Natural single amino-acid polymorphism (F19Y) in human galectin-8: detection of structural alterations and increased growth-regulatory activity on tumor cells

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Abbreviations
CRD, carbohydrate recognition domain; CD, circular dichroism; hGal-1, human galectin-1; hGal-8, human galectin-8; ITC, isothermal titration calorimetry; MMP, matrix metalloproteinase; SNP, single-nucleotide polymorphism
Abstract

Natural amino-acid substitution by single-site nucleotide polymorphism can become a valuable tool for structure-activity correlations, especially if evidence for association to disease parameters exists. Focusing on the F19Y change in human galectin-8, connected clinically to rheumatoid arthritis, we here initiate the study of consequences of a single-site substitution in the carbohydrate recognition domain of this family of cellular effectors. We apply a strategically combined set of structural and cell biological techniques for comparing properties of the wild type and variant proteins. The overall hydrodynamic behavior of the full-length protein and of the separate N-domain is not noticeably altered, but displacements in the F0 $\beta$-strand of the $\beta$-sandwich fold in the N-domain are induced, as evidenced by protein crystallography. Analysis of thermal stability by circular dichroism spectroscopy revealed perceptible differences for the full-length proteins, pointing to an impact of the substitution beyond the N-domain. In addition, small differences in thermodynamic parameters of carbohydrate binding are detected. On the level of two types of tumor cells, characteristics of binding appeared rather similar. In further comparison of influence on proliferation, the variant proved to be more active as growth regulator in the six tested lines of neuroblastoma, erythroleukemia and colon adenocarcinoma. The seemingly subtle structural change identified here thus has functional implications in vitro, encouraging further analysis in autoimmune regulation and, in a broad context, work with other natural single-site variants, using the documented combined strategy.

Introduction

The growing awareness of the unsurpassed coding capacity of carbohydrates provides increasing incentive to study the expression and structures of mammalian lectins [1]. These sugar-binding proteins translate glycan-encoded messages into responses on the level of cells [2]. By targeting distinct $\beta$-galactosides of selected glycoconjugates, the members of the galectin family do so, the formation of a lectin-glycan complex then initiating signaling, for example for growth regulation, or facilitating glycoprotein delivery to distinct cell types or within apical or axonal transport [2-4]. Obviously, these effectors are an attractive object to study the molecular route from coding by glycans to function.

A common way to delineate structure-function correlations for proteins is the strategic introduction of single-site mutations. For human galectins, the essential nature of a sequence signature, especially a strictly conserved Trp residue for ligand contact, could hereby be documented, starting with work on human galectin-1 (hGal-1) [5]. Later, long-range effects that alter thermodynamic parameters of ligand binding were detected, explicitly for the C2S and R111H mutants of this lectin [6]. These studies underline the significance of work with engineered variants, revealing that even small sequence changes can alter protein properties beyond the site of substitution. Considering the natural occurrence of single-
nucleotide polymorphisms (SNPs), the comparative analysis of the resulting mutant proteins offers an attractive approach for structure-activity research. Of particular note, these deviations from the common genotype may show correlations to clinical parameters, giving work on the variant proteins a biomedical dimension. Herein, we initiate the structural and functional characterization of SNP-based variation in the carbohydrate recognition domain (CRD) of galectins.

Up to now, SNP identification in this lectin family mostly concerned other regions of the genes. For galectins-1 and -2, these are the promoter as well as 5’- and 3’-flanking sequences and introns 1 and 2, with indications for an association to autoimmune myasthenia gravis (hGal-1) and myocardial infarction (hGal-2) [7-10]. As to SNP presence in the coding region, two allelic variations had been discovered in the sequence section for collagen-like repeats of chimera-type galectin-3, at positions 191 (P64H) and 292 (T98P) [11]. The P64H substitution, only found in humans, confers resistance to cleavage by matrix metalloproteinases (MMPs), especially MMPs-2/-9, on the level of the protein, and to doxorubicin for cells, increases sensitivity to death receptor-mediated apoptosis and appears to be associated with increased incidence of breast cancer but reduced risk for prostate carcinoma [12-14]. Occurrence of the T98P change has been reported to be increased in Taiwanese patients with rheumatoid arthritis and Chinese glioma patients as well as related to shortened survival in Chinese non-small cell lung cancer patients [15-17]. In contrast the P64H substitution showed no correlation to disease onset or progression in these three studies. Recently, four non-synonymous SNPs in exons of the gene for human galectin-8 (hGal-8) have been pinpointed. SNP rs2737713 (A>T, which causes a F19Y substitution) was selected for quantitative trait-association study and found to be significantly associated with rheumatoid arthritis [18]. Since this is the first SNP hitting a galectin’s CRD examined, which is related to a clinical parameter, we addressed the question on alterations of structure and cell biological functions using an interdisciplinary approach.

Gal-8 is a tandem-repeat-type protein, in which two different CRDs are connected by a linker [19]. Phylogenetically, its representation in the galectin network is rather conserved; e.g. the chicken genome harbors an orthologue as the only tandem-repeat-type galectin [4, 20]. It is widely present in human tissues, localized extra- and/or intracellularly (cytoplasm and/or nucleus), secreted from cells, and useful for diagnostic and prognostic assessments in histopathology, with an expression profile different from those of other galectins [21-26]. As cellular effector, it induces apoptosis or growth arrest in immune (synovial fluid cells of rheumatoid arthritis patients, CD4\textsuperscript{high}CD8\textsuperscript{high} thymocytes) and tumor cells, and also modulates cell adhesion, here by virtue of reactivity with integrins such as the \(\alpha_3\beta_1\)-heterodimer and integrin counterreceptors [27-31]. A tangible role in autoimmune regulation is further substantiated by its concentration-dependent activity to promote T cell proliferation without antigen (high concentration) and to act as co-stimulator together with antigen (low concentration) [32, 33]. Moreover, it assists Gal-1 in
differentiating mature B cells into plasma cells [34]. Fittingly, sera of patients with autoimmune disease contain autoantibodies against this galectin, whose production in patients with systemic lupus erythematosus reached the status of an independent biomarker for secondary anti-phospholipid syndrome [35, 36].

