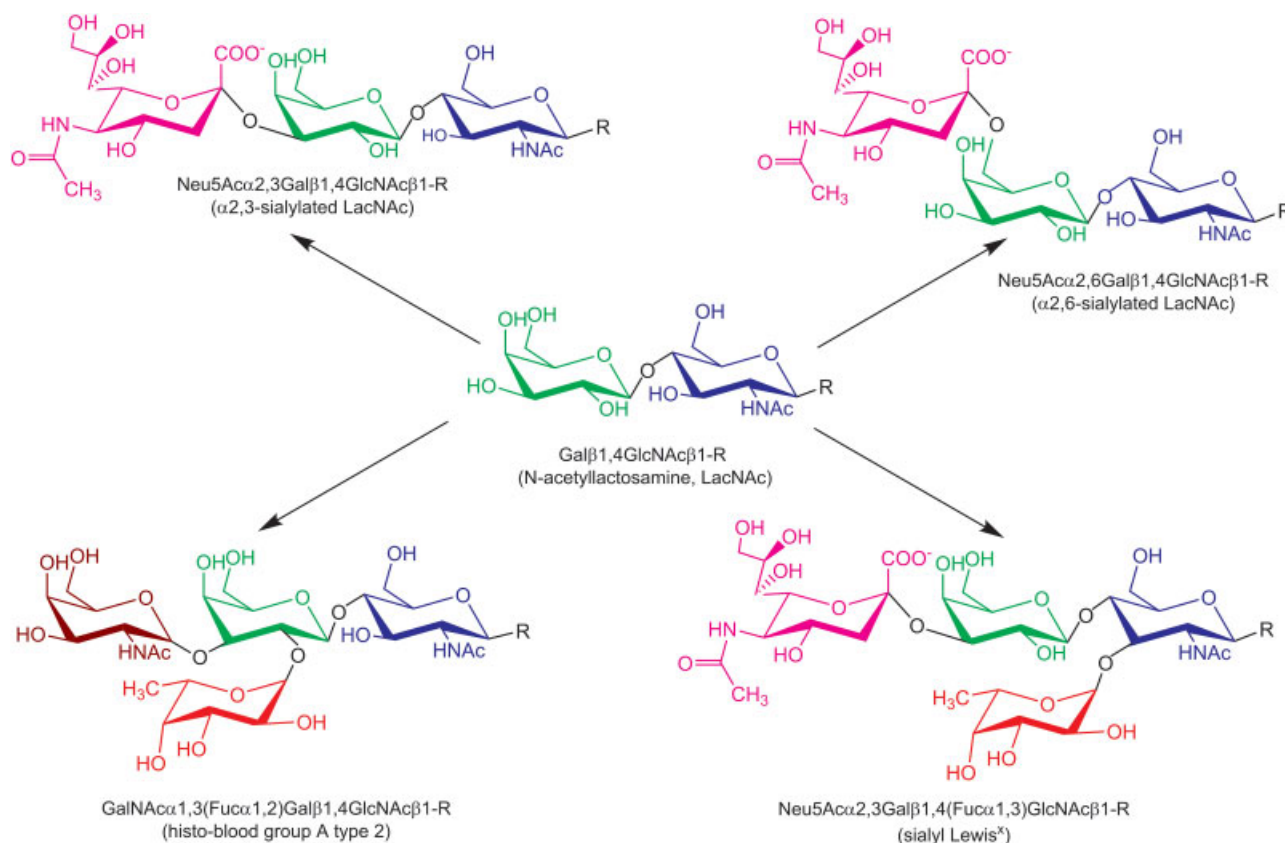


Fig. 4 Illustration of four different routes of biosynthetic elongation of a LacNAc epitope of complex-type N-glycans (centre). Specific glycosyltransferases can generate a panel of products (messages of distinct meaning), either by sialylation of terminal Gal in α 2,3- or in α 2,6-linkage (top part, left and right), by stepwise α 1,3-fucosylation (of GlcNAc) and α 2,3-sialylation (of Gal) to yield first the Lewis^x epitope (not shown) and then the shown sialyl Lewis^x tetrasaccharide (bottom part, right) or by a two-step reaction towards ABH histo-blood group (type 2) epitopes via α 1,2-fucosylation (of Gal to yield the H(0) trisaccharide; not shown) and the following α 1,3-GalNAc addition (the α 1,3-GalNAc-containing A-type tetrasaccharide is shown: bottom part, left).

