VIP36 preferentially binds to core-fucosylated N-glycans: a molecular docking study

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Abstract

Alpha 1-6 fucosyltransferase (Fut8) is known for its properties as an enhancer of nonsmall cell lung cancer metastasis and as a suppressor in hepatocellular carcinoma cells (Hep3B). Promising candidates of affected molecules include E-cadherin. In its absence, during epithelial-mesenchymal transition, the pathway triggers signaling to the nucleus via β-catenin-TCF/LEF. Contrarily, in less metastatic tumors, Fut8 stimulates cell-cell adhesion. Regulated classes of molecules could also include the sorting machinery of polarized epithelial cells, sorted ligands or both, that may be altered in cellular transformation. I have analyzed here the cargo receptor VIP36 (Vesicular-integral membrane protein of 36 kD) for carbohydrate interaction. It has been described as a lectin in the ERGIC (endoplasmic reticulum-Golgi intermediate compartment), Golgi apparatus and plasma membrane. The docking reveals top-interacting carbohydrates of the N-glycan and O-glycan class that encompass N-linked glycans of high mannose and equally complex type which likely function as sorted ligands in epithelial cells. O-glycans score lower and include core 2 residue binding. I show that fucose core modifications by Fut8 stimulate binding of N-linked glycans to VIP36, which is known to be different from binding of galectins 3 and 9. This suggests that Fut8-upregulation may directly alter the affinity of sorted cargo and may enhance the sorting to the apical pathway as exemplified in hepatocytes and traffic to bile. High affinity binding of the ganglioside GM1 carbohydrate headgroup to VIP36 suggests a linkage with protein and glycosphingolipid apical transfer in epithelial cells. Thus, this fundamental approach with large scale docking of 165 carbohydrates including 19 N-glycan high mannose, 17 N-glycan hybrid, 9 N-glycan complex, 17 O-glycan core, 27 Sialoside, 25 Fucoside and 51 other glycan residues suggests, that linked cargo-receptor apical transport may provide a path to epithelial polarization that may be modulated by core fucosylation.
Background

VIP36 was identified as a component of post-Golgi secretory vesicles (Fiedler et al 1994) and had been shown to be involved in sorting of glycosylated alpha-amylase in the exocytic pathway (Hara-Kuge et al 2002; Hara-Kuge et al 2003; Arshad et al 2013). In polarized epithelia (Mellman and Nelson 2008; Rodriguez-Boulan and Macara 2014), it had been established, that glycosylation of growth hormone was important for apical sorting (Scheiffele et al 1995); erythropoietin and apical membrane proteins were also found to require glycosylation for apical targeting (Kitagawa et al 1994; Gut et al 1998). The glycan affinities to VIP36 had been a matter of discussion.

The VIP36 protein is similar to ERGIC-53 (Itin et al 1995) and belongs to the class of eukaryotic leguminous-type lectins. It is abundantly expressed in mouse kidney, liver, intestine, lung and spleen (Fiedler and Simons 1996). In humans, it is abundantly present in testis, epididymis, seminal vesicles, placenta and thyroid gland, and also expressed in liver, gallbladder, pancreas, stomach, cervix, endometrium, fallopian tube, bone marrow, lymph node, tonsil, brain and lung (Uhlén et al 2015). In computational analyses it had been demonstrated to be related to the plant lectin in Bauhinia purpurea (Fiedler et al 1994; Fiedler and Simons 1994). Another lectin, galectin-9, has recently been implicated in apical sorting and was found to influence the cell polarity of MDCK (Madin-Darby Canine Kidney) cells (Mishra et al 2010). The relationship of both lectins, if any, remains an open question. Galectins 3 and 4 have also previously been shown to be involved in apical transport in MDCK and HT-29 (enterocyte-like) cells, respectively (Delacour et al 2006). In particular galectin-3 which is localized to the cell surface, the Golgi and the cytoplasm is found to play a role in tumor metastasis (Takenaka et al 2002). This protein is found as a binding partner of β-catenin-TCF/LEF which co-localizes to the nucleus of cells and stimulates transcriptional activity (Shimura et al 2004). Is it plausible that several lectins are involved in generating the epithelial lipid and protein polarity that leads to physiological vectorial transport while alternatively stimulating cancer progression? Wnt secretion, the molecular ligand of the Wnt/frizzled-pathway, involves N-linked glycosylation and has recently been shown to benefit from complex glycosylation (Wnt11) in the apical transport, and surprisingly may require galectins and Opossum, a p24 protein, for secretion (Fiedler et al 1996; Buechling et al 2011; Yamamoto et al 2013).

The glycoproteostasis involves ER enzymes and lectins calreticulin (CRT) and calnexin (CNX) that serve to assist the ordered glycoprotein folding and quality control and ER-associated protein degradation (ERAD)(Helenius and Aebi 2001; Satoh et al 2010; Hebert et al 2014). VIP36 had previously been demonstrated to be localized to the Golgi apparatus, exocytic carrier vesicles and the cell surface and lends itself to a sorting function in the Golgi complex (Fiedler et al 1994). The endogenous protein was then also visualized in punctate structures that had been co-localized with the ER-Golgi
intermediate compartment (ERGIC) constituent ERGIC-53. The labelling of COPI or sialyl-transferase of the trans-Golgi cisterna did further demonstrate substantial overlap as shown in MDCK cells (Fullekrug et al 1999). In contrast to MDCK cells, where in EM the Golgi apparatus and the plasma membrane were preferentially decorated with anti-VIP36 antibodies (Hara-Kuge et al 2002) it was found in other epithelial cells, that, moreover, endocytic structures and late endosomes were labeled (K. Fiedler and K. Simons, unpublished results). Immature granules and mature secretory granules as well as the trans-Golgi apparatus were decorated with antibodies in parotid secretory cells as seen in EM (Shimada et al 2003).

VIP36 and ERGIC-53 contain a single carbohydrate recognition domain (CRD). The D1 arm of mannoses linked to Asn-N-sequons was found to form extensive hydrogen bonds with VIP36 in the crystal 2DUR (Satoh et al 2007) and interestingly, the coordination of Ca\(^{2+}\) was found to orient the side chains that had previously been speculated to be involved in carbohydrate and calcium binding based on this similarity (Fiedler et al 1994), i.e. Asp131 and Asn166 as well as His190. Some variation may be introduced by the largest variability of protein sequences of VIP36s and related proteins in the glycan and calcium-binding loop that we described as region II (Fiedler et al 1994). Next to conserved amino acids (Asn166 in VIP36 and ERGIC-53, VIPL/LMAN2-L; not in mouse or human ERGL/LMAN1-L and yeast EMP46-like; Table S3) (see (Neve et al 2003)) in representative members this shows gaps. Although the loops are usually not ascribed particular functions, it could be that the large evolutionary glycan variation may be adapted by amino acid residues in the vicinity of the binding site.

Galectin-9 contains two CRDs and the N-terminal CRD was described to interact with polylactosamines and some glycosphingolipid (GSL) headgroups (Hirabayashi et al 2002; Nagae et al 2008; Nagae et al 2009; Mishra et al 2010). An activity as an eosinophil chemoattractant had also been discovered (Sato et al 2002). A link of cargo receptors to specific bilayer lipids was also described: it was found that sphingomyelin 18 interacted with the transmembrane domain of the p24 protein and regulated ER to Golgi transport (Fiedler and Rothman 1997; Contreras et al 2012). The critical glutamine residue contained within the bound signature sequence was previously found to halt export of p24 chimeras in the early secretory pathway once mutated to alanine.

Here the large scale screening of carbohydrates by computational docking (Trott and Olson 2010) replaces the glycan screens usually developed for microarrays or mass spectrometry which due to technical difficulties do usually not generate nearly complete solutions. The present data may contribute to shed light on the role of fucose in carbohydrate N-glycan binding of the L-type lectins and possible indications of branch preference of GlcNAc-transferases with respect to apical sorting of glycoproteins. The results from cognate docking suggest that GlcNAc-Transferase V (GlcNAcT-V) modification of the α6 mannose arm of N-glycan chains is not of high affinity, however, the GlcNAcT-IV
modified glycans with GlcNAc added to the α3 mannose arm may represent the preferred high affinity cargo.

N-glycan chains are modified by adding fucose residues to their terminal chains (fucosyltransferase 2; Fut2) or to the N-glycan core (fucosyltransferase 8; Fut8). Recent discovery of the influence of Fut2 activity on the protective role of fucosylation suggests a function of terminal modifications in host-microbial interactions (Pickard et al 2014; Goto et al 2014) whereas core fucosylation by Fut8 has been described as possible sorting signal in protein traffic in hepatocytes to the bile duct (Nakagawa et al 2006). Fut8 was, furthermore, shown to regulate metastasis in nonsmall cell lung cancer (NSCLC) but overexpression in hepatoma cells (Hep3B) was demonstrated to suppress intrahepatic metastasis in athymic mice (Miyoshi et al 1999; Chen et al 2013). The E-cadherin-β-catenin-TCF/LEF pathway and α5β1 integrin have been invoked in these analyses. The relation and intricate balance of cell-cell and cell-extracellular matrix adhesion to protein fucosylation (and other glycan modifications) may explain the latter two findings and glycan modifications with respect to protein traffic and polarity of epithelia.
**Results**

**Lectins**

Next to legume L-type lectins, CNX, CRT, Emp47, pentraxins (C-reactive protein, serum amyloid P) and vp4 of rotavirus also belong to L-type lectins (Fiedler and Simons 1994; Neve et al 2003; Etzler et al 2009). An overview of a structural comparison is shown in Table 1. Galectins, although only up to 10% identical in amino acid sequence, show a similar fold to VIP36. The carbohydrate binding site is, yet, removed from the site of ERGIC-53 and VIP36 glycan interaction. VIP36 an ERGIC-53 are 30.4% identical in sequence within the L-type lectin domain. Galectin-3 is only 10.1% identical in sequence to VIP36, and galectins do not contain a calcium binding site and coordinate glycans on the large β-sheet, i.e. not via loops (see Fig. 3, 7). In the present analysis I have restricted the molecular docking to VIP36.

**Cognate binding to VIP36**

The intactness of carbohydrate structures was checked and confirmed. Structural docking was simulated with PyRx (Wolf 2009), utilizing the VIP36 2E6V and 2DUR structural data (Satoh et al 2007). 2E6V was initially analyzed in Global docking (Fig. 1). Library residues of high mannose type bound with maximal affinities of up to -6.7 kcal/mol to docking sites, that in this global approach did not correspond to cognate docking. Only 13.6% of all high mannose ligands showed at least one cognate docking site, and 4.4% of high mannose ligands preferentially interacted with the authentic, apo-ligand binding pocket. None of the complex hybrid and only sialylated or fucosylated N-glycan termini, possibly of N-glycan complex type, bound preferentially in global docking, just as only some O-glycan cores would preferentially interact with the cognate binding pocket with considerate affinity. The global approach was thus abandoned and structures were, furthermore, used as energy minimized conformers generated by conjugate gradient minimization steps. The local approach resulted in 49%, 42% and 42% of all glycans interacting with the main binding pocket of 2DUR-A, 2DUR-B and 2E6V, respectively, at or better than -6.5 kcal/mol. In these models, a single calcium was preserved in the binding assay instead of adding a further calcium which would have only loosely bound by 3 coordinations in a similar position to the 2nd calcium of the VIP36-like ERGIC-53 (Velloso et al 2003). The previously determined Kₙ of VIP36 for calcium in the exoplasmic domain was found to be approx. 2.5 x 10⁴ M⁻¹ (Fiedler and Simons 1996) which is actually quite similar to the determined carbohydrate affinities in the range of 1 -2 x 10⁴ M⁻¹. Ligands interacting at or better than -6.5 kcal/mol represented possible candidates for true binding site interaction.
Top scoring high-affinity interactors of VIP36 belong to the N-glycan, Ganglioside and O-glycan class

In order to fully explore the library Carb165 the 2DUR-A conformer was used. ViewDock was utilized to analyze H-bonds in between the modeled ligand and the holo-enzyme. The top scores of 15 Carb165 (Fig. S1, I-XV) glycans were presented according to motifs included (Fig. 2); lacto-motifs were prevalent with five hits among the top 15. The interaction energies of raw, automated docking are graphed in Fig. S1 with top model numbers indicated that fulfill the structure test. The energies of these models differed slightly.

