Physicochemical characterization of C-phycocyanin from *Plectonema* sp. and elucidation of its bioactive potential through *in silico* approach

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**ABSTRACT**

C-phycocyanin (C-PC), the integral blue-green algae (BGA) constituent has been substantially delineated for its biological attributes. Numerous reports have illustrated differential extraction and purification techniques for C-PC; however, there exists paucity in a broadly accepted process of its isolation. In the present study, we reported a highly selective C-PC purification and characterization method from nontoxic, filamentous and non-heterocystous cyanobacterium *Plectonema* sp. C-PC was extracted by freeze-thawing, desalted and purified using ion-exchange chromatography. The purity of C-PC along with its concentration was found to be 4.12 and 245 µg/ml respectively. Comparative characterization of standard and purified C-PC was performed using diverse spectroscopic techniques namely Ultra Violet-visible, fluorescence spectroscopy and Fourier transform infrared (FT-IR). Sharp peaks at 620 nm and 350 nm with UV-visible and FT-IR spectroscopy respectively, confirmed amide I bands at around 1638 cm⁻¹ (C=O stretching) whereas circular dichroism (CD) spectra exhibited α-helix content of secondary structure of standard 80.59% and 84.59% of column purified C-PC, SDS-PAGE exhibited two bands of α and β subunits 17 and 19 kDa respectively. HPLC evaluation of purified C-PC also indicated a close resemblance of retention peak timing of 1.448 min, 1.233 min and 0.925 min. As a cautious approach, the purified C-PC was further lyophilized to extend its shelf life as compared to its liquid isoform. To evaluate the bioactive potential of the purified C-PC *in silico* approach was attempted. The molecular docking technique was carried out of C-PC as a ligand-protein with free radicals and α-amylase, α-glucosidase, glycogen synthase kinase-3 and glycogen phosphorylase enzymes as receptors to predict the free radical scavenging (antioxidant) and to target antidiabetic property of C-PC. In both receptors free radicals and enzymes, ligand C-PC plays an important role in establishing interactions within the cavity of active sites. These results established the antioxidant potential of C-PC and also give a clue towards its antidiabetic potential warranting further research.

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**Introduction**

Cyanobacteria are microalga that serves to be the primary producer in aquatic ecosystems which in turn are cosmopolitan within the biosphere and exhibit substantial diversification as compared to their auxiliary prokaryotic counterparts (1, 2). These prokaryotes are predominantly constituted by C-phycocyanin (C-PC), a photosynthetic pigment responsible for imparting their characteristic blue color (3). Cyanobacterial biomass as well as bioactive products derived from it have been analyzed and well documented in pharmaceuticals, cosmeceuticals, nutraceuticals, health supplements, aquaculture, bioremediation, bioenergy, biofuels and the food industry (4-7).

For a long, researchers have been attracted by the *Plectonema* sp., a filamentous cyanobacterium that is non-toxic and non-heterocystous in nature. Its fast doubling time of approximately 60 hours and minimal requirement for nitrogen sources and light has made it
an important experimental candidate (8-10). *Plectonema* sp. possesses a great variety of pigments namely carotenoids, chlorophyll and phycobiliproteins (PBPs) (11). Among all these members, carotenoids are usually considered to be essential and are structurally related to isoprenoid molecules. The carotenoids are often referred to as pigments owing to their light-absorbing properties (12). Chlorophyll within *Plectonema* sp. plays an additional chelating role which in turn is pivotal for cell repair, increases hemoglobin in the blood, fastens cell growth, aids liver cirrhosis cure an ulcer in humans (13).

PBPs are a group of pigmented proteins present in blue-green algae (cyanobacteria) and also in cryptomonads, red algae, etc (14). Generally, PBPs are mainly classified into three types, such as C-phycocerythrin (C-PE), C-phycocyanin (C-PC) and C-allophycocyanin (C-APC) (15, 16). PBPs differ in their spectral properties and are classified based on absorption spectra: C-PE (Red; \( \lambda_{\text{max}} = 540 – 570 \) nm), C-APC (Blue with a green signal; \( \lambda_{\text{max}} = 650 – 665 \) nm) and C-PC (Brilliant blue; \( \lambda_{\text{max}} = 610 – 620 \) nm) (17). PBPs are hydrophilic and stable fluorescent proteins, having high concentrations resulting in the bluish appearance of the organism and participate in energy transfer processes during photosynthesis (18).

