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Purification, Biochemical Characterization, and Amino Acid Sequence of a Novel Type of Lectin from *Aplysia dactylomela* Eggs with Antibacterial/Antibiofilm Potential

Rômulo Farias Carneiro¹ · Renato Cézar Farias Torres¹ · Renata Pinheiro Chaves¹ ·

Mayron Alves de Vasconcelos² · Bruno Lopes de Sousa³ ·

André Castelo Rodrigues Goveia¹ · Francisco Vassiliepe Arruda² ·

Maria Nágila Carneiro Matos¹ · Helena Matthews-Cascon⁴ · Valder Nogueira Freire³ ·

Edson Holanda Teixeira² · Celso Shiniti Nagano¹ · Alexandre Holanda Sampaio¹

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Abstract A new lectin from *Aplysia dactylomela* eggs (ADEL) was isolated by affinity chromatography on HClactivated SepharoseTM media. Hemagglutination caused by ADEL was inhibited by several galactosides, mainly galacturonic acid (Ka = $6.05 \times 10^6 \text{ M}^{-1}$). The primary structure of ADEL consists of 217 residues, including 11 halfcystines involved in five intrachain and one interchain disulfide bond, resulting in a molecular mass of 57,228 ± 2 Da, as determined by matrix-assisted laser desorption/ionization time of flight mass spectrometry. ADEL showed high similarity with lectins isolated from *Aplysia* eggs, but not with other known lectins, indicating that these lectins could be grouped into a new family of animal lectins. Three glycosylation sites were found in its polypeptide backbone. Data from peptide-N-

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Alexandre Holanda Sampaio alexholandasampaio@gmail.com

- ¹ Laboratório de Biotecnologia Marinha BioMar-Lab, Departamento de Engenharia de Pesca, Universidade Federal do Ceará, Campus do Pici s/n, bloco 871, Av. Mister Hull, Box 6043, Fortaleza, Ceará 60440-970, Brazil
- ² Laboratório Integrado de Biomoléculas LIBS, Departamento de Patologia e Medicina Legal, Universidade Federal do Ceará, Monsenhor Furtado, s/n, Fortaleza, Ceará 60430-160, Brazil
- ³ Departamento de Física, Universidade Federal do Ceará, Campus do Pici s/n, bloco 871, Fortaleza, Ceará 60440-970, Brazil
- ⁴ Laboratório de Invertebrados Marinhos do Ceará LIMCE, Departamento de Biologia, Universidade Federal do Ceará, Campus do Pici s/n, bloco 906, Fortaleza, CE 60455-760, Brazil

glycosidase F digestion and MS suggest that all oligosaccharides attached to ADEL are high in mannose. The secondary structure of ADEL is predominantly β -sheet, and its tertiary structure is sensitive to the presence of ligands, as observed by CD. A 3D structure model of ADEL was created and shows two domains connected by a short loop. Domain A is composed of a flat three-stranded and a curved five-stranded β sheet, while domain B presents a flat three-stranded and a curved four-stranded β -sheet. Molecular docking revealed favorable binding energies for interactions between lectin and galacturonic acid, lactose, galactosamine, and galactose. Moreover, ADEL was able to agglutinate and inhibit biofilm formation of Staphylococcus aureus, suggesting that this lectin may be a potential alternative to conventional use of antimicrobial agents in the treatment of infections caused by Staphylococcal biofilms.

Keywords Sea hare · Lectin · Galacturonic acid · Biofilm · Mass spectrometry

Introduction

Lectins are sugar-binding proteins that possess the ability to decipher the glycocode. They have been found in all living organisms. In mollusks, lectins have been detected and purified from cephalopods, gastropods, and bivalves (Alpuche et al. 2010; Bulgakov et al. 2004; Fujii et al. 2011; Song et al. 2011). Among the prospected lectins representatives of some structural families are found, such as C-, F-, and R-types; galectins; and RBLs (Chen et al. 2011; Fujii et al. 2012; Naganuma et al. 2006; Pales-Espinosa et al. 2010; Song et al. 2011). The presence of lectins in mollusks has been associated with hemolymph, mucus, hemocytes, and reproductive tissues (Chen et al. 2011; Gilboa-Garber et al. 1985; Ito et al. 2011; Zhang et al. 2014), and evidences suggest that they can act as pattern recognition receptors (PRRs), recognizing several pathogen-associated molecular patterns (PAMPs). In fact, lectin-coding genes have been found be upregulated after infection by pathogenic microorganisms (Chen et al. 2011; Mu et al. 2014). Moreover, some studies have explicitly demonstrated the antibacterial potential of lectins isolated from mollusks (Zheng et al. 2008; Takahashi et al. 2008; He et al. 2011).

Gastropoda is the most broadly distributed class of mollusk, and several lectins from gastropods have been isolated (Fujii et al. 2011; Gilboa-Garber et al. 1985; Kawsar et al. 2009). *Aplysia dactylomela* Rang, 1828 is a circumtropical aplysid gastropod and occurs worldwide in tropical to warm temperate waters (Matthews-Cascon et al. 2011). In the West Atlantic, this species is found in Florida, Mexico, the Caribbean, and Brazil. It is the biggest opisthobranch found in Northeastern Brazil and one of the main herbivores inhabiting consolidated intertidal bedrock. The egg mass of an *A. dactylomela* is a resistant gelatinous string that can reach 8 m in length. Egg mass color is variable, reaching tones of yellow, orange, and pink (Matthews-Cascon et al. 2011).

Currently, only a fetuin-binding lectin named dactylomelin-P has been isolated and characterized in some detail from the purple gland of *A. dactylomela* (Melo et al. 2000).

In the present work, we have isolated, characterized, and determined the primary structure and estimated the secondary structure of a new lectin from the eggs of *A. dactylomela*. Additionally, we created a 3D model and evaluated its antibacterial and antibiofilm potentials.

Material and Methods

Egg Collection and Extraction

Aplysia dactylomela eggs were collected during low tide in the intertidal zone at Pacheco Beach (3° 41′ 13.0″ S; 38° 38′ 21.7″ W), Caucaia, Brazil. Collections were authorized and certified by responsible environmental authorities (SISBIO ID: 33913-8).

Eggs were removed from stones and algae, washed with seawater, and transported to the lab. Immediately thereafter, eggs were crushed using mortar and pestle. Proteins from smashed eggs were extracted with ten volumes of 20 mM Tris-HCl, pH 7.6, containing 150 mM NaCl and phenylmethylsulfonyl fluoride (PMSF) 0.1 mM (Tris-buffered saline (TBS)). The mixture was then centrifuged at $8000 \times g$ for 20 min at 4 °C. The crude extract was stored at -20 °C until use. Protein content was determined by Bradford reagent, and BSA was used as protein standard (Bradford 1976).

Lectin Purification

The crude extract was applied to an HCl-activated SepharoseTM 4B column (1.0×8.0 cm), previously equilibrated with TBS. Unbound proteins were washed with the same buffer, and adsorbed proteins were recovered with TBS containing 200 mM α -lactose. Chromatography eluates were monitored by absorbance at 280 nm using a UV/Visible spectrophotometer. Retained fractions containing the purified lectin were exhaustively dialyzed against distilled water, freezedried, and stored at -20 °C until required.

