Characterization of a membrane-bound aminopeptidase purified from Acyrthosiphon pisum midgut cells

A major binding site for toxic mannose lectins

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Database
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A single membrane-bound aminopeptidase N (APN) occurs in the pea aphid (Acyrthosiphon pisum Harris) midgut, with a pH optimum of 7.0, pI of 8.1 and molecular mass of 130 kDa. This enzyme accounts for more than 15.6% of the total gut proteins. After being solubilized in detergent, APN was purified to homogeneity. The enzyme is a glycoprotein rich in mannose residues, which binds the entomotoxic lectins of the concanavalin family. The internal sequence of APN is homologous with a conservative domain in APNs, and degenerated primers of highly conserved APN motifs were used to screen a gut cDNA library. The complete sequence of APN has standard residues involved in zinc co-ordination and catalysis and a glycosyl-phosphatidylinositol anchor, as in APNs from Lepidoptera. APN has a broad specificity towards N-terminal amino acids, but does not hydrolyze acidic aminoacyl-peptides, thus resembling the mammalian enzyme (EC 3.4.11.2). The $k_{cat}/K_m$ ratios for different di-, tri-, tetra-, and penta-peptides suggest a preference for tripeptides, and that subsites $S_1$, $S'_2$ and $S'_3$ are pockets able to bind bulky aminoacyl residues. Bestatin and amastatin bound APN in a rapidly reversible mode, with $K_i$ values of 1.8 μM and 0.6 μM, respectively. EDTA inactivates this APN ($k_{obs}$ 0.14 m⁻¹·s⁻¹, reaction order of 0.44) at a rate that is reduced by competitive inhibitors. In addition to oligopeptide digestion, APN is proposed to be associated with amino-acid-absorption processes which, in contrast with aminopeptidase activity, may be hampered on lectin binding.

Aminopeptidase N (APN) is an exopeptidase that catalyzes the sequential release of N-terminal amino acids of peptides (EC 3.4.11.2). It is found in the midgut of insect larvae either as soluble enzyme or associated with the microvillar membrane. Properties of APN preparations from midgut tissue have been described for at least six orders of insects [1,2].

APNs are the major proteins in some insect midgut microvillar membranes. Probably linked to its abundance, APN is one of the targets of the insecticidal Bacillus thuringiensis δ-endotoxins [3–6]. These toxins, after binding to receptors such as APNs, form channels through which midgut cell contents leak, finally leading to insect death [7]. Also, in humans, APN is the binding site for coronavirus infection [8].

Such findings raised interest in this enzyme, leading to APN cloning from target insects in the Lepidoptera family [3,6,9–20]. All these APNs are inserted into the midgut microvillar membrane by a C-terminal glycosyl-phosphatidylinositol (GPI) anchor. Sequence comparisons with vertebrate and fungal aminopeptidases showed that their most striking similarities were in the zinc-binding motif, including residues His142, His146, and Glu166 (putative zinc ligands, putting residues in a plane to form a five-coordinate environment).

Abbreviations
APN, aminopeptidase N; ConA, concanavalin A; ConBr, concanavalin A ortholog from Canavalia brasiliensis; EST, expressed sequence tag; GPI, glycosyl-phosphatidylinositol; LeuPNA, L-leucine-p-nitroanilide; WGA, wheat germ agglutinin.
numbering according to thermolysin), and Glu143 (catalytic active residue). This conserved motif classifies the enzymes as members of the M1 family of neutral zinc metallopeptidases [21]. In spite of these research efforts, there are few detailed studies on the substrate specificity of lepidopteran microvillar APNs [1,2].

Kinetic data on a midgut APN from Coleoptera showed its similarity to mammalian APN, a family showing a broad specificity towards aminoacyl β-naphthylamides. Chemical modification experiments revealed that a metal ion, a carboxylic group, and the lateral chains of His, Arg and Tyr are important for enzyme activity [22,23].

APN sequences obtained so far are restricted to the Lepidoptera, although insect targets of B. thuringiensis toxins now include many Coleoptera (beetles) and Diptera (flies, mosquitoes). A homologue aminopeptidase has been found in the Drosophila genome [24], and several enzymes found in Rhynchosciara americana have been characterized [25,26]. No membrane-bound aminopeptidase from Hemiptera (bugs, aphids, whiteflies, scales) has been studied so far [2], nor has any truly hemipteran-active B. thuringiensis toxin yet been identified. In fact, the Hemipteran Dysdercus peruvianus has a soluble aminopeptidase [27]. Although they are key components of trophic and toxic interactions involving insects, comparative structural and functional data on insect aminopeptidases are lacking.

In aphids, APN occurs in the apical network of lamellae, which in this insect replaces the usual regularly arranged microvilli [28]. Furthermore, Sauvion et al. [29] found strong interaction of the lectin concanavalin A (ConA) with putative glycosylated receptors at the cell surface. In this paper, we describe the purification to homogeneity of the midgut membrane-bound APN from adult pea aphids Acyrthosiphon pisum (Hemiptera: Aphididae) and the cloning of its corresponding cDNA. The data show that this APN prefers tripeptides, has broad amino-acid specificity, and is the most important mannose-specific lectin-binding site in midgut membranes.

