DmCatD, a cathepsin D-like peptidase of the hematophagous insect Dipetalogaster maxima (Hemiptera: Reduviidae): Purification, bioinformatic analyses and the significance of its interaction with lipophorin in the internalization by developing oocytes

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1. Introduction

Triatomines (Hemiptera: Reduviidae) or “kissing bugs” are obligate hematophagous insects with relevance in public health since they are vectors of the protozoan Trypanosoma cruzi, the etiological agent of Chagas’ disease. Currently, about 6–7 million people worldwide are estimated as infected with the parasite (WHO, 2017), with the highest prevalence in Latin America (Hotez et al., 2008). Because T. cruzi is
mainly transmitted to people by triatomines (Miles, 2017), vector control represents the best way to reduce the incidence of the illness.

In insects, vitellogenesis is one of the most important events in reproduction (Raikhel, 2005) and, in triatomine females, it is strongly coupled to the intake of a blood meal (Stoka et al., 1987). Vitellogenesis is characterized by a rapid growth of oocytes due to a remarkable uptake and deposition of proteins, lipids and other molecules. During this process, large amounts of yolk protein precursors synthesized in the fat body and/or in the follicular epithelial cells (Izumi et al., 1994; Melo et al., 2000; Atella et al., 2005) are stored in developing oocytes. Among them, vitellogenin constitutes the main yolk protein precursor that is taken up by the oocytes by receptor-mediated endocytosis and stored as vitellin in specialized lysosomal compartments or yolk bodies (Raikhel and Dhadialla, 1992; Tufail and Takeda, 2008, 2009).

On the other hand, lipophorin is the major insect lipoprotein that carries several lipid classes in the hemolymph to the target tissues (Canavoso et al., 2001). In addition to this physiological role, there are reports indicating the ability of lipophorin to bind other molecules, such as juvenile hormone (Engelmann and Mala, 2000; Zalewska et al., 2009), proteins from the immune system, (Ma et al., 2006; Rahman et al., 2006) and morphogens (Eugster et al., 2007). In several insect species, including triatomines, it was demonstrated that lipophorin can be endocytosed by the oocytes and stored in yolk bodies with vitellin, thus functioning as a yolk protein precursor during vitellogenesis (Kawooya and Law, 1988; Ziegler and Van Antwerpen, 2006; Fruttero et al., 2011; Leyria et al., 2014).

Fertilization triggers embryogenesis, a process in which the yolk proteins are used as substrates for the growing embryo (Yamahama et al., 2005). Different peptides and acid phosphatases have been associated with the degradation of yolk proteins during embryogenesis (Nussenzevg et al., 1992; Yamamoto and Takahashi, 1993; Izumi et al., 1994; Fialho et al., 2005; Oliveira et al., 2008). Most peptides are yolk protein precursors synthesized in the fat body as well as in the ovary, released subsequently to the hemolymph as pro-enzymes and stored in the oocyte associated with the yolk bodies (Giorgi and Nordin, 2005). It was demonstrated that an acid phosphatase and cathepsin D are important regulators of yolk protein degradation during the embryogenesis of the triatomine Rhodnius prolixus (Fialho et al., 2005; Gomes et al., 2010).

Dipterogaster maxima is a triatomine species used as a model to assess biochemical, cellular and molecular events during vitellogenic and post-vitellogenic reproduction stages (Aguirre et al., 2008, 2011). Employing this insect we have demonstrated that DmCatD, a cathepsin D-like peptidase, is synthesized by the fat body and the ovary as a yolk protein precursor and stored as an inactive enzyme (pro-DmCatD) in yolk bodies (Leyria et al., 2015). It was also demonstrated that in females of D. maxima, blood deprivation promotes follicular atresia and oosorption of terminal oocytes. Such processes were characterized by an early activation of DmCatD, which seems to be part of the mechanisms regulating yolk protein degradation during this post-vitellogenic stage (Leyria et al., 2015). In spite of the physiological relevance of DmCatD in the biology of reproduction in triatomines, the mechanism involving its targeting to the oocyte membrane and posterior internalization during vitellogenesis has not been established. In this work, we have purified a DmCatD peptidase from eggs of D. maxima, reported its transcript sequence and structural properties. In addition, bioinformatic and biochemical approaches were performed to assess the interaction of DmCatD-lipoporphin and analyzed the relevance of such potential association for DmCatD internalization in developing oocytes.

2. Materials and methods

2.1. Ethics statement

Housing conditions and manipulation of hens employed in the maintenance of the insect colony followed the protocol authorized by the Animal Care Committee of the Centro de Investigaciones en Bioquímica Clínica e Inmunología (CIBICI-CONICET-Universidad Nacional de Córdoba) in accordance with the guidelines published by the Canadian Council on Animal Care with the assurance number A5802-01 delivered by the Office of Laboratory Animal Welfare (National Institutes of Health). The animal facility at the CIBICI-CONICET is a dependency of the Argentine National Ministry of Science (Sistema Nacional de Bioterios, MINCYT, http://www.bioterios.mincyt.gov.ar). No infective insect species, human blood or hen sacrifice were involved in the study. Details of the approved protocol were recently published (Leyria et al., 2015).

