The Hidden Conformation of Human Histo-blood Group Antigen is a Determinant for Recognition by Pathogen Lectins

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KEYWORDS: Oligosaccharide, Flexibility, Blood group antigen, Lectin.

ABSTRACT: Histo-blood group epitopes are fucosylated branched oligosaccharides with well-defined conformations in solution that are recognized by receptors, such as lectins from pathogens. We report here the results of a series of experimental and computational endeavours revealing the unusual distortion of histo-blood group antigens by bacterial and fungal lectins. The Lewis x trisaccharide adopts a rigid closed conformation in solution, whilst crystallography and molecular dynamics reveal several higher energy open conformations when bound to the *Ralstonia solanacearum* lectin, which is in agreement with thermodynamic and kinetic measurements. Extensive molecular dynamics simulations confirm rare transient Lex openings in solution, frequently assisted by distortion of the central N-acetyl-glucosamine ring. Additional directed molecular dynamic trajectories revealed the role of a conserved tryptophan residue in guiding the fucose into the binding site. Our findings show that conformational adaptation of oligosaccharides is of paramount importance in cell recognition and should be considered when designing anti-infective glyco-compounds.

INTRODUCTION

The current recognition of the importance of protein-glycan recognition in cellular processes is driving the efforts to elucidate the molecular basis underpinning such processes. Most carbohydrate molecules, also referred to as glycans, are considered to be flexible molecules. However, some of them, such as blood group antigens of the ABH(0) and Lewis systems (Fig. 1A), have been shown to have a well-defined conformation in solution, due to the presence of one or two fucosylated branches which restrict the number of low energy conformations that can be adopted.2 In the quest of unravelling the molecular basis dictating the recognition of carbohydrates by proteins, the present understanding is that the pre-formed conformation in solution is likely to be the bio-active one.

Despite a series of supporting evidence that the bound conformation is that occurring in solution, we raised the question whether such a hypothesis was indeed the paradigm or whether exceptions could be found. The conformations of fucosylated Lewis oligosaccharides, are considered to be rigid in solution, adopting a single shape referred to as the “Closed” conformation.24 This rigid shape is due to stacking between fucose (Fuc) and galactose (Gal) rings, by a non-conventional CH…O hydrogen bond and by steric hindrance of the N-acetyl group of GlcNAc (Fig. 1B). The crystal structure of Lewis x (Le') trisaccharide, together with NMR3 and modeling data,10,11 confirmed that the trisaccharide presents only limited conformational fluctuations around the Closed shape.

Since Le' and sialyl Lewis x (SLex') are key players in many pathologies related to inflammation, cancer and infection, and in order to understand and hence manipulate the binding process, it is fundamental to determine if the Closed conformation is maintained during interaction. Crystal structures of Le' and SLex' complexed with lectins or antibodies show the existence of the Closed conformation in the binding sites. This was also confirmed in solution by NMR with the lectins DC-SIGN13 and E-selectin.14 There is therefore a consensus for all carbohydrate-binding proteins, except for a family of fungal and bacterial lectins corresponding to the fucose-binding 6-bladed β-propeller fold. Indeed, distortion of the Le' core was reported by an NMR study of SLex' in interaction for the AAL lectin from the mushroom *Aleuria aurantia*,15 this structure being latter the first 6-bladed β-propeller lectin to be described.16 We then analyzed the conformations of the ABH and Lewis antigens when bound other β-propeller lectins from the lung pathogens *Burkholderia ambifaria* (BambL)17,18 and *Aspergillus fumigatus* (AFL1),19 respectively. While most fucosylated glycans were found in one of their low energy conformations, unexpected distortion of Le' was observed in the binding site of both lectins.

We present here a complete evaluation of the conformational behavior of Le', both in solution and in protein binding sites. In addition to the analysis of all the available complexes from lectins in the crystalline state, we describe new crystal structures of the RSL lectin from the bacterium *Ralstonia solanacearum* bound to Le' and SLex'. Several “Open” conformations of Le'
were identified in the binding sites of the β-propeller lectins. We then compared conformational dynamics of Le^x in solution to that observed in the RSL binding site. Biophysical studies were performed to characterize the thermodynamics and the kinetics of binding to this lectin. Lastly, extended molecular dynamics (MD) simulations shed light on the complete binding pathways. Overall, the present study provides unprecedented knowledge on the conformational adaptation of glycans when interacting with protein receptors. Since a precise understanding of the recognition mechanism is required to design glyco-derived compounds of therapeutic interest, the results herein have enormous potential to guide new approaches to drug design.