The series of carbohydrates that bound with considerable, high affinity to Carb165 consisted of N-glycan high mannose and N-glycan core basic structures classified as complex glycans, followed by GM1 (Simons and Ikonen 1997; Sabourdy et al 2008), N-glycan high mannose and the lactosamine motif in the third rank. Using the structural criteria, the basic order of affinities does not change with the 3rd ligand scoring number one amongst the top 15 (N-glycan high mannose) and the 13th glycan scoring number two of the top selection (N-glycan core basic: complex and fucosylated). In the 9th position of eligible carbohydrates extended O-glycan core 2 was found in rank number three using the structure tests (see Fig. S2; IX). Only the 9 top scoring models of each carbohydrate were tested. In a separate simulation by applying identical experimental parameters, β-D-GlcpNAc-(1-6)-[β-D-Galp-(1-3)]-α-D-GalpNAc-OH, the core 2 structure, as well as the T antigen, β-D-Galp-(1-3)-α-D-GalpNAc-OH, or Sialyl-Tn antigen, α-D-Neu5Ac-(2-6)-α-D-GalpNAc-OH was scoring less than the extended core, suggesting that the extensions of these O-glycans included relevant binding interactions.

The ganglio series GM1 (IV) was one of few GSL motifs in the entire database Carb165 (depleted in GSL residues). The GM1 headgroup is present, among other species, in Campylobacter jejuni as part of an epitope widely studied in human disease of autoimmune disorders caused by these bacteria (see Table S1). Further screening may later lead to precise determination of all glycosphingolipid affinities and binding specificities including their frequent H-bond acceptors and donors that may vary from other carbohydrate residues.

Hydrogen-bonding of VIP36 to carbohydrate residues

The virtual screening approach was further used to determine the exact position of amino acids in VIP36 that would provide the bulk of hydrogen bond donor and acceptor activity. Docking in this gradient approach was described to function akin to “machine learned” binding (Trott and Olson 2010) applied to statistical physics (Fig. 4). Results are summarized for measuring H-bonds in ViewDock with all 15 top scores of Carb165 against 2DUR-A. The overwhelming proximity of oxygens
(3 Å zone distance) shows the large H-bond donor capacity of carbohydrate ligands with few nitrogens in molecular proximity of the VIP36 surface (Fig. 3). The exact method used applies tolerances to H-bonding geometry that were found in crystallized structures (Mills and Dean 1996). The most probable ligand of carbohydrates in the VIP36 binding site is represented by the ND2 atom of Asn166. This ligand site is found within the central binding area at 11 o’clock and had previously been identified due to lectin homology of VIP36 to leguminous lectins (Lec_Baupu) (Fiedler et al 1994). This metal binding site appears to be conserved in leguminous lectins, ERGIC-53 and close VIP36 family members (L-type lectins). Above 41% of all 15-9 glycan model structures (Fig. 3) generated by AutoDock form hydrogen bonds wherein the aforementioned residue act as donor. This was followed by Asp261 N and the Ser96 OG located both on the perimeter of the pocket in diametrically opposite position at the bottom of the central binding cavity. 14.4% and 10.0% of all carbohydrates use these as donors. The acceptor binding sites are more heterogeneous with five residues covering the largest share of 55.3% hydrogen-bond acceptors in total, the Pro146 O residue (13.1%) as seen on the upper right of the perimeter of the central indentation, the Asn166 O (11.6%) which is bidentate (Asn) located on the upper left side above the central trough, the Glu98 OE2 (10.9%) on the right of the binding pocket, the Thr259 O (10.2%) within the central indentation and Asp131 OD1 (9.5%) located just next to Asn166 on the upper perimeter of the central indentation (Fig. 4A). Large carbohydrates that bind to the central cavity form interactions using H-bond donors that cover an area of 700 Å² coinciding with a modeled N-glycan of high mannose type (Satoh et al 2007). This was inserted in a non-flexible fashion and corresponds herein to topology 3 (Figs. 4B, S2).

**Contrasting conformers: Ranks and Differences**

The subgroup of complex N-glycans scores on average in the first position versus the grouped N-glycans of high mannose type (rank 2), Fucosides (rank 3), Sialosides (rank 4), N-glycan hybrid (rank 5), O-glycans (rank 7) and Other carbohydrates (rank 8) (Fig. 5). The plot in Fig. 5 combines the results of all conformers that have been here analyzed.

Complex N-glycans clearly differ from all other grouped glycans with significantly higher binding affinities (p<0.04)(Table S2A, B), except for the difference of these to N-glycans of high mannose type (p=0.07). Fucoside affinities also differ from some other glycan affinities at significant cutoff (p<0.04), just N-glycans high mannose, Sialosides and N-glycan hybrid fall in the same range of binding strength.

Therefore, among N-glycans significant differences can be established in this docking procedure. Provided the top scoring list prevails in cluster-multiplied approaches, some molecules previously
shown by select affinity-chromatography (N-glycans) (Kamiya et al 2005) are also revealed in this simple approach. O-glycans do significantly differ in binding strength relative to N-glycans of complex and high mannose type, Fucosides and Sialosides (p<0.02), which all show higher affinity.

**Topologies of putative glycan interactions**

Concerning the interaction of glycans of the Carb165 collection, three different topologies can be discerned. One set of glycans enters the binding cavity from Glu98 described as topology 1 or further 1b (7 / 15). Another group of Carb165 glycans approaches the binding site from His190 in topology 2 (2 /15)(Fig. 4B). Topology 3 shows the carbohydrate approaching the cavity from Tyr164 towards the centre of the binding cavity, towards the bridge in between Asp131 and Gly260 (6 / 15).

Whereas the topology seems rather simple, some O-glycans, and possibly many other low scoring Carb165 residue models, may also contact the binding site from within the central area projection with the reducing end GalNAc residue being elongated towards the centre and multiple indentations.

**Select glycan interactors**

Considering the interactions listed in Fig. S2 a number of interesting features of bound carbohydrates should be mentioned. The preferential interaction with N-glycans of high-mannose type is restricted to intermediates of the Golgi-Mannosidase (Golgi-Man) I, an enzyme which is localized to the cis-Golgi (Stanley et al 2009), and is not prevalent for Glc-containing high-mannose precursors. The latter have been implicated in protein folding and quality control (Hebert et al 1995). The final product of a fully trimmed and shortened high-mannose glycan, Man6-[Man3]-Man6-[Man3]-Man4-GlcNAc4-GlcNAc (9; Fig. S3A) binds to VIP36 with distinctly lower energy. The product of endomannosidase, that may function as a failsafe device to cleave residual Glc-Man residues of proteins in the Golgi (Roth 2002), interacts with VIP36 with some intermediate affinity (6). Also the product of α-mannosidase III binds with intermediate affinity, and is itself a substrate for lysosomal degradation (8). Alpha-mannosidase III is thus far described as an alternate backup in most cell types in mice if Golgi-ManII fails (Chui et al 1997; Moremen 2002). Yet, α-mannosidase III cannot fully replace Golgi-ManII since animals develop immune complex diseases in Golgi-ManII null mice (Haltiwanger and Lowe 2004).

A surprising finding is the high affinity of a Golgi Mannosidase II glycan (Fig. S3B) product (22) as well as its substrate (21) to VIP36. Elongated hybrid glycan chains also interact that had not previously been trimmed, although at lower affinity. However, since both, the substrate and product interact at high affinity, the reaction may have reached a “watershed” point of sorting where modified and not yet fully glycanized proteins are subjected to bimodal sorting (Fiedler et al 1996) and bound for
backward traffic or the forward route of intra-Golgi transport. How galectin interactions would affect sorting decisions in the Golgi apparatus at this transport stage remains speculative (but see below). These may be stimulated by further GlcNAcT-V (Mgat5) activity in the Golgi cisternae which may enhance glycoprotein binding.

Complex glycan preferences are also considerable with the GlcNAcT-IV product being of second highest energy of interaction (38; Fig. S3C)(kcal/mol). Simple, biantennary complex N-glycans have high but complex tetraantennary products with a bisecting GlcNAc introduced by GlcNAcT-III have lowest energies of binding. If the VIP36 receptor had a sorting role, fucosyltransferase VIII (Fut8) and GlcNAcT-IV substrate turnover may be preferential for generating the ligands for inclusion in exocytic carrier vesicles / tubules. Comparing the top row of glycan affinities (Fig. S3C) it can be seen, that the top interacting molecule displays branching at the Man3 residue (38) but not Man6 (41) and is fucosylated unlike the third ligand (44). Interestingly, LRP-1 (LDL-receptor related protein-1) endocytosis and secretion into the bile may implicate core fucosyl modification as a sorting signal in both, the endocytic and exocytic pathway (Lee et al 2006; Nakagawa et al 2006). In a separate test using the identical docking parameters it was consistently found, that the top scoring complex N-glycan (β-D-GlcpNac-(1-2)-α-D-Manp-(1-6)-[β-D-GlcpNac-(1-2)]-β-D-GlcpNac-(1-4)-(β-D-GlcpNac-(1-4)-β-D-Manp-(1-4)-β-D-GlcpNac-(1-4)]-[α-D-Fuc-(1-6)]-β-D-GlcpNac) (38) with or without the core Fuc in α1-6 linkage was scoring at least -0.7 kcal/mol better with the fucose residue added.

The terminal end preferences are displayed in Fig. S3D. Not all enzymes that lead to processing modifications are known. The lactosamine motif seems of preferential energy of interaction, yet, competitive binding or bivalent interaction of ligands with multiple VIP36s seems the sole structurally plausible solution since the N-glycan core is of equal / higher affinity (see above). Comparing the terminal glycan preferences it is surprisingly found that residues β-D-GlcpNac-(1-3)-β-D-Galp-(1-4)-β-D-GlcpNac (VI; Fig. S1) show the overall highest affinity of only three linked monosaccharides, unlike the low scoring trimannose of e.g. 2E6V or bimannose of 2DUR-A each scoring with -6.2 kcal/mol. The score allocates rank 4 to the former trisaccharide in global numbering (VI; Fig. S1). In terminal ends, the fucose residues in α1-2 linkage do not increase the binding energy as evident in comparing mono-fucosyls present or absent on trisaccharides (51/56, 53/54; Fig. S3D). This indicates that a single fucose residue as such does not elevate the binding energy in this virtual screen. Furthermore, in N-glycans containing aforementioned trimannose residues, sugars surrounding these will contribute equal to or more than 20 % to the binding energy.
Glycan interactions in specific and “average” conformers

Since different conformers have been found in the unit cell and in different experiments of crystallization, a subset of these were here compared and summarized. Only single runs of complete libraries were analyzed (without the structure test). Comparing the 3 conformers 2E6V, 2DUR-A and 2DUR-B, the top 15 Carb165 glycans display little variability on average. Summing all 3 rank 3 is filled with a N-linked glycan of high mannose type and by a complex fucosylated N-linked glycan, is preceded by a glycan with the four motif characteristics Lewis C, Lactosamine, Neo-Lactosamine, GSL Lacto series (rank 2), and a fucosylated glycan containing the lactosamine motif (rank 1).
Discussion

The purpose of this study was to demonstrate carbohydrate binding of VIP36 and to analyze high-affinity interactors in a large-scale docking with 165 glycans including 9 N-glycan complex, 19 N-glycan high mannose, 17 N-glycan hybrid, 17 O-glycan core, 27 Sialoside, 25 Fucoside and 51 other glycan residues.

N-glycan interaction

In accordance with the previously discovered homology of VIP36, a cargo receptor, with leguminous plant lectins, the present study shows that the residues Asp131, Asn166 as well as His190 are involved in calcium coordination and provide H-bonds to glycans similar to the D1 arm Manα(1-2)-Manα(1-3) of a Man-Man-Man carbohydrate (Satoh et al 2007). Mannose residues in this library of long sized carbohydrates including branched glycans of N- or O-linked glycan modifications frequently bind to the central binding indentation of VIP36 with mannose residues of the D1 or D2/D3 arms within the Asp131 – Asn166 hydrogen bonding pattern (O-linked glycans did not contain mannose). The only residue that shows top scores of binding energy of the N-linked carbohydrate type is number II (Fig. S1 and S2), a complex carbohydrate that does not contact these residues via hydrogen bonds of their mannose groups but places the GlcNAc residues of the core (II) and terminal branch in the hydrogen-bonding distances. These binding interactions may have previously been overlooked in discoveries using serial lectin affinity chromatography (Kamiya et al 2005). In these studies a M8 high mannose ligand, a product of ER mannosidase II was binding to the lectin with highest affinity, in the present study the ligands with very high mannose substitution show less affinity with at least 1.1 kcal/mol energetic difference to the top scoring M7, an intermediate in the Golgi mannosidase Golgi-Man I pathway (Fig. S3A, 10). In a second analysis by lectin affinity chromatography the M8 high mannose residues also showed the highest affinity to VIP36 closely followed by M9 and M7. Whereas the binding of VIP36 was slightly pH-dependent, VIPL showed remarkable reduced affinity at lower pH and preferably bound to M9 (Kamiya et al 2008), an early intermediate. In the present study, the subsequent pathway intermediate M7 interacted with 1.6 kcal/mol higher affinity with VIP36 than the early form M9.

