Biochemical characterization of the YBPCI miniprotein, the first carboxypeptidase inhibitor isolated from Yellow Bell Pepper (*Capsicum annuum* L.). A novel contribution to the knowledge of miniproteins stability

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https://doi.org/10.1016/j.ypep.2017.12.003

Received 9 October 2017; Received in revised form 5 December 2017; Accepted 5 December 2017

Available online 06 December 2017

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**ARTICLE INFO**

**Keywords:**
Carboxypeptidase inhibitor
Plant inhibitor
Capsicum annuum
Protease
Stable miniproteins
Gastrointestinal digestion

**ABSTRACT**

The cystine-knot metallocarboxypeptidase inhibitors (MCPIs) are peptides that contribute to control proteolytic activity, involved in storage, growth and maintenance of plants. Lately studies reported several MCPIs with potential use in biomedical applications; as anti-cancer, anti-thrombotic, anti-malaric and anti-angiogenic agents. We report the isolation, purification, chemical stability and biochemical characterization of a novel carboxypeptidase A inhibitor (YBPCI) isolated from *Capsicum annuum* L. var. Yellow Bell Pepper, the first cystine-knot miniprotein (CKM) of the species. We demonstrate the stability of YBPCI (IC₅₀ = 0.90 μg/ml) to high temperatures, high salt concentration and extreme pH values. MALDI-TOF/MS analysis detected a molecular weight of 4057 Da, and peptide mass fingerprint resulted in no matches with other protease inhibitors. In vitro gastrointestinal digestion subjecting YBPCI to pH 2 incubation and proteolytic attack resulted in complete inhibitory activity. To summarize, there are no reports to date of carboxypeptidase inhibitors in *C. annuum* species, giving our report much more relevance.

**Abbreviations:**
CKM, cystine-knot miniproteins; MCPIs, metallocarboxypeptidase inhibitors; YBPCI, Yellow Bell Pepper Carboxypeptidase Inhibitor; MALDI-TOF/MS, matrix-assisted laser desorption and ionisation time-of-flight/mass spectrometry; PMF, peptide mass fingerprint; CPs, carboxypeptidases; MCPs, metallocarboxypeptidases; PIs, protease inhibitors; PCI, Plant inhibitor; CPAU, Carboxypeptidase A inhibitor; ACI, Ascaris suum Carboxypeptidase Inhibitor; LCI, Helix aspersa Maxima Carboxypeptidase Inhibitor; TCI, Ticks Carboxypeptidase Inhibitor; HITCI, Helix pomatia Carboxypeptidase Inhibitor; NvCI, Nematodirus vectensis Carboxypeptidase Inhibitor; ECI, Endogenous Carboxypeptidase Inhibitor; SmCI, Schistosoma mansoni Carboxypeptidase Inhibitor; APA, Ascaris suum Protease Inhibitor; PII, Peptide Inhibitor from Jalapeño Pepper; CPA, Carboxypeptidase A; SDSL, sodium dodecyl sulphate; JME, β-mercaptoethanol; TEMED, N,N,N′,N′-tetramethyl ethylene diamine; BSA, bovine serum albumin; CPAU, Carboxypeptidase A unit

**1. Introduction**

Carboxypeptidases (CPs) are enzymes that cleave proteins from the C-terminus, usually one residue at a time, playing an essential role in degradation, processing, and modulation of proteins and peptides [1]. CPs were initially considered as degrading enzymes associated with protein catabolism, but over the years evidence demonstrates that some CPs play key roles in controlling a vast variety of biological processes [2]. Among CPs, metallo-carboxypeptidases (MCPS) are exopeptidases that hydrolyze the C-terminal amide bond by a zinc-dependent mechanism [3]. In humans, the dysregulated activity of some MCPS has been associated with several diseases such as blood coagulation/fibrinolysis, inflammation, carcinogenesis, epilepsy and febrile seizures, among others [1,2,4,5]. According to these studies, the use of protease inhibitors (PIs) in order to regulate MCPS action has emerged as a potential tool for the development of therapeutic strategies for a large number of diseases; as anticarcinogenic, anti-angiogenic and anti-thrombotic, among others [6–9]. Furthermore, although protease inhibitors have been considered only as antinutritional factors, they have regained interest in recent years because of their potential for drug development and positive dietary effects.

