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# **High-Mannose Oligosaccharide Hemimimetics that Recapitulate the Conformation and Binding Mode to Concanavalin A, DC-SIGN and Langerin**

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The "carbohydrate chemical mimicry" exhibited by  $sp^2$ -iminosugars has been utilized to develop practical syntheses for analogs of the branched high-mannose-type oligosaccharides (HMOs) Man<sub>3</sub> and Man<sub>5</sub>. In these compounds, the terminal nonreducing Man residues have been substituted with 5,6 oxomethylidenemannonojirimycin (OMJ) motifs. The resulting oligomannoside hemimimetic accurately reproduce the structure, configuration, and conformational behavior of the original mannooligosaccharides, as confirmed by NMR and computational techniques. Binding studies with mannose binding lectins, including concanavalin A, DC-SIGN, and langerin, by enzyme-linked lectin assay and surface plasmon resonance revealed significant variations in their ability to accommodate

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the OMJ unit in the mannose binding site. Intriguingly, OMJMan segments demonstrated "in line" heteromultivalent effects during binding to the three lectins. Similar to the mannobiose (Man<sub>2</sub>) branches in HMOs, the binding modes involving the external or internal monosaccharide unit at the carbohydrate binding-domain exist in equilibrium, facilitating sliding and recapture processes. This equilibrium, which influences the multivalent binding of HMOs, can be finely modulated upon incorporation of the OMJ sp<sup>2</sup>-iminosugar caps. As a proof of concept, the affinity and selectivity towards DC-SIGN and langerin were adjustable by presenting the OMJMan epitope in platforms with diverse architectures and valencies.

### **Introduction**

Calcium(II)-dependent lectin receptors (CLRs) play a crucial role in the innate immune response by exhibiting specificity towards high mannose oligosaccharides (HMOs) and mannose-containing polysaccharides.<sup>[1]</sup> They contribute to various functions, including driving immune cell migration to sites of inflammation or peripheral lymphatic nodes, as well as facilitating the endocytosis and digestion of potentially harmful pathogens.<sup>[2]</sup> Notable examples of this category of carbohydrate-binding proteins include the macrophage receptor (MR), mannose binding lectin (MBL), dectin-2, langerin, and dendritic cellspecific intercellular adhesion molecule 3-grabbing nonintegrin  $(DC-SIGN).$ [3]

These CLRs display a common group of amino acids at the  $Ca<sup>2+</sup>$  coordination site of the carbohydrate recognition domain (CRD).[4] Although they exhibit dissimilarities in the adjacent sequences of the CRD, leading to distinct secondary binding site profiles, they share affinity towards a broad range of natural and synthetic glycoligands. However, the downstream events upon binding can vary significantly. As an illustrative example, the binding of high mannose oligosaccharides (HMOs) on the surface of the human immunodeficiency virus (HIV) by DC-SIGN expressed in dendritic cells (DCs) results in immune system hijacking and facilitates virus propagation.<sup>[5]</sup> Conversely, binding to langerin in Langerhans cells (LCs) leads to HIV encapsulation, effectively halting the infection process.<sup>[6]</sup> These distinct outcomes highlight the significant functional implications of selective recognition of mannose-binding CLRs. Unsurprisingly,

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**Figure 1.** Structures of mannobiose (**1**) and 5,6-oxomethylidenemannonojirimycin (OMJ)-based mannobiose mimetics (**2**–**4**). Examples of cyclohexane- (**5**, **6**) or 1,2,3,–triazole-based (pseudo)mannooligosaccharide (**7**) derivatives are also shown. R represents 2-azidoethyl (for **2**–**4**) or Me (for **1**).

this has long been a sought-after goal in glycobiology, both for advancing fundamental studies on the immunomodulatory functions of these receptors and developing novel therapies.<sup>[7,8]</sup>

A compelling body of evidence supports  $\alpha(1,2)$ -linked mannobiose (Man $\alpha$ (1,2)Man, Man<sub>2</sub>, 1; Figure 1) as a preferred glycotope for mannose-specific CLRs.<sup>[9]</sup> Unlike other positional isomers of mannodisaccharides, both the reducing and nonreducing mannose units in Man<sub>2</sub> can bind to the CRD in the corresponding complex.[10] This unique behavior was initially observed in the binding of Man<sub>2</sub> to the plant lectin concanavalin A (ConA)<sup>[11]</sup> and represents the simplest case of a divalent "in-line" $[12]$  glycoligand/lectin interaction. The recognition event benefits from typical multivalent mechanisms, such as sliding and rebinding phenomena, resulting in significantly higher affinities than expected from a simple statistical effect.<sup>[13]</sup>

Man<sub>2</sub> is considered a heterodivalent compound since the two mannose constituents are not equivalent. When encountering a lectin partner, subtle differences in preference for one pose over the other may arise depending on the CRD architecture. In a preliminary communication, we suggested that such differences could be amplified when using  $Man\alpha(1,2)$ Man mimetics where one or both of the monosaccharide moieties are replaced by a sp<sup>2</sup>-iminosugar-type glycomimetic.<sup>[14]</sup> sp<sup>2</sup>-lminosugars are monosaccharide surrogates that, formally, replace the characteristic endocyclic oxygen with a pseudoamide-type nitrogen atom exhibiting substantial sp<sup>2</sup>-hybridization.<sup>[15]</sup> Unlike classical iminosugars, sp<sup>2</sup>iminosugars can participate in glycosylation reactions, yielding chemically stable  $\alpha$ -glycoside-like derivatives due to an enhanced anomeric effect.<sup>[16]</sup> Therefore, we synthesized compounds **2**–**4**, which comprise 5,6-oxomethylidenemannonojirimycin (OMJ) motifs, featuring a configurational pattern identical to that of α-D-mannopyranosides (Figure 2).<sup>[17]</sup> Encouragingly, ligands with complete discrimination capabilities towards DC-SIGN compared to langerin



**Figure 2.** Structures of the (pseudo)mannooligosaccharides **8**–**11**. R represents 2-azidoethyl.

were engineered by presenting the pseudodisaccharide OMJα(1,2)Man (OMJMan, **3**) in multiple copies on a β-cyclodextrin (βCD) platform.<sup>[14,18]</sup>

In principle, higher analogs of HMOs incorporating both mannose and OMJ units (referred to as HMO hemimimetics) can be synthesized by replicating the established procedures used for the parent oligosaccharides. In contrast to the utilization of cyclohexane<sup>[19]</sup> or 1,2,3-triazole platforms<sup>[20]</sup> as mannose mimetics for emulating Man units in Man<sub>2</sub> and HMO-related mannooligosaccharides (e.g., 5-7; Figure 1), the OMJ sp<sup>2</sup>iminosugar accurately reproduces the substitution arrangement of α-mannopyranosyl residues. We hypothesized that the presence of different OMJ moieties in the structure would have

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varying effects on the recognition of mannose-binding lectins within a series. (Hetero)multivalent effects, which are highly sensitive to ligand density and topological characteristics,<sup>[21]</sup> are anticipated to further contribute to lectin discernment when multiconjugation is employed, providing an additional level of customization. By combining the functional toolbox of HMO mimics based on  $sp^2$ -iminosugars with appropriate scaffolds, the opportunities for selective targeting of CLRs can be significantly expanded. To explore these concepts, we have conducted a comprehensive investigation encompassing synthetic, spectroscopic, computational, and biophysical studies. Specifically, our objectives were as follows:

- (a) Expansion of the  $sp^2$ -iminosugar-based HMO mimetics from  $Man<sub>2</sub>$  to include analogs of the branched mannooligosaccharide glycosides Man<sub>3</sub> (8) and Man<sub>5</sub> (10), namely compounds OMJ<sub>2</sub>Man (9) and (OMJMan)<sub>2</sub>Man (11), respectively (Figure 2).
- (b) Profiling the conformational properties and recognition abilities of the entire series of (pseudo)HMOs against the model lectin ConA and the human CLRs DC-SIGN and langerin.
- (c) Preparation of a diverse range of conjugates displaying the preferred mannobiose hemimimetic motif OMJMan on various platforms with different valencies, densities, and architectures, to enable probing assays for investigating lectin recognition variations.