On the level of glycans as ligands, 3’-sialylated/sulfated β-galactosides, N-acetyllactosamine repeats and histo-blood group ABH epitopes are preferred binding partners of the lectin, with affinity differences between the N- and C-domains, when tested separately [37-42]. Crystallographic analyses of the N-domain pinpointed Arg-59 as the most critical amino acid for the up to nM affinity toward 3’-sialylated/sulfated lactosides, and its absence in the C-domain explained the marked disparity in the respective affinity when examining both domains [43, 44] and also galectin-9’s N-domain [45]. Recently, the crystal structures of a linker-trimmed variant of hGal-8 and of the separate C-domain in complex with a peptide of the autophagy cargo receptor NDP52 have become available [46, 47]. The carbohydrate specificities of the two CRDs and their mode of presentation as pair connected by the linker determine Gal-8’s functionality as lectin. Occurrence of the SNP (in exon 1), mentioned above, may affect one or both aspects, a question answerable by structural and functional studies. Toward this end, we have devised a strategic combination of different approaches. We first report on the hydrodynamic behavior, circular dichroism (CD)-based analysis of structure and thermal stability of wild type and variant proteins, followed by presenting the crystal structures of the N-domains, then a detailed thermodynamic analysis of ligand binding, and finally function-oriented results on galectin binding to cells, regulation of growth in six different tumor cell lines and mediation of aggregation of erythrocytes.

Results

Quaternary structure and hydrodynamic properties

We started the comparative analysis of wild type hGal-8 and its F19Y mutant, working in parallel with the sets of full-length proteins and the N-domains, by characterizing their behavior in sedimentation equilibrium in the absence and presence of 0.1 M lactose. Data obtained at loading concentrations in the range of 0.3-1.3 mg/ml could invariably be fitted to a model with a single ideal component with weight-average molecular masses comparable to the theoretical masses of the monomers and to those experimentally determined by mass spectrometry (Table 1). The experimental values were independent of rotor speed, indicating homogeneity of protein preparations. Thus, in this range of concentrations, all the proteins behaved in solution predominantly as monomers.

Gel filtration analysis was performed in the presence of 0.1 M lactose to block any carbohydrate-dependent interaction with the resin. All the proteins eluted as a single sharp peak with no evidence for aggregate formation. Apparent molecular masses calculated from the elution times (Table 1) were in very
good agreement with monomer theoretical masses, showing that the hydrodynamic features of the full-length proteins and separate N-domains do not deviate significantly from those expected for globular proteins. Sedimentation velocity experiments further supported gel-filtration and sedimentation equilibrium results. The proteins were found to migrate as a main peak with almost identical $s_{20,w}^0$ values in the absence and presence of lactose (Table 1). This result excludes major changes in the hydrodynamic shape of the proteins by ligand loading, which had been encountered for hGal-1 as reduction of the gyration radius or increase of the diffusion constant [48, 49]. Frictional coefficient ratios ($f/f_o$) of about 1.1 for the separate N-domains and 1.35 for the full-length proteins were calculated. In comparison, the frictional ratio for human hGal-1 was 1.3 [50], suggesting that there are no main differences in the overall hydrodynamic shape between these two tandem-repeat-type proteins and the proto-type galectin, apart from a predictable mass-proportional alteration of the length.

Overall, the results of the gel filtration and ultracentrifugation analyses documented that the F19Y mutation does not have any traceable impact on the association state or hydrodynamic properties of either the separate N-domains or the full-length proteins. Besides, they indicated that dimers of full-length hGal-8 or its N-domain detected by chemical cross-linking at 5 µM protein concentration in the presence of a 50-fold molar excess of bis(sulfosuccinimidyl) suberate [40] can be interpreted as covalent trapping of rather transient contacts. In the quest to detect any differences between wild type and mutant proteins CD spectroscopy was next applied.

**CD-based analysis of the structure and thermal stability in solution**

The far-UV CD spectrum of hGal-8 showed the presence of a negative band centered at 218 nm (Fig. 1A), consistent with β-sheet structure. No differences were detected in the spectra of the wild type and F19Y proteins. When normalized per mole of molecule, the ellipticity signals for the full-length proteins approximately doubled those registered for the separate N-domains, in agreement with their expected total content in β-sheet structure. Interestingly, a slightly higher negative ellipticity in the 215-225 nm region was detected for the wild type N-domain compared to its F19Y mutant, hinting at a potentially distinctive feature of the β-sandwich arrangement of this domain resulting from the Phe to Tyr substitution. When monitored in the presence of 0.1 M lactose, spectra obtained for all the proteins maintained their profiles, indicating that there were no significant changes in secondary structure upon ligand binding (data not shown).

The near-UV CD spectrum of hGal-8 was characterized by the presence of a broad positive signal, from 260 to 300 nm, with well-defined bands at 264, 272, 282 and 291 nm (Fig. 1B), attributable to fine structure of Phe, Tyr and Trp residues. The spectrum was sensitive to the F19Y substitution, with a noticeable decrease in intensity of the peaks at 272 and 282 nm. Differences were proportionally higher when comparing the spectra of the N-domains, most probably due to the greater relative contribution of
residue 19 (Phe vs Tyr) to the overall spectrum of the molecule. In all cases, a significant increase in the ellipticity signals at 282 and 291 nm, in the tyrosine/tryptophan region, was observed when the spectra were obtained in the presence of lactose, while changes in the 264-272 nm region were very minor or not even present (Fig. 2A-D), suggesting that the environment of phenylalanine residues is only barely perturbed by lactose binding. However, for all the proteins, loading with 3’-sialyllactose affected the whole spectrum, including the region attributable to phenylalanine residues (Fig. 2A-D), revealing that binding of this ligand has a larger impact than lactose on the environment of aromatic residues and/or protein dynamics. Whether the noticed differences in the near-UV CD spectra are translated into different properties of the wild type and mutant proteins was first examined by measuring their thermal stability in the absence and presence of ligands.