N-glycans and its processing by these two different routes of sialylation shown in **Fig. 4** (top part), it is no surprise that, in addition to the C-type lectin fold, other types of lectins can accommodate these epitopes, especially the galectin and siglec folds shown in **Fig. 3**.

LacNAc is the canonical ligand for galectins. Its α 2,3-sialylation can enhance galectin affinity depending on the protein, most prominently for galectin-8.¹³⁰ Of note, this galectin binds VWF and coagulation factor V, the latter then imported into megakaryocytes by endocytosis, and also to the platelet integrin α IIb β ₃, a new entry to the list of platelet activators.^{131,132} Illustrating the marked impact of the site of sialic acid conjugation on ligand features, α 2,6-sialylation switches off affinity to galectins, which bind LacNAc via hydrogen bonds to the axial 4'-OH and the exocyclic 6'-OH groups of the Gal moiety (please see **Fig. 1**, second row for illustration of these positions, relative to the 3'-OH group).¹³³ In the case of α 2,6-sialylation of LacNAc shown in **Fig. 4** (top, right), the contact point at C6 is therefore occupied by a sialic acid residue. What is detrimental to galectin binding, though, is essential for association to siglecs. Presence of such a residue in α 2,3-linkage is suited for siglec-4 (myelin-associated glycoprotein); when in α 2,6-linkage, it is the docking site for siglec-2 (CD22). In sum, α 2,6-

sialylation of LacNAc extension thus abolishes affinity to galectins, can be tolerated by a C-type lectin and is the primary contact to a siglec. By the way, sialic acids can hereby implement recognition sites on glycoproteins for a siglec so that a contribution to the regulation of their plasma levels is possible, as discussed for VWF and coagulation factor VIII by binding to siglec-5 on macrophages.¹³⁴

The two other illustrated products of terminal N-glycan tailoring shown in **Fig. 4** (bottom), that is, the histo-blood group A determinant (**Fig. 4**, bottom, left) or (sialyl) Lewis^x [(s)Le^x, CD15(s)] (**Fig. 4**, bottom, right), too, have their specific meaning as ligand. For example, galectin-3 (the mentioned Mac-2 antigen; please see below) is a high-affinity receptor of the A-tetrasaccharide,¹³⁵ DC-SIGN (CD209) a receptor of epitopes of the Lewis blood group system such as Le^a or Le^x¹³⁶ and the selectins bind the shown sLe^x, an interaction of pivotal significance for the interaction of leukocytes and platelets with the vasculature under conditions of activation (please see below).^{137,138} Turning Le^x not into sLe^x but alternatively into Le^y (CD174) by terminal α 1,2-fucosylation instead of the α 2,3-sialylation shown in **Fig. 4** renders the respective N-glycan capable to interact with thrombospondin's C-type lectin-like domain, implicated in inhibiting angiogenesis.¹³⁹

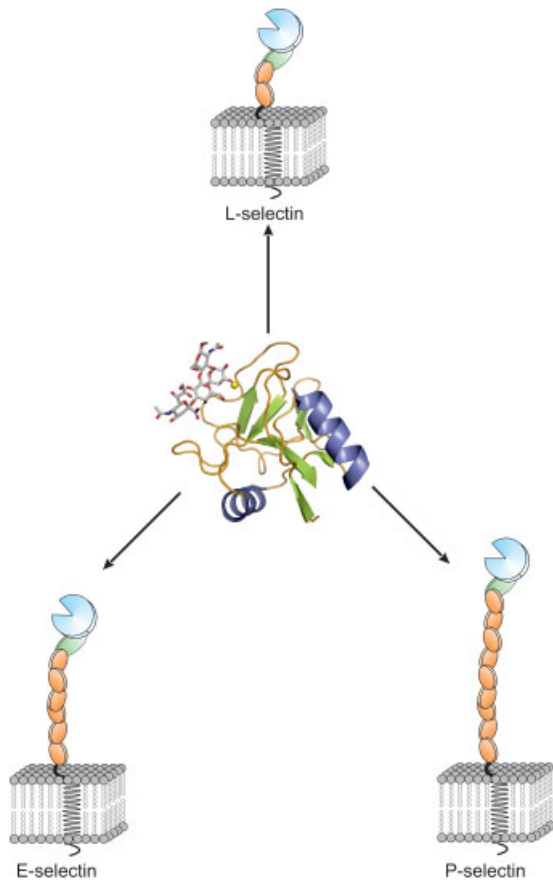


Fig. 5 Modular organization of the three selectins characterized by the C-type lectin domain (centre; please see also ►Fig. 3A) at the most prominent position for intercellular contact, followed by an epidermal growth factor (EGF)-like domain and two to nine complement-binding consensus repeats.