The fluorescent nature of PBP is exploited in many fields such as research, diagnostics and therapeutics (19, 20). In PBP, the two subunits \( \alpha \) and \( \beta \) polypeptide chains are present in equal amounts (21). Subunit \( \alpha \) has a low molecular weight (MW) of \( 12 – 19 \) kDa while \( \beta \) is 14 – 21 kDa. These subunits can assemble in various combinations of the trimer \((\alpha\beta)_3\) (about 120 kDa) or hexamer \((\alpha\beta)_6\) (about 240 kDa) forms (22, 23).

Commercially, C-PC is a highly important natural product with potential biotechnological applications in cosmetics, food, biotechnology, medicine and pharmacology (24). C-PC as non-carcinogenic and non-toxic natural food colourants are gaining remarkable prominence in view of synthetic food colorants, furthermore, their therapeutic value has also been proven (25, 26). Based upon differential purity ratio levels C-PC is classified as reactive grade (0.7), food grade (3.9) and analytical grade (ratio > 4) (27).

Diabetes mellitus is a major metabolic syndrome characterized by high blood glucose levels caused due to enhance the production of hepatic glucose, pancreatic impaired insulin production, and insulin resistance (28). Several endogenous and exogenous factors are attributed to the pathogenesis of diabetes mellitus. As an endogenous factor, oxidative stress is considered an important determinant of type 2 diabetes mellitus (29). Overproduction of free radicals such as hydroxyl radicals, superoxide radicals, hydrogen peroxide and nitric oxide radicals creates an imbalance between antioxidants inside the body resulting in oxidative stress (30, 31). On the other hand, blood sugar levels are severely regulated by the action of several enzymes such as \( \alpha \)-amylase, \( \alpha \)-glucosidase, glycogen synthase kinase-3 and glycogen phosphorylase. \( \alpha \)-amylase is responsible for breaking down long-chain carbohydrates (32), while \( \alpha \)-glucosidase directly converts carbohydrates into glucose in the small intestine. Glycogen synthase kinase-3 (GSK-3) is a serine/threonine kinase implicated in the development of insulin resistance, mainly based on its role in the regulation of glycogen synthesis (33). Whereas, glycogen phosphorylase (GP) catalyzes the breakdown of glycogen to glucose-1-phosphate in the liver and tissue to generate metabolic energy. In the liver, glucose-1-phosphate is frequently converted by phosphoglucomutase and glucose-6-phosphatase to glucose, which is released for the benefit of other tissues (34). The inhibition of \( \alpha \)-amylase, \( \alpha \)-glucosidase, GSK-3 and GP and free radical can be recognized as a therapeutic target for the control of type 2 diabetes.

Many synthetic drugs are available as an antioxidant and for the treatment of diabetes, although these are fraught with side effects (35). Therefore, recent research is directed towards treatments using natural materials that have fewer side effects (36). Cyanobacterial bioactive compounds have nowadays become an attractive source of functional compounds, including antioxidants and anti-diabetic therapeutic agents. C-PC the most dominant pigment-protein complex of cyanobacteria has also been analyzed and documented in pharmaceuticals, cosmeceuticals, nutraceuticals, health supplements and biomedical research (14). As represented in our previous study, eight cyanobacterial strains were screened to evaluate the maximum crude C-PC producing strain. *Plectonema* sp. was found to be the superior producer of C-PC (37) and thus this strain has been used further for purification and characterization of C-PC. Despite
such inherent potential of *Plectonema* sp. there exist paucity in the elucidation of optimum purification and characterization of C-PC from the same. C-PC is commonly extracted by freeze-thawing, concentrated by ammonium sulphate fractionation, purified by ion-exchange chromatography and more stabilized by lyophilization/freeze-dry method and characterized by Ultraviolet-visible spectroscopy (UV-vis), Fluorescence Spectroscopy, Fourier Transform Infrared (FT-IR) Spectroscopy, Circular Dichroism (CD) Spectroscopy and Sodium Dodecyl Sulphate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) and High-performance liquid chromatography (HPLC). In light of these critical techniques, in the present study, we tried to optimize the standard critical conditions such as differential extraction processes and their after-effects on downstream processing procedures critically mediating C-PC yield and quality from *Plectonema* sp. Further, we investigated the free radical scavenging (antioxidant) and antidiabetic potential of ligand C-PC through molecular docking techniques by inhibiting targeted free radicals and enzymes.

