

RESEARCH ARTICLE

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The serendipitous origin of chordate secretin peptide family members

João CR Cardoso*, Florbela A Vieira, Ana S Gomes and Deborah M Power

Abstract

Background: The secretin family is a pleotropic group of brain-gut peptides with affinity for class 2 G-protein coupled receptors (secretin family GPCRs) proposed to have emerged early in the metazoan radiation via gene or genome duplications. In human, 10 members exist and sequence and functional homologues and ligand-receptor pairs have been characterised in representatives of most vertebrate classes. Secretin-like family GPCR homologues have also been isolated in non-vertebrate genomes however their corresponding ligands have not been convincingly identified and their evolution remains enigmatic.

Results: *In silico* sequence comparisons failed to retrieve a non-vertebrate (porifera, cnidaria, protostome and early deuterostome) secretin family homologue. In contrast, secretin family members were identified in lamprey, several teleosts and tetrapods and comparative studies revealed that sequence and structure is in general maintained. Sequence comparisons and phylogenetic analysis revealed that PACAP, VIP and GCG are the most highly conserved members and two major peptide subfamilies exist; i) PACAP-like which includes PACAP, PRP, VIP, PH, GHRH, SCT and ii) GCG-like which includes GCG, GLP1, GLP2 and GIP. Conserved regions flanking secretin family members were established by comparative analysis of the *Takifugu*, *Xenopus*, chicken and human genomes and gene homologues were identified in nematode, *Drosophila* and *Ciona* genomes but no gene linkage occurred. However, in *Drosophila* and nematode genes which flank vertebrate secretin family members were identified in the same chromosome.

Conclusions: Receptors of the secretin-like family GPCRs are present in protostomes but no sequence homologues of the vertebrate cognate ligands have been identified. It has not been possible to determine when the ligands evolved but it seems likely that it was after the protostome-deuterostome divergence from an exon that was part of an existing gene or gene fragment by rounds of gene/genome duplication. The duplicate exon under different evolutionary pressures originated the chordate PACAP-like and GCG-like subfamily groups. This event occurred after the emergence of the metazoan secretin GPCRs and led to the establishment of novel peptide-receptor interactions that contributed to the generation of novel physiological functions in the chordate lineage.

Background

The evolution of the secretin family of brain-gut peptides remains enigmatic despite being some of the first endocrine factors ever identified. For example, in 1902 the ground breaking experiments of Bayliss and Starling with dog intestinal extracts set off the search for the active principal and, by 1961, secretin (SCT) had been isolated and sequenced [1]. Currently, 10 peptides belonging to the secretin family have been isolated in humans and include; SCT, vasoactive intestinal peptide (VIP), pitu-

itary adenylate cyclase-activating polypeptide (PACAP), peptide histidine methionine (PHM), PACAP-related peptide (PRP), growth hormone-releasing hormone (GHRH), glucagon (GCG), glucagon-like peptide (GLP 1 and 2) and glucose-dependent insulintropic peptide (GIP) [2-4]. Members of the secretin family share significant structural and conformational homology and their key metabolic and developmental functions in human make them of considerable pharmacological interest. Members of class 2 G-protein coupled receptors (a.k.a family B GPCRs), bind and are activated by the secretin family members (family B1 members or secretin family GPCRs) and specific peptide-receptor pairs have been identified in representatives of different vertebrate

* Correspondence: jccardo@ualg.pt

¹ Centre of Marine Sciences (CCMAR), Universidade do Algarve, Campus de Gambelas, 8005-139 Faro, Portugal

Full list of author information is available at the end of the article

classes. Class 2 GPCRs is a larger family of receptors and also includes members of the metazoan adhesion (B2) and insect methuselah (B3) families and secretin family GPCRs (B1) are proposed to descend from the adhesion receptors prior to protostome-deuterostome divergence [5,6].

In protostomes (nematodes, arthropods, annelids and platyhelminthes) and early deuterostomes such as *Ciona* and amphioxus, immunohistochemical (IHC) approaches using antisera raised against various mammalian secretin family members suggest they possess similar peptides to vertebrates (Table 1). PACAP-like genes (*pacap1* and *pacap2*) have only been reported in the tunicate, *Chelysoma productum* [7] and partial mRNAs (114bp) corresponding to the highly conserved PACAP coding exon [8] have been isolated in *Hydra magnipapillata* and several protostomes and deposited in public databases [9]. The existing data has been taken to indicate that an ancestral secretin family gene was probably present prior to the deuterostome-protostome divergence and most likely resembled the vertebrate PACAP precursor [2,8,10-12]. Paradoxically, in protostomes with fully sequenced genomes and extensive molecular resources (Figure 1, Additional file 1), genes encoding ligands homologous to members of the vertebrate secretin family have not been reported. In contrast, secretin-like family GPCR encoding genes which share similar sequence, structure and conserved gene environment with the vertebrate members have been identified, making ligand-receptor evolution an interesting enigma [13-16].

The present study focuses on secretin family ligands and complements previous studies aimed at identifying and characterising the evolution of family 2 GPCRs [5,6,8,14,17]. A comparative approach which takes advantage of the wealth of information currently available (genome, ESTs, peptide) for porifera, cnidaria, protostomes, early deuterostomes and vertebrates (Figure 1), is undertaken to re-evaluate the origin of the secretin family in metazoa. The sequence, gene structure and gene environment of secretin family members in vertebrates with sequenced genomes was characterised and used to search for homologue peptides, genes or genome regions in non-vertebrates.