2.2. Chemicals

Rabbit polyclonal anti-cathepsin D (catalog code sc-10725) and rabbit anti-ATP5B/β-chain of ATP synthase (β-ATPase, catalog code sc-33618) antibodies, both of human origin, were from Santa Cruz Biotechnology (Palo Alto, CA, USA). The cross-reactivity between the anti-cathepsin D antibody and DmCatD was already reported (Leyria et al., 2015). Goat anti-rabbit IgG labeled with Alexa Fluor® 568 antibody (Molecular Probes, Eugene, OR, USA); Tissue-Tek embedding medium Optimal Cutting Temperature (OCT) (Miles, Elkhart, IN, USA); MMLV reverse transcriptase (Promega, Heidelberg, Germany); Platinum® Taq DNA Polymerase High Fidelity (Thermo Fisher Scientific, Waltham, MA, USA); primers (Sigma Genosys, Houston, TX, USA); MasterPure RNA Purification Kit (Epicenter Biotechnologies, Madison, WI, USA); Fluorsave (Calbiochem, Darmstadt, Germany) and Color Prestained Protein Standard (New England Biolabs Inc., Ipswich, MA, USA) were from the indicated commercial sources. The fluorogenic peptide substrate Abz-AIAFSRQ-EDDnp (Abz, orthoaminozobenzoic acid, EDDnp, ethylenediamine-2,4-dinitrophenyl) was a kind gift from Dr. Maria Aparecida Juliano (Universidade Federal de São Paulo, Brazil). Bovine serum albumin (BSA), dimethylpimelimidate (DMP), fetal bovine serum (FBS), anti-mouse IgG conjugated to FITC antibody and all other chemicals were from Sigma-Aldrich (St. Louis, MO, USA).

2.3. Insects

Experiments were carried out with insects taken from a colony of D. maxima, maintained at 28 °C, 70% relative humidity, 8:16 h light:dark photoperiod. Insects were fed on hen blood (Canavoso and Rubiola, 1995), according to the recommendations of the National Institute of Parasitology (Health Ministry, Argentina) (Núñez and Segura, 1987). Standardized conditions of insect rearing were previously described (Aguirre et al., 2008). Briefly, fifth-instar females were separated from the males before feeding. Newly emerged females were segregated individually and placed together with two recently fed males during 48 h. Mating was checked by observation of the spermatophore. Mated females were kept in individual containers until they were able to feed a blood meal (days 10–12 post-ecdysis), which resulted in a 3.0–5.5-fold increase in the body weight of the insect. Experimental approaches were performed using vitellogenic females at days 4–6 after blood feeding (Aguirre et al., 2008; Leyria et al., 2015).

2.4. Purification of DmCatD from eggs

In a typical purification experiment, 60 eggs from D. maxima collected within 24 h post oviposition were homogenized using a Potter-Elvejem (15 strokes) in cold 20 mM sodium phosphate buffer (NaPB), pH 6.0 with the addition of protease inhibitors E64 and PMSF at final concentrations of 15 and 80 µM, respectively. The homogenate was centrifuged twice at 5000 × g (10 min each, 4 °C) and the supernatant was recovered to proceed with the protein determination (Bradford, 1976).

Fractions from purification steps were analyzed using the fluorogenic substrate for cathepsin D, Abz-AIAFSRQ-EDDnp, as reported.
previously (Aguirre et al., 2011; Leyria et al., 2015). Additional activity assays were conducted in presence of the aspartyl protease inhibitor pepstatin A.

2.4.1. Cation-exchange chromatography

The first purification step of the egg homogenate was performed by Fast Protein Liquid Chromatography (FPLC), using a column packed with 10 ml of CM-Sepharose Fast-Flow resin (GE Healthcare, Amersham Biosciences, Little Chalfont, England) which was equilibrated with 20 mM NaPB, pH 6.0. The egg homogenate was loaded and the column was washed with buffer to remove non-retained proteins, then, bound proteins were eluted with NaPB at increasing concentrations of NaCl (100, 200, 300 and 500 mM). Non-retained and eluted fractions were collected and assayed for cathepsin D activity as described previously (Aguirre et al., 2011). The fraction eluted with 200 mM NaPB showed the highest enzymatic activity and was subsequently concentrated using a 10 kDa cut off Centriprep™ Centrifugal Filter Concentrators (EMD Millipore, Billerica, MA, USA).

2.4.2. Size exclusion chromatography

Gel filtration was performed using a Superdex 75 column (1.2 × 30 cm) (GE Healthcare, Amersham Biosciences, Little Chalfont, England) equilibrated in 20 mM NaPB pH 6.0. Two milliliters of the sample eluted from the cation-exchange chromatography column at 200 mM NaCl were applied to the column and protein peaks, monitored at 280 nm, were individually collected in 0.5 ml fractions and assayed for enzymatic activity.

2.5. Electrophoresis, in-gel trypsin digestion and mass spectrometry analysis

Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was performed as described by Laemmli (1970) in a 12% separating gel to visualize the purified protein.