**Materials and methods**

**Materials**

Dialysis membrane (3.5 kDa), dipotassium hydrogen phosphate (K₂HPO₄), potassium dihydrogen phosphate (KH₂PO₄), methanol, glacial acetic acid, sodium chloride (NaCl), sodium hydroxide (NaOH), ammonium persulphate (APS), bis-acrylamide, glycerol and TEMED (N, N, N, N-tetramethylethylendiamine) were purchased from HiMedia. Sodium dodecyl sulfate (SDS) and coomassie blue dye were obtained from G-Biosciences, whereas bromophenol blue and glycerol were purchased from Rankem. A pre-stained protein marker was obtained from PUREGENE and trichloroacetic acid was purchased from Fisher Scientific. Diethylaminoethyl cellulose (DEAE-C) column matrix, ethidium bromide, EDTA and acetonitrile were purchased from Sigma-Aldrich and, glycine and acrylamide were obtained from Merck. Analytical-grade chemicals were used in this study.

**Methods**

**Experimental Cyanobacterium**

*Plectonema* sp. used during the present study was generously provided by Dr. S. M. Prasad, Department of Botany, University of Allahabad.

**Culture Maintenance**

Cyanobacterial culture (*Plectonema* sp.) was cultured under standard conditions (Temperature = 27 ± 2 °C). For regular experiments slightly modified BG-11 medium was utilized as described in our previous study, including 14/10 hrs of light and dark photoperiod respectively; pH = 7.0 along with 2400 Lux of light intensity (37-39).

**Crude Extraction and Spectrophotometric Estimation of C-PC**

Approximately two-week log phase culture was homogenized and centrifuged (5,000 rpm, 10 minutes, 4 °C). Finally, the resultant pellet was washed using phosphate buffer (10 mM; pH 7.4) having 50 mM NaCl and 0.002 M sodium azide (NaN₃) (40). The pellet was again resuspended in the same buffer with the addition of fine glass powder and then frozen at −20 °C. Repeated freeze-thawing of tubes containing pallets was done to accumulate phycobiliproteins in phosphate buffer. After freeze-thaw, crude C-PC was obtained as blue supernatant by centrifugation at 12,000 rpm for 15 minutes at 4 °C. To prevent denaturation of C-PC, 1µl / ml of PMSF (Phenyl methyl sulphonyl Fluoride) (100 µg/ml) was added (41).

Estimation of the amount of C-PC and purity were determined as described by (42, 43).

\[ C_{\text{C-PC}} = \frac{A_{(620)} - 0.474 \ A_{(652)}}{5.34} \]

Where \( C_{\text{C-PC}} \) is the concentration of C-PC in µg/ml, \( A_{(620)} \) is the absorbance of the sample at 620 nm and \( A_{(652)} \) is the absorbance at 652 nm.

**Purity** = \( A_{620}/A_{280} \)

Where \( A_{620} \) is representative of the maximum C-PC absorber peak and \( A_{280} \) is an indication of undesired protein especially aromatic amino acids.

**Purification of Crude C-PC**

**Ammonium Precipitation**

C-PC crude extracts were separated through gradual precipitation by ammonium salts up to 60 % saturation. The sample was further left undisturbed overnight (4 °C). The precipitate was further
harvested by centrifugation (10,000 rpm; 10 minutes; 4 °C temperature). The resulting supernatant (clear) was discarded and a blue pallet was collected containing C-PC (44).

**Dialysis**

To increase the level of purity of C-PC, dialysis was performed by suspending ammonium sulfate precipitated pallets in PBS (10 mM; pH = 7.4). The dialysis membrane (3.5 kDa, Himedia) was pretreated to remove membrane-bound contaminants as previously described (45). Briefly, the resuspended sample was subjected to dialysis in 100 − 200 folds of a volume of PBS (10 mM; pH = 7.4).

