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Cloning of the first cDNA encoding a putative CCRFamide precursor – identification of the brain, eyestalk ganglia, and cardiac ganglion as sites of CCRFamide expression in the American lobster, *Homarus americanus*

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Abstract

Over the past decade, many new peptide families have been identified via in silico analyses of genomic and transcriptomic datasets. While various molecular and biochemical methods have confirmed the existence of some of these new groups, others remain in silico discoveries of computationally assembled sequences only. An example of the latter are the CCRFamides, named for the predicted presence of two pairs of disulfide bonded cysteine residues and an amidated arginine-phenylalanine carboxyl-terminus in family members, which have been identified from annelid, molluscan, and arthropod genomes/transcriptomes, but for which no precursor proteinencoding cDNAs have been cloned. Using routine transcriptome mining methods, we identified four Homarus americanus (American lobster) CCRFamide transcripts that share high sequence identity across the predicted open reading frames but more limited conservation in their 5' terminal ends, suggesting the *Homarus* gene undergoes alternative splicing. RT-PCR profiling using primers designed to amplify an internal fragment common to all of the transcripts revealed expression in the supraoesophageal ganglion (brain), eyestalk ganglia, and cardiac ganglion. Variant specific profiling revealed a similar profile for variant 1, eyestalk ganglia specific expression of variant 2, and an absence of variant 3 expression in the cDNAs examined. The broad distribution of CCRFamide transcript expression in the H. americanus nervous system suggests a potential role as a locally released and/or circulating neuropeptide. This is the first report of the cloning of a CCRFamide-encoding cDNA from any species, and as such, provides the first non-in silico support for the existence of this invertebrate peptide family.

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Keywords

Decapoda; Homarus americanus, in silico transcriptome mining; RT-PCR; nervous system

1. Introduction

Recent advances in high-throughput nucleotide sequencing and in the programs used for assembling the resultant data have led to an explosion in the availability and depth of genomic and transcriptomic datasets for many organisms. *In silico* analyses of these assemblies have provided evidence supporting the existence of a number of previously unknown peptide families (*e.g.*, Christie, 2014; Conzelmann et al., 2013; Dircksen et al., 2011; Veenstra et al., 2012). While the presence of some of these groups has been confirmed via molecular cloning, mass spectrometry, and/or biochemical isolation and sequencing (*e.g.*, Bao et al., 2015; Dickinson et al., 2019), others remain *in silico* discoveries only.

The CCRFamides form one such family, which was, originally identified via the mining of an annelid, *Platynereis dumerilii*, transcriptome and named for the predicted presence of two disulfide bonded cysteine pairs and the carboxyl (C)-terminus –RFamide in the putative mature peptide (Conzelmann et al., 2013). *In silico* analyses suggest that the CCRFamide family is broadly conserved in invertebrates. In addition to the Annelida, genome and transcriptome mining has identified putative CCRFamide-like peptide-encoding sequences in multiple members of the Mollusca and Arthropoda (Conzelmann et al., 2013; Nguyen et al., 2018; Oliphant et al., 2018; Stewart et al., 2014; Veenstra, 2010, 2015, 2016a, 2016b). Despite their apparent broad phylogenetic conservation, no CCRFamide precursor has been cloned from any species, and no biochemical or mass spectral data documenting the existence of any putative mature CCRFamide peptide are published (Conzelmann et al., 2013; Nguyen et al., 2018; Oliphant et al., 2018; Stewart et al., 2014; Veenstra, 2010, 2015, 2016a, 2016b).

In the study presented here, we report the cloning of the first cDNA encoding a putative CCRFamide precursor. Using primers designed from a transcriptome-derived sequence, the complete open reading frame of an American lobster, *Homarus americanus*, CCRFamide precursor was cloned from supraoesophageal ganglion (brain) cDNAs. Using RT-PCR, we show that in addition to the brain, the prepro-CCRFamide is also expressed in at least two other portions of the lobster nervous system, specifically the eyestalk ganglia, the location of the neuroendocrine X-organ-sinus gland system (*e.g.*, Christie, 2011), and the cardiac ganglion (CG), which drives the rhythmic contractions of the heart (*e.g.*, Cooke, 2002). We further provide evidence consistent with tissue specific alternative splicing of the CCRFamide transcript.