Protein bands were manually excised from Coomassie stained gels and digested in-gel with trypsin. The preparation for mass spectrometry was performed as reported in Fruttero et al. (2014) with minor modifications. To identify proteins, a liquid chromatography (LC) separation (reversed-phase High Performance Liquid Chromatography, HPLC) coupled with tandem mass spectrometry (MS/MS) strategy was used. MS/MS analyses were performed in an electrospary ionization (ESI)-Q-Exactive mass spectrometer equipped with a High Collision Dissociation cell and an Orbitrap analyzer coupled to an EASY-nLC 1000 Liquid Chromatography system (Thermo Scientific, Waltham, MA, USA) at the CEQUIBIEM mass spectrometry facility (Centro de Estudios Químicos y Biológicos por Espectrometría de Masas, FCEyN, UBA, Argentina). The CEQUBIEM mass spectrometry facility (Centro de Estudios Químicos y Biológicos por Espectrometría de Masas, FCEyN, UBA, Argentina). The MS/MS spectra were analyzed using Proteome discoverer v. 1.4 software (Thermo Scientific, Waltham, MA, USA) and the D. maxima (AHE57676) and T. infestans (AEO94539) databases of The National Center for Biotechnology Information (NCBI) were employed for the analyses. Search parameters allowed a maximum of two missed cleavages, the carbamidomethylation of cysteine, the possible oxidation of methionine, precursor mass tolerance of 10 ppm and fragment mass tolerance of 0.05 Da. Only the matches considered of high confidence by the software were taken into account.

2.6. RT-PCR and sequencing

In order to obtain the RNA sequence of DmCatD, Reverse Transcription – Polymerase chain reaction (RT-PCR) experiments were performed. For RNA extraction, dissected fat bodies of three females were pooled and the MasterPure RNA Purification Kit was used according to the manufacturer’s protocol. To eliminate genomic DNA, samples were treated with DNase provided in the kit. RNA integrity was evaluated by electrophoresis in a 1% agarose gel, and only the band corresponding to 18S rRNA was observed. It is important to highlight that 28S rRNA of most insects contains an endogenous “hidden break” upon denaturation, the masking hydrogen bonds are disrupted, releasing two similar sized fragments that both migrate closely with 18S rRNA (Winnebeck et al., 2010). cDNA was synthesized from 2 μg of total RNA by reverse transcription reaction using oligo dT and the MMLV reverse transcriptase protocol. For PCR, Platinum® Taq DNA Polymerase High Fidelity was used. Since the genome of D. maxima is not available, the design of primers was based on the mRNA sequence of the Triatoma infestans cathepsin D (ID JN606068.1). Primers were: 5′-GCTTCTCAAATCTTTGATTACCCTC-3′ (sense) and 5′-TTATGTGTTC AACACTGCGAAGCTAC-3′ (antisense). Using this set of primers we were able to obtain a product of approximately 902 bp, encompassing about 73% of the full-length sequence of the transcript (lacking the N-terminus). PCR reactions were carried out and the products were processed using the sequencing service of the Department of Biochemistry and Molecular Biology, Oklahoma State University, OK, USA.

2.7. Bioinformatic analyses for DmCatD

The deduced amino acid sequence, molecular mass and isoelectric point prediction were assessed using tools available on ExPASy (www.expasy.org – SIB Bioinformatics Resource Portal) (Artimo et al., 2012). The active site of the peptidase was detected by ScanProsite (Sigrist et al., 2013). The bioinformatic tool from NCBI was used to analyze highly conserved regions of DmCatD. The prediction of post-translational modifications was carried out with CBS (http://www.cbs.dtu.dk/services/) and ScanProsite (http://prosite.expasy.org/scanprosite/), through multiple tools (Blom et al., 1999; Petersen et al., 2011; Sigrist et al., 2013). Modifications examined included the occurrence of phosphorylation and N-myristoylation sites. The DmCatD from D. maxima structural model was built based on crystallographic data from proteins with similar secondary structure arrangements, using Phyre2 server (Kelley et al., 2015). The structure was stereochemically evaluated using Procheck (Laskowski et al., 1993). The R. prolixus lipo- phorin structural modeling (corresponding to the apolipoporin II/I gene, VectorBase ID RPREG002125-PA) was carried out with I-TASSER (Zhang, 2008; Roy et al., 2010; Yang et al., 2015), while the DmCatD-lipophorin docking simulation was performed with two independent macromolecular docking programs with no positional biases: PatchDock (Schniedman-Duhovny et al., 2005) and PIPER (Kozakov et al., 2006)) via ClusPro 2.0 (Comeau et al., 2004).

Hydropophobity profiles were calculated with the Kyte-Doolittle scale (Kyte and Doolittle, 1982), and the electrostatic surfaces were generated with APBS tools (Baker et al., 2001) under UCSF Chimera (Pettersen et al., 2004).