The application of natural PIs offers great advantages over those of synthetic origin due to their lower toxicity and hydrophobicity [10]; however, naturally occurring metallo-carboxypeptidase inhibitors...
2.1. Materials

Carboxypeptidase A (CPA) from bovine pancreas, Pepsin from porcine gastric mucosa, Pancreatin from porcine pancreas, sodium chorate (SDS), β-mercaptoethanol (βME), Coomassie Blue G-250, N,N,N′,N′-tetramethyl ethylene diamine (TEMED) and bovine serum albumin were supplied by Sigma-Aldrich (U.S.A.). N-(4-methoxophenylazoformyl)-phenylalanine-OH potassium salt was obtained from Bachem (Switzerland).

2.2. Crude extract preparation

C. annuum seeds were collected from nearby areas around La Plata (Argentina) and washed thoroughly in distilled water. The dry seeds (30 g) were ground using a blender with addition of 3 vol. of 0.01 M phosphate buffer, 0.1 M NaCl, pH 7.4. The mixture was incubated 3 h at 4 °C and filtered with gauze. The suspension was centrifuged at 7000 g for 30 min at 4 °C, and the supernatant was collected and frozen at -20 °C until analysis.

2.3. Carboxypeptidase A inhibition activity

The CPA activity assay is based on the hydrolysis of the substrate N-(4-methoxyphenylazoformyl)-Phe-OH [30]. For the inhibition experiment a fixed concentration of CPA solution (50 nM) was pre-incubated with different concentrations of inhibitor in 0.1 M Tris-HCl buffer (pH 7.4) containing 0.2 M NaCl. After 15 min at 37 °C, the chromogenic substrate was added to the reaction mix at a concentration of 1 mM and, immediately, the absorbance at 405 nm was measured every minute for a minimum of 15 min. The protease inhibitor activity was expressed in terms of percent inhibition of CPA activity throughout the course of study, by using the comparison of absorbance changes of the test and the control, according to the equation:

\[
\text{CPA inhibition (％) = } 1 - \frac{\Delta \text{Abs (405 nm)/Δ(min)}_{\text{inhibitor}}}{\Delta \text{Abs (405 nm)/Δ(min)}_{\text{control}}} \times 100
\]

Inhibitory activity (IA) was defined according to Tellechea et al. (2016) [31]. Measurements were carried out in triplicate.

2.4. Protein estimation

Protein concentration was measured by the Bradford's method using bovine serum albumin (BSA) as standard (0-1 mg/ml). 20 μl of sample and standard were mixed with 200 μl of reagent for 10 min at room temperature. Afterwards, protein concentration was measured by absorbance at 595 nm (Tecan Infinite M200 PRO).

2.5. Purification

2.5.1. Partial purification by heat treatment

1 ml of crude extract (400 μg/ml) was subjected to 60 °C, 70 °C, 80 °C, 90 °C or 100 °C for 5 min, therally denatured proteins were removed by centrifugation (10000 g, 40 min, 4 °C). 100 μl of each supernatant -including a crude extract control- was employed as sample in the CPA inhibitory activity assay and residual activity was determined as described previously.

2.5.2. Affinity chromatography purification

A sample of heat treated extract containing 200 μg/ml of protein was loaded to a carboxypeptidase A-glyoxyl-agarose column (1.5 × 10 cm) [32] previously equilibrated with 0.1 M Tris-HCl buffer (pH 8.0) containing 0.2 M NaCl. In the first step unbound proteins were eluted with equilibration buffer, and then affinity proteins were eluted with HCl pH 2.0. The eluent was adjusted to pH 7.0 and the fractions exhibiting inhibitory activity were pooled.

2.6. Characterization of protease inhibitor

2.6.1. SDS-tricine-PAGE

SDS-Tricine-PAGE was used to determine the purification advance of the protease inhibitor during purification. Samples were mixed with sample buffer (Tris 0.13 M, SDS 2%, β-mercaptoethanol 5% v/v, glycerol 8% v/v, bromophenol blue 0.002% p/v, pH 6.8) and incubated 5 min at 100 °C, then subjected to denaturing electrophoresis at a constant current of 15 mA per gel using a Mini – Protean III dual slab
cell (Bio-Rad, Hercules, CA 94547, USA), using 4% stacking gel and 12% separating gel. After electrophoresis the gels were stained with 0.2% Coomassie Brilliant Blue G-250 and destained with distilled water.