Stepwise heating of all the proteins (full-length and N-domains) resulted in the loss of secondary structure, as evidenced by a progressive decrease in the intensity of the far-UV negative band. Therefore, the denaturation process was monitored by measuring the change in ellipticity at 218 nm as a function of temperature. A phenomenological analysis of the curves using a sigmoidal function yielded the $T_{1/2}$ values compiled in Table 1. Of note, the full-length proteins apparently denatured in a single cooperative process (Fig. 1A, inset), as also observed for the separate N-domains. However, while the thermal stability of the N-domains was very similar, the $T_{1/2}$ of the full-length form of the F19Y variant was 2.5 ºC higher than that of the wild type protein. This result identifies an impact of the F19Y substitution on the level of the full-length protein not confined to the N-terminal domain.

In all cases, thermal stability increased with lactose binding (Fig. 1A, inset and Table 2). $T_{1/2}$ values were 9.1 to 9.5 ºC higher for the N-domains and wild type hGal-8, whereas the increase for the F19Y variant was 7.4 ºC, again unveiling distinctive features for this protein. In view of the differences in near-UV CD spectra run in the presence of 3’-sialyllactose, a similar analysis of the effect of protein loading with 3’-sialyllactose was performed. It yielded a different picture. In particular, the stabilization induced by this ligand was similar for wild type and F19Y hGal-8 but the $T_{1/2}$ increase was considerably higher (more than 6 ºC) for the separate N-domains than for the full-length proteins. As only the N-terminal CRD is responsible for 3’-sialyllactose binding and the affinities are similar for all proteins (please see below), the lower degree of 3’-sialyllactose-induced stabilization of the full-length forms should be related, again, with denaturation features of regions beyond the N-domain. Having herewith delineated significant differences in thermal stability, a meticulous examination of possible alterations at the atomic level was performed using X-ray crystallography. Attempts to crystallize the full-length proteins were not successful. In fact, no crystal structure of a full-length mammalian tandem-repeat-type galectin has been solved to date, only the structure of a non-physiological linker-trimmed hGal-8 variant having been reported [44]. Thus, we focused on the N-domains for comparative analysis, as described below, the
position of the sequence variation being suited for this processing.

**Crystal structure of the N-domains in complex with lactose**

The three-dimensional structure of the N-domain of wild type hGal-8 loaded with lactose was determined at 1.35 Å resolution. The final model comprised 148 protein residues, 252 water molecules, one lactose molecule, one Zn$^{2+}$, one Na$^+$ and one molecule of glycerol (for details, please see Table 2). The structure presents the canonical “jelly roll” topology, with two sets of antiparallel β-sheets forming the β-sandwich arrangement typical of all known mammalian galectins. In addition to the common six- and five-stranded β-sheets (S1-S6, F1-F5), the structure reveals the presence of a previously unassigned extra β-strand [43, 44], termed F0 (Fig. 3A), as similarly found for the separate N-domain of human galectin-9 [45]. Strand F0 runs in antiparallel manner to the carboxy-terminal F1 strand, the two strands establishing a network of backbone hydrogen bonds (Fig. 3B). The carbohydrate-binding site is formed by residues from β-strands S4-S6 (Fig. 3C). Arg-69 interacts through hydrogen bonds with the OH-3 of the glucose unit of lactose, while the galactose unit stacks with the indole ring of Trp-86 as well as establishes direct hydrogen bonds with Arg-45, His-65, Asn-67, Asn-79 and Glu-89 and, through a water-mediated network, with Arg-59 and Trp-86. A strong electron density close to OH-3' of galactose was modeled as a Na$^+$ cation. In addition, a molecule of glycerol (the additive used for cryoprotection) was identified close to Tyr-141.

Superposition with the structure available for the same complex at lower resolution (2.33 Å, PDB entry 3AP4 [43]) showed only minor differences (Fig. 3D), as reflected by an RMSD value between Cα atoms of 0.23 Å, whereas comparison with the structure of the ligand-free form (PDB entry 3AP5, RMSD value 0.33 Å for all Cα atoms) revealed changes in the position of residues at the binding site. Arg-69 and Glu-89 move towards lactose for establishing hydrogen bonds with the sugar. Furthermore, the position of Arg-45, which exhibits a double conformation in the ligand-free form, is found to be stabilized upon lactose binding, and a small shift (around 1 Å) of Trp-86 is also noticed. Yet, the biggest changes are observed for the loop connecting strands S3 and S4, which moves 1.5 Å towards the sugar, thereby narrowing the ligand-binding site and displacing 2 Å the side chain of Arg-59 from its position in the ligand-free form (Fig. 3D). All these ligand-induced rearrangements within the carbohydrate-binding site are likely to affect the environment of Trp-86 and Tyr-141, hereby providing a structural explanation for the differences observed in the near-UV CD spectra of the hGal-8 N-domain in the absence and presence of lactose. Of note, no similar changes are detected when comparing the ligand-free and lactose-loaded crystal structures available for human galectins-1, -3, -7 and -9, the readjustment observed for the hGal-8 N-domain thus appearing as a distinctive feature of this CRD.

The crystal structure of the mutant N-domain in complex with lactose was solved in parallel at 1.9 Å resolution. A total of 291 residues (145 in chain A and 146 in chain B), 2 molecules of lactose and 99 of
water constituted the final model (Table 2). In the crystallographic dimer, the long loop connecting strands F5 and S2 of one subunit establishes 9 hydrogen bonds and 4 salt bridges with the edge of strands F2-F3-F4 from the neighbor subunit. According to the complex formation significance score of the PISA web server [51], these interactions, however, may not be sufficient for physiological homodimer formation, in agreement with the monomeric status determined in solution by gel filtration chromatography and analytical ultracentrifugation.