2. Materials and methods

2.1. Transcriptome mining and prediction of mature peptide structure

2.1.1. In silico *transcriptome mining*—Searches for putative *H. americanus* CCRFamide-encoding transcripts were conducted using a well-established protocol (*e.g.*,

Christie et al., 2015, 2017). Specifically, the database of the online program tblastn (National Center for Biotechnology Information, Bethesda, MD; http://blast.ncbi.nlm.nih.gov/Blast.cgi) was set to Transcriptome Shotgun Assembly (TSA) and restricted to data from *H. americanus* nervous system-specific transcriptomes, *i.e.*, BioProject Nos. PRJNA300643 (brain, abdominal nerve cord, CG and stomatogastric nervous system; Northcutt et al., 2016), PRJNA338672 (eyestalk ganglia-specific; Christie et al., 2017), PRJNA379629 (brain-specific; Christie et al., 2018a) and PRJNA412549 (CG-specific; Christie et al., 2018b). A CCRFamide precursor predicted from crayfish, *Procambarus clarkii*, transcriptomic data (Veenstra, 2015) was used as the query sequence for all BLAST searches. In total, four transcriptomic reads (GFDA01062303, GEBG01038526, GFUC01020658 and GFUC01020659) were identified.

2.1.2. Peptide structural prediction—The putative mature structures of a H. americanus CCRFamide isoform and several CCRFamide precursor-related peptides were predicted using a well-established workflow (e.g. Christie et al., 2015, 2017). In brief, all hits returned by a given BLAST search were translated using the ExPASy Translate tool (http://web.expasy.org/translate/) and assessed for completeness. Proteins listed as fulllength exhibit a start methionine and are flanked on their C-terminus by a stop codon, while those listed as C-terminal fragments are missing the start methionine. Next, each precursor was assessed for the presence of a signal peptide using SignalP 3.0 (http://www.cbs.dtu.dk/ services/SignalP/; Bendtsen et al., 2004). Prohormone cleavage sites were identified based on information presented in Veenstra (2000). The sulfation state of tyrosine residues was predicted using Sulfinator (https://www.expasy.org/sulfinator/; Monigatti et al., 2002), while disulfide bonding between cysteine residues was assessed using DiANNA (http:// clavius.bc.edu/~clotelab/DiANNA/; Ferrè and Clote, 2005). C-terminal amidation was predicted by homology to known arthropod peptides. Protein and peptide alignments were done using either the online program MAFFT version 7 (http://mafft.cbrc.jp/alignment/ software/; Katoh and Standley, 2013) or MUSCLE (Edgar, 2004) as implemented in Geneious Prime 2020.1.2 (https://www.geneious.com).

2.2. Molecular cloning

Total RNAs were purified from freshly dissected individual *H. americanus* brains using TRI Reagent (Life Technologies, Carlsbad, CA) and Direct-zol RNA MiniPrep spin columns (Zymo Research, Irvine, CA, USA) as described in Christie et al. (2018a). First-strand cDNAs were generated from 500 ng of DNase I-treated total RNA using Superscript III Reverse Transcriptase (Life Technologies) and custom made random pentadecamers (IDT, San Diego, CA, USA). The complete open reading frame (ORF) for the putative *H. americanus* CCRFamide precursor was PCR amplified using primers (sense – 5'-ATGGTGAGTCGTGTGGGC; antisense – 5'-CTAGTCGAGGAGTAGTTCATAACA) designed to encompass the consensus predicted start and stop sites (771 bp) in conjunction with Sapphire Amp Fast PCR Master Mix (Takara Bio USA Inc., Mountain View, CA). PCR was performed on a Biometra TRIO multiblock thermocycler (Biometra, Göttingen, Germany) in a 20-μL reaction volume containing 0.5 μL cDNA template and 0.2 μM of each primer. The thermocycler conditions consisted of: 95°C for 2 min followed by 40 cycles at 95°C for 20 s, 56°C for 20 s, 72°C for 30 s, and a final extension at 72°C for 5 min. The

resulting products were separated on a 1.5% agarose gel using a tris/acetate/EDTA buffer system with SYBR Safe (Life Technologies). The reaction products were subcloned into pCR2.1-TOPO TA (Life Technologies) and sequenced at the Arizona State University DNA Core Laboratory (Tempe, AZ).