**Results**

**Solubilization of A. pisum membrane-bound midgut APN**

About 98% of APN midgut activity [L-leucine-p-nitroanilide (LeupNA) as substrate] was found to be membrane-bound. The soluble fraction was eluted as a single peak from a Mono Q column, with a retention time similar to that of the solubilized enzyme (data not shown). The soluble enzyme was disregarded in further studies.

Acyrthosiphon pisum membrane-bound APN was well solubilized by all detergents tested (detergent concentration, % solubilization, % activity recovery): Chaps (32.7 mm, 90 ± 6%, 97 ± 8%), deoxycholate (7.3 mm, 91 ± 7%, 81 ± 9%), Triton X-100 (9.7 mm, 96 ± 5%, 116 ± 9%), Nonidet (9.7 mm, 91 ± 9%, 79 ± 8%), Control (8 ± 1% solubilization, 100 ± 8% recovery). As the best yield (solubilization) and recovery of activity were found with Triton X-100, this detergent was chosen for preparing the starting sample.

**Purification of A. pisum midgut APN**

The solubilized A. pisum APN was purified to homogeneity by one chromatographic step using a Mono Q column (Fig. 1A). From starting material
consisting of 300 guts, with total activity 2.2 U and 343 μg protein, it was possible to recover 28 μg purified APN with specific activity 40.3 U/mg \(^{-1}\). The final yield was \(\approx 50\%\), with a purification factor of 6.4. SDS/PAGE of purified APN resulted in a single 150-kDa protein band (Fig. 1B). The enzyme was found in the midgut as a major protein band and was preferentially solubilized by Triton X-100 (Fig. 1B, lane 2).

SDS/PAGE of proteins in fractions eluted from a gel-filtration column showed a correspondence between eluted activity and band intensity in stained gels (not shown), indicating homogeneity of the purified enzyme. The molecular mass calculated from gel filtration was 200 ± 30 kDa, a little higher than that obtained from SDS/PAGE.

In addition, APN can be purified using a single chromatographic step in ConA–Br-Sepharose (data not shown). The purified protein had the same mobility on SDS/PAGE and the same internal peptide sequence (see below) as APN purified on a Mono Q column.

**Properties of the purified APN from *A. pisum***

*Acrhythosiphon pisum* APN is a glycoprotein (Fig. 1C) and seems to be the major and/or most glycosylated protein from aphid midgut extracts (Fig. 1C, lane 4). It binds specifically to the lectin (*Galanthus nivalis* agglutinin) that recognizes a mannose moiety (Fig. 1C, lane 6). This agrees with the APN pattern of elution from ConA–Br-Sepharose columns (see above).

The APN purified from *A. pisum* had a pH optimum of 7.0 ± 0.5 (Fig. 2A) when assayed with LeupNA as substrate. Isoelectric focusing gave a single peak of pI 8.4 ± 0.2 (Fig. 2B), and density-gradient ultracentrifugation produced a single peak of molecular mass 130 ± 20 kDa (Fig. 2C).

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**Fig. 2.** Properties of purified midgut APN from *A. pisum*. (A) Effect of pH on enzyme activity (optimal pH 7.0 ± 0.5). Buffers used: 100 mM sodium phosphate buffer (pH 5–7) and 100 mM Tris/HCl buffer (pH 7–9.5). (B) Isoelectric focusing (pI 8.4 ± 0.2). (C) Density-gradient centrifugation. Hb, Haemoglobin; Ct, catalase. Molecular mass was calculated as 130 kDa. (D) Arrhenius plot. Activation energy was determined as \(E_a = 42.2\) kJ-mol\(^{-1}\).
The thermodynamic parameters of activation for *A. pisum* APN (Fig. 1D) were calculated by Arrhenius plot (plot of $k_{\text{cat}}$ against $1/T$). From the slope of the line, the activation energy ($E_a$) was determined to be 42.2 kJ mol$^{-1}$. Other thermodynamic parameters of activation were calculated using the relations of the transition state theory [23]. Thus, $AS^\ddagger$, $\Delta G^\ddagger$ and $\Delta H^\ddagger$ at 25 °C were estimated to be $-65.1 \text{ J mol}^{-1}\text{K}^{-1}$ ($-15.5 \text{ cal mol}^{-1}\text{K}^{-1}$), 59.0 kJ mol$^{-1}$ (14 kcal mol$^{-1}$) and 39.7 kJ mol$^{-1}$ (9.5 kcal mol$^{-1}$), respectively.

Purified APN (Mono Q column) was submitted to MS sequencing. First, it was treated with trypsin. The digested protein was separated by Q-ToF, and two of the resulting peptides were submitted to MS sequencing. The resulting sequences were (a) MDLLAIPDFR, (b) AGAMENWGMNTYK, and (c) NDSKITIYTYK. The same sequence as produced by peptide number 2 could be recovered from the protein purified by ConA–Br-Sepharose, together with a series of more than 19 mass hits covering the entire sequence (25 p.p.m. precision cut-off), including matching oxidized methionines. A Mowse score of 3.82E + 9 identified the purified and cloned sequences unambiguously (see below).