**Ion Exchange Chromatography**

The dialysed C-PC was subjected to ion-exchange chromatography to increase its purity by employing DEAE cellulose-packed columns. Dialysed C-PC binds this positive charge matrix and was thus purified according to the procedure described by (46). After incubation of dialysed C-PC with that DEAE-C column at 4 °C was briefly cleaned using appropriate buffer (10 mM; pH = 7.4) to eliminate undesirable C-PC proteins. Nearly different fractions of C-PC were subjected to UV−vis spectrophotometer for their absorbance in the range of 250 − 750 nm (eppendorf BioSpectrometer, Germany) and their absorption were specifically recorded at wavelengths i.e. 280, 562, 620 and 652 nm.

**Characterization of Purified C-PC**

**UV-visible Spectroscopy**

Absorbance spectra of purified and standard C-PC were recorded in the range of 250 − 750 nm using a UV-vis spectrophotometer (Eppendorf Biospectrometer, Germany). The purity of C-PC was calculated as stated above in section 2.2.3.

**Fluorescence Spectroscopy**

Fluorescence emission spectra of commercially available as a standard and all stages of purified C-PC samples were recorded in the range 290 − 450 nm. C-PC samples were excited at 280 nm and emissions were recorded at 350 nm on Agilent carry eclipse fluorescence spectrophotometer. Fluorescence emission at 350 nm confirmed that the purified sample was the protein that was C-PC (47).

**Fourier Transform Infrared Spectroscopy (FT-IR)**

Transmission profiles of available standard and all stages purified C-PC were recorded at 450 − 4000 cm⁻¹ on a spectrum FT-IR spectrometer (Perkin Elmer, Inc., USA) (48).

**Circular Dichroism (CD)**

CD spectra of standard, ammonium precipitated, dialysed and column purified C-PC were recorded as per the protocol described by (49). Briefly, different extracts were placed in a temperature-controlled cell holder (25 °C) enclosed with a water bath (Neslab’s RTE 110). All the samples were evaluated at a wavelength ranging from 190 to 250 nm (scan speed 20 nm per minute).

**Sodium Dodecyl Sulphate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) Analysis**

Purified cyanobacterial protein was qualitatively evaluated using SDS-PAGE for their molecular weight according to the previously described protocol (50). To confirm the purity of C-PC, 12 % SDS-PAGE was performed under non-denaturing conditions. The bands were visualized by Coomassie blue staining. The molecular weight of the purified C-PC was determined by running a pre-stained protein marker (PUREGENE) along with the sample (51).

**High-Performance Liquid Chromatography (HPLC)**

Standard and extracted column purified C-PC samples were further characterized using HPLC (51). C-18 reverse phase column was used during HPLC analysis (Shimadzu-LC-20AD, Japan). Prior to an assessment, 20% acetonitrile with 0.1% trichloroacetic acid was used to equilibrate the C-18 reverse-phase column.

**Freeze-Drying: Stepwise Process**

The column eluted C-PC protein obtained was in a dilute state and further characterization, assays for its bioactive properties due to its low concentration were inconceivable, which instigated us to find a way to concentrate the obtained protein samples.

The best available and economical method to solve this problem was to perform lyophilization or freeze-drying of C-PC proteins. Briefly, the protein samples
were allowed to freeze at −80°C for 12 hours. The frozen samples were then attached to the freeze-drying vacuum nozzle in the freeze dryer flask and the freeze dryer was switched on. A temperature of −40°C was maintained in the cooling well and with a negative pressure of 50 mTorr and left for ~20−24 hours to complete a cycle.

**Molecular docking study**

**Molecular Docking Analysis of C-PC and Free Radicals**

The 3-dimensional (3D) structure of C-PC as a ligand for scavenging and anti-diabetic agent were taken from the protein data bank (PDB) with PDB ID: 4L1E [Figure 1 (A)] for the docking study. Protein was prepared by deleting water, ligands and all the chains except A chain. The targeted structures of hydroxyl radical (CID: 961), superoxide radical (CID: 5359597), hydrogen peroxide (CID: 784), and nitric oxide radical (CID: 145068) was obtained from the PubChem database. The MMFF94 force field was used for the energy minimization of the target molecule. Gasteiger partial charges were added, non-polar hydrogen atoms were merged, and rotatable bonds were defined. Docking calculation was carried out on the protein molecule. Essential hydrogen atoms, Kollman united atom type charges, and salvation parameters were added with the aid of AutoDock tools. Grid maps of 60x60x60 Å3 grid points were generated with the help of the Auto grid program. C-PC was embedded in a 3D grid, and a probe atom was placed at each grid point. AutoDock parameter set and the distance-dependent dielectric functions were used in the calculation of the Van der Waals and the electrostatic terms, respectively. The torsion of the ligand molecule was set randomly. Each docking experiment was derived from 10 different runs that were set to terminate after a maximum of 2,500,000 energy evaluations. The population size was set to 150. The final [Figure 1 (B)] was generated with the help of Discovery Studio Visualizer (Accelrys) (30, 52).

