Extending Janus lectins architecture: Characterization and application to protocells

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Synthetic biology is a rapidly growing field with applications in biotechnology and biomedicine. Through various approaches, remarkable achievements, such as cell and tissue engineering, have been already accomplished. In synthetic glycobiology, the engineering of glycan binding proteins is being exploited for producing tools with precise topology and specificity. We developed the concept of engineered chimeric lectins, i.e., Janus lectin, with increased valency, and additional specificity. The novel engineered lectin, assembled as a fusion protein between the β-propeller domain from Ralstonia solanacearum and the β-trefoil domain from fungus Marasmius oreades, is specific for fucose and α-galactose and its unique protein architecture allows to bind these ligands simultaneously. The protein activity was tested with glycosylated giant unilamellar vesicles, resulting in the formation of proto-tissue-like structures through cross-linking of such protocells. The engineered protein recognizes and binds H1299 human lung epithelial cancer cells by its two domains. The biophysical properties of this new construct were compared with the two already existing Janus lectins, RSL-CBM40 and RSL-CBM77 Rf. Denaturation profiles of the proteins indicate that the fold of each has a significant role in protein stability and should be considered during protein engineering.

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1. Introduction

Due to their ability to crosslink various cells, such as red blood cells, lectins are originally known as agglutinins. They are generally multivalent and their selective interaction with glycoconjugates found many applications in the field of biotechnology and medicine as glycan-profiling tools [27]. Therefore, fine-tuning the specificity or the valency of lectins is a promising approach for obtaining novel tools. Synthetic biology strategies, such as the engineering of protein architecture, bring novel possibilities for lectin applications [8,13,23,24,26]. Lectin engineering is a novel domain of the synthetic glycobiology [30,32] and it can be performed at different levels, from glycan specificity to supramolecular architecture.

The trimeric oligomerization of Ralstonia solanacearum lectin (RLS) in a β-propeller shape (Fig. 1A) was recently used as a scaffold for building Janus lectins [22,25] with two faces and different specificities. The two Janus lectins, RSL-CBM40 and RSL-CBM77 Rf, were obtained by fusion of the monomeric RSL with two carbohydrate-binding modules (CBMs) with different specificity but rather similar shape, consisting of a single β-sandwich domain. This resulted in the creation of synthetic bispecific chimeras able to establish the interaction with fucose on one side and sialic acid (RLS-CBM40) or homogalacturonan (RSL-CBM77 Rf), respectively, on the other side.

Other protein domains with different specificity, but also topology diverging from the β-sandwich CBM, could be considered for building novel Janus lectins from the trimeric RSL scaffold.
β-trefoils are robust domains that contain an internal tandem forming a three-lobed architecture with almost no secondary structures [19]. They display a large variety of functions, and many show carbohydrate-binding activity. They are considered as lectins due to their multivalence, but also as CBMs (e.g., CBM13) since they sometimes occur as domains associated with glyco-active enzymes or toxins [20,21]. While their shape is well-conserved, their sequences do not show strong similarity, except for the presence of hydrophobic residues in the core and a QXW repeat found in most sequences [11]. A large number of crystal structures of β-trefoil lectins are available (212 from 63 different proteins), and based on their sequences they have been classified into 12 classes in the UniLectin3D database [2]. Datamining of genomes based on the lobe sequence signature of each class resulted in the prediction of thousands of putative β-trefoil lectins spanning all kingdoms of life [21]. The threefold symmetry results in the presence of three carbohydrate-binding sites referred to as α, β, and γ although variations in amino acid sequences sometimes result in only one or two active binding sites.

Among the known structures of β-trefoil lectin, the agglutinin from the fairy ring mushroom Marasmius oreades (MOA) presents an interesting strict specificity for αGal-containing oligosaccharides. The lectin rose attention already in the second half of the 20th century due to its high selectivity and affinity to blood group B oligosaccharide whereas very poor binding was detected to blood group A or H oligosaccharides [7,33]. MOA is specific for oligosaccharides with terminal non-reducing α1-3 linked galactose, while galactosides linked in α1-2, α1-4, and α1-6 showed no binding [33]. Due to its preference for the αGal1-3Gal disaccharide, MOA binds efficiently to blood group B (Galα1-3[Galβ1-4GlcNAc] and dimerization domain (green). Right, β-trefoil domain of MOA with different orientations (PDB code 2IHO). C) Schematic representation of Janus lectin RSL-MOA, RSL, MOAβT (β-trefoil domain of MOA), and peptide domains. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
keys due to the deactivation of the α3-galactosyltransferase during evolution [9]. Humans possess preformed antibodies directed against Gal1-3Gal, which are responsible for hyperacute rejection of animal (mainly porcine) organs in xenotransplantation attempts [4]. They are also responsible for a strong immune response against biologics such as therapeutic antibodies if produced in rodent cell cultures with active α1-3galactosyltransferase [3]. Highly selective lectins are therefore needed for the identification of such epitopes [17,29,33].

For extending the concept of Janus lectin, a fucose-specific β-propeller domain, and an α-galactose specific β-trefoil were selected (Fig. 1), first to demonstrate the possibility of utilizing a β-trefoil lectin instead of a small CBM, and second to create the dual specificity fucose/αGal that does not exist in natural or artificial glycan-binding proteins. This can be of interest in biotechnology for identifying epitopes and in synthetic biology for crosslinking cells in artificial tissues.

