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Minimal epitope for Mannitou IgM on paucimannose-carrying glycoproteins

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Abstract

Paucimannosidic glycans are restricted to the core structure [Man1-3GlcNAc2Fuc0-1] of N-glycans and are rarely found in mammalian tissues. Yet, especially [Man₂₋ 3GlcNAc₂Fuc₁] have been found significantly upregulated in tumors, including in colorectal and liver cancer. Mannitou IgM is a murine monoclonal antibody that was previously shown to recognise Man₃GlcNAc₂ with an almost exclusive selectivity. Here, we have sought the definition of the minimal glycan epitope of Mannitou IgM, initiated by screening on a newly designed paucimannosidic glycan microarray. Among the best binders were Man₃GlcNAc₂ and its α 1,6 core-fucosylated variant, Man₃GlcNAc₂Fuc₁. Unexpectedly and in contrast to earlier findings, Man₅GlcNAc₂type structures bind equally well and a large tolerance was observed for substitutions on the a1,6 arm. It was confirmed that any substitution on the single a1,3-linked mannose completely abolishes binding. Surface plasmon resonance for kinetic measurements of Mannitou IgM binding, either directly on the glycans or as presented on omega-1 and kappa-5 soluble egg antigens from the helminth parasite Schistosoma mansoni, showed submicromolar affinities. To characterize the epitope in greater and atomic detail, saturation transfer difference nuclear magnetic resonance spectroscopy was performed with the Mannitou antigen-binding fragment. The STD-NMR data demonstrated the strongest interactions with the aliphatic protons H1 and H2 of the α 1-3-linked mannose, and weaker imprints on its H3, H4 and H5 protons. In conclusion, Mannitou IgM binding requires a non-substituted a1,3-linked mannose branch of paucimannose also on proteins, making it a highly specific tool for the distinction of concurrent human tumor-associated carbohydrate antigens.

Running title: Binding by Mannitou IgM of paucimannose-carrying proteins

Keywords: Mannitou; IgM; PMG; PMP; paucimannose; N-glycan; core fucose

Introduction

The biological importance of glycosylation in health and disease has long been established and continues to grow (Lau *et al.*, 2007; Lauc *et al.*, 2010; Taganna *et al.*, 2011; Bruxelle *et al.*, 2020). The glycoproteome reflects the overall cellular status since it participates either in modulation or mediation of a wide range of processes in physiological and pathophysiological conditions, *i.e.* cell adhesion, molecular trafficking and clearance, protein folding, receptor activation, signal transduction and endocytosis (Stambuk *et al.*, 2020; Lau and Dennis, 2008). Protein glycosylation patterns are modified in various human diseases, including congenital disorders of glycosylation (Houdou and Foulquier, 2020), auto immune diseases such as rheumatoid arthritis (Magorivska *et al.*, 2018) and systemic lupus erythematosus (Szabo *et al.*, 2019), and in infectious diseases (Thaysen-Andersen *et al.*, 2015). The ubiquity of glycosylation and its fundamental significance in practically every biological process exemplifies the immense potential of glycans as biomarkers (Reily *et al.*, 2018).

The core structure of all eukaryotic *N*-glycans consists of two *N*-acetyl β glucosamine (GlcNAc) and three mannose (Man) residues, Man₃GlcNAc₂, also termed paucimannosidic glycan (PMG) (Schachter, 2009). The early stages of Nglycosylation are shared among all eukaryotes and include the generation of a lipidoligosaccharide precursor Glc₃Man₉GlcNAc₂-pyrophosphate-dolichol to linked nascent glycoproteins in the endoplasmic reticulum (Aebi et al., 2010). The second phase involves the processing of asparagine-linked Glc₃Man₉GlcNAc₂ within the lumen of the endoplasmic reticulum up to Man₅GlcNAc₂ in the Golgi apparatus. This pathway diverges evolutionary in plants, invertebrates and vertebrates, that each process the Man₅GlcNAc₂ intermediate in different ways into high-mannose, hybrid, and complex *N*-glycans. In humans, an enzyme in the Golgi apparatus, *N*-acetyl β 1-2-glucosaminyltransferase (GnT-I), transfers a GlcNAc residue on the α 1,3-linked mannose of the N-glycan. Following the trimming of the α 1,6 Man arm by α mannosidase-II, this hybrid glycan is finally truncated into a PMG by an *N*-acetyl- β hexosaminidase (Hex A or Hex B) (Tjondro et al., 2019). An alternative, GnT-I independent, truncation pathway has been shown to facilitate greatly the biogenesis of α1,6 core-fucosylated Man₄GlcNAc₂ and Man₅GlcNAc₂ in GnT-I-deficient Chinese hamster ovary cells (Lin *et al.*, 1994).

Despite the great progress achieved in the field, rapid detection of alterations in the expression of specific *N*-glycans associated with pathophysiological states is still challenging. Anti-carbohydrate antibodies lend a good diagnostic tool to this aim. Mannitou is a monoclonal mouse IgM, originally named Laz6-189, raised against a 130-kDa glycoprotein from the leech central nervous system (Flaster *et al.*, 1983). It was subsequently characterized as being able to specifically recognize paucimannose-carrying glycoproteins (PMPs) (Bajt *et al.*, 1990; Zipser *et al.*, 2012). Since, other PMPs with important neurobiological functions have be detected using Mannitou, among others human neutrophil elastase (Loke *et al.*, 2017), synapsin-1 (Simon *et al.*, 2019) and AHNAK (Becker *et al.*, 2019). Intriguing recent discoveries point to paucimannose *N*-glycans (PMGs) as facilitators of an effective immune response during physiological and pathophysiological conditions (Loke *et al.*, 2016).

A recent wide cancer glycomics analysis from human cancer cell lines and tissue samples illustrated that different cancer types and the disease progression stage can be stratified based on the distribution and ratios of paucimannosidic Nglycans (Chatterjee et al., 2019). Both the rare incidence of paucimannose Nglycosylation in the extracellular environment and its upregulation to give potent signals and communicate with the immune system have made Mannitou a useful tool for the detection of paucimannose epitopes (Chatterjee et al., 2019; Becker et al., 2019: Dahmen et al., 2015: Thaysen-Andersen et al., 2015: Loke et al., 2017: Zipser et al., 2012; Simon et al., 2019). Paucimannose-carrying proteins are exposed on the plasma membrane of transforming cells as opposed to high-mannose-carrying proteins predominantly hidden in the cytoplasm of normal cells (Loke et al., 2016), enabling their targeting by monoclonal antibodies such as Mannitou IgM. In this study, we highlight interaction regions on the Man3GlcNAc(Fuca1-6)GlcNAc PMG with paratope residues of the Mannitou variable domains and demonstrate how specificity and affinity of Mannitou is fine-tuned to differentiate between paucimannosidic and other *N*-glycans on proteins.

