Proregion of *Acanthoscelides obtectus* cysteine proteinase: 
A novel peptide with enhanced selectivity toward endogenous enzymes


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**Abstract**

*Acanthoscelides obtectus* is a devastating storage insect pest capable of causing severe bean crop losses. In order to maintain their own development, insect pest larvae feed continuously, synthesizing efficient digestive enzymes. Among them, cysteine proteinases (CPs) are commonly produced as inactive precursors (procysteines), requiring a cleavage of the peptide proregion to become active. The proregion fits tightly into the active site of procysteines, efficiently preventing their activity. In this report, a CP cDNA (*cpao*) was isolated from *A. obtectus* midgut larvae. In silico studies indicated that the complete CP sequence contains a hydrophobic signal peptide, a prodomain and a conserved catalytic region. Moreover, the encoding cDNA contains 963 bp translating into a 321 residue protein, CPAo, which was expressed in *E. coli*, fused with thioredoxin. Enzymatic assays using the recombinant protein revealed that the enzyme was catalytically active, being able to cleave the synthetic substrate Z-Phe-Arg-7-AMC. Additionally, this report also focuses the *cpao* propeptide (PCPAo) subcloning and expression. The expressed propeptide efficiently inhibited CPAo, as well as digestive CP of other bean bruchids. Little or no activity was found against proteolytic enzymes of two other coleopterans: *Rhyzopertha dominica* and *Anthonomus grandis*. The data reported here indicate the possibility of endogenous propeptides as a novel strategy on bruchids control, which could be applicable to bean improvement programs.

1. **Introduction**

Plant tissues are frequently attacked by insect pests and pathogens. This predation process can lead to severe economic loss [1]. Two species of bruchids, *Acanthoscelides obtectus* and *Zabrotes subfasciatus*, are known to cause intense damage to the common bean grains *Phaseolus vulgaris* [7,21]. The digestion process of bean bruchids is essentially based on...
proteolytic enzyme activities, which include mainly serine and cysteine proteinases (CPs). The CPs are strictly involved in intra- and extra-cellular protein degradation, playing a key role in the synthesis of free essential amino acids, which are utilized for insect growth and development [1,6].

Due to the importance of cysteine proteinases to bean weevils’ digestion mechanisms, special attention has been given to this specific enzyme. Catalytic domains of papain-like CPs share a very similar three-dimensional structure and possess a well-conserved catalytic site [19]. These enzymes are involved in a wide variety of biological processes, including housekeeping tasks within the endosomal/lysosomal system, specific functions in antigen presentation prohormone processing, as well in pathogen and insect pest predation [26]. For the potential involvement with insect pests and pathogen predation, a co-evolution process leads several organisms to synthesize several classes of compounds that are capable of reducing proteolytic activity. These compounds are known as proteinase inhibitors [20] and several reports have described high inhibitory activity of different proteinase inhibitors toward digestive enzymes, indicating their potential as biotechnological tools for insect pest control [12,22].

It was also shown that several peptides can inhibit proteolytic enzymes. Among them, the propeptides of cognate digestive enzyme sequences have been described as extremely effective [5,10]. In the papain-like family, a single propeptide is synthesized to the inactive enzyme. This propeptide production inhibits cysteine proteinases, improving the proteolytic enzymes. Among them, the propeptides of cognate digestive enzymes have been described as extremely effective [11].

The inhibitory mechanism of the prodomain comprises the active binding cleft segment and a highly flexible C-terminal segment follow this domain [2]. Both helices are not directly involved in the active side cleft occlusion, but strongly influence the propeptide affinity to the catalytic domains [14]. The inhibitory mechanism of the prodomain comprises the substrate obstruction to the active site, in which the propeptide follows the groove in the opposite direction of substrate, resulting in an inappropriate positioning of the peptide bond for hydrolysis [11].

Aiming to assess the potential use of proteinase proregion in plant biotechnology as have been proposed [27,30,32], this report describes the molecular cloning and expression of a complete cDNA sequence encoding a cathepsin L-like enzyme of A. obtectus and shows the efficiency of the N-terminal proregion domain product in inhibit bruchid cysteine enzymes, indicating its possible application in the control of bean weevils.