**Molecular Docking Analysis of C-PC and Enzymes**

Protein-protein molecular docking was performed via the ZDOCK technique to figure out the protein-protein association between C-PC as a ligand-protein molecule and α-amylase, α-glucosidase, GSK-3β and GP as targeted protein molecules. C-PC (PDB ID: 4L1E) structures were obtained from PDB. α-amylase (PDB ID: 4W93), α-glucosidase (PDB ID: 3WY1), GSK-3β (PDB ID: 3F7Z) and GP (PDB ID: 1L5Q) structures were obtained from PDB. ZDOCK is one of the most unbeaten suites that encompass large calculation facilities in Critical Assessment of Predicted Interactions (CAPRI) (53). ZDOCK is an original phase rigid body molecular docking algorithm that utilizes a fast Fourier transform (FFT) calculation to calculate progress performance for translational searching (54, 55).

**Statistical Analysis**

The data were represented as mean ± standard deviation (SD). Statistical significance of the results was determined by using one-way ANOVA followed by Tukey post-test using graph pad Prism (version 5.01), where ns = non-significant, *p < 0.05, **p < 0.01 and ***p < 0.001 were considered as significant.

**Results and discussion**

**Crude Extraction and Estimation of C-PC**

The purity ratio of the C-PC was obtained at different steps of purification by ammonium sulphate...
precipitation, dialysis and ion-exchange chromatography on the DEAE cellulose column. The purity ratio ($A_{620} / A_{280}$) showed a regular increase at each purification step. The initial crude extract showed a purity ratio ($A_{620} / A_{280}$) of 1.81. C-PC was eluted with a linear gradient of phosphate buffer (25 – 100 mM). Column chromatography of the 100 mM fraction gave the high purity ratio ($A_{620} / A_{280}$ = 4.12) as compared to standard ($A_{620} / A_{280}$ = 4.47) (Figure 2).

Figure 2. The purity of C-PC at different steps of purification and level of significance. The data are mean ± SD of three independent experiments, where, ns = non-significant, **p < 0.01 and ***p < 0.001.

The impurity of C-PC protein decreased in terms of recovery at the initial extract crude step to the final chromatography step was 99.6 ± 3.7 % and 57.2 ± 3.5 % respectively.

Characterization of Purified C-PC

UV-visible Spectroscopy

The absorption characteristics spectrum of standard, crude, ammonium precipitated dialyzed and column purified C-PC obtained from Plectonema sp. was recorded from 250 to 750 nm as represented in Figure 4. A sharp peak at 620 nm suggested high C-PC present in test samples at various stages of purification. The steepest peak was seen in the column purified phase of C-PC. No absorption peak at 650 nm or 540 nm was detected, confirming the absence of C-phycoerythrin and C-allophycocyanin in the sample.

Figure 4. UV-vis spectroscopy graph of standard, crude, ammonium precipitated, dialysed and column purified C-PC. The data are mean ± SD of three independent experiments.

Fluorescence Spectroscopy

A sharp peak in standard C-PC near 350 nm was observed which is specific for tryptophan residues, suggesting that the protein is in its folded form. All samples of C-PC exhibit a peak near 350 nm while the most prominent peak was observed in column purified extracts of C-PC indicating the pure form of C-PC (Figure 5).
**Fourier Transform Infrared Spectroscopy (FT-IR)**

Amides I is the major band in the protein IR spectrum in the region between ~ 1600 – 1700 cm\(^{-1}\). Fourier transform infrared spectra (FT-IR spectra) of standard, crude, ammonium precipitated, dialysed and column purified extract of C-PC of *Plectonema* sp. exhibited the protein-specific amide I band at 1637, 1636, 1637, 1636, 1638 cm\(^{-1}\) (C=O stretching) (Figure 6). The position and shape of the amide I bands are used for the analysis of the secondary structure of the protein. The sharp band of amide I for C-PC represents the α-helix as the main component of its secondary structure. Simultaneously, the IR spectra of C-PC ensure its purity by the absence of inorganic sulfates and phosphates (representing intensive bands at 1043 and 1015 cm\(^{-1}\)).