Results

Ligand screening with microarrays of defined paucimannose N-glycans

Initially, the characterization of carbohydrate binding profile of Mannitou IgM was performed with the help of glycan microarrays, a technique that enables the high-throughput screening of protein-carbohydrate interactions. Mannitou IgM was assayed against a microarray consisting of 135 different glycan structures (Figure S1), among which multiple paucimannose-type *N*-glycans.

This glycan microarray has been exploited previously to guantify the immune response against S. mansoni parasitic infection (Nkurunungi et al., 2019) and therefore, the glycans included comprise structures from non-mammalian origin commonly found in invertebrates and plants with substitutions such as core a1.3 fucose (Fuc) and β1,2 xylose (Xyl) (Brzezicka *et al.*, 2015). Two-fold serial dilutions of Mannitou IgM (500, 250, 125 and 62.5 nM) were incubated on the microarray at RT for one hour and interactions were revealed with Alexa Fluor 555 Goat Anti-Mouse IqM secondary antibody (Figure S2A). Previously, Mannitou antibody has been screened against the glycan array 4.0 of the Consortium for Functional Glycomics and demonstrated a highly restricted specificity for the Man₃GlcNAc₂ pentasaccharide (Zipser et al., 2012) and any substitution inhibited or significantly reduced binding. In line with these previous findings, our results show that Mana1-3(Mana1-6)ManB1-4GlcNAcB1-4GlcNAc (GL41), being the common pentasaccharide core M3GN2 of all **N-glycans, displays one of the highest RFU values**. Nevertheless, in our experiment, Mannitou binding was not exclusive for this core structure and different printed glycans exhibited important recognition (Figures 1 and S2B).

The main conclusions drawn from the microarray screening is that the printed structures efficiently recognized by Mannitou all have in common an unsubstituted mannose (also called residue 4 in high-mannose-type oligosaccharides, lacking substitution by α 1,2-linked mannoses C and D1 (Kozutsumi *et al.*, 1986)) of the α 1,3 branch in the *N*-glycan core. All substitutions on the α 1,3 Man arm either with Man α 1-2 (**GL42**) or with GlcNAc β 1-2 (**GL55**) completely abolished Mannitou IgM binding (Figure 1A). On the other hand, the substitution with core α 1,6-fucose in Man3 *N*-

glycan (**GL70**) showed similar RFU values to the non-fucosylated structure (**GL41**). In fact, Manα1-3Manβ1-4GlcNAcβ1-4GlcNAc (**GL40**), a dimannose-containing *N*-glycan lacking the α 1,6 Man arm, and its β 1,2 xylosylated counterpart (**GL2**) were also recognized, albeit with much lower intensity than their Man3 equivalent (respectively **GL41** and **GL7**, Figures 1A and S2). On the other hand, the sole α 1-6-linked dimannose-containing structure Man α 1-6(Xyl β 1-2)Man β 1-4GlcNAc β 1-4GlcNAc (**GL1**) did not bind (Figure S2), despite the absolute tolerance for β 1,2 xylosylation on the central mannose of the *N*-glycan, thus confirming the specificity of Mannitou IgM towards the nature of the α 1,3 glycosidic linkage.



■: N-Acetyl-D-glucosamine ●: D-Mannose ◀: L-Fucose

Figure 1. Mannitou IgM binding screening on the glycan microarray.

A. The binding of Mannitou IgM at different concentrations (62.5, 125, 250 and 500 nM) to selected glycan structures is represented by its mean RFU (relative fluorescence units), with the standard deviation of the mean shown as error bars. **B.** Binding curves for the binding.of Mannitou IgM to M3GN2 and M3GN2F1(6). The data were fitted according to a 1:1 Langmuir binding model.

For Mana1-3(Mana1-6)Man β 1-4GlcNAc β 1-4GlcNAc (**GL41**) and Mana1-3(Mana1-6)Man β 1-4GlcNAc β 1-4(Fuc α 1-6)GlcNAc (**GL70**), apparent equilibrium dissociation constants (K_{dsurf}) on the microarray surface were calculated by plotting the fluorescence intensities (relative fluorescence units, RFU) against the antibody concentrations and fitting the data to a one-site binding Langmuir isotherm (Liang *et al.*, 2007). Both glycans showed apparent dissociation constants in the submicromolar range (Figure 1B).



Figure 2. Dual color heat map representation of the normalized glycan microarray binding profiles of Mannitou, *Concanavalin A* (ConA) and *Galanthus nivalis* agglutinin (GNA). Hierarchical clustering is based on Euclidean distance (average linkage method). For glycan structures, see Figure S1.

The binding pattern of Mannitou IgM on the glycan microarray was compared with those of other mannose-binding lectins, namely concanavalin A (*Canavalia*

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ensiformis agglutinin, ConA) and *Galanthus nivalis* agglutinin (GNA) (Figure 2). A heat map allows to compare the performance of Mannitou IgM with the two lectin lectins to define the most suitable probe for applications. From all structures available in the glycan array, only paucimannose, high mannose and hybrid type structures are represented in the heat map. Although ConA, GNA and Mannitou all recognize mannose, important differences between the three proteins are highlighted (Figure 2). All recognition probes bind M3GN2 and M5GN2 N-glycan structures efficiently, but Mannitou recognized Xyl β 1-2-substituted M3GN2 N-glycans much more efficiently than ConA while GNA showed no binding to these structures (Brzezicka *et al.*, 2015). Additionally, both ConA and GNA bind to several high mannose and hybrid type structures (e. g. **GL65**, **GL54**, **GL45**) which are not recognized by Mannitou antibody. These results highlight the importance of defining in a precise manner the binding epitopes for the different carbohydrate-binding proteins.