2. Material and methods

2.1. CDNA cloning of A. obtectus cysteine-like proteinase

Total RNA was isolated from homogenized A. obtectus Say (Coleoptera: Bruchidae) midgut larvae following the RNeasy Mini Handbook (Qiagen®) techniques. For the reverse transcription of A. obtectus, cDNA was done using oligo d(T)-anchor primer and AMV-RT (Boehringer Mannheim®) according to the manufacturer’s instructions, using 4 μg of total RNA. RT-PCR was performed using two degenerate primers P1 (5′-TGGCGVTCVTGTGTTG-3′) and P2 (5′-ACCRCCRTYGCCARCC-3′), based on consensus sequences found in a number of CPs with an annealing temperature of 42°C. To obtain the complete cdna sequences, the 5′ and 3′ ends were amplified using a 5′/3′ RACE kit (Boehringer Mannheim®) according to the manufacturer’s instructions. The 5′ RACE was performed using primers P3 (5′-AAGTTTCTGTTATTT-3′), P4 (5′-AGATCAAAGTATTAT-3′) and P5 (5′-AACAGCACTGAAAGC-3′). 3′ RACE-PCR was carried out using primers P6 (5′-CCGGTGTCAGTGGGGCAACTA-3′) and P7 (5′-CCCATGAAAGGTCTCGCTATA-3′). The complete gene was amplified with primers P8 (5′-GGGGAATTCGTTAGGGTC-GTCGT) and P9 (5′-AAGATCCGGCTCTCATAAAATAGGA-3′) and cloned into the pGEM-T Easy Vector (Promega®). Several clones were sequenced in both strands with an ABI Prism 3700 DNA Analyzer (PE Biosystems®). Computer analysis of the DNA and derived amino acid sequences were done using the GCG package (Genetics Computer Group, Inc.), bioinformatics resources of the NCBI homepage (http://www.ncbi.nlm.nih.gov) and the EBI website (http://www.ebi.ac.uk/).

2.2. Expression and purification of PCPAo and CPAo

Coding region of A. obtectus cysteine proteinase (CPAo) and its proregion (PCPAo) were subcloned into the pET102D TOPO (Invitrogen®) and pQE40 (Qiagen®) vectors, respectively. E. coli competent cells were heat shock transformed using CPAo (strain BL21 DE3) and PCPAo (strain M15). The expression of both recombinant proteins were induced with 2.0 mM and 0.5 mM isopropyl-1-thiol-β-D-galactopyranoside (IPTG), for 3 h. Recombinant protein solubility was evaluated under different temperatures and salt concentrations. E. coli pellet cells were incubated for 1 h at 25°C in buffer A (8.0 M urea; 0.1 M NaH2PO4; 0.01 M Tris–HCl) at pH 8.0 and continuous shaking. The lysate was centrifuged at 10,000 × g for 20 min and the supernatant was loaded onto an affinity Ni-NTA (Qiagen®) column. Unbound proteins were removed with five sequential washes in buffer A at pH 6.3, then the recombinant protein was eluted from resin using buffer A at pH 4.5. CPAo was refolded by slow drop-wise dilution (≥20× dilution), stirring in the refolding buffer (50 mM Tris–HCl, pH 8.0; 5 mM EDTA; 10 mM reduced glutathione; 1 mM oxidized glutathione) and then
stirring overnight at 4 °C [10]. The renatured CPAo (10 ng per assay) was used for the proteolytic assays. The purified PCPAo was dialyzed against buffer 50 mM Tris–HCl at pH 8.0, followed by thrombin cleavage (40 h) for removal of the N-termini leader. Digested PCPAo was used in the proteolytic assays. The recombinant PCPAo (~50–100 μg) was analyzed and excised from 12% SDS–PAGE [15] and used for the production of an antiserum in mice (BALB/c, female) as previously described by Grossi-de-Sa et al. [9].

2.3. Proteolytic assays

Midguts from fourth-instar Coleoptera larvae of A. obtectus, Z. subfasciatus, C. maculatus, A. grandis, and R. dominica were dissected into ice-cold 0.1 M Tris–HCl, pH 8.0. Freshly dissected midguts were homogenized and centrifuged at 4,000 × g for 20 min at 4 °C, to remove gut walls and cellular debris. The supernatant (at a standard protein concentration of 12 ng) was used to measure the CP proteolytic activity and to analyze the inhibitory effects of PCPAo against CPs. Proteolytic inhibitory activities were tested against all enzymes described above using 10 μM of fluorogenic peptide Z-Phe-Arg-7-MCA (Sigma Co. ™). Assays were performed in 25 mM Tris–HCl, pH 6.0, according to Solomon et al. [28]. The reaction was stopped with 1.9 ml of 0.2 M Na2CO3. The endpoint reaction was measured after 30 min in a DyNA Quant 500 fluorescence reader (Amersham Pharmacia ™), with excitation at 365 nm and emission at 460 nm. In inhibitory assays, the recombinant CPAo was pre-incubated with the recombinant PCPAo (67 ng) or the synthetic inhibitor E-64 [trans-epoxysuccinyl]-l-leucylamidine-(4-guanidino) butane (10 μM), for 15 min at 37 °C. Antiserum was used as a second positive control in order to

