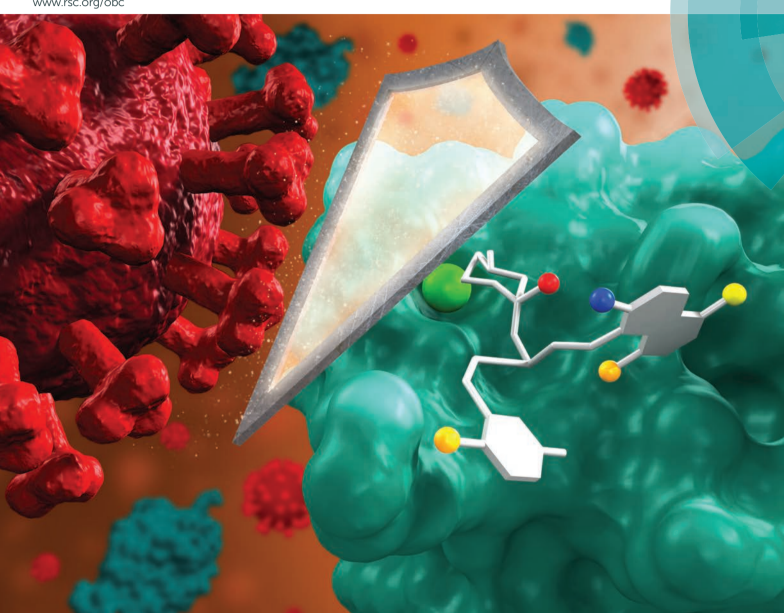
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STD NMR and molecular modelling insights into interaction of novel mannose-based ligands with DC-SIGN†

Anita Kotar, ^a Tihomir Tomašič, ^b Martina Lenarčič Živković, ^a Gregor Jug, ^b Janez Plavec*^{a,c,d} and Marko Anderluh*^b

Study of interaction of mannose-based ligands with receptor DC-SIGN using high resolution NMR in combination with molecular modelling showed that four α -p-mannoside ligands interact with the binding site predominantly through the mannose moiety. The other two aromatic groups that are bound to α -p-mannose through a glycerol linker demonstrate interaction that can be related to their substitution pattern. Ligand with naphthyl and *meta*-substituted phenyl ring exhibited the most favourable binding characteristics. In addition to the predicted hydrophobic interactions of aromatic moieties our results propose new contacts of substituted phenyl moiety in the more polar area of the flat binding site of DC-SIGN and thus offer new possibilities in further designing of novel, more potent DC-SIGN antagonists.

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Introduction

Although significant progress has been made in HIV vaccine strategies and drugs development, HIV infections remain one of the leading causes of morbidity and mortality worldwide. 1-5 Dendritic cells (DCs), host professional antigen-presenting cells that represent the first line of the innate immune system, play an important role in HIV-1 infection. The pattern recognition receptors (PRRs) located at the surface of DCs are an important factor in HIV-1 transmission that help subvert DC functions and escape immune surveillance.4,7 One of these PRRs that plays a fundamental role in cellular immunity is DC-SIGN (Dendritic Cell-Specific Intercellular adhesion molecule-3-Grabbing Non-integrin), a type II transmembrane C-type lectin located specifically on DCs.8-10 DC-SIGN acts as an adhesion molecule and binds highly glycosylated proteins present on the surface of several pathogens, including mannose glycans of the envelope protein gp120 of HIV-1. 11-13 On binding, the complex of DC-SIGN and HIV-1 is promptly internalized, degraded into smaller fragments in early endosomes and conjugated with MHC class-II proteins to initiate a

Different approaches were applied to design medicinally useful DC-SIGN antagonists. Even though reported antagonists vary in structure and properties, they mostly exploit the fact that DC-SIGN recognises various mannose- and fucose-containing glycans exposed on the surface of several viruses, bacteria, yeast and parasites. 4,8,11,23 D-Mannose ($K_i = 13.1$ mM) and L-fucose ($K_i = 6.7 \text{ mM}$) alone bind to DC-SIGN with very weak affinities, but when incorporated into suitable glycoconjugates substantially higher affinities can be obtained that offer a promise to prevent HIV-1 transmission.8,12,23,24 We recently reported a small focused library of branched mannose-based DC-SIGN antagonists bearing diaryl substituted 1,3-diaminopropanol or glycerol moieties incorporated to target the hydrophobic groove of the receptor.²³ The D-Mannose entity was presumed to serve as an anchor for interaction with the Ca2+ binding site of the DC-SIGN carbohydrate recognition domain (CRD). The designed molecules were evaluated by in vitro assay of the isolated DC-SIGN extracellular domain for their ability to compete with HIV-1 gp120 for binding to the DC-SIGN CRD. The two most potent compounds were found to inhibit the interactions between gp120 and DC-SIGN with the IC₅₀ values of 40 μ M and 50 μ M, and were among the most potent monovalent DC-SIGN antagonists

humoral immune response from T cells, but unfortunately, a small amount of virus succeeds to escape the normal lytic pathway in DCs' endosomes and avoid the host immune system. By inhibiting interaction between DC-SIGN and HIV-1 with DC-SIGN antagonists pathogen entry could be prevented causing an impediment of the infection of immune defence cells. 1,15-23

^aSlovenian NMR center, National Institute of Chemistry, Hajdrihova 19, 1000 Ljubljana, Slovenia. E-mail: janez.plavec@ki.si; Fax: +38614760300

^bFaculty of Pharmacy, University of Ljubljana, Aškerčeva 7, 1000 Ljubljana, Slovenia. E-mail: marko.anderluh@ffa.uni-lj.si; Fax: +38614258031

^cEN-FIST Centre of Excellence, Trg OF 13, 1000 Ljubljana, Slovenia

^dFaculty of Chemistry and Chemical Technology, University of Ljubljana, Večna pot 113, 1000 Ljubljana, Slovenia

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Fig. 1 Chemical structures of monovalent mannose-based ligands.

reported. Furthermore, the antagonistic effect of these compounds was evaluated by a one-point in vitro assay that measures dendritic cell adhesion to a mannan-coated surface. Again, the same compounds acted as functional antagonists of DC-SIGN-mediated DC adhesion. The binding mode of the most potent compound (4) was also studied by molecular docking and molecular dynamics simulation, which suggested that the diarylglycerol part of the molecule likely binds into the targeted hydrophobic binding site, and provides a basis for further optimization.²³ Unfortunately, the binding site of DC-SIGN is rather flat and vaguely defined and offers limited (other than van der Waals) ligand-receptor contacts.

In the present study we investigated the interactions between four monovalent mannose-based ligands 1-4 (Fig. 1) and extracellular domain of DC-SIGN (hereinafter referred as DC-SIGN). The studied ligands 1-4 consist of α -D-mannose connected by glycerol linker to different aryl moieties, which are expected to tune interactions of ligands in the hydrophobic groove of the receptor. A combination of saturation transfer difference (STD) NMR spectroscopy and molecular modelling was used to examine binding propensity of monovalent glycoconjugates for DC-SIGN at the atomic level of resolution. As expected, ligands 1-4 do not exhibit clear differences in chair conformation of D-mannose sugar moiety regardless of variation of groups attached to its anomeric carbon. On the other hand, differences in ligands' binding affinities can be correlated with the position of carboxylic group on the phenyl moiety in 1-3 as well as lipophilicity of each of the ligands. The binding mode of the ligands was additionally evaluated by molecular dynamic simulations. A good agreement between NMR and molecular modelling results provides new insights into interactions of our novel ligands with DC-SIGN and suggests new possibilities in design of novel anti-HIV therapeutics.