![Scheme 1](image)

**Scheme 1. Schematic representation of selected histo-blood group fucosylated oligosaccharides, together with convention for monosaccharide and linkages representation and different representation of the crystal structure conformation of Le^x trisaccharide with use of sticks or SweetUnityMol.**

**MATERIALS AND METHODS**

**Materials.** Recombinant RSL was produced in *Escherichia coli* as previously described and purified by affinity on mannose-sepharose column (Sigma-Aldrich). All oligosaccharides have been obtained from Elicityl (Crolles, France).

**Crystallization and structures determination.** Crystals of RSL were obtained by the hanging drop vapor diffusion method using 2 μl of drops containing a 50:50 (v/v) mix of protein and reservoir solution at 19 °C. The protein at 10 mg.ml^-1 in 20 mM Tris/HCl (pH 7.5) and 20 and 3 mM EDTA was incubated with 1h at room temperature prior to co-crystallization. The ligand was a tetrasaccharide. For the Lewis X complex, crystals plates were obtained from a solution containing 30% PEG6K and 0.28 M Tris-HCl pH 8.5 and were directly flash-frozen in a cryoloop. Diffraction data were collected at 100 K at the European Synchrotron Radiation Facility (Grenoble, France) on beamline ID144 using an ADSC Quantum Q315r detector and beamline ID23-2 using a MARCCD detector for the Le^x and Sle^x complexes respectively. The data were processed using XDS. All further computing was performed using the CCP4 suite. Data quality statistics are summarized in Table S1. The structures were solved by molecular replacement using PHASER and the trimer coordinates from 2BT9 as search model. Five percent of the observations were set aside for cross-validation analysis, and hydrogen atoms were added in their riding positions and used for geometry and structure-factor calculations. The initial model was optimized with ARP-wARP prior refinement using restrained maximum likelihood refinement in REFMAC 5.8 and iterated with manual rebuilding in Coot. Incorporation of the ligand was performed after inspection of the 2Fo–Fc weighted maps. Water molecules, introduced automatically using Coot, were inspected manually. The quality of the models was assessed using the PDB validation server (http://wwwpdb-validation.wwpdb.org/validservice/) and coordinates were deposited in the Protein Data Bank under code 5ajb and 5ajc for the Le^x and Sle^x complexes respectively.

**Titration microcalorimetry.** Recombinant lyophilized RSL was dissolved in buffer (20 mM Tris/HCl pH 7.5, NaCl 150 mM with 0.03 mM CaCl2). Oligosaccharide ligands were dissolved in the same buffer, and loaded in the injection syringe. ITC was performed with an ITC200 microcalorimeter (MicroCal Inc.) at 25°C. Titration was performed with 20 of 2 μl injections of carbohydrate ligands (1.3 to 2.0 mM) every 300 s in the lectin containing cell. Data were fitted with MicroCal Origin 7 software, according to standard procedures. Fitted data yielded the stoichiometry (n), the association constant (K_a) and the enthalpy of binding (ΔH). Other thermodynamic parameters (i.e. changes in free energy, ΔG, and entropy, ΔS) were calculated from the equation ΔG = ΔH – TΔS = -RTlnK_a, in which T is the absolute temperature and R=8.314 J mol^-1 K^-1. Two independent titrations were performed for each ligand tested. ITC figures were prepared using the Origin software provided with the apparatus, or with NITPIC and Guassi.

**Surface Plasmon Resonance.** SPR experiments were performed on a Biacore X100 instrument (GE Healthcare) at 25°C in HBS (10 mM Hepes/NaOH, pH 7.5, 150 mM NaCl, 0.05% Tween 20 and 3mM EDTA) at a flow rate of 30 μl min^-1. 354 resonance units of RSL were immobilized on a research grade CM5 chip (channel 2, 2 μg ml^-1) using amine coupling procedure; channel 1 was used as control. Experiments consisted of injection (association 180 s, dissociation 180 s) of decreasing concentration of Le^x tetrasaccharide (2-fold cascade dilutions from 200 to 0.4 μM) and H type 2 tetrasaccharide (50 to 0.05 μM). The chip was fully regenerated by two injections of 1 M fucose in running buffer for 80 s. Binding was measured as resonance units over time after blank subtraction, and data were then evaluated by using the Biacore X100 evaluation software, version 2.0.

