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Neuropeptide Degradation Produces Functional Inactivation in the Crustacean Nervous System

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The pentapeptide proctolin (Proct.; Arg-Tyr-Leu-Pro-Thr) is a modulatory transmitter found throughout the crustacean nervous system. No information is available in this system, however, as to how the actions of this peptide are terminated. To study this issue in the crab Cancer borealis, we incubated exogenous proctolin $(10^{-5}$ M) with either the thoracic ganglion (TG) or with conditioned saline (CS) that had been preincubated with the TG. We removed aliquots at standard time points for analysis by reverse-phase highperformance liquid chromatography (HPLC). We found that over time the proctolin peak became progresstvely smaller, while three novel peaks appeared and increased in size. Comigration experiments using HPLC indicated that the major novel peak was Proct. [2-51 (Tyr-Leu-Pro-Thr), while one of the two minor peaks was Proct. [3-51 (Leu-Pro-Thr). The other minor peak appeared to be Proct. [1-2] (Arg-Tyr), based on similar HPLC retention time to synthetic Proct. [1-2]. The reduction in the proctolin peak and the increase in the Proct. [2-51 peak was prevented by co-incubation of proctolin with any one of several aminopeptidase inhibitors (10 -4 M). Proct. [l-2] and Proct. [3-51 appeared to result from a diaminopeptidase-mediated cleavage of proctolin. We tested whether N-terminal cleavage functionally inactivated proctolin by coapplying proctolin (10^{-8} M) and individual aminopeptidase inhibitors $(10⁻⁵$ M) to the isolated stomatogastric ganglion (STG). We found that these inhibitors significantly enhanced the proctolin excitation of the pyloric rhythm. Furthermore, application of synthetic Proct. [2-51 to the STG had no effect unless high concentrations ($>$ 10⁻⁶ M) were used, and neither Proct. $[1-2]$ nor Proct. $[3-5]$ (10⁻⁴ M) influenced the pyloric rhythm. Our results indicate that proctolin is enzymatically degraded and thereby biologically inactivated in the crab nervous system, primarily by extracellularly located aminopeptidase activity.

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[Key words: stomatogastric ganglion, neuromodulation, aminopeptidase, pyloric rhythm, proctolin, neuropeptide, inactivation, Cancer borealis]

Considerable information is available regarding the presence, synthesis, release, and postsynaptic effects of neuropeptide transmitters (White et al., 1985; Polak and Bloom, 1987; Sossin et al., 1989; DeLoof and Schoofs, 1990; Marder and Weimann, 1992). Less is known regarding the postrelease fate of these peptides, and how that fate influences synaptic transmission. Thus far, it has been shown that intact neuropeptides can diffuse relatively long distances from their release sites (Jan and Jan, 1982; Mayeri et al., 1985), or can be enzymatically cleaved in the extracellular space (McKelvy and Blumberg, 1986; Isaac, 1987, 1988; Turner, 1987; Molineaux et al., 1988; Rose et al., 1988; Squire et al., 1991).

In some systems, extracellular cleavage functionally inactivates the neuropeptide (Turner et al., 1985; Hall and Lloyd, 1990; Owens et al., 1992). This inactivation limits the duration and distance over which the peptide influences its targets (Sigvardt et al., 1986). Sometimes, however, enzymatic cleavage either activates or enhances the physiological effects of the peptide (Turner et al., 1985; Thorsett and Wyvratt, 1987; Owens et al., 1992). While some information is available regarding how neuropeptide inactivation influences peptidergic effects at both the cellular (Sigvardt et al., 1986; Owens et al., 1992) and the behavioral level (Turner et al., 1985; Sullivan et al., 1988; Hall and Lloyd, 1990), little is known about the role played by neuropeptide inactivation in sculpting peptidergic modulation of neural network and neural network activity.
One well-studied model system for neuropeptide actions on

rhythmically active networks is the crustacean stomatogastric nervous system (STNS; Harris-Wannals et al., 1992b; Marder
nervous system (STNS; Harris-Wannals et al., 1992b; Marder $\frac{1000 \text{ N}}{1000 \text{ N}}$ Weimann et al., 19920, Martin and Weimann, 1992; Weimann et al., 1993). One particularly well-characterized rhythmic motor pattern in this system is the pyloric rhythm, generated by a neural network in the stoma-
togastric ganglion (STG; Selverston and Moulins, 1987; Harriswarrick gangholi (31 Ci, 30 iversion and Mountis, 1997, 11 answer
Warrick et al., 1992a). The pyloric rhythm is influenced by many wallier chall, $1972a$. The pyrolle inviting immedie dy many different modulatory transmitters, including at least six different neuropeptides (Harris-Warrick et al., 1992b; Marder and Weimann, 1992). Among the best-characterized neuropeptides in the STNS is the pentapeptide proctolin (Proct.; Arg-Tyr-Leu-Pro-Thr; Marder et al., 1986), which is also found throughout the crustacean CNS (Schwarz et al., 1984; Siwicki and Bishop, 1986). Exogenously applied proctolin causes a dose-dependent excitation of the pyloric rhythm (Marder et al., 1986; Hooper

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and Marder, 1987; Nusbaum and Marder, 1989a,b; Golowasch and Marder, 1992). Within the STNS, proctolin is found in the STG neuropil, in several fibers entering the STG, and in somata in the neighboring commissural (CoGs) and esophageal (OG) ganglia. In the crab *Cancer borealis* there appear to be only three pairs of proctolin-containing neurons that innervate the STG (Coleman et al., 1992). This includes the paired modulatory proctolin neuron (MPN; Nusbaum and Marder, 1989a,b) and the paired modulatory commissural neuron 1 (MCNl; Christie et al., 1993). Both of these neurons excite the pyloric rhythm.

In this article we examine the fate of exogenously applied proctolin in the nervous system of the crab Cancer borealis. We have found, in biochemical experiments using the relatively large thoracic ganglion (TG), that proctolin is enzymatically degraded primarily by an aminopeptidase (AP) inhibitor-sensitive process. In complementary electrophysiological experiments, AP inhibitors enhanced the excitatory effects of proctolin on the pyloric rhythm, suggesting that AP activity terminates the proctolin effects in the STG.

