Research Article



A New Galactose-Specific Lectin from *Clerodendrum infortunatum*.

Sukumaran Surya, Madhathilkovilakathu Haridas*

Inter University Centre for Bioscience and Department of Biotechnology and Microbiology, Kannur University, Thalassery Campus, Kannur 670661, India

**Corresponding author:* Madhathilkovilakathu Haridas, Inter University Centre for Bioscience and Department of Biotechnology and Microbiology, Kannur University, Thalassery Campus, Kannur 670661, India, Tel/ Fax: 049-0 2347394, E-mail: mharidasm@rediffmail.com

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Background: The ethno-medical significance of *Clerodendrum* genus raises the interest towards the characterization of its seed lectin by inexpensive and most effective technique.

Objective: The focus of this study is the purification, characterization, and evaluation of the antioxidant and antiproliferative potential of a galactose-specific lectin from *Clerodendrum infortunatum* L. seeds.

Materials and Methods: The crude extract, homogenized in 6 volumes of the saline containing 10 mM β -mercaptoethanol was subjected to pigment removal by Toyopeal HW-55 column prior to ammonium sulfate fractionation (40-80 %). The crude protein extract was then loaded to the gel filtration column Sephadex G-200 followed by affinity chromatography using activated galactose coupled Sepharose-4B.

Results: The SDS-PAGE analysis showed a single band of about 30 kDa which further determined by MALDI-TOF analysis. The MALDI-TOF spectra revealed that Clerodendrum infortunatum lectin (CIL) is a homo-tetramer of 120 kDa consisting of four identical subunits of 30 kDa. The haemagglutination inhibition assay was done with purified lectin by many sugars, among which N-acetyl-D-galactosmine (NAG), D-galactose and lactose exhibited high inhibition. NAG showed the highest inhibition amongst the tested sugars, having the minimum inhibitory concentration of about 0.97 mM. The lectin exhibited a moderate antioxidant activity with an IC₅₀ value of 6.1 ± 0.1 mg.mL⁻¹ and induced cell death with IC₅₀ of 82.8 µg.mL⁻¹ against human gastric cancer cell line, AGS, indicated the potential of CIL for clinical and therapeutic applications.

Conclusion: The present study demonstrated the moderate ability of the CIL to inhibit the growth of human gastric cancer cells, AGS either by causing cytotoxic or anti-proliferative effects. Thus, CIL due to its remarkable properties may be considered as a potential bio-molecule in tumor research and glycobiology.

Keywords: Anti proliferative, Clerodendrum. infortunatum Lectin, Glycoproteins, Haemagglutination, Lectins.

1. Background

Lectins are glycoproteins of non-immune origin (1, 2), possessing at least one non catalytic domain which binds reversibly to a specific monosaccharide or oligosaccharide (3). Owing to their cell agglutinating property they are commonly referred to as phyto-haemagglutinins (4, 5). The plant lectins have gained much attention because of their ubiquitous presence and applicability as glycoconjugates. Isolation and purification of lectins could be carried out through a variety of purification strategies depending on their

source, structure, physiochemical properties, specificity, and biological activity. Lectin sources of plants may be the seeds, roots, fruits, flowers or leaves. Legume lectins represent the largest and most thoroughly studied family of lectins which consists of two or four monomers, either identical or slightly different, each with a single, sugar binding site (6). Lectins play essential role in many biological processes like cell-cell recognition events such as host defense, fertilization, tumor metastasis, embryogenesis, mitogenic simulations, innate immune response, and protein trafficking (5, 7, 8). Lectins

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may serve as useful tools for characterization of the glycoproteins, histochemistry of the cells, tissues, and for examination of changes occurred on cell surfaces during pathological changes undergoing in malignancy from cell differentiation to tumor formation (7). The topology of the cell organelles with carbohydrates coat can be marked by specific surface binding property of the lectins. Such lectins are extensively used as cytochemical and histochemical probes to label mammalian cells (6). Certain plant lectins could be useful as tools to distinguish between malignant and benign tumors through their ability to determine the metastatic condition via their degree of glycosylation (9).