2.6.2. Molecular mass determination by MALDI-TOF/MS

Matrix-assisted laser desorption ionisation-time-of-flight mass spectrometry (MALDI-TOF/MS) was used for the determination of the inhibitor molecular mass, as well as its purity degree. MALDI-TOF mass spectra was acquired on a BRUKER Ultraflex spectrometer equipped with a pulsed nitrogen laser (337 nm), in linear positive ion mode, using a 19 kV acceleration voltage. Samples were prepared by mixing equal volumes of a saturated solution of matrix (3,5-dimethoxy-4-hydroxycinnamic acid–sinapinic acid) in 0.1% TFA in water/ACN 2:1, and 1 M protein solution. From this mixture, 1 μl was spotted on a sample slide (MP 384 Polished Steel, Bruker) and allowed to evaporate to dryness.

2.6.3. Protein identification by PMF

In-gel protein digestion was carried out using the In-Gel DigestZP Kit (MILLIPORE, Billerica, MA, USA). Pure protein band was excised with a scalpel, diced into 1 mm² pieces and introduced into the wells, treated with 25 mM NH₄HCO₃/5% acetonitrile (ACN) and then with 25 mM NH₄HCO₃/50% ACN (washing/destaining solutions) and finally dehydrated with pure ACN. Proteins were then reduced with 10 mM DTT in buffer 25 mM NH₄HCO₃/5% ACN for 1 h at room temperature. Cys sulphhydrils were alkylated with 50 mM iodoacetamide in 25 mM NH₄HCO₃/5% ACN for 1 h at 37 °C. Washing/destaining solutions were used for additional washing and finally, the gel pieces were dehydrated with pure ACN and then dried. Trypsin (0.02 μg) in 25 mM NH₄HCO₃/5% ACN was added to rehydrate the gel pieces in each well and the digestion was carried out for 3 h at 37 °C. The tryptic peptides obtained were extracted from the gel with 0.2% TFA and captured on the C18 resin applying vacuum. Finally, the tryptic peptides were eluted with 50% ACN/0.1% TFA.

MALDI-TOF/MS was used for protein identification by peptide mass fingerprint (PMF). Analysis was performed using an UltraFlex MALDI-TOF/MS mass spectrometer (Bruker Daltonics, Bremen, Germany). The sample was spotted on an AnchorChip target 160 and mixed with freshly prepared matrix solution (10 mg/ml of a-cyano-4-hydroxycinnamic acid in aqueous solution containing 30% ACN and 0.1% TFA). External calibration was performed using peptide calibrants. Peptide masses were acquired with a range of ca. 800–4000 m/z. PMF spectra of samples analyzed were acquired and compared. Comprehensive peak assignments were accomplished using the BioTools software package (Bruker Daltonics).

2.6.4. Effect of pH on protease inhibitor stability

The stability of protease inhibitor over extreme pH values was determined by evaluating the inhibitor activity at pH 7 after incubating the heat-treated extract in different pH (2 and 12) for 1 h, at 37 °C. After incubation, the reaction mixture was neutralized and assayed for protease inhibitor activity.

2.6.5. Effect of NaCl concentration on protease inhibitor stability

Effect of NaCl concentration on protease inhibitor stability was evaluated using sodium chloride at 0.5 M and 2 M concentrations in the reaction mixture. A ten times concentrated heat-treated extract sample was used to carry out 1 h incubation at 37 °C with sodium chloride, and then 10 μl of sample were assayed for protease inhibitory activity.