Evolutionary analyses were conducted with MEGA6 (Tamura et al., 2013). The phylogenetic tree for DmCatD was inferred using the Maximum Likelihood method (LG + G) (Le and Gascuel, 2008). Confidence was assessed by bootstrap pseudo-replications (1000 rounds). Sequences with similarity to DmCatD were searched with BlastP (Johnson et al., 2008) and employed in the tree reconstruction.

Except for the evolutionary analysis, all bioinformatics studies were conducted with a chimeric construct that included the N-terminus region of T. infestans cathepsin D.

2.8. Hemolymph collection

The hemolymph was collected with a Hamilton syringe from immobilized vitellogenic females at days 4–6 after blood feeding. Their legs were sectioned at the level of the coxa and the hemolymph was collected into cold microtubes, in the presence of 10 mM NaN3EDTA, 5 mM dithiothreitol and a cocktail of protease inhibitors: 1 mM phenylmethyl-sulfonyl fluoride (PMSF), 1 mM N-α-p-tosyl-L-lysine chloromethyl ketone (TLCK), 1 mM pepstatin A and 0.3 mM aprotinin (Fruttero et al., 2009). Samples were centrifuged at 10,000 × g for 5 min at 4 °C to remove hemocytes and then stored at −70 °C, after protein determination (Bradford, 1976).
2.9. Co-immunoprecipitation of DmCatD with lipophorin

Co-immunoprecipitation assays using hemolymph from vitellogenic females were performed as described previously (Fruttero et al., 2017). Anti-β-ATPase (control) and anti-cathepsin D antibodies (0.2 µg each) were covalently coupled to protein A Mag Sepharose beads (GE Healthcare, Little Chalfont, UK) by their incubation for 1 h at room temperature. The magnetic beads were washed with a 5-fold dilution of the sample in Tris buffered saline (TBS, 50 mM Tris, 150 mM NaCl, pH 7.5) and further recovered with a magnetic rack. After two washes, the beads were incubated with 200 mM triethanolamine buffer (pH 8.9) containing 50 mM DMP for 1 h at room temperature. The beads were washed with triethanolamine buffer as already stated and blocked with 100 mM ethanolamine buffer (pH 8.9) for 15 min at room temperature. The elution buffer (0.1 M glycine, 2 M urea, pH 2.9) was added to remove the unbound antibody and then beads were washed three times with TBS. Hemolymph or rat brain homogenates (control), both containing 60 µg of total proteins were incubated with the antibodies covalently coupled to protein A Mag Sepharose beads (anti-β-ATPase or anti-cathepsin D antibodies, 1 h at room temperature, with slow end-over-end mixing). The proteins bound to the beads were eluted with the elution buffer and protein A Mag Sepharose beads were removed using a magnetic rack. The eluted proteins, the input and the standard were subjected to Tris-Tricine-SDS gel electrophoresis as described elsewhere (Fruttero et al., 2014). The immunodetection of DmCatD-lipophorin interaction was performed by western blot, using an anti-lipophorin antibody (anti-Lp, 1:1000) obtained as described previously (Canavoso and Rubiolo, 1995). The secondary antibody, Li-Cor IRDye 800CW polyclonal goat anti-rabbit IgG (1:15,000) was incubated at room temperature for 1 h. Both antibodies were diluted in TBS-0.1% Tween 20 containing 5% non-fat milk. After washing, blots were scanned and analyzed with the Odyssey quantitative western blot near-infrared system (Li-Cor Biosciences, Lincoln, NE, USA) using default settings. DmCatD-vitellogenin interaction in the hemolymph of vitellogenic females was also tested employing a similar protocol but using an anti-vitellin antibody (anti-Vt, 1:1000) obtained as described previously (Aguirre et al., 2008).

2.10. Colocalization of DmCatD with lipophorin in the ovarian tissue

Ovaries from vitellogenic females at days 4–6 after blood feeding were dissected out in cold phosphate buffered saline (PBS, 6.6 mM Na2HPO4/KH2PO4, 150 mM NaCl, pH 7.4), using a standard stereoscope with an optic fiber light source and processed for cryostat sectioning as reported previously (Leyria et al., 2015). Tissue sections of 8 µm were obtained with a Leica CM1510 cryostat (Leica Microsystems, Wetzlar, Germany) and placed onto poly-L-lysine-treated glass slides. Ovarian sections were incubated with 1% BSA and 5% PBS in block non-specific binding sites. The slides were sequentially incubated with the anti-cathepsin D (1:100), the anti-rabbit IgG labeled with Alexa Fluor®
568 (1:400) and the anti-Lp conjugated to fluorescein isothiocyanate (FITC, 1:40) antibodies. Antibodies were diluted in 1% BSA in PBS. All incubations were performed inside a humid chamber at 37 °C for 1 h. Slides were rinsed twice with PBS for 5 min. Control experiments were carried out by omitting one or a combination of the following antibodies: anti-rabbit IgG coupled to Alexa Fluor® 568, anti-cathepsin D or anti-Lp-FITC. An additional control for autofluorescence was conducted using an anti-mouse IgG antibody conjugated to FITC (irrelevant antibody, 1:40). Slides were rinsed with PBS, air-dried, mounted in Fluorsave and observed with an Olympus FV300 laser scanning confocal microscope.