### **Results and Discussion**

### **Synthesis and conformational properties of sp2 -iminosugarterminated mannooligosaccharide hemimimetics**

After identifying OMJMan as a privileged motif for selective targeting of DC-SIGN, our synthetic efforts have been focused on preparing hemimimetics of higher HMOs with terminal (nonreducing) OMJ units. To maintain sequence homology, we installed an  $\alpha$ -oriented 2-azidoethyl aglycone. The synthesis of the pseudotrisaccharide OMJ2Man glycoside **9** was achieved by reacting the Man dual acceptor 12<sup>[22]</sup> with the OMJ pseudoglycosyl fluoride  $13^{[23]}$  ( $\rightarrow$  14, 71%) and subsequently catalytically removing the ester protecting groups (Scheme 1). A convergent strategy was employed for the synthesis of the pseudopentasaccharide (OMJMan)<sub>2</sub>Man glycoside 11, which involved the



same monosaccharide acceptor **12** and the protected OMJMan pseudodisaccharide trichloroacetimidate **17** as the glycosyl donor ( $\rightarrow$ 18, 50%), followed by deacylation (Scheme 2). Compound **17** was synthesized in three steps starting from the Man tolylthioglycoside tribenzoate **15** [24], which included glycosylation with **13** ( $\rightarrow$  **16**, 64%), removal of the *S*-tolyl group using Nbromosuccinimide (NBS), and reaction of the crude reducing intermediate with trichloroacetonitrile/1,8diazabicyclo[5.4.0]undec-7-ene (DBU; 34%). Equivalent synthetic routes were used to access the corresponding 2-azidoethyl Man<sub>3</sub><sup>[25]</sup> and Man<sub>5</sub><sup>[26]</sup>  $\alpha$ -glycosides, which were employed as controls in structural and lectin recognition experiments. These results exemplify the versatility of  $sp^2$ -iminosugars in integrating standard glycosidation methodologies, underscoring their unique "carbohydrate chemical mimic" properties.

The ability of OMJ-containing mannooligosaccharides to replicate the conformational behavior of the parent glycans was initially assessed through Monte Carlo Multiple Minimum (MCMM)[27,28] conformational searches followed by Monte Carlo/ Stochastic Dynamics  $(MC/SD)^{[29]}$  simulations at 300 K using the OPLS3e force field implemented in Maestro 12.2 (MacroModel 12.6) software.<sup>[30]</sup> To validate the method, we first examined methyl α-(1,2)-mannobioside (Man<sub>2</sub>OMe; 1-OMe). Previously reported NMR and computational data for Man<sub>2</sub>OMe in water indicated an equilibrium between a stacked (*S*) and an extended (*E*) conformation around the glycosidic linkage, with both conformations being similarly populated.[31,32] The *S* conformation is characterized by  $\phi$  (H1'-C1'-O1'-C2) and  $\psi$  (C1'-O1'-C2-H2) dihedral angle values ranging from  $-30^{\circ}$  to  $+30^{\circ}$ and  $+30^{\circ}$  to  $+50^{\circ}$ , respectively, consistent with H-1/H-1' NOE contacts (Supporting Information, Figure S49). The less flexible *E* conformation exhibits  $\phi$  and  $\psi$  values around  $-50^{\circ}$  and  $-30^{\circ}$ to  $+30^{\circ}$ , respectively, in agreement with H-5'/H-1 and H-1'/H-3 diagnostic NOE signals. We successfully reproduced this scenario (refer to the Supplementary Information for computational details).



**Scheme 1.** Synthesis of the OMJ<sub>2</sub>Man glycoside **9**. **Scheme 2.** Synthesis of the (OMJMan)<sub>2</sub>Man glycoside **11**.

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**Figure 3.** A) Structure of the OMJMan pseudodisaccharide **3** with indication of the ϕ (H1'-C1'-O1'-C2) and ψ (C1'-O1'-C2-H2) dihedral angles defined for computational experiments. B) (ϕ,ψ) map obtained by the conformational search (MCMM/OPL3e force field). C) (ϕ,ψ) map obtained after MC/SD simulation. D) Representation of the minimum energy conformations for regions I, II and III in the (ϕ,ψ) map, with indication of the theoretical diagnostic NOE contacts.

We subsequently applied the validated molecular modeling protocol in combination with NMR spectroscopy to investigate the conformational properties of the  $sp^2$ -iminosugar-based Man<sub>2</sub> mimetics 2-4 previously synthesized. The MCMM-generated  $(\phi,\psi)$  maps exhibited similar patterns for all the pseudodisaccharides. In this discussion, we will focus on the conformational analysis of the OMJMan glycoside **3** as an illustrative case (Figure 3A), while the complete dataset is provided in the Supplementary Information (Supporting Information, Figures S50–S57 and Tables S1 and S2). Three regions (I–III) with relative energies below 20 kJmol<sup>-1</sup> were identified (Figure 3B). Region I, representing the energy minimum, and region II  $(+)$ 2.4 kJmol<sup>-1</sup>) correspond to *E*-conformers and possess  $(\phi, \psi)$ values of approximately  $(-47^\circ, +17^\circ)$  and  $(-49^\circ, -34^\circ)$ , respectively. Region III (+2.4 kJmol<sup>-1</sup>) contains *S*-conformers with (φ,ψ) values of about  $(+26^{\circ},+18^{\circ})$ . Molecular dynamics simulations at 300 K for a duration of 10 ns supported a rapid conformational equilibrium around the glycosidic linkage and revealed a greater tendency to occupy *E*-conformer regions compared to Man<sub>2</sub>OMe (Figure 3C). The predicted protonproton contacts from the MCMM analysis (Figure 3D) were corroborated by NOE experiments (Supporting Information, Figures S53 and S54), which also provided experimental interproton distances that matched well with the distances calculated through MC/SD (Table 1). Comparison of such distance with the corresponding interproton distance range reported for





the mannobioside **1**-OMe further confirmed the prevalence of the *E* conformation.