**Molecular dynamics.** Molecular dynamics simulations were carried out using the AMBER12 program in the isothem isolar thermodynamic ensemble at 300K, using the h99SB force-field parameters for protein and GLYCAM06 for saccha-
Simulations were performed with the pmemd.cuda module using SHAKE algorithm on bonds involving hydrogen atoms. A time step of 2 fs was applied. Particle Mesh Ewald (PME) was used to handle long-range electrostatics interaction. The cutoff for non-bonded van der Waals interactions was set to 8 Å. The temperature and the pressure were kept constant using a Langevin thermostat with a collision frequency of 2 ps\(^{-1}\) and a weak coupling anisotropic algorithm with a relaxation time of 2 ps\(^{-1}\), respectively. The system was optimized in a step-wise manner. 10000 steps of minimization with 5000 steps in conjugate gradient were run with restraints of 20 kcal-mol\(^{-1}\)-Å\(^{-2}\) applied on all atoms of the complex. The minimization was then followed by 20 ps of molecular dynamics equilibration (NVT, 100K) with the same restraints. Restraints were then reduced by 5 kcal-mol\(^{-1}\)-Å\(^{-2}\) and another cycle of minimization-equilibration was made. Finally, the system was minimized for 10000 steps without applying any restraints. The system was then heated from 100 to 300 K using a Langevin thermostat with a collision frequency of 5 ps\(^{-1}\). Initial velocities were derived from a Maxwellian distribution at 100 K using a random seed for each simulation. 10 ns equilibration phase in the NVT and NPT ensemble were then run. Finally production phase was performed in the NPT ensemble at 300K.

Molecular dynamics simulations of Le\(^{\prime}\) in RSL. The crystal structure of RSL in trimeric state was considered for calculation therefore generating for each simulation three trajectories of Le\(^{\prime}\) in the inter-monomeric binding site and three in the intra binding site. In order to enhance sampling, two independents simulations were carrying out with different starting conformations of the glycan for 900 ns, resulting in 12 independent trajectories of Le\(^{\prime}\). The protonation state of each residue was assigned using H\(^{+}\) web server (http://biophysics.cs.vt.edu/H\(^{+}\)) resulting in a neutral charge of the protein. The only histidine residue (His60) was protonated on N\(_{\text{ε}}\) atom. The water phase was extended to a distance of 10 Å from any solute atom using a TIP3 water model. The system was optimized following the protocol previously described. Finally a production phase of 900 ns was carried out in NPT ensemble. In order to test the robustness of the results, we performed an additional 1 μs-long plain MD simulation starting from the crystal structure of RSL in complex with Le\(^{\prime}\) (pdb id: 5AJB), using the recent AMBER-ff14SB\(^{36}\) and the GLYCAM06-j force fields together with similar simulation parameters were used than those previously described. The Le\(^{\prime}\) molecules were considered in their conformation observed in the crystal structure, with one empty binding site.

Molecular dynamics simulations of Le\(^{\prime}\) in water. Three distinct initial conformations of the Le\(^{\prime}\) trisaccharide were selected to study its behaviour in water phase: Open I, Open III and Closed. In order to enhance the conformational sampling, 10 replicate MD simulations were carried out. For each structure, a random seed was used for velocity generation during the heating phase.

Simulating the unbinding Le\(^{\prime}\) to RSL. An Umbrella Sampling (US) protocol was applied to simulate Le\(^{\prime}\) unbinding event. Starting from a previously equilibrated structure of the complex RSL-Le\(^{\prime}\), a bias was applied between the C3 atom from fucose ring and the C atom from the Arg61 residue. The distance between the two atoms was increased from 9 Å to 15 Å by step of 0.5 Å. At each step, the distance was constrained with a harmonic potential and a force constant of 7 kcal-mol\(^{-1}\)-Å\(^{2}\). For each window an equilibration phase of 2 ns was run before a production of 20 ns, which led to a total simulation time of 396 ns. The potential of mean force (PMF) was calculated using WHAM software\(^{27}\) and standard deviation were calculated by bootstrapping analysis.