A superposition of the structures of the F19Y and wild type N-domains reflected a highly similar global structure (Fig. 4A), with RMSD between Cα atoms being only 0.2 Å. Furthermore, the carbohydrate-binding sites exhibit identical architecture and geometry of lactose binding (Fig. 4B). Focusing on the position 19, the lateral chain of Tyr-19 superposes with wild type Phe-19 (Fig. 4C). However, the presence of the tyrosine hydroxyl group induces a shift of β-strand F0, leading to a series of displacements of residues from position 15 towards the N-terminus, with Ile-11, Tyr-13, Asn-14 and Pro-15 being significantly displaced from the position occupied in the wild type structure. In the full-length protein, this strand precedes the linker peptide that connects the N- and C-terminal domains. Previous studies on the impact on human galectin-1 of single-site mutations introduced at some distance from the binding site revealed that the effects are not locally confined [6]. On the contrary, they are transmitted throughout the CRD even affecting the binding thermodynamics. Thus, it seems likely that the shift of the F0 strand observed in the F19Y variant may also be transmitted to other sections of the protein, even beyond the N-domain, thereby accounting for the observed differences in thermal stability. This reasoning prompted us to examine whether the thermodynamic parameters of ligand binding to the N-domains and to the full-length proteins could also be affected by the F19Y substitution. Thus, we next performed isothermal titration calorimetry (ITC), testing lactose and 3’-sialyllactose as in the CD-based experiments.

**Thermodynamic parameters of ligand binding**

Representative curves of calorimetric titrations with lactose and also with 3’-sialyllactose are shown in Fig. 5, and the thermodynamic parameters derived from the analysis are summarized in Table 3. Titration curves of full-length proteins with lactose were fitted to two-independent sets of sites (one per CRD) with binding constants of \(1.1 \times 10^4\) M\(^{-1}\) and 310-460 M\(^{-1}\). Comparison of binding parameters with those obtained for lactose binding to the N-domains enabled the unambiguous assignment of the high-affinity site to the N-terminal CRD (Table 3). By contrary, data obtained upon titration of full-length proteins with 3’-sialyllactose were consistent with the presence of only one set of binding sites per molecule; the one of the N-terminal CRD as titration of separate domains with 3’-sialyllactose proved (Fig. 5 and Table 3). Of note, the thermodynamic parameters indicate differences in \(\Delta H\) and \(\Delta S\) contributions to the affinity in a compensatory manner, with more negative values in the case of the full-length mutant and lactose (high-
In all other cases both contributions are more negative for the wild type complexes. In spite of sharing almost identical binding-site architectures, the enthalpic contribution to the binding of lactose to the separate F19Y N-domain was about 1 kcal/mol lower than for the wild type domain. This smaller $\Delta H$ was counterbalanced by a reduced entropic penalty, resulting in same binding affinities. It seems therefore reasonable to assume that observed differences in $\Delta H$ and $\Delta S$ can arise from distinct contributions of ligand-induced protein rearrangements beyond the carbohydrate-binding site. On the other hand, the affinity of both N-terminal sites for 3'-sialyllactose was comparable and one order of magnitude higher than its affinity for lactose, generally due to a substantially greater enthalpic contribution to the binding (Table 3). This increase is most probably attributable to near contacts of the sialic acid moiety with the protein, as observed in the crystal structure of the wild type N-domain – 3'-sialyllactose complex [43]. In direct comparison, the enthalpic contribution to the binding of 3'-sialyllactose to the F19Y N-domain is about 0.3 kcal/mol smaller than for the wild type domain, again being counterbalanced by a slightly lower entropy loss. The difference between binding thermodynamics is notably more pronounced for the full-length proteins than for the respective separate N-domains (Table 3). With $\Delta \Delta H$ of -1.3 kcal/mol and $\Delta \Delta S$ at -4.2 cal/(mol·K) and standard deviations of +/-0.1 kcal/mol or +/-0.2 cal/(mol·K), the F19Y substitution makes its presence clearly felt in the full-length proteins.

The lower accuracy of the thermodynamic parameters derived for lactose binding to the C-terminal CRD (low-affinity site) in full-length proteins precludes to reliably postulate a small effect of F19Y substitution at the site distant to position 19 (Table 3). The listed indications for alterations in binding parameters (but not overall affinity) of the N-domains and the full-length proteins for two types of free ligand prompted a comparative evaluation of the cell binding and effector activities of the wild type and F19Y hGal-8 proteins.

Comparison of functional properties

Since galectins are known to affect cell properties after carbohydrate-dependent surface binding, we measured affinity of association to cells and effects of this binding on cell growth in vitro for the two proteins in direct comparison. As test systems, we selected a line already studied with several other galectins (neuroblastoma), a leukemic and four carcinoma lines, thus representing different histogenetic origins. Binding to human neuroblastoma (SK-N-MC) cells was nearly completely blocked by the presence of glycan inhibitors (150 mM lactose and 0.5 mg/ml asialofetuin) in both cases. Measuring the extent of binding with increasing lectin concentrations, the two curves for the wild type protein and its variant were very similar, with calculated $K_D$-values ($B_{max}$-values) of about 118 nM (4.6 x 10^5 molecules/cell) for the wild type protein and 102 nM (4.9 x 10^5 molecules/cell) for the variant (Fig. 6).
The binding behavior at the level of cells therefore was in line with the similar binding affinities determined for simple sugars by ITC. Of note, the presence of ganglioside GM1-specific cholera toxin B-subunit only slightly reduces the extent of cell binding (Fig. 6). In contrast to homodimeric galectins-1, -2 and -7 whose sites of binding are masked by the bacterial protein, ganglioside GM1 apparently is not a key counterreceptor of hGal-8 on these cells. However, this difference in target specificity did not abolish the activity of hGal-8 as growth inhibitor. Both proteins were significantly active in this respect (Fig. 7). Besides presenting the effects of each protein relative to the control without galectin (Fig. 7A), we set the data obtained with the two proteins also into direct comparison. This data processing came up with a slightly increased inhibitory activity of the F19Y variant at \( p \leq 0.05 \) (Fig. 7B). To further examine this feature comparatively with cells of different histogenetic origin we next tested erythroleukemia (K562) cells.