For the Man<sub>3</sub> surrogate OMJ<sub>2</sub>Man glycoside 9, which contains two glycosidic linkages, namely  $\alpha(1,3)$  and  $\alpha(1,6)$ , we defined the following dihedral angles (Figure 4A):  $\phi_{C,A}$  (N5<sup>C</sup>-C1<sup>C</sup> -O3<sup>A</sup>-C3<sup>A</sup>), ψ<sub>C,A</sub> (C1<sup>C</sup>-O3<sup>A</sup>-C3<sup>A</sup>-C2<sup>A</sup>), φ<sub>B,A</sub> (N5<sup>B</sup>-C1<sup>B</sup>-O6<sup>A</sup>-C6<sup>A</sup>), ψ<sub>B,A</sub>  $(C1^{B}-CO6^{A}-CO6^{A}-CO6^{A})$ , and  $\omega_{B,A}$  (O6<sup>A</sup>-C6<sup>A</sup>-C5<sup>A</sup>-C4<sup>A</sup>). MCMM conformational searches were performed to generate ( $\phi_{C,A}, \psi_{C,A}$ ) and  $(\phi_{B,A}, \psi_{B,A})$  maps, which revealed three regions (I–III and IV–VI, respectively; Figure 4B) with relative energies below 20  $kJ$  mol<sup>-1</sup>. To facilitate comparison, the maps for each glycosidic linkage are presented in Figure 4 alongside the corresponding maps obtained for the parent Man<sub>3</sub> glycoside 8 in a parallel analysis (Figure 4C). Region I represents the energy minimum for the α(1,3) linkage at  $(\phi_{C,A}, \psi_{C,A}) = (-46^\circ, +13^\circ)$ . Rotation around  $\psi_{C,A}$ leads to region II  $(+1.2 \text{ kJ} \text{mol}^{-1})$ , while region III is obtained by rotating around  $\phi_{C,A}$  to approximately  $+30^{\circ}$  ( $+9.6$  kJmol<sup>-1</sup>). In addition to the typical through-space coupling between the protons on the carbons directly involved in the glycosidic connection (H1<sup>c</sup> and H3<sup>A</sup>), conformers located in region III are expected to exhibit a NOE contact between  $H1<sup>c</sup>$  and H2<sup>A</sup>, which was experimentally observed for both OMJ2Man (Figure 4D) and Man<sub>3</sub>. The  $\alpha(1,6)$  glycosidic linkage is predicted to be more flexible. In contrast to the relatively confined regions I–III observed in the ( $\phi_{C,A}, \psi_{C,A}$ ) map, regions IV–VI in the ( $\phi_{B,A}, \psi_{B,A}$ ) map of OMJ<sub>2</sub>Man showed a wider distribution of the  $\psi_{BA}$  angle compared to the natural mannotrisaccharide Man<sub>3</sub> (Figure 4B) and C). However, the MC/SD simulations provided similar overall conformational tendencies. The  $\phi_{C,A}$  and  $\phi_{B,A}$  angles remained close to  $-50^{\circ}$  for the majority of the simulation time, and the  $\alpha(1,3)$  and  $\alpha(1,6)$  linkages predominantly adopted conformations within regions I, II, and IV, V, respectively. A minor population of non-exoanomeric conformations, characterized by positive values of  $\phi$  and  $\psi$ , was also observed for both glycosidic linkages (Figure 4E and F).

The torsion angle around C5-C6 predominantly adopted two conformations (Figure 5A and B): *gauche-trans* (*gt*;  $\omega_{BA} = \pm$ 

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**Figure 4.** A) Structure of the OMJ<sub>2</sub>Man glycoside **9** with indication of the  $\phi_{\sf CA}$  (N5<sup>C</sup>-C1<sup>c</sup>-O3<sup>A</sup>-C3<sup>A</sup>),  $\psi_{\sf CA}$  (C1<sup>C</sup>-O3<sup>A</sup>-C3<sup>A</sup>), $\phi_{\sf BA}$  (N5<sup>B</sup>-C1<sup>B</sup>-O6<sup>A</sup>-C6<sup>A</sup>),  $\psi_{\sf BA}$  (C1<sup>B</sup>-O6<sup>A</sup>-C4<sup>B</sup>), C6<sup>A</sup>-C5<sup>A</sup>), and ω<sub>B,A</sub> (O6<sup>A</sup>-C6<sup>A</sup>-C5<sup>A</sup>-C4<sup>A</sup>) dihedral angles defined for computational experiments. B and C) (ϕ,ψ) maps obtained by the conformational search (MCMM/OPL3e force field) for the OMJ<sub>2</sub>Man and Man<sub>3</sub> (pseudo)glycosides 9 and 8, respectively. D) Expansion of the NOESY spectrum for the OMJ<sub>2</sub>Man glycoside **9**. E and F) (ϕ,ψ) maps obtained by MC/SD simulation for the OMJ2Man and Man3 (pseudo)glycosides **9** and **8**, respectively.



**Figure** 5. A and B) Evolution of the dihedral angle  $ω_{B,A}$  for the OMJ<sub>2</sub>Man and Man<sub>3</sub> (pseudo)glycosides 9 and 8, respectively, during the MC/SD simulation. C) (ψC,A,ψB,A,ωB,A) map obtained after MC/SD/MM for the OMJ2Man pseudoglycoside **9** and representation of the minimum energy conformations for clusters A, B and C. Energy values relative to the absolute minimum are given in parenthesis.

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180°) and *gauche-gauche* (*gg*;  $\omega_{BA}$ =60°). Based on the MC/SD data followed by multiple minimization (MC/SD/MM), a 3D ( $\Psi_{C.A}$ , $\Psi_{B.A}$ , $\omega_{B.A}$ ) map was constructed (Figure 5C), revealing conformers that could be classified into three clusters (A–C). Clusters A and B mainly consisted of *gt* conformers, with cluster A being the most energetically favorable (0 kJ mol<sup>-1</sup>) and cluster B slightly higher in energy (4.2 kJ mol<sup>-1</sup>). Cluster C comprised gg conformers, which were energetically less favored (5.1  $kJ$ mol<sup>-1</sup>).

The experimental interproton distances  $H1<sup>C</sup>/H3<sup>A</sup>$  and  $H1<sup>B</sup>/H3<sup>B</sup>$ H6a<sup>A</sup>, determined from NOESY experiments, exhibited good agreement with the calculated distances from either MC/SD or MS/SD/MM. However, a slight discrepancy was observed for the latter, which can be attributed to the high flexibility of the  $\alpha(1,6)$  glycosidic linkage (Table 2 and Supporting Information, Figures S58–S61 and Table S3).

The conformational behavior of the Man5 glycoside **10** can be rationalized as a combination of the previously discussed properties of the disaccharide derivative **1**-OMe concerning the two  $\alpha(1,2)$ -linked terminal mannobiose moieties, and the Man<sub>3</sub> glycoside **8**, with respect to the internal  $\alpha(1,3)$  and  $\alpha(1,6)$ glycosidic linkages. The MCMM search revealed that the linear disaccharide segments can adopt both *S* and *E* conformations. In contrast, the branched portion displayed greater flexibility, with the  $\phi$  angles oscillating around  $-50^{\circ}$  and the  $\psi$  torsions exhibiting positive and negative values. The MC/SD results supported the presence of an equilibrium among these conformations (Supporting Information, Figure S62). The hemimimetic (OMJMan)<sub>2</sub>Man glycoside 11 exhibited a similar pattern, with slightly increased flexibility in the  $\phi$  dihedral angles, resulting in a small yet significant population of nonexoanomeric conformations, as observed in the pseudotrisaccharide OMJ2Man **9**. The MC/SD simulations further confirmed the presence of rapid conformational equilibria for the glycosidic linkages, specifically the  $\omega$  torsion of the  $\alpha(1,6)$  linkage shifting between the *gg* and *gt* orientations (Figure 6).

The quantitative analysis of interproton distances obtained from ROESY experiments corroborated the calculated distances and emphasized the similarity between the conformational properties of the Man<sub>5</sub> and (OMJMan)<sub>2</sub>Man moieties in glyco-



sides **10** and **11**, respectively (Table 3 and Supporting Information, Figures S63, S64, S66 and S67). Upon MC/SD/MM minimization, the conformers of the Man<sub>5</sub> derivative 10 could be classified into five clusters (A–E; Figure 7A and Supporting Information, Figure S65), while (OMJMan)<sub>2</sub>Man derivative 11 conformers formed three clusters (A–C; Figure 7B and Supporting Information, Figure S68). Cluster A represented the global energy minimum for both compounds and exhibited very similar characteristics, with *gt* orientations around the α(1,6) ω torsion (Figure 6C and D). Conformers with *gg* orientations possessed significantly higher energies. All Man<sub>5</sub> (10) conformers adopted the *S* conformation at the mannobiose unit on the  $\alpha(1,3)$ -linked arm, while the  $\alpha(1,6)$ -linked arm displayed a mixture of *S* and *E* conformations. However, in the case of (OMJMan)<sub>2</sub>Man (11), the OMJMan residue on the  $\alpha(1,3)$ -linked arm assumed the *S* conformation exclusively in the global minimum A.