Simulating the binding Le\(^{\prime}\) to RSL. In a first attempt to model the binding event, a 1 μs-long plain MD simulation was performed starting from the apo-form of RSL surrounded by 16 Le\(^{\prime}\) molecules in solution. The AMBER-ff14SB and GLYCAM06-j force fields were used. The initial simulation box was built with 27 solvated Le\(^{\prime}\) molecules periodically distributed, and then molecules in steric clashes with the centred protein were removed. Analysis of contact between the glycans and the protein surfaces has been performed. We finally performed four targeted molecular dynamic (tMD) simulations\(^{38}\) to investigate the whole binding mechanism of Le\(^{\prime}\) into RSL binding sites, selecting two intra-monomeric and two inter-monomeric binding sites. The additional computed tMD force in comparison to a plain MD framework corresponded to the following energy term:

\[ E_{\text{tMD}} = (k/2)N\langle \text{rmsd}\rangle \]

where k is a force constant used to adjust the magnitude of the tMD force, N is a number of atoms selected for the calculation of the rmsd, and the rmsd is the mass-weighted root mean square displacement of the selected atoms in comparison to a reference structure. The reference structure was here the crystal structure of RSL-Le\(^{\prime}\) complex. As starting configuration of each simulation, a Le\(^{\prime}\) molecule was located in a random orientation at a distance of ~10 Å from the RSL binding site. We used a weak force constant of 0.1 kcal mol\(^{-1}\)-Å\(^{-2}\) in order to minimize the bias from the tMD framework. A total of 58 atoms were picked up for the calculation of the rmsd, namely the non-hydrogen atoms of the two tryptophans, the glutamate and the arginine of the binding site and those of the fucose ring of Le\(^{\prime}\). In this way, the tMD force did not affect the Le\(^{\prime}\) conformation. We note that at each step, for the calculation of the rmsd, the system was aligned on the reference structure along the backbone atoms of the protein. The AMBER-ff14SB and GLYCAM06-j force fields were used.

RESULTS AND DISCUSSION

The β-propeller lectin from R. solanacearum binds Le\(^{\prime}\) in several Open conformations. Crystals of RSL complexed with Le\(^{\prime}\) and SLε tetrasaccharides have been obtained by cocrystallization and diffracted to 1.8 and 1.7 Å resolution, respectively (see Supplementary Table 1 for statistics). The content of the asymmetric unit is a β-propeller consisting of three peptidic chains (A, B and C) and six binding sites (intra and inter monomeric) (Fig. 2A). The 3D-structure of the protein and of the fucose binding sites are in agreement with previously reported data\(^{22}\). In all sites, the fucose residue establishes hydrogen bonds to Arg, Glu, Ala and Trp residues while its methyl group locates in a hydrophobic pocket made by Trp and Ile residues.

The quality of the electron density maps allowed the location of the entire Le\(^{\prime}\) core trisaccharide in five of the binding sites of RSL/Le\(^{\prime}\) and two of RSL/SLε complexes. The whole SLε tetrasaccharide is clearly observed in one site only (Supplementary Fig. 1). While the fucose is always located with the same
orientation in the binding site, the Le’ core adopts three very different conformations, resulting in different contacts between the oligosaccharide and the protein (Fig. 2B). The conformation labelled “Open I” is observed in site A of both complexes; it presents an hydrogen bond between N-acetyl group of GlcNAc and Arg17, and only one water bridged contact for the galactose moiety. Conformation Open II is observed in site B of the complex with Le’; it brings the galactose close to the protein surface with several hydrogen bonds in particular to Trp10 and Asp32. Conformation Open III is observed in the three inter-monomeric sites of RSL/Le’ complex; the Gal and GlcNAc residues do not interact directly with the protein but are involved in a very extended hydrogen bonds network with water molecules bridging to the protein surface.

When the three conformations of the Le’ trisaccharide core are superimposed on their fucose ring in the RSL binding site, they span different regions of the large pocket above the primary binding site (Fig. 2C). All three conformations differ from the rigid Closed shape previously described. When the Closed Le’ conformation is docked in the RSL site by positioning the fucose in its canonical position, this generates a strong steric clash between the galactose residue and Trp76 or Trp 31 in intra- and inter-monomeric sites, respectively (Fig. 2D). As a Trp residue stacks to the fucose in all binding sites and as it is conserved in all lectins of the same \( \beta \)-propeller family, it may play a major role in the occurrence of the non-canonical Open conformations of Le’.