Again, the two proteins reduced cell growth significantly, and the variant clearly proved to be more potent in this respect (Fig. 8). At the highest concentration tested, the significance threshold of \( p \leq 0.005 \) was surpassed. Thus, the detected, seemingly subtle structural changes arising from the F19Y substitution do appear to have functional implication. As documented above, extent of erythroleukemia cell binding appeared to be rather similar, as was sensitivity to the presence of lactose (Fig. 9). A similar pattern for affecting cell proliferation was determined on four human colon adenocarcinoma lines (Fig. 10). In these cases, the SNP-derived variant form was more potent to reduce growth, with differences in concentration dependence between the lines. Overall, the data on proliferation support the presence of a quantitative difference between the two types of proteins in growth regulation of the tested cell systems.

Besides characterizing the effect on proliferation, likely caused by cross-linking counterreceptors in cis-configuration on the cell surface, we determined the potency of the proteins to form cell aggregates in classical haemagglutination. Herein, rabbit erythrocytes are cross-linked in trans-configuration by the bivalent proteins. The minimal concentration for a positive response was 600 ng/assay for the wild type protein, and 1000-1200 ng/assay for the variant. Sensitivity for lactose presence was higher for the variant (concentrations above 0.8 mM blocked agglutination at 1400 ng) than for the wild type protein (at least 1.5 mM required at 1000 ng).

**Discussion**

The natural occurrence of SNP-based protein variants presents an attractive model for analyzing structure-activity relationships. Whether the amino acid substitution can affect protein properties will have a notable bearing on trait prediction from genotype data, prone to diverse pitfalls [52]. We have here initiated this line of research for galectins, focusing on the F19Y substitution in the N-terminal CRD of hGal-8. The seemingly small change, i.e. addition of a hydroxyl group to an aromatic side chain, is not at a site of direct contact with the carbohydrate ligand but part of the F0 \( \beta \)-strand. Our biophysical
characterization combining the analysis of the crystallized protein and protein in solution teaches a series of lessons:

a.) All the proteins are present as monomers in solution, tested under conditions that have detected concentration-dependent changes in the status of aggregation of a plant lectin [53], and the hydrodynamic behavior is not measurably altered by the Phe to Tyr replacement.

b.) The substitution causes displacements of amino acid side chains in its vicinity, defined by crystallography.

c.) The consequences of the F19Y substitution are not locally confined. Instead, they apparently extend through the tandem-repeat-type protein, as revealed by measuring thermal denaturation and thermodynamic binding parameters. Such an effect might be facilitated by the shift of the F0 β-strand caused by the F19Y substitution, as this strand positionally precedes the linker connecting the N- and C-terminal CRDs.

d.) The characteristics of impact on ligand binding, in comparison to the wild type protein, appear to depend on the nature of the carbohydrate ligand. The measurements with two common ligands (lactose and its 3'-sialylated derivative) led to distinct data sets in terms of enthalpy/entropy compensation, suggesting that the same may likely be true for cellular activities, if involving different counterreceptors. Of note, such a case has recently been reported for an engineered mutant of human galectin-2 [54].

At this stage, the structural analysis of the variant allowed to infer that the single-site substitution has consequences beyond its local vicinity, in principle in line with the hypothesis of possible functional correlations. On the level of the tested cell types, binding properties were rather similar for the wild type and variant proteins. As seen in ITC, affinity was not affected, when quantitating extent of binding of labeled (iodinated or biotinylated) protein. Of note, the application of the cholera toxin B-subunit revealed that ganglioside GM1 is not a neuroblastoma cell counterreceptor for galectin-8, as it is for homodimeric galectins [55-57] and also for tandem-repeat-type galectin-4 [58]. As consequence, galectin-8 will then also likely not trigger ganglioside GM1-dependent anergy/apoptosis of activated effector T cells, as galectin-1 does [59, 60]. Obviously, this observation signifies that members of the galectin family can differ drastically in distinct aspects of target selection.

Probing into functional aspects, the variant proved to be more active as growth inhibitor on the three types of tumor cells than the wild type protein. Structural changes induced by the presence of the hydroxyl group thus translate into enhanced effector potency, with quantitative differences between lines of the same histogenetic origin. In addition to measuring proliferation, we determined the two proteins’capacity for cross-linking cells, using trypsin-treated, glutaraldehyde-fixed rabbit erythrocytes. Stable aggregate formation (with interactions established in trans) required less protein and was more resistant to presence of lactose with the wild type protein. These results on forming stable aggregates
point to the perspective to further examine binding properties when applying force. Equally important, since hGal-8 is known to be present in cytoplasm and nuclei (21-26), it should be kept in mind that not only glycans are physiologic counterreceptors. Galectins in general are multifunctional, and are endowed with capacity to interact with distinct peptide motifs besides their reactivity with glycans, and this at different sites [61]. A protein interacting with hGal-8’s C-domain is an autophagy cargo receptor [46, 47, 62]. In this sense, our results on actual effects of a subtle change encourage further detailed comparisons with the two proteins using various binding partners. Along this line, the other known polymorphisms of the hGal-8 gene clearly deserve attention, as do the aspects of physiological linker-length variation by alternative splicing and of sequence divergence in phylogenesis.

Beyond the given focus on (ga)lectins, our results strongly recommend the analysis in other cases beyond evident loss-of-function occurrence, as in dectin-1’s early-stop-codon (Y238X) mutant or in variants in which sites with direct ligand contact are hit, as seen for K131Q in siglec-9 [63, 64]. In addition to galectins, ficolin-L (product of the FCN2 gene), here variants exhibiting substitutions T236M/A258S in the vicinity of the contact site [65-67], or the product of the S128R polymorphism in E-selectin, with the change located in the epidermal growth factor-like domain and triggering tethering of myeloid cells [68], are other examples of candidates for respective analysis along the lines presented herein.

Considering the aspect of interplay of (ga)lectins with specific counterreceptors inspires the idea of a pairing in association studies on SNPs. In fact, functional correlations to glycogenes, via orchestrated co-regulation by a master regulator such as a tumor suppressor [69-71] or the detection of importance of distinct glycan determinants in tumor growth/host defense by Gal-1 [72, 73], can identify partner genes for setting up meaningful combinations. Case studies on core 2 N-acetylglucosaminyltransferase connect a variant (i.e. S158C) to Gal-1-induced T lymphoma growth inhibition [74] and a second variant (i.e. V152I) to increased risk of prostate cancer [75]. Collectively, our data thus reveal an unsuspected broadness of effects of the natural deviation at a single site (F19Y), establishing a precedent for further investigations, and reported information points to the potential of guided candidate selection to design paired association studies based on functional cooperation.

