Invertebrate Vasopressin/Oxytocin Homologs

CHARACTERIZATION OF PEPTIDES FROM CONUS GEOGRAPHUS AND CONUS STRIATUS VENOMS*

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From the **Department of Biology and *Howard Hughes Medical Institute, University of Utah, Salt Lake City, Utah 84112 and **Department of Biochemistry, College of Medicine and Marine Science Institute, University of the Philippines, Metro Manila, Philippines

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The vasopressin-oxytocin family of peptides is of very ancient lineage, found in organisms as diverse as hydra and man. Although these peptides have been intensively studied in vertebrates, the presumably more extensive invertebrate series is defined primarily by immunological methods. In this report, we describe the purification and structures of two peptides of the vasopressin-oxytocin family from molluscs ("Conopressins"), which were found in the venom of fish-hunting marine snails of the genus Conus. The biological activity observed when the two snail peptides are injected intracerebrally into mice is very similar to that elicited by the vertebrate neurohypophyseal hormones and presumably reflects their actions upon a common receptor in the brain. The sequences of the purified peptides reveal unique features not found in the vertebrate peptide series, most notably an additional positive charge. These are the first members of the invertebrate series of the vasopressin-oxytocin family to be characterized biochemically.

The sequences of these peptides are from: Conus geographus venom, Lys-conopressin-G, Cys-Phe-Ile-Arg-Asn-Cys-Pro-Lys-Gly-NH2; and from Conus striatus venom, Arg-conopressin-S, Cys-Ile-Ile-Arg-Asn-Cys-Pro-Arg-Gly-NH2.

The vasopressin-oxytocin peptides were first characterized as neurohypophyseal hormones in mammals. Their biological activity was defined 30 years before (1) and their structures were elucidated in the 1950s by Du Vigneaud and his coworkers (2). In recent years, the proposed physiological role and phylogenetic distribution of this family of peptides has been greatly expanded. These peptides are released in tissues other than the pituitary in mammals, and vasopressin appears to play a role as a neurotransmitter or cotransmitter in mammalian brain. In addition, compelling evidence has accumulated for the presence of these peptides in a number of invertebrate systems, including insects (3-5) and molluscs (6, 7). This evidence is primarily immunological, most often immunocytochemical, staining with antipeptide antibodies.

Recently Grimmeltikhuizen et al. (8) reported oxytocin/vasopressin-like immunoreactivity in Hydra. This coelenterate has one of the simplest nervous systems known: although synapses and secretory vesicles are present, small molecule neurotransmitters such as acetylcholine and catecholamines appear to be absent (9, 10). The high concentration of peptides in the neurons of Hydra has led to the intriguing suggestion that in primordial nervous systems the first neurotransmitters may have been exclusively peptides (10). Thus, included in the set of putative Hydra neurotransmitters is a peptide of the vasopressin-oxytocin family (8).

These results suggest that these peptides have an evolutionary lineage that dates back ~108 years. To date, only the vertebrate members of this peptide family have been purified and fully characterized, leaving the relationship between them and the ancient line of invertebrate peptides undefined. In this report we describe the first purification and sequence analysis of invertebrate vasopressin-oxytocin peptides. These peptides have been purified from an unexpected source, the venom of the fish-hunting cone snails (Conus). We propose the generic term conopressin for all vasopressin-oxytocin homologs found in Conus. The Conus venoms have proven to be rich sources of biologically active peptides (11). Peptide toxins which inhibit calcium channels (12, 13), sodium channels (14, 15), and the acetylcholine receptor (16, 17), as well as a peptide which induces a sleep-like state in mice (18, 19), have already been described.

MATERIALS AND METHODS

Venoms—Conus geographus and Conus striatus were collected around the island of Marinduque, Philippines. Venom ducts were dissected from freshly killed snails, and the crude venom was subjected to preliminary purification as described previously (12). Isolation and Bioassay of Peptides—Crude venom extract from C. geographus was fractionated on a Sephadex G-25 column as previously described (12) and the "scratcher peptide" was subsequently purified from the third major protein peak which was designated as Peak C by Olivera et al. (12). Reverse-phase HPLC1 was used to separate different peptide fractions with the appropriate biological activity as shown in Fig. 1, and the peptides were assayed by intracerebral injection into mice as previously described (12). The scratches of C. striatus venom was purified essentially in the same manner.