Based on the microarray screening results and on the reported paucimannose type structures described in pathological processes (Becker *et al.*, 2019; Chatterjee *et al.*, 2019), M3GN2 (**GL41**) and the corresponding α 1-6 core-fucosylated structure M3GN2F1(6) (**GL70**), were selected and synthesized chemo-enzymatically (Figure S3) in sufficient amounts for subsequent Surface Plasmon Resonance (SPR) kinetic binding experiments and Saturation Transfer Difference-Nuclear Magnetic Resonance (STD-NMR) structural analyses.

Molecular recognition studies of paucimannose N-glycans by Mannitou IgM

Mannitou IgM could be purified using size-exclusion chromatography due to its large molecular mass (Figure S4). The next steps involved investigating the molecular recognition of paucimannose *N*-glycans by Mannitou IgM. We deciphered their specificity for Mannitou IgM by determining the kinetic parameters of binding using SPR detection (Figure 3). The synthetic M3GN2 (**GL41**) was immobilized on a CM5 sensor chip, as it was the minimal epitope to Mannitou IgM, and M3GN2F1(6) (**GL70**) on the same chip to run the microfluidics experiment under exactly the same conditions. *N*-glycan core α 1-6-linked fucose is an important residue in carbohydrate-carbohydrate interactions, involved in complement activation and antibody-mediated

cellular cytotoxicity (Ferrara *et al.*, 2011; Sakae *et al.*, 2017). While β 1,2-linked xylose and α 1,3-linked fucose are immunogenic *N*-glycan motifs from plants and invertebrates but absent in mammals (Brzezicka *et al.*, 2015), in mammals core fucose is found exclusively as α 1-6-linked to the reducing-end *N*-acetylglucosamine (GlcNAc) moiety of the chitobiose core (Serna *et al.*, 2011).





channels. Full regerenation between cycles was obtained using 50 mM NaOH. RU: Response Units.

Mannitou IgM binds with a very fast, concentration-dependent, *on*-rate (k_{on}=k_a[C]+k_d) to these PMGs together with rather large dissociation rate (k_{off}=k_d). (Table I). Such kinetics of binding are most likely helped by the dense immobilization of the glycans and re-binding effects. Indeed, only a small portion of bound IgM "sticks" and maintains a stable interaction (Figure 3). Nanomolar affinities are obtained, with no significant difference between M3GN2 and M3GN2F1(6). We next wanted to gain insight into the influence of the presentation of PMG epitopes onto proteins, paucimannose-carrying proteins or PMPs.

Ligand	Imm. RU	R _{max} RU	SE(R _{max}) RU	ka M ⁻¹ .s ⁻¹	SE(<i>k</i> _a) M ⁻¹ .s ⁻¹	<i>k</i> _d s ⁻¹	SE(<i>k</i> _d) M ⁻¹ .s ⁻¹	K _d nM	SE(K _d) nM	Chi ²
M3GN2	237	132.5	0.5	300816	2142	2.07E-03	1.72E-05	6.87	0.08	26.86
M3GN2F1(6)	163	28.7	0.2	470885	6304	2.77E-03	3.51E-05	5.87	0.11	4.33

Table I: Immobilization rate and fitted global parameters for the kinetics of PMG binding by Mannitou IgMSE: Standard Error. RU: Response Units.

Glycoprotein differentiation by Mannitou IgM

Mannitou IgM was allowed to interact with differently glycosylated forms of omega-1 (ω 1) (van Noort *et al.*, 2020) and kappa-5 (κ 5) (Wilbers *et al.*, 2017), two major immunomodulatory Schistosoma mansoni soluble egg antigens. Mannitou IgM interacted with the omega-1 glycoproteins with much lower on- rates but also smaller off-rates than for PMGs alone (Table II). This could be explained because only two glycosylation sites are present per omega-1 protein, thus the density of PMG immobilization is much lower. Another difference is that the dextran matrix was completely shielded from interaction with Mannitou IgM by the high protein immobilization rate (Figure 4). The high-mannosylated glycoform of $\kappa 5$ (Figure S5) was found not to be bound by Mannitou IgM and it was therefore used to block aspecific binding in the reference flow channel. Regeneration between the different avoided analyte concentrations was because the paucimannose-carrying alycoproteins (PMP) could be damaged using harsh eluents, instead, a single cycle kinetic titration was used. The Mannitou IgM analyte was titrated at five different concentrations, from low (59.4 nM) to high (950 nM).

Ligand	Imm. RU	R _{max} RU	SE(R _{max}) RU	k _a M ⁻¹ .s ⁻¹	$\frac{SE(k_{a})}{M^{-1}.s^{-1}}$	<i>k</i> _d s ⁻¹	SE(k _d) M ⁻¹ .s ⁻¹	K _d nM	SE(K _d) nM	Chi ²
ω1 M3GN2	2324	66.4	1.5	3629	187	3.82E-4	8.25E-06	105	6	4.96
ω1 M3GN2F1(6)	2061	68.0	2.2	4056	298	3.10E-4	1.11E-05	76.5	6.3	16.02
ω1 M3GN2F1(3)	2505	28.8	1.4	4374	388	4.67E-4	1.84E-05	107	10	11.56
ω1 M5GN2 & Le ^x	3271	32.9	3.6	7088	1181	5.78E-4	5.53E-05	81.6	15.7	14.38

Table II: Immobilization rate and fitted global parameters for the kinetics of PMP binding by Mannitou IgMSE: Standard Error. RU: Response Units.



Figure 4. Kinetic binding experiments of Mannitou IgM to immobilized PMPs Single cycle kinetics, or kinetic titration, of Mannitou IgM onto different PMPs, of which the major glycoform(s) is/are shown(n) in the inset (see experimental procedures for full details). Sensorgrams (red) after subtraction of non-binding kappa-5 M7GN2/M8GN2, see Figure S5) in Fc1 and fitting (black) according to a 1:1 Langmuir binding model. (**A**) ω 1 M3GN2, (**B**) ω 1 M3GN2F1(6), (**C**) ω 1 M3GN2F1(3), (**D**) ω 1 M5GN2 + Le^x. RU: Response Units.