The three selectins mentioned above are classical examples of lectins with modular architecture. Their design is shown in ►Fig. 5. The terminal C-type CRD, as is the case for the V-set IgG domain of siglec rods, is optimally presented on the membrane to bridge cells. First detected by function blocking or localization using monoclonal antibodies (Mel-14) for the lymphocyte homing receptor L-selectin (CD62L),¹⁴⁰ H18/7 and H4/18 for the chemokine- and endotoxin-inducible endothelial-leukocyte adhesion molecule-1 (ELAM-1; E-selectin, CD62E)¹⁴¹ and S12 for an α -granule membrane protein of M_r 140,000 (GMP-140) that is redistributed to the plasma membrane upon platelet stimulation, what gave GMP-140 a new name: *platelet activation-dependent granule to external membrane protein* (PADGEM; P-selectin, CD62P).^{142–145} The spatial accessibility of the glycan counter-receptors at branch ends, rapid on-rates during contact formation by mostly ionic interactions and catch bonding are factors that underlie their role as selective cell adhesion molecules (explaining the origin of the term selectin¹⁴⁶), operating as anchors for cells in the blood flow. Like the mentioned enzymatic removal of Fuc from the lymphocyte surface raised evidence for the concept of glycans as postal codes in routing and delivery,¹⁸ the importance of a second sugar type present in sLe^{a/x} epitopes, that is,

sialic acids, has been revealed by treatment of sections of lymphoid organs with bacterial sialidase.¹⁴⁷ These reports converge to support the involvement of glycans in lymphocyte homing, as the blocking experiments with selectin-binding monoclonal antibodies did for the protein side. Intriguingly, α -L-fucosidase is suggested to limit leukocyte migration at late stages of inflammation (tested in murine experimental autoimmune uveoretinitis) upon induction by chemokines CCL3/5 by reducing P-selectin binding.¹⁴⁸ Besides presentation by N-glycans, the sLe^{a/x} epitopes are characteristically also a part of certain types of mucin-type O-glycans, here reaching a high density that favours to build and maintain selectin–glycoprotein contacts. In addition to the negative charge of the sialic acid, sulphate groups at C6 of Gal and/or GlcNAc moieties increase the ligand's capacity for rapid contact building via ionic charge complementarity under conditions of flow.^{149–153} Sulphation is also conducive to 'write' a routing signal for glycoprotein clearance, if GlcNAc is not conjugated with Gal but with N-acetylgalactosamine (GalNAc), a further mode of N-glycan tailoring.¹⁵⁴

Before turning to mucin-type O-glycans in more detail, a second salient lesson should be drawn from ►Fig. 4, besides the documented sets of structure–ligand relationships: a glycan epitope (such as LacNAc), that is, a certain acceptor, can be converted into more than one product. This opens opportunities for highly versatile regulation of glycome representation by modulating enzyme and/or substrate availability. Exemplifying the intimate connection of a distinct biomedically relevant process with glycosylation, the malignancy-associated high-level sialylation can be interpreted, among other implications, as a protection against growth inhibition. Along this line, the tumour suppressor p16^{INK4a} has been shown to counteract malignancy at this level. This protein is essential to induce anoikis in human pancreatic carcinoma (Capan-1) cells. It works via orchestrated down-regulation of α 2,6-sialylation (by reducing sialic acid biosynthesis) teamed up with up-regulation of homobivalent galectin-1, which is the functional receptor 'reading' the increase in terminal LacNAc and cross-linking the similarly up-regulated glycoprotein counter-receptor $\alpha_5\beta_1$ -integrin via this recognition to trigger caspase-8 activation.^{155,156}

