Cellular/Molecular

Peptidomics of a Single Identified Neuron Reveals Diversity of Multiple Neuropeptides with Convergent Actions on Cellular Excitability

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In contrast to classical transmitters, the detailed structures and cellular and synaptic actions of neuropeptides are less well described. Peptide mass profiling of single identified neurons of the mollusc Lymnaea stagnalis indicated the presence of 17 abundant neuropeptides in the cardiorespiratory neuron, visceral dorsal 1 (VD1), and a subset of 14 peptides in its electrically coupled counterpart, right parietal dorsal 2. Altogether, based on this and previous work, we showed that the high number of peptides arises from the expression and processing of four distinct peptide precursor proteins, including a novel one. Second, we established a variety of posttranslational modifications of the generated peptides, including phosphorylation, disulphide linkage, glycosylation, hydroxylation, N-terminal pyro-glutamylation, and C-terminal amidation. Specific synapses between VD1 and its muscle targets were formed, and their synaptic physiology was investigated. Whole-cell voltage-clamp analysis of dissociated heart muscle cells revealed, as tested for a selection of representative family members and their modifications, that the peptides of VD1 exhibit convergent activation of a high-voltage-activated Ca current. Moreover, the differentially glycosylated and hydroxylated α2 peptides were more potent than the unmodified α2 peptide in enhancing these currents. Together, this study is the first to demonstrate that single neurons exhibit such a complex pattern of peptide gene expression, precursor processing, and differential peptide modifications along with a remarkable degree of convergence of neuromodulatory actions. This study thus underscores the importance of a detailed mass spectrometric analysis of neuronal peptide content and peptide modifications related to neuromodulatory function.

Key words: mollusk; MALDI-TOF mass spectrometry; glycopeptide; posttranslational modification; HVA calcium channel; neuromodulation

Introduction

Neuropeptides play many important roles in the generation of behavior and the regulation of physiological processes in both vertebrates and invertebrates. Often, multiple neuropeptides are coexpressed in single cells, and their combinatorial action through divergent as well as convergent pathways enhance the information processing capacity of neurons in the nervous system (Kupfermann, 1991; Peaker, 1992; Zupanc, 1996; Brezina and Weiss, 1997; Hokfelt et al., 2000; Kupfermann and Weiss, 2001; Nusbaum et al., 2001; Merighi, 2002). This emphasizes the importance of detailed knowledge of the full complement of the chemical diversity of the coexpressed neuropeptides in relation to their possible action as cotransmitters in neuronal communication.

In principle, peptide diversity can be generated via several mechanisms, namely (1) cell-specific coexpression of distinct peptide precursor genes (Cropper et al., 1987; Church and Lloyd, 1991; Hekimi et al., 1991; Vilim et al., 1996; Jimenez et al., 1998; Wood et al., 2000), (2) alternative splicing of a pre-mRNA encoding different precursors (Saunders et al., 1992; Bogerd et al., 1993), (3) alternative proteolytic processing of a precursor protein (Eipper and Mains, 1980) and (4) posttranslational modifications of the peptides (Loh, 1992; Dockray et al., 1996). In all instances, a peptidomics approach based on mass spectrometry has proven to be superior to other classic biochemical approaches for defining peptide diversity at the level of single cells and small biopsies (Jimenez and Burlingame, 1998; Li et al., 2000; Schrader and Schulz-Knappe, 2001). Although studies using this methodology have revealed that the coexistence of multiple neuropeptides in single (invertebrate) neurons is a general phenomenon, the physiological significance of the full chemical diversity of
peptides, including that generated by posttranslational modifications, has not been determined.

The identifiable neuron visceral dorsal 1 (VD1) and the electrically coupled right parietal dorsal 2 (RPD2) in the brain of the mollusc *Lymnaea stagnalis* provide a convenient model preparation in which to examine the generation of peptide diversity in the context of biological function, namely, the modulation of cardiorespiratory behavior (Janse et al., 1985; Kerkhoven et al., 1991). Previous studies have identified three peptides in VD1 and RPD2, of which two were shown to possess cardioexcitatory effects (Bogerd et al., 1994). Subsequent mass spectrometric analysis of single VD1 and RPD2 neurons (Jimenez et al., 1994, 1998) revealed overlapping peptide profiles and the presence of a total of 17 abundant mass peaks, representing putative peptides. This finding suggested that the peptide phenotype of VD1 and RPD2 is far more complex than reported previously (Bogerd et al., 1994) because of the expression of hitherto unknown neuropeptides and/or posttranslational processing.

To begin understanding the significance of neuropeptide diversity in VD1 and RPD2 and its functional correlate on target cells, we first performed a peptidomics approach to characterize the unidentified neuropeptides present in VD1 and RPD2, including their posttranslational modifications. Subsequent functional analysis of VD1 and RPD2 peptides on its target cells, i.e., heart muscle cells, showed that peptides derived from distinct precursor proteins and differentially modified forms exhibit convergent yet differential activation of the same calcium channel current in heart muscle fibers.

**Materials and Methods**

**Animals**

Adult specimens of *L. stagnalis* (shell height, 30–35 mm) bred in the laboratory under standard conditions were used. The animals were kept at 20°C and a 12 h light/dark cycle and were fed lettuce ad libitum.