Affinities (K_d) fit near 100 nM, with a small favour of Mannitou IgM for ω1 M3GN2F1(6) and M5GN2 (Table II and Figure 5). For another omega-1 PMP glycan-engineered with a single Le[×] epitope (second *N*-glycan shown in Figure 4D) as the major glycoform (Wilbers *et al.*, 2017), no binding was detected using SPR. In line with this observation, Le[×] epitopes presented on the glycan microarray (e. g. **GL91** and **GL92**) were not recognized by Mannitou IgM (Figure S2A). Therefore, the affinity for omega-1 presenting mixed major glycoforms of M5GN2 and Le[×] (Figure 4D) could be most probably attributed to oligomannose-5 binding. This finding is again congruent with the results of IgM binding to M5GN2 (**GL43**) of the glycan microarray (Figure 1A). Possibly, the binding of M5GN2 (**GL43**) is a little stabilized over M3GN2 (**GL41**) consistently between microarray screening and SPR measurements (Table II) and judged based on the maximal binding capacity (Rmax) and affinity (Figure 5).



Figure 5. Comparison of PMP with PMG binding of Mannitou IgM

Maximal binding (Rmax, light grey) and affinities (K_a, dark grey) of Mannitou IgM analyte for the two M3GN2 and M3GN2F1(6) PMG ligands, coupled via the amino-pentyl spacer at their reducing end on the sensor chip (Figure S3), and comparison with their presentation as asparagine-linked *N*-glycans on the immobilized omega-1 proteins.

Epitope mapping by STD-NMR spectroscopy

STD-NMR spectroscopy is often used to determine molecular surface patches/regions on glycan epitopes that are directly involved in binding proteins such as lectins and carbohydrate-binding antibodies (Arda and Jimenez-Barbero, 2018; Henriques *et al.*, 2020). In spectra collected in aqueous, i.e. polar, buffers, only aliphatic and aromatic epitope proton resonances are visible, while the polar/acidic set of H atoms in glycans (those that are part of hydroxyl, amine and carboxy groups) rapidly exchange with the surrounding solvent molecules (Blaum *et al.*, 2018).

For the STD-NMR experiments, the antigen-binding fragment (Fab) of Mannitou was generated by transient transfection in mammalian HEK293^T cells and purified using nickel affinity chromatography (Figure S6A). A mammalian expression system was chosen because it allows an efficient post-translational processing in the secretory pathway that is vital for the correct folding of antibodies and its fragments

(Aricescu *et al.*, 2006). The bioactivity of Mannitou Fab was verified prior to STD-NMR experiments by means of SPR that followed complex formation of the Fab with immobilized ω -1 M3GN2 (Figure S6B). The observed fast *on*- and *off*-rates indicated it to be in the right range of affinity for STD-NMR measurements to be feasible.





¹H-STD experiments for Mannitou Fab (25 μ M) with glycan epitope (1 mM) (**A**) Man₃GlcNAc₂ and (**B**) Man₃GlcNAc₂Fuc. In (**A**), protons of the glycan giving STD signals are labeled (black). Stars (*) indicate potentially overlapping protons from the α 1-3- and α 1-6-linked mannose. Protons that do not give any STD (non-overlapping) signals are labelled (grey) in the reference spectrum. A 3D-structural model of the interaction regions (indicated by thick grey lines) of M3GN2F1(6) was derived from the STDDD spectrum (top) and the off-resonance spectrum (bottom). It can be seen that the STDDD spectra in (**A**) and (**B**), respectively without and with the α 1-6 core fucose, are almost identical.

¹H-STD NMR experiments were performed using the same two PMGs selected from the microarray screening (Figure 6). The STD spectra at the aliphatic and the aromatic irradiation frequencies for the non-fucosylated M3GN2 (**GL41**) (Figure 6A) and fucosylated M3GN2F1(6) (**GL70**) (Figure 6B) were essentially the same. All protons belonging to the α 1-3-linked mannose gave STD signals, where H1 and H2 were the protons with the strongest STD effect. The same protons from the mannose on the α 1,6 arm demonstrated no STD effect. In addition, there was a very weak STD effect of H2 of the central β Man residue and a weak STD signal of the acetyl (Ac) group on the GlcNAc1 residue (Figure 6).

In the more crowded region of 3.6 - 3.9 ppm, an STD effect is also observed for H3, H4 and H5 protons of the aMan residue. These protons resonate at the same frequency for a1,6 and a1,3-linked mannose, however, the above-mentioned results involving H1 and H2 suggest that they correspond to the a1,3-linked mannose. The protons of GlcNAc2 exhibited no STD effect, in contrast to the protons on the acetyl (Ac) group of the reducing end GlcNAc (GlcNAc1). This probably results from the conformation of the PMG in the protein-bound form, where the acetyl group of GlcNAc1 points to the same side as the a1,3-linked mannose, while the one of GlcNAc2 points in the opposite direction (Figure 6).

A molecular model for Mannitou Fab in complex with its minimal epitope

Molecular docking simulations indicate that the M3GN2 (GL41) PMG epitope occupies a binding site shaped by Complementary Determining Region (CDR) loops as predicted using the Paratome program (Kunik *et al.*, 2012) on the sequence of Mannitou Fab. The amino acid sequences of the purified IgM were for the large majority covered using MALDI-TOF MS peptide fingerprinting within a 1% precision m/z rate and confirmed the full presence of the heavy and light chains of Mannitou IgM (Figure S7). Prediction of the binding mode of M3GN2 onto the homology model of Mannitou Fab shows the α 1-3-linked mannose inserted in a snug pocket on the protein surface, between Trp33 and Tyr104 respectively belonging to the heavy chain CDR1 and CDR3 (Figure 7). Running a docking simulation with the α 1-6 corefucosylated PMG, M3GN2F1(6) (GL70), does not change this pattern of binding.

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However, as soon as an additional residue is substituted onto the α 1-3-linked mannose, this interaction with Mannitou Fab is impeded. This is congruent with what was observed in the microarray study and is presumably due to the complementary shape of the cavity, where a substitution on the axial 2-position of the α 1-3-linked mannose of M3GN2 may force it to leave its snug pocket.



Figure 7. Positioning of M3GN2 into the Mannitou Fab homology model

Molecular docking between the Mannitou Fab 3D-model and Man₃GlcNAc₂ demonstrates the docking of the α1-3 mannose into a snug pocket. Residues are colored according to the CDR they belong to and identically as depicted on the sequenced IgM (Figure S7). Molecular presentation prepared using The PyMOL Molecular Graphics System, Version 2.0, from Schrödinger, LLC.