Hemagglutinating Activity and Inhibition Assays

Hemagglutinating activity (HA) and inhibition assays were performed following pre-established methods (Sampaio et al. 1998), using human (A, B, and O) and rabbit erythrocytes, both treated with proteases and untreated.

The following sugars and glycoproteins were used in the inhibition assay: D-xylose, D-ribose, L-fucose, L-arabinose, Lrhamnose, D-galactose (Gal), D-mannose, D-glucose (Glc), Dglucosamine, D-galactosamine, *N*-acetyl-D-glucosamine (GlcNAc), *N*-acetyl-D-galactosamine (GalNAc), D-galacturonic acid, D-fructose, D-sucrose, D-melibiose, α -D-lactose, β -D-lactose, D-lactulose, D-maltose, D-raffinose, methyl- α -Dgalactopiranoside, methyl- β -D-galactopiranoside, methyl- β -Dthiogalactose, phenyl- β -D-galactopiranoside, 4-nitrophenyl- α -D-galactopiranoside, 4-nitrophenyl- β -D-galactopiranoside, 2nitrophenyl- β -D-galactopiranoside, O-nitrophenyl- β -Dgalactopiranoside, and type 2 porcine stomach mucin (PSM).

Biochemical Characterization

The effects of pH, temperature, divalent cations, and EDTA on HA were determined as described by Sampaio et al. (1998). Molecular mass of the lectin was estimated by SDS-PAGE, as described by Laemmli (1970). LMW-SDS marker kit (GE Healthcare, IL, USA) was used as standard.

Native molecular mass was estimated by size exclusion chromatography (SEC) on BioSuite 250 HR SEC column (0.78 \times 30 cm, 5-µm particle size, Waters Corp., MA, USA), coupled to an Acquity UPLC system (Waters Corp., MA, USA), which had been equilibrated with Tris-HCl 20 mM, pH 7.2, containing NaCl 500 mM. The column was previously calibrated with a mixture of standard proteins: conalbumin (75 kDa), ovalbumin (45 kDa), carbonic anhydrase (29 kDa), ribonuclease A (14 kDa), and aprotinin (6.5 kDa).

Molecular Mass Determination

Molecular mass was determined on an AutoFlex III matrixassisted laser desorption/ionization time of flight (MALDI-TOF)/TOF mass spectrometer (Bruker Daltonics, Germany). The samples were dissolved in deionized water at variable concentration, mixed (1:3 v/v) in an α -cyano-4-hydroxycinnamic acid (CHCA) matrix solution (5 mg/mL of CHCA on acetonitrile, water, and trifluoracetic acid, 50, 40, and 10% v/v), spotted onto a sample plate, and then dried at room temperature. The spectra were acquired in linear positive mode and were processed with Flex Analysis 3.4 software (Bruker Daltonics, Germany).

N-Terminal Sequencing

An aliquot of 15 µg of purified lectin was solubilized in 15 µL of acetonitrile 10% and centrifuged at $8000 \times g$ for 5 min at room temperature. The solution was applied into a glass fiber disk and submitted to N-terminal sequencing.

Automated Edman degradations were performed in the Shimadzu model PPSQ-31A protein sequencer (Shimadzu Corp., Japan). PTH-amino acids from the N-terminus sequence were separated on a 2.0×250 mm Wakosil ODS column (Wako Pure Chemical Corp., Osaka, Japan) connected to a model LC-20AT pump. The absorbance was detected at 269 nm with a UV–Vis SPD-20A detector.

Primary Structure Determination by Tandem Mass Spectrometry

Purified lectin was submitted to SDS-PAGE as described above. After staining, electrophoretic bands were excised using a plastic pipette tip. Spots were reduced with dithiothreitol, and the lectin was carboxamidomethylated (CAM-lectin) with iodoacetamide, following pre-established methods (Shevchenko et al. 2006).

Treated spots were digested with trypsin, chymotrypsin, and pepsin. Digestions with trypsin and chymotrypsin were performed in 50 mM ammonium bicarbonate at 1:50 w/w (enzyme/substrate). Digestion with pepsin was performed in 100 mM HCl at 1:50 w/w (enzyme/substrate). All digestions were maintained for 18 h and then stopped with 2 μ L of 2% formic acid. Peptides were extracted from gel according to Shevchenko et al. (2006).

Peptide solution was loaded onto a C-18 ($0.075 \times 100 \text{ mm}$) nanocolumn coupled to a nanoAcquity system (Waters Corp., MA, USA). The eluates were directly infused into a nanoelectrospray source. Chromatography and mass spectrometry analysis were performed according to established methods (Carneiro et al. 2013).

Deglycosylation of tryptic peptides was carried out by GlycoprofileTM II, Enzymatic In-Solution N-Deglycosylation Kit (Sigma-Aldrich, St. Louis, MO, USA). First, the lectin (1 mg/mL) was solubilized in RapiGestTM SF surfactant 0.1% (Waters Corp., MA, USA), followed by digestion of CAM-lectin with trypsin 1:100 w/w (enzyme/substrate) for 18 h at 37 °C. RapiGestTM SF surfactant was pelleted by addition of TFA at final concentration of 0.2%.

Tryptic peptides were digested with peptide-N-glycosidase F (PNGase F) (1 U) (Sigma-Aldrich, St. Louis, MO, USA) for 18 h at 37 °C. Deglycosylated peptides were loaded onto a C-18 nanocolumn, and the eluates were directly infused into a nanoelectrospray source. The mass spectrometer was operated in positive mode with a source temperature of 373 K and a capillary voltage at 3.0 kV. MS/MS was performed according to the data-dependent acquisition (DDA) method. The lock mass used in acquisition was m/z 785.84 ion of the [Glu1] fibrinopeptide B. The selected precursor ions were fragmented by collision-induced dissociation (CID) using argon as the collision gas. All CID spectra were manually interpreted.

Circular Dichroism

Circular dichroism spectroscopic measurement was performed on a Jasco J-815 spectropolarimeter (Jasco International Co., Tokyo, Japan) connected to a peltier with controlled temperature. The lectin (0.2 mg/mL in 20 mM phosphate buffer, pH 7.0, containing 150 mM NaCl) was placed in a rectangular quartz cuvette with 0.5-mm path length. Spectra were acquired at a scan speed of 50 nm/min with a bandwidth of 1 nm. The acquisitions were performed at 190–250 nm (far-UV) and different temperatures (20–80 °C) with eight accumulations per temperature. The analysis of structural data was performed by DICHROWEB web server (Whitmore and Wallace 2008).

For measurement at 250–330 nm (near-UV), lectin at 0.5 mg/mL was previously incubated with D-galactose, α -D-lactose, D-galactosamine, and galacturonic acid at 10 mM each and placed in a rectangular cuvette with 5-mm path length. Spectra were acquired as described above.