2.6.6. In vitro simulation of gastrointestinal digestion

In order to evaluate the effect of gastrointestinal digestion on protease inhibitor stability, an aliquot of 1 ml of 100 °C treated extract (200 μg/ml) was subjected to gastric simulation. First the sample was adjusted to pH 2 with 1 M HCl, then pepsin (3400 IU/mg) was added on a rate of 182 IU per mg of peptide and incubated 2 hs at 37 °C and 200 rpm. After incubation, sample was adjusted to pH 5.3 with 0.5 M NaHCO₃ and then to pH 7.5 with 2 M NaOH. Pancreatin from porcine pancreas was added on a rate of 35 IU per mg of peptide and incubated 3 hs at 37 °C and 200 rpm. Finally, the reaction was stopped by heating at 100 °C for 10 min and assayed for protease inhibitory activity.

3. Results and discussion

3.1. Isolation of carboxypeptidase A inhibitor from Yellow Bell Pepper seeds

MCPIs have been identified in Solanaceae members such as potato and tomato, nevertheless there are no reports to date of MCPIs from Capsicum annuum members. In order to isolate a novel CPA inhibitor, crude extract preparation was achieved by pulverizing Yellow Bell Pepper (Capsicum annuum L.) seeds with addition of phosphate buffer, posterior incubation and centrifugation. Protein concentration determined by Bradford’s method using a BSA (0-1 mg/ml) calibration curve resulted in 400 μg/ml. Inhibitory activity was detected through kinetic measurements employing a simple microplate method [31]. If we define a CPA unit (CPAU) as the amount of enzyme required to produce 0.01 absorbance units diminution at 340 nm per minute, employing N-(4-methoxyphenilazoformyl)-Phe-OH as substrate at 37 °C and pH 7.4, then we observe 0.76 CPAU for the control curve without inhibitor, and 0.42 CPAU with crude extract addition. Thus, the crude extract presents a 45% CPA inhibition.

3.2. Purification of carboxypeptidase A inhibitor from Yellow Bell Pepper

3.2.1. Partial purification by heat treatment

3.2.1.1. Carboxypeptidase A inhibition activity. Previous studies suggest PIs are heat-stable molecules due to disulfide bonds that stabilize their structure. Based on these results, we proposed a simple and rapid purification method that consisted in evaluation of the inhibitory activity after treating the crude extract to different temperatures. We observe that treatment at 80 °C during 5 min yields a product with higher activity than treatment at 60 °C. Surprisingly, treatment at 100 °C gives a product with inhibitory activity as high as treatment at 80 °C (Fig. 1), which indicates that the carboxypeptidase inhibitor from Yellow Bell Pepper is a very stable molecule. Although PIs are heat-stable molecules, we found no reports on MCPIs with resistance to 100 °C, such thermal treatment usually produce a reduction on inhibitory activity. In this case we are in presence of an unpublished result; the 100 °C treatment results in an inhibitory activity equal to the observed at 80 °C, opening new insights in PIs stability.

As observed in Fig. 1, the CPA inhibitory activity is not only
maintained with temperature increase; it also increment gradually to the different thermal treatments. It is expected that, due to the great stability of these inhibitors, the same inhibitory activity observed in the crude extract will be maintained after thermal treatments. However, in this case the inhibitory activity presents a considerable increase. This would indicate the presence in the crude extract of CPA activating biomolecules that are affected by the thermal treatments, the presence of thermolabile biomolecules that reversibly inactivate the CPA inhibitor (YBPCI), or the possibility that in crude extract the inhibitor is in an inactivated isomeric that is reverted by thermal treatment [33–36].

3.2.2. Protein pattern by SDS-PAGE. Proteins from each purification step were separated under reducing conditions and stained with Coomassie Blue after electrophoresis (Fig. 2). Based on the protein patterns we observe that the crude extract contains a variety of protein with different molecular weight. Heat treatment at 60 °C produces no significant changes in protein levels. Proteins with molecular weight around 20 and 30 kDa and proteins with molecular weight higher than 60 kDa were largely removed by incubation at 80 °C. After heat treatment at 100 °C, total removal of proteins with molecular weight over 6 kDa is achieved [37], suggesting that those proteins that disappeared with the heating process are presumed to be heat labile proteins. These results, in addition to the elevated inhibitory activity of the treatment at 100 °C agree with previous studies that indicate PIs are heat-stable molecules. In this particular case, 100 °C treatment is proposed as a first purification step because it removes high protein proportion that remained intact in 80 °C treatment (as observed in Fig. 2) preserving carboxypeptidase inhibitory activity.