2.3. RT-PCR expression profiling

To examine the expression of CCRFamide precursor-encoding transcripts, RNAs were purified from four biological replicates of *H. americanus* brain (one brain per replicate), eyestalk ganglia (one eyestalk pair per replicate), CG (ten ganglia per replicate), and cardiac muscle (~25 mg of cardiac muscle per replicate), as described in several recent publications (Christie et al., 2017; 2018a, 2018b). cDNAs were generated as described above from ~100 ng (CG) or 500 ng (brain, eyestalk ganglia, and cardiac muscle) of DNase I-treated total RNAs. RT-PCR was performed in 20-μL reaction volumes containing either 0.5 μL of brain, eyestalk ganglia, or cardiac muscle cDNA templates or 1 µL of CG-derived cDNAs. Primers (sense – 5'- GTCTTGGCCAATCACAGC and antisense – 5'-GTGAGGACTTCGACGACT) were designed to generate a 528-bp internal fragment (nt 64-591) common to all of the *H. americanus* CCRFamide precursor transcripts. To assess the expression of potential transcript splice variants, the consensus ORF antisense primer described above was used in conjunction with specific sense primers: variant 1 (GFDA01062303) - 5'-ACTTGACTTTCGCACTTCTC; variant 2 (GEBG01038526) - 5'-TGAAGGCGTGGGGGTGA; and variant 3 (GFUC01020659) - 5'-CCTCTCACTTCACAAATATTAC. Primers (sense - 5'- GGTCGTACCACCGGTATT and antisense – 5'- CATCCTGTCGGCAATTCC) were also designed to amplify nt 439-942 of H. americanus actin (FJ217215). PCR was performed with each of the replicated cDNA sets and used thermocycler conditions identical to those described above. The resulting products were visualized on 1.5% agarose gels with images obtained using an AlphaImager Gel Documentation System (ProteinSimple, San Jose, CA) and processed in Photoshop CS6 v13.0 (Adobe Systems Inc., San Jose, CA).

3. Results

3.1. In silico identification of H. americanus CCRFamide-encoding transcripts and prediction of CCRFamide precursor proteins and peptides

To identify putative *H. americanus* CCRFamide precursor-encoding transcripts, the sequence of a previously described crayfish, *P. clarkii*, prepro-CCRFamide (Veenstra, 2015) was used to search nervous system-specific TSA datasets for sequences encoding putative homologs. The assemblies searched included one produced using multiple nervous system regions as the source of starting RNA (Northcutt et al., 2016), and three region-specific transcriptomes, *i.e.*, those for the eyestalk ganglia (Christie et al., 2017), the brain (Christie et al., 2018a), and the CG (Christie et al., 2018b). While no transcripts encoding putative CCRFamide precursors were found in the CG assembly, four were identified from the other transcriptomes, one each from the eyestalk ganglia and mixed nervous system region datasets (Accession Nos. GFDA01062303 and GEBG01038526, respectively) and two from the brain transcriptome (Accession Nos. GFUC01020658 and GFUC01020659). Although the transcripts are largely identical across a significant portion of the open reading frame

(Fig. S1), the regions immediately upstream of the putative start codons differ (Fig. 1A). Only the **GFDA01062303** transcript encodes a full-length precursor peptide with a predicted signal peptide; the other transcripts either lack a defined start site (*i.e.* first in-frame ATG downstream of a stop codon) or have an alternative start relative to **GFDA01062303**. This apparent transcript specific variation could indicate alternative splicing of the *H. americanus* CCRFamide gene or may reflect assembly issues that arise with repetitive sequences common to untranslated regions (Lima et al., 2017). Significant sequence variation specific to the 5'terminal ends of CCRFamide-encoding transcripts (GGLH01033349.1, GGLH01033500.1) in *Penaeus monodon* (giant tiger prawn), however, supports the presence of a conserved splicing mechanism.