The thermodynamic parameters of lectin folding and unfolding in the presence and absence of ligands were calculated by monitoring the changes in ellipticity at 293 nm as a function of temperature (Greenfield 2007). The lectin (8 μ M) was submitted to temperatures ranging from 70 to 95 °C with a ramp rate of 4 °C/min, and sampling was conducted at 1 °C each in the presence and absence of D-galactose, D-galacturonic acid, D-galactosamine, and α -lactose at 80 μ M each.

As proposed by Greenfield (2006, 2007), ligand-induced changes in the molar ellipticities of the lectin at 368 K were employed to find the affinity constant (Ka) for binding of carbohydrates to the lectin.

Structural Modeling

Structural Prediction

The structural prediction of the lectin was performed through interplay among the MODELER (Webb and Sali 2014) suite, I-TASSER (Yang and Zhang 2015), and QUARK (Xu and Zhang 2013) servers. Initially, the complete protein sequence (217 amino acids) was submitted to I-TASSER and QUARK servers for overall modeling. In parallel, three sequence fragments from the lectin (amino acid residues 1–70, 71–141, and 142–217) were also submitted to these servers in order to obtain the individual folding of each protein domain. The best models were selected based on their C- and TM-scores². Afterwards, the selected models were used as input to the MODELER suite, v.9.16. For structural modeling of the lectin, MODELER default parameters were used, and sequence alignment corrections were manually edited.

Initially, 30 theoretical models were generated, which were ranked based on their discrete optimized protein energy (DOPE) scores (Shen and Sali 2006). The models were then selected and analyzed for their stereochemical properties with the Protein Structure Validation Suite (PSVS) server, which integrates consolidated validation platforms, such as Procheck (Laskowski et al. 1993) and Molprobity (Davis et al. 2004). Finally, the best model was submitted to ModRefiner (Xu and Zhang 2013) (http://zhanglab.ccmb.med.umich.edu/ModRefiner).

Binding Site Prediction

A meta-server approach (http://zhanglab.ccmb.med.umich. edu) using the COACH (Yang et al. 2013) server was used to identify the most reliable carbohydrate-binding site.

Molecular Docking Calculations

The structural basis for carbohydrate recognition was explored through molecular docking calculations. For this purpose, four carbohydrates previously analyzed on hemagglutination inhibition assays were selected, including D-galactose (CID: 6036), D-galactosamine (CID: 24154), D-galacturonic acid (CID: 439215), and α -lactose (CID: 84571). All carbohydrate structures were obtained from the PubChem Substance and Compound database (Wang et al. 2009).

Calculations were performed with AutoDock Vina, version 1.1.2 (Trott and Olson 2009). The Autodock graphical interface AutoDockTools, version 1.5.6, was used to keep polar hydrogens and add partial charges to the proteins and ligands using Kollman United charges (Morris and Huey 2009). The protein and carbohydrate ligands were treated as rigid and flexible molecules, respectively.

Initially, a blind docking strategy was applied using a search space defined by a 40 Å \times 60 Å \times 40 Å cube covering the whole protein surface. Then, a refined search was performed using a search space defined by a 20 Å \times 20 Å \times 20 Å cube centered on the spot previously predicted by the COACH server as the most suitable carbohydrate-binding site. For all calculations, exhaustiveness was set to 15, and all other parameters were used as default. For each docking, the ten top-ranked generations based on the predicted binding affinity (in kilocalories per mole) were analyzed.

The solutions were first chosen based on the presence of important interactions commonly visualized for carbohydrate recognition by lectins, such as the stacking of pyranosidic rings against aromatic or aliphatic amino acid residues and the proper disposal of hydrogen bonds around hydroxyl groups (Wittmann and Pieters 2013). Since all selected ligands presented similar chemical structures, a common binding pattern was sought. The most suitable results were further ranked based on the theoretical binding energy, which was given as a negative score in kcal/mol.

Antibacterial Activity

Microorganisms and Culture Conditions

The assays to evaluate antimicrobial activity were performed on *Staphylococcus aureus* ATCC 25923 and *Escherichia coli* ATCC 11303. Bacteria were grown in trypticase soy agar medium (TSA; Himedia, India) and incubated at 37 °C for 24 h. After growth on the solid medium, some isolated colonies were removed, inoculated into 10 mL trypticase soy broth (TSB; Himedia, India), and incubated for 24 h at 37 °C under constant agitation. Prior to use, the cell density of each bacterial suspension was adjusted to a final concentration of 2×10^6 cells/mL.

Bacterial Aggregation Assay

After bacterial growth, bacterial suspensions were harvested by centrifugation at $2000 \times g$ for 10 min, washed twice with TBS/Ca²⁺, suspended in TBS/Ca²⁺ containing formaldehyde 4% and then kept for 16 h at 4 °C. Afterwards, the cells were washed twice with TBS/Ca²⁺, and suspended in the same buffer. Bacterial count was estimated by the serial dilution method, and optical density at 620 nm (OD₆₂₀) was maintained around 1.0. Bacterial agglutination was performed as described by Melo et al. (2014).

Effects of the Lectin on Planktonic Cells

The effects of lectin on planktonic cells were assessed by the broth microdilution method. Briefly, the lectin was diluted in 150 mM NaCl at concentrations ranging from 7.8 to 250 μ g/mL. The assay was performed in 96-well polystyrene plates by addition of 100 μ L of each bacterium at 2 × 10⁶ cells/mL to each well. The microplates were then incubated at 37 °C for 24 h in constant agitation, and the optical density was measured at 620 nm using a microplate reader (SpectraMax® I3, Molecular Devices LLC, CA, USA).

Biofilm Formation Assay

The effects on biofilm formation were evaluated as proposed by Vasconcelos et al. (2014). Briefly, sterile 96-well polystyrene plates were prepared following the same steps as previously established for planktonic cell growth. However, two plates were used for this purpose: one for biomass quantification by crystal violet and the other for enumeration of biofilm viable cells. In order to verify the role of the carbohydrate recognition domain (CRD) on antibiofilm activity exerted by the lectin, it was tested after previous incubation with 12.5 mM α -D-lactose. The influence of protein folding on antibiofilm activity was also investigated. For this purpose, the lectin was tested after denaturation by heating 1 h at 100 °C.

Biofilm Mass Quantification Quantification of biofilm mass was determined by crystal violet staining. After biofilm development, the content of each well was removed, and the wells were washed twice with 200 μ L of 150 mM NaCl to remove weakly adherent cells. For fixation of biofilms, 200 μ L of 99% methanol were added to each well. After 15 min, the methanol was removed, and the plates were allowed to dry at 25 °C. Then, 200 μ L of crystal violet were added to each well. After 5 min, the excess of crystal violet was removed, and the plates were washed twice with water. Finally, 200 μ L of acetic acid (33%, *v*/*v*) were added to wells to dissolve the crystal violet bound to the biofilm mass, and the absorbance was measured at 590 nm (OD₅₉₀) using a microplate reader (SpectraMax® I3).

Quantification of Number of Biofilm Viable Cells In order to determine the number of viable cells, the wells were washed twice with 150 mM NaCl, and then 200 μ L of the same solution were added to each well containing biofilms. The plates were placed in an ultrasonic bath for 10 min to release biofilmentrapped cells. Serial decimal dilutions from the obtained suspensions were plated on TSA to allow viability determination. The plates were then incubated for 24 h at 37 °C, and the number of colony forming units per milliliter (CFU/mL) was determined.