Elucidation of lectin-carbohydrate interactions at the molecular level (10) and the anti-tumor potential of lectins like Concanavalin A (Con A), Phaseolus vulgaris, phytohaemagglutinin-L (PHA-L), Wheat germ agglutinin (WGA), Polygonatum odoratum lectin (POL), Soyabean lectin and Griffonia simplicifolia agglutinin (GSA) by inhibiting the proliferation of the different malignant cell lines (11-17) has made the lectins extensively useful elements in the research and diagnostics. Con A, the first reported legume lectin, shows apoptosis in the human melanoma A375 cells and autophagy inducing activities mediated by the cytochrome c release. Con A entered the preclinical trials due to its apoptotic and autophagic properties (18). The Clerodendrum infortunatum (Family-Lamiaceae) is a flowering shrub found rich in constituents, such as clerodolone, clerodone, clerosterol, flavonoids, terpenoids, saponins, phenolics, and sugars (20,21). The genus is widely distributed in the tropical and subtropical regions. Moreover, it is used in many systems of the medicine such as Ayurveda, Homeopathy, Siddha, and Unani, for ailments like alopecia, diarrhea, skin disorders, asthma, and postnatal complications. It is well-known for its hepato-protective and antimicrobial activities (22-24). Consequently, the ethno-medical significance of the diverse species of the Clerodendrum genus raises the interest towards characterization of its seed lectin. Hence, the present work aimed to study a lectin from Clerodendrum infortunatum using a simple and most effective technique, mainly focusing on further characterization via determination of its antioxidant and antiproliferative potential towards gastric-carcinoma cell lines. The early reports on Clerodendrum trichotomum agglutinin (CTA) illustrates the purification and the primary characterization of a lectin from clerodendrum genus shown an insight towards this goal (25). In light of the earlier findings on plant lectins, we undertook

the present study to purify and characterize a novel lectin from *Clerodendrum infortunatum* (CIL) with the potential of the applications.

2. Objectives

A galactose specific lectin from *Clerodendrum infortunatum* was purified and characterized via biochemical means and its antioxidant and antiproliferative potential was discussed.

3. Materials and Methods

3.1. Preparation of Crude Protein Extract

Clerodendrum infortunatum seeds were collected from the nearby locality of Kannur University Campus in Kannur district, Kerala, India and authenticated by the authority of the Botany department, Govt. Brennen College, Dharmadam, Thalassery. A voucher specimen (MH/DBM 005 2012) was prepared and deposited at the Department of Biotechnology, Kannur University, Kerala for future reference. Samples were washed, air dried and stored at 4 °C. Seeds of Clerodendrum infortunatum (100 g) were homogenized in 6 volumes of the saline containing 10 mM β-mercaptoethanol in a mortar and pestle. In order to alleviate the enzymatic browning of the extract a small amount of polyvinylpyrolidine was added. The sonicated slurry was stirred at 4 °C overnight and filtered through cheese cloth. The slurry was subjected to centrifugation at 10,000 \times g at 4 °C for 30 minutes and filtered through Whatman No.1 filter paper for removal of floating lipids. The dark brown colored supernatant obtained after centrifugation was subjected to the pigment removal.

3.2. Pigment Removal from the Crude Extract

Pigment removal from the crude extract performed by the Toyopeal HW-55 column (Sigma-Aldrich, United States). The dark brown color of the extract, due to the presence of pigments, was found to be interfering with the conventional galactose specific columns. Removal of the pigments before performing size exclusion or affinity column chromatography was essential for the protein purification.