### 3.3. Characterization of carboxypeptidase a inhibitor from Yellow Bell Pepper (YBPCI)

#### 3.3.1. Characterization of partially purified YBPCI

3.3.1.1. Dose-response curve of partially purified YBPCI. A dose-response curve was performed to determine the potency of inhibition of carboxypeptidase A, and IC50 values were calculated according to Tellechea et al. (2016) [31] (Fig. 3A). The heat treatment at 100 °C of the extract from Yellow Bell Pepper produces an IC50 value of 6.7 μg/ml in accordance to remotion of heat labile proteins, which results in a higher specific inhibitory activity.

3.3.1.2. Effect of pH on stability of partially purified YBPCI. At extreme pH values, strong intramolecular electrostatic repulsion caused by high net charge results in swelling and unfolding of the protein molecules [38]. Intramolecular disulphide bridges are presumably responsible for the functional stability of some PIs in the presence of various physical and chemical denaturants [39], such as pH, temperature, and reducing agents. The pH stability of 100 °C treated extract was investigated by incubation at extreme pH values (Fig. 4A.). The partially purified YBPCI shows high stability at extreme pH values, being slightly more stable at pH 2. Similar results have been reported for *Solanum* trypsin inhibitors [27], but there is no published data referred to pH stability of CPA inhibitors therefore we present the first study on pH stability of MCPIs.

3.3.1.3. Effect of salt on stability of protease inhibitor. The effect of NaCl on the inhibitory activity of 100 °C treated extract is depicted in Fig. 4B. No marked changes in the inhibitory activity were found when NaCl was added at 0.5 M and 2 M concentrations. From the result, the partially purified YBPCI shows high salt stability, representing a new proof of the molecule’s great stability.

3.3.1.4. Effect of gastrointestinal digestion on stability of protease inhibitor. It has been recognized that dietary proteins and peptides are susceptible to hydrolysis during the different stages of gastrointestinal digestion, namely ingestion, digestion and absorption [40]. Once ingested, these proteins and peptides are subjected to hydrolysis by different enzymes present in the gastrointestinal tract such as pepsin, trypsin, chymotrypsin and peptidases at the surface of epithelial cells to release peptides of various lengths [41]. Some of these peptides may exert a direct function at the gastrointestinal tract. However, other peptides can be absorbed to reach target organs and tissues through systemic circulation [42].

In order to examine the effect of gastrointestinal proteases on YBPCI

#### Table 1

| Purification step | Total protein amount (mg) | Protein concentration (μg/ml) | IC50 (μg/ml) | Total Inhibitory activity, IA (mU) | Specific inhibitory activity, IA (mU/mg) | Purity (fold) | Yield (%) |
|------------------|--------------------------|-------------------------------|-------------|------------------------------------|----------------------------------------|--------------|-----------|
| Extract          | 40                       | 400                           | 46.4        | 756                                | 19                                    | 1            | 100       |
| 100 °C Treatment | 5                        | 200                           | 6.7         | 647                                | 129                                   | 6.8          | 85        |
| Affinity Cromatography | 1                   | 16                            | 0.9         | 218                                | 218                                   | 11.5         | 28        |
stability, digestion with pepsin and pancreatin was carried out according to Hernández-Ledesma et al. (2007) [43] (Fig. 4C.). The 100 °C treatment shows fully inhibitory activity after gastric simulation, suggesting that YBPCI reaches the colon in intact form. Experiments with cystine-knot miniproteins [44] and the Bowman-Birk Inhibitor [45] showed that these molecules enters the circulation through intestinal epithelial cells or the paracellular mechanism, opening new perspectives on potential clinical applications for oral administration of IPs.

In our work, we demonstrate the stability of partially purified YBPCI towards gastrointestinal digestion, through kinetic inhibition assays and dose-response curves on digested YBPCI. Even though in vitro gastrointestinal digestion has been assayed on other cystine-knot miniprotein [44], our work represents the first evidence of intact inhibitory activity of a cystine-knot miniprotein after gastric simulation, contributing to the understanding of IP's stability.