**Experimental procedures**

**Protein preparation**

Wild type and variant proteins (full-length proteins with 34-amino-acid linker and the separate N-domains) were obtained by recombinant production followed by chromatographic purification and rigorous quality controls as described [18, 41]. Protein concentrations were routinely determined using the theoretical molar adsorption coefficients.
Mass spectrometry

MALDI-MS measurements were performed on a Voyager DE-PRO mass spectrometer (Applied Biosystems, Foster City, CA) equipped with a pulsed nitrogen laser (λ = 337 nm, 3 ns pulse width, and 3 Hz frequency) and a delayed extraction ion source. Ions generated by laser desorption of samples were introduced into a time of flight analyzer (1.3 m flight path) with an acceleration voltage of 25 kV, 90-93% grid voltage, 0.01-0.1% ion guide wire voltage, and a delay time of 350 ns in the linear positive ion mode. Mass spectra of the separate N-domains were obtained over the m/z range 10-25 ku, using thioredoxin (Escherichia coli, oxidized) and apomyoglobin (Calibration Mixture 3 of Sequazyme Peptide Mass Standards Kit; Applied Biosystems) for external mass calibration. Spectra of the full-length proteins were obtained over the m/z range 25-50 ku. Carbonic anhydrase and enolase from Saccharomyces cerevisiae were used in this case for external mass calibration. Sinapinic acid at 10 mg/mL in TFA 0.3%:acetonitrile 70:30 (vol/vol) was used as matrix. Samples were mixed with the matrix at a ratio of ~1:4 (vol/vol), and 1 µl of this solution was spotted onto a flat stainless-steel sample plate and dried in air.

Analytical gel filtration and ultracentrifugation

Gel filtration was performed with a Superose 12 10/300 GL column (void volume: 7.8 ml; GE Healthcare) at a flow rate of 0.5 ml/min in 5 mM sodium phosphate buffer (pH 7.2) containing 0.2 M NaCl (PBS), 4 mM β-mercaptoethanol, 0.02% NaN₃, and 0.1 M lactose. BSA (66 kDa), ovalbumin (42.7 kDa), carbonic anhydrase (29 kDa), cytochrome C (12.4 kDa), aprotinin (6.5 kDa), and vitamin B₁₂ (1.35 kDa) were chromatographed under similar conditions for column calibration. Sedimentation equilibrium experiments were carried out by centrifugation of 80 µl samples, adjusted to different protein concentrations, at 15,200, 18,000 and 25,000 rev/min for the N-domains and 10,000, 11,800 and 20,400 rev/min for the full-length proteins, as described previously [53]. Sedimentation velocity experiments were run at 45,000 rev/min for 5 h using 400 µl samples. All measurements were performed at 20 °C in an Optima XL-A analytical ultracentrifuge (Beckman Coulter, Krefeld, Germany) with an AN50-Ti rotor. Weight-average molecular weights were calculated using the HeteroAnalysis program, version 1.1.2 (http://www.biotech.uconn.edu/au/) Differential sedimentation coefficients were calculated by least-squares boundary modeling of the experimental data with the program SEDFIT [76]. Solvent density and viscosity at 20 °C were computed using the Sednterp software. The partial specific volume and degree of hydration of the proteins were calculated from the amino acid composition using the same program.

Circular dichroism

CD spectra were measured with a J-810 spectropolarimeter, equipped with a Peltier temperature control system, using a bandwidth of 1 nm and a response time of 4 s. Far-UV spectra were recorded in 0.1 cm
path-length quartz cells at a protein concentration of 0.2 mg/ml, and near-UV spectra at 1.0 mg/ml in 1 cm path-length cuvettes. Routinely, the corresponding buffer baseline was subtracted. The spectra were representative of three to four independent measurements. Thermal denaturation experiments were carried out by increasing the temperature from 20 °C to 90 °C at a scanning rate of 0.66 °C/min. Variations in ellipticity at 218 nm were monitored at steps of 0.2 °C. Thermal denaturation profiles were described in terms of the following sigmoidal function:

$$\Theta(T) = \Theta_D(T) - \left[\Theta_D(T) + \Theta_N(T)\right] / \left\{1 + \exp\left[A(T-T_{1/2}) / RT_{1/2}\right]\right\}$$

where $T$ is the absolute temperature, $T_{1/2}$ is the half-transition temperature, $R$ is the gas constant, $A$ is the constant accounting for the variation with temperature of native and denatured states, and $\Theta_D(T)$ and $\Theta_N(T)$ denote the ellipticity of the denatured and native states at temperature $T$. $\Theta_D$ and $\Theta_N$ were approximated as linear functions of temperature, as described previously [77]. The fitting of the sigmoid to the experimental data was carried out by non-linear minimum-squares, using the Origin software.

Crystallization, data collection and processing

The N-domains of wild type and F19Y proteins were concentrated to 20 mg/ml using Amicon Ultra centrifugal filter units (Millipore). Crystals were grown using the sitting drop vapor diffusion method at 22 °C, by mixing 1 μl of protein solution with 1 μl of reservoir solution. The wild type N-domain was crystallized in 20% w/v PEG 4000, 100 mM HEPES sodium salt (pH 7.2) and 10 mM zinc chloride. The N-domain of the F19Y mutant was crystallized in 30% w/v PEG 4000, 100 mM MES sodium salt (pH 6.5). Crystals were transferred to a cryoprotectant solution consisting in the reservoir solution supplemented with 20% v/v glycerol (wild type N-domain) or 20% ethylene glycol (F19Y mutant N-domain) and subsequently flash-cooled in liquid nitrogen.