Collectively, the computational and NMR data discussed above highlight that the substitution of peripheral  $\alpha$ -Dmannopyranoside units with sp<sup>2</sup>-iminosugar motifs in Man<sub>2</sub>,  $Man<sub>3</sub>$ , and  $Man<sub>5</sub>$  does not significantly alter their conformational properties. Although these hemimimetics exhibit slightly higher conformational flexibility, the relative distribution of conformers and their corresponding equilibria remain quite similar, establishing them as true conformational mimics.

### **Comparative evaluation of the ConA binding abilities of HMO hemimimetics by enzyme-linked lectin assay (ELLA)**

The above commented studies have shown that all accessible conformations of the natural mannooligosaccharides Man<sub>2</sub>,  $Man_{3}$ , and  $Man_{5}$  are equally accessible for the synthesized OMJbased analogues. Therefore, it is anticipated that the analogues can adopt the necessary conformation for binding to lectin receptors that recognize canonical HMO-related partners. The configurational pattern and the secondary hydroxyl group profile are identical. However, there are notable structural differences in the primary position region between Man and OMJ units, such as the anchor of the C5-C6 dihedral angle in the *gt* conformation due to the presence of a five-membered carbamate ring and the absence of the 6-OH, which is replaced by a carbonyl group. These modifications may modulate the interactions with amino acids at the primary and/or secondary carbohydrate binding sites in lectin partners.



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Figure 6. A) Structure of the (OMJMan)<sub>2</sub>Man derivative 11 with indication of the intermonosaccharide connectivity. (ϕ,ψ) maps obtained by the conformational search (MCMM/OPLS3e force field). B) ( $φ,ψ$ ) map obtained by MC/SD simulation. C) Evolution of the dihedral angle  $ω_{B,A}$  during the MC/SD simulation.



**Figure 7.** A and B) ( $\psi_{CA}$ , $\psi_{BA}$ , $\omega_{BA}$ ) maps obtained by MC/SD/MM calculations for the Man<sub>5</sub> and (OMJMan)<sub>2</sub>Man (pseudo)glycosides **10** and **11**, respectively. C and D) Representation of the absolute minimum energy conformations (clusters A) from the above ( $\psi_{C,A},\psi_{B,A},\omega_{B,A}$ ) maps for **10** and **11**.

ConA, an extensively studied plant lectin, has played a significant role in advancing our understanding of multivalency. Many concepts derived from studies on multimannoside binding to ConA can be extrapolated to mannose-specific lectins in animals, despite their structural differences. This highlights the universal mechanisms underlying affinity enhancement through the multivalent effect. Notably, ConA, like DC-SIGN and langerin, can bind to Man<sub>2</sub> motifs in two distinct modes, positioning either the nonreducing or the reducing unit at the CRD. In this study, the enzyme-linked lectin assay (ELLA)<sup>[33]</sup> was employed to investigate the impact of replacing mannose units with OMJ motifs on complex formation with ConA. ELLA determines the ability of the tested glycoligand to compete with a reference ligand, which is immobilized on the microplate well (in this case, yeast mannan), for binding to the target lectin, which is labelled with an enzyme that catalyzes a reaction providing a suitable readout. In our case, we used horseradish peroxidase-labelled ConA (HRP-ConA). The use of horseradish peroxidase as a label mitigates crosslinking phenomena due to its large size. Therefore, ELLA is expected to provide information on the interaction with a single carbohydrate binding site in the lectin, eliminating potential aggregation effects.[34]

Figure 8 presents the ELLA plots and the corresponding  $IC_{50}$ values obtained for the series of (pseudo)di-, tri-, and pentasaccharides against HRP-ConA included in this work. It is evident that ConA does not recognize the pseudodisaccharide composed exclusively of the sp<sup>2</sup>-iminosugar, namely  $OMJ_2$ . Indeed, a control experiment using the monosaccharide analog OMJ confirmed that this motif is not a ligand for ConA (data not shown). This finding aligns with the established understanding of the minimal structural requirements for ConA binding, which necessitate the presence of the diequatorial OH3/OH4 hydroxyls and the primary OH6 hydroxyl.<sup>[35]</sup> Surprisingly, the hemimimetic OMJMan demonstrated significantly enhanced ConA ligand efficacy compared to the monovalent reference, ManOMe, approaching the binding efficiency of Man<sub>2</sub>. In other words, the nonreducing OMJ unit reinforces the recognition of the reducing Man unit. However, docking experiments did not reveal favorable secondary interactions between the  $sp^2$ iminosugar and the neighboring amino acids within the carbohydrate recognition domain (CRD) (Supporting Information, Figure S82). One possible explanation is that once the Man moiety is situated at the CRD, rapid exchange with the OMJ unit at the protein surface becomes facilitated, even if the latter is a very poor ligand. This "reduction of dimensionality" mechanism has also been proposed to operate in heteromultivalent biological systems, leading to an increase in the half-life of the bound state.<sup>[18,36]</sup>

In ELLA conditions, the Man<sub>3</sub> glycoside 8, for which ConA possesses an extended binding site,<sup>[37]</sup> exhibits 20-fold higher HRP-ConA binding affinity than ManOMe. However, binding is completely abolished for the hemimimetic OMJ<sub>2</sub>Man glycoside derivative **9**, which lacks the primary hydroxyls. In sharp contrast, the Man<sub>5</sub> and (OMJMan)<sub>2</sub>Man glycosides 10 and 11 demonstrate similar behavior in ELLA, displaying approximately 100-fold higher potency than the Man<sub>2</sub> and OMJMan glycosides **1** and **3**, respectively. This consistency can be attributed to the divalent presentation of the (pseudo)mannobiose motif. In other words, both the  $\alpha(1,3)$ - and  $\alpha(1,6)$ -linked arms in Man<sub>5</sub> and  $(OMJMan)_2$ Man maintain the  $(hetero)$ divalent "in line" ligand character. Molecular docking experiments conducted on the Man<sub>5</sub> glycoside **10**/ConA complex support the notion that a nonreducing Man unit is positioned at the CRD, preferably the one on the  $\alpha(1,3)$ -linked arm. A similar situation is observed for the (OMJMan)<sub>2</sub>Man glycoside 11 (Supporting Information, Figure S83). These findings align with the crystal structure reported for the corresponding complex with the N-acetylglucosamineterminated pentasaccharide  $\beta$ -GlcNAc-(1  $\rightarrow$  2)- $\alpha$ -Man-(1  $\rightarrow$  3)-[β-GlcNAc-(1- $\rightarrow$ 2)- $\alpha$ -Man-(1- $\rightarrow$ 6)]-Man (GlcNAc<sub>2</sub>Man<sub>3</sub>).<sup>[38]</sup>

### **Comparative evaluation of DC-SIGN and langerin binding to**  $Man_3/OMJ_2Man$  and  $Man_5/(OMJMan)_2Man$ **(pseudo)oligosaccharides by NMR, molecular docking and surface plasmon resonance (SPR)**

In the context of the interaction between the  $Man_2$  or OMJMan glycosides **1** or **3** and DC-SIGN and langerin, previous studies utilizing NMR transfer techniques, including transferred-NOESY (tr-NOESY) and saturation-transfer difference (STD) NMR, and computational techniques have revealed that the CRD of these lectins can be occupied by either the nonreducing Man/OMJ unit or the reducing Man unit. Binding involves coordination to the  $Ca^{2+}$  cofactor through two adjacent hydroxyl groups in a trans diequatorial disposition and, unlike ConA, the presence of the primary OH6 hydroxyl is not essential for efficient binding. DC-SIGN exhibits a preference for the binding mode involving the reducing monosaccharide, while langerin favors the opposite pose.[14]