Consistent with the previous analysis, in this unbiased screening I also found the low affinity interactions of the monoglycosylated M9. Moreover, two-fold glucosylated high mannose glycans, substrates of endomannosidase, display less interaction with VIP36 than M9. Quality control mechanisms proposed for VIP36 in the early secretory pathway would here be unconfirmed and likely be unrelated to glucosyl-groups to maintain a transient regulatory cycle of protein folding quality control. This is proposed for the CNX/CRT ER proteins and their glyco-ligands in the secretory pathway.
While this glucosylation pathway seems excluded for VIP36 based on carbohydrate affinity, a dual function of VIP36 could include the “watershed” point of sorting with both, the Golgi Mannosidase II substrate/intermediate and Golgi Mannosidase II product of equal affinity (Fig. S3B). These two carbohydrates contain one GlcNAc residue each and the VIP36 lectin affinity may be used to halt the secretion of transmembrane / secreted forms of glycoproteins in the medial Golgi up until further modifications ensue. These modifier reactions could install the second GlcNAc group on the α1-6 mannose branch by GlcNAcT-II and could then allow for transport and processing in the Golgi. This mechanism would ensure that accessible N-glycans become more highly glycosylated and are converted to complex-type glycan forms. A number of other scenarios could be envisioned using fucosyl modifications and further addition of carbohydrate termini as modifiers of the bulk transport in quality control. It is expected, that later extended modifications (Stanley and Cummings 2009) are primarily added to the α1-6 branch of the N-glycan since α1-3 branches shows less mobility as demonstrated in the recent RMSD analysis (Jo et al 2013). The proper installed GlcNAc on the α1-6 branch would then, once further processed from bi-antennary to tri-antennary preferentially on the other branch (see below), once more bind to the VIP36 and be transported or apically sorted since this complex glycan displays highest affinity (Fig. S1).

Another interesting family of proteins are the p24 proteins (Fiedler et al 1996; Fiedler and Rothman 1997). These, also known as EMP24/GP25L/Erp, are another class of cargo-receptors that have been described in COPI-coated vesicles and in the Golgi complex, and newly described functions include cargo clustering and recognition of glycosylphosphatidylinositol (GPI)-anchored proteins (Theiler et al 2014). Thus multiple lectins are likely found in the secretory pathway that each may regulate inclusion into transport vesicles and mediate sorting. In Drosophila Wnt proteins require Evenness interrupted (Evi/Wls) for transfer to the plasma membrane while at the same time p24 proteins are necessary for anterograde traffic - glycosylation is furthermore shown to be the targeting signal in apical transport: modifications of Wnt11 include complex N-glycosylation, it is secreted apically whereas Wnt3a which is only high-mannose modified is transferred to the basolateral cell surface in MDCK cells (Buechling et al 2011; Yamamoto et al 2013). Wnt3/5 in Drosophila is also N-glycosylated, yet, the role of its glycan modification in the early secretory pathway has not been clarified.

**Glycosphingolipids in development**

The third scoring glycan included the GSL ganglioside GM1 in this screen. Assuming that glycosphingolipids are involved in apical delivery as originally envisioned (Holthuis et al 2001), it is tempting to speculate that this interaction is causally related to raft sorting at the exit stations of the Golgi apparatus. The benign effects of mouse β1-4-GalNAc-Transferase knockouts with abolished GM1 synthe-
sis, that only showed effects late in mouse life would have to be explained in view of high affinity GM1 interaction. If sorting platforms would use these together with VIP36 as adaptors, the mouse would alternate these with other raft GSLs. In this model of hexosaminidase deficient and specific knockout mice, abundant GSL GM3 and GD3 were found, but no ganglioside GM1 production (Proia 2003; Furukawa et al 2004). Advances in subcellular localization of glycosphingolipid biosynthetic reactions have pin-pointed to the trans-Golgi for ganglioside and globoside synthetic machinery, and vesicular intra-Golgi transport been ascribed to the former but not to the latter: the TGN was ascribed a role for FAPP2-mediated glucosylceramide (GlcCer) import which is the substrate for specific globoside biosynthesis (Gb3 synthase etc.) and lactosylceramide (LacCer) is synthesized earlier in the Golgi complex if channeled into the ganglioside pathway (D’Angelo et al 2013).

It is rather unlikely that globosides may replace gangliosides in VIP36 binding since globosides are not essential for mouse development and e.g. neither is GM3 - in a VIP36 or raft-based model of sorting this may be proposed. Yet, infants bearing a point mutation in the GM3 synthase gene develop an early-onset epilepsy syndrome (Jennemann and Groene 2013). Most importantly, it has been shown that the Lc3 synthase (β1-3-GlcNACT-V) gene is developmentally lethal in pre-implantation in mice (Biellmann et al 2008), presumably affecting events as early as adhesion of cells (Hakomori 2008). This would suggest the pivotal role of lacto-series / neo-lacto series of glycosphingolipids. Later analysis showed no essential requirement of the Lc3 synthase in transgenic mice deletion, since mice survived to term (Kuan et al 2010). It is possible that in these three studies (see Review (Jennemann and Groene 2013)) a switch from lacto- to globo- and ganglio-series of GSLs had impact on the cellular growth and differentiation or strain differences contributed to the different results.

Moreover, it is possible that complications affect interpretation of genetic differences and variation possibly due to the different contribution of N-linked glycan chains and their modification since these or both, glycans of lipids and proteins, are used as primarily sorted products. Thus, could VIP36 provide the adaptor for linking the “sorted glycan” and the sorting coat? The raft model includes the sorting of proteins and these may attach to lipids as co-sorted cargo. On the other hand, a raft may be distinguished from other membrane areas and sorted via receptors recognizing a lipid attaching to VIP36, which as sorting platform gathers co-sorted proteins among other cargo. Whether, in this model, VIP36 could provide a sink for 2 sorted ligand classes collectively segregated from other molecules remains to be studied. Since the same enzyme likely adds GlcNAc residues to nascent lacto- and neo-lacto series of GSL and to N-glycan chains it is difficult to predict the outcome of above described experiments with respect to sorting and availability of ligand. Furthermore, of interest are recent findings, that in addition to VIP36, galectins among other cargo receptors are involved in clustering and apical transport of glycoproteins (Mishra et al 2010).
The most likely candidate for an essential function of glycosphingolipids and their headgroup is glucosylceramide demonstrated with a gene Ugcg-knockout mouse. These mice are embryonically lethal and display defects in the ectodermal cell layer at E7.5 due to malfunctioning in GlcCer synthase (Yamashita et al 1999). GlcCer synthase expression is widely found and seems enhanced in this cell layer. It has previously been established that GlcCer is differentially sorted in the apical and basal pathways and this provides evidence for the lipid-based sorting machinery (vanIJzendoorn and Hoekstra 1999; Holthuis et al 2001). It is possible that only the combined effect of loss of all GSLs of the GlcCer series have these detrimental outcomes. The role of galactosylceramide (GalCer) synthase has been controversially discussed and studies in Ugta8a knockout mice have found survival for 90 days with severe demyelination of the central nervous system or death at 25-30 days after birth. Further into this branch of GalCers, it has been found that sulfatide formation by the sulfotransferase (Gal3st1) will also affect the nervous system in myelination as well as male fertility although later and less severe (Honke et al 2002).

Current insight would suggest that although there is an exquisite order of gene-regulation in development and sequential up-regulation of specific glycolipid biosynthesis at distinct time points the fundamental nature of the lipid bilayer and its sorting machinery depends on very few carbohydrates that are added to GSLs or functional redundancy may explain most observations that have been made so far.

**Branches of N-glycans and their lectin affinity**

One important result from this screen is the lower interaction of GlcNAcT-V (Mgat5)-branched N-linked glycans (Partridge et al 2004; Boscher and Nabi 2013) that show less binding than the top-binding GlcNAcT-IV-branched glycans, corresponding to the D3 versus D1 mannose arm without GlcNAc (see overview Figs. 6, 7). Since previous experiments in animals in tumor growth have pinpointed to the role of these enzymes and products as decisive modifiers of metastasis and focal adhesion signaling, it may be tempting to speculate on the function of carbohydrate branching with respect to conformational freedom, glycoprotein sorting and, as shown, membrane organizing functions (Granovsky et al 2000; Christiansen et al 2014; Stowell et al 2015). The conformational space of singly branched versus doubly or even triply branched polymers is restricted at each branch point and thus their wandering in space is limited to a smaller number of interactions. In this regard the Mgat5 branch is no exception but since the number of neo- or lactosamine groups added preferentially to the Mgat5 terminus is high, the number of interactions cannot be predicted but may change in their selectivity. If the sorting function that is proposed is fulfilled, VIP36 could bind to the GlcNAcT-IV branch of modified glycoproteins but less to the GlcNAcT-V branch of processed glycans and may properly sort the former but less so the latter. In contrast, the focal contact interactions...
that have been shown to involve the lectin galectin-3 may be stimulated by increased Mgst5 expression, and this may lead to maintaining a longer residence of Mgst5-modified glycans (Partridge et al 2004). Therefore, assembled complexes of epidermal growth factor (EGF) and transforming growth factor (TGF) receptor and galectin-3 may signal longer and promote the invasion of cells stimulating cell motility. The aforementioned role of galectin-3 in nuclear signaling via the β-catenin pathway that was demonstrated in tumor cell lines would furthermore contribute to the epithelial-mesenchymal transition (Shimura et al 2004). In tumour sections, galectin-3 is detected with the peroxidase-antiperoxidase reaction (Mac-2) and in contrast to its expression in the cell lines, the decrease of nuclear localization is associated with tumour progression (Lotz et al 1993). In colon carcinoma the interplay with other galectin family members or the stage specific alterations of cell growth, adhesion or attachment may be different.

While complex N-glycans are certainly important for growth since in GlcNAcT-I knockouts mice do not survive to term (Ioffe and Stanley 1994) not all of their functions were appraised. Interestingly, homozygous GlcNAcT-I/- cells did not participate in growth and the lung epithelial cell layer in the bronchus demonstrating the dependence on complex/hybrid N-glycans in adhesion (Ioffe et al 1996). Yet, recent results suggest that the oligomannose form e.g. of modified E-cadherin produces stronger cell-cell interactions than the complex or bisecting GlcNAc-modified form (Hall et al 2013) and that an explanation of the phenotype of the mice will require more molecular research. In the analysis of Lec1 versus Pro-5 and LEC10B cells it was suggested that the information content of glycans is being utilized in surface expression and organization. With respect to sorting it is likely that in polarized cells the glycome would be utilized for polarized apical targeting.
Conclusions

With respect to the sorting signals it is predicted that the core α1-6 linked fucose would enhance apical transport of N-glycosylated proteins once bound to the sorting machinery. Interestingly, it has been found that in hepatocytes this core fucose may be involved in apical transport (Lee et al 2006; Nakagawa et al 2006). Although speculative, it is possible that the core fucose modification is essential, but loss-adaptation can upregulate GlcNAc bisection by GlcNAcT-III (Wang et al 2005; Kurimoto et al 2014) that in glycan-receptor interaction would exert a similar function as the added core fucose.

The GlcNAcT-V products of N-linked glycans could be anchored to the cytoplasmic coat via soluble clustering receptors and ensuing cargo aggregation which may function akin to coincidence detection on the cytoplasmic face, including cytoplasmic domains of cargo-receptors and luminal soluble galectin-9. Since neither galectin-9 nor galectin-3 have previously been shown to preferentially interact with core-fucosylated N-glycans (Hirabayashi et al 2002) VIP36 may carry out its function by directly interacting with the fucose core. Carbohydrates interacting with each family member of L-type lectins in vivo remain to be studied in detail in the future.

The role of VIP36 or VIP36-like proteins in phagocytosis (Shirakabe et al 2011; Huang et al 2014) and its lectin function in clearance of *Staphylococcus aureus*, *Vibrio parahaemolyticus*, and *Aeromonas hydrophila*, or *Campylobacter jejuni* suggested based on the current results, may be studied in a further project.

Large scale screening including carbohydrate headgroups of glycolipids will require supercomputing in particular if multiple lectins and/or conformers are studied. It could be expected that cancer treatment and the understanding of cancer metastasis would benefit from this undertaking.
Methods

Structural comparisons were carried through with PDBeFold v.2.59 (Krissinel and Henrick 2004) (searched 8. Feb. 2015).