X-Ray diffraction data were collected at Beamlines ID23-1 and ID29 of the European Synchrotron Radiation Facility (ESRF) and at the PROXIMA-1 beamline SOLEIL Synchrotron (France). Crystallographic data were processed using XDS [78] and Aimless [79]. The space group of crystals and unit cell dimensions are specified in Table 2. The structures have been solved in two different crystal forms. For the wild type N-domain, the tetragonal crystals contained one monomer, while the orthorhombic crystals of the F19Y variant contained a dimer in the asymmetric unit. Likewise, previously reported crystal structures of the wild type N-domain in complex with several ligands were solved in three crystal forms with dissimilar number of protein molecules (from 1 to 4) in the asymmetric unit [43], the different crystal packing interactions not affecting the overall fold. The Mathews coefficient for a monomer in the asymmetric unit was 2.74 Å³/Da for the wild type N-domain and 2.15 Å³/Da for a dimer of the F19Y mutant, which correspond to a solvent content of 55.2% and 42.7%, respectively [80].

The atomic coordinates and structure factors (codes 4BMB, 4BME) have been deposited in the Protein Data Bank.
Data Bank.

Structure determination and refinement

The structures were determined with the molecular replacement method implemented in the Phenix software suite [81], using the structure at 1.92 Å-resolution available for the N-domain of wild type hGal-8 (Protein Data Bank entry code 2YV8) as model. The translation-libration-screw (TLS) groups were defined using the TLMSD server [82]. Repetitive cycles of refinement and manual building were done using Phenix [81] and Coot [83]. Water molecules were added to the model during the last steps of refinement. Final R/R\textsubscript{free} values were 0.16/0.18 and 0.23/0.28 for the wild type and F19Y mutant structures, respectively. Validation of final models was performed using the Molprobity software [84]. Protein-protein interactions were analyzed using the PISA web server [51]. Details of structure refinement and quality of final models are presented in Table 2. Structural figures were generated using Pymol [85].

Isothermal titration calorimetry

Calorimetric titrations were performed at 25 ºC with a Microcal VP-ITC microcalorimeter (GE Healthcare, Northampton, MA, USA), as recently described for chicken galectins [86]. The proteins were exhaustively dialyzed against PBS containing 4 mM β-mercaptoethanol (PBS\textsubscript{β}) and sugar solutions were prepared with the final dialysate. Titrations were performed by stepwise injections of the sugar-containing solution into the reaction cell loaded with the protein solution at concentrations of 165-170 µM. The heat of ligand dilution was determined separately and subtracted from the total heat produced following each injection. Titration data were analyzed assuming binding models with either one or two classes of binding sites using the Microcal-ITC Origin software.

Cell assays

Haemagglutination was performed with trypsin-treated glutaraldehyde-fixed rabbit erythrocytes in V-shaped microtiter plates in V-shaped microtiter plates using 25 µL of a 5% suspension and 25 µL galectin-containing solution (serial two-fold dilution of freshly prepared stock solutions of different initial concentrations). Cells for the proliferation assays were grown in RPMI 1640 medium supplemented with 1.75% fetal calf serum (erythroleukemia (K562) cells), Eagle’s minimal essential medium containing 10% fetal calf serum (neuroblastoma (SK-N-MC) cells) and Dulbecco’s minimal essential medium containing 10% fetal calf serum (colon adenocarcinoma cell lines derived from microsatellite-instable (HCT116, Colo60H) or microsatellite-stable (SW480, SW707) tumors). All media additionally contained antibiotics (100 U/ml penicillin and 100 µg/ml streptomycin). Lectin binding to cell surfaces was analyzed either with biotinylated proteins in FACScan analysis using fluorescent streptavidin-R-phycoerythrin as reporter
or with iodinated proteins (wild type protein: 195 KBq/µg, F19Y variant: 219 KBq/µg, prepared with carrier-free Na\textsuperscript{125}I (Hartmann Analytic, Braunschweig, Germany) and Iodobeads (Pierce, Bonn, Germany)) following optimized protocols [55, 56]. Competition assays with cholera toxin B-subunit (Sigma, Munich, Germany) were performed with iodinated galectins, adding the ganglioside GM\textsubscript{1}-specific probe 1 h prior to the labeled protein as described [58]. Cell growth was assessed in parallel assays using the blue chromogen 3-(4,5-dimethyl-thiazol-3-yl)-2,5-diphenyl-tetrazolium bromide (0.5 mg/ml; Sigma) or a commercial kit (CellTiter 96; Promega, Mannheim, Germany) [58].

Acknowledgments

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References


Figure Legends

**Fig. 1.** CD spectra of wild type and F19Y mutant proteins. Far-UV (A) and near-UV (B) CD spectra obtained for the full-length wild type hGal-8 (continuous lines), its F19Y mutant (dash-dot lines), and for the respective wild type (dash lines) and mutant (dot lines) N-domains. Ellipticity values are normalized per mole of molecule. Spectra were recorded at 20 °C for 0.2 mg/ml (A) or 1 mg/ml (B) protein solutions in PBSβ. *Inset* in A, changes with temperature in the ellipticity at 218 nm of the full-length wild type (squares) and F19Y mutant (circles) proteins in the absence (open symbols) and presence (filled symbols) of 0.1 M lactose. The continuous lines correspond to the fit of a sigmoidal function to experimental data (see Experimental procedures).

**Fig. 2.** Effect of ligand binding on the near-UV CD spectra of wild type and F19Y proteins. Spectra of 1 mg/ml solutions in PBSβ of full-length wild type hGal-8 (A), its F19Y mutant (B), and the respective wild type (C) and mutant (D) N-domains were recorded at 20 °C in the absence (solid lines) and presence of 0.1 M lactose (dash lines) or 1 mM 3’-sialyllactose (dot lines).

**Fig. 3.** Crystallographic structure of the N-domain of wild type hGal-8. A, ribbon model of the domain in complex with lactose, showing the two sets of antiparallel β-strands (F0-F5 and S1-S6) that form the β-sandwich motif. B, close view of F0 and F1 β-strands highlighting interactions shorter than 3 Å which stabilize the N-terminal F0 β-strand. C, architecture of the carbohydrate-binding site, and D, superposition of our structure (light grey) with known structures of the N-domain in complex with lactose (dark green, PDB 3AP5 (43)) and in ligand-free form (light green, PDB 3AP4 (43)).