Only \( \beta \)-propeller lectins are able to open the solution conformation of Lewis oligosaccharides. The conformational analysis of Le’ in protein binding sites was first applied to the other fucose-specific \( \beta \)-propeller lectins for which crystal structures are available. Crystals of AFL have been obtained as a complex with Lewis Y (Le’ - a tetrasaccharide with two fucose residues) (code 4D4U)\(^{39}\). A new shape of the Lewis oligosaccharide, referred to as Open IV, is observed in two of the binding sites. The BambL/Lex complex (code 3ZW1)\(^{18}\) presents electron density for oligosaccharides in two sites; one corresponds to the Open III conformation and the other to a different one with distortion of GlcNAc ring in skew-boat ‘S’, which is referred as Open V. All of these observed conformations are displayed in Figure 3A using SweetUnityMol\(^{21}\) for clearer representation of the different shapes.

Oligosaccharides conformations are primarily defined by the relative orientations of their constituting monosaccharides at their glycosidic linkages. When reporting the values of torsion angles at each linkage on the corresponding potential energy map previously calculated with MM3 force-field\(^{11}\) (Fig. 3B), the Closed shape falls into the two main low energy regions of both \( \alpha \)Fuc1-3GlcNAc and \( \beta \)Gal1-4GlcNAc maps. On the opposite, all of the open conformations have at least one of their glycosidic linkage conformation lying in secondary minima. When compared to the canonical Closed conformation observed in solution, Open I is characterized by a large change of angle for both \( \alpha \)Fuc1-3GlcNAc and \( \beta \)Gal1-4GlcNAc linkages, whereas Open II and Open III are mostly distorted about the \( \beta \)Gal1-4GlcNAc one. Open IV exhibits variation for both \( \Phi \) and \( \Psi \)angles at the \( \alpha \)Fuc1-3GlcNAc linkage. Open V is not represented on the map as its GlcNAc ring is distorted, a situation that was not considered while calculating these potential energy maps.

In order to simplify the description of the different shapes, a 2D-representation was created based on the relative distance and orientation between the fucose and galactose residues. A polar coordinate system (or radar graph) has been designed with the radius \( r \) as the distance between C4Fuc and O4Gal atoms and the polar angle \( \Theta \) representing the dihedral angle between fucose and galactose ring planes (Fig. 3C). This representation enables a clear discrimination between the different shapes observed in the crystal structures. The Closed conformation with stacked galactose and fucose rings is characterized by a
small value of $r$ (<5 Å) and a dihedral angle $\Theta$ close to 0°, whereas all open conformations brings the two rings farther apart.

Figure 3. Analysis of the 60 conformations of Le$^e$ moiety from 30 crystal structures of protein complexes at resolution better than 2.5 Å (see Supplementary Table 2 and Table 3). A. Sweet-unity representation of the different shapes observed with same color coding as used in Fig. 1. B. Analysis of glycosidic linkages conformation as a function of the torsion angles $\Phi$ (O5-C1-O1-CX) and $\Psi$ (C1-O1-CX-CX+1) and superimposition on the MM3 energy map of each disaccharide 11. C. Analysis of the shape of the observed conformations using a polar coordinate system graph with the radius $r$(Å) representing the distance between C4 Fuc and O4 Gal atoms and the polar angle $\Theta$(°) representing the dihedral angle between fucose and galactose ring planes defined by C2 Gal-C5Gal-O5Gal and O5 Fuc-C5Fuc-C2Fuc, respectively.

The analysis of crystal structures of proteins complexed with Le$^e$ or Le$^e$-containing glycans (Le$^e$, Sle$^e$ and sulfo Le$^e$) was extended to all lectins and antibodies, by searching the 3D-Lectin and MAbs databases available in Glyco3D40 (http://glyco3d.cermav.cnrs.fr), and taking into account only the structures with a resolution better than 2.5 Å. Including the three β-propeller lectins described above, the search resulted in 30 crystal structures (Supplementary Table 2) of lectins and antibodies originating from animals, plants, fungi, bacteria and virus. 60 oligosaccharides could be analyzed from these crystal structures (Supplementary Table 3) and for 48 of them the Le$^e$ moiety was found to adopt the Closed conformation as represented in Figure 3C. The Open conformations are strictly restricted to the β-propeller family, and all β-propeller binding sites contain an Open conformation, except for one site of AFL1 complexed with Le$^e$. In this particular case, the second fucose of the tetrasaccharide (αFuc1-2Gal, and not αFuc1-3GlcNAc) is engaged in the binding site. This analyze confirms that all proteins except the β-propellers bind Le$^e$ in its Closed conformation, and that only the binding site of fucose-specific β-propellers, with stacking Trp residue, is able to select or to induce the Open conformations that are specifically observed for this family of lectins.