3.2.1. Activation of the Toyopearl HW-55 Column

The Toyopearl HW-55 (25 mL) was suspended in 40 mL aniline in the presence of sodium cyanoborohydride (450 mg) and was incubated at 37 °C for 24 hours. The unreacted formyl groups were reduced by adding sodium borohydride (250 mg). The extract was then passed through the activated Toyopearl HW-55 column

pre equilibrated with 6 % saline for effective removal of pigments hindering the lectin-binding capacity to the affinity adsorbent. Then early colorless filtrate obtained after passing through the absorbent column was used for purification of the lectin (25, 26).

3.3. Purification of the Lectin

Lectin from the crude seed extract of the *Clerodendrum infortunatum* was purified based on the protocol reported by Kitagaki *et al.* (25) with slight modifications. The crude seed extract was subjected to ammonium sulfate precipitation method and most of the lectins was precipitated at 40-80 % of ammonium sulfate solvent. Then the fraction was subjected to the centrifugation at 10,000 ×g at 4 °C for 15 min. The supernatant was discarded and the precipitate was collected and dissolved in 20 mM phosphate buffered saline (PBS), pH 7.2 and was subjected to the dialysis (dialysis tube cutoff- 20 kDa) with the same buffer for further purification.

3.3.1. Gel Filtration - Sephadex G-100

The crude extract was loaded onto the 20 mM PBS equilibrated Sephadex G-100 column and eluted at a flow rate of 8 mL per hour with the same buffer, at 4 °C. Fractions were collected and checked for haemagglutinating activity. The protein was monitored by measuring absorbance at 280 nm. The fractions with haemagglutination activity were pooled out and subjected to further purification events.

3.3.2. Affinity Chromatography - Galactose Coupled Sepharose-4B

For further purification of the protein, the pooled protein fractions showing haemagglutination, obtained after gel filtration were subjected to the affinity chromatography with galactose coupled Sepharose-4B as per the protocol of Franco-Fraguas et al. (27) with slight modifications. Sepharose-4B (20 g) was incubated in 0.4 M NaOH and 5% epichlorohydrin at 40 °C for 2 hours. The activated matrix was washed and incubated in 0.1 M NaOH and 10% galactose for 2 days at room temperature. Reactive epoxy groups were blocked by incubating for 4 hours at room temperature in 1 M ethanolamine. The fractions obtained after dialysis against 20 mM PBS, was applied to activated, galactose-coupled Sepharose-4B column, previously equilibrated with the same PBS at 4 °C. The adsorbed proteins were eluted from the column with the PBS containing 0.2 M galactose and checked for haemagglutinating activity. The fractions exhibiting haemagglutination were pooled, dialyzed, and subjected to the characterization. The end point of the assay, visually estimated after an hour and the minimum protein concentration required for agglutinating human Red Blood Cells (RBCs), was expressed as the haemagglutination titer (28, 29). The reciprocal of the highest dilution of the lectin exhibiting haemagglutination was defined as the haemagglutination titer. The specific activity is the number of haemagglutination titers per milligram of protein (30).

3.4. Biophysical and Biochemical Characterization of the Lectin and Molecular Weight Determination

3.4.1. SDS-Polyacrylamide Gel Electrophoresis (SDS PAGE)

The molecular weight of the purified lectin was determined by SDS-PAGE, using the broad range molecular weight marker on 12% polyacrylamide gel containing 1% SDS in the presence of β -mercaptoethanol at a constant voltage of 120 V (31, 32).

3.4.2. MALDI-TOF Mass Spectrum and Iso-Electric Focusing (IEF) Analysis

The mass spectrum was recorded on Shimadzu Biotech Axima-CFRTM Plus MALDI mass spectrometer (Shimadzu, Japan) in the linear positive ion mode using sinapinic acid as the matrix. The mass spectrum was acquired by using 19 KV acceleration voltage. A total of 500 scans were averaged for the final spectrum.