The docking approach was carried through with structures obtained from PDB (http://www.pdb.org) using the 3-dimensional (3D) coordinates published for 2DUR and 2E6V. The programme used for docking was AutoDock VINA (Trott and Olson 2010). The input and output uses PDBQT structure format. Since the coordinates for 2DUR-A provided ambiguous side-chain conformations that showed disorder in the X-ray analyses, the chosen conformers for aforementioned approach consisted of Glu98(A)-Asn183(A)-(Gly184(A))-Ser185(B) and Thr203(A) rotamers. Within the AutoDock implementation of PyRx version 0.8 from Sargis Dallakyan (http://pyrx.scripps.edu) the grid was set as indicated, which reached a minimal resolution of 0.375 Å for all local docking approaches. Only continuous runs of docking for all Carb165 residues in one session were evaluated (unless otherwise indicated in control simulations). The structural criteria chosen were the formation of double H-bonds to Asp131 and Asn166 for some panels as indicated; this negated often cited limitations of top scores to RMSDs of 0.0. The global approach necessitated a slightly lower resolution for the grid (sizes 51x51x40 Å). The programme utilizes OpenBabel (O’Boyle et al 2011) and AutoDock 4/VINA (Forli and Olson 2012) (http://autodock.scripps.edu) and an uff (united force field) for energy minimization, conjugate gradients with 200 steps and a cut-off for energy minimization of 0.1. Partial charges were added to receptors using PyBabel (MGL Tools; http://mgltools.scripps.edu). No limits to torsions were allowed which resulted in a maximum of attempted adjustments. Single CPU time was up to 150 hours for longest/branched ligands.

The carbohydrate collection “Carb165” was downloaded from the laboratory of Prof. Woods (Tschampel et al 2006) (http://www.ccr.c.uga.edu) (with multiple conformers), was selected and filtered for uniqueness against a laboratory collection of glycosphingolipid-glycan headgroups (GSL) that was built using the chemical descriptor-library of LIPIDMAPS and downloaded (http://www.lipidmaps.org) or the Glycosciences website (http://www.glycosciences.de).

H-bonding was determined with ViewDock and with tolerances 0.4 Å, 20° (Mills and Dean 1996). Annotation of carbohydrates was obtained from descriptions and species annotations from http://www.glycome-db.org including the MCS search routine (Ranzinger et al 2009). Drawing utilities were from GlycoWorkbench (Ceroni et al 2008). Prosite searches were carried through with prosite.expasy.org (Sigrist et al 2002). Panther search with http://www.pantherdb.org was used for visualizing multiple sequence alignments and phylogenetic tree results (Thomas et al 2003).
Deep View Ver.4.1 (http://www.expasy.org/spdbv) and Chimera (http://www.cgl.ucsf.edu/chimera) were used for further structural analyses. Energy minimization of structures was carried through with a minimum of 100 steps of conjugate gradients applying the amber ff12sp force field (Wang et al 2006; Case et al 2012) using Gasteiger charges (Gasteiger and Marsili 1980). "Holo enzymes" were thus constructed from the apo-liganded structures by converting side chains rotamers within the binding site as well is within the entire structure for advanced tracking of docked ligands. Ligands were analyzed with ViewDock. Hydrogen-bonding files were further processed in Excel (Microsoft) and sqlite data were analyzed using SQLite (Hipp, D. R.).

Further graphics were calculated or presented with SPSS Statistics (IBM) 21 and Excel (Microsoft). T-tests were carried through with post-hoc testing including skewness, Kolmogorov-Smirnov and Levene tests.
List of abbreviations

CNX  calnexin
CRD  carbohydrate recognition domain
CRT  calreticulin
EGF  epidermal growth factor
ER  endoplasmic reticulum
ERAD  endoplasmic reticulum associated degradation
ERGIC  endoplasmic reticulum-Golgi intermediate compartment
Evi  evenness interrupted
Fuc  fucose
Fut  fucosyltransferase
G  ganglioside (with class indicator)
Gal  galactose
GalCer  galactosylceramide
GalNAc  N-acetylgalactosamine
Gb  globoside
Glc  glucose
GlcCer  glucosylceramide
GlcNAc  N-acetylgalcosamine
GlcNAcT-V  GlcNAc-Transferase V (Mgat5)
Golgi-Man  Golgi-mannosidase
GPI-anchor  glycosylphosphatidylinositol-anchor
GSL  glycosphingolipid
Hep3B  hepatocellular carcinoma cells
Lac  lactose
LacCer  lactosylceramide
Lc  lacto- (with numerator for the number of carbohydrates)
LDL  low-density lipoprotein
LEF  lymphoid enhancer binding factor
Man  mannose
MCFD2  multiple coagulation factor deficiency protein 2
MDCK  Madin-Darby Canine Kidney
NeuAc  N-acetylenuraminic acid
NSCLC  nonsmall cell lung cancer
TCF  T-cell transcription factor
| Abbreviation | Description |
|--------------|-------------|
| TGF          | transforming growth factor |
| Ugcg         | UDP-glucose ceramide glucosyltransferase |
| Ugt          | UDP-galactose ceramide galactosyltransferase |
| VIP36        | vesicular-integral membrane protein of 36 kD |
| Wnt          | wingless |
Competing interests

The author declares no competing financial or non-financial interests.

Authors' contributions

KF conceived and carried out the study, drafted the manuscript and prepared tables, figures and graphics. The author read and approved the final manuscript.

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References

Arshad N, Ballal S, Visweswariah SS (2013) Site-specific N-linked glycosylation of receptor guanylyl cyclase C regulates ligand binding, ligand-mediated activation and interaction with vesicular integral membrane protein 36, VIP36. J Biol Chem 288:3907–3917. doi: 10.1074/jbc.M112.413906

Biellmann F, Huelsmeier AJ, Zhou D, et al (2008) The Lc3-synthase gene B3gnt5 is essential to pre-implantation development of the murine embryo. BMC Developmental Biology. doi: 10.1186/1471-213X-8-109

Boscher C, Nabi IR (2013) Galectin-3-and phospho-caveolin-1-dependent outside-in integrin signaling mediates the EGF motogenic response in mammary cancer cells. Mol Biol Cell 24:2134–2145. doi: 10.1091/mbc.E13-02-0095

Buechling T, Chaudhary V, Spirohn K, et al (2011) p24 proteins are required for secretion of Wnt ligands. EMBO reports 12:1265–1272. doi: 10.1038/embor.2011.212

Case DA, Darden TA, T.E. Cheatham I, et al (2012) AMBER 12. University of California, San Francisco.

Ceroni A, Maass K, Geyer H, et al (2008) GlycoWorkbench: A tool for the computer-assisted annotation of mass spectra of glycans. Journal of proteome research 7:1650–1659.

Chen C-Y, Jan Y-H, Juan Y-H, et al (2013) Fucosyltransferase 8 as a functional regulator of nonsmall cell lung cancer. Proceedings of the National Academy of Sciences 110:630–635. doi: 10.1073/pnas.1220425110

Christiansen MN, Chik J, Lee L, et al (2014) Cell surface protein glycosylation in cancer. Proteomics 14:525–546. doi: 10.1002/pmic.201300387

Chui D, Oh-Eda M, Liao Y-F, et al (1997) Alpha-mannosidase II deficiency results in dyserythropoiesis and unveils an alternate pathway in oligosaccharaide biosynthesis. Cell 90:157–167.

Contreras F-X, Ernst AM, Haberkant P, et al (2012) Molecular recognition of a single sphingolipid species by a protein’s transmembrane domain. Nature 481:525–529. doi: 10.1038/nature10742

D’Angelo G, Uemura T, Chuang C-C, et al (2013) Vesicular and non-vesicular transport feed distinct glycosylation pathways in the Golgi. Nature 501:116–120.

Delacour D, Cramm-Behrens CI, Drobecq H, et al (2006) Requirement for Galectin-3 in Apical Protein Sorting. Current Biology 16:408–414. doi: http://dx.doi.org/10.1016/j.cub.2005.12.046

Ellgaard L, Molinari M, Helenius A (1999) Setting the standards: Quality control in the secretory pathway. Science 286:1882–1888.

Etzler ME, Surolia A, Cummings RD (2009) L-type lectins. Essentials of Glycobiology

Fiedler K, Parton RG, Kellner R, et al (1994) VIP36, a novel component of glycolipid rafts and exocytic carrier vesicles in epithelial cells. EMBO J 13:1729–1740.
Fiedler K, Rothman JE (1997) Sorting determinants in the transmembrane domain of p24 proteins. J Biol Chem 272:24739–24742.

Fiedler K, Simons K (1996) Characterization of VIP36, an animal lectin homologous to leguminous lectins. Journal of Cell Science 109:271–276.

Fiedler K, Simons K (1994) A putative novel class of animal lectins in the secretory pathway homologous to leguminous lectins. Cell 77:625–626.

Fiedler K, Veit M, Stamnes MA, Rothman JE (1996) Bimodal interaction of coatomer with the p24 family of putative cargo receptors. Science 273:1396–1399.

Forli S, Olson AJ (2012) A force field with discrete displaceable waters and desolvations entropy for hydrated ligand docking. J Med Chem 55:623–638.

Fullekrug J, Scheiffele P, Simons K (1999) VIP36 localisation to the early secretory pathway. Journal of Cell Science 112:2813–2821.

Furukawa K, Tokuda N, Okuda T, et al (2004) Glycosphingolipids in engineered mice: insights into function. Seminars in Cell & Developmental Biology 15:389–396. doi: http://dx.doi.org/10.1016/j.semcdb.2004.03.006

Gasteiger J, Marsili M (1980) Iterative partial equalization of orbital electronegativity - a rapid access to atomic charges. Tetrahedron 36:3219–3228.

Goto Y, Obata T, Kunisawa J, et al (2014) Innate lymphoid cells regulate intestinal epithelial cell glycosylation. Science. doi: 10.1126/science.1254009

Granovsky M, Fata J, Pawling J, et al (2000) Suppression of tumor growth and metastasis in Mgat5-deficient mice. Nature Medicine 6:306–312.

Gut A, Kappeler F, Hyka N, et al (1998) Carbohydrate-mediated Golgi to cell surface transport and apical targeting of membrane proteins. EMBO J 17:1919–1929.

Hakomori S (2008) Structure and function of glycosphingolipids and sphingolipids: Recollections and future trends. Biochim Biophys Acta 1780:325–346.

Hall MK, Weidner DA, Chen JM, et al (2013) Glycan Structures Contain Information for the Spatial Arrangement of Glycoproteins in the Plasma Membrane. PLOS ONE. doi: 10.1371/journal.pone.0075013

Haltiwanger RS, Lowe JB (2004) Role of glycosylation in development. ANNUAL REVIEW OF BIOCHEMISTRY 73:491–537. doi: 10.1146/annurev.biochem.73.011303.074043

Hara-Kuge S, Ohkura T, Ideo H, et al (2002) Involvement of VIP36 in intracellular transport and secretion of glycoproteins in polarized Madin-Darby canine kidney (MDCK) cells. J Biol Chem 277:16332–16339.

Hara-Kuge S, Seko A, Yamashita K (2003) Carbohydrate recognition of vesicular integral protein of 36 kDa (VIP36) in intracellular transport of newly synthesized glycoproteins. Recognition of Carbohydrates in Biological Systems, Part B: Specific Applications. pp 525–532
Hebert DN, Foellmer B, Helenius A (1995) Glucose trimming and reglucosylation determine glycoprotein association with calnexin in the endoplasmic reticulum. Cell 81:425–433.

Hebert DN, Foellmer B, Helenius A (1996) Calnexin and calreticulin promote folding, delay oligomerization and suppress degradation of influenza hemagglutinin in microsomes. EMBO Journal 15:2961–2968.

Hebert DN, Lamriben L, Powers ET, Kelly JW (2014) The intrinsic and extrinsic effects of N-linked glycans on glycoproteostasis. Nat Chem Biol 10:902–910.

Helenius A, Aebi M (2001) Intracellular functions of N-linked glycans. Science 291:2364–2369.

Hirabayashi J, Hashidate T, Arata Y, et al (2002) Oligosaccharide specificity of galectins: a search by frontal affinity chromatography. Biochimica et Biophysica Acta (BBA) - General Subjects 1572:232–254. doi: http://dx.doi.org/10.1016/S0304-4165(02)00311-2

Holthuis JC, Pomorski T, Raggers RJ, et al (2001) The organizing potential of sphingolipids in intracellular membrane transport. Physiol Rev 81:1689–1723.