The CD spectra of all the C-PC samples showed two minima at 208 and 222 nm, a feature of α-helix structure proteins, which are in accordance with the CD spectra of standard C-PC (Figure 7). Similarities in the relative proportions of secondary structure in all C-PC samples suggest the purity of the isolated C-PC. To better assess the structural transition, the CD spectra of C-PC were de-convoluted through the K2D3 software. The α-helix content in standard C-PC was 80.59 % while ammonium precipitated, dialysed and column purified extract of C-PC exhibited 84.59 % α-helix content.

**Sodium Dodecyl Sulphate-Polyacrylamide Gel Electrophoresis (SDS-PAGE)**

The purity of purified C-PC was analysed by 12 % SDS-PAGE (lane-1 crude extraction, lane-2 high-intensity standard, lane-3 ammonium precipitated, lane-4 dialysed and lane-5 ion-exchanged column chromatography samples respectively). The SDS-PAGE analysis of all purified C-PC showed two main bands corresponds to α and β subunits of C-PC, further validating its homogeneity and purity by comparing with standard C-PC (lane-2). Additionally, the calculated molecular mass of α and β subunit was found to be 17 kDa and 19 kDa (Figure 8).
Figure 8. 12% SDS-PAGE analysis of purified C-PC from Plectonema sp., M- Protein molecular marker; 1- Crude extract; 2- Standard; 3- Ammonium sulphate precipitated; 4- Dialysed and 5- Ion-exchange column chromatography.

High-performance liquid chromatography (HPLC)

HPLC analysis of standard C-PC and purified column C-PC was performed at 620 nm. Purified column C-PC of Plectonema sp., showed peaks with the retention time of 1.465 min, 1.234 min, 1.097 min and 0.905 min, equivalent to the standard peak at 1.448 min, 1.233 min and 0.925 min. [Figure 9 (A) and (B)] confirming the identity of the sample as C-PC.

Figure 9. HPLC chromatogram of (A) standard C-PC and (B) purified column C-PC from Plectonema sp.

Molecular Docking Analysis of C-PC with Free Radicals

The docked complex of C-PC ligand molecule with targeted molecule hydroxyl radical showed that the complex was formed by the hydrogen bond interaction of targeted molecule hydroxyl radical with ARG30 and SER37 residues of C-PC. The hydrophobic interaction was observed between targeted molecule hydroxyl radical and GLY102 and ARG30 residues of C-PC [Figure 10 (A)], the docked complex had binding energy (ΔG) of −2.5 kcal/mol.

The docked complex of C-PC ligand with targeted molecule superoxide radical was formed by a hydrogen bond between superoxide radicals and LYS1 residue of C-PC, hydrophobic interaction was observed between targeted superoxide radical and GLY105 residue of C-PC [Figure 10 (B)], the binding energy was −2.19 kcal/mol. In the docked complex of C-PC and hydrogen peroxide, hydrogen bonds and hydrophobic interaction were found between hydrogen peroxide THR104 residue of C-PC [Figure 10 (C)], the binding energy was −2.24 kcal/mol.

The docked complex of C-PC with nitric oxide radical was formed by the hydrogen bond and hydrophobic interaction between nitric oxide radical and TYR129 residue of C-PC [Figure 10 (D)], the binding energy was found to be −3.44 kcal/mol (Table 1). Complexes with the highest binding were selected for analyzing the stability of protein-target interaction.