**Kinetic parameters of *A. pisum* APN**

The purified aphid APN showed a broad specificity towards N-terminal aminoacyl residues, although it was unable to hydrolyze L-aspartic acid β-(β-naphthylamide) (Table 1). The preferred substrates (higher $k_{\text{cat}}/K_m$ values) were those bearing leucine or methionine at the N-terminus, and the least preferable those presenting a proline at the N-terminus (Table 1). There was a slight preference for tripeptides (Table 1), as judged by a comparison of $k_{\text{cat}}/K_m$ values for peptides of the Leu-(Gly)$_n$ series which differ only in the number of Gly residues (Table 1).

Leucine hydroxamate is a simple intersecting linear competitive inhibitor of APN (Fig. 3), with $K_i = 5 \pm 1 \mu M$; the same is true for arginine hydroxamate ($K_i = 34 \pm 7 \mu M$). $K_i$ values for aminocyl hydroxamates depend on the hydroxamate used, not on the substrate used (l-leucine-β-naphthylamide or l-arginine-β-naphthylamide) (not shown). This indicates that l-leucine-β-naphthylamide and l-arginine-β-naphthylamide are hydrolyzed at the same active site.

*Acrithosiphon pism* APN inhibition by amastatin and bestatin are rapidly reversible by dilution (not shown), as observed with microsomal aminopeptidase [30]. Their pattern of inhibition is an intersecting, competitive, linear type, with $K_i = 1.8 \mu M$ for bestatin and $K_i = 0.6 \mu M$ for amastatin (not shown).

Inactivation of *A. pisum* APN by EDTA follows pseudo-first-order kinetics with $k_{\text{obs}} = 0.14 \text{ m}^{-1}\text{s}^{-1}$, which is virtually completely suppressed by the competitive inhibitor arginine hydroxamate at a concentr-

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**Table 1.** Kinetic parameters for purified APN from *A. pisum*. Relative values of $k_{\text{cat}}/K_m$ were calculated using LeupNA as reference for synthetic substrate. For peptide substrates, PheGlyGlyPhe was used as reference. The values were determined at least twice by 10 independent determinations with different substrate concentrations. SEM values were calculated by fitting data by a weighted linear regression using the software SigmaPlot®. AlaβNA, L-alanine-β-naphthylamide; AlapNA, L-alanine-$p$-nitroanilide; ArgpNA, L-arginine-$p$-nitroanilide; MetpNA, L-methionine-β-naphthylamide; MetpβNA, L-methionine-$p$-nitroanilide; ProβNA, L-proline-β-naphthylamide.

| Substrate         | $K_m$ (mM) | $k_{\text{cat}}$ (s$^{-1}$) | $k_{\text{cat}}/K_m$ (m$^{-1}$s$^{-1}$) | $k_{\text{cat}}/K_m$ (relative) |
|-------------------|------------|-----------------------------|----------------------------------------|---------------------------------|
| LeupNA            | 0.057 ± 0.007 | 119 ± 15                    | 2090 ± 360                             | 100                              |
| MetpNA            | 0.026 ± 0.008 | 45 ± 3                      | 1720 ± 540                             | 80.2                             |
| ArgpNA            | 0.19 ± 0.02  | 158 ± 31                    | 832 ± 180                              | 40.5                             |
| AlapNA            | 1.1 ± 0.04   | 222 ± 4                     | 202 ± 9                                | 9.9                              |
| LeuβNA            | 0.038 ± 0.003 | 105 ± 2                     | 2760 ± 220                             | 133                              |
| ArgβNA            | 0.058 ± 0.06  | 72 ± 2                      | 1220 ± 130                             | 59.0                             |
| MetβNA            | 0.043 ± 0.005 | 28 ± 1                      | 642 ± 76                               | 30.9                             |
| AlaβNA            | 0.28 ± 0.04  | 67 ± 2                      | 240 ± 8                                | 11.4                             |
| ProβNA            | 1.6 ± 0.2    | 6.8 ± 0.2                   | 4.3 ± 0.5                              | 0.2                              |
| AspβNA            | –           | < 0.19                      |                                        |                                  |
| LeuGly            | 0.26 ± 0.08  | 5.6 ± 0.4                   | 22 ± 7                                 | 11.1                             |
| LeuGlyGly         | 0.28 ± 0.06  | 26 ± 2                     | 92 ± 20                                | 48.9                             |
| LeuGlyGlyGly      | 0.9 ± 0.1    | 16 ± 1                     | 19 ± 2                                 | 10                               |
| LeuGlyGlyGlyGly   | 1.1 ± 0.1    | 17 ± 1                     | 15 ± 1                                 | 7.8                              |
| LeuLeuLeu         | 0.070 ± 0.02 | 6.4 ± 0.3                   | 91 ± 23                                | 47.8                             |
| PheGly            | 0.60 ± 0.08  | 13 ± 4                     | 22 ± 8                                 | 11.1                             |
| PheGlyGly         | 1.1 ± 0.3    | 23 ± 2                     | 22 ± 6                                 | 11.1                             |
| PheGlyPheGly      | 0.71 ± 0.07  | 92 ± 2                     | 130 ± 13                               | 68.8                             |
| PheGlyGlyPhe      | 0.27 ± 0.03  | 50 ± 2                     | 188 ± 21                               | 100                              |
tion corresponding to 25-fold its $K_i$ value (Fig. 4). The reaction order with respect to EDTA was 0.44. As EDTA has two metal-binding sites, the data support the conclusion that removal of only one metal ion is sufficient to inactivate the enzyme. In agreement with this, the partially EDTA-inactivated enzyme has the same $K_m$ and pH optimum as native aminopeptidase (not shown).