**Fig. 4.** Comparison of crystallographic structures of the F19Y mutant and wild type N-domains of hGal-8. A, ribbon model of the F19Y mutant (yellow) in complex with lactose (A chain) superposed to the structure of the respective complex of the wild type protein (blue). B, comparison of the carbohydrate-binding sites of the F19Y mutant (C atoms in yellow) and wild type (C atoms in light grey) N-domains. C, detailed comparison of the relative positioning of F0 and S1 strands in the F19Y mutant (in yellow) and wild type N-domains (in light grey). Residue 19 is labeled with an asterisk, and water molecules in the structure of the mutant protein are shown as pink spheres.

**Fig. 5.** Calorimetric titration of wild type and F19Y mutant proteins with lactose and 3’-sialyllactose. A, representative plots of the heat released per mole of lactose injected during the titration of wild type (□) and F19Y (○) hGal-8 as a function of the sugar/protein molar ratio. B, titration of wild type hGal-8 (□) and its N-terminal domain (△) with 3’-sialyllactose. Solid lines correspond to the best fit of the experimental data using two-sets-of-sites (A) and one-set-of-sites (B) models.
**Fig. 6.** Binding analysis of $^{125}$I-labeled proteins to human neuroblastoma cells. Carbohydrate-dependent binding of labeled wild type (circles, solid/short dash lines) and variant (triangles, solid/long dash lines) proteins was measured in series with increasing galectin concentration using cells cultured for five days (final density: $10^5$ cells per well), in the absence (filled symbols, solid lines) or presence of 250 µg/ml cholera toxin B-subunit (open symbols, dash lines). Results are the means of three independent experiments. Inset: Scatchard analysis of galectin binding. Measurements were performed at 37 °C.

**Fig. 7.** Effect of lectins on neuroblastoma cell growth. Proliferation was measured after 48 h incubation in the presence of wild type (black bar) and variant (grey bar) proteins, using untreated cultures (white bar), set to 100%, in parallel as control and reference for statistical data processing (A). Results of calculating statistical significance of differences between the two data sets in direct comparison between wild type and mutant proteins are given in (B). Presented data are the means of four independent experiments ± SD, level of significance is graded into three categories: * $0.05 \geq p > 0.01$, ** $0.01 \geq p \geq 0.005$ and *** $p < 0.005$.

**Fig. 8.** Effect of lectins on erythroleukemia cell growth. Proliferation was measured after indicated periods of time in the presence of given concentrations of wild type (black bar) and variant (grey bar) proteins using untreated cultures in parallel as control, set to 100%, and as reference (A). Statistical evaluation of the two data sets in direct comparison is given in (B). Results are the means of five independent experiments ± SD and two independently prepared protein batches. For grading of statistical significance, please see legend to Fig. 7.

**Fig. 9.** Cytofluorometric analysis of galectin binding to human erythroleukemia cells and its sensitivity to lactose. Cell surface staining (percentage of positive cells/mean fluorescence intensity) by labeled wild type (A) and F19Y mutant (B) proteins at 2 µg/ml and sensitivity to presence of 1 mM, 5 mM, 10 mM and 20 mM lactose (grey area: background in the absence of lectin), all numbers given from bottom (no lactose added) to top (highest lactose concentration at 20 mM and finally the background control). Aliquots of cell suspensions from the same batch were tested in parallel in three independent series with SD-values not exceeding 12.4% after data normalization.

**Fig. 10.** Effect of lectins on colon adenocarcinoma cell growth. Proliferation of cells of four lines derived from microsatellite-instable (Colo60H, HCT116) or microsatellite-stable (SW480, SW707) tumors was measured after 48 h incubation in the presence of wild type (black bar) and variant (grey bar) proteins, using untreated cultures in parallel as control and as reference, set to 100% (A). Statistical evaluation of the two data sets in direct comparison is given in (B). Results are the means of four independent experiments ± SD. For grading by statistical significance, please see legend to Fig. 7.
Table 1. Parameters of wild type version and F19Y variant of full-length hGal-8 and separate N-domains. Thermal stability in the absence and presence of ligands was analyzed using circular dichroism. * $M_w$, weight-average molecular mass; ** $M_{app}$, apparent molecular mass.

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</tr>
<tr>
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<td></td>
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<tr>
<td>$M_w$ * (kDa)</td>
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<td>33 ± 2</td>
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<tr>
<td>0.1 M lactose</td>
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<td>$s_{20,w}^0$ (S)</td>
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<td>$f/f_0$</td>
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<td>$T_{1/2}$ (ºC)</td>
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<td>0.1 M lactose</td>
<td>60.6 ± 0.1</td>
<td>61.3 ± 0.1</td>
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<td>1 mM 3'-sialyllactose</td>
<td>55.6 ± 0.1</td>
<td>58.3 ± 0.1</td>
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Table 2. X-Ray diffraction data and refinement statistics for the wild type version and F19Y variant of the N-terminal hGal-8 domain

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*values in parentheses are for outer-resolution shell
Table 3. ITC-derived thermodynamic parameters for the binding of lactose and 3'-sialyllactose to the wild type version and F19Y variant of full-length hGal-8 and separate N-domains

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<td>$\Delta H_1$</td>
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<td>(M$^{-1}$)</td>
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Figure 1

(A) Graph showing a plot of protein concentration against wavelength, with a inset graph showing a similar plot against temperature.

(B) Graph showing a plot of protein concentration against wavelength, with a peak at different wavelengths.
Figure 3
Figure 6
Figure 7

A

B

relative proliferation (% of control)

25 100

25 100

galectin concentration (µg/ml)

galectin concentration (µg/ml)
Figure 10

A

Colo60H

HCT116

SW480

SW707

Relative proliferation (% of control)

galactin concentration (μg/ml)

B

relative proliferation (% of control)

galactin concentration (μg/ml)