Translation of the identified transcripts revealed three distinct putative CCRFamide precursors: two full-length proteins, one from **GFDA01062303** (*H. americanus* prepro-CCRFamide variant 1 [Homam-prepro-CCRFa-v1; 256 amino acids]) and the other from **GEBG01038526** (*H. americanus* prepro-CCRFamide variant 2 [Homam-prepro-CCRFa-v2; 245 amino acids]), and one C-terminal fragment (*H. americanus* prepro-CCRFamide variant 3 [Homam-prepro-CCRFa-v3; 262 amino acids]), which was deduced from both **GFUC01020658** and **GFUC01020659**. The three CCRFamide precursors differ only in the region of their signal peptides and at one conservatively substituted residue (Fig. 1B). A total of six peptides were predicted from the collective set of preprohormones, including one isoform of CCRFamide (Table 1). The putative mature CCRFamide is 36 amino acids in length and is predicted to be C-terminally amidated with disulfide bridging between its first and fifth, and second and fourth cysteines (Table 1). A pair of sulfated tyrosine residues is predicted for CCRFamide precursor-related peptides 2a and 2b (Table 1), which are derived from Homam-prepro-CCRFa-v1/3 and Homam-prepro-CCRFa-v2, respectively (Fig. 1B).

Multiple sequence alignment of HomamCCRFa with CCRFamides from diverse arthropod species reveals significant C-terminal conservation, with most peptides exhibiting a characteristic P-M/V/I-R-F-amide tetrapeptide motif and four equally spaced Cys residues (Fig. 1C). Interestingly, decapod specific atypical amino acid usage was observed with a charged Arg residue at position -7 (relative to the C-terminus) rather than the polar or small hydrophobic residue present in species from other Orders. In addition, the two Nephropidae in our analysis were distinguished by an atypical Ala at position -20, which, in sequences other than the *Stegodyphus mimosarum* A (African social velvet spider) CCRFamide, is occupied by amino acids with larger sidechains (*e.g.* Phe, Ser, Gln, Asn, or His).

3.2. Cloning of a CCRFamide-encoding cDNA from the brain of Homarus americanus

While a number of studies, including the one presented here, have provided *in silico* evidence for the existence of CCRFamides in invertebrates (Conzelmann et al., 2013; Nguyen et al., 2018; Oliphant et al., 2018; Stewart et al., 2014; Veenstra, 2010, 2015, 2016a, 2016b), there has not yet been confirmation of the CCRFamide family using methodologies such as molecular cloning. To generate a complete ORF for the longest of the putative full-length *H. americanus* CCRFamide precursors, *i.e.*, Homam-prepro-CCRFa-v1, primers were designed based on the **GFDA01062303** transcript and used to amplify products from brain cDNAs. Sanger sequencing of the resultant amplicons showed them to be >99% identical in

nucleotide sequence to that of the transcript used for primer design. The protein deduced from the cloned cDNA, which was submitted to GenBank under **Accession No.** MN661244, is identical in amino acid sequence to that of Homam-prepro-CCRFa-v1.

3.3. Expression profiling of CCRFamide expression in select portions of the Homarus americanus nervous system

Transcriptomic data suggest that the CCRFamide precursor is expressed in the eyestalk ganglia, in addition to the brain, of *H. americanus*. To confirm this expression, and to further investigate whether or not this peptide might also be present in the CG, an offshoot of the central nervous system that controls the rhythmic contractions of the heart (*e.g.*, Cooke, 2002), RT-PCR profiling of these portions of the nervous system was conducted; cardiac muscle was used as a negative control for these analyses. PCR products were consistently amplified from multiple brain (n=3 of 4), eyestalk ganglia (n=3 of 4), and CG (n=2 of 4) cDNA samples (Fig. 2); no amplicon was generated from any of the cardiac muscle cDNAs (n=3 independent samples). The inconsistent amplification from CG cDNAs could be an indication that the CCRFamide transcript is conditionally regulated in this tissue.

To assess the validity of the transcript assemblies and to determine if the putative transcript variants exhibit differential expression, sense primers were designed to the variant specific sequences upstream of the respective start codons. The preproCCRFa-v1 primer consistently generated full-length ORF amplimers of the expected size (800 nt) from brain (n=3 of 4) and eyestalk ganglia (n=3 of 4) but was more limited in CG (n=1 of 4). A preproCCRFa-v2 amplimer (785 nt) was only observed in eyestalk ganglia, albeit faintly (n=3 of 4), and no preproCCRFa-v3 specific product (791 nt) was amplified from any of the cDNA sets (Fig. 2). As with the 500-bp CCRFamide fragment in the CG, the inconsistent amplification of the specific variants could indicate condition-based transcriptional activity.