**Extraction and purification of peptides**

Dissected VD1 and RPD2, which shares the same set of peptides as VD1 with the exception of the small cardioactive peptides (SCPs) (Jimenez et al., 1998), were collected on solid carbon dioxide and stored at −80°C. Pooled neurons (total, 800) were extracted by boiling in 0.1 m acetic acid for 8 min and centrifuged for 10 min at 4°C, and the supernatant was fractionated by HPLC using a Nucleosil C18 column (5 μm; 250 × 2.1 mm) with an increasing concentration of acetonitrile in 7 mM trifluoroacetic acid (TFA). The flow rate was 300 μl/min, and 1 min fractions were collected. Each fraction was screened by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF-MS) to search for the molecular ion species that were detected in the single VD1 mass spectrum. In some cases, fractions that contained the molecules of interest were further separated using the same Nucleosil column with increasing acetonitrile concentration in 0.05% HCl.

**Mass spectrometry**

For MALDI-TOF-MS, an aliquot (0.5 μl) of each sample (HPLC fraction or enzymatic digest) was mixed with 1 μl of matrix (10 mg 2,5-dihydroxy-benzoic acid and 1 mg 5-methoxy-2-benzoic acid dissolved in 1 ml of 7.5 mM trifluoroacetic acid in 30% acetonitrile). After drying of the sample, the target was placed into a laboratory-built laser desorption reflectron time-of-flight mass spectrometer equipped with a pulsed nitrogen laser (337 nm; pulse width, 3 ns) for analysis, as described previously (Jimenez et al., 1998). The spectra were externally calibrated using a peptide mixture, yielding an accuracy of 0.05%.

To obtain peptide sequence information, post-source decay mass spectra were recorded on a VG ToFspec SE time-of-flight mass spectrometer (Micromass, Manchester, UK) equipped with a nitrogen laser operated in reflectron mode, as described previously (Matsui et al., 1997). Samples were prepared as described above.

Electrospray MS and fragmentation spectra were recorded using a triple-quadrupole instrument (API300) from MDS-Sciei (Concord, Ontario, Canada) with a standard ion spray source that was infused at 5 μl/min. The collisionally induced dissociation gas used was nitrogen, and the collision energy was 35–50 eV, 0.2 Da step size, and 5–10 s total scan time.

**Amino acid sequence analysis**

Amino acid sequencing was performed by automated Edman degradation in a model 473 pulse liquid phase sequencer (Applied Biosystems, Foster City, CA), using the sequencing program recommended by the company. The sequence data were also used for quantification of the amount of peptides purified.

**Enzymatic digestions**

Peptide sequences were incubated with 1 μg of endo-lys-c (Sigma, St. Louis, MO) in 100 μl of 50 mM ammonium acetate buffer containing 1 mM dithiothreitol for 18 h at 20°C. Purified peptide K and α2 peptides were incubated with 1 μg of trypsin (Promega, Madison, WI) in 100 μl of 50 mM ammonium carbonate buffer for 18 h at 20°C. Dephosphorylation of peptide K was performed by incubation with calf intestine alkaline phosphatase (Promega) in 100 μl of 50 μl of 50 mM ammonium carbonate buffer containing 1 mM magnesium chloride for 30 min at 30°C. Digested peptides were directly analyzed by MALDI-TOF-MS and in some cases purified by reversed-phase HPLC using trifluoroacetic acid as counterion.

**Cloning and sequencing of pro-I cDNA, Northern blotting, and in situ hybridization**

To obtain the cDNA sequence of the precursor of peptide I (pro-I) first two degenerate nested antisense primers oligo VD2 (OLVD2) [5′-GGAAATTCGCGAANCAGGCACGAGAA(TC)TCNGC(TG)GATC-3′] and OLVD3 [5′-GGAAATTCGAIAC(GA)AA(TC)TCNGC(TG)GATC/TC(TC)-GAGTANGC-3′] (Isogen Bioscience, Ijsselstein, The Netherlands) containing 5′-EcoRI restriction sites, designed on the partial peptide I sequence KAYED RefV PVVPVK were used. Then, two specific nested sense primers, I-peptide query sense (IQS3) (5′-GGGACGTCA-GGTATATCCAGCTGCG-3′) and IQS4 (5′-GGGATCCAgAC-TGCAAGCCGAG-3′) (Isogen Bioscience) containing 5′-BamHI restriction sites, designed on the 5′ end of the obtained 5′ rapid amplification of cDNA ends (RACE) I-cloned, were used. The specific peptide I primers were used in combination with an anchored dT primer [adapter oligo A (ADA)-dT30] and its respective adaptor primers ADA-I and ADA-II (Tensen et al., 1994) in 5′ RACE and 3′ RACE PCR on hexanucleotide primed cDNA from single RPD2 neurons as described previously (Spijker et al., 1999) with the following modifications. Three individual RPD2s were isolated using a fine glass pipette (tip diameter, ±100 μm) and transferred to a glass plate. After removal of superfluous liquid, 11 μl of water containing 400 pmol of hexanucleotides was added and mixed, and the sample was heat treated and processed for cDNA synthesis. After incubation, the cDNA was phenolized and precipitated. After dATP tailing, each cDNA synthesis was used for a first, a second, and a third round of amplification using 50 pmol of I-primer (OLVD2 or IQS3) in combination with ADA-dT30, 50 pmol of I-primer (OLVD2 or IQS3) in combination with 50 pmol of ADA-I, and 50 pmol of nested I-primer (OLVD3 or IQS4) in combination with 50 pmol of nested adaptor primer ADA-II. Amplified cDNA of 5′ and 3′ RACE was digested with EcoRI/SaeI or with BamHI/SaeI and separated on an agarose gel. Fragments were cloned and sequenced in both orientations according to the dideoxy chain termination method (Sanger et al., 1977), using T7 DNA polymerase.