Fig. 1 ADEL purification. Affinity chromatography on HClactivated SepharoseTM column (1.0×8.0 cm). The column was equilibrated and washed with TBS. Retained fractions were eluted with 0.2 M lactose in TBS. Approximately 20 mL of the crude extract were applied. Flow was 1 mL/min. SDS PAGE 15%. *M* molecular marker. ADEL in absence and presence of β mercaptoethanol

Statistical Analysis

Statistical analyzes were performed by GraphPad Prism® version 5.0 from Microsoft Windows®. The data from all assays were compared using one-way analysis of variance (ANOVA), with Bonferroni post hoc test. The data were considered significant when p < 0.05.

Results

Lectin Purification

Crude extract of *A. dactylomela* eggs showed an evident and strong HA against human and rabbit erythrocytes. The HA induced by the treatment with extract was inhibited by the monosaccharide D-galactose and the disaccharide α -D-lactose. Affinity chromatography on HCl-activated SepharoseTM 4B was performed, and a novel lectin from sea hare eggs was successfully purified after elution with 0.2 M lactose (Fig. 1). In order to simplify the nomenclature, the new lectin was termed ADEL (derived from *A. dactylomela* eggs lectin). It was purified eight times and represents 81% of the total activity of the crude extract (Table 1).

Hemagglutinating Activity and Inhibition Assays

ADEL showed agglutinating activity in all tested erythrocytes in native and protease-treated forms. The lectin showed a slight preference for trypsin-treated rabbit erythrocytes (data not shown).

Several galactose-related carbohydrates showed inhibitory activity at distinct levels, with HA promoted by ADEL (Table 2). Galacturonic acid was the most potent inhibitor, followed by Dgalactosamine and α -methyl-D-galactopyranoside. The



 Table 1
 ADEL Purification

Fraction	Protein total	Titer (HU/ mL)	Specific activity		Yield (%)	Purification (fold)	MAC ^a (µg/
	(iiig)		(HU/ mg)	Total			
Crude	19.4	64	355	6887	100	1	2.80
ADEL	2.0	256	2844	5688	81	8	0.35

^a Minimal concentration of lectin able to cause hemagglutination

minimum inhibitory concentrations (MIC) for these sugars were 0.2, 5, and 5 mM, respectively.

Biochemical Characterization

HA promoted by ADEL was optimal at pH 6 and 7. The lectin was stable in temperatures lower than 60 °C, and a complete loss of activity was achieved at 80 °C. EDTA and CaCl₂ did not affect lectin activity (data not shown).

SDS-PAGE of ADEL resulted in 60 and 29 kDa bands in the absence and presence of β -mercaptoethanol, respectively, suggesting that ADEL is a dimeric protein formed by two identical polypeptide chains linked by a disulfide bond

 Table 2
 Inhibition of the hemagglutinating activity of ADEL by sugars and glycoproteins

	MIC
Sugar	
D-galactose	12.5 mM
α -methyl-D-galactopyranoside	10 mM
β-methyl-D-galactopyranoside	5 mM
D-galactosamine	5 mM
N-acetyl-D-galactosamine	>100 mM
Phenyl-B-D-galactopyranoside	6.25 mM
Phneyl-B-D-thiogalacopyranoside	>50 mM
4-nitrophenyl- α -D-galactoside	10 mM
4-nitrophenyl-β-D-galactoside	5 mM
2-nitrophenyl-	10 mM
O-nitrophenyl-β-D-galactoside	10 mM
D-galacturonic acid	0.2 µM
L-rhamnose	100 mM
L-arabinose	100 mM
α -lactose (gal $\beta(1\rightarrow 4)\alpha$ -glc)	12.5 mM
β -lactose (gal $\beta(1\rightarrow 4)\beta$ -glc)	6.25 mM
Lactulose (gal $\beta(1\rightarrow 4)$ fru)	6.25 mM
Melibiose (gal α (1 \rightarrow 6)glc)	6.25 mM
Rafinose $(gal\alpha(1\rightarrow 6)glc\alpha(1\rightarrow 4)fru)$	6.25 mM
Glycoprotein	
PSM type III	0.125 mg/mI
	-

(Fig. 1). Native molecular mass estimated by SEC was 14 kDa, suggesting an anomalous behavior of the lectin in gel filtration (Fig. 2).

MALDI-TOF mass spectrometric analysis showed two distinct ions: $[M + 1H]^{1+}$ at m/z 57,228 and $[M + 2H]^{2+}$ at m/z 28,613, which agrees with the molecular mass estimated by SDS PAGE (data not shown).

The first 29 amino acid residues of ADEL were identified by amino acid sequencing. The N-terminal sequence ¹DPDKCKTIRVESWSYKYAEKVVEDASYVL²⁹ showed high similarity with a galacturonic acid-binding lectin from *Aplysia kurodai*.

Amino Acid Sequencing by MS/MS

The primary structure of ADEL was determined by overlapping of amino acid sequences of peptides sequenced by MS/ MS deposited in Uniprot under accession number C0HK25. The amino acid sequence of ADEL consists of 217 residues, including 11 half-cystines involved in at least five intrachain disulfide bonds and one interchain disulfide bond (Fig. 3). According to the DiANNA 1.1 web server (Ferre and Clote 2005), cysteines are paired in the following form: ⁵Cys-Cys¹⁸⁷, ⁴²Cys-Cys⁶⁸, ⁶¹Cys-Cys⁷⁷, ¹¹⁴Cys-Cys¹³⁵, and ¹⁴²Cys-Cys²⁰⁶, whereas ¹⁷²Cys is involved in maintenance of the dimer. The theoretical isoeletric point was 5.41.

Theoretical molecular mass for the amino acid sequence totaled 24,450 Da. However, this value is lower than the molecular mass determined by MALDI-TOF. This divergence is most likely a result of glycosylation in the polypeptide chain.

ADEL has four glycosylation sites, of which three contain attached glycans, as indicated by tryptic glycopeptides (Table 3). These peptides showed differences in their observed and calculated molecular mass. These differences were attributed to glycans with distinct masses.

To confirm the existence of glycans attached to the polypeptide backbone, a deglycosylation assay was performed. Peptides presented in Table 3 were already deglycosylated after treatment with PNGase F. For instance, peptide T-9 at m/z 1076.45 has a sequence of ¹¹⁸SFGFYNQNSTVWVNHGCR¹³⁶, and its calculated molecular mass was 2171.95, whereas mass observed by MS was 3226.25. The difference between observed and calculated mass was 1054.3, corresponding to the glycan with composition HexNAc₂Hex₄. After PNGase F digestion, T-9 showed m/z 726.31 and observed molecular mass of 2172.9, indicating the release of the glycan (Fig. 4).