Portions of this work were published previously in abstract form (Coleman et al., 1991).

Materials and Methods

Animals. Cancer borealis were obtained from Neptune Seafood & Lobster Co. (Boston, MA). Crabs were maintained in refrigerated (10–12°C), aerated aquaria with filtered, circulating artificial seawater. Data were obtained from 72 male crabs weighing between 300 and 800 gm.

Solutions. C. borealis physiological saline contained (mmol/liter) NaCl, $440. VCl$ 11: MeCl. $26. CoCl$ 12: Trigma base. 10: and maleic acid. 5.6 , $15.74.7.5$. Decotatin, and the AD inhibitors amastatin, bestating 5 (pH 7.4–7.5). Proctolin and the AP inhibitors amastatin, bestatin, epiamastatin, and puromycin were obtained from Sigma Chemical Co (St. Louis, MO). The AP inhibitors arphamenine B and actinonin were obtained from Peninsula Labs (Belmont, CA). Proct. [2-51 (Tyr-Leu-Pro-Thr) and Proct. [3-51 (Leu-Pro-Thr) were provided by Dr. James M. Weimann (Department of Biology, Brandeis University), and were synthesized using the procedure for peptide synthesis described in Wei $m_{\rm H}$ (1992). Procedure for peptide symmesis described in Wei s_1 is a synthesized by Research Genetics, Inc. (Huntsville, AL). All peptides and peptidase inhibitors were stored either as frozen or lyophilized and peptidase infinitions were stored entired as frozen or typerinzed
alianzate of stock solution (10+2 m or 10+2 m) that were diluted to the anyweis of stock solution (10 \pm M of 10 \pm M) that were unuted to the \pm working concentration in saline immediately before they were used.

Ganglionic incubations. Because a large source of neural tissue was necessary to cleave amounts of proctolin large enough to be detected by absorbance at 206 nm, we used the TG in these experiments. After each dissection, the sheathed TG was pinned to a Sylgard (Dow Corning)coated petri dish and preincubated for at least 30 min in 3 ml saline. Following several rinses, the TG was then incubated with either (1) saline, (2) saline plus Proct. or Proct. $[2-5]$ (Peptide; 10^{-5} M), or (3) saline plus peptide (10^{-5} M) and peptidase inhibitor (10^{-4} M). Each TG incubation was maintained for at least 2 hr (11–15°C) and 220 μ l aliquots were removed at $T = 1$, 60, and 120 min. At $T = 0$ min, the aliquot were removed at $t = 1$, ou, and 120 min. At $t = 0$ min, the anguot contained 2.2×10^{-7} mot of proctotin. Aliquots immediately received 25 μ l of HCl (1.0 m) to inactivate enzymatic activity and were then analyzed by reverse-phase high-performance liquid chromatography (HPLC; see below). FLC , see Delow). Include saling include the $\frac{1}{\sqrt{1-\frac{1}{n}}}\int_{-\infty}^{\infty}$

 α conditioned saline incubations. Incubations of $1\,\mathrm{G}$ to generate conditioned saline (CS) involved a 30-60 min preincubation with saline, which was replaced with fresh saline (3 ml) for a subsequent 2 hr incubation. This CS was then removed and incubated with the same test solutions described above for TG incubations. All incubations of CS aliquots were performed at 13°C. Each CS aliquot was a 700 μ l solution of CS (560 μ l) and peptide (10⁻⁴ M, 70 μ l) plus either peptidase inhibitor $(10^{-3}$ M) or saline (70 μ l). Final concentrations in each aliquot were 10⁻⁵ M peptide and 10^{-4} M inhibitor (when present). From each aliquot, HPLC samples (220 μ) were removed at $T = 1, 60$, and 120 min, and immediately inactivated by adding HCl (1.0 μ , 25 μ l).

HPLC. All HPLC analyses were performed using a Rabbit HPX Dual Pump Delivery System with a Dynamic Mixer (Rainin) controlled by a Macintosh SE computer using Dynamax HPLC METHOD MANAGER (version 1.3, Rainin). Sample volumes (200 μ l) contained 180 μ l of CS

solution plus 20 μ l of HCI. These samples were analyzed with a Vydac C_{18} HPLC analytical column (0.46 \times 25 cm, 300 Å pore size) to which was fitted a Vydac C_{18} guard column and a precolumn filter. Samples were analyzed with 0.1% trifluoroacetic acid (TFA, pH 2; Aldrich) using a linear gradient of acetonitrile (CH,CN) of 0.2-39% in 13 min (3%/ min), at a flow rate of 1.0 ml/min. Peaks were detected at 206 nm using METHOD MANAGER software. Control injections were made of synthetic a Knauer variable wavelength UV detector that was controlled by the peptide and peptidase inhibitors to verify retention time and to approximate the amount of material producing each peak of interest. Detection threshold for proctolin was approximately 2×10^{-12} mol.

In comigration experiments, material corresponding to peaks of interest was collected during the linear gradient using a Dynamax FC-2 fraction collector (Rainin). Purified samples were dried (SpeedVac Concentrator, Savant) and resuspended in HPLC solution A (99.7% H,O, 0.2% CH,CN, and 0.1% TFA). These samples were analyzed by HPLC using the same C_{18} column and solvents, but under isocratic conditions. The isocractic condition for Proct. [2-51 was 18% CH,CN, while for Proct. [3-5] it was 14% CH₃CN. Each resuspended fraction was halved and the following injections were made to examine each peak of interest: (1) half of the collected peak alone, (2) an approximately equivalent amount of the putative synthetic analog, and (3) a mixture containing the other half of the collected peak and an approximately equal amount of the synthetic analog. To confirm sample identity and purity, after isocratic elution, each peak of interest was analyzed also using the original linear gradient. As a final test of sample purity, the Proct. [2-51 peak was collected again, dried, and resuspended for HPLC analysis under isocratic conditions using a second solvent system with triethylamine (TEA, pH 4.5; Pierce, Inc.) rather than TFA as the counter ion. With this solvent system, Proct. [2–5] was assayed at 13% CH₃HN.