Results

Chemistry

DC-SIGN antagonists 1 and 4 were synthesized as reported, 23,25 while synthesis of 2 and 3 is outlined in Scheme 1. 3-Hydroxybenzoic acid (5) was first converted to its methyl ester 6 in methanol under acidic conditions. Potassium salt of compound 6, which was prepared using potassium hydroxide in methanol, was used for epoxide ring opening of 7 to obtain racemic diarylglycerol 8. Mannosylation of 8 with 2,3,4,6-tetra-O-acetyl-α-D-mannopyranosyl trichloroacetimidate²³ and with trimethylsilyl trifluoromethanesulfonate (TMSOTf) promoter yielded α -D-mannoside 9 stereospecifically. Since 8 was used as a racemate, 9 was obtained as a mixture of two diastereomers that could not be separated using flash column chromatography. Deprotection of 9 using the Zemplén method gave final compound 3, which was further hydrolysed under alkaline conditions to yield the final compound 2.

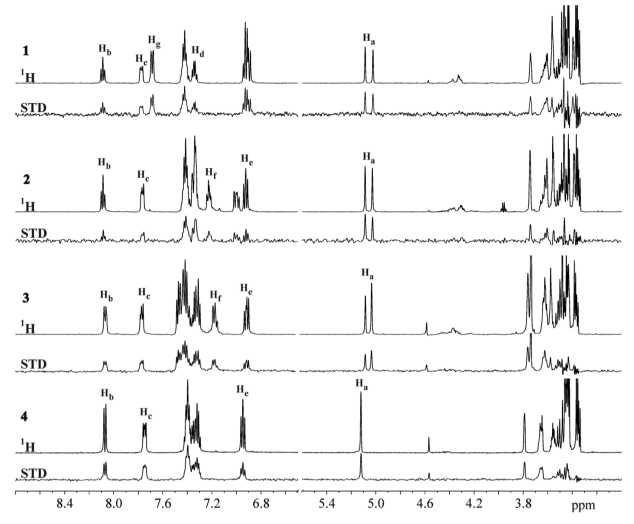
STD NMR experiments and epitope mapping

The STD NMR experiments were performed to explore the binding nature and propensity of ligands 1-4 to interact with DC-SIGN. Reference and STD spectra exhibited narrow and well resolved lines (Fig. 2). Significant STD effects observed for ligands 1-4 unequivocally demonstrated their binding to the DC-SIGN. In order to increase accuracy of epitope mapping we used only well resolved and isolated NMR resonances of the individual ligand, avoiding uncertain contributions in the case

HO COOME

$$R_2O$$
 R_2O
 $R_$

Scheme 1 Reagents and conditions: (a) SOCl₂, MeOH, r.t., 24 h; (b) (i) KOH, MeOH, r.t., 20 min; (ii) tetrabutylammonium bromide, toluene/N,N-dimethylformamide, 90 °C, 20 h; (c) 2,3,4,6-tetra-O-acetyl-α-p-mannopyranosyl trichloroacetimidate, TMSOTf, CH₂Cl₂, 0 °C, then r.t., 24 h; (d) NaOMe, MeOH, 1 h, then Amberlite® IR120 H, 10 min; (e) 1 M NaOH, EtOH, r.t., 24 h.



¹H and STD NMR spectra of 1–4 in the presence of DC-SIGN. NMR spectra were recorded at 25 °C, 2.0 s saturation time, 400 μM concentration of ligand and 1:50 receptor-to-ligand ratio in 25 mM Tris-d₁₁ (pD = 8.0), 150 mM NaCl, 4 mM CaCl₂ in ²H₂O and 25% DMSO-d₆ at 600 MHz. Resolved signals are labelled according to atom numbering shown in Fig. 3.

of signal overlap. Quantitative analyses of the STD spectra were done at receptor-to-ligand ratio of 1:50. Rather high ratio was used to reduce the effect of cross-rebinding, where a previously saturated ligand re-enters a binding pocket.

For 1-4 the most intense STD signals were observed between δ 3.3 and 3.9 ppm, which suggested that the mannose moiety was closest in contact with the surface of DC-SIGN. However, quantitative interpretation of STD effects of sugar protons was hampered due to spectral overlap with the exception of anomeric H_a proton. H_a proton exhibited well-resolved signals for all four ligands 1-4. Moreover, it showed the highest relative STD effect among protons that were analysed in epitope mapping. Consequently, the signal of H_a proton was set as a reference with 100% STD effect. Intensities of other signals were expressed relative to the H_a signal (Fig. 3).

The aromatic groups exhibited weaker STD effects in comparison to the mannose moiety regardless of variations in the attached aryl entities. Both phenyl Hg protons of 1 together

showed STD effect of 51%. However, in the STD spectra one cannot distinguish between individual contributions of two H_g atoms due to their symmetric arrangement at meta-position of phenyl group. H_b, H_c and H_d protons of naphthyl group of 1 exhibited STD effects of 61%, 43% and 53%, respectively. Obvious STD effects were seen for other aromatic protons as well. Unfortunately their quantitative elucidation was prevented due to spectral overlap. In 2 the phenyl H_f proton showed STD effect of 68% and H_b, H_c and H_e protons from the naphthyl moiety exhibited the STD effects of 78%, 54% and 50%, respectively. However, direct comparison of ligands has to be done with certain degree of caution because of monitoring of protons on different positions on aromatic groups of ligands, namely H_d in 1 and H_e in 2-4 on naphthyl ring, and H_g in 1 and H_f in 2-4 on phenyl moiety. In 3 H_b and H_c protons from naphthyl and H_f from phenyl group showed similar STD effects ranging from 73% to 77%. He proton in 3 exhibited STD effects of 65%. Experimentally observed STD

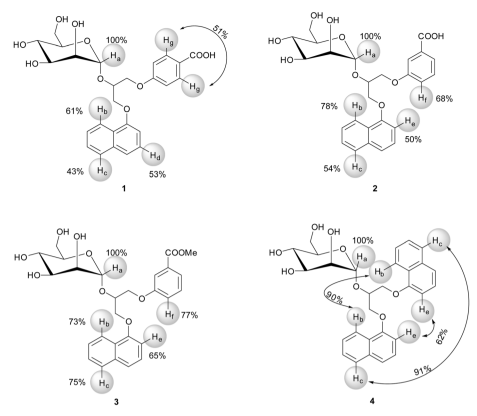


Fig. 3 Epitope mapping for 1-4. The relative degrees of saturation of marked H_a-H_g protons expressed in % are mapped onto the structure and normalized with respect to H_a proton. STD effect of H_a was arbitrarily set to 100% in the case of all four ligands. Arrows are used to report the overall STD effects for symmetric groups.

effects for H_b and H_c protons on both naphthyl rings in 4 were 90% and 91%, respectively. STD effects of H_e protons were markedly weaker (62%). However, one should keep in mind that H_b , H_c and H_e protons in 4 could not be resolved due to symmetry of the ligand. Observed STD effects therefore represented the overall contribution to the binding for both protons.