The crystal structure of MOA revealed that the lectin assembles as a homodimer with the monomer composed of two distinct domains, adopting β-trefoil fold at the N-terminus and α/β fold at the C-terminus (Fig. 1B). The C-terminal domain serves as a dimerization interface and retains a proteolytic function [5,15]. The lectin β-trefoil domain has a threefold symmetry, and the conserved motif (Gln-X-Trp)_3 is involved in the hydrophobic core of the structure. Co-crystals of MOA with Galα1-3Galβ1-4GlcNAc revealed that each binding site has different ligand occupancy, emphasizing the fact that slight differences in amino acids might affect the binding (Fig. 1B) [10].

In the present study, we designed a Janus lectin as a fusion protein of monomeric RSL at the N-terminus and the MOA β-trefoil domain at the C-terminus (Fig. 1C). We produced and characterized the Janus lectin RSL-MOA with double specificity toward fuco- and 1-3Gal, which are responsible for hyperacute rejection of animal (mainly porcine) organs in xenotransplantation attempts [9]. Humans possess preformed antibodies directed against Gal1-3Gal, which are responsible for hyperacute rejection of animal (mainly porcine) organs in xenotransplantation attempts [4]. They are also responsible for a strong immune response against biologics such as therapeutic antibodies if produced in rodent cell cultures with active α1-3galactosyltransferase [3]. Highly selective lectins are therefore needed for the identification of such epitopes [17,29,33].

2. Methods

2.1. Gene design and cloning

2.1.1. MOAβT

The gene for the β-trefoil domain of MOA was obtained by polymerase chain reaction where the plasmid pET-25b+RSL-MOA was used as a template. The primers, TTACATATGGAGCTTACTGAGCTG (Forward) and ATACATTTCAGTACGGTCTTACAC (Reverse) were designed to align to the full sequence of the β-trefoil domain of MOA and were ordered from Eurofins Genomics (Ebersberg, Germany). The restriction enzyme sites of Ndel and Xhol were added at 5’ and 3’ ends, respectively. Subsequently, the gene moajt and plasmid pET-TEV vector [12] were digested by RE Ndel and Xhol and ligated resulting in the pET-TEV-MOAβT vector. After transformation by heat shock in the E. coli DH5α strain, a colony screening was performed, and the positive plasmids were amplified and controlled by sequencing.

2.1.2. RSL-MOA

The original amino acid sequence of the lectin MOA from Marasmius oreades was obtained from the PDB database. The gene rsl-moa was designed as a fusion chimera with RSL at the N-terminus and the β-trefoil domain of MOA at the C-terminus via the linker PNGELSS (Supplementary Fig. 1). The gene was ordered from Eurofins Genomics (Ebersberg, Germany) after codon optimization for the expression in the bacteria Escherichia coli. The restriction enzyme sites of Ndel and Xhol were added at 5’ and 3’ ends, respectively. The synthesized gene was delivered in plasmid pEX-A2-RSL-MOA. Subsequently, plasmid pEX-A2-RSL-MOA and the pET-25b+ were digested by Ndel and Xhol restriction enzymes to ligate rsl-moa in pET-25b+. After transformation by heat shock in the E. coli DH5α strain, a colony screening was performed, and the positive plasmids were amplified and controlled by sequencing.

2.2. Protein expression

E. coli BL21(DE3) cells were transformed by heat shock with pET-TEV-MOAβT plasmid prior pre-culture in Luria Broth (LB) media with 25 μg/mL kanamycin at 37 °C under agitation at 180 rpm overnight. The next day, 10 mL of preculture was used to inoculate 1 L LB medium with 25 μg/mL kanamycin at 37 °C and agitation at 180 rpm. When the culture reached OD600 of 0.6–0.8, the protein expression was induced by adding 0.1 mM isopropyl β-D-thiogalactoside (IPTG), and the cells were cultured at 16 °C for 20 h.

E. coli KRX (Promega) cells were transformed by heat shock with the pET-25b+RSL-MOA plasmid and pre-cultured in LB media substituted with 50 μg/mL ampicillin at 37 °C under agitation at 180 rpm overnight. The following day, 10 mL of preculture was used to inoculate 1 L LB medium with 50 μg/mL ampicillin at 37 °C and agitation at 180 rpm. When reached an OD600 of 0.6–0.8, the protein expression was induced by adding 1 % rhamnose, and the cells were cultured at 16 °C for 20 h.

The cells were harvested by centrifugation at 14000 × g for 20 min at 4 °C and the cell paste was resuspended in 20 mM Tris/HCl pH 7.5, 100 mM NaCl (Buffer A), and lysed by a pressure cell disruptor (Constant Cell Disruption System) with a pressure of 1.9 kBar. The lysate was centrifuged at 24 000 × g for 30 min at 4 °C and filtered on a 0.45 μm syringe filter prior to loading on an affinity column.

2.3. Protein purification

2.3.1. MOAβT

The cell lysate was loaded on 1 mL HisTrap column (Cytiva) pre-equilibrated with Buffer A. The column was washed with Buffer A to remove all contaminants and unbound proteins. The MOA was eluted by Buffer A in steps during which the concentration of imidazole was increased from 25 mM to 500 mM. The fractions were analyzed by 12 % SDS PAGE and those containing MOAβT were collected and deprived of imidazole by dialysis in Buffer A. The N-terminal His-tag was removed by TEV cleavage with the ratio 1:50 mg of TEV:protein in the presence of 0.5 mM EDTA and 1 mM TCEP over night at 19 °C. After, the protein mixture was repurified on 1 ml HisTrap column (Cytiva) and the pure protein was concentrated by Pall centrifugal device with MWCO 3 kDa and stored at 4 °C.