**Carbohydrate and lectin interactions with A. pisum APN**

The enzyme strongly interacts with lectins that bind mannose-like *Galanthus nivalis* agglutinin and ConA, as observed in blotting assays (Fig. 1C) and in the purification steps (see above). The interaction with lectins was evaluated by density-gradient ultracentrifugation (Fig. 5). After 30 min of preincubation of APN with the lectins, wheat germ agglutinin (WGA), which binds to sialic acid and N-acetylglucosamine moieties, or ConA, which binds to glucose and mannose moieties, the samples were submitted to density-gradient ultracentrifugation. APN with WGA results in a single peak (Fig. 5A), as observed in Fig. 2C, which corresponds to the APN without bound lectin (Fig. 5A). Mixing ConA with APN resulted in all the activity being at the bottom of the tube, meaning a molecular mass higher than 400 kDa (Fig. 5B), resulting from lectin binding and agglutination. When a competitive monosaccharide ($\alpha$-methyl mannose) was added to the incubation mixture of ConA with APN at a concentration of...
500 mM, a peak of intermediate molecular mass was observed (Fig. 5C), corresponding to the partially aggregated form of APN with ConA (the molecular mass of ConA is 73 kDa under the density-gradient conditions).

The activity recovered from the density gradients was similar with and without lectins (data not shown). The kinetics parameters of APN (kcat and Km) associated with ConA were unaffected, indicating that the catalytic site and the mannosylated site(s) are quite far apart on the enzyme molecule.

Sequence coding the A. pisum APN

Full-length cDNA was obtained for A. pisum APN with 3234 bp (GenBank accession number DQ440823). This sequence codes for a protein of 973 amino acids with 3234 bp (GenBank accession number DQ440823). This full-length cDNA contains a short 5¢-UTR from 1 to 132 bp and 3¢-UTR from 3055 to 3234 bp.

The protein encoded by this cDNA contains the three peptide sequences obtained from the purified enzyme, showing identity between the purified enzyme and the cDNA sequence, as well as MS peaks with high Mowse score (see above) which unambiguously identified the cloned sequence.

The coding protein has high similarity to other aminopeptidases. It possesses the domain HEXXH + G, characteristic of gluzincins, and the domain GAMEN, found in many of these enzymes (Fig. 6). Putative N-glucosylation sites (predicted at http://www.cbs.dtu.dk/services/NetNGlyc/) are assigned in Fig. 6, as well as the GPI anchor site in its C-terminal domain, as predicted by the DGPI software [32]. The presence of the signal peptide and GPI anchor signal are consistent with the known characteristics of insect APNs. O-glucosylation sites were predicted to be present in the region (close to the C-terminus) of the APN using the NetOGlyc 3.1 server [33]. clustalw sequence alignment with other insect aminopeptidases (Fig. 7) showed that A. pisum APN has a weak similarity to class 2 aminopeptidases from Lepidoptera [20].

Discussion

Occurrence, properties and sequence of A. pisum APN

Acyrthosiphon pisum has a membrane-bound and a soluble aminopeptidase corresponding to 98% and 2% of the midgut aminopeptidase activity, respectively, when LeupNA is used as substrate. There is a single molecular species of A. pisum membrane-bound aminopeptidase in this tissue, as judged by Mono Q chromatography after solubilization of almost 100% of its activity.

The membrane-bound APN from A. pisum was purified to homogeneity, with a yield of 51.9% and specific activity of 40.3 U mg⁻¹. Taking into account that the specific activity of the homogenized mid gut of this insect is 6.3 U mg⁻¹ per animal and that each midgut has 19 µg protein [27], it is possible to calculate that there are about 3 µg APN per midgut and that APN amounts to 15.6% of midgut protein. This is confirmed by SDS/PAGE of the midgut homogenate, where it is possible to recognize APN as a major protein in the preparation.

APN is a glycosylated protein of molecular mass 130 kDa (density-gradient centrifugation) and pI 8.4. Molecular masses determined by SDS/PAGE (150 kDa) or gel filtration (200 kDa) are probably artifacts. The molecular mass of the unglycosylated protein is 109 kDa and pI 5.3 (predicted from the amino-acid sequence). The data led us to conclude that ≈16% of the molecular mass of APN is carbohydrate.