4. Discussion

In the study presented here, a cDNA for a lobster, *H. americanus*, CCRFamide precursor is reported. This is the first cloning of cDNA encoding a prepro-CCRFamide from any species, and thus represents the only non-*in silico* evidence for the existence of this invertebrate peptide family. RT-PCR profiling suggests that CCRFamide is broadly expressed in the *H. americanus* nervous system, including in the brain, eyestalk ganglia, and CG. While the function(s) of CCRFamide remain unknown, it is highly likely that it serves as a locally released and/or circulating neuromodulator in the lobster. Detection of CCRFamide-encoding transcripts in the eyestalk ganglia suggests that it might be present in the X-organsinus gland system, a major neuroendocrine control center in decapod species (Christie, 2011). If so, the peptide could be delivered via the circulatory system to a diverse array of potential targets. The presence of CCRFamide in the CG suggests a potential role in modulating cardiac output, a role played by a number of –RFamide-containing peptides in *H. americanus* (Dickinson et al., 2015, 2016; Stevens et al., 2009).

As with the biological role and target tissue, the cognate CCRFamide receptor has yet to be elucidated. Although most regulatory peptides and neuromodulators signal through class A G protein-coupled receptors (GPCRs) (Tikhonova et al., 2019), disulfide-bridged peptides

like CCRFamide frequently act on other signaling systems such as insulin-like peptides and prothoracicotropic hormone (PTTH) with tyrosine kinases (Nässel and Zandawala, 2019), eclosion hormone with a guanylate cyclase (Nässel and Zandawala, 2019), or, as proposed for the agatoxin-like peptides, with ion channels (Sturm et al., 2016). Thus, specific identification of the CCRFamide receptor is potentially complicated by the breadth of potential molecular targets. Further complicating receptor identification and peptide functionality is the fact that the CCRFamide gene has been lost in a number of the more genetically malleable model organisms (*e.g. Drosophila melanogaster*) typically employed for functional studies (Veenstra, 2019). However, given that the neural circuit contained within the CG, the neuromuscular junction between the ganglion and cardiac muscle, and the muscle itself are all potential targets for CCRFamide modulation (e.g., Dickinson et al., 2015, 2016; Fort et al., 2007a, 2007b; Stevens et al., 2009), additional transcriptomic studies may provide insights into CCRFamide function and receptor elucidation.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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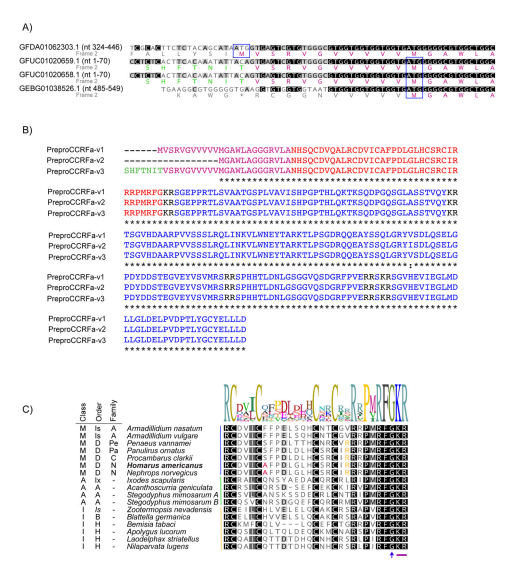


Figure 1.