Honke K, Hirahara Y, Dupree J, et al (2002) Paranodal junction formation and spermatogenesis require sulfoglycolipids. Proceedings of the National Academy of Sciences of the United States of America 99:4227–4232. doi: 10.1073/pnas.032068299

Huang Y, Tan J-M, Wang Z, et al (2014) Cloning and characterization of two different L-type lectin genes from the Chinese mitten crab Eriocheir sinensis. Developmental & Comparative Immunology 46:255–266. doi: http://dx.doi.org/10.1016/j.dci.2014.04.015

Ioffe E, Liu Y, Stanley P (1996) Essential role for complex N-glycans in forming an organized layer of bronchial epithelium. Proceedings of the National Academy of Sciences USA 93:11041–11046. doi: 10.1073/pnas.93.20.11041

Ioffe E, Stanley P (1994) Mice lacking N-acetylglucosaminyltransferase I activity die at mid-gestation, revealing an essential role for complex of hybrid N-linked carbohydrates. Proc Natl Acad Sci 91:728–732.

Itin C, Schindler R, Hauri HP (1995) Targeting of protein ERGIC-53 to the ER/ERGIC/cis-Golgi recycling pathway. J Cell Biol 131:57–67.

Jennemann R, Groene H-J (2013) Cell-specific in vivo functions of glycosphingolipids: Lessons from genetic deletions of enzymes involved in glycosphingolipid synthesis. Progress in Lipid Research 52:231–248. doi: 10.1016/j.plipres.2013.02.001

Jo S, Lee HS, Skolnick J, Im W (2013) Restricted N-glycan conformational space in the PDB and its implication in glycan structure modeling. PLOS Computational Biology. doi: 10.1371/journal.pcbi.1002946

Kamiya Y, Kamiya D, Yamamoto K, et al (2008) Molecular basis of sugar recognition by the human L-type lectins ERGIC-53, VIPL, and VIP36. Journal of Biological Chemistry 283:1857–1861. doi: 10.1074/jbc.M709384200
Kamiya Y, Yamaguchi Y, Takahashi N, et al (2005) Sugar-binding properties of VIP36, an intracellular animal lectin operating as a cargo receptor. Journal of Biological Chemistry 280:37178–37182. doi: 10.1074/jbc.M505757200

Kitagawa Y, Sano Y, Ueda M, et al (1994) N-glycosylation of erythropoietin is critical for apical secretion by Madin-Darby canine kidney cells. Exp Cell Res, 213:449–457.

Krissinel E, Henrick K (2004) Secondary-structure matching (SSM), a new tool for fast protein structure alignment in three dimensions. Acta Crystallographica Section D 60:2256–2268. doi: 10.1107/S0907444904026460

Kuan C-T, Chang J, Mansson J-E, et al (2010) Multiple phenotypic changes in mice after knockout of the B3gnt5 gene, encoding Lc3 synthase—a key enzyme in lacto-neolacto ganglioside synthesis. BMC Developmental Biology. doi: 10.1186/1471-213X-10-114

Kurimoto A, Kitazume S, Kizuka Y, et al (2014) The absence of core fucose up-regulates GnT-III and Wnt target genes: a possible mechanism for an adaptive response of glycan function. Journal of Biological Chemistry 289:11704–11714. doi: 10.1074/jbc.M113.502542

Lee SH, Takahashi M, Honke K, et al (2006) Loss of core fucosylation of low-density lipoprotein receptor–related protein-1 impairs its function, leading to the upregulation of serum levels of insulin-like growth factor–binding protein 3 in Fut8−/− mice. Journal of Biochemistry 139:391–398. doi: 10.1093/jb/mvj039

Lotz MM, Andrews CW, Korzelius CA, et al (1993) Decreased expression of Mac-2 (carbohydrate binding protein 35) and loss of its nuclear localization are associated with the neoplastic progression of colon carcinoma. Proceedings of the National Academy of Sciences 90:3466–3470. doi: 10.1073/pnas.90.8.3466

Mellman I, Nelson WJ (2008) Coordinated protein sorting, targeting and distribution in polarized cells. Nat Rev Mol Cell Biol 9:833–845.

Mills J, Dean P (1996) Three-dimensional hydrogen-bond geometry and probability information from a crystal survey. J Comput Aided Mol Des 10:607–622.

Mishra R, Grzybek M, Niki T, et al (2010) Galectin-9 trafficking regulates apical-basal polarity in Madin–Darby canine kidney epithelial cells. Proceedings of the National Academy of Sciences 107:17633–17638. doi: 10.1073/pnas.1012424107

Miyoshi E, Noda K, Ko JH, et al (1999) Overexpression of α1-6 Fucosyltransferase in Hepatoma Cells Suppresses Intrahepatic Metastasis after Splenic Injection in Athymic Mice. Cancer Research 59:2237–2243.

Moremen KW (2002) Golgi alpha-mannosidase II deficiency in vertebrate systems: implications for asparagine-linked oligosaccharide processing in mammals. Biochimica et biophysica acta 1573:225–235.

Nagae M, Nishi N, Murata T, et al (2009) Structural analysis of the recognition mechanism of poly-N-acetyllactosamine by the human galectin-9 N-terminal carbohydrate recognition domain. Glycobiology 19:112–117. doi: 10.1093/glycob/cwn121
Nagae M, Nishi N, Nakamura-Tsuruta S, et al (2008) Structural Analysis of the Human Galectin-9 N-terminal Carbohydrate Recognition Domain Reveals Unexpected Properties that Differ from the Mouse Orthologue. Journal of Molecular Biology 375:119–135. doi: http://dx.doi.org/10.1016/j.jmb.2007.09.060

Nakagawa T, Uozumi N, Nakano M, et al (2006) Fucosylation of N-glycans regulates the secretion of hepatic glycoproteins into bile ducts. Journal of Biological Chemistry 281:29797–29806. doi: 10.1074/jbc.M605697200

Neve EPA, Svensson K, Fuxe J, Pettersson RF (2003) VIP36-like membrane protein with a putative function in the export of glycoproteins from the endoplasmic reticulum. Experimental Cell Research 288:70–83. doi: 10.1016/S0014-4827(03)00161-7

Nishio M, Kamiya Y, Mizushima T, et al (2010) Structural basis for the cooperative interplay between the two causative gene products of combined factor V and factor VIII deficiency. Proc Natl Acad Sci U S A 107:4034–4039. doi: 10.1073/pnas.0908526107

O'Boyle NM, Banck M, James CA, et al (2011) Open Babel: An open chemical toolbox. J Cheminf 3:33.

Partridge EA, Le Roy C, Di Guglielmo GM, et al (2004) Regulation of cytokine receptors by Golgi N-glycan processing and endocytosis. Science 306:120–124. doi: 10.1126/science.1102109

Pickard JM, Maurice CF, Kinnebrew MA, et al (2014) Rapid fucosylation of intestinal epithelium sustains host-commensal symbiosis in sickness. Nature 514:638–641.

Proia RL (2003) Glycosphingolipid functions: insights from engineered mouse models. Philosophical Transactions of the Royal Society of London Series B: Biological Sciences 358 :879–883. doi: 10.1098/rstb.2003.1268

Ranzer R, Frank M, Von der Lieth CW, Herget S (2009) Glycome-DB.org: a portal for querying across the digital world of carbohydrate sequences. Glycobiology 19:1563–1567.

Rodriguez-Boulan E, Macara IG (2014) Organization and execution of the epithelial polarity programme. Nature Reviews Molecular Cell Biology 15:225–242. doi: 10.1038/nrm3775

Roth J (2002) Protein N-Glycosylation along the Secretory Pathway: Relationship to Organelle Topography and Function, Protein Quality Control, and Cell Interactions. Chem Rev 102:285–303.

Sabourdy F, Kedjoura B, Sorli SC, Colié S (2008) Functions of sphingolipid metabolism in mammals — Lessons from genetic defects. Biochimica et Biophysica Acta 1781:145–183. doi: 10.1016/j.bbalip.2008.01.004

Sato M, Nishi N, Shoji H, et al (2002) Functional analysis of the carbohydrate recognition domains and a linker peptide of galectin-9 as to eosinophil chemoattractant activity. Glycobiology 12 :191–197. doi: 10.1093/glycob/12.3.191

Satoh T, Chen Y, Hu D, et al (2010) Structural Basis for Oligosaccharide Recognition of Misfolded Glycoproteins by OS-9 in ER-Associated Degradation. Molecular Cell 40:905–916. doi: 10.1016/j.molcel.2010.11.017
Satoh T, Cowieson NP, Hakamata W, et al (2007) Structural basis for recognition of high mannose type glycoproteins by mammalian transport lectin VIP36. The Journal of biological chemistry 282:28246–55. doi: 10.1074/jbc.M703064200

Satoh T, Suzuki K, Yamaguchi T, Kato K (2014) Structural Basis for Disparate Sugar-Binding Specificities in the Homologous Cargo Receptors ERGIC-53 and VIP36. PLOS ONE 9:e87963. doi: 10.1371/journal.pone.0087963

Scheiffele P, Peranen J, Simons K (1995) N-glycans as apical sorting signals in epithelial cells. Nature 378:96–98.

Shimada O, Hara-Kuge S, Yamashita K, et al (2003) Localization of VIP36 in the post-Golgi secretory pathway also of rat parotid acinar cells. J Histochem Cytochem 51:1057–1063.

Shimura T, Takenaka Y, Tsutsumi S, et al (2004) Galectin-3, a Novel Binding Partner of β-Catenin. Cancer Research 64:6363–6367. doi: 10.1158/0008-5472.CAN-04-1816

Shirakabe K, Hattori S, Seiki M, et al (2011) VIP36 protein is a target of ectodomain shedding and regulates phagocytosis in macrophage raw 264.7 cells. Journal of Biological Chemistry 286:43154–43163.

Sigrist CIA, Cerutti L, Hulo N, et al (2002) PROSITE: A documented database using patterns and profiles as motif descriptors. Briefings in Bioinformatics 3:265–274.

Simons K, Ikonen E (1997) Functional rafts in cell membranes. Nature 387:569–572.

Stanley P, Cummings RD (2009) Structures common to different glycans. Essentials of Glycobiology

Stanley P, Schachter H, Taniguchi N (2009) N-glycans. Essentials of Glycobiology

Stowell SR, Ju T, Cummings RD (2015) Protein Glycosylation in Cancer. Annual Review of Pathology: Mechanisms of Disease 10:473–510. doi: 10.1146/annurev-pathol-012414-040438

Svensson L, Hult AK, Stamps R, et al (2012) Forssman expression on human erythrocytes: biochemical and genetic evidence of a new histo-blood group system. Blood 121:1459–1468.

Takenaka Y, Fukumori T, Raz A (2002) Galectin-3 and metastasis. Glycoconjugate Journal 19:543–549. doi: 10.1023/B:GLYC.0000014084.01324.15

Theiler R, Fujita M, Naga M, et al (2014) The α-Helical Region in p24γ2 Subunit of p24 Protein Cargo Receptor Is Pivotal for the Recognition and Transport of Glycosylphosphatidylinositol-anchored Proteins. Journal of Biological Chemistry 289:16835–16843. doi: 10.1074/jbc.M114.568311

Thomas PD, Campbell MJ, Kejariwal A, et al (2003) PANTHER: A Library of Protein Families and Subfamilies Indexed by Function. Genome Research 13:2129–2141. doi: 10.1101/gr.772403

Trott O, Olson AJ (2010) AutoDock Vina: improving the speed and accuracy of docking with a new scoring function, efficient optimization and multithreading. Journal of Computational Chemistry 31:455–461.
Tschampel SM, Kirschner KN, Woods RJ (2006) Incorporation of carbohydrates into macromolecular force fields: Development and validation. Nmr Spectroscopy and Computer Modeling of Carbohydrates: Recent Advances (Amer Chemical Soc) 930:235–257.

Uhlén M, Fagerberg L, Hallström BM, et al (2015) Tissue-based map of the human proteome. Science. doi: 10.1126/science.1260419

van Ijzendoorn SCD, Hoekstra D (1999) Polarized sphingolipid transport from the subapical compartment: Evidence for distinct sphingolipid domains. Mol Biol Cell 10:3449–3461.

Velloso LM, Svensson K, Pettersson RF, Lindqvist Y (2003) The crystal structure of the carbohydrate-recognition domain of the glycoprotein sorting receptor p58/ERGIC-53 reveals an unpredicted metal-binding site and conformational changes associated with calcium ion binding. Journal of molecular biology 334:845–51.

Wang J, Wang W, Kollman PA, Case DA (2006) Automatic atom type and bond type perception in molecular mechanical calculations. Journal of molecular graphics & modelling 25:247–260. doi: 10.1016/j.jmgm.2005.12.005

Wang X, Inoue S, Gu J, et al (2005) Dysregulation of TGF-β1 receptor activation leads to abnormal lung development and emphysema-like phenotype in core fucose-deficient mice. Proc Natl Acad Sci USA 102:15791–15796. doi: 10.1073/pnas.0507375102

Wolf L (2009) digital briefs: New software and websites for the chemical enterprise. C&EN 87:48.

