A novel and efficient and low-cost methodology for purification of *Macrotyloma axillare* (Leguminosae) seed lectin

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**ABSTRACT**

The \(\text{N}-\)acetyl-galactosamine specific lectin from *Macrotyloma axillare* seeds (LMA) was purified by precipitation and ion exchange chromatography. The LMA 0.2 mol L\(^{-1}\) fraction showed hemagglutinating activity on erythrocytes A1. The results for molecular mass determinations were about 28 kDa. The LMA pH-dependent assays showed best hemagglutinating activity at pH 6.0–8.0; being decreased at acidic/alkaline conditions and by EDTA treatment. LMA is a tetramer at pH 8.2 and a dimer at pH 4.0. Human erythrocytes from ABO system confirmed the A1 specificly for LMA. This new methodology is useful and easy, with low costs, for lectin purification in large amounts.

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1. Introduction

Lectins are a class with several structure-related proteins, which possess considerable specific binding capacity for carbohydrates molecules [1]. They are binding proteins of non-immune origin, with highly ordered three-dimensional structures that can be associated as dimeric or tetrameric complexes [2]. The leguminous plants, particularly their seeds, are recognized lectins sources [3]. Generally, the lectin basic monomer is described as containing two major \(\beta\)-sheets and another one smaller (about to 20–30 kDa). This folding and the features of the carbohydrate-binding site, that consists of a shallow groove on the superimposed \(\beta\)-sheets, are very similar in all leguminous lectins [4–6]. Due to the specific binding properties, the lectins can be considered important biotechnological tools with wide applications, such as the ability of agglutinate complex carbohydrates, glycoproteins, erythrocytes, vegetative cells, lymphocytes, fibroblasts, spermatozoids, fungi and bacteria [7,5]. Thus, the lectin in vitro functions are well documented and the molecular basis of this interaction has been studied with a variety of biophysical techniques, including X-ray crystallography, NMR and isothermal titration calorimetry (ITC) [8–10].

The lectin of *Macrotyloma axillare* (LMA) was previously isolated by affinity chromatography using antigen A1 + substance H coupled to Sepharose 4B\(^{\text{TM}}\) resin. LMA has \(\text{N}-\)acetylgalactosamine (GalNac) specificity, similar amino acid composition and the same N-terminal sequence [11] as the lectin isolated from Dolichos biflorus (DBL) [12,13]. Such as DBL, LMA is specific for A1 human erythrocytes, being therefore useful for routine blood group identification at blood banks [14,15], in addition to other lectin biotechnology applications that include the GalNac recognition [16,17].

The affinity chromatography techniques generally are the first choice for lectin purification, in accordance to their binding features [18] but this method can be very expensive and, sometimes, the procedure does not supply enough amounts for industrial applications. The affinity chromatography maintenance is a hardy task when plant crude extracts are loaded into columns because such samples contain pigments, oily components, proteolytic enzymes and other complex substances that could damage the affinity chromatography system and impair the purification. Considering such problems and the difficulties observed for affinity systems and their importance for lectin purification, the present work proposes a new preparative methodology for the purification of *M. axillare* seed lectin, based on thermal and ethanolic precipitation followed by ion exchange chromatography. Such simple and versatile methodology (accepted patent [19]) yields active lectin amounts applicable for preparative and industrial purposes. Some LMA properties were
studied, such as thermal and pH-dependent stability; molecular mass determination by mass spectrometry (ESI-TOF), SDS-PAGE (denatured form) and by molecular exclusion (native form), since such studies and properties were not fully explored for LMA.

2. Materials and methods

2.1. Reagents and materials

The M. axillare seeds ("Java" cultivar) were purchased from Mat-suda Seeds, Brazil. Q-Sepharose™ ion exchange and Sephacryl™ S-200 HR 16/60 size exclusion column were purchased from Amersham Pharmacia Biotech (Uppsala, Sweden). Molecular mass standards were obtained from Sigma–Aldrich (St. Louis, MO, USA). All other reagents were of analytical grade or of high quality and purchased from reputed commercial suppliers.