3.3.2. Characterization of purified YBPCI

3.3.2.1. Molecular mass determination. The molecular mass of the protease inhibitor purified by affinity chromatography was estimated based on SDS-PAGE analysis (about 4 kDa) (data not shown). Matrix-assisted laser desorption ionisation-time-of-flight mass spectrometry (MALDI-TOF/MS) was used for the determination of the inhibitor molecular mass (Fig. 5.). Thus, the molecular weight is confirmed and precisely determined (4057 Da) of the order of the masses of

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**Fig. 3.** Dose-response inhibitory curves. (A) Dose-response relationship for partially purified YBPCI. (B) Dose-response relationship for purified YBPCI. Data points represent the mean value of three determinations.

**Fig. 4.** Stability assays on partially purified YBPCI. (A) Effect of pH on stability of partially purified YBPCI. (B) Effect of NaCl addition on stability of partially purified YBPCI. (C) Effect of gastric simulation on stability of partially purified YBPCI. Error bar indicates standard deviation from triplicate determination.
carboxypeptidase inhibitors already studied (imaPCI: 4234 Da, [12]; PCI: 4295 Da, [11]; NvCI: 5945 Da, [17]; LCI: 7326 Da, [14]; ACI: 7724 Da, [13]; TCI: 7935 Da, [15]; H1TCI: 8400 Da [16]).

3.3.2.2. Protein identification. Trypsin digestion of the inhibitor was analyzed by MALDI-TOF MS. MASCOT search tool (URL http://www.matrixscience.com) was used for identification of tryptic maps. The range of molecular weighs employed for the analysis of peptide mass fingerprint (PMF) of tryptic digested peptides from YBPCI, was between 1000 Da and 4000 Da. No matches are found with other plant inhibitor which confirms the unique nature of the YBPCI and represents the first MCPI isolated from Capsicum annuum. Protein identification and differentiation by PMF has been adopted in our group as an excellent tool to differentiate, in fast and unequivocal way, proteases and protease inhibitors with very similar physicochemical and functional properties [32].

3.3.2.3. Dose-response curve of purified YBPCI. A dose-response curve was performed to determine the potency of inhibition of carboxypeptidase A, and IC50 value was calculated according to Tellechea et al. (2016) [31] (Fig. 3B.). The purified YBPCI produces an IC50 value of 0.9 μg/ml, which results in a higher specific inhibitory activity than partially purified YBPCI.

4. Conclusion

The discovery of natural protease inhibitors is becoming more important to food and medicinal scientists. Herein, we report the first metalloprotease inhibitor isolated from Capsicum annuum, a world-class vegetable, and demonstrate the stability of the inhibitor to high temperatures, salt concentration and extreme pH values. Although this characteristic has been studied in other T-knot peptides, mostly on cyclotides of other mechanistic type, the results from the present research represents the first published study on stable miniproteins MCPI-type. Also, it is the first report on gastrointestinal digestion stability of these molecules, suggesting that PIs could reach the colon in intact form, and encouraging the study of possible pharmaceutical applications as natural anti-parasitic and/or anti-bacterial formulations. Our work contributes to the understanding of IP's stability, representing the first demonstration of a cystine-knot miniprotein that conserves intact inhibitory activity after in-vitro gastrointestinal digestion.

Based on previous research that demonstrate some PIs reach systemic circulation through intestinal epithelial cells or the paracellular mechanism, the development of organic formulations or the controlled ingest of MCPIs-rich vegetables could represent a natural strategy for the treatment of multiple pathologies; angiogenesis, thrombosis, cancer, malaria, parasites. Also it could represent a natural way of health control in general population. To ascertain whether dietary consumption of MCPIs-rich foods is compatible with biological activity of these miniproteins in vivo as we found in vitro requires further and specific experimental evaluation.

Acknowledgments

This study forms part of the DSc degree thesis of J. Cotabarren, which was carried out at the UNLP. J. Cotabarren and M. Tellechea are doctoral fellows from UNLP and W.D. Obregón is a member of the Researcher Career Program of the Argentine Council of Scientific and Technical Research (CONICET). The authors acknowledge support of the Ministerio de Economía y Competitividad (MINECO) of Spain.