HPLC data analysis. All chromatograms were stored for analysis using Dynamax METHOD MANAGER software. The peak areas for the four peaks of interest-proctolin and the three degradation products-were entered into spreadsheets for further analysis using LOTUS l-2-3, and then were imported into SIGMAPLOT (version 4.1) for graphic representation. In most cases, data were normalized by representing each peak of interest as a percentage of the summed peak area of all four peaks. However, this analysis was complicated by the fact that three of the peptidase inhibitors used for these studies shared retention times with one or another of the four peaks of interest. Thus, arphamenine B had one of another of the four peaks of interest. Thus, af phaneling D hau the same retention time as Proct. $[3-5]$, puromycin the same as Proct. $[2-5]$, and epiamastatin the same as proctolin on the TFA linear gradient. Therefore, some data are also represented as the absolute peak area for each peak of interest. All background peaks with the same retention time as Proct. and the Proct. fragments were subtracted from the peaks of interest before analysis and graphic representation. Background peaks were identified as those peaks that were present, with invariant size at all time points, in CS alone aliquots.

Electrophysiology. Electrophysiological experiments were performed on the isolated STNS, which includes the CoGs, OG, and STG, plus their connecting and motor nerves (Fig. 1). The nervous system was pinned down in a Sylgard-coated petri dish, and was superfused continuously with chilled $(10-14^{\circ}C)$ physiological saline at a rate of $7-12$ μ muousiy with chined (10–14°C) physiological sanne at a rate of $/12$ miximum, implies activity from the COOs to the $\frac{51}{10}$ was blocked to applying isotome sucrose (7.5×10^{14} M) to a vaseline well surrounding α and all negative order (Fig. 1). The STG was superfused separately from the OG and all nerves by surrounding the STG with a Vaseline well that held approximately 0.5 ml saline (Fig. 1). Bath volume was approximately 10 ml. α activity of the pyroduced by the pyroduced by the α

ine activity of the pyloric motor pattern produced by the STG was monitored continuously by extracellular recordings from identified STG motor nerves (Nusbaum and Marder, 1989a). Extracellular recordings were made using stainless steel wire electrodes that were pressed into the Sylgard alongside individual motor nerves and isolated from the bath by Vaseline.

To examine the effects of coapplication of proctolin and peptidase inhibitor to the STG, saline superfusion in the STG compartment was stopped, and the saline was first replaced with 500 μ l of either saline alone or peptidase inhibitor (10^{-5} m) dissolved in saline, and then incubated for 10 min (incubation 1). Saline alone was then replaced by either saline alone or proctolin (10^{-8} M or 10^{-7} M) dissolved in saline (500 μ l) and again incubated for 10 min (incubation 2). Similarly, peptidase inhibitor saline was replaced by a saline solution (500 μ I) containing the same peptidase inhibitor (10^{-5} m) plus proctolin (10^{-8} m) , and incubation 2). After the second incubation,

saline superfusion was resumed for 45 min. During test incubations, the temperature in the STG compartment never increased by more than 3°C relative to the temperature in the outer compartment or that occurring during saline superfusion of the STG compartment.

Electrophysiology data analysis. Data were analyzed during the final minute of each application. We characterized how proctolin influenced the pyloric motor pattern by monitoring the pyloric cycle frequency and the number of spikes per burst in the lateral pyloric (LP) motor neuron. Cycle frequency was calculated as the reciprocal of the duration from the first spike in a pyloric dilator (PD) neuron burst to the first spike in the next PD neuron burst. Each analyzed segment is an average of either the cycle frequency or the number of LP spikes per burst over 10 consecutive cycles. In some experiments ($N = 5$), proctolin (10^{-7} M) caused little $(0.1 Hz) or no increase in pyloric cycle frequency. We$ did not include the results from these experiments in our analysis of changes in cycle frequency during application of different peptidase inhibitors. All data were entered into SIGMAPLOT for graphic representation.

Statistics. All statistical analyses were performed using GBSTAT (version 3.0). Statistical significance was determined by using either Student's t test or, for three or more test conditions, analysis of variance (ANOVA). Tukey's t test was used for post hoc multiple comparisons of ANOVA, with a critical level of $\alpha = 0.05$. All data are presented as mean \pm standard deviation (SD). Statistical analyses of the data in Tables 2-5 were performed by comparing the difference between incubation 2 and incubation 1 for each experimental condition relative to this difference for the control condition (10^{-8} M or 10^{-7} M proctolin in saline without any inhibitors, as indicated in each table).

The proctolin data from Tables 2 and 3, and those from Tables 4 and 5, were obtained from separate experiments. In each case, the proctolin data were obtained from the same preparations as were the other data in the same table.

Results

Proctolin degradation

Incubations of proctolin with the TG, followed by HPLC analysis of aliquots of the incubate at 1,60, and 120 min time points, revealed that the amount of proctolin present in the incubate diminished over the course of 2 hr ($N = 4$; Fig. 2A). In contrast, when the same amount of proctolin was incubated under the same conditions with saline alone, the size of the HPLC-detected proctolin peak remained constant over a 2 hr period (not shown). In parallel with the reduction in the size of the proctolin peak during TG incubations there appeared three novel HPLC-detected peaks, called putative proctolin fragments 1-3 (PPF-1, PPF-2, PPF-3), which increased in size with incubation time (Fig. 2A). Thus, at $T = 1$ min, nearly all (97%) of the analyzed peak area resulted from proctolin. At $T = 120$ min, only 82% of this area was due to the proctolin peak and the remaining 18% resulted from the PPF peaks. The concomitant decrease in the proctolin peak and increase in three novel peaks suggested th the proctolin peak and increase in three hover peaks suggested
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PPF-1, the largest of the three novel HPLC-detected peaks, and the three novel HPLC-detected peaks, and the th