STD build-up studies

To precisely map ligand epitopes in close contact with the receptor we acquired STD build-up curves for 1–4 by collecting STD spectra at twelve different saturation times ranging from 0.1 to 5.0 s at receptor-to-ligand ratio of 1:50. The maximal STD intensities, $\mathrm{STD}_{\mathrm{max}}$, and the saturation rate constants, k_{sat} , were obtained by fitting experimental data to eqn (2). Experimental data as well as best-fit curves for 1 are shown in Fig. 4, whereas the corresponding figures for 2–4 can be found in the ESI (Fig. S2–S4†). The agreement between the experimental data and the fitted model was good with typically χ^2 values below 0.1. A clear build-up of the STD effects with increasing saturation time can be observed, reflecting the amplification of the STD (Fig. 4 and S4–S6†).

At longer saturation times flattening of the build-up curves can be observed, because of the relaxation rate of saturated ligand in the free state. For 1 the most prominent STD_{max}

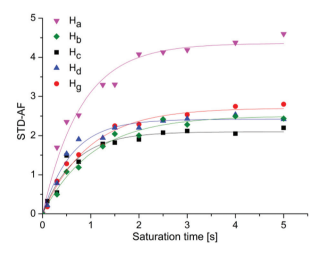


Fig. 4 STD-amplification factor of 1 as a function of saturation time. Experimental data were fitted to a rising exponential eqn (2), to obtain the STD_{max} and k_{sat} . H_a: STD_{max} = 4.46 \pm 0.1, k_{sat} = 1.1 \pm 0.1 s⁻¹, R^2 = 0.973; H_b: STD_{max} = 2.50 \pm 0.07, k_{sat} = 0.96 \pm 0.08 s⁻¹, R^2 = 0.983, H_c: STD_{max} = 2.06 \pm 0.07, k_{sat} = 1.5 \pm 0.2 s⁻¹, R^2 = 0.961; H_d: STD_{max} = 2.49 \pm 0.06, k_{sat} = 1.5 \pm 0.2 s⁻¹, R^2 = 0.982; H_g: STD_{max} = 2.66 \pm 0.08, k_{sat} = 1.1 \pm 0.1 s⁻¹, R^2 = 0.978). Symbols represent data, whereas solids lines are the mathematical least-square fits.

belonged to the H_a proton (4.46), while the corresponding values for aromatic H_b, H_c, H_d and H_g protons were markedly lower (2.50, 2.06, 2.49 and 2.66, respectively). Accordingly, H_a proton should be the one most directly involved in binding and closer to the receptor, followed by aromatic protons that are approximately equidistant from the protein surface. Different k_{sat} values spanning from 0.96 to 1.54 s⁻¹ were calculated for protons of 1. In the case of 2 the STD_{max} values of H_a, H_b, H_c, H_e, H_f protons were 3.71, 1.30, 1.87, 1.48 and 1.61, respectively. It is interesting that k_{sat} values among protons of 2 did not vary remarkably and were all around 1.30 s^{-1} . For 3 the highest STD_{max} of 3.44 was obtained for H_a proton. Significantly lower values were acquired for aromatic protons, H_b (2.01), H_c (1.58), H_e (1.08) and H_f (1.77). Great differences in $k_{\rm sat}$ from 1.19 to 2.45 s⁻¹ were determined among protons of 3. Similarly, protons of 4 exhibited k_{sat} values ranging between 1.59 and 2.44 $\ensuremath{\text{s}^{-1}}.$ The STD_{max} values of 4 were 3.41 for H_a proton and 1.53, 1.69 and 1.30 for naphthyl H_b , H_c and H_e protons, respectively. Once again, results inferred that the Ha proton was in close contact with the protein and that the mannose moiety contributed primarily to the binding process. Furthermore, initial build-up rates were analysed as well. Initial slopes of the curves at zero saturation time were derived by multiplication of STD_{max} and k_{sat} and correspond to the STD effects (STD₀) in the absence of influence of T₁ longitudinal relaxation time bias and rebinding effects. We calculated STD₀ for H_a-H_{\sigma} protons and compared these results with the STD intensities at 2.0 s saturation time (STD_{2s}) from epitope mapping. For ease of comparison, the values were normalized with respect to H_a as a 100 (Table 1).

The STD₀ values were lower than STD_{2s} effects. The two exceptions were H_c and H_d protons of 1, which exhibited markedly higher STD₀ values. STD_{2s} and STD₀ values for H_g of 1 were comparable. The same observations were noticed also for H_c and H_e protons of ligand 2. On the contrary, H_b and H_f protons of 2 showed significant differences in STD effects. For ligand 3, identical STD_{2s} and STD₀ values were observed for H_e

Table 1 Normalized STD intensities^a

		1	2	3	4
STD _{2s}	H_b	61	78	73	90
	H_{c}	43	54	75	91
	H_d	53	nd	nd	nd
	H_{e}	nd^b	50	65	62
	H_{f}	nd	68	77	c
	H_{g}	51	nd	nd	_
STD_0	$H_{\rm b}^{\circ}$	49	46	59	42
	H_{c}	62	48	53	38
	H_d	73	nd	nd	nd
	H_{e}	nd	45	65	45
	H_{f}	nd	49	58	_
	$H_{ m g}$	46	nd	nd	_

^a Normalized intensities for H_b-H_g protons at 2.0 s saturation time with respect to STD of H_a which was arbitrarily given a value of 100. ^b Not determined because of peak overlap. ^c Ligand does not contain this proton.

proton. On the other hand, significantly lower STD₀ effects were detected for H_b, H_c and H_f protons in comparison to STD_{2s} values. Surprisingly, the lowest STD₀ values and the greatest variances among STD effects were observed for protons of 4.

Characterisation and comparison of binding conformations

The normalized build-up curves of STD effects for individual protons were compared to characterise differences in binding modes among ligands 1-4 (Fig. 5). STD effects of analysed protons within individual ligand at different saturation times were normalized to STD effect of Ha proton at saturation time

Good correspondence in overall trend between STD buildup curves was observed for H_a proton in 1-4 (Fig. 5a). One can observe slight deviation of data points among the ligands at saturation times between 1.0 and 2.0 s (Fig. 5a). Similar buildup curves suggest comparable binding conformation of mannose moiety in 1-4 in the binding site of DC-SIGN.

Normalized STD-AF values for Ha proton were the most pronounced in all four cases of interaction analysis of 1-4 with DC-SIGN. These results suggest that the Ha proton and associated mannose sugar serve as a binding initiator and are crucial for interaction. Comparison of normalized build-up curves for H_b proton indicated different rate of saturation of this proton within individual ligand. 1 and 3 exhibited comparable outline of build-up curves and reach almost identical plateau value. Build-up curve of 2 grew in the same way, but reached the plateau at a lower value. 4 experienced different build-up mode of H_b protons with steeper initial slope that flattens at value comparable to H_b proton of 2. Differences in saturation trend originate from contribution of both Hb protons of 4, which obviously get saturated in shorter time but do not contribute predominantly to the binding. The normalized STD-AF values for H_c proton showed excellent correspondence with deviation of individual experiments data points within experimental error (Fig. 5c). These observations confirmed analogous binding mode of H_c proton of 1-4. H_c protons of 1-3 participate approximately equally in the interaction with the receptor. Both H_c protons of 4 together showed similar plateau value as 1-3. However, one should keep in mind that the contribution of individual H_c proton would be markedly reduced, suggesting lower binding propensity of ligand 4. As in the case of H_b protons of 4 we observed analogous slope outline for H_c protons of 4 indicating faster rate of saturation in comparison to 1-3. The major differences in build-up curves were observed for $H_{\text{d/e}}$ and $H_{\text{f/g}}$ protons. The discrepancy for H_d and H_g protons of 1 was expected due to their different position on aromatic rings in comparison to H_f and H_e protons of 2-4. On the other hand, significant alterations in STD effects of He proton of 2 and 3 imply different binding properties of the naphthyl group and highlight sensitivity of binding characteristics of ligands. Although, He protons of 2 and 4 exhibited comparable STD build-up curves, direct comparison can be misleading since the curve of 4 rep-