The RADAR tool (http://www.ebi.ac.uk/Tools/pfa/radar/) was employed to detect the presence of three tandem-repeat domains along the polypeptide chain, which share distinct levels of identity (Fig. 5). Domain A (2-70) has identity of 63 and 55% to domain B (74-144) and C (152-214), respectively, whereas domain B showed identity of 49% to domain

2 00

1.50

1.00

0.50

0.00

Absorbance at 280 nm



18.00

21.00

15.00 Time

12.00

Fig. 2 Size exclusion chromatography of ADEL. The column BioSuite 250 HR SEC (0.78×30 cm, 5-um particle size, Waters Corp.) was equilibrated and eluted with Tris-HCl, 50 mM, pH 7.2, containing NaCl

6.00

9.00

3.00

C. Furthermore, ADEL presented microheterogeneities in three positions: ⁴K/M, ⁵³D/A, and ¹⁹⁹A/T.

ADEL showed high sequence similarity with AKL (*A. kurodai* eggs lectin, GenBank: AB968312.1) and with a hypothetical lectin isoform X1 (NCBI RefSeq XP_005101558.1) from *Aplysia californica* (Fig. 6).

Circular Dichroism

Spectra of native ADEL in far-UV exhibited minimum absorption at 215 nm and maximum at 202 and 229 nm, suggesting a predominance of β -conformation in its secondary structure (Fig. 7a).

According to the CONTIN prediction method (Van Stokkum et al. 1990), available online as part of the server DICROWEB (http://dichroweb.cryst.bbk.ac.uk), the

500 mM. Approximately 200 μg of ADEL were loaded. UPLC operated at flow of 0.5 mL/min

24.00

27.00

30.00

theoretical secondary structure of ADEL consisted of 4% α -helix, 44% β -sheet, 21% turn, and 32% coil.

Changes in the secondary structure of ADEL were examined by monitoring modifications in CD spectra as a function of temperature. Between 20 and 70 °C, no significant changes in secondary structure were observed, but after heating at 80 °C, some modifications occurred in maximum and minimal absorptions, indicating loss of secondary structure (Fig. 7b).

The spectra acquired in near-UV in the presence of some ligands revealed mild changes in tertiary structure of the lectin in comparison with native form (Fig. 7c). D-galactose (10 mM) caused a slight variation in tertiary structure of ADEL, while galacturonic acid (10 mM) exhibited strong effect on ADEL conformation.

Thermodynamics parameters of ADEL folding and unfolding in the presence and absence of carbohydrates are