2.3.2. RSL-MOA

The cell lysate was loaded on 10 mL ω-mannose-agarose resin (Merck) pre-equilibrated with Buffer A. The column was washed with Buffer A to remove all contaminants and unbound proteins and the flow-through was collected. RSL-MOA was eluted by Buffer A with the addition of 100 mM ω-mannose or 100 mM l-fucose in one step. Due to not sufficient binding capacity of the column, the flow-through was reloaded on the column several times and the protein was eluted as described previously. The fractions were analyzed by 12 % SDS PAGE and those containing RSL-MOA were collected and dialyzed against Buffer A. The protein was concentrated by Pall centrifugal device with MWCO 30 kDa and the pure protein fractions were pooled, concentrated, and stored at 4 °C.
2.4. Isothermal titration calorimetry (ITC)

ITC experiments were performed with MicroCal ITCC200 (Malvern Panalytical). Experiments were carried out at 25 °C ± 0.1 °C. Protein and ligand samples were prepared in Buffer A. The ITC cell contained proteins in a concentration range from 0.05 mM to 0.2 mM. The syringe contained the ligands in solutions in a concentration from 50 µM to 10 mM. 2 µL of ligands solutions were injected into the sample cell at intervals of 120 s while stirring at 750 rpm. Integrated heat effects were analyzed by nonlinear regression using one site binding model (MicroCal PEAQ-ITC Analysis software). The experimental data were fitted to a theoretical curve, which gave the dissociation constant (Kd) and the enthalpy of binding (∆H).

2.5. Surface plasmon resonance (SPR)

The SPR experiments were performed using a Biacore X100 biosensor instrument (GE Healthcare) at 25 °C. Biotinylated polyacrylamide-attached (PAA) sugars, such as PAA-α-fucose, PAA-α-galactose, and PAA-β-galactose (Lectinity) were immobilized on CMS chips (GE Healthcare) pre-coated with streptavidin, as previously described [25]. In the sample cell, the PAA-sugars were immobilized either as a mixture for low-density chips (1:9 of sugar of interest:non-interacting sugar) or as an individual PAA-sugar. The immobilization levels of chip CM5 LD PAA-α-fucose se: PAA-β-galactose (1:9) were FC1: streptavidin 3694 RU, PAA-β-galactose 49 RU, FC2: streptavidin 3541 RU, PAA-α-fucose: PAA-β-galactose 596 RU. The solutions were prepared at a concentration 200 µg/mL in 10 mM HEPES buffer pH 7.5 with 100 mM NaCl and 0.05 % Tween 20 (Buffer S). The reference cell was prepared in the same way as described for the sample cell, however, this time only non-interacting sugars were used. All the experiments were carried out in Buffer S. For the titration experiments, different concentrations of protein were injected onto the chip with the flow of 10 or 30 µL/min, and the protocol included steps: association time 300 s, dissociation time 300 s, 2x regeneration step for 180 s by 1 M fucose. The single-cycle kinetics experiment was carried out with the flow of 10 µL/min with an association and dissociation time of 120 s, while the regeneration was done only at the end of the run. For the inhibition assay, RSL-MAO with the concentration of 50 nM was pre-incubated with various concentrations of α-fucose for a minimum of 1 h at 4 °C and subsequently injected onto the fucose chip. The used protocol was identical to the one usage according to the manufacturers protocol. For the labeling, Biotinylated GUS (Sigma Aldrich) or Cy5 (GE Healthcare) were used. Fluorescent dyes were fluorophore-conjugated biotin (Thermo Scientific) and re-seeded for a subculture or for experiments. For experiments, cells were incubated with different concentrations of RSL-MAO for indicated time points.

2.6. Protein labeling

RSL-MAO and MOAβT were dissolved at 1 mg/mL in Dulbecco’s phosphate-buffered saline (PBS) and stored at 4 °C prior to usage. For fluorescent labeling, NHS-ester conjugated Atto488 (Thermo Fisher) or Cy5 (GE Healthcare) were used. Fluorescent dyes were dissolved at a final concentration of 10 mg/mL in water-free DMSO (Carl Roth GmbH & Co), aliquoted, and stored at −20 °C before usage according to the manufacturers protocol. For the labeling reaction, 100 µL of lectin (1 mg/mL) was supplemented with 10 µL of a 1 M NaHCO3 (pH 9.0) solution. Hereby, the molar ratio between dye and lectin was 5:1 for RSL-MAO (cell assays) and 2:1 for RSL-MAO and MOAβT for GUV assays. The labeling mixture was incubated at 4 °C for 90 min, and uncoupled dyes were separated using Zeba Spin™ desalting columns (7 kDa MWCO, 0.5 mL, Thermo Fischer). Labeled lectins were stored at 4 °C, protected from light.

2.7. Composition and preparation of giant unilamellar vesicles (GUVs)

GUVs were composed of 1,2-distearoyl-sn-glycero-3-phosphocholine (DOPC), cholesterol (both Avanti Polar Lipids, United States), Atto647N 1,2-distearoyl-sn-glycero-3-phosphoethanolamine (DOPE; Sigma Aldrich) or Atto488 1,2-distearoyl-sn-glycero-3-phosphoethanolamine (DOPE; Sigma Aldrich) and one of the following glycolipids at a molar ratio of 64:7:30:0.3:5: The glycolipids are FSL-A (Function-Spacer-Lipid with blood group A trisaccharide) (Sigma Aldrich), FSL-B (Function-Spacer-Lipid with blood group B trisaccharide) (Sigma Aldrich) or FSL-isoGb3 (Function-Spacer-Lipid with isoglobotriaosyl saccharide).