Total RNA from visceral and parietal ganglia was isolated (Chomczynski and Sacchi, 1987), and ~10 μg of RNA was glyoxylated, fractionated on a 1.8% agarose gel, and transferred to charged nylon membranes. A peptide-I-specific 3′ RACE clone (nucleotides 79–633) was used to make a randomly primed probe, labeled with [α-32P]dATP (specific activity >108 cpm/μg) by primer extension. The membrane was hybridized in 6× SSC, 0.2% SDS, 5× Denhardt’s solution, 10 μg/ml RNA, and 10 μg/ml herring sperm DNA, at 65°C for 16 h, subsequently washed in 0.5X SSC and 0.2% SDS at 65°C for 45 min, and then autoradiographed.

The cellular localization of mRNA encoding peptide I was studied in sections of the visceral and right parietal ganglia of *L. stagnalis* in situ.
hybridization using a digoxigenin–dUTP-labeled cRNA. Parallel sections were used to confirm the simultaneous presence of α1 peptide in VD1 and RPD2 using immunocytochemistry. Tissue was fixed in 1% paraformaldehyde and 1% acetic acid (Smit et al., 1996). cRNA probes were synthesized from 100 ng of a PCR fragment of the M13 vector containing a partial I-peptide cDNA (nucleotides 1–260), including the T3 and T7 RNA polymerase promoter. Hybridization, washing, and visualization were done as described by Smit et al. (1996).

Cardiac muscle dissociation and neuron isolation
Muscle cells were dissociated from dissected atria and ventricles under sterile conditions using 0.2% dispase/collagenase in defined medium (Ridgway et al., 1991) (one atrium or ventricle per ml) for 4 h at 30°C while gently shaking. The dissociated cells were subsequently washed three times with defined medium and sedimented at low speed, before plating. Dissociated muscle fibers were kept in defined medium in Costar (Cambridge, MA) dishes at 20°C for 1–4 d (Lee et al., 2002). All neuronal culture procedures were performed as described previously (Syed et al., 1996). In brief, VD1 and right pedal dorsal 1 (RPD1) were mechanically isolated under sterile conditions from the brain by applying gentle suction through a fire-polished pipette. Isolated cells were plated directly on poly-l-lysine-coated dishes containing brain-conditioned medium. After 24 h in culture, muscle cells were plated on top of neurites of VD1, and synapses were allowed to form during 24 h.

Electrophysiology
Simultaneous intracellular recordings from cocultured cells. For intracellular recordings, conventional electrophysiological techniques were used as described previously (Syed et al., 1996). Glass microelectrodes (resistance of 20–40 MΩ) were filled with a saturated solution of potassium sulfate (K₂SO₄), and neurons were impaled using Narishige (Tokyo, Japan) micromanipulators (models M202 and M204). The electrophysiological signals were amplified (amplifier model IR-283; Neurodata Instruments, New York, NY), displayed on a model IR-283; Neurodata Instruments, New York, NY), displayed on a Gould Instruments (Valley View, OH) chart recorder (model TA240S).

Voltage-clamp recordings. Dissociated ventricle and auricle cells were bathed in HEPES-buffered saline (HBS) (see above). To record calcium currents, we used Ba²⁺ as charge carrier. HBS was washed out and replaced under continuous perfusion by a solution containing the following (in mM): 40 NaCl, 2 BaCl₂, 10 HEPES, and 2 4-aminopyridine, pH 7.8

0.1 GΩ feedback resistance. Pipettes (2–6 MΩ) were pulled on a Flaming/Brown P-87 (Sutter Instruments, Novato, CA) horizontal microelectrode puller from Clarke Electromedical Instruments (Pangbourne, UK) GC-150 glass. In the cell-attached configuration (seal resistance of ~1 GΩ or more), the fast capacitive transients of the patch pipette were compensated. After disruption of the patch membrane, series resistance (~7 MΩ) and cell capacitance (~100 pF) were compensated. In the whole-cell mode, we waited ~15 min before measuring barium currents, to equilibrate the interior of the cell with the pipette solution. Data acquisition was controlled by a CED 1401 analog-to-digital converter (Cambridge Electronics Design, Cambridge, UK) connected to an Intel 80486-based computer, run with voltage-clamp software developed in our laboratory. The current recordings were filtered at 1–5 kHz, sampled at >2 kHz, and stored on-line. This system allowed simultaneous application of voltage steps, acquisition of current recordings, and timed application of peptides.