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DPDMCKTIRV ESWSYKYAEK VVEDASYVLN MTVVDRQSAA ACTLGESFGY QKDTLWVDHG CRADFKVCYL
                              |-----T1<sup>(5210.27)</sup>-----||-T3<sup>(1257.46)</sup>|
 |--01<sup>(1635.64)</sup>--|
                                                                                                 |---
                                                    |----T2<sup>(1716.74)</sup>----|
                   |---02 (1650.71) --- |
                                                                                                 |---
      -----N-TERMINAL------
                                                           |---P1<sup>(1688.76)</sup>---| |--Q3<sup>(1625.64)</sup>---|
 PVMPTECQTL RVESWNYKYA EKVVKGAALF INMTVEDRQS EASCDLDKSF GFYNQNSTVW VNHGCRADFN
 --T4<sup>(1865.76)</sup>-||--T5-|
                                     |--T7<sup>(2652.16)</sup>--||-T8<sup>(1151.38)</sup>||----T9<sup>(3226.25)</sup>-----||---
 ----T6 (2772.30) <u>924.38</u>
                                          |-----|
                                                                                       |-Q6<sup>(1624.63)</sup>-
                         |--04 (1423.78) -- |
 ICYLKGAVTT STINVSSWNY QYATKVLPAA SCIYSMRVVN QQSAAPCTLG TTYGFVANTM WVDDGCRADF
 T10-| |----T11<sup>(2062.02)</sup>-----||-T12<sup>(1366.60)</sup>-||-----T13<sup>(3317.48)</sup>-------||--
1142.51
---Q7 (1562.76) ---- |
                             |--08<sup>(1292.64)</sup>--| |----P2<sup>(1921.93)</sup>-----|
                                                                                         09 (1053.54)
 NPTYYSP
 -T14-- | 1173.53
 |-P3--| 840.39
```

Fig. 3 Amino acid sequence of ADEL. Primary structure of ADEL was determined by overlapping of peptides obtained by digestions with trypsin (T), chymotrypsin (Q), and pepsin (P). *Asterisks* represent

sequence microheterogeneites. Molecular mass determined by mass spectrometry are in parenthesis. Leucine and Isoleucine were assigned according with similarity between ADEL and AKL

Peptide	m/z	Amino acid sequence	Molecular mass		Δ	Glycan prediction	Glycan mass
			Observed	Calculated			
T-1	1303.57	²¹ VVEDASYVLNMTVVDRQSAAACTLGESFGYQK	5210.27	3507.66	1702.61	HexNAc ₂ Hex ₈	1702.58
T-7	1246.05	⁹⁶ GAALFINMTVEDR	2490.10	1435.71	1054.39	HexNAc ₂ Hex ₄	1054.37
T-7.1	1165.04	⁹⁶ GAALFINMTVEDR	2328.07	1435.71	892.36	HexNAc ₂ Hex ₃	892.31
T-7.2	1327.09	⁹⁶ GAALFINMTVEDR	2652.16	1435.71	1216.45	HexNAc ₂ Hex ₅	1216.42
T-9	1076.45	¹¹⁹ SFGFYNQNSTVWVNHGCR	3226.25	2171.95	1054.30	HexNAc ₂ Hex ₄	1054.37
T-9.1	1400.59	¹¹⁹ SFGFYNQNSTVWVNHGCR	4198.66	2171.95	2026.70	HexNAc2Hex10	2026.68
Q-5	1236.83	100FINMTVEDRQSEASCDLDKSF	3707.49	2491.10	1216.39	HexNAc ₂ Hex ₅	1216.42

 Table 3
 Amino acid sequence and mass differences of glycopeptides of ADEL

N Asn residue that bound N-linked glycans

presented in Table 4. As demonstrated in the hemagglutinating assay, galacturonic acid is the most important ligand of ADEL; the interaction between galacturonic acid and ADEL shows Ka = $1.5 \times 10^7 \text{ M}^{-1}$. Furthermore, the presence of galacturonic acid increases ADEL Tm by 9 K in comparison to apo-lectin (data not shown).

ADEL Structural Prediction and Docking Calculation

Prior to molecular modeling, the ADEL sequence was used as input data for a PSI-BLAST (Ding et al. 2014) search against the protein data bank (PDB) to identify suitable template structures (http://blast.ncbi.nlm.nih.gov/Blast.cgi). No significant sequence



Fig. 4 Glycosylated peptide of ADEL. Mass spectra of the ion $[M + 2H]^{2+}$ at m/z 1246.05. N-acetyl-hexosamines and hexoses are represented by squares and circles, respectively (a). Deglycosylated peptide of ADEL, after PNGase F digestion (b)



Fig. 5 Alignment of the repeated domains of ADEL. Alignment realized by ESPript 3.0. *Black* and *white boxes* represent identical and non-identical amino acids, respectively. Consensus >50 represents amino acids present in at least 50% of the aligned sequences

similarity was found to any experimentally determined structure. Based on that, interplay between different modeling servers and software was performed to produce the most reliable structural prediction of ADEL.

The overall starting ADEL structure was obtained by I-TASSER. However, several poorly modeled patches were identified since no template structures were available on PDB. To overcome this issue, the structural predictions of each individual ADEL domain performed by QUARK were used to fill the structural gaps. The selected structures were used by MODELER to produce a complete ADEL model, with 99.1% of residues placed in allowed regions (90.7% on favorable regions) of the Ramachandran plot (Supplementary Fig. 1).

The obtained ADEL model (217 amino acids) possesses two independent domains connected by a short loop (Phe100-Val105), which were named domains A (Asp1-Leu99) and B (Glu106-Pro217) (Fig. 8a). Domain A is composed of a flat three-stranded and a curved five-stranded β - sheet, while domain B presents a flat three-stranded and a curved four-stranded β -sheet. According to I-TASSER, the crystal structure of the polymeric immunoglobulin receptor ectodomain from the rainbow trout *Oncorhynchus mykiss* (PDB 5F1S) is the structure most closely related to ADEL on PDB, despite its low sequence identity.

The predicted binding site is located at the interface between domains A and B, mainly built by residues Lys6, Ile8, Tyr17, Glu106, Asp107, Gln109, and Arg207 (Fig. 8b).

Refined docking calculations resulted in an increasing order of theoretical binding energy among the analyzed carbohydrates, as follows: D-galactose < D-galactosamine < α lactose < D-galacturonic acid (Table 5). Docking calculations also revealed a conserved binding pattern for the analyzed monosaccharides over the ADEL carbohydrate-binding site. It is possible to observe that the side chains of residues Lys6 and Ile8 act to stabilize the pyranosidic ring of the monosaccharides through hydrophobic stacking interactions. On the other hand, the hydroxyl groups of these ligands establish an



Fig. 6 Alignment of ADEL and other *Aplysia* eggs lectins. Amino acid sequence of ADEL was aligned with *Aplysia kurodai* eggs lectin (AKL) and a putative lectin from *Aplysia californica* (isoform X). Alignment

realized by ESPript 3.0. *Black and white boxes* represent identical and non-identical amino acids, respectively. Consensus >50 represents amino acids present in at least 50% of the aligned sequences



Fig. 7 CD spectra of ADEL. a Far-UV CD spectra (190–250 nm) of ADEL. The cuvette path length was 0.05 cm; protein concentration was 8 μ M in PBS, pH 7. b Far-UV CD spectra of ADEL in different

intricate hydrogen bond pattern with the main chain carbonyl mass groups from residues Tyr17 and Asp107 and with the side *S. au* chain polar groups from residues Glu106, Gln109, and 40%.

In the case of α -lactose, the binding pattern seems to be slightly different. The pyranosidic rings from the D-galactose and D-glucose residues are stabilized not only through hydrophobic interactions with residues Lys6 and Ile8, but also with the side chain of residue Thr7. Further, the hydrogen-binding pattern also changes to accommodate this bulkier ligand (Fig. 9d and Table 5). It is worth mentioning that the ADEL carbohydrate-binding site is disposed in a groove between domains A and B surrounded by other solvent-exposed charged and hydrophobic amino acid residues that may comprise extended sites for the anchorage of complex ligands.

Agglutination and Antibacterial Activity

Arg207 (Fig. 9 and Table 5).

ADEL agglutinated *S. aureus* cells, but it was not able to agglutinate *E. coli* (Figure 10). Despite its agglutinating activity, ADEL was not able to inhibit the growth of either species (data not shown).

Effects of ADEL on Biofilm Formation

The effect of ADEL on the prevention of biofilm formation was assessed by two different assays: determination of biofilm

Table 4 Affinity constants and free energy of the binding of sugars to ADEL at 358 K $\,$

Sugar	$10^{-5} \times \mathrm{Ka}$ (M^{-1})	α	$-\Delta G_F$ (kJ mol ⁻¹)	<i>T_M</i> (K)	
Native	_	0.338	361.5	356	
α-D-lactose	0.4	0.594	1060	359	
D-galactose	6.1	0.788	2697.5	361	
D-galactosamine	6.8	0.793	2775	361	
D-galacturonic acid	150	0.873	4989.2	365	

 α fraction folded at 358 K, $\Delta G_{\rm F}$ free energy of folding, $T_{\rm M}$ unfolding transition