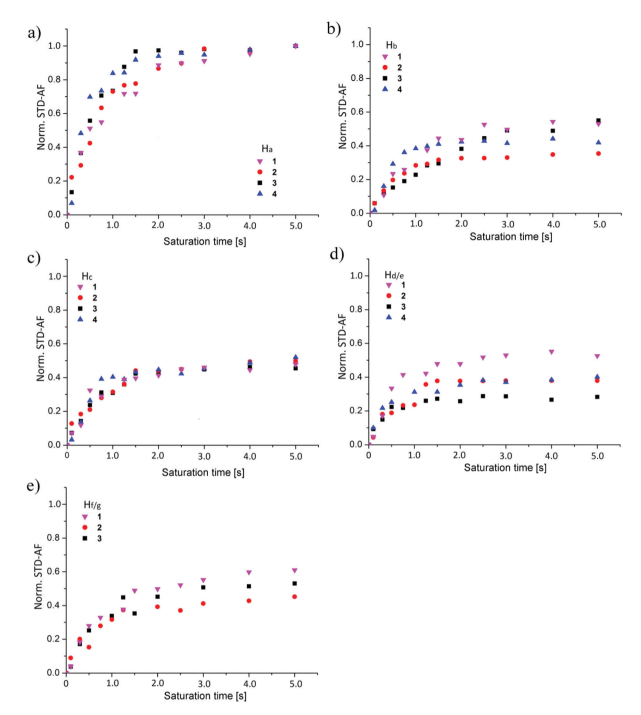


Fig. 5 Overlaid normalized STD build-up curves of individual protons in 1–4. The normalized STD effects of ligand protons are shown as a function of saturation time of (a) H_a , (b) H_b , (c) H_c , (d) $H_{d/e}$ and (e) $H_{f/g}$.

resents saturation of two He protons (Fig. 5d). When comparing structural setup and position of H_d proton of 1 and H_e of 2-4 it is evident that H_d proton was closer to the surface of receptor with respect to He. Furthermore, the comparison of curves of H_f and H_g protons of the phenyl ring from 1-3 showed similar STD build-up (Fig. 5e). Because of the inseparability of two Hg protons of 1 and their joint contribution to STD values we believe that the H_f of 2 and 3 are still spatially

closer to the hydrophobic groove of DC-SIGN in comparison to H_g of 1. Comparative analysis of 2 and 3 additionally revealed that H_f proton from 3 lies closer to the receptor with respect to H_f of 2. Higher lipophilicity of 3 evidently enables stronger interactions with the receptor. STD build-up curves comparison for 1-4 led us to conclusion that variations in chemical structures of ligands influence their bound conformations as well as binding modes in the binding site of DC-SIGN.

Evaluation of binding affinities

Amplification of STD effects with increasing concentration of 1-4 was used to determine the binding affinities. Receptorligand dissociation constants (K_D) were determined with the use of single-ligand titration experiments that were performed at five different ligand concentrations from 80 to 800 µM at constant saturation time of 2.0 s. By monitoring the growth of the STD-AF upon titration K_D values were calculated with the use of the eqn (3) (Fig. 6). Experimental data showed that the maximum STD-AF was not reached for any of ligands even at 800 µM ligand concentration. This observation is in agreement with fast exchange of ligands from the bound to the free state. The variation in exchange kinetics among the ligands is reflected in different slopes of hyperbolic curves (Fig. 6).

Consistently, these results suggest that exchange kinetics was the fastest for 4 and the slowest for 3. Thus 3 stayed in the binding site of DC-SIGN for a longer period of time in comparison to other ligands. The binding affinity of 3 and 2 was in lower milimolar range with K_D values of 0.45 mM and 0.96 mM, respectively. For 1 and 4 weaker binding with K_D values of 1.6 mM and 4.9 mM was observed, respectively.

The direct comparison of K_D values is difficult as many experimental factors, such as saturation time, ligand residence time in the complex, intensity of the signal and accumulation of saturation in the free ligand affect determination of receptor-ligand binding affinities by STD NMR spectroscopy. Therefore, we used binding isotherm of STD-AF initial growth approach that allows K_D evaluation without aforementioned spurious factors. To verify the sensitivity of the system of individual ligand and DC-SIGN to saturation transfer, we analysed

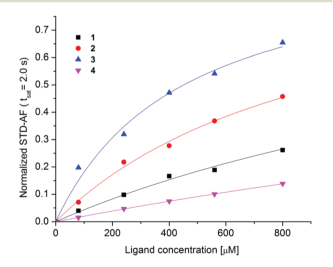


Fig. 6 Normalized STD-AF curves as a function of ligand concentration in the range from 80 to 800 μM . Experimental data were collected at 25 °C, 2.0 s saturation time on a 600 MHz NMR spectrometer with 8 μ M DC-SIGN concentration in 25 mM Tris-d₁₁ (pD = 8.0), 150 mM NaCl, 4 mM CaCl₂ and 25% DMSO-d₆ in ²H₂O. Experimental data were fitted to the eqn (3) (1: K_D = 1.6 \pm 0.5 mM, R^2 = 0.992, 2: K_D = 0.96 \pm 0.08 mM, R^2 = 0.999, 3: K_D = 0.45 \pm 0.05 mM, R^2 = 0.997, 4: K_D = 4.9 \pm $0.7 \text{ mM}, R^2 = 0.999$).

STD-AF of 2-4 at different ligand concentrations from 80 to 800 µM at saturation times between 1.0 and 4.0 s and obtained STD build-up curves. The STD₀ values as a function of the ligand concentration were fitted to eqn (3) to determine K_D values without dependence on the spectral properties of the observed signals (Fig. S8†). Binding isotherm protocol gave similar K_D values in comparison to the titration experiments at a single saturation time. K_D values were 0.90 mM for 2, 0.51 mM for 3 and 3.0 mM for 4. These K_D values confirmed that 3 exhibits the lowest k_{off} rate and thus spends the longest time as part of a complex with DC-SIGN.