Table 1. Binding Energy, Intermolecular Energy, Inhibitory Constant, Binding residues and Hydrogen bonds length comparison table of C-PC with free radicals molecular docking; target (A), Binding Energy (kcal/mol) (B), Intermolecular Energy (kcal/mol) (C), Inhibitory Constant (mM) (D), Binding residues (E), Hydrogen bonds length (Å) (F)

<table>
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<tr>
<th></th>
<th>A</th>
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<td>2.99</td>
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Molecular Docking Analysis of C-PC with Enzymes

The ZDOCK generated the top 10 docked predictions. All of the complexes were evaluated using the ZDOCK score. ZDOCK Score (force field) = -(ligand/receptor interaction energy + ligand internal energy) (Xiong et al., 2013). Complexes with the highest ZDOCK scores were selected for analyzing the stability of protein-protein interaction. The ZDOCK score for the interaction of C-PC with the residue of α-amylase and α-glucosidase interaction was found to be 1588.66 and 1823.19 respectively. The ZDOCK score of the C-PC with GSK-3β and GP was found to be 1525.48 and 1794.53, respectively (Figure 11).

**Figure 10.** Molecular docking interaction of C-PC with free radical. C-PC (Ligand) is represented in sticks and free radical (Target) is represented in stick and ball form, hydrogen bonds are represented in green colour, hydrophobic interactions are represented in red colour, (A) Molecular docked model of C-PC with hydroxyl radical, (B) Molecular docked model of C-PC with superoxide radical, (C) Molecular docked model of C-PC with hydrogen peroxide and (D) Molecular docked model of C-PC with nitric oxide radical.
Cyanobacterial phycobiliproteins have gained importance due to their potential applications in different commercially important sectors. One such phycobiliprotein is C-PC which Beholds numerous applications and is isolated, purified and characterized from Plectonema sp. (56). Different purification procedures affect C-PC concentration because the efficiency of one procedure on particular Plectonema sp. may not be efficient for the others. To overcome this drawback our study was designed to elucidate a broad applicable C-PC isolation and purification method for Plectonema sp. The four major steps employed for the isolation and purification of C-PC from Plectonema sp. were (i) crude extract preparation, (ii) ammonium sulfate precipitation, (iii) dialysis, and (iv) anion exchange chromatography. The competence of extraction methods was determined by calculating the concentration and purity ratio of isolated C-PC. The purity of C-PC plays a considerable role in commercial applications (56) and is generally estimated using the absorbance ratio of A620/A280 (57). The purity of C-PC after ammonium sulphate precipitation was found to be 1.81 folds enhanced in comparison to the crude extract having a concentration of 347.67 µg/ml with an initial recovery of 99.6%. Moreover, the dialysis of ammonium sulphate precipitated fraction further augmented it purity content by 1.5 folds having a concentration of 317 µg/ml and a decreased recovery of 86.5%. This step is predominantly useful in salting-out unwanted proteins simultaneously concentrating C-PC. Furthermore, C-PC was finally purified by DEAE Cellulose column, which further increased the purity level by 1.25 folds as compared to dialyzed C-PC having a concentration of 245 µg/ml with further decreased recovery of 57.2%. The highest purity ratio of 4.12 at the final purification step of column chromatography was also found to be nearly equivalent to the purity ratio of a standard that is 4.47.

The absorption spectra of the column purified C-PC exhibited a prominent peak at 620 nm which was in accordance with the peak of standard C-PC. Additionally, the C-PC of Plectonema sp. was further characterized for its intrinsic fluorescence properties using fluorescence spectroscopy. The sharp emission peak at 350 nm of column purified C-PC elucidated the presence of C-PC which has been earlier specified for the native form of this protein thereby validating its purity (58). The absence of peaks at 1043 and 1015 cm\(^{-1}\) in FT-IR spectra confirms the absence of inorganic impurities, while the sharp amide I band in standard along with column purified C-PC indicates α-helix as the chief constituent of its secondary structure (59). CD spectra of standard and column purified C-PC exhibit two negative minima in the ultraviolet region at 208 nm and 222 nm, which is a characteristic of the α-helical structure of the protein. The two negative peaks both contributed to an n→π* transition for the peptide bond with α-helicity (60) demonstrated the α-helical structure of C-PC, devoid of β-sheets. In the array to characterize C-PC, HPLC was performed using C18 reverse-phase column, exhibiting peaks with retention times of 1.465 min, 1.234 min, 1.097 min and 0.909 min. which was, equivalent to the peaks of standard C-PC of 1.448 min, 1.233 min and 0.925 min. Moreover, SDS-PAGE also confirmed the two highly intensified bands from column purified C-PC in comparison with the standard and revealed its α and β subunits with a molecular weight of 17 and 19 kDa respectively, thereby confirming the purity of C-PC.