GUVs were prepared by the electroformation method as earlier described [34]. Briefly, lipids dissolved in chloroform of a total concentration of 0.5 mg/mL were spread on indium tin oxide-covered (ITO) glass slides and dried in a vacuum for at least one hour or overnight. Two ITO slides were assembled to create a chamber filled with sucrose solution adapted to the osmolarity of the imaging buffer of choice, either HBSS (for live-cell imaging) or PBS (for GUV-only imaging). Then, an alternating electrical field with a field strength of 1 V/mm was implemented for 2.5 h at RT. Later we observed the GUVs in chambers manually built as described [34].

2.8. Imaging of RSL-MAO and MOAβT binding to GUVs

Samples containing GUVs and lectins were imaged using a confocal laser scanning microscope (Nikon Eclipse Ti-E inverted microscope equipped with Nikon A1R confocal laser scanning system, 60x oil immersion objective, NA = 1.49, and four laser lines: 405 nm, 488 nm, 561 nm, and 640 nm). Image acquisition and processing were made using the software NIS-Elements (version 4.5, Nikon) and open-source Fiji software (https://imagej.net/soft-ware/fiji/).

2.9. Cell culture

The human lung epithelial cell line H1299 (American Type Culture Collection, CRL-S803) was cultured in Roswell Park Memorial Institute (RPMI) medium supplemented with 10% fetal calf serum (FCS) and 4 mM l-glutamine at 37 °C and 5% CO2, under sterile conditions. Cells were cultivated in standard TC-dishes 100 (Sarstedt AG & Co KG) until 90 % confluence, detached with trypsin (0.05 % trypsin-EDTA (1x) solution; Gibco, Thermo Fischer Scientific) and re-seeded for a subculture or for experiments. For experiments, cells were incubated with different concentrations of RSL-MAO for indicated time points.

2.10. Flow cytometry analysis

H1299 cells were detached with 2 mL of 1.5 mM EDTA in PBS, and 1 × 10^5 cells were counted and transferred to a U-bottom 96 well plate (Sarstedt AG & Co KG). To quantify protein binding to cell surface receptors, cells were incubated with different concentrations of fluorescently labeled RSL-MAO-Cy5 lectin for 30 min at 4 °C and protected from light compared to PBS-treated cells as a negative control. For the saturation of glycan-binding sites, 180 nM RSL-MAO-Cy5 was pre-incubated with 100 mM soluble α-fucose or with 100 mM 4-Nitrophenol α-α-galactopyranoside (PNPG; Sigma-Aldrich, Chemie GmbH), for 30 min at RT and in the absence of light. At the end of pre-incubation, the solution was diluted 100 times and added to cells for 30 min at 4 °C. Subsequently, cells were centrifuged at 1600 × g for 3 min at 4 °C and washed twice with FACS buffer (PBS supplemented with 3 % FCS v/v). After the last washing step, the cells were resuspended with FACS buffer and transferred to FACS tubes.
(Kisker Biotech GmbH Co. KG) on ice and protected from light. The fluorescence intensity of treated cells was monitored at FACS Gal-\textit{lios} (Beckman Coulter Inc.) and further analyzed using FlowJo V.10.5.3.

2.11. Cell proliferation (MTT) assay

Cell viability of H1299 cells upon treatment with RSL-MOA was investigated in a standard MTT assay. Cells were treated with increasing concentrations of RSL-MOA for 24 h to monitor cell proliferation or cytotoxicity. \(1 \times 10^4\) cells per well were transferred to a 96-well plate with a U-bottom. The cells were centrifuged at 1600 \(\times\) g for 3 min at RT. The cell pellet was re-suspended in 100 \(\mu L\) of variously concentrated protein solutions (9, 18, 36, 90, 180, 360, 720, 1000 nM) and transferred to a 96-well flat-bottomed plate. The cells were incubated for 24 h at 37 °C. Subsequently, 10 \(\mu L\) of MTT labelling solution (MTT Cell Proliferation Kit, Roche) was added to each well, and the cells were incubated for 4 h at 37 °C. At the end of incubation, 100 \(\mu L\) of the solubilisation reagent was added to each well, and the plate was further incubated at 37 °C overnight. The next day, the absorbance of the samples was measured at 550 nm using a BioTek microplate reader. The reference wavelength was set at 690 nm. The data was further analyzed using Microsoft Excel and GraphPad Prism software.

2.12. Statistical analysis

All data in graphs are presented as mean ± standard deviation (SD) and were calculated from the results of independent experiments. Statistical testing was performed with GraphPad Prism software and Microsoft Excel using data of \(\geq\) 3 biological replicates. Statistical differences in independent, identical samples were determined with a two-tailed, unpaired \(t\)-test. Non-significant results are not highlighted.

2.13. Differential scanning calorimetry (DSC)

The DSC experiments were carried out in Micro-Cal PEAQ DSC instrument (Malvern Panalytical). The protein samples were prepared in Buffer A, which could be substituted with the addition of a ligand. The sample cell contained the protein solutions in the concentration range of 9–29 mM. The reference cell contained the same buffer as present in the sample but without protein. The increase in temperature was measured from 20 to 130 °C with a scan rate of 200 °C/min. The data were analyzed by Micro-Cal PEAQ DSC software with a non-two-state and progress baseline method fitting model.