Amino Acid Analysis—Peptide samples were hydrolyzed in vacuo in 6 N HCl, 1% phenol for 18 h at 105 °C. For C. striatus, amino acid analysis was done by reverse-phase HPLC of phenyl-thiocarbamyl derivatives (20).

Peptide Sequencing—Reduction and carboxymethylation was carried out as previously described (16) except that a volatile buffer (0.02 M N-ethylmorpholine) was used. Sequencing of reduced and carboxymethylated peptides was carried out by sequential Edman degradation in a Beckman 890D spinning cup sequenator, using 0.1 M Quadrol buffer and Polybrene carrier (21). Phenylthiohydantoin-derivatives were identified by HPLC on a Hewlett-Packard 1041B instrument using an Ultrasphere ODS column (0.46 x 15 cm, 5 μ particle size, endcapped) eluted with a gradient of acetonitrile in 0.05 M sodium acetate, pH 4.5 (16).

1 The abbreviation used is: HPLC, high performance liquid chromatography.

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Fig. 1. Purification of the scratcher peptide from Conus geographus venom. One gram of crude lyophilized venom was extracted and fractionated on a Sephadex G-25 column as previously described (12). The first 6 of 21 successive fractions containing “scratcher activity” (now known as the calcium channel blockers, w-conotoxins) were pooled and chromatographed on a Vydac C18 semi-preparative column eluted with a gradient of acetonitrile in 0.1% trifluoroacetic acid (A). The peak indicated by an arrow elicted both shaking and scratching activities. Chromatography of material from this peak on a Vydac C18 analytical column eluted with a gradient of acetonitrile in 0.05% heptafluorobutyric acid (B) separated the “scratcher” from the “shaker” activity. The scratcher peak, when rechromatographed under the same conditions as in B, appears homogeneous (C). The absorbance plotted is at 214 nm. Dashed lines indicate the gradient of acetonitrile.

RESULTS

Assay and Purification of Conopressins—Intracerebral injection of these peptides at levels of >0.5 nmol causes mice to scratch themselves within a few minutes. The “scratcher” activity is masked in crude venom by the actions of other toxins, particularly the lethal ones. The first conopressin was detected in Conus venom fractions from Sephadex G-25 filtration, where its elution profile overlapped extensively with that of the Ca2+ channel toxins (w-conotoxins). On reverse-phase HPLC in 0.1% trifluoroacetic acid/acetonitrile (Fig. 1A), the C. geographus activity was associated with the leading edge of the major peak of w-conotoxin GVIA or the “shaker peptide.” On rerunning in 0.05% heptafluorobutyric acid/acetonitrile (Fig. 1B) a clean separation was obtained, the scratching effects being elicited only by material from a minor peak. This material, which was apparently homogeneous, was then analyzed by sequencing. Purification of conopressin from C. striatus venom followed essentially the same steps.

Peptide Structural Analysis—The peptides were reduced and carboxymethylated as described under “Materials and Methods” (16) and then applied to the sequenator cup. In both cases the sequences were unambiguous, although there were heavy losses of the COOH-terminal Gly, a common experience with amided peptides. Results of the analysis for both conopressins are given in Table I, and sequences of the two conopressins are shown in Table II.

Amino acid analysis of conopressin S was in agreement with the sequence, though the Ile-Ile dipeptide was only then analyzed by sequencing. Purification of conopressin from both cases the sequences were unambiguous, although there presence of a basic residue, Arg, at position 8 is strongly correlated with conopressin activity in vertebrate hormones and in hundreds of synthetic analogues (27). Since both of the Conus peptides share this feature, we propose the name “conopressin” for the group. The presence of Ile at position 3 could be used to argue in favor of “conotocin,” but this is confusingly similar to the well-established “conotoxin.” Both peptides have neutral hydrophobic residues at position 2, Phe in conopressin G and Ile in conopressin S. All the vertebrate peptides have Tyr at this position, with the sole exception of phenypressin from marsupials (see Table II). An aromatic residue at position 2 is believed to be specifically required for interaction with neurophysin in vertebrate systems (23). The presence of Ile in conopressin S implies either that there are no endogenous neurophysins for conopressin(s) or that Conus neurophysins are more tolerant of substitutions at position 2.