2.2. Lectin extraction from M. axillare seeds—crude extract

M. axillare seeds were powdered with semi-industrial grinder and the powder obtained was stored at −20 °C. For lectin extraction, the crude extract was prepared with 80 g of seed powder and sufficient volume of 0.15 mol L⁻¹ NaCl solution with 5 mmol L⁻¹ of MnCl₂ and 5.0 mmol L⁻¹ of CaCl₂ to complete a final suspension about 20% (w/v) (400 mL), and incubated at 4 °C for 24 h. After incubation, the suspension was filtered and the retained precipitate was washed with the same solution described above until adding up the volume to 400 mL. Then, the M. axillare crude extract obtained was centrifuged at 4800 × g by 30 min at 4 °C for the removal of insoluble fibers. The supernatant was submitted to the next step.

2.3. Thermal precipitation of the crude extract

The M. axillare 20% (w/v) crude extract was submitted to thermal precipitation at 85 °C in water bath for 30 min and was soon transferred to a cold ice bath. After this cooling, the thermally treated crude extract was submitted to centrifugation at 4800 × g for 30 min at 4 °C. The precipitate material was discarded and the supernatant was submitted to the next precipitation step.

2.4. Precipitation by cold ethanol

Commercial ethanol was cooled to −20 °C previously to perform the precipitation step. The thermally treated crude extract was submitted to precipitation at 4 °C by addition of increasing ethanol volumes from 20 to 60% (v/v), regarding the initial crude extract volume and the ethanol volume added. The precipitate obtained in the range of 20–60% (v/v) of ethanol was submitted to centrifugation at 4800 × g for 30 min at 4 °C. The pellet was resuspended at 0.01 mol L⁻¹ of NaHCO₃, pH 8.3, drop by drop. The redissolved fraction was submitted to centrifugation to discard denatured and insoluble proteins. The supernatant was freeze-dried and stored until the next ion exchange chromatography step.

2.5. Ion exchange chromatography of the fraction precipitated with 20–60% ethanol

The 20–60% ethanol precipitated fraction (500–800 mg aliquots) was dissolved in 10 mL of 0.01 mol L⁻¹ of NH₄HCO₃, pH 8.3 buffer and loaded on a HiloAff™ 16/10 Q-Sepharose™ anion exchange chromatography column (10.0 cm × 1.6 cm) coupled to a Fast Performance Liquid Chromatography System (FPLC™ Pharmacia™) equilibrated with the same buffer. The elution was performed by stepwise NaCl gradients 0.10; 0.20; 0.30; 0.40; 0.50 and 1.0 mol L⁻¹, all of them in 0.01 mol L⁻¹ of NH₄HCO₃ buffer, pH 8.3. The chromatographic flow rate was 3.0 mL min⁻¹. The eluate was monitored at 280 nm and the hemagglutinating activity was determined (described in Section 2.6). The fractions with A1 hemagglutinating activity were collected and submitted to dialysis against 0.01 mmol L⁻¹ NH₄HCO₃ buffer, pH 8.3. After an exhaustive dialysis procedure (10,000 dilution factor), the active fractions were freeze dried and stored at −20 °C.

2.6. Protein assays, hemagglutination activity (HA) determination and SDS-PAGE analysis

The protein determinations were carried out by the method described by Lowry et al. [20] using bovine serum albumin (BSA) as a standard. The hemagglutination activity (HA) was measured by twofold serial dilutions of the samples on microtiter plates (96 × 96 wells) using 2.5% (v/v) of A1 human erythrocytes suspension (treated or not treated with trypsin) in 0.15 mol L⁻¹ of NaCl. The lectin samples were incubated with erythrocytes suspension for 60 min at room temperature and the hemagglutination activity units (HAU) were expressed by the reciprocal of the highest dilution with visible positive hemagglutination. The specific activity (HAU mg⁻¹) was defined as the ratio between HAU mL⁻¹ and the protein concentration mg mL⁻¹. The total activity was calculated by the product between the total lectin mass (mg) and the specific activity. The purification factor was determined by the ratio of the initial specific activity to the final specific activity.

The samples from each purification step were analyzed on 12.5% SDS-PAGE (10 cm × 14 cm plate), in accordance to the methodology previously described [21]. The gels were revealed, by silver [22] or Coomassie Blue staining [23] for purification evaluation and relevant molecular mass determination, respectively. For glycoprotein gel staining, the Schiff periodic acid (PAS) method was used [24].