The pI of the lectin was determined by using Rotofor (preparative iso electric focusing cell-Bio-Rad). About 0.5 mL of the lectin extract was mixed with 1 mL of ampholyte (pH 3-10 Bio-Rad) and 18.5 mL of the distilled water. After degassing, the mixture was loaded onto a mini chamber with a syringe, and supplied a 12 W to the chamber. A gradual increase in voltage (from 300 V to 500 V) for 3 hours was monitored. After attaining resultant voltage stability, the separated samples were vacuum harvested as 2 mL fractions. The pH of collected fractions was read subsequently and assayed for haemagglutination activity.

3.4.3. Haemagglutination Activity Study

Haemagglutination assay was performed on microtitre plate using trypsinised human RBCs. The RBCs were washed twice in PBS to get a cell suspension of 2% in the same buffer. The haemagglutination activity of the protein solution after affinity column in PBS was determined. About 0.2 mL (2 mg.mL⁻¹) of the lectin solution was serially diluted with PBS and each dilution was mixed with equal volume of 2% RBCs suspension and incubated for about 1 hour at room temperature. Results were examined visually (33).

3.4.4. Haemagglutination Inhibition Study

The haemagglutination inhibition test was done in the presence of series of sugars including NAG, D-galactose, lactose, D-fructose, D-mannose, D-galactosamine, D-arabinose, D-glucose, maltose, D-fucose, D-ribose, and sucrose of about 0.2 M concentration. The assay was performed on the microtitre plate by adding 50 μ L of the sugar solution (0.2 M) serially diluted by PBS followed by the addition of 50 μ L of the purified lectin (2 mg.mL⁻¹), needed for visible agglutination, to each well. After incubation at 37 °C for an hour, 50 µL of the 4% human RBC was added and the mixture was allowed to stand for further 1 hour at room temperature. Reactions were compared with a negative control (50 μ L sugar solution + 50 μ L PBS + 50 μ L 4% human RBC). The minimum concentration of sugar required to inhibit haemagglutination was recorded. The IC_{100} is the minimum concentration required for 100% inhibition of the CIL (34).

3.4.5. Effect of Temperature and pH on the Lectin-Induced Haemagglutination

Thermal denaturation study was performed by incubating the purified lectin (2 mg.mL⁻¹) at various temperatures, ranging from 10 °C to 100 °C for 30 minutes. The pH dependent haemagglutination activities of CIL were performed by pre-incubating the samples at different pHs, ranging from 2 to 10. A buffer cocktail including 50 mM glycine-HCl (pH 2-3), 50 mM sodium acetate/acetic acid (pH 4-5), 50 mM sodium phosphate/HCl (pH 6-7), 50 Mm Tris-HCl (pH 8) and 50 mM glycine-NaOH (pH 9-10) were used for testing pH stability. The thermal and pH dependent haemagglutination treatments were performed with 50 μ L aliquots of the lectin solution at room temperature. A graph was plotted for pH/ temperature vs. haemagglutination activity.

3.5. Determination of Total Sugar Content of the Lectin The total neutral sugar content of the purified lectin was estimated by anthrone method using D-glucose as the standard (35). CIL solution (2 mL, 2 mg.mL⁻¹) was hydrolyzed/denatured into its monomers by keeping it in a boiling water bath for 3 hours in the presence of 5 mL of 2.5 N HCl. It was then neutralized by adding pinches of solid sodium carbonate until the effervescence ceased. The supernatant after centrifugation was used for further analysis. Anthrone gave colored compound with simple sugars, the digested products. The amount of total carbohydrate present in the test sample was estimated from the absorbance plotted against a glucose standard curve at 578 nm.