Data analysis. The data points of the dose–response curve were fitted by a four-parameter Hill function (r = 0.9908) using Sigma Plot 2000 software (SPSS, Chicago, IL). A one-way ANOVA in conjunction with a post hoc Tukey–Kramer test was used to assess the significance of dose dependence. A Student’s t test was used (p < 0.05) for comparison of the effect of the differentially modified α2 peptides.

Results
Purification and characterization of VD1 peptides
Profiling of single VD1 and RPD2 neurons by MALDI-TOF-MS revealed a multitude of major molecular ions that are named
molecules A–L (Fig. 1A) (Jimenez et al., 1998). Closer examination of the mass region containing molecules D–H (Fig. 1B) revealed that mass peaks D–H are each accompanied by an extra peak at a mass increment of 16 Da, which are named accordingly as molecules D2, E2, F2, G2, and H2 (Fig. 1B). Previous studies have identified some of the VD1 and RPD2 molecules as bioactive peptides; peptides A, B, and L are uniquely expressed in VD1 and represent the small cardioactive peptides A, B, and the C-terminal peptide, respectively, that are all derived from the small cardioactive peptide precursor (Jimenez et al., 1998). Peptides D, E, and J correspond to the α1, α2, and β peptides, respectively, derived from the alternatively spliced α1 and α2 precursors expressed in both VD1 and RPD2 (Bogerd et al., 1994), with the β peptide contained in both precursors. The remaining mass peaks, i.e., molecules C, D2, E2, F, F2, G, G2, H, H2, I, and K, are present in both VD1 and RPD2 (Jimenez et al., 1998) and do not correspond to previously characterized peptides, suggesting that they may represent novel and/or posttranslationally modified peptides. Moreover, these peptides are present in all single-cell VD1 spectra analyzed to date (up to a few hundred), indicating the robustness of their expression. To gain additional insight into these molecules, we purified them from 800 individually dissected VD1 and RPD2 using a C18 column with increasing acetonitrile concentration in TFA (Fig. 2). Screening of each HPLC fraction with MALDI-TOF-MS allowed us to localize the fractions with the molecules of interest (Figs. 1, 2), which were then subjected to additional purification using reversed-phase HPLC (if necessary), structural characterization, and functional studies.

Table 1 contains a summary of the mass spectrometry and sequencing data of the peptides identified in this study.

MALDI-TOF-MS screening of the HPLC fractions revealed that molecule C [mass-to-charge ratio (m/z) of 1160.1] (Table 1) eluted in fraction 23, and the structure was established by Edman degradation as DRRNLGDTVN. This peptide corresponds to residues 38–47 of the N-terminal of the α peptide prohormones (Bogerd et al., 1993). Peptide material in fractions eluting at 42, 43, 45, 48, 51, 53, 54, and 55 min (Fig. 2) contained molecular ions corresponding to molecules H2 (m/z of 4149.5), H (m/z of 4133.6), G2 (m/z of 3770.8), G (m/z of 3754.8), F2 (m/z of 3391.8), F (m/z of 3375.7), D2 (m/z of 2417.0), and E2 (m/z of 3012.7), respectively, as those detected in the single-cell mass spectrum of VD1 (Fig. 1, Table 1). Some of these peptides were further resolved using reversed-phase HPLC with HCl as a counterion (data not shown). Molecule D2, which occurs at a mass increment of 16 Da to α1 peptide, yielded an α1 peptide sequence (DMYEGLAGRCQHPRNCPGFEN), indicating a modification (see below). Molecules F, G, and H are 379 Da (molecule F), two times 379 Da (molecule G), and three times 379 Da (molecule H) heavier than α2 peptide, suggesting that they represent multiple posttranslationally modified forms of this peptide. Indeed, Edman sequencing of purified molecules F, G, and H all yielded the same amino acid sequence, namely that of the α2 peptide (DMVTTTRIGTGGLAGRCQHHPRNCPGFEN). In addition, F2, G2, and H2, which differ each by 16 Da to F, G, and H, respectively, also yielded an α2 peptide sequence.

To locate the site(s) of the posttranslational modification(s), ~10% of the purified peptides F, G, and H and their +16 Da variants were pooled together and digested with trypsin. The remainder was kept for bioassay experiments. For comparison, synthetic α2 peptide was also digested with trypsin (Fig. 3A). As expected, trypsin cleavage of synthetic α2 peptide yields three fragments with the peptides determined, namely that of the α2 peptide (DMVTTTRIGTGGLAGRCQHHPRNCPGFEN). In addition, F2, G2, and H2, which differ each by 16 Da to F, G, and H, respectively, also yielded an α2 peptide sequence.

To compare the site(s) of the posttranslational modification(s), ~10% of the purified peptides F, G, and H and their +16 Da variants were pooled together and digested with trypsin. The remainder was kept for bioassay experiments. For comparison, synthetic α2 peptide was also digested with trypsin (Fig. 3A). As expected, trypsin cleavage of synthetic α2 peptide yields three fragments with the peptides determined, namely that of the α2 peptide (DMVTTTRIGTGGLAGRCQHHPRNCPGFEN). In addition, F2, G2, and H2, which differ each by 16 Da to F, G, and H, respectively, also yielded an α2 peptide sequence.