Yamamoto H, Awada C, Hanaki H, et al (2013) The apical and basolateral secretion of Wnt11 and Wnt3a in polarized epithelial cells is regulated by different mechanisms. JOURNAL OF CELL SCIENCE 126:2931–2943. doi: 10.1242/jcs.126052

Yamashita T, Wada R, Sasaki T, et al (1999) A vital role for glycosphingolipid synthesis during development and differentiation. Proc Natl Acad Sci USA 96:9142–9147.
**Figure legends**

**Figure 1**
The CARB 165 library was constructed for this experimental approach. CARB 165 was tested against 2E6V, the structure of VIP36 chosen for a virtual ligand screen. 2E6V is established as “Apo” structure in “Global” screening, the structural studies resolved mannose residues in crystallization. The CARB 165 library was subsequently screened in the Local approach with 2DUR-A, 2DUR-B and towards binding to 2E6V, all three molecularly relaxed. "Holo enzyme" behavior is inferred from side chain rotation. The number of database entries in the respective carbohydrate group is indicated.

**Figure 2**
Summary of newly discovered glycan ligands of VIP36. The Carb165 library was screened for interactions with 2DUR-A. Number of glycan motifs among top scores are indicated (no double counts of extended structures or multiple motifs)(15 top scoring glycans).

**Figure 3**
Carbohydrate interaction. Amino acids of VIP36 implicated in H-bonding to carbohydrate ligands are presented with surface-proximate groups, the legend indicates the IUPAC coloring of carbohydrates and their respective surface imprint at 3 Å. Few residues are covered behind E168. The binding site is shown with determined H-bond donors in black and acceptors in green for Carb165 (15·9 Models).

**Figure 4**
H-bonding of glycan binding and topology of reducing end interaction with VIP36. A Comparison of H-bonding. The glycans Carb165 identified in the docking approach were used to establish the binding site geometry with top scoring ligands 9x15. Ligands were evaluated in H-bonding (0.4 Å, 20° relaxed) and the % of bonds formed is graphed. Amino acids are indicated. B Topology of the VIP binding site measured by virtual screen. Top 15 glycans of Carb165 were analyzed for binding and found geometries are indicated and labeled with 1-3. The base of the arrow indicates the reducing end of the carbohydrate, the arrowhead the direction of the antennae of N-glycans. Fig. 3 is shown in similar molecular orientation. For comparison also Fig. S2 can be used.
Figure 5
Graphical presentation of Carb165 interactors of VIP36. A Three dimensional plot of all conformers 2E6V, 2DUR-A and 2DUR-B of VIP36 in all-against-all manner (within sub-libraries). Carbohydrates are listed in groups of N-glycan hybrid (violet box), N-glycan high mannose (green box), N-glycan complex (light violet box), Sialoside (light violet diamond), Fucoside (orange triangle) and O-glycan core (yellow square). Other glycans that were not included in this nomenclature are indicated (n=44)(green thin ring). 2DUR-A was graphed against 2DUR-B and 2E6V (each holo). Sialosides include carbohydrates of O-glycan feature if these are sialylated, O-glycans include glycans that are fucosylated. The graph uses scatter plot axes definitions. The runs consisted of 3 sessions with 159 carbohydrates each. The group centers are indicated in large symbols.

Figure 6
Model of transport and sorting: Overview of N-glycan structure, VIP36 lectin affinity is shown here for the first time for the high mannose and complex type N-glycans. Some enzymes involved in N-glycan processing are indicated. MAN1B1 has recently been identified in the Golgi complex (see (Hebert et al 2014)). ERGIC-53 (green), VIPL (VIP36-like protein), VIP36 (red) and Galectins are indicated and are localized to the ER, ERGIC (yellow; co-localization), Golgi apparatus, RE or PM. The residue labeled Manα2 has not been included in a previous model of ERGIC-53 interaction (Satoh et al 2014) and is indicated for clarity since the M9 (Man9) carbohydrate residue is the described substrate of (1) and (2). Serial lectin affinity chromatography points to monoglucosylated M7 as the high affinity interactor of ERGIC-53. VIPL is described to interact with M9 and M8 high mannose residues with high affinity. Galectin-9 (Gal-9) is a high affinity interactor of polylactosamines. Forssman antigen glycosphingolipid (GSL) was found to interact as well, the exact GSL that binds to galectin-9 in polarized sorting in humans remains to be determined (Svensson et al 2012). Galectin-3 (Gal-3) was found to be involved in a non-raft pathway in apical sorting. VIP36 is modeled to bind to docked core residues of N-glycans involving a fucose residue (1-6 linked), mannoses and N-acetylg glucosamines. Preferential interaction of VIP36 with the mannose (1-3) branch makes additional clustering interactions likely for the mannose (1-6) branch. Apical sorting signals would involve both branches of N-glycans and GSLs. No preferential binding site of core fucose 1-6 of all complex N-glycans (Fig. S3C; 38, 39, 41) with VIP36 could as yet be demonstrated. Calnexin (CNX) or Calreticulin (CRT) and lectin ERAD (ER-associated protein degradation) receptors OS-9 or XTP3-B are also indicated. OS-9 binds to M8 - M5 high-mannose N-glycans, XTP3-B binding may be similar (Hebert et al 2014). Both proteins may also directly interact with the membrane protein SEL1L of the ERAD complex via its own glycans. Further information on ER quality control of protein folding and degradation is found in the cited
review. The p24 proteins are introduced to mention their interaction with the glycosylphosphatidylinositol (GPI)-anchor of GPI-anchored proteins (p24\(\gamma\)) likely, indirectly or directly, through their membrane proximal \(\alpha\)-helical domain (Theiler et al 2014). Abbreviations: ER (endoplasmic reticulum), ERGIC (ER-Golgi intermediate compartment), PM (plasma membrane), RE (recycling endosome).

**Figure 7**
Structural comparison of L-type lectin, Galectin and P-type lectin. VIP36 (2DUR-A; residues 51-301), ERGIC-53 (4GKX-A; residues 31-270) and Galectin-9 (3LSE-A; residues 6-148) were analyzed and aligned in three dimensions with Chimera. P-type lectin OS-9 (3AIH-A; residues 109-228) is shown in similar aspect for comparison. Carbohydrates are indicated and shown in the binding pocket (orange) if present. Calcium is shown in green, top-scoring entries of the similarity search (Table 1) were aligned. Galectin-9 binds carbohydrates in similar position to e.g. galectins 3 and 4. The P-type lectin OS-9 and OS-9-like proteins recognize carbohydrates primarily via strand residues, OS-9 binding involves a Trp-Trp motif. Further P-type lectins/mannose 6-phosphate receptor (MPR) homology domain-containing lectins (including the cation-independent and cation-dependent MPR) have been omitted. The glycan binding site of VIP36 and galectin-9 is non-similar, whereas primarily loop residues Ser96, Asp131, Asn166, Gly260, Asp261, Leu262 bind to the carbohydrate in VIP36 it is strand residues in galectin-9 that interact including His61, Asn63, Arg65, Asn75 and Arg87 that do not align. Resolved carbohydrates in the structures corresponded to \(\alpha\)-D-man-\(\alpha\)-D-man (VIP36), \(\alpha\)-D-man-\(\alpha\)-D-man (ERGIC-53), \(\beta\)-lactose (Galectin-9) and \(\beta\)-D-man-\(\alpha\)-D-man-\(\alpha\)-D-man (OS-9). ERGIC-53 has been crystallized in the presence of the recruitment protein MCFD2 (Multiple Coagulation factor Deficiency protein 2)(Nishio et al 2010). The binding face is indicated. MCFD2 binds coagulation factors V and VIII and in an autosomal recessive disorder in mutants causes reduced serum levels. Independent ERGIC-53 binding of glycoprotein ligands without MCFD2 has also been shown.
Tables

Table I
Structural comparison of leguminous-type lectins. VIP36 (2DUR) was used in the database search with PDB. Structural hits were ranked according to PDBEfold Q-Score. RMSDs, each Structure found and the Protein name is indicated. All retrieved structures were derived from eukaryotic proteins unless otherwise indicated (Species), some proteins were crystallized in mutant form and top-scoring protein isoforms of each entry were given. The results were collated from 2860 structure files; the lowest match was adjusted to 60%. Entries that had previously been ranked and determined to be structurally similar to L-type lectins (1JHN, 3FBY, 1IKQ) were added for non-inclusive ranking. Proteins shown in bold are discussed in the text.
Library of Carbohydrates

N-glycan high mannose 19
N-glycan hybrid 17
N-glycan complex 9
Sialoside 27
Fucoside 25
O-glycan core 17
Other glycan 51
Figure 2

Top scoring library glycans

- Lactosamine motif
- O-glycan
- N-glycan complex
- N-glycan high mannose
- N-glycan hybrid
- GSL
top scores:
topologies
Figure 5

- N-glycan high mannose
- N-glycan hybrid
- N-glycan complex
- Sialoside
- Fucoside
- O-glycan core
- Other glycan
Figure 6

**Trafficking: N-Glycans in the trans-Golgi**

1: ER Mannosidase I (MAN1B1); 2: ER Glucosidase II; 3: GlcNAcT-V; 4: GlcNAcT-II; 5: GlcNAcT-IV; 6: GlcNAcT-I; 7: GlcNAcT-III; 8: FucT-VIII

**Recognition of N-Glycans in the ER**

Manα2 → Asn

1. CNX, CRT, ERGIC-53
2. Asn

**OS-9 / XTP3-B** → **SEL1L**
| Protein               | Structure | Q-Score | RMSD  | Species                          |
|-----------------------|-----------|---------|-------|----------------------------------|
| VIP36                 | 2dur      | 1       | 0.00  | Canis lupus familiaris           |
| ERGIC-53              | 4gkx      | 0.7     | 1.08  | Homo sapiens                     |
| EMP47                 | 2a71      | 0.52    | 1.80  | Saccharomyces cerevisiae         |
| EMP46                 | 2a6w      | 0.48    | 1.95  | Saccharomyces cerevisiae         |
| Pea Lectin            | 2bqp      | 0.43    | 2.01  | Pisum sativum                    |
| Soybean Agglutinin    | 2sba      | 0.41    | 1.88  | Glycine max                      |
| Basic Agglutinin      | 2zml      | 0.41    | 1.97  | Psophocarpus tetragonolobus      |
| Seed Lectin           | 4u2a      | 0.40    | 2.07  | Vatairea macrocarpa              |
| Isolectin B4          | 1n47      | 0.40    | 1.94  | Vicia villosa                    |
| Lectin DB58           | 1ul       | 0.40    | 1.92  | Vigna unguiculata                |
| Lectin                | 2eig      | 0.40    | 2.21  | Lotus tetragonolobus             |
| ...                   | 1oq1      | 0.33    | 2.39  | Prokaryotic, Bacillus subtilis    |
| Galectin-9            | 3lse      | 0.31    | 2.41  | Homo sapiens                     |
| Galectin-7            | 5gal      | 0.31    | 2.07  | Homo sapiens                     |
| Galectin-8            | 3ap4      | 0.30    | 2.31  | Homo sapiens                     |
| Galectin-3            | 3sli      | 0.30    | 2.26  | Homo sapiens                     |
| Galectin-4            | 3lt8      | 0.29    | 2.44  | Mus musculus                     |
| Galectin-1*           | 4no4      | 0.29    | 2.51  | Rattus norvegicus                |
| Galectin LEC-6        | 3vv1      | 0.29    | 2.24  | Caenorhabditis elegans           |
| Galectin-1            | 1sla      | 0.29    | 2.53  | Bos taurus                       |
| Galectin-2            | 2ymz      | 0.29    | 2.52  | Gallus gallus                    |
| Galectin-10 Charcot-Leyden | 1g86 | 0.28    | 2.49  | Homo sapiens                     |
| ...                   | 3i4i      | 0.27    | 2.48  | Prokaryotic, uncultured murine large bowel |
| Secreted putative glycosylhydrolase | 3imm | 0.27 | 2.16 | Prokaryotic, Parabacteroides distasonis |
| Calreticulin          | 3o0w      | 0.26    | 2.63  | Mus musculus                     |
| SPRY domain-containing protein | 2yyo | 0.26 | 2.98 | Homo sapiens                     |
| Sialidase             | 2w68      | 0.24    | 2.33  | Prokaryotic, Vibrio cholerae     |
| Protein neutralized   | 4kg0      | 0.23    | 3.04  | Drosophila melanogaster          |
| ...                   | 3i4i      | 0.27    | 2.48  | Prokaryotic, uncultured murine large bowel |
| VAL-1                 | 3im0      | 0.23    | 2.91  | Virus, Chlorella virus           |
| Ephrin type-A receptor| 4et7      | 0.20    | 2.81  | Homo sapiens                     |
| Sex hormone binding globulin | 1d2s | 0.20 | 2.87 | Homo sapiens                     |
| Ephrin type-B receptor| 3pli      | 0.19    | 2.69  | Homo sapiens                     |
| Thrombospondin 1      | 1z78      | 0.19    | 3.44  | Homo sapiens                     |
| beta-Neurexin         | 3bop      | 0.19    | 2.72  | Mus musculus                     |
| Agrin                 | 1pz8      | 0.19    | 2.52  | Gallus gallus                    |
| Collagen NC4 domain (CO9A1) | 2uur | 0.19 | 3.02 | Homo sapiens                     |
| alpha-Neurexin        | 2h0b      | 0.18    | 2.48  | Bos taurus                       |
| Perlecan LG3          | 3sh5      | 0.18    | 2.73  | Homo sapiens                     |
| C-reactive protein    | 1lj7      | 0.18    | 3.38  | Homo sapiens                     |
| Serum amyloid P       | 2a3x      | 0.18    | 3.43  | Homo sapiens                     |
| XRCC1                 | 3k75      | 0.17    | 3.37  | Homo sapiens                     |
| Tripartite motif-containing | 3uv9 | 0.16 | 3.06 | Macaca mulatta                  |
| ...                   | 1jhn      | 0.15    | 2.99  | Canis lupus familiaris           |
| Calnexin              | 1ikq      | 0.13    | 3.98  | Virus, Rhesus rotavirus          |
| VP4                   | 1qam      | 0.12    | 1.91  | Canavalia ensiformis             |
| Concanavalin A        | 3by       | 0.11    | 2.62  | Homo sapiens                     |
| Laminin alpha LG4-LG5*| 1okq     | 0.10    | 2.55  | Mus musculus                     |
| Exotoxin A            | 1ikq      | 0.06    | 3.41  | Prokaryotic, Pseudomonas aeruginosa |