3.6. DPPH Free Radical-Scavenging Assay

The 2,2-Diphenyl-1-picrylhydrazyl (DPPH) free radical scavenging activity of CIL was performed according to Paixao et al. (36). CIL solutions of the various molarity (i.e., 2, 4, 6, 8, and 10 mg.mL⁻¹ in 20 mM PBS, pH 7.2) were added to an equal volume of (10 mL) methanolic solution of DPPH (400 µg.mL⁻¹). Ascorbic acid (2-10 mg.mL⁻¹) was considered as reference standard. After incubation at room temperature in dark for 30 min, the absorbance of the resulting solution was measured at 517 nm using UV-Vis Spectrophotometer (Model. U.2800, Hitachi, Japan) against the control. An equal volume of DPPH solution was taken as the control for the radical scavenging assay. All experimental measurements were carried out in triplicate and expressed as average of three analyses \pm standard deviation. The free radical scavenging activity was measured by the following equation. IC50 value, defined as the concentration of antioxidant, required for 50% scavenging of DPPH radicals in the specified time period, was determined graphically (37, 38).

DPPH radical-scavenging activity (%) =

 $(A_c - A_s) / A_c \times 100$ Equation 1 where, $A_c =$ Absorbance of control, $A_s =$ Absorbance of sample.

3.7. Assay of Anti-Proliferative Activity on Gastric Cancer Cell Line

3-(4, 5-dimethylthiazol-2-yl)-2,5-diphenylt The etrazolium bromide (MTT) assay was performed with human gastric cancer cell line, AGS obtained from ATCC. The AGS cell line was cultured in Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% fetal bovine serum. For assay on AGS cell lines the cells were adjusted to a cell density of 8×10^4 cells/mL in DMEM. The cells (100 μ L) were seeded onto each well of a 96-well plate and incubated overnight at 37 °C in the presence of 5% CO₂. Varying concentrations of CIL prepared using 2-fold serial dilution of about 3.1-200 µg.mL⁻¹ were added to each well followed by incubation at 37 °C, 5% CO₂ for 48 hours. After incubation, the medium was discarded and 25 µL of MTT (5 mg.mL⁻¹) was added to each well, followed by incubation for 3 hours. Dimethyl sulfoxide (DMSO) of about 150 µL was added to each well to dissolve the MTT formazan formed at the bottom of the wells. All measurements were carried out in triplicate and are expressed as average of three analyses \pm standard deviation. The optical density (OD) was measured at 570 nm and the percentage inhibition of the cells was calculated by the following equation (39).

Fractions	Total Protein (mg)	Haemagglutination activity (titer)	Specific activity (titer.mg ⁻¹)	Yield (%)	Purification fold
1. Crude extract	600	4500	7.5	100	1
2. After Toyopeal HW-55 column	552	4250	7.7	94.4	1.023
3. After 80% $(NH_4)_2SO_4$ fraction	210	3281	15.62	72.91	2.082
4. After Sephadex G-100 column	62	1937	31.24	43.04	4.165
5. After affinity column	12.5	895	71.6	20.00	9.54

Table 1. Purification yields of various chromatographic fractions of CIL.

The percentage of inhibition (%) = Treated OD value / OD value of the untreated control $\times 100$. Equation 2

4. Results

4.1. Purification of CIL Lectin

The 80% ammonium sulfate fractionated crude protein extract after dialysis against 20 mM PBS was applied to an activated pigment absorbent column Toyopeal HW-55. The brown colored pigment in the crude protein extract was effectively adsorbed by Toyopeal HW-55. Almost a colorless crude protein effluent was subjected to the Sephadex G-100 column at 4 °C which was previously equilibrated with the same buffer. The active fractions showing haemagglutination activity were pooled and purified further by affinity chromatography on activated galactose coupled Sepharose-4B. The other fractions devoid of haemagglutination were not used for further analysis as they contained a small amount of protein having significant biological activity. The overall purification data was summarized in Table 1.

4.2. SDS-PAGE

The fractions obtained after affinity chromatography on galactose coupled Sepharose-4B column gave a single band of about 30 kDa on 12% SDS-polyacrylamide gel electrophoresis in the presence of β -mercaptoethanol indicating that it was a pure protein (**Fig. 1**).