Analysis of the tr-NOESY spectra for Man<sub>3</sub> glycoside 8 in the presence of DC-SIGN and langerin revealed the characteristic sign change of cross-peaks, indicating the transient formation of the high molecular weight complex and therefore the interaction with the protein (Supporting Information, Figure S69A). This phenomenon was also observed for the OMJ2Man glycoside **9**/DC-SIGN pair (Supporting Information, Figure S69B). However, in the case of **9** in the presence of langerin, only a subset of cross-peaks exhibited sign changes. This suggests a weaker interaction that involves a smaller area of the pseudotrisaccharide molecule accompanied by a variation on local flexibility (Supporting Information, Figure S69B). Binding of both **8** and **9** to DC-SIGN was confirmed by STD experiments (Figure 9A and C and Supporting Information, Figures S71 and S72). Due to extensive signal overlapping, it



**Figure 8.** ELLA plots for the inhibition of HRP-ConA to yeast mannan by compounds **1**–**4** and **8**–**11**, with indication of the corresponding IC50 values and relative potencies (RP; referred to ManOMe). Data for methyl α-D-mannopyranoside (ManOMe) are included as a monovalent Man reference.

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Figure 9. A and B) STD plots for the Man<sub>3</sub> glycoside 8 faced to DC-SIGN and langerin, respectively. C and D) STD plots for the OMJ<sub>2</sub>Man pseudoglycoside 9 faced to DC-SIGN and langerin, respectively. The structures of the ligands, with indication of the normalized STD (%; color scale) derived from the initial slopes, are shown. Protons that are observed in the STD spectra but cannot be integrated due to overlapping are labelled with an asterisk.

was challenging to map the complete epitopes; however, the data indicate that the nonreducing residues are primarily involved in the recognition process. Transfer of saturation was found to be larger for the  $\alpha(1,3)$ -linked residue in both cases, suggesting that this unit interacts more favorably with the lectin. Nevertheless, the existence of a dynamic equilibrium between two alternative binding modes, with the  $\alpha(1,3)$ - or  $\alpha(1,6)$ -linked Man or OMJ moiety coordinating the Ca<sup>2+</sup> cation through OH3 and OH4, cannot be ruled out. In the case of STD experiments with langerin, saturation values of the ligand protons were significantly lower compared to the analogous experiments with DC-SIGN (Figure 9B and D). Nonetheless, the data are consistent with the existence of a similar equilibrium between the two binding modes.

The examination of supramolecular complex structures obtained through molecular docking, which are consistent with experimental NOE contacts in the bound state, provides insights to rationalize the observations. In the most favorable binding pose, Man3 glycoside **8** binds to DC-SIGN in a pincer-like fashion, where both nonreducing residues interact simultaneously with the protein, while the branched reducing residue does not significantly participate in binding. The  $\alpha$ (1,3)-linked residue occupies the metal coordination site, whereas the  $\alpha(1,6)$ -linked residue establishes hydrogen bonds and van der Waals interactions with neighboring amino acids, such as Asn344, Arg345, and Asn362 (Figure 10A). This binding mode lines up with previous reports on  $Man<sub>3</sub>/DC-SIGN$  interactions<sup>[39]</sup> and is supported by crystal structures of the GlcNAc<sub>2</sub>Man<sub>3</sub>/DC- $SIGN<sup>[40]</sup>$  and  $Man<sub>4</sub>/DC-SIGN$  complexes.<sup>[41]</sup> A similar binding mode is observed for the trisaccharide mimetic OMJ<sub>2</sub>Man glycoside **9**, although the interactions of the α(1,6)-linked

residue with Arg345, which involve the primary hydroxyl in Man<sub>3</sub>, are not observed (Figure 10B). In contrast, when considering langerin, the docking results suggest binding modes for **8** and 9 that resemble those reported for  $Man_{2r}^{[42]}$  with only one of the nonreducing residues occupying the CRD. Additional interactions at secondary sites do not contribute to binding in either case (Figure 10C and D).

tr-NOESY spectra for the free Man<sub>5</sub> and (OMJMan)<sub>2</sub>Man glycosides **10** and **11** exhibited cross-peaks with the same sign as those in the diagonal, which is compatible with large molecules. In addition, the NOESY peaks in presence of lectin remained unchanged compared with the free ligands, consequently we were unable to provide qualitative information on the formation of lectin complexes (see the Supporting Information for a detailed description of the NOESY NMR experiments). In the case of the **10**/DC-SIGN pair, STD results demonstrated significant saturation of all protons in the two mannobiose antennae, supporting complex formation (Figure 11A). Although some signals experienced overlapping, it was evident that the intensity of the STD signals from the internal Man residues, particularly the  $\alpha(1,3)$ -linked residue, was higher. The best binding pose obtained through molecular docking closely resembled that reported for the GlcNAc<sub>2</sub>Man<sub>3</sub>/ DC-SIGN complex,<sup>[40]</sup> with the  $\alpha(1,3)$ -linked Man coordinated to the metal center through OH3 and OH4. The external Man unit in this branch was accommodated on Val351, similar to the binding observed for Man<sub>2</sub><sup>[42]</sup> while the  $\alpha(1,6)$ -linked mannobiose segment oriented towards Phe313 and Ser360, establishing several van der Waals and hydrogen bond contacts (Figure 11B and Supporting Information, Figure S73). The corresponding STD and molecular docking experiments for the



**Figure 10.** Predicted binding mode for ligands A) **8** and B) **9** to DC-SIGN (coordinates from PDB 1 K9I) obtained by docking. Predicted binding mode for ligands C) 8 and D) 9 to langerin (coordinates from PDB 3P5F) obtained by docking, superimposed with the crystal structure of the Man<sub>2</sub>/langerin complex<sup>[42]</sup> (carbons are represented in green for Man<sub>2</sub> and in lilac or light blue for 9).



Figure 11. A) STD growth plots for the Man<sub>5</sub> glycoside 10 faced to DC-SIGN and epitope map. B) Predicted binding mode for ligand 10 to DC-SIGN (coordinates from PDB 2IT5) obtained by docking. C) STD growth plots for the (OMJMan)2Man pseudopentasaccharide **11** faced to DC-SIGN and epitope map. D) Predicted binding mode for ligand **11** to DC-SIGN (coordinates from PDB 2IT5) obtained by docking. The structures of the ligands, with indication of the normalized STD (%, color scale) derived from the initial slopes, are shown. Protons that are observed in the STD spectra but cannot be integrated due to overlapping are labelled with an asterisk.

(OMJMan)2Man pseudopentasaccharide **11** revealed a nearly identical binding mode to DC-SIGN, indicating that the  $sp^2$ - iminosugar hemimimetic mirrors the natural HMO **10** (Figure 11C and D and Supporting Information, Figure S74).



Figure 12. A) STD growth plots for the Man<sub>5</sub> glycoside 10 faced to langerin and epitope map. B) Predicted binding mode for ligand 10 to langerin (coordinates from PDB 3P5D) obtained by docking, superimposed with X-RAY Man<sub>2</sub>/langerin. C) STD growth plots for the (OMJMan)<sub>2</sub>Man pseudopentasaccharide 11 faced to langerin and epitope map. D) Predicted binding mode for ligand **11** to langerin (coordinates from PDB 3P5D) obtained by docking, superimposed with X-RAY Man<sub>2</sub>/langerin. The structures of the ligands, with indication of the normalized STD (%) derived from the initial slopes, are shown. Protons that are observed in the STD spectra but cannot be integrated due to overlapping are labelled with an asterisk.