3. Results

3.1. Purification of DmCatD from eggs of D. Maxima

The activity of cathepsin D peptidase in egg homogenates from D. maxima was confirmed, in line with our previous report on the fat body, Table 1. Identification of the purified protein by tandem mass spectrometry (MS/MS). The band obtained from gel filtration chromatography was excised from the gels and identified by LC-ESI-Orbitrap. The results were analyzed employing the search engine of the Proteome Discoverer (version 1.4) software, using the databases of D. maxima (AHE57676) and T. infestans (AEO94539) available at the National Center for Biotechnology Information (NCBI).

<table>
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<th>Identified peptides</th>
<th>Sequence coverage (%)</th>
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<td>Dipetalogaster maxima</td>
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<td>Triatoma infestans</td>
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</table>

Fig. 2. (A), Nucleotide and the corresponding deduced partial amino acid sequences of DmCatD. (B), In order to perform bioinformatic studies, the homologue region from Triatoma infestans cathepsin D was used to complete the N-terminal region of the sequence (amino acids in bold). An arrow indicates the catalytic residue in the active site; underlined regions show a catalytic motif of the active site; asterisks indicate an active site flap; the post-translational cleavage site is shown between the two highlighted blocks; the stars indicate the cysteine residues and the white block shows the proline-loop region. (C), Putative phosphorylation and N-myristoylation sites.
ovarian tissue and hemolymph (Leyria et al., 2015). Based on this result, purification of DmCatD was conducted by a combination of cation-exchange and gel filtration chromatography, employing homogenates of eggs collected within 24 h post oviposition as starting biological material. After the first chromatographic step, the activity of cathepsin D peptidase was found to be highest in the fraction eluted with 200 mM NaCl (data not shown). After gel filtration, a major peak of cathepsin D peptidase activity was eluted at 10.5 ml (Fig. 1A). When this fraction was subjected to SDS-PAGE, a single band of approximately 43 kDa was detected (Fig. 1B). As shown in Fig. 1C, the yield of purified DmCatD peptidase was 6.3% achieving 64-fold of purification.

The 43 kDa band obtained from gel filtration was excised from the gel, digested with trypsin and the resulting peptides were analyzed by MS/MS. When searched against the NCBI database, the peptide profiles matched the cathepsins D from D. maxima and T. infestans (Table 1). Three independent samples from different purification batches yield the same set of results.

3.2. Sequence analysis and structural properties of DmCatD

When the RT-PCR assays were performed to obtain the sequence of DmCatD, a single PCR product of the expected size and sequence was amplified. As expected, approximately 73% of full-length DmCatD transcript was obtained, sequenced and annotated in the GenBank under the ID KF724683.1. The protein has a predicted sequence of 284 amino acids (GenBank: AHE57676.1) (Fig. 2A). Since the N-terminus is necessary to perform specific bioinformatic analyses and the DmCatD sequence was not complete, the homologue region from T. infestans Cathepsin D was used to form a chimeric full-length sequence (amino acids in bold, Fig. 2B).

Bioinformatic analysis performed using NCBI tools showed that in the chimeric cathepsin D construction, the sequence corresponding to DmCatD presented a catalytic aspartic peptidase active site motif, DTGS, in the C-terminus lobe (Fig. 2B, underlined region) and displayed a conserved catalytic site of aspartate (Fig. 2B, arrow). In addition, DmCatD showed an extended loop, projecting over the cleft to form a flap of 11 residues in the active site (RLVYGKSMVG, Fig. 2B, asterisks), which is conserved in cathepsin D family (Metcalf and Fusek, 1993). It was also found that in DmCatD, 5 of the 6 cysteine residues, which are characteristic of the family of aspartic peptidases (Shewale and Tang, 1984; Fusek and Větvička, 2005), were conserved (Fig. 2B, stars).

DmCatD also showed a post-translational cleavage site that varies in length and amino acid composition in different species (Fig. 2B, shaded regions) (Zaidi et al., 2008).

Bioinformatics approaches that included the prediction of putative phosphorylation and N-myristoylation sites occurring in the DmCatD sequence are summarized in Fig. 2C. On the other hand, the structural properties of DmCatD peptidase were assessed in silico employing the chimeric construct that included the N-terminus sequence of T. infestans cathepsin D. The analysis of the structural model for DmCatD (Fig. 3A) indicated that the protein has a positively charged, hydrophilic surface as one of its main features (Fig. 3B and C, respectively).
3.3. Molecular phylogenetic analysis of DmCatD

According to the phylogenetic tree obtained upon the evolutionary analysis carried out with the DmCatD sequence deposited in NCBI database (ID KF724683.1), cathepsin D from the hemipterans *D. maxima*, *T. infestans* and *Riptortus pedestris* but not from *Halyomorpha halys* clustered together even though they are distantly related to the other sequences, including those from insects (Fig. 4). Cathepsin D from *H. halys* is shown as an intermediate between such a cluster and the other inspected sequences. Thus, cathepsin D from these four hemipteran species constitutes a unique group of enzymes, with peculiar features that suggest distancing from "classical" cathepsin D peptidases. Most of the insect sequences also clustered following their orders as can be seen for the hymenopterans *Bombus impatiens*, *Melipona quadrifasciata*, *Apis florea*, *Apis dorsata*, *Apis mellifera*, *Polyrhachis vicina*, *Vollenhovia emeryi* and *Wasmannia auropunctata*. However, the louse *Pediculus humanus* was placed together with the non-insect copepods *Caligus clemensi* and *Lepeophtheirus salmonis*. The sequences of the remaining non-insect groups analyzed, including bivalves, ticks and mammals also clustered together.