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specific peptides (at m/z of 802.1, 825.1, and 1426.3) that cover the full peptide sequence (Fig. 3A, right, B). The molecular ion at m/z of 1426.3 represents tryptic peptides 3 and 4, which are still connected via a disulphide bridge. After trypsin cleavage of the pool of modified \( /H9251 \) peptides F, F2, G, G2, H, and H2, three specific peptides were detected, namely those corresponding to tryptic peptides 2, disulphide-linked peptides 3 \( /H11001 \) and disulphide-linked peptides 3 \( /H11001 \) (\( +16 \) Da) (see below) (Fig. 3A, left). Tryptic peptide 1 (m/z of 825.1) was absent from the spectrum, suggesting that the site of the 379 Da modification resides in the N-terminal part of the \( /H9251 \) peptide. However, the expected posttranslationally modified tryptic peptide 1 was not detected at mass increments of 379 Da of the native peptide. Therefore, under the present MS conditions, the structure of modified tryptic peptide 1 may be unfavorable for the desorption/ionization process and so cannot be effectively detected and further analyzed.

Alternatively, to obtain structural information on the 379 Da modification, we subjected the intact variant \( /H9251 \) peptides to low-energy collisions on a triple–quadrupole instrument. Figure 3C shows the MS/MS spectrum of peptide F2. The low-mass region contains numerous carbohydrate-related fragments, indicating
the presence of N-acetyl hexosamine (HexNac) (m/z of 204.0) and HexNac-related ions (m/z of 186.0, 168.0, 144.3, 137.8, and 126.5). The ions at m/z of 186 and 168 may be explained as water losses. The ion at 144 most likely arises from protonated HexNac by a formal loss of acetic acid, i.e., loss of water and additional loss of the amide-linked acetyl group. The ion at 126 would then correspond to loss of acetyl and loss of two water molecules. The same ions were detected in the low-mass regions of G2 and H2 (data not shown).

The spectrum of the modified /H9251 2 peptides digested with trypsin also indicated that the site of the 16 Da modification on /H9251 2 peptides F2, G2, and H2 is at the C terminal of the peptides. This can be deduced from the presence of an additional peak with a mass corresponding to that of the disulfide-linked tryptic peptides 3 and 4, /H11001 16 Da (Fig. 3A, left). To further characterize this fragment, we subjected it to post-source decay analysis (Fig. 3D), which yielded y ions at m/z of 175 (y1), 272 (y2), 409 (y3), and 546 (y4). Cleavage at the disulfide bond yielded the fragment ions at m/z of 809 and 743. The 16 Da mass increment was located on the C-terminal fragment PGFN (at m/z of 435). No additional sequence information was observed on the PGFN moiety.

Among these four amino acid residues, a modification of 16 Da is most likely located on the proline residue, because it corresponds to the commonly occurring hydroxy-proline residue. Furthermore, during Edman sequencing, an extra peak in addition to proline was also observed, which eluted in a position expected for hydroxy-proline (data not shown).

In summary, the above sequencing and peptide mapping data of the /H9251 peptide family members identified peptide E2 as hydroxy-proline-containing /H9251 peptide, peptide F as monoglycosylated /H9251 peptide, peptide F2 as monoglycosylated HyP-/H9251 peptide, peptide G as diglycosylated /H9251 peptide, peptide G2 as diglycosylated HyP-/H9251 peptide, peptide H as triglycosylated /H9251 peptide, peptide H2 as triglycosylated HyP-/H9251 peptide, and peptide D2 tentatively as HyP-/H9251 peptide.

Edman degradation of the molecule of 6455.5 Da in fraction 113 (Fig. 2) corresponding to molecule K in the VD1 spectrum (Fig. 1) revealed the N-terminal sequence of the /H9252 peptide (GSPYEPSLTGKGDQI...). This suggests that peptide K and the /H9252 peptide have the same peptide backbone and differ by a posttranslational modification of 80 Da, presumably the phosphorylation of a single amino acid residue. To confirm this assumption, peptide K was digested with alkaline phosphatase, which removed the phosphate group from the peptide, resulting...
in a reduced mass of peptide K by 80 Da (Fig. 4A). Using software to predict phosphorylation sites (NetPhos 2.0; http://www.cbs.dtu.dk/services/NetPhos/) (Blom et al., 1999), the θ peptide sequence contains eight predicted phosphorylation sites (Fig. 4B). To locate the site of phosphorylation, peptide K was digested with trypsin, and the tryptic fragments were detected by mass spectrometry. Figure 4C shows that the θ-terminal tryptic peptide is detected at a mass increment of 80 Da, suggesting that either the Thr at position 50 (NetPhos score of 0.623) or the Ser at position 53 (NetPhos score of 0.992) (Fig. 4B) is phosphorylated, with the latter being more likely.