temperatures. **c** Near-UV CD spectra (250–330 nm) of ADEL with some ligands. The cuvette path length was 0.5 cm; protein concentration was 20 μ M in PBS, pH 7

mass and enumeration of biofilm viable cells. The treatment of *S. aureus* biofilms with ADEL reduced biomass by almost 40%. The effect was seen in all concentrations evaluated (Fig. 11a). Regarding the number of biofilm-entrapped viable cells, ADEL reduced *S. aureus* viability mainly at higher concentrations (250–62.5 μ g/mL) (Fig. 11b). Interestingly, the biofilm formation of *E. coli* was not affected (Fig. 11).

The antibiofilm activity on *S. aureus* was abolished when ADEL was denatured by heating at 100 °C. On the other hand, treatment with α -D-lactose neither interfered with biofilm formation nor did it inhibit the antibiofilm activity of ADEL (data not shown).

Discussion

A new lectin from eggs of the sea hare *A. dactylomela* was successfully isolated and purified by affinity chromatography using SepharoseTM chromatography media. The use of SepharoseTM for purification of galactophilic lectins is a classical method, and several lectins have been successfully isolated by this procedure (Gasparini et al. 2008; Jimbo et al. 2000; Moura et al. 2006; Vasta and Marchalonis 1986). The activation of SepharoseTM by HCl constitutes an essential step to expose galactosides of the internal matrix, as long as these residues are available for specific lectin interaction.

The literature reports three lectins isolated from reproductive tissues of sea hares, which were isolated from *Aplysia depilans* (AGL) and *A. kurodai* (AKL-1 and AKL-2). AGL and AKL-1 are dimeric proteins with 30 kDa per subunit linked by disulfide bonds, as is ADEL. AKL-2 is also a dimeric protein, but with 16 kDa per subunit (Gilboa-Garber et al. 1985; Kawsar et al. 2009, 2011).

The literature shows that AKL-1, AKL-2, and AGL all have a strong affinity for galacturonic acid (Gilboa-Garber et al. 1985; Kawsar et al. 2009, 2011). ADEL also presents high specificity for the same carbohydrate. In fact, the specificity of lectins from sea hare eggs is a characteristic that deserves closer attention based on the fact that the species are spread far apart, but share the same sugar specificity.



Fig. 8 ADEL overall structure. a Monomer structure of ADEL represented as *colored cartoon* and *lines with a transparent surface* delineating the protein shape. The carbohydrate-binding site location is

highlighted by a *red dashed circle*. **b** Carbohydrate-binding site representation, which shows the coordination of D-galacturonic acid by the main site residues (represented as *green sticks*)

Thus, galacturonic acid should be considered as one of the preferred targets for the sea hare eggs lectins.

In addition to galacturonic acid, ADEL was also inhibited by D-galactose and its derivatives. ADEL showed slight preference by β -galactosides, thus suggesting the involvement of the hydroxyl at C-1 in the interaction between lectin and sugar. For instance, methyl- β -D-galactopyranoside and 4nitrophenyl- β -D-galactopyranoside are two times more effective than methyl- α -D-galactopyranoside and 4-nitrophenyl- α -D-galactopyranoside, respectively. Moreover, β -lactose was more effective in inhibiting ADEL than α -lactose. The presence of substituents at C-1, such as methyl, phenyl, and nitrophenyl, increases inhibitory efficiency when compared to Dgalactose. However, methyl- β -D-thiogalactopyranoside

Table 5 Comparative analysis based on docking calculations of the main interactions and bond distances involved in the coordination of the selected carbohydrate ligands by ADEL. The theoretical binding energies are presented at the bottom. The selected cutoff for hydrogen bonds and hydrophobic interactions were 3.5 and 4.5 Å, respectively

ADEL		Galacturonic acid		Lactose		Galactosamine		Galactose	
Atom	Residue	Atom	Distance (Å)	Atom	Distance (Å)	Atom	Distance (Å)	Atom	Distance (Å)
CE	Lys6	C1	4.5	C1 (Glc)	4.5	C1	4.2	C1	4.2
0	Thr7	_	-	O4 (Gal)	2.7	_	_	-	_
		_	-	O6 (Gal)	2.8	-	-	-	_
CG1	Ile8	C1	4.1	C2 (Glc)	4.1	C1	4.0	C1	4.0
0	Tyr17	O1	2.8	O1 (Glc)	3.2	01	2.7	01	2.7
		O2	3.0	O2 (Glc)	3.1	Ν	3.1	Ν	3.1
OE1	Glu106	O4	3.1	O1 (Glc)	3.5	O4	3.0	O4	3.0
		05	3.1	-	-	O5	3.4	O5	3.4
		_	-	-	-	-	3.0	O6	3.1
0	Asp107	O3	2.8	O6 (Glc)	2.8	O3	2.8	O3	2.8
		O4	2.9	-	-	O4	3.2	O4	3.2
OE1	Gln109	O7	3.0	O2 (Gal)	2.8	_	—	-	_
NE2	Gln109	O7	3.3	O1 (Gal)	3.2	_	—	-	_
		-	—	O2 (Gal)	3.3	_	—	-	_
		-	—	O6 (Glc)	3.0	_	-	-	_
NH1	Arg207	O6	3.2	-	-	O6	3.0	O6	2.9
NH2	Arg207	O4	3.1	-	-	O4	3.0	O4	3.0
		O6	3.3	-	-	O6	3.1	O6	3.1
		O7	3.0	-	_	_	—	_	_
Total energy		-5.6 kca	l/mol	-5.4 kcal/mol		-5.2 kcal/mol		-5.1 kcal/mol	

Fig. 9 Ligand coordination at the carbohydrate-binding site. a Coordination of D-galactose, represented as vellow sticks. b Coordination of D-galactosamine, represented as magenta sticks. c Coordination of D-galacturonic acid, represented as green sticks. **d** Coordination of α -lactose, represented as purple sticks. The protein backbone and the main amino acid residues involved in carbohydrate coordination by ADEL are represented as grav cartoon and sticks, respectively. Hydrogen bonds are represented as vellow dashes. A few interactions were omitted for the purpose of clarity (see Table 5)



showed no inhibitory effect, indicating steric hindrance, most likely from the presence of a thiol group. Although C-1 seems to be an important recognition point, a similar grade of inhibition by melibiose (Gal α 1 \rightarrow 6Glc), α -D-lactose (Gal β 1 \rightarrow Glc), and raffinose (Gal α 1 \rightarrow 6Glc α 1 \rightarrow 4Fru) indicates that C-1 cannot be considered as a key point in the recognition by ADEL.

Since D-glucose, L-fucose, and D-mannose are not recognized by ADEL, the orientation of hydroxyls at C-2 and C-4 seems to be more important than that at C-1. The axial orientation of C-2 and equatorial orientation of C-4 present in D-galactose seem to be essential for interaction. ADEL was inhibited to a greater degree by D-galactosamine compared to D-galactose, and it was not inhibited by GalNAc, indicating that a small uncharged substituent at C-2 (i.e., amine) increases affinity, whereas a charged substituent (i.e., acetyl) abolishes it.

The importance of C-5 was indicated by strong inhibition by the presence of a hydroxymethyl group in D-galacturonic acid when compared to the inhibitory effect produced by Dgalactose, L-arabinose, and L-rhamnose, even though the first two sugars show the same orientation of hydroxyls at C-2, C-3, and C-4. The presence of the hydroxymethyl group strongly improves lectin-sugar binding since galacturonic acid is 6×10^5 times more effective than D-galactose.

The glycoprotein PSM has a terminal GalNAc and Dgalactose as internal residue (Slomiany and Meyer 1972),



Fig. 10 Agglutination of bacteria by ADEL. *S. aureus* incubated with TBS (a) and ADEL (b); *E. coli* incubated with TBS (c) and ADEL (d). *Arrows* indicate bacterial agglutination



Fig. 11 Biofilm formation of *S. aureus* and *E. coli* in the presence of ADEL. **a** Biofilm biomass and **b** number of viable cells present in the biofilms. *p < 0.05 compared to control

and it exhibited inhibitory effect on ADEL. Like other sea hare lectins, ADEL was inhibited by small amounts of PSM (Kawsar et al. 2009). The presence of T-antigen (Gal β 1 \rightarrow 3GalNAc) in PSM could account for the inhibitory effect based on the strong recognition of T-antigen by AGL (Gilboa-Garber and Sudakevitz 2001).

Consistent with data from the hemagglutination inhibition assay, CD spectrum of ADEL acquired in the near-UV spectral region revealed that the interaction with galacturonic acid plays an important role in ADEL conformation. D-galactose, D-galactosamine, and α -D-lactose also exerted some influence on the tertiary structure of ADEL, but to a lesser extent. These sugars showed distinct Ka values in the following order: galacturonic acid > D-galactosamine > D-galactose > α -D-lactose.

Curiously, the order of affinity seen in molecular docking was different in comparison to that observed in CD and HA inhibition. In the computational approach, galacturonic acid remains a sugar with higher binding energy, but α -D-lactose comes in second, followed by D-galactosamine and D-galactose.