STD experiments likewise were consistent with Man<sub>5</sub> and (OMJMan)2Man glycosides **10** and **11** binding to langerin. Unfortunately, only partial binding epitope maps could be drawn, due to substantial signal overlap (Figure 12A and C and Supporting Information, Figure S77 and S78). Since the integrable signals differ for **10** and **11**, a direct comparison can be misleading. Indeed, the <sup>1</sup>H-STD NMR spectra (Supporting Information, Figure S77 and S78) showed that the  $\alpha(1,2)$ -linked Man or OMJ units were the ones primarily involved in complex formation in both cases. This is consistent with docking calculations (Figure 12B and D). Nevertheless, participation of the internal Man units cannot be discarded. The ligands were found to exhibit less contacts with langerin as compared with DC-SIGN, which is in accord with the greater challenges faced by langerin in accommodating larger ligands.

The biomimetic properties of the hemimimetic compounds **9** and **11** were further validated through surface plasmon resonance (SPR) studies, (see the Supporting Information for the corresponding sensorgrams) which aimed to assess their competitive inhibition of DC-SIGN and langerin binding to mannosylated bovine serum albumin (Man-BSA). The reference mannooligosaccharide derivatives **8** and **10** were employed as controls in the experiments. Table 4 displays the corresponding  $IC_{50}$  values for the inhibitory activity of the compounds. In order to facilitate comparison,  $IC_{50}$  values for Man<sub>2</sub> and OMJMan glycosides **1** and **3** are also provided in the table.

In terms of DC-SIGN binding, the pseudotrisaccharide OMJ<sub>2</sub>Man glycoside **9** (IC<sub>50</sub> = 593 μM) exhibited higher affinity towards this lectin compared to the mannotrisaccharide derivative **8** ( $IC_{50}$  = 700  $µ$ M). This enhanced binding can be attributed to the increased conformational flexibility of the





former, which enables a better adaptation to the pincer-like binding mode mentioned earlier. Additionally, the dynamic interconversion between the  $\alpha(1,3)$ -linked and  $\alpha(1,6)$ -linked OMJ residues coordinating the  $Ca^{2+}$  cation may occur more readily in OMJ<sub>2</sub>Man. Concerning Man<sub>5</sub> and (OMJMan)<sub>2</sub>Man glycosides 10 and 11, their affinity trend and IC<sub>50</sub> values (209 and 265 μM, respectively) can be interpreted in terms of their divalent nature compared to Man2 and OMJMan glycosides **1** and **3** (IC<sub>50</sub> values 590 and 1060  $\mu$ M, respectively). The overall conclusion is that substituting the nonreducing Man residue with OMJ in the oligosaccharides does not significantly impact the binding properties to DC-SIGN, which aligns with the NMR and computational data.

The inhibitory potencies of the (pseudo)glycoligands **1**, **3**, **8**–**11** towards langerin, as assessed by SPR, were lower compared to DC-SIGN for all ligands tested. Notably, substituting Man with OMJ at the nonreducing ends of the ligands consistently resulted in impaired langerin binding. However, despite this detrimental effect, the observed  $IC_{50}$  trend for the hemimimetic ligands **3**, **9** and **11** closely mirrored that of the parent mannooligosaccharides **1**, **8** and **10**, indicating that (hetero)multivalent effects operate similarly in both ligand series.

The ratio between the  $IC_{50}$  values obtained for langerin and DC-SIGN provides valuable insights into the selectivity of each ligand towards these receptors. Our data suggests that as the size of the HMO-related pseudooligosaccharides increases, the selectivity of DC-SIGN over langerin decreases. Specifically, when considering the (pseudo)di- (**1** and **3**) and (pseudo)pentasaccharides (**9** and **10**), which can be regarded as mono- and divalent species of Man<sub>2</sub> (or OMJMan), an enhancement in binding efficiency is accompanied by a reduction in selectivity. No significant impact of replacing the peripheral Man units into OMJ motifs is seen.

### **Synthesis of multivalent OMJMan conjugates with varied architectures and comparative evaluation of their ConA binding abilities by ELLA**

In our previous communication,<sup>[14]</sup> we used a  $βCD$  scaffold to access the (pseudo)disaccharide multiconjugates OMJMan-βCD (20 and 22) and Man<sub>2</sub>-βCD (19 and 21) (Figure 13). In that case, the presence of OMJ instead of Man at the periphery of the heptavalent tetradecavalent glycocluster resulted in total DC-SIGN over langerin selectivity. This is sharply different to the results above discussed for the (pseudo)penta-saccharides **9** and **10**. The discrepancy can be attributed to the fact that the influence of valency and architectural features on (hetero)multivalent effects is substantial. In order to gain a

more comprehensive understanding of how these parameters differentially affect Man<sub>2</sub> and OMJMan displays binding to the lectins, we next undertook the synthesis and evaluation of an expanded library of OMJMan multivalent derivatives on platforms with varying topologies.

The new (pseudo)glycoclusters prepared in this work are depicted in Figure 14 (first generation) and Figure 15 (second generation derivatives). First generation compounds **23**–**28** were synthesized using copper(I)-catalyzed azide-alkyne coupling (CuAAC) reactions employing three different poly(alkyne) equipped scaffolds: (a) the cyclodecapeptide scaffold 37,<sup>[43]</sup> (b) the polylysine dendron **38**[43] and (c) the planar cyclophosphazene 39.<sup>[44]</sup> The multiconjugation reactions of 37–39 with Man<sub>2</sub> or OMJMan α-azidoethyl glycosides (**1** or **3**) were performed in dimethylformamide (DMF)-phosphate buffered saline (PBS; pH 7.4) in the presence of tris(3-hydroxypropyl-triazolylmethyl)amine (THPTA). A combination of cupric sulfate and ascorbic acid was used to generate the Cu(I) catalyst, resulting in the formation of tetravalent (**23**–**26**) and hexavalent (**27** and **28**) displays (Scheme 3).

Next, second-generation conjugates **29** and **30** were obtained by incorporating an N'-pentynoyl appendage on the unsubstituted lysine residue in (pseudo)glycocyclodecapeptides **23** and **24** (see Supporting Information, p. S15). The resulting alkyne-functionalized derivatives **40** and **41** were subjected to CuAAC with the structurally related tetrazide **42**[45] to yield the desired hexadecavalent compounds (Scheme 4). Similar transformations were carried out with the alkyne-functionalized polylysine (pseudo)glycodendrons **43** and **44**, obtained by Npentynoylation of **25** and **26** (Supporting Information, p. S17), and tetrazides **42** and **45**, [46] resulting in the corresponding homologous hexadecavalent derivatives **31**, **32** and **33**, **34**, respectively (Scheme 4). Finally, N'-azidoacetylation of **25** and **26** with azidoacetic acid N-hydroxysuccinimide (NHS) ester (Supporting Information, p. S22) and CuAAC of the resulting



**Figure 13.** Structures of the βCD-scafolded (pseudo)glycoclusters previously synthesized **19**, **20** (heptavalent) and **20**, **21** (tetradecavalent).



**Figure 14.** Structures of the first generation (pseudo)glycoclusters built on cyclodecapeptide (**23** and **24**; tetravalent), polylysine dendron (**24** and **26**; tetravalent) and cyclophosphazene (**27** and **28**; hexavalent) scaffolds.



**Scheme 3.** Synthesis of the first generation (pseudo)glycoclusters **23**–**28**.



**Figure 15.** Structures of the second generation (pseudo)glycoclusters combining cyclodecapeptide/cyclodecapeptide (**29** and **30**; hexadecavalent), cyclodecapeptide/polylysine dendron (**31** and **32**; hexadecavalent), polylysine dendron/polylysine dendron (**33** and **34** (hexadecavalent) and polylysine dendron/cyclophosphazene (**35** and **36**; tetraeicosavalent) scaffolds.