Immunoblot for identification of glycosylated proteins recognized APN as the most abundant glycoprotein in the midgut and thus as an important target for lectins. Taking into account that more than one lectin can bind a single APN molecule (Fig. 5), and the abundance of APN in microvillar membrane, this enzyme is potentially the most important lectin-binding site in aphid midgut. The calculated amount of aminopeptidase in microvillar membrane may explain the capacity of each aphid to feed on a diet containing lectins amounting to as much as 1 µg of ConA in 48 h [29]. Also immunohistochemical observations of lectin binding on the midgut demonstrated that the stomach (ventriculus I) cell membranes are the primary target for ConA, followed by the intestine (remaining midgut chambers) cell membranes [29]. Activity measurements found APN along the midgut, and immunolocalization with APN antibodies showed that APN is associated with a specialized plasma membrane associated with the apical lamellae. These consist of a complex network of lamellae, linked one to another by trabeculae to resist the osmotic pressure caused by high-sugar phloem-sap ingestion [28]. The apical lamellae replace the regularly arranged microvilli observed in most midgut cells. APN localization data are in agreement with the lectin-binding site found by Sauvion et al. [29]. However, as presented here, APN activity is not affected by glucose/mannose-binding lectin.
Fig. 6. cDNA coding sequence of *A. pisum* APN and its deduced sequence (GenBank accession number DQ440823). The predicted signal peptide is underlined, and the C-terminal GPI cleavage signal sequence is dotted underlined. The characteristic zinc binding/gluzincin motif, 
\[ \text{HEXXH} + E \], and the gluzincin aminopeptidase motif, GAMEN, are highlighted in a bold/gray box. Peptides identified by MS analysis are double underlined. Boxed residues correspond to the MS sequenced peptides. Putative N-glycosylated asparagine residues are dark-shaded using the NetNGlyc 1.0 server (http://www.cbs.dtu.dk/services/NetNGlyc/), and putative O-glycosylated threonines residues identified by the NetOGlyc 3.1 server (http://www.cbs.dtu.dk/services/NetOGlyc/) are also shaded.

Aphid midgut aminopeptidase

P. T. Cristofoletti et al.

5580
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Acyrthosiphon pisum APN has a broad specificity towards the N-terminal amino-acid residues of peptides, but it does not hydrolyze acidic aminoacyl-peptides, and, although it is by no means proven, it appears to prefer peptides longer than dipeptides. Thus, *A. pisum* APN resembles the vertebrate enzyme (EC 3.4.11.2) [34] and the following insect midgut enzymes: microvillar membrane APN from *Tenebrio molitor* [22], soluble *Tineola bisselliella* (Lepidoptera) aminopeptidase [35,36], soluble *Attagenus megatoma* (Coleoptera) aminopeptidase [37], soluble and microvillar *R. americana* (Diptera) aminopeptidases [25,26], and membrane-bound *Spodoptera littoralis* (Lepidoptera) aminopeptidase [38]. Another resemblance between *A. pisum* APN and the vertebrate enzyme are both inhibited by bestatin and amastatin, which in both cases is rapidly reversible [30]. It should be noted, however, that *A. pisum* APN has a $K_m$ value for peptides much smaller than those for *T. molitor* APN [22].

Substrates with different N-terminal amino-acid residues are hydrolyzed at the same site of *A. pisum* APN, as hydroxamate $K_i$ values do not depend on the aminoacyl-$b$-naphthylamide used as substrate. Substrates with a bulky aminoacyl residue in position P1 (numbering of Schechter & Berger [39]) are better substrates for APN. The same is true for the P2¢ position (compare Leu-Gly-Gly with Leu-Leu-Leu in Table 1), but not for the P1¢ position (compare aminoacyl-naphthylamide with aminoacyl-$p$-nitroanilide). This suggests that the subsites S1, S2, and probably S3¢ of the enzyme are pockets able to bind bulky aminoacyl residues, and this hypothesis agrees with the fact that amastatin is a better inhibitor of *A. pisum* APN than bestatin. Bestatin has bulky residues putatively able to interact with S1¢ and S2¢ of the enzyme (see above), and amastatin with S1, S1¢ and S2¢ [40].

*Acyrthosiphon pisum* APN is the first insect digestive aminopeptidase that does not belong to the order Lepidoptera to be fully characterized and sequenced.
A rapid survey of the *A. pisum* databank (http://urgi.inforubigen.fr/cgi-bin/annotation_form.pl?organism = apism) allows the identification of more than 25 contigs with some relation to the word ‘aminopeptidase’, and similarly, a high number of ‘aminopeptidases’ are encoded in the *Drosophila* genome (http://flybase.org). A total of 29 *A. pisum* expressed sequence tags (ESTs) have full complementarities with the cloned APN in almost 60 000 ESTs. From these 29 ESTs, 22 belong to libraries from the digestive tract, and seven from libraries from whole insect (none from other tissue libraries), meaning that this aminopeptidase is potentially very specific to the aphid midgut.

The APN sequence has all identified residues essential for zinc binding and catalysis. In the sequence, it was easy to recognize the signal peptide, several potential glycosylation sites, as well as a GPI anchor at its C-terminus. This anchor is possibly an adaptation to a phloem-based diet, avoiding excretion of the enzyme into the honeydew, as the phylogenetically related Hemipteran *Dysdercus peruvianus* has a soluble aminopeptidase, in spite of the fact that, in this case, the enzyme is trapped between the microvillar and perimicrovillar membranes [27,28].