The signal (access to LacNAc), the 'reader' (galectin-1) and the downstream effector ($\alpha_5\beta_1$ -integrin) are thus co-regulated towards anoikis induction, giving the glycophenotypic change a functional meaning. Of note, a similar team building to attain T cell apoptosis has been reported for α 2,6-sialyltransferase, CD45 and galectin-1.¹⁵⁷ The respective delineation of regulatory pathways for generating selectin counter-receptors^{158–160} has proven to be a hallmark of supporting the concept of functionally dynamic glycomics in lymphocyte homing and platelet aggregation. On the level of O-glycans, counter-receptor occurrence can be as tightly controlled in a coordinated manner by competition between different routes and terminal tailoring, following the same principles as shown in ►Fig. 4 for N-glycans. Routes of mucin-type O-glycan processing after the initial incorporation of a GalNAc residue into the protein are illustrated in ►Fig. 6. The presence of the α -linked GalNAc residue

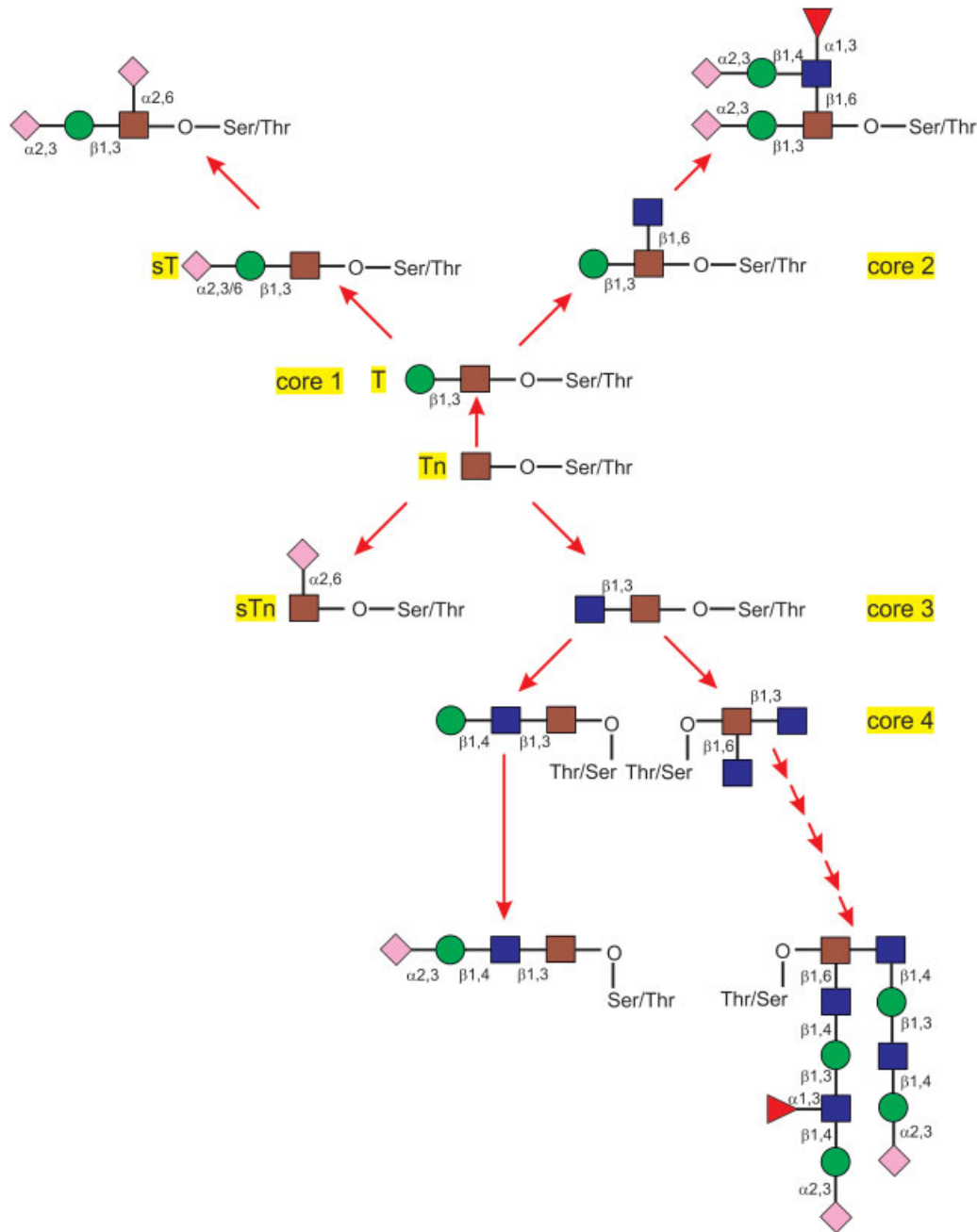


Fig. 6 Illustration of six different routes of biosynthetic elongation of the first product of mucin-type O-glycosylation, i.e., T_n antigen (centre). α 2,6-Sialylation of the GalNAc residue results in the sialyl (s)T_n determinant that cannot undergo any further processing. The activity of the respective sialyltransferases underlies occurrence of sT_n at the expense of all other O-glycans. Extension of the T_n antigen towards branched structures can alternatively proceed along two routes that establish either the core 1 or the core 3 structures. Enzymatic β 1,3-galactosylation produces the core 1 disaccharide (T antigen) that can be sialylated stepwise in up to two positions, first in α 2,3-linkage at the Gal moiety to yield the sialyl (s)T antigen, finally in α 2,6-linkage at the GalNAc residue to produce disialylated T (top part, left). Alternatively, the T antigen can be subject to core 2 trisaccharide generation by adding GlcNAc at the central GalNAc residue in β 1,6-linkage (top part, right). As shown in ► Fig. 2 and in ► Fig. 4 for N-glycans, this GlcNAc residue is the starting point of branch elongation that can encompass LacNAc repeats and the Le/sLe determinants. This trisaccharide therefore is the platform for obtaining various types of oligosaccharides, depending on the actual availability of enzymes and substrates for the competing routes of processing. Mechanistically, the same holds true for the case of transition of the T_n antigen to the core 3 disaccharide and then core 4 structures (bottom part, right), in close analogy to core 1 and core 2 synthesis. Correspondingly, without branch introduction, the core 3 disaccharide can be extended in two steps to a sialylated tetrasaccharide (bottom part, centre), in analogy to core 1 oligosaccharide processing. Occurrence of α 2,3-sialylated LacNAc in fully processed core 1/3 structures resembles that of a complex-type N-glycan branch ends (please see ► Fig. 4, top part, left). When considering that the processing of a T_n antigen can take diverse routes, it becomes obvious that the system of mucin-type O-glycosylation has manifold possibilities to generate a large number of O-glycans by regulating acceptor, enzyme and/or substrate levels. For explanation of symbols for sugars, please see legend to ► Fig. 1.