(pseudo)glycodendritic azides **46** and **47** with the hexaalkynylphosphazene platform 39 provided the tetraeicosa Man<sub>2</sub> and OMJMan conjugates **35** and **36** (Scheme 4) The structure, homogeneity, and purity of all new compounds were confirmed by mass spectrometry (MS), reversed-phase (RP)-HPLC, <sup>1</sup>H, <sup>13</sup>C,

and, when applicable,  $31P$  NMR (Supporting Information, Figures S1–S48).

The  $IC_{50}$  values obtained by ELLA for the inhibition of yeast mannan/HRP-ConA binding by the whole series of Man<sub>2</sub> and OMJMan multiconjugates **19**–**36** are summarized in Table 5 (see



**Scheme 4.** Synthesis of the second generation (pseudo)glycoclusters **29**–**36**.

the Supporting Information, Figure S79–S81, for the corresponding inhibition plots). It should be noted that ELLA data tends to reach a plateau for strong lectin binders, primarily due to the inherent limitations of the technique in accounting for crosslinking and chelate contributions to the multivalent effect. Nevertheless, the overall results indicate that OMJMan benefits from multivalency similarly to Man<sub>2</sub>. The extend of the inhibitory activity enhancement (relative potency, RP), normalized to that of the monovalent pseudodisaccharide (RP/n), depends on the valency and on the platform used. The hexadecavalent cyclodecapeptide-based OMJMan cluster **30** stood out among the second generation derivatives (RP/n  $=$ 10.6), whereas the hexavalent cyclophophazene conjugate **28** showed the highest RP/n value (6.64) among the first generation derivatives. By comparison, the increase in affinity with the number of epitopes is somewhat less pronounced for βCD conjugates (RP/ $n=1.44$  and 3.31, for the hepta and tetradecavalent OMJManclusters **20** and **22**, respectively; even much lower, 0.48 and 0.38, for the Man<sub>2</sub> conjugates 19 and 21). This can be attributed to the higher degree of compaction of the substituents on the βCD rims, which may hinder the approachability of HRP-ConA and limit accessibility of the internal (nonreducing) Man unit to the CRD. $[47]$ 

### **Comparative evaluation of DC-SIGN and langerin binding to OMJMan pseudoglycoclusters by surface plasmon resonance (SPR)**

Subsequently, we investigated the binding abilities of OMJMan pseudoglycoclusters towards DC-SIGN and langerin using SPR analysis. The obtained apparent dissociation constant (App  $K_D$ ) values from the SPR sensorgrams are presented in Table 6. For the cyclodecapeptide-scaffolded tetravalent compound **24**, satisfactory curve fittings could not be achieved. Instead, we determined the  $IC_{50}$  values for the inhibition of the corresponding lectin binding to Man-BSA. It should be emphasized that in this case the binding of the ligand to the lectin occurs in the solution phase, whereas for  $K_D$  determinations it takes place on the surface. Therefore, any direct comparison between these two sets of data should be approached with caution.

The SPR data obtained for the first-generation clusters revealed App  $K_D$  (or  $IC_{50}$ ) values in the micromolar range, whereas the second generation exhibited  $K_D$  values in the nanomolar range. Notably, considering that the valency is fourfold higher in the second generation series, the significant enhancement in binding affinity is consistent with the ability of the mannobioside mimetic OMJMan to leverage the multivalent





 $^{\circ}$  Folds of increase relative to the corresponding IC<sub>50</sub> value for the monovalent Man<sub>2</sub> or OMJMan glycoside (1 or 3, respectively).

Table 6. SPR-derived apparent dissociation constant (App K<sub>D</sub>) values for the OMJMan pseudoglycoclusters 26, 28, 30, 32, and 36 binding to DC-SIGN and langerin, and IC<sub>50</sub> values for the inhibition of DC-SIGN and langerin binding to surface-immobilized BSA-Man by compound 24. DC-SIGN vs langerin selectivity values are also shown. Data for the OMJMan/βCD conjugates 20 and 22 are included for comparative purposes.<sup>[1</sup>]



<sup>a</sup> Defined as App K<sub>D</sub> (langerin)/App K<sub>D</sub> (DC-SIGN) or IC<sub>50</sub> (langerin)/ IC<sub>50</sub> (DC-SIGN). <sup>b</sup> No binding detected at 500 µM. <sup>c</sup> IC<sub>50</sub> value for the inhibition of the lectin binding to mannosylated bovine serum albumin (Man-BSA).

effect, as previously observed for ConA binding. Furthermore, the increase in valency had a remarkable impact on the selectivity between DC-SIGN and langerin. While the binding affinity for DC-SIGN was only slightly more favorable than for langerin for the smaller pseudoglycoclusters **24**, **26** and **28** (1.41 to 1.45-fold), the App  $K_D$  (langerin)/App  $K_D(DC-SIGN)$  ratio increased to 33.3, 43.3, and 26.9 for compounds **32**, **34**, and **36**, respectively. Interestingly, compound **30** exhibited reversed selectivity, demonstrating it to be an almost two-fold better ligand for langerin compared to DC-SIGN. This unexpected result could be attributed to non-specific interactions between the scaffold itself and the langerin-functionalized surface. Previous studies have shown that the contribution of the naked double cyclopeptide-based scaffold, without any glycoligand, accounts for up to one-third of the binding observed for a tetradecavalent mannobioside mimetic constructed on the same system. In contrast, this platform is only minimally recognized by DC-SIGN.<sup>[48]</sup>

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It is also interesting to compare the findings presented in this study with previous data obtained for the hepta- and tetradeca-valent OMJMan/βCD conjugates **20** and **22**. In the latter case, the increase in valency did not lead to a strengthened multivalent effect against DC-SIGN. This result aligns with the observations made for ConA, reinforcing the idea that the access of the glycomimetic motif to the lectin CRD is significantly hindered when presented on the more compact βCD platform compared to other scaffolds. The less accessible CRD of langerin renders this lectin more susceptible to steric hindrance, resulting in a complete lack of binding.

Given the good performance of the Force Fields at reproducing the conformational properties of the HMO-related compounds **8**–**11**, we decided to apply a similar approach to model the more complex multivalent derivatives. Experimental details are given in the supplementary information. Figure 16 depicts representative structures for the heptavalent and tetradecavalent OMJMan-βCD (20 and 22) and Man<sub>2</sub>-βCD (19 and **21**) glycoclusters. Based on crystallographic structures we

construct two models for the extracellular domains of DC-SIGN and langerin, with the CRD and the first module of the extracellular alpha helix connecting it with the membrane, E and F, respectively. The increase of the App  $K<sub>D</sub>$  value on going from the heptavalent to the tetradecavalent derivatives can be ascribed to a classical multivalent effect: several situations in fast interconversion can display almost similar interactions (fast up-down rotation). Interestingly, close examination of the molecular models reveals that the rigid βCD platform hinders the access of the internal ("reducing") Man residue to the protein CRD (Figure 16). This provides a rational for the observed differences in the behaviour of the ligands relative to the two lectins: while the more open tetrameric arrangement of DC-SIGN enables the interaction with the external Man or OMJ residue as well as with the internal Man unit, trimeric langerin can only accommodate external Man residues. Consequently, langerin binding is banned for the OMJMan conjugates **20** and **22**.



**Figure 16.** A–D) Representative molecular models of compounds **19**–**22**. E) Structure of DC-SIGN. F) Structure of langerin. Protein structures are taken from the PDB entries 1k9i and 3PQG with minor automatic modifications, while in the ligands the βCD ring is represented as CPK and the substituents as sticks.