Discussion

The biological importance of glycosylation in physiological and pathophysiological conditions is broadly acknowledged. Human protein paucimannosylation characterization was pioneered by the group of Nicola Parker, who discovered an unconventional form of α - and β -mannose epitope-rich human Nglycosylation, on proteins in sputum from inflamed and bacteria-infected individuals (Thaysen-Andersen et al., 2015; Venkatakrishnan et al., 2015). Namely neutrophil

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elastase, proteinase 3 and cathepsin G had been reported earlier to be abundantly enriched in paucimannose on the plasma membrane of human neutrophils upon their activation (Campbell and Owen, 2007) and had been held responsible of driving cluster formation on the neutrophil cell surface (Hajjar *et al.*, 2008). Mammalian PMP biogenesis has a hexosaminidase-dependent pathway, where an N-acetyl- β hexosaminidase trims immature complex *N*-glycans retroactively down to PMGs (Tjondro *et al.*, 2019). This same strategy we have applied in the chemo-enzymatic synthesis of M3GN2F (Figure S3).

Human PMG products of the hexosaminidase activity were recently characterized by quantitative proteomics (Chatterjee et al., 2019). The thus truncated α -mannose-terminating structures spanned the monosaccharide compositions of [Man₁₋₃GlcNAc₂ Fuc₀₋₁] on human paucimannose-carrying proteins (PMPs), with a prevalence of one of the two dimannose isomers, namely PMG was not printed on the glycan array (Figure S1) and the latter only contains the dimannose Mano1-6Man with a β 1,2-linked xylose substitution on the central β Man (GL1) (Figure S1A). Dimannose GL1 with a1,6 mannose residues, but lacking a1,3 residues, does not show any binding to Mannitou IgM, while GL4, with a GlcNAc extension on the a1.6 Man arm is tolerated and demonstrates binding to Mannitou (Figure S2). The binding of the GL1 glycan can be directly compared with the Mana1-3Man dimannose equivalent, GL2. GL2, GL8 and GL40 are all Mana1-3Man dimannose PMGs with similar weak binding signals, whereas the M3GN2 structures GL7 and GL41 showed strong binding to Mannitou IgM, independent of the presence of β 1,2-linked xylose on the central mannose. For all these reasons, the absence of signal for the α 1-6-linked dimannose PMG (**GL1**) was interpreted to be due to nonbinding of the α 1,6-arm and this *N*-glycan was not included in subsequent analyses. Among other PMGs found in human paucimannosylation, M1GN2 (GL39) displaying a single mannose residue, did also not bind Mannitou IgM (Figure S2A).

Mannitou only displays significant affinities from oligomannose-3 on, represented by the minimal epitope M3GN2. Measured affinities are near 100 nM (K_d) both in glycan array and SPR experiments (Figure 1B and Table II). ¹H-STD NMR experiments were performed using two PMGs selected from the microarray screening.

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Non-fucosylated M3GN2 (**GL41**) (Figure 6A) and fucosylated M3GN2F1(6) (**GL70**) (Figure 6B), PMG epitopes for Mannitou Fab, displayed very similar STDDD spectra in which the same protons showed analogous ¹H-STD NMR effects. The two epitopes are thus estimated the same. What is more, the intensities of their STD signals were the same, which indicates that their interactions are also the same in terms of affinity. Likewise, when printed on the microarray (Figure 1), immobilized on the SPR sensor chip (Table I) or presented on one of the omega-1 glycosylation sites (Table II), there appears no significant difference in affinity between the two PMGs (Figure 5). That suggests that no hindrance, nor significant contribution to affinity, is made by the fucose α 1,6-linked to the core GlcNAc1 (Figure 6). It has been described that core fucose may lead to a different presentation of the *N*-glycan and introduce dynamics in the glycan chain (Sakae *et al.*, 2017). Yet, the presence of the α 1-6-linked core fucose does not hinder binding to Mannitou, which is a positive point for the use of the antibody in diagnosis of PMGs and PMPs, as M3GN2 and M3GN2F1(6) were both found to be very important PMGs in human cancer samples (Chatterjee *et al.*, 2019).

STD-NMR spectroscopy revealed that the protons strongest implicated in interactions with Mannitou were H1 and H2 on the α 1,3-linked mannose, with weaker signals from the other protons of the same arm, from the H2 proton of the central βMan and from the acetyl group of GlcNAc1 at the reducing end of the N-glycan (Figure 6). Interestingly, the STDDD spectra at the aliphatic and the aromatic irradiation frequencies were essentially identical. Coherent with the ¹H-STD NMR measurements, molecular simulations indicate that Mana1-3 might dock into a pocket, or hydrophobic groove, formed by aromatic residues of the predicted CDR loops of the Mannitou Fab (Figure 7). These conclusions are fully in line with the results from the microarray and from the docking, which showed that any substitution at the 2position of α Man on the α 1,3 Man arm, either with Man α 1-2 (**GL42**) or with GlcNAc β 1-2 (GL55), abolishes binding (Figure 1A). Once the requirement of the unsubstituted α 1,3 Man arm is fulfilled, the substitution of the α 1,6 Man arm showed an important permissiveness in the structure (GL23, GL27, GL34, Figure S2). Additionally, different non-mammalian substitutions such as core Xylβ1-2 (G2, G7, Figure S2), core Fucα1-3 (G72, G90, Figure S2) and core-bis fucosylation (GL10, GL75, Figure S2) did not impede binding. As these structures are not part of the mammalian glycome, they were not considered for further studies, but this data could open new applications of Mannitou antibody in invertebrate and parasitic glycomic studies.