defines the T_n antigen (n for nouvelle; CD175), the target of serum auto-antibodies en route to spontaneous polyagglutinability of erythrocytes in the cold causing the T_n syndrome.^{161–163} This determinant is not only a docking site for antibodies but also for human lectins.

Functional Pairing: Mucin-Type O-Glycans

This type of protein glycosylation is characterized by reaching a high local density, ideal for a defensive barrier and also for high-affinity receptor binding.^{164–167} The committed step for mucin-type O-glycosylation is performed by members of the family of polypeptide N-acetylgalactosaminyltransferases (GalNAcTs), most of the 20 enzymes harbouring a β-trefoil lectin domain that binds GalNAc in addition to the enzymatically active protein part.^{32,33,168} They cooperate to let GalNAc incorporation reach diverse levels of local density in their substrates, and more than 80% of the proteins passing through the Golgi are predicted to be processed this way. Interestingly, each enzyme can well fulfil a particular functional role, as revealed by KO mouse analysis. Despite integrity of all other enzymes, a deficiency in one of the N-acetylgalactosaminyltransferases, that is, GalNAcT-1, caused a moderate to severe bleeding disorder due to reduced blood levels of coagulation factors and decreased level of recruitment of leukocytes to sites of inflammation, highlighting O-glycan relevance to haemostasis and vascular biology.¹⁶⁹ Binding studies of T_n-presenting glycopeptides demonstrated that O-linked GalNAc is a ligand for endogenous lectins. The macrophage Gal(-binding C)-type lectin (MGL, CD301, CLEC10A) and galectin-4 are tissue receptors of the T_n antigen.^{170,171} Present in Kupffer cells that also express the mentioned α_Mβ₂-integrin for β-GlcNAc-dependent platelet clearance, a C-type lectin (CLEC4F) is responsible for efficient uptake of T_n-presenting mucins and of platelets of genetically engineered mice made defective in O-glycan maturation so that this process is entirely arrested at the stage of O-GalNAc presentation.^{172,173} Physiologically, this epitope is obliterated for this process by α2,6-sialylation. The resulting disaccharide sialyl T_n (sT_n; CD175s) shown in ►Fig. 6 maintains affinity for human MGL,¹⁷⁴ as is the case for α2,6-sialylated LacNAc at N-glycan termini and the closely related hepatic C-type lectin (please see above). At the same time, the sialylation step enables a gain-of-function, because it confers binding to siglec-15 on macrophages.^{175,176} A V-set module resembling that of siglecs in a paired Ig-like receptor, that is, PILRα, also binds the sT_n antigen.^{177,178}