The lack of selectivity in the case of multivalent OMJMan glycoclusters built on RAFT cyclodecapeptide, lysine-based dendrimer or cyclophosphazene platforms probably implies the participation of the internal Man residue on langerin binding. Inspection of the molecular models for the first-generation compounds **24**, **26** and **28** as well as for the second generation conjugates **32**, **34** and **36** indicates a much less compact architecture as compared with the βCD derivatives, supporting this hypothesis (Figure 17).

### **Conclusions**

The results discussed in this manuscript provide a practical approach for the synthesis of HMO hemimimetics that faithfully replicate the substitution and configuration pattern of their natural counterparts. This general strategy is based on the concept of chemical mimicry, leveraging the unique stereoelectronic properties of sp<sup>2</sup>-iminosugars. Specifically, the cyclic acylimine functionality exhibits glycosylation-like reactivity, controlled by the anomeric effect, enabling the implementation of synthetic schemes that mirror established methods for accessing the mimicked oligosaccharides. NMR and computational investigations confirm that replacing mannopyranoside units in HMOs with the  $sp^2$ -iminosugar analog motif OMJ maintains the available conformational space in aqueous solution. Moreover, the OMJ-capped HMO hemimimetics studied in this work exhibit binding to the mannose-specific lectins ConA, DC-SIGN, and langerin, adopting essentially the same bound conformation as the emulated HMO. Interestingly, the OMJMan glycotope, a mannobiose-like structure, behaves as a heterodivalent ligand for all three lectins, similar to Man<sub>2</sub>. Notably, the behavior of the pseudopentasaccharide (OMJ- $Man$ )<sub>2</sub>Man can be rationalized as a divalent OMJMan entity, analogous to Man $_5$  as a divalent mannobiose derivative. Furthermore, OMJMan clusters benefit from classical multivalent effects, leading to additional enhancements in binding affinity, highlighting their functional biomimetic character. The results also stress that the subtle structural differences between Man and OMJ units can result in distinct preferences for accommodation in the carbohydrate recognition domain (CRD) of specific lectins, especially when presented in multiple copies. Most importantly, it is shown that the selectivity towards the biomedically relevant CLRs DC-SIGN and langerin can be biased with the OMJMan glycotope surrogate. Our findings demonstrate that the overall architecture of the (pseudo)glycocluster, not just the valency, plays a crucial role in this context. We envision that the chemical mimicry strategy can be expanded to access hemimimetics of virtually any natural oligosaccharide containing  $\alpha$ -linked monosaccharide residues, thereby opening up new avenues to intervene in carbohydrate-mediated recognition processes.

### **Experimental Section**

**General Methods**: All commercial chemicals were of reagent grade and were used without further purification. Optical rotations were measured at 20 $\pm$ 2 °C in 1-dm tubes on a Jasco P-2000 polarimeter.  ${}^{1}$ H (and  ${}^{13}$ C NMR) spectra were recorded at 500 (125.7) MHz with Bruker 500 DRX o BrukerAvance III spectrometers and chemical shifts (δ) were reported in parts per million (ppm). 1D TOCSY, 2D COSY, HMQC and HSQC experiments were used to assist on NMR assignments. Thin-layer chromatography (TLC) was carried out on aluminum sheets coated with silica gel 60  $F_{254}$  Merck with visualization by UV light and by charring with ethanolic 10%  $H_2SO_4$  and 0.1% ninhydrin. Column chromatography was carried out on Silica Gel 60. ESI mass spectra were recorded on a Bruker Daltonics Esquire6000™ ion-trap mass spectrometer. Elemental analyses were carried out at the Instituto de Investigaciones Químicas (Sevilla, Spain).

The compounds 2-azidoethyl *O*-(α-D-mannopyranosyl)-(1→2)-α-Dmannopyranoside (Man<sub>2</sub>, 1),<sup>[24]</sup> 2-azidoethyl α-d-mannopyranosyl-(1!2)-α-D-5 *N*,6*O*-oxomethylidenemannonojirimycin (ManOMJ, **2**), 2-azidoethyl α-D-5 *N*,6*O*-oxomethylidenenojirimycinyl-(1→2)-α-Dmannopyranoside (OMJMan, **3**,) 2-azidoethyl α-d-5 *N*,6*O*-oxomethylidenemannonojirimycinyl- (1→2)-α-D-5 *N*,6*O*-oxomethylidenemannonojirimycin (OMJOMJ, **4**), 2-azidoethyl 3,6-di-*O*-α-Dmannopyranosyl-α-D-mannopyranoside (Man<sub>3</sub>, 8),<sup>[25]</sup> 2-azidoethyl<br>2,4-di-O-benzoyl-α-D-mannopyranoside (12),<sup>[22]</sup> (1*R*)-2,3,4-tri-O-2,4-di-*O*-benzoyl-α-D-mannopyranoside (**12**),[22] (1*R*)-2,3,4-tri-*O*acetyl-1-fluoro-5 *N*,6*O*-oxomethylidene-1-deoxymannojirimycin<br>(13),<sup>[23]</sup> S-tolyl-3,4,6-tri-O-benzoyl-a-D-mannopyranoside (15).<sup>[24]</sup> (**13**),[23] *S*-tolyl-3,4,6-tri-*O-*benzoyl-α-D-mannopyranoside (**15**),[24] heptavalent Man-Man βCD7 derivative (**19**),[14] heptavalent OMJ-Man βCD7 derivative (**20**),[14] tetradecavalent Man-Man βCD7 derivative (21),<sup>[14]</sup> tetradecavalent OMJ-Man βCD7 derivative (22),<sup>[14]</sup> tetrapropargylated RAFT (37),<sup>[43]</sup> tetrapropargylated polylisine dendron (38),<sup>[43]</sup> hexapropargylated cyclotriphosphazene (39),<sup>[44]</sup> RAFT-N3 (**42**),[45] polylysine scaffold-N3 (**45**),[46] 2,3,4,6-tetra-*O*-benzoyl-α-Dmannopyranosyl trichloroacetimidate (**48**; Supporting information, p S8)[49] and 2-*O*-(2,3,4,6-tetra-*O*-benzoyl-α-D-mannopyranosyl-2,3,4 tri-*O-*benzoyl-α-D-mannopyranosyl trichloroacetimidate (**50**; Supporting information, p S13) were prepared according to described procedures.

### **Supporting Information**

Experimental protocols for all glycosylation, protection/deprotection and multi-CuAAC reactions, compound characterization for all substrates and products, NMR spectra for substrates and products, HPLC chromatograms, detailed protocols for conformational and lectin binding studies by computational and spectroscopic techniques, ELLA, and SPR, including molecular representations, data plots and sensorgrams. The authors have cited additional references within the Supporting Information.[51–64]

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**Figure 17.** A–F) Representative molecular models of compounds **24, 26, 28, 32, 34**, and **36**. G) Structure of DC-SIGN. H) Structure of langerin. Protein structures are taken from the PDB entries 1k9i and 3PQG with minor automatic modifications, while the ligands are represented as sticks.

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### *Conflict of Interests*

The authors declare no conflict of interest.

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# **RESEARCH ARTICLE**

**The "carbohydrate chemical mimicry"** of sp 2 -iminosugars enables the synthesis of high-mannose-type oligosaccharide analogs whose solution and lectin bound conformations match those of the natural partners. Upon multivalent presentation, enhanced affinities towards ConA, DC-SIGN and langerin are achieved, with distinct selectivity profiles.



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**High-Mannose Oligosaccharide Hemimimetics that Recapitulate the Conformation and Binding Mode to Concanavalin A, DC-SIGN and Langerin**