**Function of *A. pisum* APN and lectin toxicity**

The role of the microvillar aminopeptidase is postulated to be hydrolysis of oligopeptides formed by the action of luminal proteinases [1,2,22]. In aphids, a cathepsin L was found to be partially associated with modified perimicrovillar membranes and is possibly involved in degradation of toxic proteins found in the phloem sap [28,41]. APN is certainly responsible for the final digestion of peptides generated by cathepsin L. Another possibility is that APN is somehow associated with putative amino-acid-binding sites at the plasma membranes associated with the apical lamellae (modified perimicrovillar membranes). These are thought to increase the amino-acid concentration (usually low in the aphid diet) [28], thus facilitating absorption. APN may also be directly linked to absorptive sites in apical lamellae, as has been suggested for Lepidoptera [42]. Finally, APN may serve as the primary digestive enzyme responsible for the assimilation of the phloem sap small peptide fraction, chemical components largely unexplored at the moment.

As *B. thuringiensis* is not effective in aphid control, lectins have been used as insecticidal agents against aphids [43]. The soluble protein, ferritin, is the snowdrop lectin-binding protein in the planthopper *Nilaparvata lugens* [44]. The authors postulated that alteration of iron metabolism might be related to its lectin toxicity. Although ferritin was not the most abundant protein in midgut preparations, this protein was the most specifically recognized in *N. lugens*. However, none of the 2D PAGE protein spots observed in pea aphid homogenates as binding to ConBr was identified as corresponding to ferritin (F. A. Mendonca de Sousa & Y. Rabbé, unpublished), although the ferritin gene is largely transcribed in *A. pisum* midguts [45]. It is possible that the mechanism of toxicity found in planthopper is different from that found in aphids.

In *A. pisum*, the lectin, ConA, is a potent toxin affecting survival and growth, but WGA is relatively ineffective [46]. These data agree with the fact that aphids do not possess a peritrophic membrane [28]. Consequently, this toxicity must result from lectin binding to target proteins in the apical membranes from the midgut, although not related to the inhibition of APN activity. One explanation of this effect is a decrease in amino-acid absorption caused by ConA binding to APN, with deleterious effects on the putative associated proteins thought to bind to amino acids (see above). Indeed, ConA-intoxicated aphids have been shown to display altered hemolymph free amino-acid profiles and modified excretion of asparagine in their honeydew [47]. It is still possible that a reduction in membrane protein lateral mobility or its resistance to phloem osmotic pressure is the major cause of lectin toxicity to aphids. These possibilities need to be evaluated.

**Experimental procedures**

**Animals**

*Acrithosiphon pisum* Harris aphids (Hemiptera: Aphididae), clone Ap-LL01, were maintained in the laboratory on broad bean seedlings (*Vicia faba*) in ventilated plexiglass cages (21 °C; 70% relative humidity; 16 h light/8 h darkness). For the experiments, a limited number of mass-reared adults were allowed to lay eggs for 24 h on young *Vicia* plants, and the resulting apterous insects were used as 9-day-old adults.

**Chemicals**

Buffer salts, detergents, molecular-mass markers, protein inhibitors, and most substrates were purchased from Sigma-Aldrich (St Louis, MO, USA). Glycoprotein detection kits came from Boehringer-Mannheim (Mannheim, Germany). The peptides Leu-Gly-Gly-Gly and Leu-Gly-Gly-Gly-Gly-Gly were gifts from Dr L. Juliano (Unifesp, São Paulo, Brazil).
Preparation of samples

Adult apterous aphids were immobilized on a flat surface, using adhesive tape, and their guts were removed under a stereomicroscope in Yeager’s physiological solution [48]. The midguts were separated and homogenized in double-distilled water with the aid of a Potter-Elvehjem homogenizer. The homogenates were labeled crude homogenate and stored. Crude homogenates were used to assay APN or were centrifuged at 100 000 g for 1 h at 4 °C, resulting in a supernatant (labeled midgut soluble fraction) and a pellet (midgut cell membranes). Washed midgut cell membranes were prepared by dispersing the midgut cell membranes in water, followed by three freezing and thawing cycles, and re-centrifugation at 100 000 g for 1 h at 4 °C. All centrifugations were performed on a Hitachi Ultracentrifuge model Himac 70P-72 with an RPS 40T rotor.

Protein determination and enzymatic assays

Protein was determined as described by Bradford [49] using ovalbumin as standard. When samples contained detergent, protein was determined by the method of Smith et al. [50], as modified by Morton & Evans [51], using BSA as standard.

Routine assays of APN were performed using 1 mM LeupNA as substrate (initially solubilized in dimethyl sulfoxide) in 100 mM Tris/HCl buffer, pH 7.0, at 30 °C. Unless otherwise specified, the same conditions were used for all other substrates. Naphthylamine liberated from aminoacyl-β-naphthylamides, nitroaniline from aminoacyl-p-nitroanilides, and phenylalanine and leucine from the different peptides were determined spectrophotometrically by the methods of Hopsu et al. [52], Erlanger et al. [53] and Nicholson & Kim [54], respectively. In each determination, incubations were continued for at least four different periods of time, and the initial rates were calculated. All assays were performed so that the measured activity was proportional to protein and incubation time. Controls without enzyme or without substrate were included. One enzyme unit (U) is defined as the amount that hydrolyzes 1 μmol substrate min⁻¹, at 30 °C.