3. Results

3.1. Design and production of the \(\beta\)-trefoil domain of MOA (MOA\(\beta\)T)

MOA occurs naturally as a dimer, assembled by a strong association between the two C-terminal domains. Monomeric MOA composed only of the \(\beta\)-trefoil domain would be of interest for biotechnology applications. The gene of the MOA \(\beta\)-trefoil domain was defined as the 156 N-terminal AAs of MOA full sequence and was amplified by PCR and subcloned into the vector PET TEV as a fusion with an N-terminal His-tag sequence. The resulting protein was named MOA\(\beta\)T and was recombinantly produced in soluble form in the bacterium \textit{E. coli} BL21(DE3). The protein was purified by immobilized metal ion chromatography followed by His-tag cleavage by TEV (tobacco etch virus) protease. The resulting protein has an estimated molecular weight of 17.2 kDa. TEV cleavage followed by additional immobilized metal ion chromatography reduced the sample contaminants and the protein purity was verified by 12 % SDS PAGE electrophoresis (Supplementary Fig. 2A).

3.2. Design and production of Janus lectin RSL-MOA

Janus lectin RSL-MOA was designed as a gene fusion of monomeric RSL (\(N\)-terminus) and \(\beta\)-trefoil domain of lectin MOA (\(C\)-terminus). The fusion of the two protein domains by the eight AAs linker PNGELLSS results in the protein sequence of 255 amino acids (Supplementary Fig. 1). The gene sequence was optimized for bacterial expression, and the synthesized gene was subcloned into the plasmid pET25b+. The protein was produced in soluble form in the bacterium \textit{E. coli} KRX. The protein was subsequently purified by affinity chromatography on an agarose-mannose column due to the interaction between the RSL domain and mannose residues. Protein analysis by 12 % SDS PAGE electrophoresis showed that Janus lectin RSL-MOA has an apparent size of 83 kDa, which corresponds to a trimer. RSL oligomerization appears to be resistant to denaturation conditions. A smaller amount of dimers and monomers were also visible, probably because of partial oligomer denaturation (Supplementary Fig. 2B).

3.3. Biophysical characterization of MOA\(\beta\)T and RSL-MOA by isothermal titration calorimetry (ITC)

MOA\(\beta\)T affinity towards Gal\(1\text{-}3\)Gal disaccharide, the terminal disaccharide of Gal\(\text{lii} epitope, was assayed by titration microcalorimetry. Large exothermic peaks were obtained at the beginning of the titration (Fig. 2). The affinity was not strong enough to obtain a sigmoidal curve, and therefore, as recommended in such cases [31], the stoichiometry (\(N\)) was fixed, using a value of \(N = 2\) since, as previously shown by Grahn et al., [10], MOA binding sites have different affinities, and we expected at least two of them to be active. A dissociation constant (\(K_d\)) of 150 \(\mu M\) was obtained, which confirms the functionality of the isolated \(\beta\)-trefoil domain, and is in good agreement with the \(K_d\) measured for the whole MOA protein, i.e., 182 \(\mu M\) [33] (Fig. 2A).

RSL-MOA was designed to possess six binding sites for fucose and up to nine binding sites for \(\alpha\)-galactose on the opposite faces. The functionality of both binding interfaces was tested by various biophysical approaches. To this end, we designed an ITC experiment with a consecutive injection of both ligands. First, Gal\(1\text{-}3\)Gal was titrated into the cell containing RSL-MOA. Subsequently, the cell content from the first experiment (complex RSL-MOA/ Gal\(1\text{-}3\)Gal) was titrated by \(\alpha\)-methyl fucoside (\(\alpha\)-MeFuc) (Fig. 2B).

The thermogram obtained by titrating Gal\(1\text{-}3\)Gal in RSL-MOA solution was very similar to the one obtained for MOA\(\beta\)T. The dissociation constant (\(K_d = 226 \mu M\)) was comparable to the \(K_d\) for the isolated \(\beta\)-trefoil domain (as described above) and for the whole MOA [33]. The subsequent titration by \(\alpha\)-MeFuc resulted in a sigmoid shape, due to stronger affinity, with a measured stoichiometry of \(N = 2\), corresponding to the presence of two fucose binding sites per RSL monomer. The \(K_d\) was measured to be 0.4 \(\mu M\), in excellent agreement with the previously measured affinity (0.7 \(\mu M\)) for RSL [16]. This experiment proved that both parts of RSL-MOA are functional and able to bind their ligands at the same time.

3.4. Biophysical characterization of RSL-MOA by surface plasmon resonance

To evaluate the effect of multivalency and therefore measure avidity instead of affinity, surface plasmon resonance (SPR) was used. Streptavidin CM5 chips were functionalized by different
ligands, i.e., biotinylated PAA-α-fucose, PAA-α-galactose, and PAA-β-galactose.

No binding of RSL-MOA could be observed with α-galactose chip, even with high density surface (200 µg/mL biotinylated PAA-α-Gal), probably because of its very low affinity for the monosaccharide α-galactose ($K_D = 8$ mM) [33]. On the opposite, RSL-MOA bound efficiently to fucosylated chips. In order to avoid mass transfer [6], low density (LD) CM5 chip was prepared with mixture of PAA-α-fucose/PAA-β-galactose in the ratio 1:9 (200 µg/mL) and RSL-MOA was injected in increasing concentrations (5 to 1000 nM). The regeneration step was performed between each injection using 1 M fucose solution. As shown in Fig. 3A, a dose-dependent response was obtained, with a steep association phase and very weak dissociation event. The steady-state analysis showed that even at the highest protein concentration (1000 nM) the chip surface was not saturated by RSL-MOA, i.e., the plateau phase was not reached (Fig. 3A). Nonetheless, the fitting of the kinetics constant could be performed and values for affinity ($K_D = 4.9 \times 10^{-10}$ M) and kinetics ($k_{on}$ and $k_{off}$) were estimated (Table 1).