2.7. Molecular mass determination

The lectin molecular mass was determined by electrospray mass spectrometry (ESI Q-TOF MicroTM, Micromass, UK) in the positive ion mode. Mass spectrometer calibrations were conducted by using sodium iodide in the 100–3000 m/z range. Lectin samples from the ion exchange final step (20–25 µg) was solubilized in 50 µL of 50% (v/v) acetonitrile in water solution plus 0.2% (v/v) formic acid and applied to the mass spectrometer by a syringe pump system at 10 µL min⁻¹ flow rate. The capillary voltage was 3.5 kV and cone voltage was 60 V. The spectrum data obtained was the result from 30 scans (2.5 s) combined. Original data (m/z) were treated (base-line subtraction, smoothing and centering) and analyzed by the Mass Lynx™ 4.0 software.

2.8. Determination of the relative mass of the native protein by size exclusion chromatography

The LMA native molecular mass determination was performed by size exclusion chromatography using a Sephacryl™ HR S-200 column (1.6 cm × 60 cm) coupled to a FPLC™ System at flow rate of 1.0 mL min⁻¹. Lectin sample solutions of 500 µL (about 1.5–3.0 mg mL⁻¹) were loaded in the chromatography system at pH 4.0 (0.05 mol L⁻¹ acetate buffer) and pH 8.3 (0.01 mol L⁻¹ NaH₂CO₃) for the native molecular mass determination and the oligomeric behavior at different pH values. The molecular mass standards used for column calibration were β-amylase (205 kDa); alcohol dehydrogenase (150 kDa); bovine serum albumin (66 kDa); ovalbumin (46 kDa); trypsinogen (25 kDa) and cytochrome C (12.5 kDa). The retention times plotted as a function of the logarithm of molecular mass and the respective linear regression obtained were used for
molecular mass and aggregation states determination for LMA at pH 4.0 and 8.0.

2.9. Heat and pH stability for LMA

The heat stability of the LMA was determined by incubation of aliquots of lectin solution (1.3–1.4 mg mL⁻¹) at different temperatures (70, 75, 80, 85 and 90 °C) for 10, 20, 30, and 90 min and the remaining hemagglutinating activity (HA) was determined. The results were expressed by relative percentage to a control sample (not thermally treated). For pH stability, 5.0 mL of purified active LMA (2.5–2.8 mg mL⁻¹) were dialyzed for 24 h against buffers with different pH: pH 2.0–3.0 (0.1 mol L⁻¹ glyceine buffer); pH 5–6 (0.1 mol L⁻¹ sodium acetate buffer); pH 7.0–9.0 (0.1 mol L⁻¹ of Tris-buffer); pH 10.0 and 12.0 (0.1 mol L⁻¹ glycine buffer); a 0.15-mol L⁻¹ NaCl solution was employed as control and reference of 100% of activity. After dialysis, the pH of all LMA samples was adjusted to 7.4 by another dialysis procedure for 24 h against 0.1 mol L⁻¹ of ammonium acetate buffer at pH 7.4. The hemagglutination assay was conducted at pH 7.4 and the results were expressed as percentage of residual activity relative to the control.

2.10. Influence of EDTA, Ca(II) and Mn(II) on LMA activity

The LMA purified (1 mg mL⁻¹) was incubated for 10 h against 50 mmol L⁻¹ EDTA solution at 4 °C with continuous shaking. After the test, the samples treated with EDTA were dialyzed exhaustively (dilution factor of 10,000) against 0.15 mol L⁻¹ of NaCl at 4 °C, and the hemagglutinating activity was assessed before and after the addition of 50 mmol L⁻¹ of CaCl₂ and MnCl₂. As a HA control sample, one aliquot of the same LMA solution was submitted to dialysis procedure against aqueous solution of NaCl 0.15 mol L⁻¹, 5 mmol L⁻¹ CaCl₂ and MnCl₂ (no EDTA treatment, only dialysis effect control).

2.11. Hemagglutinating activity of LMA on the O, A, B and AB human erythrocytes

The LMA specificity on the ABO system (human blood groups A1, A2, B and O) was assessed by using erythrocytes, treated or not with trypsin, from voluntary healthy donors. The HA experiment with native cells (not treated with trypsin) was performed as mentioned before. For the erythrocyte trypsin treatment, a 10% (v/v) erythrocyte suspension in 0.15 mol L⁻¹ NaCl was prepared with 0.1% (w/v) of commercial trypsin (Merck®). The trypsic suspension was incubated at 37 °C for 70 min with occasional stirring. The trypsin-treated erythrocytes (TTE) was washed three times with 0.15 mol L⁻¹ of NaCl solution by centrifugation at 1100 × g for 15 min, each time, followed by suspension at 0.15 mol L⁻¹ of NaCl. The final sediment of TTE was used for 2.5% (v/v) erythrocytes suspension (0.15 mol L⁻¹ of NaCl) for HA determination of the LMA purified samples.