Solubilization of APN by detergents

In order to evaluate the solubilizing efficiency of detergents, samples of 200 μL midgut homogenate at a concentration of 13 guts per mL (which contains ≈50 μg protein) of midgut cell membranes were suspended in 10 mM Hepes buffer, pH 7.4, in the presence and absence of several detergents. After 17 h at 4 °C with shaking, the suspensions were centrifuged at 100 000 g for 1 h at 4 °C, and supernatants were assayed for APN. APN activity was determined in the resulting supernatants and referred to the original preparation of cell membranes (as percentage solubilization). Recovery is the percentage of the sum of solubilized plus nonsolubilized activity referred to the original preparation of cell membranes. Data are mean ± SEM calculated from determinations carried out in three different preparations.

For routine solubilization of APN, midgut cell membranes were suspended in 10 mM Hepes buffer, pH 7.4, containing 10 mM Triton X-100. After 1 h at 4 °C, the suspension was centrifuged at 25 000 g for 30 min at 4 °C, and the supernatant used as a source of enzyme.

Purification of detergent-solubilized APN

Cell membranes corresponding to ≈300 A. pisum midguts (wet weight ≈6 mg) were solubilized with Triton X-100 as described above, and applied to a Mono Q HR 5/5 column (0.5 cm internal diameter × 5 cm) equilibrated with 20 mM Tris/HCl buffer, pH 7.0, containing 0.1% Triton X-100 in an FPLC system. Controls showed that protease inhibitors are not necessary. Elution was carried out with a gradient of 0–0.6 M NaCl in the same buffer. The flux was 1.0 mL min⁻¹, and fractions of 0.4 mL were collected. Fractions showing activity with LeupNA were pooled, and purification was checked by SDS/PAGE.

Alternatively, the APN was purified on a 3-mL ConA–Br-Sepharose column (6 × 30 mm). The column was washed with 15 mL 20 mM acetate buffer, pH 4.2, containing 0.5 M NaCl, then with 15 mL 100 mM citrate/phosphate buffer, pH 6.0, containing 2 mM CaCl₂, before being equilibrated with 15 mL 20 mM Tris/HCl buffer, pH 7.0, with 0.1% Triton X-100. The solubilized samples were applied to the column and eluted with 20 mM Tris/HCl buffer, pH 7.0, containing 0.1% Triton X-100 and 0.5 M α-methyl mannoside. Fractions of 1 mL were collected at a flow rate of 1 mL min⁻¹.

SDS/PAGE

Electrophoresis of A. pisum samples in denaturing conditions (SDS/PAGE) was carried out on 7.5% (w/v) polyacrylamide gels containing 0.1% (w/v) SDS, on a discontinuous pH system [55], using Mini Protean II cells (Bio-Rad, Hercules, CA, USA). Samples were lyophilized and suspended in sample buffer containing 60 mM Tris/HCl buffer, pH 6.8, 2.0% (w/v) SDS, 5% (v/v) 2-mercaptoethanol, 10% glycerol and 0.2% (w/v) bromophenol blue and heated for 3 min at 95 °C in a water bath before being loaded on to the gels. Electrophoresis was carried out at 200 V until the tracking dye reached the bottom of the gel. The gel was then silver-stained [56] or stained using 0.1% (w/v) Coomassie Blue R in 10% acetic acid/40% methanol for 30 min. In the last case, destaining was achieved with several washes in a solution containing 40% methanol and 10% acetic acid.
Aphid midgut aminopeptidase

P. T. Cristofoletti et al.

Isolelectric focusing

Isolelectric focusing was performed as described by Terra et al. [57], in columns of 7.5% polyacrylamide gel containing 10% ampholytes pH 3–10 (Pharmalyte 3–10, Pharmacia, Uppsala, Sweden). Samples were applied after polymerization and prefocusing (30 min at 31 V cm⁻¹) on the top of the alkaline side of the gels. For samples in detergent, 0.1% Triton X-100 was added to the gels and fractionation buffer. Recoveries of activities applied to the gels were 80–100%.

Density-gradient centrifugation

Samples of purified APN (100 μL) containing ≈1 μg purified protein with or without 100 μg lectins were layered on the top of 10-mL glycerol gradients (10–30%, w/v) made up in 100 mM Tris/HCl buffer, pH 7.0, in the presence or absence of 500 mM α-methyl mannoside. Centrifugation and collection of fractions were performed as described previously [58]. The molecular mass of APN was calculated by the method of Martin & Ames [59], using the sedimentation rates of hemoglobin (64.5 kDa) and bovine liver catalase (232 kDa) as reference standards. Activity recovery was 80–100%.

Determination of molecular mass by gel filtration

Samples of 200 μL, containing 2–4 μg purified protein, were applied to a gel-filtration column in an FPLC system (Pharmacia-LKB Biotechnology, Uppsala, Sweden) by using a Superose HR 10/30 column (1.0 cm internal diameter × 30 cm) equilibrated and eluted in 100 mM Tris/HCl buffer, pH 7.0, containing 0.1% Triton X-100. Fractions of 0.4 mL were collected at a flow rate of 0.5 mL min⁻¹. Molecular masses were calculated using the following proteins as standards: aprotinin (6.5 kDa), cytochrome c (12.4 kDa), ovalbumin (45 kDa), BSA (65 kDa), and β-amylase (200 kDa). Recoveries were 80–100%.