Molecular docking

Plausible binding modes of compounds (S)-1, (R)-1, (S)-2, (R)-12, (S)-3 and (R)-3 were first studied by molecular docking calculations using our previously reported and validated FlexX protocol.23 The docking binding mode of compound 4 has already been described.²³ DC-SIGN CRD in complex with Man₄ (PDB entry: 1SL4) was selected as receptor for ligand docking under defined pharmacophore type constraints. In detail, to place the core p-mannose residue correctly, the side chain carboxylate groups of Glu354 and Glu347 were defined as hydrogen bond acceptors, which interact with 3- and 4-hydroxyl groups of the core D-mannose in the crystal structure. Additionally, a pharmacophore with spherical coordination around Ca2+ ion was defined to correctly consider the complex interactions between Ca2+ ion and 3- and 4-hydroxyl group of D-mannose. The D-mannose residue of the studied antagonists 1-4 is predicted to bind to the DC-SIGN Ca²⁺-binding site. In the case of (S)-3, (R)-3 (Fig. 7) and 4 the binding of the D-mannose residue is in agreement with the defined pharmacophore type constraints, whereas in the case of benzoic acid derivatives (S)-1, (R)-1, (S)-2 and (R)-2, the D-mannose residue is rotated in a way that only the 2-hydroxyl group is in contact with the Ca2+ ion. Such a D-mannose binding mode does not allow for octacoordination of the Ca2+ ion in the binding site and has, to the best of our knowledge, not been observed in the crystal structures of DC-SIGN CRD-ligand complexes. The FlexX-calculated binding modes of both diastereoisomers of compound 3 show the same binding mode of the D-mannose residue, which interacts with Ca2+ ion and forms hydrogen bonds with Glu347, Asn349, Glu354, Asn365 and Lys368 side chains (Fig. 7). The main difference is in the orientation of the diarylglycerol moiety. In the case of (R)-3 phenyl ring makes hydrophobic contacts with Phe313 and forms a hydrogen bond between ester carbonyl group and Ser360 side chain hydroxyl group, whereas the naphthalene ring points towards the solvent (Fig. 7a). The opposite can be observed in the binding mode of (S)-3, where the naphthalene ring contacts the Phe313 side chain, while the ester carbonyl group forms hydrogen bond with the Asn344 side chain (Fig. 7b). In contrast to (R)-3 and (S)-3 both naphthalene rings of compound 4 are predicted to form hydrophobic and/or π - π interactions with the side chain of Phe313.²³

In order to evaluate whether other docking tools can correctly predict the D-mannose binding pose of compounds 1

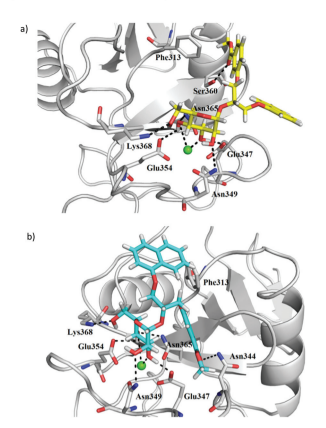


Fig. 7 FlexX docking poses of compounds (a) (R)-3 (in yellow sticks) and (b) (S)-3 (in cyan sticks) in DC-SIGN CRD Ca2+-binding site. Protein (PDB entry: 1SL4) is presented in grey cartoon and Ca²⁺ ion as a green sphere. For clarity only side chains of amino acid residues predicted to interact with the docked ligands are shown as grey sticks. Hydrogen bonds are shown as black dashed lines. Figure was prepared by PyMOL.²⁶

and 2, the docking of compounds 1-4 to DC-SIGN CRD was repeated with our recently reported docking protocol²⁷ using FRED 3.0.1, ²⁸⁻³⁰ as available in OEDocking software (Release 3.0.1, OpenEye Scientific Software, Inc., Santa Fe, NM, USA; http://www.eyesopen.com/). In this case, the Glu347 and Glu354 side chains were defined as hydrogen bond acceptors and the Asn365 side chain as a hydrogen bond donor to constrain the placement of the D-mannose moiety. FRED docking poses for compounds (R)-3 and (S)-3 were found to be very similar to those obtained by FlexX, positioning the naphthalene and phenyl rings to the same binding pockets of the binding site (Fig. S9†). In contrast to FlexX, FRED was able to correctly place the D-mannose residue of (S)-1, (R)-1, (S)-2 and (R)-2, allowing the octacoordination of Ca^{2+} by interactions with D-mannose 3- and 4-hydroxyl groups (Fig. S10 \dagger). Both aromatic moieties of (S)-1, (R)-1, (S)-2 and (R)-2 form hydrophobic contacts and/or π - π interactions with Phe313 side chain (Fig. S10†), also observed in the case of compound 4.23 Carboxylic acid moiety of these ligands points toward the solvent and is not in contact with the protein.

Molecular dynamics simulations

Recently, we have shown that DC-SIGN CRD is a challenging target for docking, because of its flat binding site, and careful selection of the docking software and parameters is needed to correctly predict the binding of DC-SIGN antagonists.²⁷ Considering the flat DC-SIGN binding site, the use of semi-flexible docking protocols, where the protein is treated as rigid and the ligand as flexible, is not sufficient to represent the dynamic nature of DC-SIGN-ligand interactions in solution. Moreover, superposition of crystal structures of DC-SIGN CRD in complex with different ligands shows that some binding site residues, including Phe313, which is targeted by the hydrophobic substituents of 1-4, are rather flexible.

Therefore, to get a better understanding of the binding of 3 to DC-SIGN CRD, molecular dynamics simulations of DC-SIGN CRD in complex with (R)-3 or (S)-3 in water were performed. Considering similar docking poses obtained by FRED and FlexX, the highest ranked FlexX docking pose for each ligand in complex with DC-SIGN CRD, as shown in Fig. 7, was selected to be used as an input for molecular dynamics simulation. The complexes were first solvated using TIP3P water model and were then subjected to molecular dynamics simulation, which consisted of three consecutive steps: (i) solvent equilibration for 0.5 ns with ligand and protein constrained harmonically around the initial structure; (ii) equilibration of the complete system for 1 ns with ligand and protein released; (iii) an unconstrained 10 ns production run to allow the protein and the ligand to position themselves according to physical forces between them. Physical stability of the simulations was monitored by temperature, volume and total energy vs. time plots. All these parameters of the system were stable during the production run of the simulation. Structural stability of the system during the simulation was monitored by measuring the RMSD values of protein backbone atoms during the simulation (Fig. 8a and 9a).

Interestingly, R and S isomers of 3 in complex with DC-SIGN showed significantly different behaviour during the molecular dynamics simulations. After 0.3 ns of the production run p-mannose 4-hydroxyl group of (R)-3 slowly began to lose contact with the Ca2+ ion (Fig. 8c). For the next 0.5 ns only D-mannose 3-hydroxyl group of (R)-3 was in contact with Ca²⁺ ion (Fig. 8d). Since Ca²⁺ ion was not octacoordinated anymore, the complex became unstable and the ligand left the binding site (Fig. 8e), which is also reflected by the large RMSD values of the ligand (Fig. 8b). In contrast, molecular dynamics simulation showed greater stability of the DC-SIGN CRD-(S)-3 complex (Fig. 9). Although quite high RMSD values for the ligand were observed (Fig. 9b), p-mannose moiety of (S)-3 remained tightly bound to Ca²⁺ ion and other interacting residues, as described above. Glycerol linker with the two aromatic moieties showed much greater flexibility than D-mannose. At the beginning of the production run, glycerol linker slightly rotated and oriented the phenyl ring of the (S)-3 toward Arg345, while the naphthyl ring remained in contact

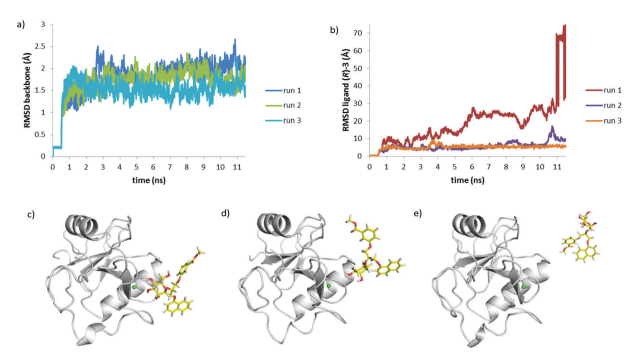


Fig. 8 (a) Backbone RMSD and (b) ligand RMSD vs. time plots for molecular dynamics simulations of DC-SIGN CRD in complex with (R)-3. Molecular dynamics snapshot (run 1) of (R)-3 in complex with DC-SIGN CRD at (c) 0.3 ns, (d) 0.8 ns and (e) 4.5 ns of the production run.