Alignment of the proline-loop region of all the sequences employed in the phylogenetic inference and the consensus sequence for this motif showed that cathepsin D from hemipterans had a nonconserved proline loop (Fig. 5).

3.4. DmCatD and lipophorin interaction: co-immunoprecipitation, colocalization and docking analysis

Taking into account the relevant function of DmCatD in the reproductive biology of *D. maxima* (Aguirre et al., 2011; Leyria et al., 2015) we performed approaches to address the role of lipophorin in carrying DmCatD to developing oocytes. Co-immunoprecipitation assays were carried out to test the interaction between DmCatD-lipophorin and DmCatD-vitellogenin in the hemolymph of vitellogenic females of *D. maxima*. In our experimental conditions, the results showed that DmCatD co-immunoprecipitated with endogenous lipophorin but not with vitellogenin (Fig. 6A and B). Furthermore, the colocalization of DmCatD and lipophorin was investigated by immunofluorescence, analyzing vitellogenic follicles. Colocalization of lipophorin with DmCatD was detected in the perioocytic space, likely bound to the oocyte membrane, and in the yolk bodies (Fig. 7, merged images, upper panel). Negative controls corresponding to immunofluorescence assays show the specificity of the antibodies (Fig. 7, lower panel). Taken together, the results support the interaction between DmCatD and lipophorin in the hemolymph from vitellogenic females, and strongly suggest that such a lipoprotein acts as a DmCatD carrier cooperating with its targeting and posterior internalization by the oocytes.

In addition, computational assays were undertaken to analyze the feasibility of a binding between DmCatD and lipophorin. For that
Fig. 5. Alignment of the proline-loop region. All sequences employed in the phylogenetic inference were aligned by the proline-loop region. The consensus sequence for this motif is shown below the alignment, the conserved amino acids are highlighted using the same color and the sequences corresponding to the hemipteran species are indicated by a bracket.

<table>
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<th>Species</th>
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<td>Bos Taurus</td>
<td>G P M G M D I P P P G G P L M</td>
</tr>
<tr>
<td>Ixodes scapularis</td>
<td>G P M G L G M P Q - - S P L M</td>
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<tr>
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<td>G P M S L E M P Q - - S P L M</td>
</tr>
<tr>
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<td>G P T G M T L P Q - - - I D M</td>
</tr>
<tr>
<td>Ripoptopus pedestris</td>
<td>G P E P T - - - - - - - - Y A F</td>
</tr>
<tr>
<td>Dipetalogaster maxima</td>
<td>G P A T M - - - - P R A P Q F P W</td>
</tr>
<tr>
<td>Triatoma infestans</td>
<td>G P T T L - - - - P S A P Q F P W</td>
</tr>
</tbody>
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Hemiptera

Fig. 6. (A), Co-immunoprecipitation (Co-IP) of endogenous Lipophorin (Lp) and DmCatD. The hemolymph of vitellogenic females or rat brain homogenates (control) were incubated with 0.2 µg of anti-cathepsin D or 0.2 µg of anti-ATP5b antibodies (control). Samples were transferred to nitrocellulose membranes and then probed with an anti-lipophorin antibody (1:1000). Arrow indicates apoLp-II subunit (~ 80 kDa). Lp line: purified Lp loaded as reference. The presence of endogenous Lp was visualized by loading hemolymph and probing the sample with the anti-lipophorin antibody (Input). (B), Co-immunoprecipitation of endogenous vitellogenin (Vg) and DmCatD. The hemolymph of vitellogenic females was incubated with 0.2 µg of anti-cathepsin D antibodies. Samples were transferred to nitrocellulose membranes and then probed with anti-vitellin antibody (1:1000). The main subunits of Vg (Mr ~ 170 kDa and 174 kDa), visualized as a single immunoreactive band, are shown with an arrow. Vg lane: purified Vg loaded as a reference. The presence of endogenous Vg was visualized by loading hemolymph and probing the sample with the anti-vitellogenin antibody (Input).
purpose, we modeled the lipophorin structure employing the apolipo-
phorin II/I sequence from R. prolixus. The resulting model was then
used to perform docking assays involving lipophorin and DmCatD. As
shown in Fig. 8, the obtained complexes were convergent regarding
their binding location and orientation. Since these are docking ar-
rangements obtained from two independent software programs,
without any spatial restriction for the docking search space, their
common focal localization for the DmCatD-lipophorin complex forma-
tion highlights its higher probability as a physiological complex in the
insect.