Molecular dynamics demonstrates drastic differences in the conformational behavior of Le$^e$ conformations from solution to RSL binding sites. In order to analyze the conformational dynamics of the Le$^e$ trisaccharide bound to RSL, the crystal structure of the complex was submitted to two MD simulations with explicit hydration using the AMBER12 program31 with the ff99SB force field parameters for protein32 and GLYCAM06 for saccharides33 (Fig. 4A and Supplementary Fig. 2). Simulations were started either from conformation Open III or Open I in all sites, with duration of 1 and 0.85 μs, respectively. During the simulations, no change was observed for the location of the fucose in the primary binding site (Supplementary Fig. 3), hence confirming the stability of the hydrogen bond network. On the opposite, GlcNAc and Gal residues displayed large fluctuations exploring at least two among the five Open conformations. These fluctuations were often correlated with variation of the GlcNAc ring pucker away from the 4C1 as followed by analysis of three intra-ring torsion angles41 (Supplementary Fig. 3). Therefore the variety of conformations observed in the crystal structures truly reflects the flexibility of the trisaccharide in the binding site. This conformational dynamics was also confirmed using the most recent force field AMBER-ff14SB36 with Glycam06j (Supplementary Fig. 4).

MD simulations were also performed for Le$^e$ in solution for 30 independent trajectories of 1 to 10 μs, starting either from the canonical Closed conformation, or from two of the Open conformations observed in RSL binding sites (Fig. 4B and Supplementary Fig. 5). For the shorter simulation (1 μs) starting from the Closed shape, no conformational change was observed. When the simulation started from an Open state, the conformation went to the Closed one after less than 0.2 μs and remained then stable. However, 5 simulations among the set of 30 displayed short rearrangements from Closed to Open conformations (on a ns timescale) and only one of them showed a longer stay of 0.2 μs in the open state with a sampling of different shapes (Fig. 4B). Interestingly, a twist of the GlcNAc ring was observed in most of the opening events. Overall our data show that opening of the Le$^e$ trisaccharide can also occurs in solution, but as a rare event that can be detected providing that multiple long MD simulations are performed. Such an occurrence may nevertheless be too rare to be detected by NMR methods.
Experiments demonstrate conformational enthalpy cost and shorter residence time for Le\(^\text{e}\) in RSL binding site when compared to linear oligosaccharides. In order to get experimental data on the binding mechanisms, the thermodynamics and kinetics of RSL binding to several oligosaccharides were investigated through isothermal titration micro-calorimetry (ITC) and surface plasmon resonance (SPR). The titration thermogram as displayed in Figure 5A, is in agreement with the binding of two Le\(^\text{e}\) molecules per RSL monomer, and it exhibits the classical exothermic peaks, a characteristic feature of enthalpy-driven interactions (Table 1). The affinity for RLS is in the medium range (K\(_d\) = 25.7 \(\mu\)M) and comparison with other fucose-containing glycans (Supplementary Fig. 6) indicates a higher affinity for 2’fucosyllactose (H type 5 epitope) and 2’-fucosyllactosamine (H type 2 epitope). From previous structural studies of Fuc1-2Gal-containing oligosaccharides (H type series) complexed with RSL or with the related BamB\(^\text{11,22}\), no deviation from low energy conformation is observed upon binding for these two oligosaccharides. The lower affinity of Le\(^\text{e}\) is mostly due to a less negative enthalpy of binding, whereas the entropy change is in the same order of magnitude than those measured for other oligosaccharides. From the thermodynamic analysis, the high-energy Open conformations adopted by Le\(^\text{e}\) in the binding site is reflected by an enthalpy cost, resulting therefore in a relatively lower affinity, when compared to linear unconstrained oligosaccharides.