 Γ reters, the largest of the time hover Γ reterint of 13.3 min, and had a retention time on the linear gradient of 13.3 min, and thus eluted immediately before the proctolin peak, which had a retention time of 13.7 min (Fig. 2). PPF-2 and PPF-3 had retention times of 11.9 min and 10.6 min, respectively (Fig. 2). FEIGHT HILLS OF THE 2 MINUTED IN THE TEST CONTROL CONTROL IN THE T α is α in contrast, the T in all α min alignous routine α is α in all α in all α is α in all α is α is aliquot. In contrast, the $T = 1$ min aliquot routinely showed a small but discernable peak at the PPF-1 retention time (Fig. 2). This peak, however, resulted from background absorbance as it was also present in control (CS alone) HPLC injections (Fig. $2B$, top trace). W_2 , top trace).

we examined the extent to which the proctom-degrading

Figure 1. Schematic diagram of the stomatogastric nervous system (STNS) of the crab Cancer borealis. The paired commissural ganglia (CoG) were surrounded by Vaseline wells (heavy lines) and the saline inside the wells was replaced with isotonic sucrose (7.5 \times 10⁻¹ M). The stomatogastric ganglion (STG) was surrounded by a separate Vaseline well. The STG was superfused separately from the remainder of the STNS. Shaded region in each ganglion represents the neuropil. In the STG, the neuropil is surround by the neuronal somata. Three lines entering the neuropil represent the axons of neurons innervating the STG from CoGs and OG. dgn, dorsal gastric nerve; dvn, dorsal ventricular nerve; ion, inferior esophageal nerve; lvn, lateral ventricular nerve; mvn, medial ventricular nerve; OG, esophageal ganglion; son, superior esophageal nerve; stn, stomatogastric nerve.

conditioned saline from the TG, after which this CS was incubated with proctolin (see Materials and Methods; $N = 20$). Similar to its incubation with the TG, the proctolin peak diminished over time with CS incubation and the same three novel peaks were the only nonproctolin peaks to change with time (Fig. 2B). Surprisingly, the CS was as effective in degrading proctolin as was incubating the peptide directly with the TG itself ($p > 0.05$ Table 1). As shown by the aggregate data ($N =$ 20) in Figure 3, the amount of each PPF produced in CS at $T = 60$ min and 120 min was significantly greater than that produced at $T = 1$ min. At all time points, the size of the PPF-1 produced at $T = 1$ lillier. At all three points, the size of the $T T T T$ peak was greater than critici or the other two novel peaks. conpeak at these later time points. Thus, the amount of the initial peak at these fater third points. Thus, the amount of the finite proctom anquot (1.6 \pm 0.15 nmol) remaining at $T = 00$ mm was 1.66 \pm 0.12 nmol, while there was 1.55 \pm 0.14 nmol remaining at $T = 120$ min. For these experiments, during the 120 min incubations, the proctolin degradation rate was approximately 0.19 nmol/liter/sec alely 0.19 hindi/iner/sec $T₁$ above conditioned saling that $T₂$ and $T₃$ and $T₄$ and $T₅$ and $T₆$ and $T₇$ and $T₈$ and $T₉$ and $T₉$ and $T₉$ and $T<$

The ability of the TC to produce conditioned same that \mathbb{R} effectively degraded proctolin was not limited to the first 2 hr incubation. In three experiments, three successive conditioned

Figure 2. Proctolin (10⁻⁵ M) degradation resulting from incubation with the TG (A) and conditioned saline (CS, B). In each case, HPLC chromatograms show proctolin degradation at different time points. With increasing incubation time, the putative proctolin fragment (PPF) peaks increased in size. All unlabeled peaks are background peaks that are unrelated

salines were produced, each one involving a 2 hr incubation with the same TG (Fig. 4). In each of these experiments, we found that CS 3 was essentially as effective as CS 1 in reducing the size of the proctolin peak and increasing the size of the PPF peaks on HPLC.

salines were produced, each one involving a 2 hr incubation

Idenbjication of proctolin degradation products rae against the procedure aegregation products

We reasoned, by comparing the above results with those of earlier studies of proctolin degradation in insect neural tissue (Quistad et al., 1984; Isaac, 1987), that the PPFs were likely to

Figure 3. Mean proctolin and PPF peak areas with increasing incubation time in CS ($N = 20$). With increasing incubation time, the percentage of the total peak area due to proctolin decreased while it increased for the PPFs. Note the different scale on the ordinate for the PPF peaks and the proctolin peak. Data are plotted as percentage of total peak area, with 100% equal to the sum of the peak areas of PPF-3, PPF-2, PPF-1, and Proct.; **, $p < 0.01$ compared to $T = 1$ min (ANOVA), for each peak of interest. Abbreviations are the same as in Figure 2.

be Proct. [2-51 (PPF-1), Proct. [3-51 (PPF-2) and Proct. [l-2] (PPF-3). To determine if this was indeed the case, we performed HPLC comigration experiments and compared the retention times of purified PPF-1 and PPF-2 with synthetic analogs (see Materials and Methods). It was difficult to collect and purify sufficient PPF-3 for comigrations, so we compared the HPLC retention time of PPF-3 and synthetic Proct. [l-2] (see below). Purified PPF-1 did indeed consiguity with synthetic Proct. $[2-2]$ (See 00.000). $\frac{1}{5}$, both under isocratic conditions (N = 5; Fig. 5.4) and with a 5], both under isocratic conditions $(N = 5$; Fig. 5A) and with a linear gradient (data not shown). Similarly, PPF-2 comigrated mical gradient (data not shown), similarly, $\text{Tr} \div 2$ comigrated interwith symmetric Froct. $[3-3]$ under isocratic conditions $(N = 3, 7)$ Fig. $5B$). To verify further the identity of PPF-1, we used a second solvent system that used TEA in place of TFA as the counter-ion (see Materials and Methods). Under these conditions, PPF-1 again comigrated with Proct $[2-5]$ (not shown; *n* $=$ 3). Recovery of PPF-2 was insufficient for accurate analysis with the TEA solvent system. In parallel HPLC injections, using the TFA linear gradient, synthetic Proct. $[1-2]$ had the same retention time as PPF-3. ention time as $Pyr-3$.