Sialylation of the T_n epitope is not the only way to use it as substrate. As shown in ►Fig. 6, an alternative pathway, competing with the synthesis of the sT_n antigen, is the generation of the T(F) disaccharide (CD176; first described as antigen by O. Thomsen and his assistant V. Friedenreich, thus termed Thomsen–Friedenreich antigen¹⁷⁹), the core 1 disaccharide of mucin-type O-glycans. Dense clustering favours binding of galectins-3 and -4.^{180,181} Its sialylation in α2,3-linkage renders binding to galectin-8¹³⁰ and siglecs, especially sialoadhesin (siglec-1)¹⁸² and siglec-9,¹⁸³ possible, disialylation (at the Gal and GalNAc moieties) to siglec-4.¹⁸⁴ In the framework of the system of O-glycosylation, the α2,3-sialyla-

tion of the core 1 disaccharide precludes its branching to yield core 2 glycans, as core 1 disaccharide synthesis makes core 4 production impossible (►Fig. 6). Of fundamental importance, the decisions made by committing steps at the various stages highlight the broadness of the glycan panel and the enormous adaptability of the glycan profile, strongly suggesting their influence on regulating cellular activities. Core 2 branching and sLe^x production, as shown in ►Fig. 6, are steps to build selectin counter-receptors such as sLe^x noted above. Interestingly, the Golgi-resident sialyltransferase-IV-dependent final step of sLe^x synthesis has also been revealed to be important for chemokine (CCL3 and CCL4) binding to the CCR5 receptor¹⁸⁵ and for CXCR-2-triggered firm leukocyte arrest after CXCL1/CXCL8 injection in cremaster muscle venules.^{186,187}

Looking at regulation of galectin binding to mucin-type O-glycans more closely, an analogy can be drawn to the concerted actions described above for N-glycan α2,6-sialylation and galectin-1. Decisions between routes of synthesis explain how growth-regulatory processes are switched on or off.^{181,188,189} The role as a metastasis suppressor of the glycosyltransferase N-acetylgalactosaminide α2,6-sialyltransferase-2 (ST6GalNAcT-2) in breast cancer had been attributed to reduced T(F) disaccharide presence that lowers breast tumour cell aggregation and retention at metastatic sites via galectin-3.¹⁹⁰ Effectively, this type of α2,6-sialylation diverts product generation from reaching, for example, core 1 to sT_n, as shown in ►Fig. 6 and already mentioned above. Convergent effects on extent of presence of this disaccharide by altering its status of α2,6-sialylation and core 2 branching, too, favour lung cancer formation.¹⁹¹

Next, like selectins, galectins can home in on branched O-glycans, selectins to the sLe^{a/x} termini, especially galectins-1 and -3 to the internal LacNAc repeats of the β1,6-branch. Susceptibility of activated T cells,¹⁹² lymphoma cells¹⁹³ and prostate (LNCaP) carcinoma cells¹⁹⁴ to apoptosis induction by galectin-1 and impairment of natural killer cell activation by the binding of galectin-3 to tumour cells¹⁹⁵ are examples for the significance of galectin pairing with this part of the β1,6-linked branch of extended core 2 O-glycans shown in ►Fig. 6. Branch extension (and terminal tailoring by fucosylation, sialylation and sulphation to obtain sulphated sLe^{a/x} epitopes) is thus crucial to convey the signal for lectin binding.