![Figure 4](image1.png)

**Figure 4.** Selected MD trajectories of Le\(^\text{e}\). A: Two trajectories of Le\(^\text{e}\) in RSL binding site C (top) and AB (bottom). B: Two trajectories of Le\(^\text{e}\) in water. Left panel: Time evolution of rmsd (Å) from canonical Closed structure of Le\(^\text{e}\) (grey dots represent the observed values, while the black line reports the running average of the individual values). Right panel: Time evolution of Le\(^\text{e}\) shape as the function of fucose-galactose distance r (Å) and ring-planes angle Q\(\Theta\) (in °) defined as in Fig. 3 (color coding as relative evolution of time in trajectory). All trajectories are available in supp. info.

![Figure 5](image2.png)

**Figure 5.** A. Titration calorimetry of Le\(^\text{e}\) tetrasaccharide (5 mM) in cell containing RSL (172 \(\mu\)M) at 25°C with thermogram resulting from injection and fit of integrated heat. B. Comparison of SPR sensorgrams when circulating Le\(^\text{e}\) tetrasaccharide (top) and H type 2 tetrasaccharide (bottom) at various concentrations on a RSL-functionalized chip. The arrow indicates the slower dissociation phase for H type 2 compared to Le\(^\text{e}\).

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[a] from\(^\text{22}\); [b] N fixed to 2

SPR was previously performed to investigate the binding of all Lewis and ABH blood group oligosaccharides to RSL\(^\text{22}\), the measured affinities were very close to the ones reported in the present study, as measured by ITC. Another series of SPR experiments were performed with a low density RSL chip in order to reach a precise determination of the kinetics of interaction. The Le\(^\text{e}\) and Le\(^\text{a}\) branched structure could then be compared with H type 2 tetrasaccharide (Table 2). The resulting affinity
results were in agreement with the ITC measurements, and with previous literature data\textsuperscript{22}, but some differences could be observed when looking at details of kinetics (Fig. 5B). Indeed, the 13 fold affinity increase for H type 2 tetrasaccharide when compared to Le\textsuperscript{a} is mainly due to a slower dissociation rate (6 fold) and in a lesser way to a faster association rate (2 fold). Le\textsuperscript{e} appears to bind at the same speed than linear H-type 2 does, but exits much more rapidly from the site. This observation is in agreement with the strained conformation that Lewis oligosaccharides have to adopt in RSL binding site (spring effect). Whereas the present experimental data could bring some information about the process of dissociation between RSL and the constrained Le\textsuperscript{e}, it is more difficult to gain insights into the association process.

<table>
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<th>Table 2. Kinetics data obtained from SPR</th>
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Modeling the association and dissociation processes reveals the conformational rearrangement. In order to determine the mechanism involved in the distortion-associated binding of Le\textsuperscript{e} to RSL, simulations were set to follow the exit and entry pathways. The RSL/Le\textsuperscript{e} complexes are very stable and no dissociation was observed for microsecond simulations (Supplementary Fig. S3). A directed MD simulation was therefore performed with Umbrella Sampling approach in order to pull the Le\textsuperscript{e} from the site in equilibrium at each step (Fig. 6A and Supplementary Movie 1). The observed motion corresponds to the sliding of the fucose residue along the Trp indole ring, while maintaining contact with methyl group at C6 and CH groups at C4 and C5. The protein binding site did not display any conformational flexibility, except for a limited motion of the Trp-containing loop, which resulted in a slight opening of the Trp (limited to 2\textdegree). The other residues, Gal and GlcNAc, moved freely, exploring the different Open shapes of the trisaccharide. It was only after 270 ns and after the release of the fucose from the binding site that the Le\textsuperscript{e} could reach its Closed conformation. Again we observed that some conformational changes between the different Open conformations were accompanied by significant distortion of the GlcNAc ring.

The first attempts to observe the entry of Le\textsuperscript{e} in the binding site were performed throughout standard MD simulation on a system containing the RSL molecule surrounded by 16 Le\textsuperscript{e} trisaccharides located in solution far from the protein. (Supplementary Fig. 7). No complete binding of Le\textsuperscript{e} to RSL could be observed during this simulation. In some instances, some trisaccharides came in close contact to the binding site, the fucose moiety establishing stable transient interactions (20-50\textmu s) with the Trp residue. These trisaccharides remained in the Closed conformation and no opening was observed that could have yield to entry in the binding site. Given the low frequency of Le\textsuperscript{e} opening in solution, the probability to observe the complete binding process on this timescale was likely to be too low.