3. Results

3.1. First steps of partial purification (crude extract, thermal precipitation of the crude extract and precipitation by cold ethanol)

The comparative results of partial purification of lectins from crude extract, thermal precipitation and ethanol precipitation are shown in Table 1. These procedures were considered satisfactory for the next step of purification.

3.2. Ion exchange chromatography of the 20–60% ethanol precipitated fraction

The resultant profile of purification by ion exchange chromatography (Fig. 1) showed four main peaks with satisfactory resolution among them. More data about this purification step and its efficiency are presented in Table 1.

3.3. Protein assay, hemagglutination activity determination and SDS-PAGE analysis

LMA peak at elution NaCl concentration of 0.2 mol L⁻¹ was chosen for all hemagglutination activity tests and characterization assays, because it comprises about 70% of the LMA total activity (Fig. 1) and the largest mass percentage, if compared to the other fractions (Table 1). The SDS-PAGE analyses for all fractions have shown the same double-band profiles (Fig. 2), with an average molecular mass value of 28 kDa.

![Fig. 1. Ion exchange chromatography (HiLoad™ 16/10 Q-Sepharose™ coupled in FPLC system) profile of the LMA crude extract precipitated at 20–60 % ethanol precipitation. Flow rate of 3.0 mL min⁻¹ and (●) detection at 280 nm (—) salt gradient of NaCl and (•••) log₂ of (HU).](image-url)
3.4. Molecular mass determination by size exclusion chromatography and mass spectrometry

Molecular mass of LMA determined by ESI-MS showed single Gaussians curves that, after being processed, give rise to molecular masses from 27 to 28 kDa (Table 2). By size exclusion chromatography some molecular masses values from 55 to 110 kDa (Table 3) were found depending on the experimental conditions (e.g., pH variation).

<table>
<thead>
<tr>
<th>Condition</th>
<th>Molecular mass (kDa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH 4.0</td>
<td>54 ± 1</td>
</tr>
<tr>
<td>pH 6.0</td>
<td>106 ± 5</td>
</tr>
<tr>
<td>pH 8.0</td>
<td>116 ± 3</td>
</tr>
</tbody>
</table>

Flow rate was 1.0 mL min⁻¹ and monitored by UV absorbance at 280 nm. The linear fitting obtained from molecular mass standards was defined by linear regression yielding the following equation: \( y = -0.0515x + 5.4978 \) (\( R^2 = 0.9915 \)). The data are expressed as the average of three independent experiments and one standard deviation.

Table 2

<table>
<thead>
<tr>
<th>Condition</th>
<th>Molecular mass (kDa)</th>
<th>DBL[a]</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;Truncated&quot; monomer (processed)</td>
<td>27,137.8 ± 3.5</td>
<td>27,139</td>
</tr>
<tr>
<td></td>
<td>27,186.3 ± 1.7</td>
<td>27,188</td>
</tr>
<tr>
<td></td>
<td>27,297.0 ± 8.7</td>
<td>27,349</td>
</tr>
<tr>
<td>Intact monomer (not processed)</td>
<td>28,269.8 ± 1.7</td>
<td>28,273</td>
</tr>
<tr>
<td></td>
<td>28,316.0 ± 1.3</td>
<td>28,314</td>
</tr>
<tr>
<td></td>
<td></td>
<td>28,432</td>
</tr>
</tbody>
</table>

[a] The molecular mass data for LMA are represented by average ± 1 standard deviation (\( n = 5 \)).

3.5. Heat and pH stability for LMA

Results for thermal stability of LMA as a function of the incubation time are shown in Fig. 3. It can be noticed that after 10 min incubation none differences are noted for all temperatures tested, but prolonged incubation led to significant differences between the samples in the assay in Fig. 4 the residual activity dependence as a function of pH is presented.

3.6. Influence of EDTA, calcium(II) and manganese(II) on LMA activity

The LMA hemagglutination activity on EDTA treatment and in the presence Ca(II) and Mn(II) metal ions are shown in Table 4. The LMA activity is null before EDTA treatment and the hemagglutination activity recovery is uncompleted after addition of any divalent metal ions.