In aggregate, any modulation in enzyme, substrate and acceptor availability can shift the relative proportions of the products, as can be deduced from ►Figs. 4 and 6 (for gangliosides as counter-receptors of tissue lectins, please see Ref. 196; for glycan–glycan interactions, please see Ref. 197). Considering occurrence of competition for an acceptor at different sites of the illustrated pathways of N- and mucin-type O-glycosylation, the extent of impact is not fixed but regulatable. As a consequence, cellular responsiveness to tissue lectins is intimately tunable, giving a functional dimension to the noted complexity of the glycome. Clearly, the potential for regulatory events will be broadened, if the lectin side is also a platform for fine-tuning, at the levels of presence and of structural design. The mentioned diversification of an ancestral gene is a main route towards this aim, and ►Fig. 5 documents changes in the length of the stalk

among the three selectins. Molecular diversity can also be achieved (1) by sequence changes in the CRD affecting the profile of ligands among members of a family and (2) by alterations of the modular design. Hereby, emergence of functional antagonism due to differences in modular design is possible. For this case, galectins-1 and -3 serve as role model, as already mentioned above.^{101,198} This emerging concept of functional crosstalk is a driving force to characterize in detail the lectins' structures, and this work raises awareness to address a fundamental question: what is the significance of modular architecture? Whereas lectins with trans-membrane sections such as selectins (please see ►Fig. 5) are ideal for cell–cell bridging, the situation is much less clear for lectins without such a module.

Lectin Diversity: The Example of Galectins

Vertebrate adhesion/growth-regulatory galectins are organized in three types of modular architecture (►Fig. 7). The manifestation of a strict selection process towards the three illustrated forms serves as instructive case for further work to answer the given question. As common approach of structural analysis, crystallography has been brought to its limits for the galectins.

Whereas the prototype (homodimeric) galectins and most CRDs obtained by engineering could readily be analysed by crystallography, often in complex with ligands and even within a broad range of pH values,^{133,199} intra-molecular dynamics of the tandem-repeat- and chimera-type proteins is a likely reason that precluded obtaining crystals for the full-length proteins of these two groups. Recently, partial truncation of the N-terminal segment of human galectin-3 by engineering facilitated to take structural analysis of crystals beyond the CRD.^{200,201} Like the three selectins, this galectin had first been detected by monoclonal antibodies (clones M3/31 and M3/38),²⁰² immunohistochemically present in macrophages, dendritic cells and in epithelium, thus termed the Mac-2

antigen.^{203–205} Its ligand spectrum covers glycans and proteins,²⁰⁶ its trimodular design shown in ►Fig. 7 enables aggregation to oligomers via contacts between the N-terminal tails and/or the CRDs in the presence of multivalent ligand^{207–210}; please see Flores-Ibarra et al²⁰¹ for review of the literature). This manifestation of a strict selection process towards the three forms shown in ►Fig. 7 is not only a suited test model, but it also inspires routes to redesign nature to clarify structure–activity relationships.

Looking at an involvement of galectin-3 in thrombosis, together with its counter-receptor galectin-3-binding protein (Mac-2 BP/90K; an adhesion mediator interacting with β_1 -integrin sub-units, fibronectin, nidogen and collagens IV–VI),^{211–213} it has been suggested to play a critical role in venous thrombosis.²¹⁴ The appearance of Mac-2 BP/90K as contaminant in preparations of recombinant coagulation factor IX was therefore judged to be cause of 'unforeseen consequences'.²¹⁵ Its pro-inflammatory signalling function is not only active in osteoarthritis pathogenesis,¹⁰² but, for example, also locally in unstable plaque regions of carotid endarterectomy²¹⁶ or rheumatoid synovium.²¹⁷ The molecular and topological nature of the active galectin-counter-receptor lattice for outside-in signalling remains to be defined. As the role of the recently discovered appearance of galectin-3' CRD in heterodimers with prototype CRDs²¹⁸ warrants elucidation.

When comparing the design of the homo-/hetero-bivalent family members, the structural difference between the non-covalent homodimers and the heterodimers with linker peptides between the two different CRDs is clear. However, the relationship from structure (e.g. type of CRD or length of linker) to function is again largely unexplored. Here, the combination of rational protein engineering, for example, turning galectin-1 into a covalently associated variant connected by the linker of a tandem-repeat-type galectin,²¹⁹ with activity assays, for example, platelet activation with galectin-8 as positive control, offers an attractive approach