 Frot. [1-2] and Proct. [3-5] could have been coproduced by a single diaminopeptidase (DAP)-mediated cleavage. Proct. [3-5], however, could also have been produced by AP cleavage of Proct. $[2-5]$. To study this latter possibility, we incubated synthetic Proct. [2-5] (7×10^{-9} mol) both with the TG and with CS, using the same protocols and HPLC analysis that were used for proctolin incubations. Both TG ($N = 3$; data not shown) and $CS (N = 4; Fig. 6) degraded Proof. [2–5] and produced Proct.$ [3-5]. After 2 hr CS incubations, $94.3 \pm 2.2\%$ of the total peak area was Proct. $[2-5]$ (Fig. 6), as compared with approximately 88% being due to the proctolin peak (Table 1) after comparable

Figure 4. Proctolin (10⁻⁵ M) degradation by sequentially generated conditioned salines from the TG, assayed after incubations of 120 min. Data are plotted as the percentage of the total peak area of the proctolin and PPF peaks. All three conditioned salines were from normal Cancer saline. $N = 3$; $p > 0.05$ (ANOVA), for comparisons of corresponding peaks in all three incubations.

proctolin incubations. Thus, after these 2 hr incubations, aliquots contained 1.59 ± 0.07 nmol of Proct. [2-5], while aliquots at $T = 0$ min contained 1.80 \pm 0.19 nmol. This resulted in a degradation rate of approximately 0.22 nmol/liter/sec, which was similar to the proctolin degradation rate (0.19 nmol/liter/ sec). As shown in Figure 6, there were increasing amounts of Proct. [3-51 produced over the course of the 2 hr incubations. Thus, Proct. [3-51 could be produced by serial N-terminal cleav-Thus, Proct. [3-5] could be produced by serial N -terminal cleavage of proctolin. In contrast to the proctolin incubation experiments, no Proct. $[1-2]$ was produced from the Proct. $[2-5]$ incubations.

Aminopeptidase inhibitors diminish proctolin degradation Proctolin had been shown to be sensitive to AP activity (Quistad

Proctom had been shown to be sensitive to AP activity (Quistad) et al., 1984; Isaac, 1987), so we tested the ability of six different AP inhibitors to prevent proctolin degradation by CS. As shown in Figures 7 and 8, each AP inhibitor that we tested was able to prevent at least partially the reduction in the size of the proctolin peak and/or the appearance of one or more of the proctolin fragment peaks. For the inhibitors amastatin, acti-

 $p > 0.05$, Student's t test, for comparison of each peak in TG versus CS.

Figure 5. Comigration of proctolin degradation products and their synthetic analogs: HPLC chromatograms under CH,CN isocratic conditions, following collection from CH,CN linear gradient using TFA as the counter-ion (see Materials and Methods) for synthetic Proct. $[2-5]$ and PPF-1 (A) , and synthetic Proct. $[3-5]$ and PPF-2 (B) . AI , Half of synthetic Proct. $[2-5]$ fraction collected from linear gradient. A2, Half of PPF-1 fraction collected from linear gradient following proctolin incubation in CS. A3, Combination of second half of synthetic Proct. [2-5] and second half of PPF-1 from proctolin degradation in CS. B1, Half of synthetic Proct. [3-51 fraction collected from pre- vious TFA linear gradient. B2, Half of PPF-2 fraction collected from previous TFA linear gradient following proctolin incubation in CS. B3, Combination of half of synthetic Proct. [3–5] and PPF-2 fractions. Arrows indicate peaks of interest. Unlabeled peaks are solvent front and background peak.

nonin, and bestatin, we quantitatively analyzed the fraction of the initial amount of proctolin that was found in the form of proctolin, Proct. $[2-5]$, Proct. $[3-5]$, and Proct. $[1-2]$ after 2 hr co-incubations of CS with proctolin and each inhibitor (Fig. 8A). Because of overlap of HPLC retention times with peaks of interest, data for the other three inhibitors could only be analyzed as peak areas (see Materials and Methods; Fig. 8B). When we compared these peaks with those resulting from incubation of proctolin without inhibitor, we found that each of these inhibitors significantly decreased the amount of proctolin degradation (Fig. 8). In terms of the size of either the proctolin peak or that of Proct. [2-51, amastatin and actinonin were equally effective and were more effective than bestatin. For example, after 2 hr incubations, 88% of the total peak area was proctolin when no inhibitors were present, while 93% was proctolin when bestatin was also present and 98% was proctolin when either amastatin or actinonin was present (Fig. 8A). Interestingly, the ability of each inhibitor to prevent the ap-
Interestingly, the ability of each inhibitor to prevent the ap-

perception of Processes of Processes and not necessarily correlate include the material correlation of the theory pearance of Proct. $[2-5]$ did not necessarily correlate with the ability of that inhibitor to block the appearance of Proct. $[3-5]$ and Proct. [l-2]. Thus, relative to CS alone, only actinonin and TTOUL $1-2$. Thus, relative to US alone, only actificantly $\frac{1}{2}$ and $\frac{1}{2}$ (Figure and $\frac{1}{2}$ and \frac $[1-2]$ (Figs. 7, 8). While amastatin also significantly reduced the size of the Proct. $[3-5]$ peak, it did not change the size of the Proct. $[1-2]$ peak relative to CS alone (Figs. 7, 8). Additionally, whereas bestatin and arphamenine had little influence on Proct. $[3-5]$, their presence produced a slight increase in the Proct. $[1-$ 2] peak size (Fig. $8B$). Puromycin did not alter the size of either the Proct. [3–5] or Proct. [1–2] peaks.
Like proctolin degradation, the Proct. [2–5] degradation was

also amastatin sensitive. Thus, after 120 min co-incubations of Proct. [2-5] plus amastatin in CS, the Proct. [2-5] peak was indistinguishable from that of controls (percentage total peak area, 99.1 \pm 0.2%; N = 4) and only trace amounts (<1%) of Proct. [3-51 were detected. In contrast, approximately 6% of the peak area was represented by the Proct. [3-51 peak after 120 min incubations of Proct. [2-51 with CS alone (Fig. 6). This amastatin-sensitive degradation of Proct. [2-51 presumably accounts for the decrease in the Proct. [3-51 peak that occurred from proctolin incubations in CS plus amastatin. No other inhibitor was tested on Proct. [2-5] degradation.