As an alternative, a single-cycle kinetic experiment was performed. RSL-MOA was injected at four concentrations (10 to 500 nM) (Fig. 3B). The advantage of this method is that no regeneration step is needed. The estimated $K_D$ of $4.9 \times 10^{-12}$ M was two orders lower if compared to the titration experiment; however, such a variation is acceptable in binding events characterized by avidity. In order to evaluate the capacity of monosaccharides to inhibit the multivalent binding and compete with protein binding to the chip surface, 50 nM RSL-MOA was pre-incubated with various concentrations of fucose (5 to 5000 µM). As shown in Fig. 3C, the complete inhibition was achieved in the presence of high concentrations of fucose (2500 and 5000 µM), and a IC$_{50}$ value of 7.8 µM was obtained.

The ability of RSL-MOA to bind the glycan-decorated surface of SPR chips was confirmed by several experimental setups. The variations in the affinity constant just illustrate the difficulty to quantify binding when avidity is the major event. These results are in agreement with previous ones evaluating RSL avidity for the fucose chip [1]. This avidity constant is at least 1000-fold higher than the affinity measured for fucose monosaccharide in solution, confirming the very strong cluster effect resulting from the topology of the RSL β-propeller with the presentation of six binding sites on the same face, binding very efficiently to fucose presented in a multivalent manner on a surface.

3.5. MOAβT binds, while RSL-MOA binds and crosslinks differentially glyco-decorated giant unilamellar vesicles

The binding properties of MOAβT and RSL-MOA were tested with glyco-decorated giant unilamellar vesicles (GUVs). The GUVs were fluorescently labelled with Atto647N- or Atto488-DOPE lipid and contained functional spacer lipid (FSL) with different terminal saccharides attached, i.e., FSL-isoglobotriose (iGb3), FSL-A, and FSL-B carrying Galα1-3Galβ1-4Glc, blood group A and blood group B terminal trisaccharides, respectively (Fig. 4). These glyco-decorated GUVs were incubated with MOAβT and RSL-MOA lectins, which were both fluorescently labeled with same Atto488 fluorophore (Fig. 4 A-D).

MOAβT-Atto488 (500 nM, green) showed almost no binding to FSL-iGb3-containing GUVs (red) (Fig. 4A). While, on the contrary,
b MOAb4T-Atto488 bound FSL-B-containing GUVs and induced the tubule-like structures (Fig. 4B) indicating its ability to bind glycan structures on the membranes.

The Janus lectin RSL-MOA-Atto488 (green) binds efficiently to the surface of FSL-iGb3-GUVs (red), even at a concentration of 200 nM (Fig. 4C). The difference is striking when compared to the absence of binding of MOAb4T-Atto488 (500 nM) (Fig. 4A).

The super-multivalency resulting from the presence of three β-trefoil domains, displaying a total of nine possible αGal binding sites, makes a very strong difference in terms of ability to bind to the glycosylated surfaces.

The capacity of RSL-MOA to crosslink GUVs carrying different oligosaccharides was tested using two populations of fluorescently labeled vesicles, FSL-iGb3-GUVs (labelled with Atto647N-DOPE, red) and FSL-A-GUVs (labelled with Atto488-DOPE, green). When the unlabeled RSL-MOA was incubated with the liposomes, cross-linking was observed between red and green GUVs, as well as, cross-linking between GUVs of the same color. The cross-linking between the same population GUVs is possibly due to lectin topology and multivalency, β-trefoil of MOA, and β-propeller of RSL. Herein, due to the double specificity of lectin RSL-MOA, we confirmed that multivalent and two-site-oriented topology has

**Table 1**

| Experimental condition                        | $k_{on}$ (1/Ms) | $k_{off}$ (1/s) | $K_D$ (M) | $R_{MAX}$ (RU) | Chi² |
|----------------------------------------------|-----------------|-----------------|-----------|----------------|------|
| Chip CMS PAA-α-fuc LD Classical kinetics     | 4.42*10⁷        | 2.08*10⁻⁷       | 4.9*10⁻¹⁰ | 3030           | 8.23 |
| Chip CMS PAA-α-fuc LD Single cycle kinetics | 1.84*10⁴        | 1.09*10⁻⁷       | 5.9*10⁻¹² | 1960           | 40.4 |

Fig. 3. SPR sensorgrams of Janus lectin RSL-MOA on CMS-PAA-α-fuc LD chip. A) Titration experiment of different concentrations of RSL-MOA (blue) and fitting curves (1:1 binding fit) (black) (left) and steady-state analysis of RSL-MOA titration (right). B) Single-cycle kinetics of RSL-MOA with concentrations of 10, 50, 250, and 500 nM. C) Inhibition of 50 nM RSL-MOA by various concentration of fucose. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
3.6. RSL-MOA binds to human epithelial cells

The ability of RSL-MOA to recognize and bind to glycans on the surface of human cells was investigated by flow cytometry. RSL-MOA was fluorescently labeled with a Cy5 dye (RSL-MOA-Cy5) and incubated with the non-small cell lung cancer cell line H1299, for 30 min at 4°C. As depicted in the histograms of fluorescence intensity (Fig. 5A), cells treated with increasing concentrations of RSL-MOA-Cy5 (0.07 – 0.7 μM) showed a dose-dependent binding of the lectin to the cell surface. In order to confirm lectin specificity and glycan-driven binding to receptors at the plasma membrane of H1299 cells, a number of inhibition assays was performed (Fig. 5B, C). RSL-MOA-Cy5 was pre-incubated with 100 mM fucose or 100 mM synthetic analogue of α-Gal epitope, p-nitrophenyl-α-D-galactopyranoside (PNPG), respectively. The high concentrations of ligands were chosen due to previous observations with lectin RSL, for which a complete inhibition of binding to fucosylated receptors on the surface of H1299 cells was achieved only in the presence of 100 mM fucose [28].