In order to trigger the whole binding process, targeted MD simulations were performed by driving the Le\textsuperscript{e} trisaccharide in four binding sites. This procedure was successful in all cases but one, with opening of Le\textsuperscript{e} as the Trp pushed the Gal away from the Fuc ring (Supplementary Fig. 8). In site A, the trisaccharide remained closed and could not enter completely in the site, while in sites B and AB it could open to adopt shapes Open II and Open IV with fucose stacked to Trp. A two steps trajectory could be observed in site BC with rapid opening of Le\textsuperscript{e}, allowing the fucose to establish weak contact with Trp, followed by the sliding motion along the Trp until complete entry in the site (Fig. 6B and Supplementary Movie 2).

Figure 6. A Simulations of exit and entry of Le\textsuperscript{e} in RSL binding site. A. Umbrella sampling simulation of the Le\textsuperscript{e} structure exiting the inter-monomeric AB binding site of RSL. B. Targeted MD simulation of the Le\textsuperscript{e} structure entering the inter-monomeric BC binding site of RSL. Top-left panels: time evolution of rmsd (Å) from canonical Closed structure of Le\textsuperscript{e} (grey dots represent the observed values, while the black line reports the running average of the individual values). Bottom-left panels: time evolution of Le\textsuperscript{e} shape as the function of fucose-galactose distance r (Å) and ring-planes angle (°) defined as in Figure 3 (color coding as relative evolution of time in trajectory). Top-right panels: rmsd of the non-hydrogen atoms of both the fucose ring and the binding site (Trp, Arg, Glu) aligned in comparison to the crystal structure of the Le\textsuperscript{e}-RSL complex, as a function of the simulation time. Bottom-right panels: Time evolution of the Le\textsuperscript{e} structure in vicinity of the RSL binding site. Carbon, nitrogen, oxygen atoms are represented in cyan, blue and red, respectively. For sake of clarity, hydrogens are not shown.
The entry and exit simulations therefore highlighted the role of the Trp residue in the intermediate stage: the flat indole ring surface of this aromatic amino acid is able to make first van der Waals interaction with the apolar patch on fucose. From this, the opening of the trisaccharide, often accompanied by deformation of GlcNAc, is necessary for complete entering in the binding site.

**DISCUSSION AND CONCLUSION**

The rigidity of the Le' core in solution is a widely accepted paradigm. However, a recent simulation study, in conjunction with NMR experiments, suggested that the Le' core has some flexibility, but is limited to fluctuation of the N-acetyl group orientation coupled with small variations at each glycosidic linkages. The present work therefore reports the first unambiguous experimental evidence from x-ray structure confirmed by computer simulation of the large-range flexibility of Le'. In solution, the transition events are rare and could be observed only by performing extended MD calculations. From a methodological point of view, this confirms that MD simulations of microsecond timescale are required to deconvolute the conformational behavior of oligosaccharides as previously demonstrated for N-glycan oligosaccharides and heparan sulfate fragments.

The conformational route that the oligosaccharide could use to change from the closed conformation to an open one is not obvious as the stacking of fucose and galactose limits the flexibility of each linkage. The changes in conformation has to involve either concerted motions at both linkages, or more likely, some distortion of the GlcNAc ring. Of course, the question arises as to whether the frequency of such ring distortions may depend on the parametrization of ring shape in the force-field. Nevertheless, several studies based on high resolution crystal structures of protein complexes, AFM experiments and simulations have already pointed towards alternative ring puckering besides the C1 for GlcNAc. Indeed, the distortion of GlcNAc is confirmed, as observed in the crystal structure of BambL/Le' complex where the ring adopts a S2 pucker.

From the ensemble of experimental and theoretical studies described here, it is proposed that a Trp residue of RSL and related lectins plays a crucial role in the stabilization of the open shape of Le' and guides its trajectory into the binding site. Aromatic residues play a key role in protein-carbohydrate interactions, and in particular Trp residues been found to have an enthalpy cost to European Synchrotron Radiation Facility, Grenoble, France for access and technical support to beamlines ID23-2 and ID144. MD calculations were performed on the CECIC Platform of ICMG. Technical help of Emilie Gillon is warmly appreciated as well as careful reading by Claire Doherty.

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