*mutant
Supplementary Figures/Tables

Figure S1
Top Carbohydrates interacting with VIP36. The structurally identified mannosyl ligand of 2DUR-A was removed, the structure was energy minimized and virtually screened. CARB residues interacting with highest binding site affinity are indicated and presented in IUPAC nomenclature. All ligands cover the central binding pocket that provides binding indentations that are seen (geometrically) between Tyr164, Asp261 and Pro 146, Glu 98 covering the central area Asp131, Asn166 (see Fig. 3). Energies indicated above are listed with values for molecules interacting with both, Asp131 and Asn166 via (0.4 Å, 20° relaxed) H-bonding. Model ranking is given for these doubly (or multiply) H-bonded ligands selected. Gobal score refers to ranking which combines CARB and GSL residues neglecting above mentioned structural requirement of forming 2 H-bonds to Asp131, Asn166. Sequential labeling in roman numbers.

Figure S2
Structure and docking of Carb165-library VIP36 ligands. The ligands identified with highest affinity to VIP36 (2DUR-A) are listed with IUPAC coloring (Structure) and imprint onto the VIP36. Heteroatom coloring was utilized to show potential H-bonding interactions. The topography (Topo) of the ligand binding reveals orientations of headgroup and cores that corresponded to three axes numbered 1-3 (see Fig. 4B). IUPAC formulas (IUPAC) and sequential labeling in roman numbers are indicated.

Figure S3
Glycan preference. The carbohydrate interaction of VIP36 (2DUR-A) was classified according to the energy (kcal/mol) and top scoring of each glycan group; Δ values are indicated. A High-mannose glycan preference. B Hybrid glycan preference. Standard enzyme product and substrates are indicated. C Glycan complex preference. D Glycan terminal end preferences. Some enzymes with (reference to the text) that are involved in the reaction paths are indicated with edduct (orange), product (blue) and intermediate (green). Sequential labeling is indicated in arabic numbers.
Table S1
Information on glycans and expression. Species shown (Species) were collected from Glycosciences.de websources using MCS. Glycan top-hits were scored in virtual binding to VIP36 conformer 2DUR-A. The top 15 are presented similarly in Figs. S1 and S3. Sequential labeling is in roman numbers.

Table S2
Conformers of 2E6V, 2DUR-A and 2DUR-B were compared for carbohydrate binding in virtual screening with the Carb165 library. A T-test with ligands of 2E6V, 2DUR-A and 2DUR-B (n=159-3). The glycans grouped for comparisons were subjected to one-sided T-test. Statistical significances of group differences are plotted. Post-hoc tests show that the values are normally distributed at p>0.08 levels (Kolmogorov-Smirnov and Lilliefors). A Levene statistics of 3.98 indicates the possibility of inflation of error (N=477). For comparison, p values are marked with * if at p<0.05 level with tests with unequal variances; these only showed differences with Fucosides. B The average of binding energies to the 2E6V, 2DUR-A and 2DUR-B combined is listed. The confidence interval (CI) is indicated.

Table S3
Clustal (Higgins et al 1996) was used to align the Prosite entries (Sigrist et al 2002) that were retrieved with the Canis familiaris VIP36 protein. Regions previously described in comparison to leguminous plant lectin Lec_Baupu are labeled I-III.

References
Higgins D, Thompson J, Gibson T (1996) Using CLUSTAL for multiple sequence alignments. Methods Enzymol 266:383–402.

Sigrist CJA, Cerutti L, Hulo N, et al (2002) PROSITE: A documented database using patterns and profiles as motif descriptors. Briefings in Bioinformatics 3:265–274.
Figure S1

Energy kcal/mol

Model No.

|   |   | -7.4 | -6.7 |   | -7.1 | -6.4 | -6.6 | -7.2 | -6.7 | -7.1 |
|---|---|-----|-----|---|------|------|------|------|------|------|
| I | II |     |     | III|     |      |      |      |      |      |
| IV| V  | VI  | VII | VIII| IX  | X    | XI   | XII  | XIII | XIV  | XV   |

Global Score
### Carbohydrates interacting with VIP36: Top scores

| Name and Comment                  | Structure | Topo | No./IUPAC                                                                 |
|-----------------------------------|-----------|------|---------------------------------------------------------------------------|
| **I**                             |           |      | A-D-Manp-(1-2)-A-D-Manp-(1-6)-[A-D-Manp-(1-2)-A-D-Manp-(1-3)]-A-D-Manp-(1-6)-[A-D-Manp-(1-3)]-B-D-Manp-(1-4)-B-D-GlcNAc-(1-4)-B-D-GlcNAc-OH |
| N-glycan high mannose             |           | 1    |                                                                           |
| Location: medial Golgi            |           |      |                                                                           |
| **II**                            |           | 2    | B-D-GlcNAc-(1-2)-A-D-Manp-(1-6)-[B-D-GlcNAc-(1-4)]-A-D-Manp-(1-3)]-B-D-Manp-(1-4)-B-D-GlcNAc-(1-4)-[A-D-Fuc-(1-6)]-B-D-GlcNAc-OH |
| N-glycan core basic               |           |      |                                                                           |
| Location: medial/trans Golgi      |           |      |                                                                           |
| **III**                           |           | 3    | A-D-Manp-(1-6)-[A-D-Manp-(1-2)]-A-D-Manp-(1-3)]-A-D-Manp-(1-6)-[A-D-Manp-(1-2)]-A-D-Manp-(1-3)]-B-D-Manp-(1-4)-B-D-GlcNAc-(1-4)-B-D-GlcNAc-OH |
| N-glycan high mannose             |           |      |                                                                           |
| Location: medial Golgi            |           |      |                                                                           |
| **IV**                            |           | 2    | A-D-Neu5Ac-(2-3)-[B-D-Galp-(1-3)]-B-D-GalpNAc-(1-4)]-B-D-Galp-(1-4)-B-D-GlcP-OH |
| Glycosphingolipid-ganglio series  |           |      |                                                                           |
| GM1, location: Medial/trans-Golgi |           |      |                                                                           |
Lactosamine motif

intermediate LacNAc to PolyLacNAc

N-glycan core basic
precursor of paucimannose-N-glycans* or complex N-glycans; Golgi hexosaminidase removes GlcNAc / GlcNAcT-I required for viability

Lactosamine motif, Polylactosamine, Glycosphingolipid Neo-lacto series
LacNAc in N-glycans; type 2

O-glycan core 2, Galα1-3Gal epitope, Lactosamine motif
frequent in many tissues
Lactosamine motif, LacDiNAc

LacDiNAc in N-glycans, frequently sialylated
Location: trans-Golgi

Lewis C, Lactosamine motif, Neo Lactosamine, Glyco-sphingolipid Lacto series
LacNAc variation also in N-glycans; terminal type 1

N-glycan core basic
Location: medial Golgi

N-glycan core basic
Location: medial/trans-Golgi

O-glycan core 2, Lactosamine motif
O-glycan core 6, Lactosamine motif

less frequent

*X Frequent in insects"
Figure S3A: Glycan high-mannose preferences

1. Golgi-Man I
2. Golgi-Man I
3. Golgi-Man I
4. Golgi-Man I
5. Golgi-Man I
6. Endomannosidase
7. Golgi-Man I
8. GlcNAcT-I, Lys Man
9. α-mannosidase III
10. Golgi-Man I
11. α-mannosidase III
12. Golgi-Man I
13. ER Glucosidase II
14. ER Glucosidase I
15. ER Man II
16. ER Glucosidase II
17. ER Glucosidase II
18. ER Man I
19. Endomannosidase
20. ER Man I
Figure S3B: Glycan hybrid preferences

| high | affinity | low |
|------|----------|-----|
| 21   | Golgi-Man II | 0.1 |
| 22   | Golgi-Man II | 0.1 |
| 23   | β1-4GalT | 0.2 |
| 24   | GlcNAcT-III | 0.6 |
| 25   | GlcNAcT-III | 0.7 |
| 26   | Golgi-Man II | 0.7 |
| 27   | GlcNAcT-I | 0.8 |
| 28   | β1-4GalT | 0.8 |
| 29   | β1-4GalT | 0.9 |
| 30   | β1-4GalT | 1.0 |
| 31   | ST6Gal-I | 1.0 |
| 32   | Golgi-Man II | 1.1 |
| 33   | ST6Gal-I | 1.3 |
| 34   | ST6Gal-I | 1.5 |
| 35   | ST6Gal-I | 1.6 |
| 36   | GlcNAcT-III | 1.8 |
| 37   | ST6Gal-I | 1.8 |
Figure S3C: Glycan complex preferences

| Glycan Complex | Affinity | High | Low |
|---------------|---------|------|-----|
| GlcNAcT-IV    | -0      | 38   | 41  |
|               | -0.3    | 39   | 42  |
| GlcNAcT-V     | -0.6    | 40   | 43  |
| FucosylT-VIII | -0.9    | 41   | 44  |
|               | -2.0    | 44   | 45  |
| β1-4GalT      | -1.1    | 42   | 45  |
|               | -2.3    | 45   | 45  |
| GlcNAcT-III   | -1.3    | 43   | 45  |
Figure S3D: Glycan terminal preferences

| high | affinity | low |
|------|----------|-----|
| 46   | α1-4FucT | -0  |
| 47   | β1-3GlcNAcT | -0.1 |
| 48   | β1-4GalT | -0.2 |
| 49   | ST6Gal-I | -0.7 |
| 50   | β1-3GalT | -0.2 |
| 51   | β1-3GalT | -0.5 |
| 52   | ST6Gal-I | -0.5 |
| 53   | β1-3GalT | -0.7 |
| 54   | α1-2FucT | -0.8 |
| 55   | ST3Gal-III | -0.8 |
| 56   | α1-2FucT | -1.3 |
| 57   | α1-3GalT | -1.5 |