The specificity study towards human ABO erythrocytes (Fig. 5) demonstrated that LMA has hemagglutinating activity only for A1 erythrocytes, as it was also verified for DBL [11,12].

4. Discussion

The M. axillare seed lectin (LMA) was isolated firstly by Haylett and Swart [11]. The purification procedure adopted was affinity
chromatography using the antigen A1 + substance H coupled to Sepharose 6B (Pharmacia™). Such procedure was the same technique used by Etzler and Kabat [12] to purify the *D. biflorus* lectin (DBL).

The affinity chromatography procedures are frequently used for lectin purification because of the relative facility to bind (covalently link) carbohydrates to insoluble resins employed for this technique. Thus, fast and efficient procedures can be performed for lectins purification in only one step. The isolation by affinity is possible in this case because the lectins do not suffer any modification after binding to carbohydrates [18].

However, the affinity chromatography processes are very expensive methods because they need specific materials with high purification degree that elevate substantially the process costs. Besides, the affinity column maintenance can be a difficult task when crude extracts from plant are loaded into the columns. Such samples contain, frequently, many pigments and oily components that can impregnate the resin and damage the affinity columns.

The LMA has the same specific properties of DBL, in this way the procedure based on the precipitation by ammonium sulfate and molecular exclusion chromatography in Sephadex G100™ followed by ion exchange in Carboxymethyl Cellulose [25] could be a simple alternative method for preparative purification of LMA. However, such process comprises three purification steps with difficult intermediate procedures, requiring desalination and concentration of large volumes of samples.

The new development strategy of purification resulted in a fast, simple and low-cost procedure that could obtain large amounts of the purified lectin (Table 1). The methodology consists of two precipitation steps (by temperature and by ethanol, Table 1) followed by ion exchange chromatography in Q-Sepharose™ pH 8.3 (Fig. 1). According to the proposed methodology, the LMA purification yields an average around 0.6% (w/w) (lectin/seed powder), while Haylett and Swart [11] isolated LMA by affinity chromatography, obtaining a 0.9% (w/w) yield of LMA. However, taking into account the readiness to obtain LMA by the new methodology proposed, this apparent yield deficit can be balanced by the methodology easiness and the low cost per purification cycle performed.

The inclusion of the thermal treatment as a purification step turned the crude extract (CE) protein profile simpler and increased the lectin specific activity, because the total mass protein was decreased. Subsequently, the ethanol precipitation step contributed for additional enrichment, as can be seen in the purification table (Table 1) and SDS-PAGE analysis (Fig. 2). The choice of commercial ethanol for protein precipitation (range of 20–60%) allowed some experimental advantages, such as the recovering of about 95% hemagglutinating activity and the dialysis suppression because ethanol can be removed by evaporation at reduced pressure. Furthermore, the first precipitation steps inactivate proteinases (trypsin-like) of the crude extract, thus preventing a possible LMA degradation.

As can be verified in Fig. 2, the protein profile from this step revealed just the LMA bands (relative mass = M_r = 28 ± 1 kDa). The ion exchange chromatography provides the final purification step, yielding three active LMA fractions named LMA 0.2, 0.3 and 0.4 mol L^{-1} elution salt concentration (Fig. 1).

Among them, LMA 0.2 mol L^{-1} was chosen for all tests and characterization assays, because it comprises about 70% of the LMA total activity and the largest mass percentage, if compared to the other fractions. Besides, the LMA 0.3 and 0.4 mol L^{-1} revealed some pigments that, possibly, interfered with the protein assay and with the hemagglutinating activity determination. Moreover, such latter fractions showed not reproducible data when they were submitted to different blood group assays using trypsin-treated erythrocytes (data not shown) while the LMA 0.2 mol L^{-1} had shown the expected results. On the other hand, the SDS-PAGE analyses for all fractions have shown the same double-band profiles (M_r = 28 ± 1 kDa, Fig. 2), which indicates to be the same LMA but with different charge density, considering ion exchange profile.

Thus, only LMA 0.2 mol L^{-1} was used for all assays and from now, it will be referred just as LMA.