4. Discussion

Cathepsin D is a soluble lysosomal aspartic endopeptidase involved
in the degradation and/or activation of proteins, hormones and en-
zymes, among other functions (Benes et al., 2008). Regarding triato-
mines, cathepsin D is implicated in yolk protein degradation during the
embryonic development of R. prolixus (Fialho et al., 2005; Gomes et al.,
2010) as well as in the digestion of ingested blood proteins in T. in-
festans (Balczun et al., 2012). In D. maxima, DmCatD peptidase was
necessary for promoting early degradation of vitellin if follicular atresia
was triggered by deprivation of blood meals (Aguirre et al., 2011;
Leyria et al., 2015). In this work, we have obtained original information
about DmCatD by achieving its purifi-
cation from egg homogenates. By
ion-exchange chromatography and gel
fi-
ltration (Fig. 1A), a standard
protocol for purifying cathepsin D as well as other peptidases
(Defferrari et al., 2011), we have found that DmCatD from eggs dis-
played a molecular mass of 43 kDa (Fig. 1B), similar to those reported
for pro-DmCatD in the fat body and ovaries by western blots (Leyria
et al., 2015). The MS/MS analysis of the 43 kDa band obtained after
puriﬁcation retrieved peptide proﬁles matching those of the cathepsin
D from D. maxima (AHE57676) and T. infestans (AEO94539), con-
ﬁrming thus its identity (Table 1).

In this work, the predicted protein sequence of DmCatD
(AHE57676.1) shares an 86% identity with cathepsin D protein of T.
infectans (ADK47877.1) (Fig. 2). On the other hand, a pro-cathepsin of
Putative binding arrangement for the DmCatD-lipophorin complex

Fig. 8. Arrangement for putative binding of the complex between chimeric cathepsin D and lipophorin. Ten top-ranked docking solutions are shown as colored cathepsin D monomers bound to lipophorin, depicted in grey cartoon with superposed molecular surface. The solutions are densely concentrated in the same spot of the protein, reinforcing its high probability as a binding site.

~42 kDa molecular mass in the chimeric cathepsin D construction is close to the mass of the purified DmCatD from eggs and MS/MS analysis reported here (Fig. 1), and it is also in agreement with the pro-DmCatD described in Leyria et al. (2015) and to the findings in other insects (Padilha et al., 2009; Gui et al., 2006; Kang et al., 2017).

Bioinformatic analysis revealed that the chimeric cathepsin D displays a conserved catalytic site consisting in two aspartate residues, one of them found in the DmCatD sequence (Fig. 2B). These aspartate residues are known to play a key catalytic role in cathepsins D and E of the pepsin family as well as in renins (Dunn, 2002; Fusek and Větvička, 2005). On the contrary, the aspartic peptidase BYC from the tick Rhizophalus (Boophilus) microplus eggs, which is responsible for the yolk degradation during embryogenesis, lacks the second aspartate residue from the catalytic site (Nascimento-Silva et al., 2008). Eukaryotic aspartic peptidases contain two conserved DTGS/T motifs, each one provides a catalytic aspartate residue to the active site (Davies, 1990). In DmCatD peptidase sequence, the DTGS motif in the C-terminus lobe was strictly conserved (Fig. 2B).

The deduced DmCatD peptidase sequence showed putative phosphorylation and N-myristoylation sites (Fig. 2C), that could be relevant in the post-translational modification of the protein and, consequently, in the regulation of its binding and enzymatic properties.

The structural model for the chimeric construct of cathepsin D assessed in silico indicated that this peptidase exhibited a positively charged, hydrophilic surface as main features (Fig. 3), being the latter a characteristic expected for a soluble protein. These properties might contribute to its interaction with lipophorin, discussed below, which in turn has predominantly a negative surface charge (Roosendaal et al., 2009). However, the contribution of the hydrophobic and negative patches of DmCatD surface to the binding capacity of the lipoprotein cannot be disregarded (Smith and Davidson, 2010).

Molecular phylogenetic analysis from selected taxa indicated that in Hemiptera, cathepsin D clusters together but not in the expected pattern from species-level phylogeny (Fig. 4). Within the cladogram, hemipteran cathepsin D is clearly separated from that of other insects and vertebrates. This clustering pattern has been observed recently in a survey for cathepsin D from the midgut of the hemipteran Dysdercus peruvianus (Pimentel et al., 2017). The main reason for this separation seems to be the non-lysosomal nature of the majority of cathepsin D peptidases in hemipterans, as indicated by their non-conserved proline loop (Fig. 5). In Musca domestica, three cathepsin D peptidases were reported in the midgut, two of them lacking the proline loop. It was suggested that this characteristic could be somehow associated with an extracellular role for these enzymes (Padilha et al., 2009). In the hemipteran D. peruvianus, nine of ten cathepsin D transcribing genes that were specifically expressed in the midgut lack the proline loop (Pimentel et al., 2017). Additionally, two cathepsins D having or not a proline loop, were found to be secreted into the midgut lumen of T. infestans (Balcun et al., 2012). The authors proposed that such a finding could be an adaptation limited to triatomines. In this work, the results indicated that DmCatD displayed features in common with secreted peptidases. Moreover, we recently reported that DmCatD was localized in yolk bodies, which are specialized compartments in developing oocytes (Leyria et al., 2015). These findings can be related to the physiological role of DmCatD assigned in D. maxima, which functions as a yolk protein precursor during vitellogenesis but also as an acid peptidase that regulates vitellin degradation.