Oxidative stress through the production of free radicals has been proposed as the root cause of the
development of type 2 DM (61). An antioxidant is known to exert a protective effect against oxidative damage and is linked with a reduced risk of chronic diseases (62). Consequently, molecular docking studies were performed to analyze the association between ligand C-PC and its antioxidant property. C-PC was docked with free radicals: Hydroxyl radicals, superoxide radicals, hydrogen peroxide and nitric oxide radicals using AutoDock 4.2. Molecular docking analysis between C-PC and free radicals demonstrated that C-PC binds with all four free radicals. Our results were in accordance with previous findings of in vitro antioxidant activity of C-PC (63). C-PC was found to possess greater interaction (binding energy = −3.44 kcal/mol) with nitric oxide radical as compared to hydroxyl radical (binding energy = −2.5 kcal/mol), superoxide radical (binding energy = −2.19 kcal/mol), hydrogen peroxide (binding energy = −2.24 kcal/mol). These results demonstrate that C-PC is an efficient scavenger of free radicals; therefore, the therapeutic use of C-PC as a natural antioxidant appears promising.

Highly involved mechanisms for antidiabetic therapy are inhibition of intestinal alpha-amylase and alpha-glucosidase, altering the activity of GSK3 and glycogen phosphorylase (64). Alpha-amylase and alpha-glucosidase are the essential enzymes that catalyses the primary step in hydrolysis of starch to maltose and finally glucose. Inhibition of these enzymes in intestine results in delayed carbohydrates digestion (65). GSK3 negatively regulates numerous features of insulin signaling (66) while GP is a significant therapeutic target for type 2 diabetes with a direct effect on blood sugar levels via the glycogenolysis pathway (67). Similar in vitro alpha-amylase and alpha-glucosidase inhibition activity were recorded by (68), who investigated the alpha-amylase and alpha-glucosidase inhibition activity was recorded in concentration-dependent manner with increasing concentration of C-PC. Similarly, (69) investigated the in vitro antidiabetic activity of purified pigments from three cyanobacterial species; Microcoleus, Lyngbya, and Synechocystis and found that purified C-PC and C-phycocerythrin from all species inhibited the enzyme the most (96.62 %), while crude solvent extracted compounds had the least activity. Hence, to elucidate the antidiabetic potential of the C-PC molecular docking technique was performed. Z-dock technique of protein-protein interactions was executed between C-PC and enzymes (70). All enzymes bind significantly with C-PC, the docking conformations of C-PC with alpha-amylase, alpha-glucosidase, GSK-3β and GP deciphering the inhibitory potential of C-PC on enzymes. Our findings were in accordance with an in silico study conducted by (71-73), which demonstrated that C-PC inhibits alpha-amylase and alpha-glucosidase by binding to the active site and disrupting substrate-enzyme interactions. The highest ZDOCK score reveals the highest protein-protein interaction. The maximum ZDOCK score was observed by the interaction of C-PC and alpha-glucosidase (ZDOCK score: 1823.19), while the ZDOCK score of interaction C-PC with alpha-amylase, GSK3- β and GP was found to be 1588.66, 1525.48 and 1794.53 respectively.

Conclusion

Apart from establishing C-PC purification through sequential usage of ammonium sulphate and ion-exchange chromatography, the present study provides an insight for augmenting the relative stability of purified C-PC using lyophilization. Characterization of purified C-PC by various techniques as demonstrated in this study would certainly aid in studies dedicated to the improvement in the stability and the purity of C-PC. The molecular docking outcomes recommend substantial antioxidant and antidiabetic potential of C-PC by interacting with free radicals and enzymes. Furthermore, the in silico results not only confirm the findings of the other in vitro experiments but also provide light on the overall molecular interaction between C-PC and carbohydrate-metabolism enzymes. To further comprehend the therapeutic potential of C-PC and establish the mechanism of action of C-PC in reducing hyperglycemia in diabetic patients, more scientific validation is required. Indeed, in silico screening will play a key role in identifying active ligands for the novel protein targets that are on the horizon.

Conflict of interest

The authors declare that they have no conflicts of interest with the contents of this article.

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