Detection of carbohydrates in purified APN

Protein samples were blotted on to nitrocellulose sheets after SDS/PAGE [60]. Detection of carbohydrates was performed with the DIG Glycan Detection kit, and identification of carbohydrate moieties was accomplished with lectins by using the DIG Glycan Differentiation kit. The procedures followed the supplier’s instructions (Boehringer Mannheim).

Kinetic studies

The effect of substrate concentration on the activity of purified APN was determined using at least 10 different substrate concentrations. Kinetics were determined by the method of Martin & Ames [59], using the sedimentation rates of hemoglobin (64.5 kDa) and bovine liver catalase (232 kDa) as reference standards. Activity recovery was 80–100%.

EDTA inactivation

Purified APN was incubated with EDTA (1–50 mM) for different times at 40 °C in 100 mM citrate/phosphate buffer pH 6. The EDTA-inactivation reactions were stopped by 100-fold dilution of reaction mixtures with 100 mM Tris/HCl buffer (pH 7.0)/0.1% Triton X-100. The remaining activity was measured using LeuPNA as substrate, under the conditions described above. Protection against inactivation by EDTA was investigated with aminoacyl hydroxamates, which are simple linear competitive inhibitors of A. pisum APN. The reaction order was determined by incubation of APN with different concentrations of EDTA.

Microsequencing of purified APN

Purified APN was electroblotted onto poly(vinylidene difluoride) membranes after SDS/PAGE [62]. The membranes were stained for protein using 0.1% Coomassie Blue R-250 in a 50% (v/v) methanol, and were destained with 50% methanol. Dried poly(vinylidene difluoride) membranes were submitted to tryptic digestion, and the resulting peptides (two peptides) were submitted to MS sequencing at the sequencing facility of the Pasteur Institute (Paris).

Alternatively, in-gel digestion was performed for protein identification: spots were excised from preparative gels using pipette tips. The spots were washed with 100 μL 25 mM NH₄HCO₃ for 30 min, twice destained for 30 min with 100 μL 25 mM NH₄HCO₃/acetoni-trile (v/v), and dehydrated in acetonitrile. Gel spots were completely dried using a vacuum centrifuge before trypsin digestion. The dried gel volume was evaluated, and 3 vol. trypsin (V5111; Promega, Madison, WI, USA; 10 ng μL⁻¹ in 25 mM NH₄HCO₃) was added. Digestion was performed at 37 °C over 5 h. The gel pieces were centrifuged, and 8–12 μL acetonitrile (depending on gel volume) was added to extracted peptides. The mixture was sonicated for 5 min and centrifuged. For MALDI-TOF MS analysis, 1 μL supernatant was loaded directly on to the MALDI target. The matrix solution (5 mg mL⁻¹ α-cyano-4-hydroxycinnamic acid in 50% acetonitrile containing 0.1% trifluoroacetic acid) was added immediately and allowed to dry at room temperature. A Voyager DE-Pro model MALDI-TOF mass spectrometer (Perseptive BioSystems, Farmingham, MA, USA) was used in positive-ion reflector mode for peptide mass fingerprinting. External calibration was performed with a
standard peptide solution (Proteomix; LaserBio Laboratories, Sophia-Antipolis, France). Internal calibration was performed using peptides resulting from autodigestion of porcine trypsin. Monoisotopic peptide masses were assigned and used from NCBI database searches (plus A. pism APN sequence) with the ‘MS-FIT’ software.

Cloning of APN from A. pism

Total RNA was extracted from midgut epithelium of A. pism with Trizol following the instructions of the manufacturer, Invitrogen, which are based on those of Chomczynski & Sacchi [63]. mRNA was purified with Qiagen mRNA purification kit, and a cDNA library was constructed with the kit Smart (Clontech, Mountain View, CA, USA), following the instructions of the manufacturer. A partial sequence of a cDNA coding for APN was amplified using degenerated primers for the APN consensus sequence that contains the peptide AGAMENWGM identified in MS analysis (primers APN-U538: 5′-TTYCCITGY TITYGAYGARCC-3′, based on peptide ‘TGLYRSS’; APN-L1051: 5′-RTTICRAACCACWKRTG-5′, based on peptide ‘THQWFGN’). PCR was performed using Taq DNA polymerase (Invitrogen) using the standard method. The PCR product was cloned in pGEM-T Easy Vector (Promega), sequenced, and the identified fragment sequence had high identity with that of APN.

This sequenced cDNA fragment contains the two identified peptides sequenced from purified protein, and was BLASTed (BLASTN) against the A. pism ESTs deposited at NCBI [45]. The recovered fragments were clustered and BLASTed again against the A. pism ESTs until the N-terminus was completed with signal peptide and 5′-UTR sequence. Putative 3′ reads were recovered from A. pism ESTs using BLAST (BLASTN) with the best full-length BLAST hit as driver (Apis mellifera, accession number XP 366261). Also, ESTs corresponding to A. pism APN can be recovered along the sequence using MS peaks with Mascot engine (http://www.matrixscience.com/). Reads were clustered into two contigs covering the 3′ region and the 5′ region. The final gap between these contigs was recovered using PC with forward primers 5′-GGCATGGTGAG GACTATGTTGGCCG-3′ combined with reverse primer 5′-GCCATGCAGCGTCGTATGGCG-3′, and the complete sequence was obtained and deposited at GenBank with the accession number DQ440823.

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