Molecule I at m/z of 6020.2 in the single-cell spectrum of VD1 (Fig. 1A) was located in fraction 67 (m/z of 6019.1) of the HPLC fractions (Fig. 2). Edman sequencing revealed the N-terminal sequence of this peptide as GPGAKKF, which is a unique sequence not present in the National Center for Biotechnology Information (NCBI) database. To obtain internal sequence information, the purified molecule I was digested with endo-lys-c, and the fragments were purified by a C18 column (Fig. 5A). Amino acid sequence analysis of the main fragment at m/z of 1451.2 Da in fraction 36 yielded the sequence AYEDREFVPVPK, which is also not present in the NCBI database. The obtained amino acid sequences indicate that peptide I is a novel peptide. These sequences were subsequently used to synthesize degenerate oligonucleotides for a PCR experiment to characterize the neuron peptide precursor dubbed the peptide I precursor.

Together, the peptide purification and characterization data explained all of the major unidentified molecular ions from the VD1 single-cell spectrum and revealed both novel peptide expression as well as extensive post translational modifications of previously identified peptides.

Characterization of the novel peptide I precursor and cellular localization of the transcript
To elucidate the structure of the precursor encoding peptide θ, a PCR strategy was used on cDNA of a single neuron, i.e., RPD2, which also contains peptide I (Jiménez et al., 1998). Using triplicate reactions of 5‘ RACE and 3‘ RACE, a 298 nt 3‘ untranslated region. The longest open reading frame of 222 nt is flanked by a 113 nt 5‘ leader and a 298 nt 3‘ untranslated region. The proposed initiation codon Met-1 is preceded by an upstream in-frame stop codon. Taking into account that Northern blot analysis detected a transcript of ~1.8 kb (Fig. 5C), we conclude that the peptide I encoding cDNA of 633 nt represents a cDNA that is not full length. Nevertheless, only 5‘ and 3‘ untranslated sequences are missing because the cDNA clone comprises the complete open reading frame. The deduced amino acid sequence of the peptide I cDNA contains a hydrophobic leader sequence that is predicted to be cleaved after Ala-20 (von Heijne, 1983) and a single peptide domain of 53 amino acids. The predicted protonated mass of the peptide I is 6027 Da, whereas the measured protonated mass is 6019.1/6020.2 Da (mass accuracy of ~0.05%). Therefore, most likely the six Cys residues are connected via three disulphide bridges, yielding a protonated molecular weight of 6021 Da (Table 1).

The cellular localization of mRNA of peptide I was studied in sections of the visceral and right parietal ganglia of L. stagnalis by in situ hybridization (Fig. 6A, B, respectively). VD1 and RPD2 neurons were identified on alternating sections by immunocytochemistry using an anti-α peptide antibody (Fig. 6C, D, respectively). The transcripts of peptide I are present in both neurons, confirming the peptide profiles obtained by MALDI-TOF-MS of single VD1 and RPD2 (Jiménez et al., 1998).

Functional connectivity of VD1 to the heart
Previous morphological (Kerkhoven et al., 1991), physiological, pharmacological (Bogerd et al., 1994), and peptide chemical (Jiménez et al., 1998) studies indicated that VD1 and RPD2 may modulate heart beat via the use of its cardioactive peptide transmitters. These in vivo studies do not, however, provide unequivocal evidence that these effects were indeed mediated directly. To further investigate the role of VD1 in modulating heart cell activity, we cocultured VD1, isolated from the brain, with enzymatically dissociated auricle and ventricle cells. First, we confirmed that the dissociated auricle and ventricle muscle fibers were functionally viable. Specifically, muscle contractions were induced by applying α2 peptide exogenously. An example of the effect of α2 peptide on the ventricle membrane potential is shown in Figure 7A (n = 12). Next, the neuron VD1 was isolated and cocultured with the dissociated muscle fibers. After 24 h of coculture, simultaneous intracellular recordings clearly show the presence of synaptic connections between VD1 and the heart muscle fibers (Fig. 7B–F) (n = 17). Spiking activity in VD1 induced by depolarizing current resulted in 1:1 excitatory junction potential (EJP) in the muscle cells (Fig. 7), indicating the presence of a chemical synapse. These synapses were clearly chemical because hyperpolarizing current pulses in VD1 (Fig. 7B) were not registered in the muscle fiber and the synaptic transmission was completely blocked by zero Ca2+ saline (data not shown). Moreover, in the in vitro coculture situation, the effects of VD1 on isolated heart muscle cells were voltage dependent. As the resting membrane potential of the postsynaptic muscle was brought closer to its resting membrane, the VD1-induced EJPs in the target became smaller, albeit generating spikes in the muscle fiber (Fig. 7D–F) (n = 7). To test for the specificity of synapse formation, the heart muscle was paired with RPeD1 (does not innervate heart), and synapses were tested electrophysiologically. We found that, despite extensive physical contacts, RPeD1 failed to establish synapse with the ventricular muscle fiber (Fig. 7G) (n = 5).
Convergence of VD1 peptides onto the same calcium channel in dissociated heart muscle cells