Among the high-mannose structures printed, only Man5 N-glycan (GL43, Figure 1A) and core fucose substituted Man5 *N*-glycans (**GL74**, **GL134** in Figure S2B) were recognized by the antibody efficiently. Considering these results, there is a quantitative difference in recognition between our screening and previously published data (Zipser et al., 2012), where on the glycan array from the Consortium for Functional Glycomics a small fluorescence signal was detected for pentamannose (Man5, without the chitobiose core) and only a very minor fluorescence signal for M5GN2. The Man5 N-glycan M5GN2 (GL43) was therefore previously described to have only a residual binding towards Mannitou antibody, while in our data it was recognized at least as efficiently as M3GN2 (GL41). For the same PMGs presented on a glycoprotein, omega-1 M5GN2 & Lex, a similar affinity was observed as for omega-1 M3GN2 (Table II, Figure 5). Glycan densities may play an important role in the binding of the multivalent IgM, and we also observed this effect using SPR in a significantly higher affinity for PMGs, that were much more densely immobilized, than for PMP binding (Figure 5). It is perhaps not so surprising for carbohydrate-binding proteins specific for the α 1,3 Man arm of the N-glycan core structure to have a tolerance for substitutions on the α 1,6 Man arm. It is not unusual either, for instance a similar non-distinction between M3GN2 and M5GN2 has been found for the FimH lectin from Escherichia coli (Bouckaert et al., 2006). In contrast, the kappa-5 glycoprotein carrying predominantly M7GN2 and M8GN2 (Figure S5) did not bind Mannitou IgM and for this reason it was used as the baseline reference in the SPR experiments (Figure 4). In conclusion, our results indicate that Mannitou antibody has submicromolar affinities also for oligomannose-5 N-glycans, that are generally not called paucimannosidic.

Alpha-mannose determinants are relative rare glycan-epitopes in physiological extracellular environments (Dahmen *et al.*, 2015), but may be actively secreted or leaked from cells to transmit potent signals when required (Thaysen-Andersen *et al.*,

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2015; Loke *et al.*, 2017). Our understanding of the role of these signals in mounting an effective immune response during physiological and pathophysiological conditions is rapidly advancing. The functional implications of Mannitou IgM binding M3GN2 and M5GN2 epitopes will likewise need to be further addressed. In conclusion, our multidisciplinary approach of the molecular recognition of paucimannose *N*-glycans by Mannitou antibody by glycan microarrays, SPR, STD-NMR, modelling and docking allowed to pinpoint the minimal epitope-determining regions of the glycan and shed light on the paratope of Mannitou. Our study highlights the rational for the "forbidden" substitution on the α 1,3-linked mannose branch of the *N*-glycan pentasaccharide core Man3GlcNAc2. Gaining a better structural insight into the selectivity of the monoclonal antibody warrants to improve its diagnostic qualities in specifically targeting distinct types of paucimannosylation.

Materials and Methods

Gene sequencing of Mannitou IgM

The genomic sequences of the variable domains (VL and VH) of Mannitou antibody were determined by Fusion Antibodies Ltd (Belfast, Northern Ireland) by mRNA extraction from hybridoma cells (Laz6-189/Mannitou mAb available from the DSHB, deposited by B. Zipser) and cDNA synthesis by RT-PCR. The positive PCR products were identified by agarose gel electrophoresis and sequenced on an ABI3130xl Genetic Analyzer (Applied Biosystems).

Mannitou IgM expression, purification and variable region sequence analysis

In order to produce Mannitou IgM, hybridoma cells were cultured in serum-free medium (SFM) for three weeks and allow to secrete mAb in the medium. Subsequently, the supernatant was recovered and the Mannitou IgM was purified using a HiLoadTM 16-600 SuperoseTM 6 prep grade size exclusion chromatography column on an ÄKTA Pure (Cytiva). The running buffer consisted of 0.2 M sodium bicarbonate at pH = 8 and 0.5 M NaCl. The separation was carried out at a maximum flow rate of 0.5 ml/min.

Peptide mass fingerprinting using MALDI-TOF was performed upon in-gel digestion of Mannitou IgM using trypsin. The protein bands for heavy and light chain were excised and subjected to an in-gel digestion protocol consisting of dithiothreitol reduction at 56 °C (10 mM, 20 min), followed by iodoacetamide alkylation (55 mM, 20 min in a dark environment) and finally trypsin incubation (12.5 μ g/ μ L) overnight at 37 °C. After digestion, peptides were extracted with ammonium bicarbonate (25 mM) and trifluoroacetic acid (0.1% in water). The sequences for the variable domains were uploaded in the data base together with the constant regions (UniProtKB P01872 or entry IGHM_MOUSE for the heavy constant μ chain, P01837 or entry IGKC_MOUSE for the light κ constant domain) to enable proteomics analysis of the peptide mass fingerprinting MALDI-TOF spectrometric data. The Complementarity Determining Regions or CDRs were predicted using the program Paratome (Kunik *et al.*, 2012) and glycosylation was predicted based on programs NetNGlyc (Gupta and Brunak, 2002) and NetOGlyc (Steentoft *et al.*, 2013).

Microarray preparation and screening with Mannitou IgM and plant lectins

The glycan microarrays were prepared as described earlier (Brzezicka et al., 2015). Briefly, 50 μ M ligand solutions (1.25 nL, 5 drops, 250 pL drop volume) in sodium phosphate buffer (300 mM, 0.005% Tween-20, pH=8.4) were spatially arrayed employing a robotic non-contact piezoelectric spotter (SciFLEXARRAYER S11, Scienion) onto N-hydroxysuccinimide (NHS) activated glass slides (Nexterion H, Schott AG). After printing, the slides were placed in a 75 % humidity chamber for 18 hours at 25°C. The remaining NHS groups were quenched with 50 mM solution of ethanolamine in sodium borate buffer (50 mM, pH=9.0) for 1h. The slides were washed with PBST (PBS/0.05% Tween-20), PBS and water, then dried in a slide spinner and stored at -20°C until use. Mannitou monoclonal antibody, secreted by the hybridoma cells in the serum-free medium, was diluted to approximately 25 μ g/mL in PBS (1% BSA, 0.01% Tween-20). The antibody solution (200 μ L) was incubated on the microarrays for 1h at RT. The slides were washed with PBST and PBS. Next, they were incubated with Alexa Fluor 555 Goat Anti-Mouse IgM (1:1000) (Thermo Fischer Scientific) in PBS (1% BSA, 0.01% Tween-20) for 1 h in the dark. The microarrays were washed from the unbound secondary antibody with PBST, PBS and water. Solutions of fluorescently labelled ConA-555 (10 µg/mL) and GNA-555 (25 µg/mL) in Page 21 of 29

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TSM buffer (Tris 25 mM, 150 mM NaCl, pH=7.5) containing 2 mM CaCl₂ and 2 mM MgCl₂ were incubated in the glycan microarray at RT for 1 hour in the dark. The slides were washed with TSM buffer containing 0.01% Tween-20 and water. Slides were subsequently dried in a slide spinner dried. The fluorescence measurements were performed on Agilent G2565BA Microarray Scanner (Agilent Technologies) at 10 μ m resolution. The quantification of fluorescence was done using ProScanArray Express software (Perkin Elmer) employing an adaptive circle quantification method from 50 μ m (minimum spot diameter) to 300 μ m (maximum spot diameter). Average RFU (Relative Fluorescence Unit) values with local background subtraction of four spots and standard deviation of the mean were reported using Microsoft Excel and GraphPad Prism software.