Fig. 1 – Nucleotide and deduced amino acid sequence of the cpao cDNA. The propeptide sequence is indicated in bold. The three residues involved in the active site (Cys, His, Asn) are boxed. Dots indicate the residues of the interspersed ERFNIN motif and triangles indicate residues of the interspersed GNFD motif. Underlines indicate potential N-glycosylation site in the sequence. Circles indicate conserved cysteine residues that could form disulfide bridges. Sequence data have been submitted to GenBank under accession number AY345219.1.
abolish PCPao inhibitory activity. The inhibitory activities were calculated using the fluorescence reduction. The inhibitory activities were directly compared to free MCA produced by cysteine-like enzymes. One relative unit corresponds to 0.5 mM of free MCA produced for 30 \mu g ml\(^{-1}\) of proteinases utilized, after 30 min of reaction. The blank fluorescence readings (minus substrate) were subtracted. Assays were carried out in triplicate, with variability in endpoint fluorescence values not exceeding 10%.

Fig. 2 – Sequence alignment of cathepsin L-like pro-mature sequence from several species. Pre-regions were detected by SignalP but omitted in the alignment. Sequences sources are: A. obtectus (AY345219.1), C. maculatus (AAQ11970.1), A. gossypii (CAD33266.1), S. zeamais (BAA24424.1), R. prolixus (AAL34984.1), H. sapiens (NP_001903), D. virgifera virgifera (AAQ17127.1), A. grandis (AAR 02406.1), T. mollitor (AAP94048.2), and papain from C. papaya (P05994). Amino acid residues of the catalytic triad are marked by an asterisk. Arrows indicate conserved cysteine residues that could form disulfide bridges and the line divides pro- and mature regions.
3. Results and discussion

3.1. Molecular cloning of CPAo

The full-length cpao cDNA (963 bp) has an open reading frame of 321 amino acid residues length (Fig. 1). Multiple alignment analysis of the predicted protein sequence shows high homology to cathepsins L-like proteinases from several sources as visualized in Fig. 2. As other cathepsins from papain superfamily, CPAo is synthesized as prepro-proteinase (Fig. 1), and a direct comparison to other cathepsins indicated that Ile_{110} is the probable pro-sequence cleavage site showing a pro-sequence of 89 residues followed by a mature protein composed by 211 residues (Fig. 1). The molecular mass of deduced pro-protein (34.0 kDa) was determined by the Protein Machine software available at the Expasy site (http://us.expasy.org/tools/). Further, primary structure analysis strictly showed homology of CPAo to other mature cysteine-like proteinases. The structure of cloned CPAo was proposed by threading analyses (BioInfo Meta Server) revealing extreme similarities to egg white cystatin, which have their structure solved by X-ray diffraction (data not shown) [3]. This model revealed a wedge shape composed of a five-stranded antiparallel b-sheet wrapped around a central long a-helix folded into a wedge, with the partially flexible N-terminal segment and the first and a second b-hairpin loop forming a molecular edge [3]. Moreover, detailed information is focused on the catalytic triad residues (Cys_{25}, His_{159} and Asn_{175}) denoted by box in Fig. 1) that were conserved in papain-like proteinase family. In addition, the CPAo an active site contains four conserved half-cysteines probably involved in disulfide bond formation (Cys_{131}–Cys_{165} and Cys_{263}–Cys_{310}) (Fig. 1). Therefore, the propeptide showed homology to interspersed ERFNIN motif (dots in Fig. 1) present in cathepsins L, H, and S and totally absent in cathepsin B-like enzymes [13]. The other conserved motif GNFD that is limited to the papain superfamily [19] is also present in propeptides of CPAo, which are strictly involved in contact to the enzyme surface (triangles in Fig. 1). According to Czaplewski et al. [6], both motifs and the prosegment-binding loop (PBL) of enzyme core (H_{40}–D_{125} in cathepsin L) are essential for the N-termini folding into a conserved structural core and also to direct the propeptide fragment at opposite side of substrate into the binding site cleft. Experiments carried out with recombinant truncated propeptide of cathepsin L have demonstrated that two N-terminal a-helices are essential for propeptide binding, greatly contributing to stability [33,8]. In relation to homology searches using BLASTP, the deduced amino acid sequence of isolated cDNA clearly showed that CPAo belongs to the C1 peptidase papain superfamily as defined by Rawlings and Barrett [24]. Furthermore, CPAo showed high identity (50%) to putative gut cathepsin L-like CP from C. maculatus (Fig. 2). Significant identities were also found to Diabrotica virgifera virgifera (Coleoptera) cathepsin L-like (44%), human cathepsin L (43%), cathepsins L-like of Sitophilus zeamais (Coleoptera), Tenebrio molitor (Diptera), and Rhodnius prolixus (Hemiptera) (41%), cathepsins L-like of A. grandis (Coleoptera) and Aphis gossypii (Hemiptera) (40%) (Fig. 2). These data suggest a clear primary structure relation between insect cysteine proteinases. Similar studies were previously carried out with five different CPs synthesized by Paragonimus westermani adult worms, which showed that insect cysteine proteinases could be grouped together [23].