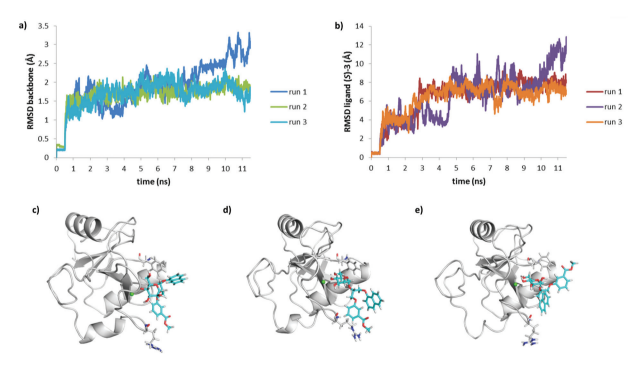


Fig. 9 (a) Backbone RMSD and (b) ligand RMSD vs. time plots for molecular dynamics simulations of DC-SIGN CRD in complex with (s)-3. Molecular dynamics snapshot (run 1) of (5)-3 in complex with DC-SIGN CRD at (c) the beginning (d) after 1.3 ns and (e) 7.5 ns of the production run.

with Phe313 side chain (Fig. 9c), as observed in the docking conformation (Fig. 7b). During the next few nanoseconds of the simulation, the naphthyl ring of (S)-3 started moving away from the Phe313 (Fig. 9d) and finally reached Asn344/Arg345

site (Fig. 9e). Such a rotation of the glycerol linker resulted also in reorientation of the phenyl ring, which left the Asn344/ Arg345 site and moved toward Phe313 (Fig. 9d and e); however, it was mainly in contact with solvent.

To evaluate whether the observed differences in (R)-3 and (S)-3 behaviour during the first MD simulation are relevant, the MD was repeated twice more for each DC-SIGN-ligand complex. In all cases the protein backbone was structurally stable, reflected by the low RMSD values (Fig. 8a and 9a). Ligand RMSD values and visual inspection of the trajectory during different MD runs of DC-SIGN-(S)-3 complex showed that the D-mannose moiety is tightly bound to Ca²⁺ and neighbouring amino acid residues, while the diaryl substituted glycerol linker finally adopts conformation similar to that presented in Fig. 9e. In contrast to the first MD simulation run of DC-SIGN-(R)-3 complex, the second and third runs showed its better stability. Towards the end of the second production run the D-mannose moiety of (R)-3 began to lose contact with Ca²⁺, which might eventually lead to dissociation of the complex, as presented in Fig. 8d and e. In the third run, the complex remained fairly stable, reflected by the relatively low RMSD values (Fig. 8b). D-Mannose residue of (R)-3 was constantly in contact with Ca2+, while the diarylglycerol moiety showed conformational flexibility.

The final pose of production run was very similar to that of (S)-3 presented in Fig. 9d, with the naphthalene ring pointing toward solvent and the phenyl ring interacting with Arg345 side chain.

Discussion

The current study presents new structural and mechanistic details at the atomic level of binding interactions of novel monovalent antagonists with DC-SIGN. Acquired STD NMR data on 1-4 unequivocally demonstrated their interactions in the binding site of the receptor. The STD group epitope mapping revealed that the mannose group formed the strongest interactions with the DC-SIGN and was in the closest contact to the receptor in the case of ligands 1-4 (Fig. 3). Individual STD contributions and interaction involvements of sugar protons were inseparable due to spectral overlap of signals between δ 3.3 and 3.9 ppm. On the other hand, anomeric H_a proton of 1-4 displayed well-resolved signals with δ between 5.0 and 5.2 ppm. Additionally, most effective STD transfer to Ha proton of 1-4 was observed among all investigated protons (i.e. H_a-H_g). Consequently, H_a proton was considered as an adequate representative of strong mannose group interaction. These results correlate well with the known binding specificity of DC-SIGN for D-mannose sugar. 23,24,31

In accord with group epitope mapping results, STD build-up studies revealed the tightest involvement of H_a (and thus mannose group) in DC-SIGN interaction. Normalized build-up curves for H_a – H_g protons of 1–4 exposed structure modification-related differences in binding modes. These curves were consistent in the case of H_a proton, indicating that binding configuration of mannose sugar was very similar for all studied ligands. The latter was confirmed for H_c proton on the naphthyl ring as well. Normalized STD build-up curves with great agreement confirmed analogous binding mode of H_c

proton regardless of structure variations of ligands. On the other hand, build-up curves of Hb proton indicated altered involvement of this proton in ligand-receptor interaction. Results suggest that H_b proton is in closer contact with the receptor in 1 and 3 in comparison to 2 and 4. Additionally, from comparison of H_{d/e} protons it is obvious that H_d of 1 was closer to the surface of the receptor with respect to H_e of 2-4. One should keep in mind that in the case of all naphthyl protons in 4 individual contributions would be markedly reduced because of overall symmetry-related STD effects of H_b, H_c and H_e protons. Although, build-up curves of H_b and H_c of 4 indicating faster rate of saturation in comparison to 1-3, maximum STD-AF were reached at lower values. Rapid saturation at low STD-AF plateau values is an indicator of fast exchange kinetics, which is in agreement with exchange properties of 4. The latter was also confirmed with the highest K_D value among 1-4 which supports our observations of lowest binding affinity of 4.

While two naphthyl constituents demonstrated low interaction tendency, substitution one of naphthyl ring with phenyl moiety resulted in moderate improvement of binding propensity. Our results indicate that a different position of the carboxylic group on the phenyl ring influences the binding conformation not only of modified part of ligand but also of intact naphthyl ring. When comparing 1 and 2 it is obvious that the position of the carboxylic group on the phenyl moiety affects the binding propensity of the ligand. Although, STDbuild up curves show that H_b and H_c protons of 1 lay closer to the receptor in comparison to 2, the opposite was observed for H_g of 1 and H_f of 2. Contribution of individual H_g proton of 1 is believed to be lower than of H_f of 2. para-Position of the carboxylic group on the phenyl ring in 1 displays better binding properties of adjacent naphthyl moiety in comparison to 2. However, shift of the carboxylic group from para- to meta-position improved binding propensity of the phenyl constituent as confirmed by STD group epitope mapping and evaluation of $K_{\rm D}$ values. Lower $K_{\rm D}$ value determined for 2 (0.96 mM) in comparison to 1 (1.6 mM) indicates better binding tendency of overall ligand. First, these results imply that, between both constituents, the phenyl moiety primarily contributes to binding affinity and plays a more important role in complex formation. And second, the carboxylic group at meta-position on the phenyl ring prolongs the half-life of ligand-receptor complex.