Proctolin actions are enhanced by aminopeptidase inhibitors

To test the physiological role of the AP activity identified in our biochemical studies, we examined the ability of the AP inhibitors used in these biochemical studies to influence the minoriors used in these orochemical studies to immente the the structure galaxies of proctome on the pytone motor pattern in the stomatogastric ganglion. The experimental setup is illustrated in Figure 1 and is described in the Materials and Methods. We reasoned that if the AP inhibitors effectively prevented proctolin degradation, then they should enhance the proctolin-mediated excitation of the pyloric motor pattern in the STG. To test this hypothesis, we applied each AP inhibitor (10^{-5} M) with proctolin (10^{-8} M). As predicted, many of these inhibitors did enhance the proctolin excitation of the pyloric motor pattern (Figs. 9, 10; Tables 2, 3: incubation 2 vs 1). In contrast, none of these inhibitors influenced the pyloric motor pattern when applied alone (Tables 2, 3; incubation 1). The enhanced excitation that occurred when these inhibitors were coapplied with proctolin included (1) a faster pyloric motor pattern, and (2)

Figure 6. Proct. $[2-5]$ (10⁻⁵ M) is degraded during incubations with CS from the TG. The percentage of the total peak area (Proct. [2-51 plus Proct. [3-51) due to Proct. [2-51 decreased while that due to Proct. [3-5] increased with increasing incubation time. $N = 4$; **, $p < 0.01$; $i, p < 0.05$; compared to $T = 1$ min (ANOVA) for each peak of interest. Data are plotted as in Figure 3.

more intense impulse bursts in several pyloric neurons. For example, in the experiment shown in Figure 9, bestatin only slightly enhanced the proctolin-mediated excitation of the pyloric rhythm, but amastatin increased this activity such that it was equivalent to that occurring in the presence of a IO-fold higher concentration of proctolin. Only the peptidase inhibitors amastatin and arphamenine B significantly increased the pyloric cycle frequency relative to proctolin (10^{-8} M) alone (Table 2). Cycle inequency relative to proctomic (row M) alone (rable 2). while the checks of actificating on eye requested ω be comparably large, they were only studied in two experiments (Table 2). In contrast, all five inhibitors enhanced the number of LP neuron impulses per burst as compared with proctolin (10-8 μ) alone (Fig. 10, Table 3). Bestatin and ambanonine B, (10-8 μ). Bestatin and ambanonine B, (10 \degree M) alone (Fig. 10, Table 3). Destatin and arphamentic β , α and α and α and α and α and α mass of α and α α mass of α m $\sum_{n=1}^{\infty}$ and $\sum_{n=1}^{\infty}$ is a set it was individual individual that it was indictinguishable from on LP neuron activity such that it was indistinguishable from that resulting from application of 10^{-7} M proctolin (Fig. 10, Table 3).

Physiological actions of proctolin degradation products T

I o test whether the observed proctoll cleavages were macuvating the peptide, we studied the effects on the pyloric motor pattern of bath applying each identified proctolin degradation product to the STG. As shown in Figure 11, when we applied Proct. $[2-5]$ in the concentration range where the parent peptide effectively excites the pyloric rhythm $(10^{-9}$ M to 10^{-6} M), there was no effect on this motor pattern. However, higher doses of Proct. $[2-5]$ (>10⁻⁶ M) did excite the pyloric rhythm ($n = 3-5$). This effect was moderately strong for applications of 10^{-5} M Proct. $[2-5]$ and it was particularly strong for applications of 5 \times 10⁻⁵ M (Fig. 11). These latter effects were indistinguishable from those of lower doses of the parent peptide $(10^{-7}$ M proctolin; Fig. 1, Tables 4, 5). In contrast to the results with Proct.

Figure 7. Proctolin degradation in the presence of different aminopeptidase inhibitors: HPLC chromatograms of aliquots of proctolin (10⁻⁵ M) plus or minus aminopeptidase inhibitor (10⁻⁴ M) in CS, after incubation for 120 min ($T = 120$ min). The size of the proctolin peak and that of peak I were both reduced with aminopeptidase inhibitors present when compared with CS plus proctolin alone. Additionally, peak 2 was reduced by amastatin and actinonin, while peak 3 was reduced by actinonin. Best., bestatin; Actin., actinonin; Amast., amastatin; Proct., proctolin.

[2-5], bath application of even higher doses (10^{-4} M) of Proct. $[2-3]$, bath application of even higher doses (10 \pm M) of Flock. $[1-2]$ and 1 roct. $[3-3]$

Discussion

Proctolin degradation by conditioned saline

 I foctor and the pentapel shown that the pentapeptide process is the pentapeptide process in the pentapeptide process I In this study we have shown that the pentapeptide proctor is \mathbb{R}^n . cleaved by the crab nervous system. This cleavage is mediated primarily by AP activity, although there appears to be some DAP-like activity as well. Surprisingly, we found that saline preincubated with the TG had nearly as much proctolin-cleaving activity as the TG. Thus, nearly all of the proctolin-cleaving activity was probably released into the extracellular space rather than remaining tightly bound to ganglionic membranes.

There are several possible explanations for the generation of proctolin-degrading activity in the CS. For example, this activity might have resulted from exocytosis. Alternatively, the activity might have been tethered to the extracellular surface of ganglionic membranes by a labile linkage that was continually cleaved.
Neither of these possibilities was examined in our experiments.