3.2. Expression and purification of recombinant proteins (PCPAo and CPAo)

In order to study the effects of the PCPAo toward its cognate enzyme (CPAo), the propeptide, as well as the mature enzyme, were expressed in the E. coli system. For the PCPAo, the nucleotide sequence encoding the CPAo proregion was amplified by PCR using specific flanking primers and cloned in frame with dihydrofolate reductase (DHFR) site
in the pQE40\textsuperscript{R} vector (Fig. 3C). The sequence encoding mature protein (CPAo) was cloned in frame at its 5’ end with thioredoxin in the pET102D\textsuperscript{R} TOPO (Fig. 3B). The fusion with DHR/thioredoxin was chosen in both cases to increase solubility, stability, and avoid degradation of recombinant proteins. SDS–PAGE analysis of total protein extracts revealed a major band of 51.0 kDa. Apparently, this is the molecular mass for CPAo plus fusion protein (Fig. 3A, lane 3). Additionally, a molecular mass of 41.0 kDa was observed for the propeptide plus fusion protein (PCPAo) (Fig. 3A, lane 6). No expressed protein band was found at uninduced cell extracts (Fig. 3A, lanes 2 and 5). Both recombinant proteins were detected in the insoluble cellular fraction (data not shown) and inclusion bodies were solubilized in 8.0 M urea and further purified using affinity chromatography by using a Ni-NTA affinity column (Fig. 3A, lanes 4 and 7). A thrombin digestion process was utilized to purify recombinant fusion protein. At first, recombinant CPAo expressed in E. coli was inactive and the proteolytic activity was recovered by slow serial dilution in buffer containing glutathione oxidized/reduced. Several proteins have been renatured by this procedure, including enzymes and also proteinaceous inhibitors [10,18].

3.3. Proteolytic assays

In vitro inhibitory fluorimetric assays showed that CPAo was totally inhibited by the PCPAo, as well by the CP-specific synthetic inhibitor E-64 (Fig. 4). Furthermore, previous incubation of PCPAo with polyclonal antibodies, raised against the propeptide, was able to abolish PCPAo inhibitory activity (Fig. 4). The prodomain specificities were also evaluated against several sources of digestive enzymes, which included A. obtectus, Z. subfasciatus, C. maculatus, A. grandis, and R. dominica (Fig. 5). PCPAo efficiently inhibited the CP proteolytic activity of bean bruchids extracts. By contrast, PCPAo produced low inhibitory activity against R. dominica CPs (32%) and no activity was identified toward A. grandis CPs (Fig. 5). Similar data were obtained with H. glycines CP propeptide, in which the prodomain product strongly inhibited a mature cathepsin L-like proteinase and other CPs of other related phytonematode species, but no effect was observed against insect digestive proteinases [27]. Besides, the procongopain chimeric peptide was also able to inhibit its cognate enzyme, congopain, but was unable to inhibit human cathepsins B and L. The authors suggest that procongopain propeptide is specific to protozoa enzymes, being unspecific to other kingdoms [16]. In order to shed some light over the specificity question, many authors tried to correlate the primary structure with its molecular specificity [25–29]. Nevertheless, this correlation was not elucidated.

4. Conclusion

In this report, a recombinant propeptide from an insect cysteine proteinase was investigated for its ability to inhibit the mature enzyme in order to be biotechnologically used as inhibitor target of specific proteinases. Our data showed that the product of A. obtectus CP prodomain preferentially inhibited midgut CPs from Bruchidae family. Surprisingly, no inhibitory activity against other digestive CP proteinases from other coleopteran genus as R. dominica and A. grandis was found. Understanding the propeptide inhibitory mechanism and its specificity toward the digestive CPs from insect species could be useful for the development of molecular strategies for insect pest control tailored for the inhibition of specific target systems. In summary, this report proposes a different strategy to control bean bruchids by using a novel endogenous
prodomain peptide. In the near future, this strategy could be used as a biotechnological tool, possibly reducing pest infestation and decreasing crop losses, offering several benefits to farmers and consumers.

REFERENCES