Furthermore, esterification of the carboxylic group at *meta*-position on the phenyl moiety additionally decreased $K_{\rm D}$ value. Notable differences in binding properties among 2 and 3 were observed not only for $H_{\rm f}$ proton but also for the whole set of protons on the naphthyl ring. $H_{\rm f}$ proton was after carboxylic group esterification closer to the surface of the receptor, since STD effect increased from 68% in 2 to 77% in 3. Greater spatial proximity was additionally confirmed with higher normalized STD-AF values for $H_{\rm f}$ of 3 in regard to 2. While $H_{\rm c}$ protons of 2 and 3 demonstrated analogous binding involvements, $H_{\rm b}$ of 3 was shown to be closer to the receptor surface. On the other hand, in the case of $H_{\rm e}$ proton of 3 we observed

less intense contact to the receptor in comparison to 2. It is evident that esterification of the carboxylic group enhanced the binding features of 3 which was reflected in lowest K_D value among 1-4 (0.45 mM). Additionally, we confirmed that esterification affected the position of the naphthyl ring in the binding site of DC-SIGN as well.

Our molecular docking and molecular dynamics simulations showed that structure-based design of novel DC-SIGN antagonists based only on docking calculations remains a great challenge and provides very limited information about the protein-ligand interactions in solution. Docking protocol is usually able to correctly predict the binding of the D-mannose moiety, which is in most cases the most tightly bound part of the DC-SIGN antagonist. Because of the rather flat DC-SIGN binding site, there are only few residues that could be targeted with the ligand to increase its affinity. In our case, aromatic moieties for interaction with the Phe313 side chain were introduced to improve the binding by possible π - π stacking/hydrophobic interactions, while carboxylates were introduced to explore possible formation of additional ionic interaction with Arg345 (see Fig. 10). While the docking calculations predicted the targeted hydrophobic interaction between the ligand and DC-SIGN (Fig. 7), molecular dynamics simulations showed that these interactions are probably quite weak given the observed flexibility of the ligands and low stability of the complexes during the molecular dynamics simulations (Fig. 8 and 9).

Therefore, taken together the results of STD NMR and molecular dynamics simulations, we can conclude that our ligands bind rather tightly through D-mannose residue. We

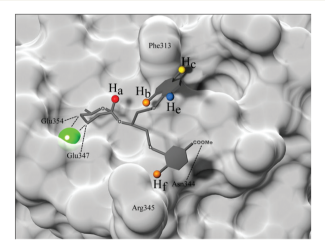


Fig. 10 Binding model of DC-SIGN-3 complex predicted based on measured STD effects and molecular modelling results. Surface of DC-SIGN CRD Ca²⁺-binding site is shown in the back (PDB code 1SL4²⁴). Ca²⁺ ion is shown as bigger green sphere, while observed ligand protons, namely H_a , H_b , H_c , H_e and H_f , are shown as smaller spheres. Ligand protons are coloured based on ligand-receptor interproton distance, red displaying the closest and blue presenting the most distant spatial proximity. Hydrogen and coordination bonds are marked with

proposed a binding model of DC-SIGN-3 complex based on combined results obtained by NMR and molecular modelling (Fig. 10). Interaction of the 3- and 4-hydroxyl groups of D-mannose with DC-SIGN CRD Ca²⁺-binding site is driven by four key bonds, namely two coordination bonds with Ca2+ ion and two H-bonds with side chains of Glu347 and Glu354. Additionally, H-bonds between other D-mannose hydroxyl groups and Asn349, Asn365 and Lys368 side chains of DC-SIGN were predicted as well. Strongest interaction of p-mannose was confirmed by highest STD effects for Ha proton, marked with red (Fig. 10). Other parts of the ligands show moderate flexibility, especially glycerol linker that enables rather free conformations and binding possibilities of attached aryl moieties. However, the naphthyl ring was detected in the spatial proximity to Phe313 where it forms hydrophobic interactions as predicted.

STD NMR results suggest that the naphthyl moiety adopts a conformation where H_b proton (orange on Fig. 10) is the closest to the receptor followed by H_c proton (yellow on Fig. 10). Weakest interaction and thus greatest receptor-ligand interproton distance was observed for H_e proton, marked with blue. On the other hand, flexibility resulting from glycerol linker allowed unforeseen engagement of phenyl group in more polar area, where it forms hydrogen bond with Asn344 and contacts with Arg345 (possible cation- π interaction) of DC-SIGN. Conformation of the phenyl entity thus apparently depends on quite prominent H_f proton-receptor proximity and formation of hydrogen bond between esterified carboxylic group and Asn344 side chain. Observation of phenyl moiety retention in polar environment of the binding site offers new insights in ligand behaviour within flat binding sites and additionally emphasizing the need for further designing of novel and more potent DC-SIGN antagonists.

Experimental

Chemistry

Detailed information is given in the ESI.†

STD experiments. STD NMR samples were prepared in 25 mM TRIS- d_{11} (pD = 8.0), 150 mM NaCl, 4 mM CaCl₂ in ²H₂O and 25% of DMSO-d₆. In all STD experiments the concentration of receptor DC-SIGN was 8 µM. DC-SIGN ECD protein (residues 66-404) were overexpressed and purified according to the already published procedure. 20,22,32 For the STD NMR studies, a pseudo-2D version of STD NMR sequence with DPFGSE water suppression was used for the interleave acquisition of on- and off-resonance spectra at 25 °C with 512 scans. For selective saturation cascades of Gaussian pulses with a length of 50 ms and 10 dB spaced by 0.1 ms delays were employed. The on-resonance irradiation was performed at 0.9 ppm; the off-resonance frequency was 30 ppm. For the protein signal suppression a 30 ms spin lock filter was used. The STD spectrum was obtained by subtraction of saturated spectra from reference spectra. The fractional STD effect was calculated by $(I_0 - I_{sat})/I_0$, where I_{off} stands for intensity of

signal in off-resonance spectrum. $I_{\rm sat}$ is intensity of signal in on-resonance spectrum. $I_{\rm off}-I_{\rm sat}$ represents the intensity of the STD NMR spectrum. The appropriate blank experiments, in the absence of the protein, were performed to assure the absence of direct irradiation of the ligand. Saturation times to obtain the STD build-up curves were 0.1, 0.3, 0.5, 0.75, 1.0, 1.25, 1.5, 2.0, 2.5, 3.0, 4.0, 5.0 s. In titration study, five STD experiments were performed with varying concentrations, 80, 240, 400, 560 and 800 μ M.

STD-amplification factor is determined according to eqn (1), in which STD-AF is a STD amplification factor. $I_{\rm off}$ stands for intensity of signal in off-resonance spectrum. $I_{\rm sat}$ is intensity of signal in on-resonance spectrum. $I_{\rm off}-I_{\rm sat}$ represents the intensity of the STD NMR spectrum. [L] is the concentration of ligand and [P] stands for the concentration of receptor.

$$STD-AF = \frac{I_{off} - I_{sat}}{I_{off}} \times \frac{[L]}{[P]}$$
(1)

The build-up curves are fitted to the monoexponential eqn (2), where STD-AF stands for the STD amplification factor of a given proton at saturation time t, STD_{max} is the maximal STD intensity obtainable when long saturation times are used, and $k_{\rm sat}$ is the observed saturation rate constant.

$$STD-AF = STD_{\max} (1 - e^{(-k_{\text{sat}}t)})$$
 (2)

The hyperbolic behaviour of the curves obtained from the plot of STD-AF as a function of ligand concentration is appropriately described by eqn (3), where STD-AF is a STD amplification factor, α_{STD} represents the maximum amplification factor, [L] is a ligand concentration and K_{D} is dissociation constant.

$$STD-AF = \frac{\alpha_{STD} \times [L]}{K_D + [L]}$$
 (3)

For the non-linear least-squares curve-fitting to the eqn (2) and (3) was used Origin 8.1 software.³³ All experimental data were presented with standard errors bars.