46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57
Table S1

| Carbohydrates          | No. / Species |
|------------------------|---------------|
| N-glycan high mannose  | I             |
|                        | *Bos taurus*  |
|                        | *Coturnix japonica*  |
|                        | *Homo sapiens*  |
|                        | *Sus scrofa*    |
| N-glycan core basic    | II            |
|                        | *Homo sapiens*  |
|                        | *Sus scrofa*    |
| N-glycan high mannose  | III           |
|                        | *Bos taurus*  |
|                        | *Felis catus*  |
|                        | *Homo sapiens*  |
|                        | *Mus musculus*  |
|                        | *Ricinus communis*  |
|                        | *Sus scrofa*    |
| Glycosphingolipid-ganglio series | IV |
|                        | *Balaenoptera acutorostrata*, *Balaenoptera brydei*, *Balaenoptera edeni*, *Bos taurus*, *Campylobacter jejuni*, *Canis lupus familiaris*, *Carassius auratus*, *Cavia porcellus*, *Chinchilla*, *Chinchillidae*, *Chionodraco myersi*, *Cichlidae*, *Cricetinae*, *Cricetulus griseus*, *Cyprinus carpio*, *Delphinus delphis*, *Discopyge tschudii*, *Equus caballus*, *Felis catus*, *Gadus*, *Gallus gallus*, *Glóberephala macrorhynchus*, *Homo sapiens*, *Leishmania amazonensis*, *Macaca fuscata*, *Macaca mulatta*, *Megathura crenulata*, *Mus musculus*, *Orcinus orca*, *Oreochromis mossambicus*, *Oryctolagus cuniculus*, *Ovis aries*, *Papio hamadryas*, *Pelophylax esculentus*, *Phocoenoides dalli*, *Phodopus sungorus*, *Physeter catodon*, *Rattus*, *Scardinius erythrophthalmus*, *Soricidae*, *Stenella coeruleoalba*, *Suncus murinus*, *Sus scrofa*, *Torpedo marmorata*, *Tursiops truncatus*, *Xenopus laevis* |
| Lactosamine motif      | V             |
| Lactosamine motif      | VI            |
|                        | *Homo sapiens*  |
| N-glycan core basic    | VII           |
|                        | *Bos taurus*  |
| Lactosamine motif, Poly lactosamine, Glycosphingolipid Neo-lacto series | VIII |
|                        | IX            |
|                        | *Rattus norvegicus*  |
Lactosamine motif, LacDiNAc

Lewis C, Lactosamine motif, Neo Lactosamine, Glycosphingolipid Lacto series

N-glycan core basic

O-glycan core 1, Lactosamine motif

O-glycan core 6, Lactosamine motif

X
Bos taurus
Homo sapiens

XI
Homo sapiens

XII
Bos taurus
Cricetinae
Mus musculus
Nicotiana alata

XIII
Inferred based on extension:
Bos taurus, Cricetulus griseus, Fowl plague virus, Gallus gallus, Homo sapiens, Mus musculus, Rattus, Sendai virus, Sus scrofa, Torpedo californica, unidentified influenza virus

XIV

XV
Bos taurus
Homo sapiens
### A

**Statistical Probabilities**

| 2DUR-A 2DUR-B 2E6V | Sialoside | O-glycan core | Fucoside | N-glycan high mannose | N-glycan hybrid | N-glycan complex | Other glycan |
|---------------------|-----------|---------------|----------|-----------------------|----------------|-----------------|-------------|
| Sialoside           | 0.008*    | 0.328         | 0.334    | 0.111                 | 0.011*         | 0.001*          |             |
| O-glycan core       |           | 0.007*        | 0.015*   | 0.199                 | 0.002*         | 0.336           |             |
| Fucoside            |           |               | 0.484    | 0.079                 | 0.036          | 0*              |             |
| N-glycan high mannose|           |               |          | 0.105                 | 0.066          | 0.001*          |             |
| N-glycan hybrid     |           |               |          |                       | 0.013*         | 0.081           |             |
| N-glycan complex    |           |               |          |                       |                | 0*              |             |
| Other glycan        |           |               |          |                       |                |                 |             |

### B

**Glycan Interactions**

| Glycan               | Average (kcal/mol) | CI         |
|----------------------|--------------------|------------|
| N-glycan complex     | -6.69              | -6.41 - -6.95 |
| N-glycan high mannose| -6.46              | -6.31 - -6.61 |
| Fucoside             | -6.45              | -6.34 - -6.56 |
| Sialoside            | -6.42              | -6.33 - -6.51 |
| N-glycan hybrid      | -6.32              | -6.15 - -6.48 |
| O-glycan core        | -6.21              | -6.05 - -6.38 |
| Other                | -6.17              | -6.05 - -6.28 |
Table S3

CLUSTAL regions (I – III)

| Region                  | Sequence                        |
|-------------------------|---------------------------------|
| EMP46_YEAST/52-269      | DELKWNKGYSLNLLEvtQdQQKELSQTTLGDVKLEERGFVLTG...-KNTCSCIILKEPE |
| EMP47_YEAST/36-254      | DASKLSSDYSLCLDIN...ARKVPNQTGEQASLEEGRIVLTS...KQNSKCSITLQG |
| LMA1L_HUMAN/31-252      | RRFYEKLFSKGRGLA...PGAPIFSSHDAIGGLEERVL...MNRSRGAWSRAS |
| LMA1L_MOUSE/32-254      | RRFYEKLFSKGRGLAV...PGAPIFSSHDAIGGLEERVL...MNRSRGAWSNIS |
| LMA1L_RAT/32-253        | RRFYEKLFSKGRGLA...PGAPIFSSHDAIGGLEERVL...MNRSRGAWSRIS |
| LMA2L_BOVIN/49-274      | EYLKRELSKYQGVGTSSSLNLNMGANMVMTQYR...MQSOKGACGNVP |
| LMA2L_HUMAN/49-274      | EYLKRELSK...GTGSSSLNLNMGANMVMTQYR...MQSOKGACGNVP |
| LMA2L_MOUSE/40-273      | EYLKRELSK...GTGSSSLNLNMGANMVMTQYR...MQSOKGACGNVP |
| LMA2L_PONAB/49-274      | EYLKRELSK...GTGSSSLNLNMGANMVMTQYR...MQSOKGACGNVP |
| LMAN1_CHLAE/44-267      | RRFYEKYSFKGHLVQ...SDGTVPF...NANPSSDQIRVAG...LKSQGCVMTKAK |
| LMAN1_HUMAN/44-267      | RRFYEKYSFKGHLVQ...SDGTVPF...NANPSSDQIRVAG...LKSQGCVMTKAK |
| LMAN1_MOUSE/52-275      | RRFYEKYSFKGHLVQ...SDGTVPF...NANPSSDQIRVAG...LKSQGCVMTKAK |
| LMAN1_RAT/52-275        | RRFYEKYSFKGHLVQ...SDGTVPF...NANPSSDQIRVAG...LKSQGCVMTKAK |
| LMAN2_CANFA/52-276      | EHLKRELSK...GSSSMLTD...STLTSQYR...ERSKECNSTNQP |
| LMAN2_HUMAN/52-276      | EHLKRELSK...GSSSMLTD...STLTSQYR...ERSKECNSTNQP |
| LMAN2_MOUSE/54-278      | EHLKRELS...GSSSMLTD...STLTSQYR...ERSKECNSTNQP |
| YOC5_SCHPO/17-224       | AAYVPKQSLQPLQHSL...SNAIPWEK...SVE...GULYTTSKdNEARSGD...STSV |
| YQF8_SCHPO/24-248       | TSHLERL...ESAYINH...GMRNLWWEYGC...STVDRKNGIFTQd...VSNQGCVWSTRLP |

| Region                  | Sequence                        |
|-------------------------|---------------------------------|
| EMP46_YEAST/52-269      | YSTKdAMTIEETTFSFSFCPRG..STKECGILAEWLKQGNEq...DSTELFqGSSKKGGMILLR |
| EMP47_YEAST/36-254      | FDLKdSTMEWTFR...CVS...QTQGCIQRPPFRQDSNvpRDQKLYN...GPNVCGLQLLLV |
| LMA1L_HUMAN/31-252      | VPFS...A...TEVEVQ...MTCLGR...RGAQCMAV...TRGRG...HVCCYL...GLASDGICIFF |
| LMA1L_MOUSE/32-254      | VSFP...STEVQCMNTCPGR...RGAQCMAV...MTKDR...QVSY...LGEASDGIGIFF |
| LMA1L_RAT/32-253        | VSFP...STEVQCMNTCPGR...RGAQCMAV...MTKDR...QVSY...LGEASDGIGIFF |
| LMA2L_BOVIN/49-274      | CFLR.D...AEIQVFHR...QGGKlnLHGD...ATL...TKDR...QPC...FG...NMDKGVLGGVFD |
| LMA2L_HUMAN/49-274      | CFLR.D...AEIQVFHR...QGGKlnLHGD...ATL...TKDR...QPC...FG...NMDKGVLGGVFD |
| LMA2L_MOUSE/40-273      | CFLR.D...AEIQVFHR...QGGKlnLHGD...ATL...TKDR...QPC...FG...NMDKGVLGGVFD |
| LMA2L_PONAB/49-274      | CFLR.D...AEIQVFHR...QGGKlnLHGD...ATL...TKDR...QPC...FG...NMDKGVLGGVFD |
| LMAN1_CHLAE/44-267      | AAFE...NVEVEVTNRVGR...IGACDAILYTNQQ...LCP...FG...SADLNGVIFF |
| LMAN1_HUMAN/44-267      | AAFE...NVEVEVTNRVGR...IGACDAILYTNQQ...LCP...FG...SADLNGVIFF |
| LMAN1_MOUSE/52-275      | AAFE...NVEVEVTNRVGR...IGACDAILYTNQQ...LCP...FG...SADLNGVIFF |
| LMAN1_RAT/52-275        | AAFE...NVEVEVTNRVGR...IGACDAILYTNQQ...LCP...FG...SADLNGVIFF |
| LMAN2_CANFA/52-276      | CFLR.D...AEIQVFHR...QGGKlnLHGD...ATL...TKDR...QPC...FG...NMDKGVLGGVFD |
| LMAN2_HUMAN/52-276      | CFLR.D...AEIQVFHR...QGGKlnLHGD...ATL...TKDR...QPC...FG...NMDKGVLGGVFD |
| LMAN2_MOUSE/54-278      | CFLR.D...AEIQVFHR...QGGKlnLHGD...ATL...TKDR...QPC...FG...NMDKGVLGGVFD |
| YOC5_SCHPO/17-224       | LRQV...GQLSTS...VAHVSe...NENTFFAIRY...TA...AVG...SE...PP...FG...ASD...LISQ |
| YQF8_SCHPO/24-248       | TPSS...SDYFQFFIT...INRENT...LFGD...AFLAARA...KPC...FG...FTD...NGVIFF |
EMP46_YEAST/52-269
EMP47_YEAST/36-254
LMA1L_HUMAN/31-252
LMA1L_MOUSE/32-254
LMA1L_RAT/32-253
LMA2L_BOVIN/49-274
LMA2L_HUMAN/49-274
LMA2L_MOUSE/48-273
LMA2L_PONAB/49-274
LMA1L_CHLAE/44-267
LMA1L_HUMAN/44-267
LMA1L_MOUSE/52-275
LMA1L_RAT/52-275
LMA2L_CANFA/52-276
LMA2L_HUMAN/52-276
LMA2L_MOUSE/52-278
YOC5_SCHFO/17-224
YQF8_SCHFO/24-248

LDN--.....KLGESVRAYLDGCKDLDIES---PYF---SELFQYQDSMVPSTLRTLTPNP
NNG--.....PLGTLRGLNDGQKPVRKTKI-TYDQSFPSLGMCYQDSVPSTIRTVYDL
SPAEDT--....QDSRAIRLSDHCEPQGCSAQGGLCSWDIEFRPHFPRARITYWG
SSTSBDV--....QDSRAIRLSDHDLQCSGQGRVRELSCRDFRNFPRPHPRARITYWR
SSSDNV--....QGSNARILSDHDLQCSQGQGRVRELSTRDFRNFPRPHPRARITYWR
TYPNEEkqgeRVFgYSASVMNGLSYSEHERPCTELEGCTAIARNLHYDFTLVRYVK
TYPNEEkqgeRVFgYSASVMNGLSYSEHERPCTELEGCTAIARNLHYDFTLVRYVK
TYPNEEkqgeRVFgYSASVMNGLSYSEHERPCTELEGCTAIARNLHYDFTLVRYVK
SFDNDGk...KNNP-AIVINTGGNRQHYBDQCASCQALSCQDRFKPFPVFRAKITYQ
SFDNDGk...KNNP-AIVINTGGNRQHYBDQCASCQALSCQDRFKPFPVFRAKITYQ
SFDNDGk...KNNP-AIVINTGGNRQHYBDQCASCQALSCQDRFKPFPVFRAKITYQ
SFDNDGk...KNNP-AIVINTGGNRQHYBDQCASCQALSCQDRFKPFPVFRAKITYQ
TYPNEDet.eRVFgYSASVMNGLSYSEHERPCTELEGCTAIARNLHYDFTLVRYVK
TYPNEDet.eRVFgYSASVMNGLSYSEHERPCTELEGCTAIARNLHYDFTLVRYVK
TYPNEDet.eRVFgYSASVMNGLSYSEHERPCTELEGCTAIARNLHYDFTLVRYVK
TYPNEDet.eRVFgYSASVMNGLSYSEHERPCTELEGCTAIARNLHYDFTLVRYVK
IDQGK----...IFVRGYNLDSKFLAQFTPDLPFPKCTIESSPEA-LNNILLLHD
TYNNHr...gTLELETVYMKGQLHTPYEDKNASEE...CSALNRGNDYNLGLKDK

KLYDG
QFTNG
LTESL
LTESL
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