The total LMA activity recovery from four repetitions of this purification methodology demonstrated similar results, and Table 1 is a representative and reproducible example of the purification. The LMA hemagglutinating activity remained unaffected at temperatures up to 50 °C for 90 min. This finding is in agreement with the high values of the denaturation temperature observed for LMA at differential scanning calorimeter (DSC) experiments (melting temperature-T_m, about 100 °C at pH 6.2, data not shown). This results show that LMA has high thermal stability. The thermally treated crude extract did not suffer significant HA decrease on heating up to 85 °C for 30 min (Fig. 3). It is important to point out that the temperature test around 85 °C is 15 °C lower than the T_m obtained by DSC experiments (100 °C), therefore the LMA at 85 °C is 100% of native state. Moreover, the SDS-PAGE of this purification step revealed good refinements at the protein profile (Fig. 2) with increased specific activity (Table 1).

The investigation of LMA activity (HA) after exposure to different pH values demonstrated that it presented 100% of activity between pH 6 and 8 (Fig. 4), but the HA is gradually reduced below pH 6.0 and above pH 8.0 with complete loss of activity at pH values between 10 and 12. These findings explain the results of isothermal titration calorimetry experiments performed with LMA and GalNac (data not shown), whose results showed lower binding constants.
at acid pH range (<5.5), as well as at basic pH range (>8.0). On LMA purification using affinity chromatography, Haylett and Swart [11] proposed the decreasing of pH up to 5.0 as an alternative to perform LMA elution, because to remove *D. biflorus* lectin (DBL) from the affinity column [12]. GalNac is useful in analytical scale but more expensive and not applicable for preparative chromatography. In the present study, it was verified that the pH decreasing affects the HA when LMA is incubated for long periods (24 h), thus the purification procedures for LMA must be made near neutral pH (6.0–8.0) for total HA maintenance.

The LMA activity shows to be dependent of divalent metal ions (Table 4). The EDTA treatment completely removed the LMA hemagglutinating activity in all the experiments performed (n = 4). The HA assessed before the treatment with CaCl$_2$ and MnCl$_2$ at 50 mmol L$^{-1}$ (dilution factor of 10,000), both together or separated, revealed that the total activity is not recovered, remaining about 20% of the total activity if compared with the control samples (n = 4). Interestingly, the Ca(II) alone induction alone induced a recovering of about 11% of LMA activity as compared to the control, which is similar to that previously observed with the DBL activity [26]. Such results suggest that the LMA requires both ions for optimal hemagglutinating activity. As discussed previously [4,27], the Ca(II) and Mn(II) metal ions, although not directly involved, are necessary for carbohydrate binding to leguminous lectins (metalproteins). Indeed, the metal ion removal process for *Erythrina speciosa* lectin [28], a Gal/GalNac lectin, led to a complete loss of hemagglutinating activity, which is in agreement with our LMA results, but the HA recovery was complete when Ca(II) and Mn(II)$^{2+}$ were added to *E. speciosa* lectin. Such different results could be related to LMA incapacity to recover the proper topological carbohydrate binding site, thus the complete activity could not be measured. In fact, Haylett and Swart [11] showed that the LMA has other divalent metal ion species, such as Mg(II) and Zn(II) present at different proportions, however not structurally identified. However the addition of these metal ions (at 5 mmol L$^{-1}$) did not cause a full recovering of LMA activity after the EDTA treatment, even with Ca(II) and Mn(II) present. Additionally, it must be pointed out that long dialysis procedures (>36 h) decreased the LMA activity by up to 10%. Thus, the LMA HA recovery could not be determined with real accuracy, even regarding control samples that had gone through the same dialysis procedure.

The molecular mass of the DBL monomers, determined by ESI-TOF, showed two distinct mass groups with mass difference about 1129, on average [29]. Such difference is due to the processing that takes place on the monomeric C-terminal and this cleavage is necessary for the proper DBL assembly. This processing removes about 10–11 amino acids from the C-terminus (an α-helix segment) [29]. The LMA molecular mass determination by ESI-TOF showed the same profile as that determined earlier for DBL [31] (Table 2).

The mass spectra data from LMA were grouped in the same way as performed for the DBL data, i.e., the molecular mass data were separated according to their dimensions considering 50 as a minimal relative mass difference. In this way, LMA can be considered identical to DBL. In fact, Haylett [11] demonstrated 100% of homology between the first fifty amino acid that comprise the N-terminal (an α-helix segment) [29]. The LMA molecular mass determination by ESI-TOF showed the same profile as that determined earlier for DBL [31] (Table 2).

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