4.3. MALDI-TOF Mass Spectrum and Iso-Electric Focusing (IEF) Analysis

The MALDI-TOF mass spectrum analysis of the CIL showed prominent peaks at 30 kDa and 120 kDa indicated the presence of homodimers. Therefore, it was revealed

that CIL is a homo dimer and form tetrameric association in solution with the apparent molecular weight of 120 kDa (Fig. 2).

After running preparative IEF, the pH of the harvested protein was monitored and recorded. IEF revealed that the



Figure 1. Determination of molecular weight of CIL by SDS-PAGE (12% gel with 1% SDS in the presence of β -mercaptoethanol). Lane A: Broad range molecular weight markers, Lane (B-D): 10, 15 and 20 μ L of purified CIL. A single band corresponding to 30 kDa was obtained for CIL.



Figure 2. Determination of molecular weight of CIL by MALDI-TOF analysis using sinapinic acid as the matrix.

CIL is a basic lectin of pI 8.4. Subsequently the fractions were also assayed for haemagglutinationactivity. The haemagglutinationactivity vs pHs of the corresponding Rotofor fractions were summarized in the graph (Fig. 3).



Figure 3. Iso-electric focusing profile of CIL by using Rotofor with gradual supply of a voltage from 300 V to 500 V for three hours.

4.4. Haemagglutination and Haemagglutination Inhibition Studies

The purified CIL agglutinated the trypsinised human RBCs in microtitre plate with visible agglutination mass. The NAG was strong inhibitor amongst the studied sugars with the minimum inhibitory concentration of about 0.97 mM. It was found that D-galactose and lactose showed haemagglutination inhibition at 5.60 and 7.80 mM concentrations. Other sugars such as D-fructose, D-mannose, D-galactosamine and D-arabinose showed characteristic minimal inhibitions when compared to non inhibitory sugars (NI) mentioned in **Table 2**. Glucose, maltose, D-fucose, D-ribose and sucrose were NI up to 300 mM concentrations.

4.5. Effect of Temperature and pH on Lectin-Induced Haemagglutination

Thermal denaturation studies showed that CIL lost its agglutinating activity completely at 100 °C and 50% of activity lost at 50 °C. The pH dependent studies on haemagglutination activity of CIL revealed that the

Table 2. Haemagglutination inhibition assay of CIL with different sugars (0.2 M) and their corresponding IC100 values.

Sugars studied	IC ₁₀₀ (mM)	
N-acetyl-D-galactosamine	0.97	
D-galactose	5.6	
Lactose	7.8	
D-fructose	122.00	
D-mannose	124.00	
D-galactosamine	250.00	
D-arabinose	250.00	
D-glucose, Maltose, D-fucose, D-ribose, Sucrose	NI	

optimum pH for lectin activity was between pH 7 and 8 respectively. In acidic of < 7 and basic pH of > 8 the haemagglutinating activity of lectin was greatly reduced (Fig. 4).

4.6. Total Sugar Content of Lectin

Total sugar estimation by Anthrone method gave sugar content of 8.8%, at the initial protein concentration of 0.5 mg.mL⁻¹. The sugar concentration derived from the standard glucose plot was 0.044 mg.mL⁻¹.

4.7. Antioxidant Activity Estimation by the DPPH Radical-Scavenging Assay

The DPPH radical-scavenging ability of the CIL was determined and compared with the standard antioxidant, ascorbic acid. The free radical scavenging activity of CIL showed the IC₅₀ value of 6.1 ± 0.1 mg.mL⁻¹ while the standard ascorbic acid exhibited 5.7 ± 0.1 mg.mL⁻¹.



Figure 4. Effect of temperature and pH on the haemagglutination activity of CIL (2 mg.mL⁻¹). The hemagglutination activity was preserved at or below 50 $^{\circ}$ C and at pH 7 and 8



Figure 5. Estimation of antioxidant activity of CIL by using ascorbic acid (2-10 mg.mL⁻¹) as standard measured spectophotometrically at 517 nm.