Synthesis of M3GN2 and M3GN2F1(6) paucimannosidic glycans

M3GN2 (GL41) and M3GN2F1(6) (GL70), each figuring an amino-pentyl spacer at their reducing end (Figure S3), were prepared using a modular synthesis starting from biantennary G0, as described earlier (Serna et al., 2010). Core (a1-6) fucosylation on G0 (5.2 mg, 3.68 μ mol) was effected using the GDP-fucose (3.3 mg, 5.52 µmol) donor and the Anopheles gambiae FUT6 enzyme (Serna et al., 2013) in MES buffer (80 mM, pH=6.5) including MnCl₂ (20 mM), by stirring at RT until complete consumption of the starting material. Next, core (a1-6) fucosylated G0 was treated with β-N-acetylglucosaminidase from Streptococcus pneumoniae (New England Biolabs) in sodium acetate buffer (50 mM, pH=5.5) including CaCl₂ (5 mM) at 37°C until complete consumption of the starting material. The crude product was purified on Bond Elute carbon graphitized cartridges (Agilent) to vield 2.0 mq Man3GlcNAc(Fuc α 1-6)GlcNAc (1.76 μ mol) or a 48% yield over the two steps (Figure S3): ¹H NMR (500 MHz, D₂O) δ 5.12 (d, J = 1.7 Hz, 1H), 4.93 (d, J = 1.6 Hz, 1H), 4.91 (d, J = 4.0 Hz, 1H), 4.68 (d, J = 7.9 Hz, 1H), 4.51 (d, J = 8.1 Hz, 1H), 4.27 (d, J = 1.9 Hz)Hz, 1H), 4.14 (q, J = 6.8 Hz, 1H), 4.08 (dd, J = 3.4, 1.5 Hz, 1H), 3.98 (dd, J = 3.4, 1.7 Hz, 1H), 3.97 – 3.56 (m, 30H), 3.05 – 2.94 (m, 2H), 2.11 (s, 3H), 2.04 (s, 3H), 1.73 – 1.55 (m, 4H), 1.47 – 1.35 (m, 2H), 1.24 (d, *J* = 6.6 Hz, 3H). The HRMS (MALDI-TOF) m/z [M+Na]⁺ calculated for C₄₅H₇₉N₃NaO₃₀ 1164.4646, was found to be 1164.4664.

PMG - Mannitou IgM kinetics of binding detected using SPR

The chip for the multi-cycle analysis using Mannitou IgM was prepared by direct immobilization of Man₃GlcNAc₂ (**GL41**) and Man₃GlcNAcFuc1 (**GL70**). Man₃GlcNAc₂ was covalently immobilized via amine coupling at 237 RU (237 pg ligand/mm² sensor surface) in Fc2 and Man₃GlcNAc₂Fuc1 at 163 RU in Fc4 on a CM5 sensor chip using 10 mM NaAc at pH=4.5 as immobilization buffer. Fc1 and Fc3 were blocked immediately after activation and served as reference surfaces. The binding interaction was studied using increasing concentrations of Mannitou mAb (0 nM, 0.8 nM, 0.16 nM, 0.33 nM, 0.66 nM, 1.31 nM, 2.63 nM, 5.25 nM, 10.5 nM, 21 nM, 42 nM, 84 nM, 168 nM, 336 nM) in 200 mM NaHCO₃, 500 mM NaCl, pH=7.8 buffer, at a flow rate of 30 μ l/min at 10°C. Each sample injection (1 min contact time, 6.5 min dissociation time) was followed by a regeneration using with 50 mM NaOH.

PMP - Mannitou interactions and affinities using SPR detection

Glyco-engineered omega-1 (ω 1, UniProt Q2Y2H4) helminth glycoproteins were plantproduced in Nicotiana benthamiana and purified using cation exchange chromatography, as described previously (Wilbers et al., 2017). The sensor chip for single-cycle analysis using Mannitou IgM was prepared by immobilizing $\Delta XT/FT \omega 1$ carrying predominantly the carrying the N-glycan of choice. 10 µg/ml of glycoprotein was covalently immobilized via amine coupling in 10 mM NaOOCCH3 at pH = 4.5. M3GN2 was immobilized at 2300 RU (2300 pg PMG/mm² sensor surface) in Fc2, w1 M3GN2F1(6) at 2000 RU in Fc3, and 001 M3GN2F1(3) at 2500 RU in Fc4. For these PMPs, MALDI-TOF glycosylation profiles are available respectively from Figures 2D. 2E and 2G in a recent publication (van Noort et al., 2020). The other tested omega-1 glycoproteins w1 M5GN2 & Lex and w1 single Lex were immobilized at 3270.6 and 2604.8 RU, respectively. The w1 M5GN2 & Lex protein carries a mixture of oligomannose-5 (GL43 counterpart) and Lex (GL91 counterpart) N-glycans upon coexpression of d35S:sialFucT and d35S:sialGalT transferases, whereas the ω1 single Le^x protein was expressed using a weaker, constitutive, Gpa2 promoter to reduce sialGalT expression, resulting in an almost complete lack of hybrid Lex-type N-glycans and enabling the synthesis of relatively homogeneous N-glycans carrying a single Le^x motif (GL91 counterpart). The MALDI-TOF glycosylation profiles are available from

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respectively Figures 3A and 3C, in (Wilbers *et al.*, 2017). Engineered kappa-5 (κ 5) is another glycoprotein from *S. mansoni* (Wilbers *et al.*, 2017) that displays highmannose *N*-glycans (M7GN2/M8GN2) larger than oligomannose-5 (Figure S5), produced using kappa-5 (strain AGL) + P19 (strain AGL) with co-expression of the SmFucTD and pHYG-GPAII-DrGaIT transferases (van Noort *et al.*, 2020). Since no Mannitou IgM binding could be detected to the engineered kappa-5, even upon an immobilization rate of 3000 RU, this protein served as a blank. It was covalently immobilized in the reference flow channel Fc1 in order to block the sensor surface from a-specific binding and prepare Fc2-Fc1, Fc3-Fc1 and Fc4-Fc1 difference sensorgrams. The binding was studied using increasing concentrations of Mannitou IgM (59.375 nM, 118.75 nM, 237.5 nM, 475 nM, 950 nM) in 200 mM NaHCO₃, 500 mM NaCl, pH=7.8 buffer, at a flow rate of 30 μ l/min at 10°C. Each sample injection (3 min. association) was followed by 4 min. dissociation. All SPR data analysis was performed with BIA Evaluation Software.