To unequivocally establish the presence of receptors for the different colocalized VD1 and RPD2 peptides on muscle cells of the heart and to study their functional role (including the glycosylated forms of the α2 peptide), we measured the electrophysiological effects of the differentially modified α2 peptides, SCPb, peptide 1, and the phosphorylated β peptide on dissociated heart muscle cells. Under whole-cell voltage-clamp control conditions, muscle cells from the auricle responded to step depolarizations with variable time- and voltage-dependent outward currents. Ventricle muscle cells displayed reproducible responses to step depolarizations. Therefore, the effects of the VD1/RPD2 peptides were compared on dissociated ventricle cells. All VD1/RPD2 peptides tested caused most prominent effects on the inward currents (Figs. 8, 9). Using saline containing barium ions, we observed both a low-voltage-activated (LVA) and a high-voltage-activated (HVA) current in the ventricle cells (Fig. 8A,B). The LVA current was rapidly inactivating and insensitive to dihydropyridines and appeared not to be modulated by the VD1/RPD2 peptides (Fig. 8B). The HVA current was suppressed by the dihydropyridine nicardipine (81 ± 12% block by 1 μM; means ± SEM; n = 4) (data not shown). Therefore, the HVA current can be classified as a typical L-current, in agreement with a previous study in which this current was analyzed in more detail (Yeoman et al., 1999).

The α2 peptide caused a consistent and dose-dependent increase in the amplitude of the L-type current (Fig. 8). We did not observe any changes in kinetic properties or voltage dependence. The dose dependence was determined for the range of 10⁻⁸ to 10⁻⁶ M (Fig. 8C). Between 10⁻⁷ and 10⁻⁶ M, the dose–effect curve was very steep, with a maximal effect of the barium current observed at 5 × 10⁻⁶ M (Fig. 8C). The dose–response curve can be described by a four-parametric Hill function (r = 0.9908), and the effect of concentration is significant (one-way ANOVA, p = 0.0037). Because saturation of the response did not yet occur within this concentration range, the EC₅₀ cannot be determined with certainty but is estimated to be ~2 × 10⁻⁷ M. Responses to higher concentration of α2 peptide could not be recorded because of the contractions of the muscle fibers. Next, we investigated whether the posttranslationally modified α2 peptides mimicked the effect of the unmodified α2 peptide. Like unmodified α2 peptide, the different modified α2 peptides (HyP-α2 peptide E2, monoglycosylated HyP-α2 peptide F2, diglycosylated HyP-α2 peptide G2, and triglycosylated HyP-α2 peptide H2) all increased the amplitude of the HVA-barium current without altering the kinetics or voltage dependence. An example of the response of the monoglycosylated HyP-α2 peptide F2 is shown in Figure 9A. Interestingly, the modified α2 peptides yielded larger responses than the α2 peptide itself (Student’s t test, p < 0.05). Comparing the effect of the same concentration (2.5 × 10⁻⁶ M) demonstrates that the differentially glycosylated HyP-α2 peptides F2, G2, and H2 are equipotent and approximately twofold more potent than HyP-α2 peptide E2 (Student’s t test, p < 0.05), which in turn is approximately twofold more potent than unmodified α2 peptide (Student’s t test, p < 0.05) (Fig. 8D) (n = 5 for each peptide).

Testing the effect of SCPb on the isolated barium currents showed a striking effect of SCPb on the HVA-barium current (Fig. 9B). The threshold concentration for this effect was 1 μM (n = 4). At 10 μM SCPb, the increase of this current amounted to 300% (n = 3)
Peptide diversity generated by posttranslational modification

We characterized multiple posttranslational modifications of the α and β peptides derived from the α peptide precursors. The α1 and α2 peptides are highly similar and differ only in the N terminal, in which the Tyr-Glu residues at positions 3 and 4 in α1 are replaced by Val-Thr-Thr-Arg-Ile-Gly-Thr-Gly in α2. Using tryptic peptide mapping and tandem MS analysis, we demonstrated the presence of three differentially glycosylated α2 peptides, most likely differentially modified on the three Thr residues in the N terminal. Absence of these residues in the α1 peptide explains the lack of glycosylation of α1. Tandem MS of modified α2 peptides demonstrated the presence of HexNac, which is commonly found in O-linked sugars attached to Ser and Thr residues. The carbohydrate of 379 Da may consist of HexNac-hexuronic acid, because this is the only structure predicted by the GlycoMod tool. Glycosylation of protein hormones has been implicated in protein folding, stability (Baudys et al., 1995; Kihlberg et al., 1995; Haneda et al., 2001), intracellular transport, enhanced receptor binding, and biological potency (Flack et al., 1994; Nissen, 1994; Furuhashi et al., 1995). However, few reports exist on the presence and functions of naturally occurring glycosylated peptides (Ivell et al., 1981; Nakakura et al., 1992). Therefore, in addition to their increased signaling properties, the differentially glycosylated α2 peptides may have several other features than the unmodified α2 peptide.

Besides differential glycosylation, the Pro residue 25 of each α2 peptide is differentially modified into a HyP variant. Hydroxylation is a common posttranslational modification of Pro residues that occur in collagens (McCormack et al., 1994; Gaill et al., 1995). Because the HyP residues are especially important for noncovalent interactions between proteins, it may be speculated that the higher potency of the HyP-containing α2 peptide is caused by increased receptor binding.