Figure 8. Summary of effectiveness of different aminopeptidase inhibitors on CS-mediated proctolin degradation. A, Percentage of total peak area represented by each of the proctolin-related peaks. Data were collected following proctolin incubation (10^{-5} M) with CS plus or minus an aminopeptidase inhibitor (10⁻⁴ M) at $T = 120$ min. For each condition, data are plotted as the mean peak area for each peak divided by the mean total area for the four proctolin-related peaks. B, Mean peak area of proctolin and each proctolin fragment after CS incubation of proctolin (10^{-5} M), plus or minus an aminopeptidase inhibitor (10⁻⁴ M; $T = 120$ min). Data for no inhibitors, bestatin, actinonin, and amastatin are replotted from A. Missing bars indicate values for which the inhibitor shared the same retention time as the peak of interest. Overlapping peaks are arphamenine B and Proct. [3-51, puromycin and Proct. [2-S], and epiamastatin and proctolin. Statistical analysis indicates significantly different than CS with no inhibitors within each peak of interest: **, $p < 0.01$; *, $p < 0.05$; ANOVA.

However, there is precedence for the latter possibility in that other cell surface enzymes, including acetylcholinesterase, are membrane anchored via a phospholipase C-sensitive, glycosylphosphatidylinositol linkage (Futerman et al., 1985; Haas et al., 1988; Low and Saltiel, 1988). Such a linkage also exists for at least one AP, aminopeptidase P (AP-P, EC 3.4.11.9; Hooper and Turner, 1988).

Multiple proctolin-degrading enzymes?

Our results are consistent with proctolin cleavage in the crab nervous system being mediated by at least two different ennet vous system being inequated by at least two different enzymes. The major proctolin-degrading activity appears to be an AP, since most of the proctolin cleavage is suppressed by AP Ar_1 , since most of the proctom cleavage is suppressed by Ar_1 the major AP is the major and the completion experiments intreate the the major AP inhibitor-sensitive degradation product is Proct. [2-5]. The second proctolin-degrading activity appears to be a DAP.

The second proctom-degrading activity appears to be a DAF . This conclusion is based primarily on the coappearance of Proct. $[1-2]$ and Proct. $[3-5]$ in proctolin incubations with either CS or TG. While Proct. [3-5] was also produced via serial, AP inhibitor-sensitive cleavages of proctolin, the larger source of Proct. [3–5] generation appeared to be cleavage of the Tyr-Leu bond via DAP activity. This interpretation is supported by the observation that in most of the proctolin co-incubations with AP inhibitors, the same amounts of Proct $[3-5]$ and Proct. $[1-$ 2] were generated as without any inhibitor present, even though Proct. [2-5] production was reduced significantly. The DAP-like activity, however, can only account for about 20% of the total

proctolin degradation, because approximately 80% of the proctolin degradation is eliminated in the presence of the AP-inhibitor amastatin.

Identity of the proctolin-cleaving peptidases

Previous workers (McDonald and Barrett, 1986; Turner et al., 1989) have characterized four distinct mammalian APs that show some similarity to the proctolin-cleaving activity that we have described in the crab nervous system. These include leutiave described in the crab fiel vous system, These mendo feu-
sine AP (LAP; EC 2.4.11.1), aminopeptidase M (AP-M; EC CHE AT $(LA, LC, J, 11.1)$, aminopeptidase B $(LT, LC, 3.4.11.6)$, and amino-3.4.11.2), aminopeptidase B (AP-B; EC 3.4.11.6), and amino-
peptidase A (AP-A; EC 3.4.11.7). Each of these peptidases exhibits different substrate and inhibitor specificities. Of these enzymes, the likeliest ones to degrade proctolin effectively are LAP, AP-M, and AP-B. Purified mammalian LAP has been S_{max} , AP-M, and AP-D. Furnical manimianal LAT has been shown to cleave proctome electricity in solution, annough the cleavage products were not identified (Brown, 1977; Davis et al., 1989). The preferred substrates for both AP-M and AP-B include peptides, such as proctolin, with an N -terminal arginine (McDonald and Barrett, 1986; Turner et al., 1989). Each of these APs, however, has been reported to have an inhibitor-sensitivity profile that is distinct from that of the proctolin-sensitive AP activity seen by us in the crab (Umezawa and Aoyagi, 1983; McDonald and Barrett, 1986). It is, however, possible that the AP activity in our study represents several different enzymes with overlapping inhibitor sensitivities, rather than a single AP.
Supporting this possibility is our finding that the crab AP ac-

Figure 9. Effects of bath-applied aminopeptidase inhibitors (10⁻⁵ M) on the excitatory effects of proctolin (10⁻⁸ M) on the pyloric motor pattern. Activity of an ongoing pyloric motor pattern from a single experiment is monitored by extracellular recordings of a pyloric motor nerve, the dvn (see Fig. 1). LP, lateral pyloric constrictor motor neuron; PY, pyloric constrictor motor neuron; PD, pyloric dilator motor neuron.

tivity was approximately equally inhibited by amastatin, actinonin, puromycin, and epiamastatin.