Homarus americanus CCRFamide precursor alignments. A) Nucleotide sequence alignment of the region surrounding the predicted start sites in the four identified *H. americanus* CCRFamide precursor-encoding transcripts. Start codons are shown in blue boxes. Amino acids predicted to comprise a signal peptide are shown in pink, residues upstream of the predicted signal peptide are shown in green, and the untranslated region is shown in grey. B) MAFFT alignment of putative Homarus americanus CCRFamide precursors deduced from nervous system-specific transcriptomic data. PreproCCRFa-v1 corresponds to GFDA01062303, preproCCRFa-v2 to GEBG01038526, and preproCCRFa-v3 to GFUC01020659. In each protein, the signal peptide is shown in pink, residues upstream of the predicted signal peptide are shown in green, the CCRFamide isoform is shown in red, precursor-related peptides are shown in blue, and consensus dibasic cleavage sites are shown in black. In the line immediately below each sequence grouping, the symbol "*" indicates amino acids that are identical in all three proteins, while ":" denotes amino acids that are

similar in structure among all three sequences. C) Multiple sequence alignment of arthropod

CCRFamides. Sequences are based on conceptual translation of transcripts from species representing three Classes - Malacostraca (M, blue line), Arachnida (A, green line), and Insecta (I, yellow line). Atypical amino acid usages are shown in red (predominant Nephropidae-based) or orange (decapod specific). Abbreviations are: Order - Isopoda (Is), Decapoda (D), Ixodida (Ix), Araneae (A), Isoptera (*Is*), Blattodea (B), and Hemiptera (H); Family - Armadillidiidae (A), Penaeidae (Pe), Palinuridae (Pa), Cambaridae (C), and Nephropidae (N). The predicted amidation site is indicated by the arrowhead and the consensus C-terminal dibasic convertase cleavage site is indicated by the purple line. The *H. americanus* sequence is shown in bold. Sequence accession numbers are listed in Table S1.

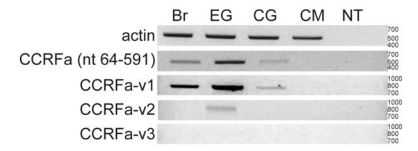


Figure 2. PCR-based expression profiling of CCRFamide-encoding transcripts in select portions of the *Homarus americanus* nervous system and heart muscle. Primers designed to amplify an internal 528-bp fragment of the CCRFamide precursor as well as variant specific transcripts encompassing the complete ORF were used with supraoesophageal ganglion (brain; Br), eyestalk ganglia (EG), cardiac ganglion (CG), and cardiac muscle (CM) cDNAs. As a control for cDNA integrity, a 503-bp fragment of actin was likewise amplified. Reactions were also performed in the absence of a template (no template; NT). The gel images shown are representative of positive amplification in at least one of four biological replicates.

 Table 1.

 Peptides predicted from putative *Homarus americanus* CCRFamide precursors

Peptide	Structure
CCRFa	NHSQ <u>C</u> DVQALR <u>C</u> DVICAFPDLGLH <u>C</u> SR <u>C</u> IRRRPMRFa
CCRFa- PRP-1	SGEPPRTLSVAATGSPLVAVISHPGPTHLQKTKSQDPGQSGLASSTVQY
CCRFa- PRP-2a	$TSGVHDAARPVVSSSLRQLINKVLWNEYTARKTLPSGDRQQEAYSSQLGRYVSDLQSELGPDY_{(SO3H)}DDSTEGVEY_{(SO3H)}VSVMRS$
CCRFa- PRP-2b	$TSGVHDAARPVVSSSLRQLINKVLWNEYTARKTLPSGDRQQEAYSSQLGRYISDLQSELGPDY_{(SO3H)}DDSTEGVEY_{(SO3H)}VSVMRS$
CCRFa- PRP-3	SPHHTLDNLGSGGVQSDGRFPVE
CCRFa- PRP-4	SGVHEVIEGLMDLLGLDELPVDPTLYGCYELLLD

Peptide abbreviations: CCRFa, CCRFamide; CCRFa-PRP, CCRFamide precursor-related peptide.

Abbreviations in peptide structures: Y(SO3H), sulfated tyrosine; \underline{C} , cysteine residue participating in a disulfide bond; a, carboxyl-terminal amide group.

Numbering of CCRFamide precursor-related peptides is based on their order within the preprohormone.

CCRFamide precursor-related peptide 2a was predicted from both prepro-CCRFamide variants 1 and 3, while CCRFamide precursor-related peptide 2b was predicted from prepro-CCRFamide variant 2; the two isoforms differ from one another in valine vs. isoleucine at position 52.

In the putative mature CCRFamide isoform, disulfide bridging is predicted between the first and fifth and second and fourth cysteine residues.