Edman degradation in conjunction with phosphatase treatment and tryptic peptide mass mapping demonstrated that the majority of...
β peptide is phosphorylated. In the C-terminal part of the β peptide, the casein kinase consensus site (i.e., Ser/Thr-X-X-Glu/Asp-) at the Ser residue at position 53 is most likely phosphorylated (NetPhos score of 0.992). Casein kinases are present at the luminal side of the Golgi apparatus (Vegh and Varro, 1994), and, therefore, they may phosphorylate the β peptide during passage through the Golgi apparatus.

**VD1 modulates heartbeat via a direct neuromuscular connection**

*In vitro* experiments in this study support the idea that VD1 modulates heartbeat via the use of its peptides. Coculture experiments of VD1 and RPDeD1 and dissociated heart auricle and ventricle cells show that excitatory synapses are selectively formed in *vitro* between VD1 and heart cells. The successful synapse formation between VD1 and heart muscle cells is in line with previous immunocytochemical findings (Kerkhoven et al., 1991) that showed the presence of numerous α1 peptide-positive varicosities terminating on auricle trabecular muscle fibers, which are reminiscent of synapses. Moreover, VD1 and RPDeD2 peptides derived from all prohormones can be detected in biopsies of heart tissue using mass spectrometry (Jimenez et al., 1998). However, VD1 is not the only neuron that sends projections to the heart. Several peptidergic and nonpeptidergic heart motor neurons have been described in the past (Buckett et al., 1990). Recently, the possible role of this current was investigated in detail in heart ventricle cells of *Lymnaea* by Yeoman and Benjamin (1999). They showed that the L-type current is important for pacemaking in generating action potentials and in providing calcium for contraction-excitation coupling. Therefore, modulation of the L-type current by VD1 peptides may alter the myogenic beating rate of the heart.

**Convergent and divergent neuromodulation**

The frequent occurrence of colocalized neurotransmitters suggests that the combination of transmitters released by a neuron attributes specificity to its actions (Kupfermann, 1991; Brezina and Weiss, 1997). The results reported here are interesting and perhaps surprising for several reasons: first, because VD1 and RPDeD2 cosynthesize such a large array of chemically diverse peptides; and second, because most of these peptides converge onto a single type of calcium channel to modulate a single type of Ca current. To the best of our knowledge, this study is the first to demonstrate that single neurons exhibit such a complex pattern of peptide gene expression, precursor processing, and modifications. The present results indicate that neuropeptides derived from three different genes (α peptides, SCPS, and LyCCAP) can exhibit activation of the same effector protein, the L-type HVA calcium channel. This remarkable degree of convergence in neuromodulation was also observed in the crustacean somatogastric ganglion (Swensen and Marder, 2000; Nusbaum et al., 2001). Convergence might prevent overmodulation by multiple modulatory inputs.

In addition to their convergent effects, it is feasible that α peptides and SCPs (and possibly LyCCAP) may also exert divergent effects. Based on the complexity of the whole auricle response and the differential effects of α2 peptide and SCPS application (Jimenez et al., 1998), it is likely that these peptides influence more than one channel type in *Lymnaea* heart muscle cells. This notion is consistent with the observation that the synaptic effects of VD1 in the coculture were evident even at varying hyperpolarizing potentials, suggesting the involvement of other ion channels, in addition to the HVA calcium channel. The potential discrepancy between sharp electrode and patch-clamp recordings may also be attributable to differential sensitivities of

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**Figure 9.** Selective increase of the HVA-barium current in ventricle cells by monoglycosylated HyP-α2 peptide F2, SCPh, and peptide I/LyCCAP. A–C. Family of current traces at −40 to +20 mV (10 mV steps) in the absence (left) and presence (middle) of monoglycosylated HyP-α2 peptide F2 (A), SCPh (B), and peptide I/LyCCAP (C). Voltage step protocols were applied as indicated in the right panel. Currents of representative ventricle cells are shown. Application of each peptide caused a strong increase in the HVA current.

**Differentially modified α2 peptides, SCPh, and LyCCAP converge onto calcium channels in dissociated heart ventricle cells**

We compared the effects of selected peptides of VD1 and RPDeD2 on membrane currents of heart ventricle muscle fibers, because they displayed more reproducible voltage-activated currents than the auricle fibers. The voltage-clamp recordings showed that a dihydropyridine-sensitive HVA current belonging to the L-type family is modulated by all tested variant α2 peptides, but significant quantitative differences were observed between them, with the posttranslationally modified forms being more potent. The SCPh peptide modulated the same HVA L-type calcium current as the α2 peptides, but the threshold of the response was higher than that for α2 and the modulatory effect reversed very slowly. Interestingly, the novel peptide LyCCAP also activated this current, with kinetics similar to the α2 peptides. Together, our voltage-clamp experiments show that seven peptides, representing three peptide families, exhibit convergent activation of a HVA L-type calcium current. Interestingly, this current is also modulated by FMRFamide (Brezden et al., 1999), from heart motor neurons (Buckett et al., 1990). Recently, the possible role of this current was investigated in detail in heart ventricle cells of *Lymnaea* by Yeoman and Benjamin (1999). They showed that the L-type current is important for pacemaking in generating action potentials and in providing calcium for contraction-excitation coupling. Therefore, modulation of the L-type current by VD1 peptides may alter the myogenic beating rate of the heart.
Functionality of Peptide Diversity in a Single Neuron