In addition to clarifying the interactions between ADEL and sugars, CD measurement was also used to estimate secondary structure. According to the CD spectra, ADEL has a high content of β -conformations (sheets and turns), a characteristic of many lectins (Vasta et al. 2004). Indeed, the 3D model of ADEL is predominantly composed of β -structures, but interestingly, it presents a singular structure for lectins.

Animal lectins present a wide structural diversity. For instance, galectins have a typical jelly roll β -sandwich conformation (Vasta et al. 2004). The C-type lectin domain (CTLD) consists of a double loop stabilized by two disulfide bonds highly conserved in the base of the second loop (Zelensky and Gready 2005), while R-type lectins showed a β -trefoil (Uchida et al. 2004). However, none of these folds resembles the ADEL model. Obviously, for a more accurate comparison, the determination of three-dimensional structure of ADEL by experimental approaches is required.

ADEL, *A. kurodai* egg lectin and AGL are independent of divalent ions (Gilboa-Garber et al. 1985; Kawsar et al. 2009). AKL-1 and AGL are glycoproteins (Gilboa-Garber et al. 1985; Kawsar et al. 2009). In addition to evidence that AKL and AGL are glycoproteins, our study could provide accurate glycan structures of ADEL by MS analysis. ADEL is also a glycoprotein with three distinct glycans attached to three distinct N-glycosylation sites.

N-glycosylation is one of the most common post-translational modifications (PTMs) present in proteins, and several functions are attributed to its presence, including correct protein folding, protein localization, and regulation of protein degradation. Glycosylation events typically occur in the endoplasmatic reticulum (ER), and oligosaccharide maturation occurs in the Golgi apparatus (Ghazarian et al. 2011). According to the type of carbohydrate attached to polypeptide chain, N-oligosaccharides can be classified into three groups: high mannose, complex, and hybrid oligosaccharides (Kukuruzinska and Lennon 1998). Generally, all N-glycans share a common pentasaccharide core: Man α 3(Man α 6)Man β 4GlcNAc β 4GlcNAc. Our findings indicated that this precursor is also present in the ADEL structure.

In the first N-glycosylation site of ADEL (30 NMT), an oligosaccharide of 1702.61 Da was found with mass corresponding to the pentasaccharide precursor (892 Da) with addition of five hexose residues (810 Da). In the second site (102 NMT), molecular mass of the oligosaccharide ranges from 892.36 to 1216.45 Da, corresponding to the pentasaccharide precursor alone, and the precursor with two hexose residues (324 Da), respectively. Finally, the third site (126 NST) has an oligosaccharide of 1054 Da that consists of the precursor and one hexose residue.

All oligosaccharides found in ADEL consist of a pentasaccharide precursor increased by some hexose residues. Among N-glycan groups, the only one that contains only hexoses, besides the precursor, is the high-mannose oligosaccharide.

Therefore, MS analysis strongly suggests that all oligosaccharides attached to the ADEL polypeptide chain are highmannose-type oligosaccharides. To the best of our knowledge, this is the first time that glycan composition has been characterized for a lectin from opisthobranch (Gastropoda).

Although several efforts were made to remove oligosaccharides attached to ADEL, they were unsuccessful. Digestion of ADEL by PNGase F showed no effect, since no shift was observed in SDS PAGE and MS (data not shown). We speculate that the native folding of ADEL was maintained, even when using heat treatment and surfactant reagents during digestion. However, PNGase F could digest tryptic peptides of ADEL. All glycosylated peptides were digested by PNGase, indicating that no deoxyhexoses are attached to chitobiose precursors.

Besides glycosylation, specificity for carbohydrates, divalent ion independence, and quaternary arrangement, ADEL shares a high level of sequence similarity with AKL-1. ADEL showed 91 and 88% of identity and 97 and 93% of similarity with AKL-1 (GenBank: AB968312.1) and isoform X1 of *A. californica* (NCBI RefSeq XP_005101558.1), respectively.

Lectins isolated from the *Aplysia* genus showed no similarity to other known lectins, but the presence of a repeated domain in their structures is a common property of certain families of lectins. In particular, lectins that showed repeated domain in their primary structures and shared similarity of sequence with B-chain of Ricin were grouped in the R-type lectin family (Hirabayashi et al. 1998).

Lectins from sea hare eggs showed low similarity with Ricin, did not share conserved residues, and had no β -trefoil domain typical of R-type lectins, according to structural modeling. Thus, ADEL, AKL, and isoform X1 could not be grouped in this family, but could be grouped in a new family of lectins considering their similarities of sequence and biochemical characteristics. However, further experiments should be performed to better characterize these lectins.

ADEL was effective in agglutinating planktonic cells and inhibiting biofilm formation by *S. aureus* ATCC 25923. In fact, several galactophilic lectins can agglutinate bacterial cells (Carneiro et al. 2015; Melo et al. 2014; Moura et al. 2006), recognize glycans on the cell surface and distinguish Gram-positive from Gram-negative bacteria (Sun et al. 2007; Moura et al. 2006). The surface of Gram-positive bacteria is constituted by several structures containing carbohydrates, such as peptidoglycan, capsular polysaccharides, and teichoic acid (Weidenmaier and Peschel 2008), which may serve as possible binding sites for ADEL.

Lectins from marine organisms have been exploited for several biological applications, including antimicrobial activity (Moura et al. 2006; Roué et al. 2012; Cheung et al. 2015). Although some studies have focused on the antimicrobial potential of marine lectins, only a few are directly related to their effect on pathogenic biofilms. In the present work, we also evaluated the antibacterial potential of ADEL on planktonic growth and biofilm formation by *S. aureus*. ADEL does not inhibit bacterial growth, but rather inhibits biofilm formation. According to Liljemark et al. (1981), the formation of bacterial aggregates can cause a decrease in the number of adherent cells, which may explain why ADEL does not inhibit bacterial growth, but is, instead, effective on biofilm formation. Similar results were obtained by Vasconcelos et al. (2014),

using plant and red algae lectins on Gram-positive and Gramnegative bacteria, as well as on yeasts biofilms.

In order to investigate the involvement of CRD in antibiofilm activity, the lectin was preincubated with α -Dlactose at MIC (12.5 mM). Interestingly, the activity was not abolished. This is not surprising since differences in affinity can be achieved between monosaccharides and structured glycans. Therefore, it is possible that ADEL has a higher affinity to complex glycans located on the *S. aureus* surface. Indeed, lectins can preferentially recognize carbohydrate complexes (Ghazarian et al. 2011). Furthermore, the inhibition of biofilm formation by ADEL is dependent on its structural integrity, in particular because denaturation by heating inhibited its antibiofilm activity.

Biofilms are described as structured communities of surfaceassociated microbial cells enclosed in a self-produced polymeric matrix (Flemming and Wingender 2010). Since biofilms are associated with increased resistance to conventional antimicrobial agents, ADEL could be considered as an alternative to preventing biofilm formation.

In addition, the antibiofilm activity exhibited by ADEL allows us to speculate about its physiological roles. For instance, *Aplysia* eggs are deposited on the surface of stones and algae where they remain for a few days until hatching. During incubation, eggs are exposed to several biological stresses, including bacterial infection. Therefore, the endogenous production of ADEL may be induced by the presence of pathogenic bacteria, and the lectin might act in the defense of eggs.

In conclusion, we herein report a novel type of lectin isolated from *A. dactylomela*, which is highly specific for galacturonic acid and has three high-mannose oligosaccharides attached to its polypeptide chain. Moreover, ADEL can inhibit biofilm formation by *S. aureus*. Thus, ADEL could be considered as an alternative to conventional antimicrobial therapy used to combat Staphylococcal biofilms.

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