We have reported that DmCatD, which is synthesized as a yolk protein precursor in the fat body and the ovarian tissue, is involved in vitellin degradation during follicular atresia (Leyria et al., 2015). Studies in A. aegypti demonstrated that a vitellogenic carboxypeptidase (VCB) and a thiol cathepsin B pro-protease (VCP), which are both involved in yolk degradation during embryogenesis, are synthesized in the fat body as yolk protein precursors and stored in developing oocytes along with vitellin (Cho et al., 1991, 1999). However, how these yolk protein precursors are targeted from the fat body to oocytes through the hemolymph and further internalized remains to be elucidated. Swevers et al. (2005) proposed that lipophorin serves as a carrier for VCB and VCP, and that the internalization of these yolk protein precursors probably occurs via a “piggyback” mechanism. Up to date, no experimental evidence supporting this pathway was reported.

Lipophorin mostly delivers its hydrophobic cargo to developing
The interaction of DmCatD with lipophorin demonstrated in this work complex at the oocyte membrane as a step for its further internalization (Fruttero et al., 2017), contributes in docking the lipophorin-DmCatD membrane of the cells (Ziegler and Van Antwerpen, 2006). However, in other tissues mediates the internalization of lipophorin particles. Taking into account the relevant role of DmCatD in the reproductive biology of D. maxima (Aguirre et al., 2011; Leyria et al., 2015) and considering that the mechanism involved in its targeting to developing oocytes is unknown, we performed experiments to address the role of lipophorin as a carrier of DmCatD in vitellogenic females. Co-immunoprecipitation assays (Fig. 6) and colocalization of lipophorin and DmCatD (Fig. 7) support their endogenous association. The association between cathepsin D and a lipoprotein has been described previously. Thus, it was reported that cathepsin D associated with both, lipid-free recombinant full-length human apolipoprotein E and lipidated human plasma full-length apolipoprotein E, playing a possible role in Alzheimer’s disease (Zhou et al., 2006). Additionally, the results from computational docking assays favored the lipophorin-DmCatD interaction (Fig. 8).

From a physiological point of view, this work provides evidence indicating that lipophorin acts as a carrier for DmCatD, which in turn could be part of a mechanism of functional relevance for its transport to developing oocytes and further internalization. In contrast to lipophorin, which binds to several molecules such as proteins and hormones (Ma et al., 2006; Rahman et al., 2006; Zalewska et al., 2009), vitellogenin has been reported to interact with few partners (Engelmann and Mala, 2000). Therefore, it seems very unlikely that vitellogenin participates in the transport of DmCatD through circulation since under our experimental conditions no co-immunoprecipitation between these proteins in the hemolymph of vitellogenic females was observed. On the other hand, even though hemolymph pH is always near neutrality (Harrison, 2001), the microenvironment associated to the extracellular side of the plasma membrane may exhibit pH changes (Maouyo et al., 2000) that could lead to cathepsin D activation. Thus, binding of lipophorin with DmCatD may also be important in preserving such a peptide from both activation and/or proteolytic events in the hemolymph.

Until now, the mechanisms involved in the internalization of peptides within the oocytes of insects have not been reported. In mammals, intracellular transport of pro-cathepsin D is mediated by the mannose-6-phosphate receptors (M6Pr), which capture and target the proenzyme to lysosomal compartments (Fusek and Větvička, 2005). However, alternative M6Pr-independent mechanisms have also been reported (Laurent-Matha et al., 1998). For instance, in mammalian fibroblasts, secreted pro-cathepsin D is partly endocytosed by LDL receptor-related protein-1 (LRP1) (Deroç et al., 2012). This finding suggests a role of LpR in the endocytosis of pro-DmCatD since both, LRP and LpR are members of the LDL receptor family (Ziegler and Van Antwerpen, 2006). Furthermore, the colocalization of lipophorin and DmCatD in yolk bodies (Fig. 7) also suggests the LpR participation. It is also important to analyze if the non-endocytic lipophorin receptor, the β subunit of the ATP synthase complex (β-ATPase), which is expressed in the oocytes of a related triatomine Panstrongylus megistus (Fruttero et al., 2017), contributes in docking the lipophorin-DmCatD complex at the oocyte membrane as a step for its further internalization.

In summary, DmCatD is part of a physiological mechanism that regulates yolk protein degradation in D. maxima (Leyria et al., 2015). The interaction of DmCatD with lipophorin demonstrated in this work offers a unique scenario to unveil the participation of different lipophorin receptors, which in turn may modulate each other or function in coordination by promoting the internalization of DmCatD by developing oocytes.

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