Fig. 4. The binding properties of MOAβT and RSL-MOA to glyco-decorated GUVs. A) 500 nM MOAβT-Atto488 (green) shows almost no binding to the FSL-iGb3-GUVs (red, labelled with fluorescent lipid Atto647N-DOPE). B) 500 nM MOAβT-Atto488 (green) binds to FSL-B-GUVs (red, labelled with fluorescent lipid Atto647N-DOPE). C) 200 nM RSL-MOA-Atto488 (green) binds to FSL-iGb3-GUVs (red, labelled with fluorescent lipid Atto647N-DOPE). D) Unlabeled 200 nM RSL-MOA binds two different populations GUVs FSL-iGb3-GUVs (red, labelled with fluorescent lipid Atto647N-DOPE) and FSL-A-GUVs (green, labelled with fluorescent lipid Atto488-DOPE) and crosslinks them. The GUVs were composed of DOPE, cholesterol, glycolipid of choice, and membrane dye to the proportion of 64.7:30.5:0.3 mol%, respectively. Scale bars are 10 μm. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
The presence of 100 mM fucose significantly decreased RSL-MOA-Cy5 binding to treated cells by approx. 50% compared to unblocked RSL-MOA-Cy5 (Fig. 5B), but did not abolish it entirely. Increasing the concentration up to 500 mM of fucose did not result in a further decrease in protein binding. The interaction of the engineered lectin with the cell surface was therefore not completely inhibited upon saturation of the fucose-binding sites, suggesting that the MOA domain partially compensates for the binding of Janus lectin to H1299 cells. On the other hand, 100 mM PNPG had almost no effect on RSL-MOA-Cy5 binding to treated cells, as indicated by the histograms in Fig. 5C. Fluorescence of cells incubated with RSL-MOA-Cy5 or RSL-MOA-Cy5 pre-treated with PNPG exhibited a similar intensity, as illustrated by the overlapping histograms (Fig. 5C). Reasonably, inhibition of protein binding was not achieved due to the low affinity of MOA toward monosaccharides [33].

These results suggest that both RSL and MOA domains of Janus lectin are able to bind to glycans present on H1299 cancer cells. Furthermore, we evaluated the potential cytotoxicity induced by RSL-MOA in treated cells. As illustrated in Supplementary Fig. 3, cell viability and proliferation were preserved at 24 h post-treatment with increasing concentrations of RSL-MOA (in the range 0 – 1 μM), confirming the potential of this tool in biomedical research and applications.

3.7. Comparison of thermal stability of Janus lectins and their distinct domains

Janus lectins are created by assembling protein domains with different biophysical properties. It is therefore of interest to evaluate the effect of the fusion of these modules, and also to compare the biophysical properties of Janus lectins.

The thermal stability of RSL-MOA and its constituting domains was measured with differential scanning calorimetry (DSC). Thermograms of MOAβT and RSL are displayed in Fig. 6. These two modules have very different thermal stability with a denaturation/midpoint temperature (Tm) of 42 °C for the MOAβT and 92 °C (with two Tm at 90.5 °C and 93.8 °C) for the β-propeller RSL. It could be hypothesized that MOAβT is easily denatured as it represents only a single domain of the whole original protein. The very strong thermal resistance of RSL is related to the compact and robust architecture of the β-propeller, and the two slightly different Tms extracted from the curve may correspond to the dissociation of the trimer followed by the unfolding of each monomer. As expected, adding α-MeFuc as a ligand to the protein solution increased the Tm of RSL to values of approximately 100 °C and 103 °C, for sugar concentrations of 125 and 250 μM, respectively.

The thermal unfolding of RSL-MOA and other Janus lectins was carried out in the presence of a high concentration of fucose since the protein was tested after the purification procedure. RSL-MOA displayed two events of denaturation at very different temperatures, i.e., 40 °C and 115 °C. From the results obtained on the separated domains, it is clear that the lower Tm corresponds to the denaturation of RSL. The Tm of the MOA moiety is not different when isolated or linked to RSL. The Tm of the RSL moiety is higher than for the β-propeller alone, which is probably due to the high concentration of fucose in the medium.