Previous work in the insect nervous system showed that extracellular proctolin is a substrate of AP activity and that its degradation was most effectively inhibited by amastatin (Quis-

Table 2. The effect of aminopeptidase inhibitors on the proctolinmediated excitation of the pyloric rhythm cycle frequency (Hz)

	Saline superfusion		Incubation 1 Incubation 2
Saline $(n = 5)$	0.45 ± 0.2	0.47 ± 0.2	0.46 ± 0.2 †
10^{-8} M proctolin $(n = 9)$	0.46 ± 0.2	0.47 ± 0.1	0.54 ± 0.2
Bestatin ($n = 7$)	0.39 ± 0.2	0.44 ± 0.2	0.50 ± 0.2 †
Epiamastatin $(n = 5)$	0.48 ± 0.1	0.48 ± 0.2	0.55 ± 0.2 †
Arphamenine B $(n = 5)$	0.39 ± 0.1	0.38 ± 0.2	$0.49 \pm 0.2^*$
Actinonin $(n = 2)$	0.54 ± 0.4	0.63 ± 0.4	0.72 ± 0.2
Amastatin $(n = 5)$	0.47 ± 0.1	0.47 ± 0.1	$0.61 \pm 0.1*$
10^{-7} м proctolin (n = 5)	0.42 ± 0.2	0.45 ± 0.2	$0.66 \pm 0.2^*$

"Incubation 1" indicates STG incubation with either saline (for saline, 10^{-8} M proctolin, and 10^{-7} M proctolin) or aminopeptidase inhibitor $(10^{-5}$ M, for all others). "Incubation 2" indicates STG incubation with incubate 1 plus either saline (for saline), 10^{-7} M proctolin (for 10^{-7} M proctolin), or 10^{-8} M proctolin (for all others). See Materials and Methods for further details.

* Value is significantly different from 10^{-8} M proctolin application: $p < 0.05$, Student's t test.

† Value is significantly different from 10^{-7} M proctolin application: $p < 0.05$, Student's t test.

tad et al., 1984; Isaac, 1987). However, other aspects of the inhibitor profile determined by Isaac (1987) for the proctolincleaving AP activity of insect neural membrane fractions differed from that of the AP activity described in the present study. Proctolin was also degraded by DAP activity in these previous studies, although the particular DAP responsible for this degradation was not identified (Quistad et al., 1984; Isaac, 1987). However, Hui (1988) reported that proctolin is readily cleaved by DAP V. DAP V, purified from rat brain membranes, strongly prefers peptides with an arginine residue at the N-terminus, including proctolin.

Details are as for Table 2.

Figure 10. Cumulative results of the influence of different aminopeptidase inhibitors on the proctolin-mediated changes in the number of LP neuron impulses per burst. Data represent the mean difference from application of either saline (saline, 10^{-8} M Proct., 10^{-7} M Proct.) or the relevant inhibitor. Each inhibitor (10^{-5} M) was first applied alone, and then immediately coapplied with proctolin (10⁻⁸ M). *S*, saline (*n* = 8);
 10^{-8} M Pr, proctolin (*n* = 12); *B*, bestatin (*n* = 8); *Ar*, arphamenine (*n* = 8); *Ac*, actinonin (*n* = 4); *E*, epiamastatin (*n* = 5 Proct.); $+$, $p < 0.05$ (relative to 10^{-7} M Proct.); Student's t test.

Physiological role of proctolin-cleaving peptidases

Our results are consistent with there being extracellularly located AP activity within the STG that terminates the neural actions of proctolin. This conclusion is based on the fact that the same AP inhibitors that effectively suppressed proctolin degradation in our biochemical experiments were able to enhance the excitatory effects of bath-applied proctolin in our electrophysiological experiments. Although we analyzed the enzymatic degradation of proctolin using a different ganglion than we used to monitor its electrophysiological effects, the order of effectiveness of the peptidase inhibitors used in both types of experiments was similar.

In addition to the proctolin-degrading activity reported by us to reside within the crab nervous system, there is previously

Figure 11. Effects of Proct. [2-5] on the pyloric motor pattern. Bath application of Proct. $[2-5]$ at $\leq 10^{-5}$ M to the isolated STG did not influence the pyloric motor pattern. However, higher concentrations of Proct. $[2-5]$ (5 × 10⁻⁵ M) excited the pyloric motor pattern in a manner similar to bath application of proctolin (10^{-7} M) . Pyloric motor pattern was monitored by an extracellular recording of the $\frac{dv}{dx}$. All recordings were from the same preparation.

reported proctolin-degrading activity within the crustacean hemolymph. Schwarz et al. (1984) reported that, in preliminary experiments where they injected exogenous proctolin in the hemolymph of the lobster Homarus americanus, proctolin was degraded with a half-life of 6-10 min. Proctolin cleavage also occurs in insect hemolymph (Steele and Starratt, 1985). Because the STG is located within a major artery, any neurally released proctolin that was not degraded by ganglionic enzyme activity would likely be degraded in the hemolymph. Our experiments

Details are as for Table 2.

Details are as for Table 2.

do not enable us to predict which of these two sources of proctolin-degrading activity would be larger.

We were surprised to find that proctolin was only approximately 500-fold more effective than Proct. [2-51 in exciting the pyloric motor pattern. An earlier study by Starratt and Brown (1979) using a proctolin-sensitive muscle assay, showed that proctolin was $> 10⁶ \times$ more effective than Proct. [2–5]. Whether this difference results from different proctolin receptors, differential sensitivities to resident peptidases, or differential access to receptors remains to be determined.

The proctolin degradation reported in the present study may well contribute to the relatively short-lasting excitatory effects of two proctolin-containing neurons, MPN and MCNl, on the pyloric motor pattern in the crab STG (Nusbaum and Marder, 1989a,b; Nusbaum et al., 1992; Christie et al., 1993; Coleman and Nusbaum, 1994). Whereas some modulatory neurons have effects that long outlast their period of impulse activity (Harris-Warrick et al., 1992b), the effects of both MPN and MCNl subside within seconds after their activity is terminated (Nusbaum and Marder, 1989a,b; Nusbaum et al., 1992). Because the effects of the proctolin neurons on the pyloric rhythm are equivalent to bath application of 10^{-7} M to 10^{-6} M proctolin and threshold for proctolin actions on this system is approximately 1O-9 M (Marder et al., 1986; Nusbaum and Marder, 1989b), the local concentration of neurally released proctolin probably decreases by 2-3 orders of magnitude within several seconds or less. Such a rapid decrease in local proctolin concentration would be facilitated by the proctolin-cleaving actions identified in the present study. Finally, since many extracellularly located peptidases cleave more than one neuropeptide (Turner et al., 1985; Turner, 1987), it is possible that the peptidase activities documented in this study also degrade other neurally released pep t_{t} in the state and depends system.

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