Mannitou Fab expression and purification

cDNA fragments encoding the light chain (LC), and the heavy-chain variable domain (VH) with the first domain of the heavy-chain constant region, referred to as HC, were codon-optimized, synthesized and subcloned into a pUC57 E. coli expression vector (GenScript). The DNA of interest was amplified by PCR using primers HC_FP: GACTAGTACCGGTGAGGTGAAGCTTCTCGAGTCTGG, HC_RP: GACTAGTGGTACCTTAGTGATGGTGATGG, LC_FP: GACTAGTACCGGTGATGTTGTGGTGACTCAAACTCCACTC and LC RP: GACTAGTGGTACCTTAACACTCTTTCCTGTTG (FP is forward primer and RP is reverse primer). The PCR products were digested with Agel and Kpnl restriction enzymes, purified, and ligated into pHL-sec expression vectors. The HC insert was cloned between a secretion signal sequence and in frame with a C-terminal 6-histidine tag, whereas the LC fragment carried a stop codon before the Kpnl cleavage site (Aricescu et al., 2006). The expression of Mannitou Fab was achieved by transient transfection of adherent human embryonic kidney (HEK293T) cells, cultured in highglucose DMEM (Dulbecco's Modified Eagle's Medium) supplemented with Lglutamine, non-essential amino acids and 2% FBS (Thermo Fisher Scientific). Equal amounts of plasmid DNA encoding the heavy and light chains were used for transfection - 1 μ g of each vector per 1 ml of transfection volume. The DNA-PEI MAX (Transfection Grade Linear Polyethylenimine Hydrochloride, MW = 40,000 Da) solution (1:2) was incubated for 10 min at RT to allow the complex formation prior transfection. The cells were grown in expanded-surface polystyrene roller bottles (2125 cm², CELLMASTER, Greiner Bio-One) at 37 °C, in a humidified atmosphere of 5% CO₂ in air, inside a roller bottle incubator. The cell-culture supernatant was harvested 7 days post-transfection, clarified by centrifugation at 6 000 x g for 20 min, then filtered through a 0.45 µm membrane and diluted threefold with PBS. The Fab was by affinity chromatography using Ni-Sepharose 6 Fast Flow resin (Cytiva) exploiting the selective His6-tag at the *C*-terminus of the Fab HC. The binding buffer consisted of 50 mM NaH₂PO₄ at pH=8.0 and 300 mM NaCl. Two consecutive washes with ascending concentrations of imidazole (10 mM and 30 mM, pH=7.0) were performed to eliminate as many impurities as possible. Mannitou Fab was eluted with 250 mM imidazole in a buffer of 50 mM NaH₂PO₄ at pH=7.0 and 300 mM NaCl. The antibody was further purified by size-exclusion chromatography on a Superose 6 Increase 10/300 GL column (Cytiva) using buffer consisting of 20 mM 4-(2hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) at pH=7.4and 150 mM NaCl.

Epitope mapping in STD-NMR Spectroscopy

¹H-STD NMR experiments were performed at 25°C on a Bruker AVANCE III 800 MHz spectrometer equipped with a cryoprobe. Samples were prepared in PBS buffer in D₂O (pD=7.4) in 3 mm NMR tubes. The measurements were taken using a 40:1 molar ratio with a 1 mM concentration of the oligosaccharide epitope and a 25 μ M concentration of Mannitou Fab. Chemical shifts for the epitope were recorded at one off-resonance (no signals of protein or ligand) irradiation frequency set at 100 ppm and two on-resonance irradiation frequencies, one set on the aliphatic region of the protein signals (0.8 ppm) and one set on the aromatic region of the protein signals (6.9 ppm). Protein saturation was achieved with a train of Gaussian-shaped pulses of 50 ms duration, with a total irradiation time of 2 s. STD spectra were obtained by subtracting the on-resonance from the off-resonance (reference) spectrum. Subtraction of ¹H-STD NMR spectra of the free ligands at 1 mm in D₂O was also Page 25 of 29

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acquired to ensure that no direct irradiation of the oligosaccharide was taking place. Finally, because some signals were present in STD spectra from the antibody, that interfere in the analysis of the epitope protons, a second STD experiment was acquired on a sample of the protein alone. This spectrum serves as a blank that can be subtracted from the STDD spectra in the presence of the oligosaccharide epitope to obtain the final, cleaner STDDD spectra. Data acquisition and processing were performed with TopSpin 3.0 software (Bruker) and the figures were built using MestReNova v.8.0.2.

Homology modelling and molecular docking

A model of Mannitou Fab was obtained using homology modelling by supplying the crystal structures of the heavy chain of human monoclonal antibody CR4354 (1.4 Å, PDB ID: 3N9G, sequence identity: 31%) and the light chain of IGG1-K B13I2 (2.8 Å, PDB ID: 1IGF, sequence identity: 79% to the program Modeller (Sali and Blundell, 1993). Subsequently, energy minimization of the global model was done applying AMBER forcefield in order to remove putative steric clashes between the side chains. Simulation of possible binding modes of the studied paucimannosidic *N*-glycans to this three-dimensional model of Mannitou Fab was performed using GOLD docking program (Jones *et al.*, 1997). GOLD is based on a genetic algorithm and, in this case, considers the ligands as highly flexible as they are glycosidic compounds, while the side chains of most of the residues are kept rigid. For the search procedure, a sphere of 10 Å was centered on the CDRs of the variable region of the Fab. The different binding poses were scored with the ChemPLP scoring function and the pose with the best ChemPLP docking score was presented on the molecular surface of the Mannitou Fab homology model for further structural analysis.

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