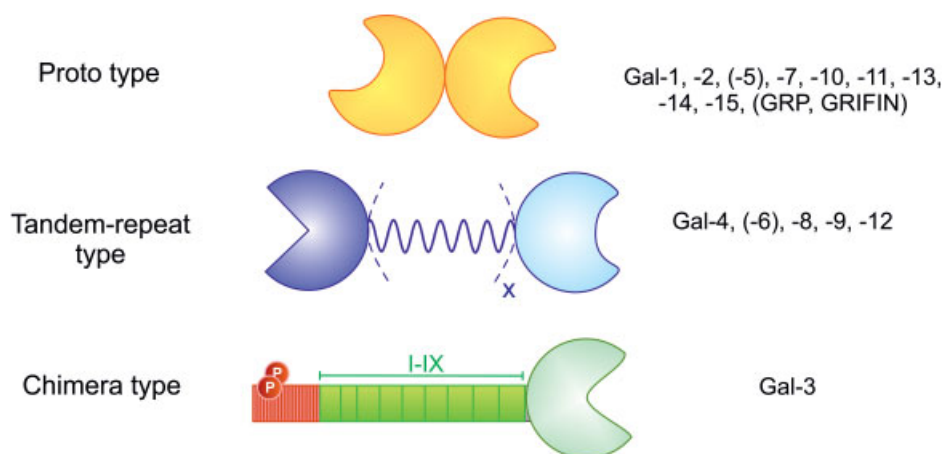


Fig. 7 Illustration of the three types of modular architecture of vertebrate galectins, i.e., non-covalent association of two identical carbohydrate recognition domains (CRDs) to a homodimer (prototype), covalent association (by a linker of distinct length; it differs between galectins, and alternative splicing brings about length variation in certain cases, e.g. galectin-8 with two lengths at 33 or 75 amino acids) of two different CRDs (tandem-repeat type) and the trimodular product of N-terminal association of a peptide with two sites for serine phosphorylation, a repeat region of non-triple-helical collagen-like peptides (nine in human galectin-3) and the canonical CRD, termed chimera type, with listing of the respective numbers/acronyms in the galectin nomenclature in each group (right side).

for thorough analysis, with the perspective to develop innovative antagonists or agonists on the platform of human galectins and beyond.

Conclusion

In retrospect, there has been ‘the widely held belief that carbohydrates are dull compounds, and that they serve only as structural or protective materials (e.g. cellulose in plants and chitin in insects) and as an energy source (glycogen in animals), but lack any biological specificity. The possibility that living organisms form a myriad of compounds in which carbohydrate is covalently linked to protein, with the carbohydrate having manifold functions, was ignored by most chemists and biochemists alike’.²²⁰ Ironically, when it was realized, the exceptionally demanding character of the task to handle analysis of glycans, called complex carbohydrates for a long time, required enormous efforts to begin reaching firm conclusions on structure–activity relations.

Our introductory focus on the amazing structural talents of carbohydrates to accomplish high-density coding has laid the foundation to understand the paradigmatic change in the way glycans are viewed. The convergence of the great strides made in glycan analysis with discovering a matching diversity of endogenous receptors (lectins) explains why the field of functional glycomics evolved rapidly in the last decades. Much as glycophenotyping with plant lectins, now also done with tissue lectins, has been a boon for sensing the dynamics and spatio-temporal control of representation of distinct glycan epitopes,^{221–227} the concept of their functional pairing with endogenous receptors is becoming a gateway towards understanding the workings of this aspect of the sugar code. The case study of the selectins has already taught the amazing lesson that platelet activation and inflammatory stimuli set genetic reprogramming and transport processes in motion to let functional pairing happen at the right place and at the right time, and selectins as representatives of C-type lectins are not alone.

The range of tissue lectins revealed to be involved in the mentioned processes is steadily growing, underscoring the nature of glycans as biochemical messages, as noted for inflammation.²²⁸ The co-evolution of messages (glycans) and readers (lectins) as well as their apparently intimately orchestrated expression justify the term ‘third alphabet of life’ for carbohydrates shown in ► **Fig. 1**.

With these central points made, it is reasonable to advise that the possibility of functional activity of the glycans should be considered, when dealing with a glycoprotein. Looking, for example, at the glycoprotein VWF, C-type lectins, a siglec and galectins are capable to home in on its glycan(s) (for overview of VWF binders, please see Ref. 229), and new aspects in the field of lectins are being connected to haemostasis, for example, the lectin pathway of complement activation in crosstalk to coagulation.^{230,231} Among the open challenges thus are a delineation of counter-receptors for lectins as well as of lectin networking within and between families, hereby deservedly putting members of families beyond selectins firmly on the map of receptors acting in concert in haemostasis and thrombosis.

Conflict of Interest

None declared.

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