Other Janus lectins were also assayed by DSC for comparison. The proteins RSL-CBM40 and RSL-CBM77_Rf were obtained as described previously [22,25] and the buffer was supplemented by fucose as in the case of RSL-MOA. As already experienced with RSL-MOA, two separate events of denaturation were observed (Fig. 6) (Table 2). The high-temperature Tm, corresponding to the denaturation of RSL, is observed, with a value as high as 121 °C for RSL-CBM40. CBM77_Rf and CBM40 have a similar β-sandwich fold, and they present rather similar melting temperatures of 67 °C and 65 °C, respectively. This value is more than 25 °C higher than the one observed for MOA, indicating a clear difference in thermal stability between β-sandwich and β-trefoil.
4. Discussion and conclusion

In this work, we extended the concept of Janus lectin and therefore developed a universal strategy for increasing lectin valency and introducing an additional specificity [22,25]. Janus lectins are engineered as fusion chimeras and due to the presence of the lectin RSL, these synthetic proteins assemble as trimers, resulting in the multiplication of lectin binding sites and thus expected higher affinity toward ligands. The first two Janus lectins, RSL-CBM40 [25,28] and RSL-CBM77 Rf [22] were designed as fusion chimeras of a lectin and CBM domain with a possible application as drug carriers or protein crosslinker in plant cell wall engineering, respectively. However, here we propose an alternative where the Janus lectin RSL-MOA is composed of two individual lectin domains, β-trefoil from lectin MOA and β-propeller of RSL.

The biophysical properties of RSL-MOA were compared with MOAβT, the engineered β-trefoil domain of MOA lectin. Both proteins showed the same affinity toward ligands in solution (ITC experiments) confirming their activity. However, their behavior resulted differently when tested with glyco-decorated liposomes (GUV experiments). We observed that MOAβT showed almost no binding to FSL-iGb3-GUVs while RSL-MOA interacts with such vesicles and even mediates their crosslinking. These observations are probably conditioned by the super-multivalency of engineered RSL-MOA that displays nine sugar-binding sites, instead of only three in MOAβT. On the other hand, MOAβT displayed active binding with blood group B GUVs (FSL-B-containing GUVs) confirming that the lectin domain is able to bind glycoconjugates on the membrane surface. RSL-MOA also retains the ability to bind two differently glyco-decorated GUVs and crosslink them. This confirms that the engineered topology does not affect protein binding and such a lectin is of interest for the development of a highly selective tool for the construction of proto-tissues.

The SPR analysis of RSL-MOA revealed the strong dose-dependent binding of the β-propeller RSL toward low-density PAA-α-Fuc CM5 chips. Even though almost no dissociation event was observed, the fitting procedure allowed for the estimation of binding constants. Depending of the experimental design, some variations are observed in affinity and kinetics, due to the strong avidity of the system, generating complexity. Nonetheless,

![Image](MOAβT, RSL, RSL-CBM40, RSL-CBM77 Rf with or without ligands. The figure displays experimental conditions, fitted data, and schematic representation of proteins and ligands.)

**Table 2**

| Protein     | Ligand       | Protein (g/L & μM) | Tm1 (°C) | Tm2 (°C) |
|-------------|--------------|--------------------|----------|----------|
| MOAβT       | –            | 0.5 (29)           | 41.8     | –        |
| RSL         | 125 μM αMeFuc| 0.25 (25)          | 90.5     | 93.8     |
| RSL         | 250 μM αMeFuc| 0.25 (25)          | 98.9     | 102.5    |
| RSL-MOA     | 100 mM Fuc   | 0.25 (9)           | 101.6    | 105.3    |
| RSL-CBM77 Rf| 100 mM Fuc   | 0.5 (18)           | 39.5     | 114.5    |
| RSL-CBM40   | 100 mM Fuc   | 0.5 (16)           | 67.4     | 116.6    |
RSL-MOA showed at least 1000 higher affinity for surface-presented oligosaccharides compared to the solution state, supporting the fact that the addition of MOA does not affect its binding properties.

The ability of RSL-MOA to recognize glycans exposed at the surface of H1299 cells was analyzed by flow cytometry, demonstrating a strong dose-dependent binding. H1299 cells are characterized by the presence of highly fucosylated glycoconjugates, among others, on their surface [14]. Furthermore, the effect of saturation of RSL-MOA sugar-binding sites with 100 mM fucose or 100 mM PNPG, was investigated and resulted in a decrease in the interaction between lectin and cells. We believe that the presence of the galactose-specific MOA domain partially compensates for the inhibition of fucose-dependent interaction between the lectin and the cell surface-exposed glycans. Our hypothesis is supported by previous observations with the lectin RSL, which showed similar binding properties to H1299 cells that could be fully inhibited by 100 mM fucose. On the other hand, 100 mM PNPG did not affect RSL-MOA binding, suggesting that the synthetic analogue of α-galactose (PNPG) did not bind MOA with sufficient affinity, as already observed previously with α-galactose monosaccharide [33] and confirmed by SPR observations in the present work. Additionally, RSL-MOA does not show any cytotoxicity toward H1299 cells making it a suitable candidate for the further development in biomedical applications. However, the probable ability of this lectin to agglutinate erythrocytes is a limitation for in vivo use, and the main application would be as a research tool, before further developments are carried out.

The thermal stabilities of three Janus lectins, i.e., RSL-MOA, RSL-CBM40, and RSL-CBMTM7a, were compared by DSC. During protein denaturation, two events of unfolding are observed for all Janus lectins implying the fact that each protein domain has a different stability. Structurally similar CBMs (β-sandwich fold) share almost the same Tm, while β-trefoil is more thermally unstable. On the other hand, the β-propeller RSL showed extremely high Tm, whereas the addition of the ligand has a tremendous effect on its stability. Our findings, therefore, suggest that the structural fold of the individual domains should be taken into account when designing novel Janus lectins if stability is to be considered.

Until now, three Janus lectin with different specificities and architecture have been engineered. Therefore, we are confident that with this strategy, the novel bispecific lectins with improved valency can be obtained and find their applications in numerous fields of biotechnology and biomedicine.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jsb.2022.11.005.

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