The CIL exhibited significant antioxidant activity when compared to the standard ascorbic acid (Fig. 5).

4.8. Assay of Anti-Proliferative Activity on a Gastric Cancer Cell Line

The in vitro MTT assay disclosed that treatment with CIL displayed anti-proliferative activity against gastric cancer cell line, AGS with IC_{50} of 82.8 µg. mL⁻¹ showing inhibition of cell growth indicating a moderate anti-proliferative activity after incubation for about 48 hours (**Fig. 6**).



Figure 6. Determination of anti-proliferative activity of CIL against gastric cancer cell line, AGS and the optical density measured at 570 nm.

5. Discussion

The legume lectins represent the largest and most thoroughly studied family of lectins with high sequence similarities, very similar tertiary structures and biophysical properties (45). Some species of Lamiaceae are used in folk medicines and lectins are widely distributed in them also (46, 47). Since the *Clerodendrum infortunatum* belongs to the Lamiaceae family, its lectin also sheds new incrementing light on the specificity, physiological role and evolution of the classical legume lectins. The purification strategy of CTA provoke us to characterize a novel lectin from *Clerodendrum infortunatum* with therapeutic potential (25). According to the literature, most of the CIL got precipitated in 40-80% of ammonium sulphate fraction with effective pigment removal by activated Toyopearl HW-55 (25). The SDS-PAGE and MALDI-TOF spectra revealed that CIL is a homotetramer of 120 kDa consisting of four identical subunits of 30 kDa. The optimum pH of CIL was between 7 and 8 whereas its 50 % of activity lost at 50 °C, which correlates well with the previous reports (46).

The antioxidant activity reported for diverse species of Clerodendrum genus encouraged the assessment of free radical scavenging potential with CIL (20). Reports on the antioxidant activity of lectins are scares. The CIL exhibited moderate antioxidant activity with IC_{50} value of 6.1 ± 0.1 mg.mL⁻¹. The trend showed that CIL moderately inhibited proliferation of AGS cell lines in a dose- and time-dependent manner and is obviously in the range with IC_{50} of 82.8 µg. mL⁻¹ when compared with early known lectin reports. In fact, CIL is one among the few lectins by far presenting antitumor properties, as assumed from its anti-proliferative potential. The antitumor potential of Chinese pinto bean lectin (CPBL) against nasopharyngeal carcinoma, (HONE-1 cells), Lotus corniculatus lectin against human lung cancer cells (HCT116) and Phaseolus acutifolius agglutinin (PAA) against human colon adenocarcinoma cell line (Sw480) with IC₅₀ values of 30.8 μ M, 60 and 84.2 μ g. mL⁻¹ respectively has been reported earlier (29, 40-43).

6. Conclusion

The critical part involved in the purification of thenew lectin, CIL was the removal of pigments coexisting in the fruit, along with the lectin by Toyopearl HW-55. The lectin got precipitated in 40-80% of ammonium sulphate fractionation, followed by the effective purification with the Sephadex G-100 and the galactose coupled affinity column Sepharose-4B. The MALDI-TOF mass spectrum analysis, demonstrated CIL as a homo tetramer of 120 kDa. The purified CIL showed specificities towards different sugars like NAG, galactose and lactose in the decreasing order. The CIL exhibited DPPH radical-scavenging ability with IC_{50} value of 6.1 \pm 0.1 mg.mL⁻¹.The present study demonstrated the moderate ability of CIL to inhibit the growth of human gastric cancer cells, AGS either by causing cytotoxic or anti-proliferative effects with

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 IC_{50} of 82.8 µg.mL⁻¹. Thus, CIL may be considered as a potential candidate with biomedical applications in immunomodulation and tumor research. This was a basic approach and still more research, including *in vivo* analysis is desirable.

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