The unique feature shared by both snail peptides is the presence of a basic residue, Arg, at position 4. This gives them a net charge of +3, higher than that of the vasopressins (+2) or oxytocins (+1). Whether this contributes unique biological properties will be of great interest. An arginine vasopressin analogue with this substitution has been synthesized (29) and shown to be similar to vasopressin in its antidiuretic and pressor activities.

The structural similarity with vasopressin and other neuropehysel hormone is obvious from Table II. Thus, the peptide from C. geographus is designated Lys-conopressin-G and the peptide from C. striatus is designated Arg-conopressin-S. This nomenclature follows both the commonly used system for vasopressins, as well as for Conus peptides (22).

Biological Effects of Intracerebral Injection—The biological responses elicited by intracerebral injection of conopressin G were compared with those elicited by Arg-vasopressin and oxytocin (Table III). They were similar in all cases, with conopressin being intermediate in potency between the vertebrate hormones. The scratching response is dose-dependent with respect to time of onset and duration of symptoms. At very low doses, the mice have the appearance of grooming their faces. The symptomatology elicited by these peptides upon intracerebral injection into mice is highly characteristic (23-25), presumably diagnostic of a specific central nervous system receptor. This may be homologous to the receptor which controls the stereotypic “flanking marking” response characterized in Golden hamsters (26); scratching is one feature of this territorial behavior.

DISCUSSION

The peptides isolated from C. geographus and C. striatus are clearly members of the vasopressin-oxytocin family (Tables II and III), yet they have distinctive features. A basic residue, Lys or Arg, at position 8 is strongly correlated with pressor activity in vertebrate hormones and in hundreds of synthetic analogues (27). Since both of the Conus peptides share this feature, we propose the name “conopressin” for the group. The presence of Ile at position 3 could be used to argue in favor of “conotocin,” but this is confusingly similar to the well-established “conotoxin.” Both peptides have neutral hydrophobic residues at position 2, Phe in conopressin G and Ile in conopressin S. All the vertebrate peptides have Tyr at this position, with the sole exception of phenypressin from marsupials (see Table II). An aromatic residue at position 2 is believed to be specifically required for interaction with neurophysin in vertebrate systems (23). The presence of Ile in conopressin S implies either that there are no endogenous neurophysins for conopressin(s) or that Conus neurophysins are more tolerant of substitutions at position 2.

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Peptide | Amino acid sequence | Charge | Animal group
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A. Piscivorous Conus scratcher peptides  
Lys-conopressin-G | Cys-Phe-Ile-Arg-Asn-Cys-Pro-Cys-Leu-Gly-(NH₂) | +3 | Mollusc  
Arg-conopressin-S | Cys-Ile-Arg-Asn-Cys-Pro-Arg-Gly-(NH₂) | +3 | Mollusc
B. Vertebrate neurohypophysal hormones  
Lys-vasopressin | Cys-Tyr-Phe-Gln-Asn-Cys-Pro-Cys-Leu-Gly-(NH₂) | +2 | Suiformes, peccaries, wart hogs, and hippopotami  
Arg-vasopressin | Cys-Tyr-Phe-Gln-Asn-Cys-Pro-Arg-Gly-(NH₂) | +2 | Marsupials  
Phenopressin | Cys-Phe-Phe-Gln-Asn-Cys-Pro-Cys-Leu-Gly-(NH₂) | +2 | Birds, reptiles, amphibians, bony fishes, and possibly cartilaginous fishes  
Arg-vasotocin | Cys-Tyr-Ile-Gln-Asn-Cys-Pro-Leu-Gly-(NH₂) | +1 | Mammals  
Oxytocin | Cys-Tyr-Ile-Asn-Cys-Pro-Cys-Pro-Glu-Leu-Gly-(NH₂) | +1 | Cartilaginous fishes (ray)
Gluotocin | Cys-Tyr-Ile-Ser-Asn-Cys-Pro-Gln-Gly-(NH₂) | +1 | Mammals
Ancestral vertebrate peptide | Cys-Tyr-X-X-Asn-Cys-Pro-X-Gly-(NH₂) | +2 | Mammals

We propose to call all peptides of this homologous peptide set "conopressins." The nomenclature is adapted from that used for conotoxins, i.e. G (for C. geographus), S (for C. striatus) (22). Thus, the first conopressin from Conus magus (if different in sequence) would be conopressin M.

Adapted from Archer (31) and Hadley (32). The cysteine residues form internal disulfide bonds.

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**REFERENCES**

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