Molecular modelling

Computer hardware and software. All the computational work was performed on a workstation with four octal core AMD Opteron CPU processors, 16 GB RAM, two 750 GB hard drives, running 64-bit Scientific Linux 6.0.

Ligand and protein preparation. The three-dimensional models of compounds **1–4** were built in ChemBio3D Ultra 13.0.³⁴ The geometries and charges of the molecules were optimized using the MMFF94³⁵ force field and partial atomic charges assigned in ChemBio3D Ultra 13.0. The energy was minimized until the gradient value was smaller than 0.001 kcal (mol Å)⁻¹. The optimized structure was further refined with GAMESS interface in ChemBio3D Ultra 13.0, using the semi-empirical PM3 method, QA optimization algorithm and Gasteiger Hückel charges for all atoms for 100 steps.

The crystal structure of DC-SIGN CRD in complex with tetramannoside Man₄ (PDB entry: 1SL4) was taken as a receptor for

docking calculations using FlexX36,37 as available in LeadIT.38 Receptor was prepared in LeadIT graphical user interface using the Receptor wizard. Amino acid residues within a radius of 7 Å around Man4 were defined as the binding site. Hydrogen atoms were added to the binding site residues and correct tautomers and protonation states were assigned. Water molecules and the ligand were deleted from the crystal structure, while Ca²⁺ ion in the Ca²⁺ binding site was defined as part of the receptor. Pharmacophore type constraints in FlexX-Pharm were defined in order to correctly place the core mannose residue during the docking procedure. First, Pharm-Metal pharmacophore with spherical coordination around Ca²⁺ ion was defined to account correctly for complex interactions between Ca2+ ion (406) and D-mannose residue of the designed ligands. Secondly, side chain carboxylate groups of Glu347 and Glu354 were defined as hydrogen bond acceptors, which form hydrogen bonds with 3- and 4-hydroxyl groups of D-mannose residue in the crystal structure. All three selected pharmacophore type constraints were specified as essential.

For docking with FRED software (Release 3.0.1, OpenEye Scientific Software, Inc., Santa Fe, NM, USA; http://www.eyesopen.com), the protein was prepared using MAKE RECEPTOR (Release 3.0.1, OpenEye Scientific Software, Inc., Santa Fe, NM, USA; http://www.eyesopen.com). 28-30 We determined the grid box size; inner and outer contours of the grid box were calculated automatically. The outer contour size was 6195 Å³, while the grid box size was 9216 Å³. The setup of contours was set as "Balanced". Amino acid residues Asn365, Glu347 and Glu354 and Ca²⁺ ion were defined as pharmacophore type constraints. FRED requires the preparation of input conformers of each ligand prior to docking. These conformers were generated using OMEGA 2.4.6³⁹ from OpenEye Scientific Software.

Ligand docking. The FlexX molecular docking program, as available in LeadIT, was used for ligand docking. Hybrid algorithm (enthalpy and entropy driven ligand binding) was used to place the 'base fragment'. The Maximum number of solutions per iteration and the Maximum number of solutions per fragmentation parameter values were increased to 1000, while other parameters were set at their default values. Proposed binding modes and scoring function scores of the top ten highest scored docking poses per ligand were evaluated and the highest ranked binding pose was used for graphical representation in PyMOL.

Ligands 1–4 were also docked using FRED (version 3.0.1. OpenEye Scientific Software, Inc., Santa Fe, NM, USA; http://www.eyesopen.com). Docking resolution was set to high, other settings were set as default. Twenty-five docking solutions were inspected visually and the best ranked FRED-calculated conformation was used for analysis and representation.

Molecular dynamics. The molecular dynamics package NAMD (version 2.9)⁴⁰ and CHARMM22 force field⁴¹ were used for molecular dynamics simulations of DC-SIGN-ligand complexes. The FlexX best scored docking pose of compounds (R)-3 or (S)-3 in complex with DC-SIGN CRD was used as the initial structure for MD simulation. Molecular mechanics parameters for compounds (R)-3 and (S)-3 were estimated using

ParamChem tool. 42-44 Steepest descent (10 000 steps) and adopted basis Newton-Raphson (10 000 steps) energy minimizations were first performed to remove atomic clashes and to optimize the atomic coordinates of the protein - ligand complex. The structure of the energy minimized complex for MD simulation was prepared using psfgen in VMD (version 1.9.1.). 45 The complex was then embedded in a box of water, which was modelled explicitly by a TIP3P model. 46 The system was neutralized by addition of KCl at 0.4 M concentration. The MD simulation was carried out in the NPT ensemble employing periodic boundary conditions. Langevin dynamics and Langevin piston methods were used for temperature (300 K) and pressure (1 atm) control, respectively. Short- and longrange forces were calculated every 1 and 2 timesteps, respectively, with a timestep of 2.0 ps. The smooth particle mesh Ewald method47 was used to calculate electrostatic interactions. The short-range interactions were cut off at 12 Å. All chemical bonds between hydrogen and heavy atoms were held fixed using SHAKE algorithm. 48 The simulation consisted of three consecutive steps: (i) solvent equilibration for 0.5 ns with ligand and protein constrained harmonically around the initial structure; (ii) equilibration of the complete system for 1 ns with ligand and protein released; (iii) an unconstrained 10 ns production run to allow the protein and the ligand to position themselves according to physical forces between them. The trajectory of the equilibration and production run was used for analysis in VMD. The MD simulation of each complex was repeated three times.

Conclusions

In the present study we investigated the interactions between four monovalent mannose-based ligands and extracellular domain of DC-SIGN. The studied ligands, some of which are novel entities, consist of α-D-mannose connected by glycerol linker to different aryl moieties, which were expected to tune interactions of ligands in hydrophobic groove of the receptor. The combination of saturation transfer difference (STD) NMR spectroscopy and molecular modelling has been used to examine binding affinity of monovalent glycomimetics for DC-SIGN at the atomic level of resolution. Our study showed that four α-D-mannoside ligands interact with the binding site predominantly with the mannose moiety. Importantly, the other two aromatic groups that are bound to α-D-mannose through glycerol linker demonstrate interaction that can be related to their substituent pattern. Ligand with naphthyl and meta-substituted phenyl ring exhibited the most favourable binding characteristics, which might imply involvement of new contacts of substituted phenyl moiety in more polar area of flat binding site of DC-SIGN. Although our first impression was that carboxylate in the studied ligands should interact primarily with Arg345, it seems that the Asn344 residue forms H-bond with the polar group and thus dictates the main interactions with the polar substituent at phenyl ring, while Arg345 possibly forms cation– π interaction with the phenyl moiety.

Our findings point that besides lipophilic residues, possible involvement of H-bonds and/or cation-π might boost the interaction of small molecule DC-SIGN ligands and thus offer new possibilities in further designing of novel, stronger DC-SIGN antagonists.

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