Results

Vertebrate secretin family members

Sequence database searches using the nucleotide and mature peptide sequences of human and zebrafish secretin family members, readily identified homologues in lamprey, teleost and tetrapod (*Xenopus*, lizard, chicken) genomes. This is due to the high sequence conservation of the mature peptide region between the vertebrate members which facilitates their identification (Table 2) (see reviews [2,4,8,18,19]).

The tetrapod members

In humans, 10 peptides encoded by six genes have been isolated. In *Aves*, homologues of the mammalian members have been identified and peptides and corresponding transcripts were isolated in duck (*Anas platyrhynchos*), chicken (*Gallus gallus*) and turkey (*Meleagris gallopavo*) [20-27]. *In silico* analysis of the chicken genome identified six genes encoding secretin family members which share similar organisation to the human homologues and comparative analysis revealed they correspond to the peptides and nucleotide precursors previously described (Table 1 and Table 2). Searches in the reptile and amphibia genomes identified homologues for all human members with the exception of SCT and it remains to be established if the failure is due to the incomplete nature of their genome assemblies or to the absence of this gene. In reptiles, few members of this family have been reported to date. A VIP and GCG peptides were isolated from the *Alligator mississippiensis*, the nucleotide precursor of the latter peptide reported from the *Heloderma suspectum* [28-30] and a PRP/PACAP mRNA was recently characterised from the Italian wall lizard *Podarcis sicula* [31,32]. Sequence database searches on the lizard *Anolis carolinensis* genome identified for the first time the genes encoding the reptile PHI/VIP (ENSACAG00000005619), PRP/PACAP (ENSACAG00000008729), GHRH (ENSACA G00000011836), GCG/GLP (ENSACAG00000014182) and GIP (ENSACAG00000006291) (Table 2). The predicted gene organisation suggests the coding exons for the mature peptides share identical structure with other metazoan genes and to date the exon encoding the lizard GLP2 remains to be identified.

Homologues of human secretin members have previously been reported in amphibians. VIP and PACAP and were isolated in *Rana ridibunda* [33,34]; GCG from *Rana catesbeiana* [35] and in *Xenopus laevis* single transcripts for PRP/PACAP [36], PHI/VIP, GHRH [37], GCG/GLP [38] and GIP [27] have been described. The genome of *Xenopus tropicalis* contains secretin family homologues which share high sequence conservation with the tetrapod genes (Table 2), but one difference was the presence of three GLP1 exons (GLP1a, b and c) within the GCG/GLP gene structure as a result of a species-specific exon duplication.

The fish members

Peptides, transcripts and genes of the secretin family have also been isolated from the most diverse vertebrate clade, the teleosts. VIP was isolated from cod (*Gadus morhua*) [39,40] and goldfish (*Carassius auratus*) [41], PACAP from the Japanese stargazer (*Uranoscopus japonicus*) [42] and GCG from the channel catfish (*Ictalurus punctatus*) [43,44] (Table 1). The identification of two transcripts for PHI/VIP in goldfish *Carassius auratus* [45] and zebrafish

Table 1: Molecular and expression data available for the secretin members in metazoa.

	Species	PRP/ PACAP	PH/VIP	GHRH	GCG/GLP	GIP	SCT	References
DEUTEROSTOMES								
VERTEBRATES								
Mammal	<i>Homo sapiens</i>	N	N, P	N, P	N, P	N, P	N, P	[100-109]
Ave	<i>Anas platyrhynchos</i>	N	N	N	P			[9,21,22]
	<i>Gallus gallus</i>	N,P	N,P	N	N,P	N	P	[24-27,35,110-112],
	<i>Meleagris gallopavo</i>		N		N			[23,113]
Reptile	<i>Alligator mississippiensis</i>		P		P			[28,29]
	<i>Heloderma suspectum</i>				N			[30]
	<i>Podarcis sicula</i>	N						[31]
Amphibia	<i>Rana catesbeiana</i>				P			[114]
	<i>Rana ridibunda</i>	N,P	P					[33,34]
	<i>Xenopus laevis</i>	N	N	N	N	N		[9,27,36-38]
	<i>Xenopus tropicalis</i>				N	N		[9,27],
Teleost	<i>Carassius auratus</i>		N,P	N	N			[37,41,45,115],
	<i>Clarias macrocephalus</i>	N			N			[9,116],
	<i>Danio rerio</i>	N	N	N	N	N		[9,27,37,47,83,117,118]
	<i>Gadus morhua</i>	N	P					[39,40]
	<i>Ictalurus punctatus</i>	N			N,P			[9,43,44]
	<i>Oncorhynchus nerka</i>	N						[119]
	<i>Takifugu rubripes</i>	N	N					[8]
	<i>Uranoscopus japonicus</i>	P						[42]
Chondrichthyes	<i>Torpedo marmota</i>	N, IHC						[120]
	<i>Dasyatis akajei</i>	IHC						[121]
	<i>Callorhynchus milii</i>				P			[122]
Agnatha	<i>Lampetra fluviatilis</i>				P			[123]
	<i>Lampetra japonica</i>	N						[9]
	<i>Petromyzon marinus</i>				N,P			[52,124]
UROCHORDATES	<i>Chelyosoma productum</i>	N, IS						[7]
	<i>Ciona intestinalis</i>		IHC				IHC	[125-127]
	<i>Halocynthia roretzi</i>	N*						[9]
	<i>Styela plicata</i>		IHC		IHC		IHC	[125,127-129]
CEPHALOCHORDATES	<i>Branchiostomata lanceolatum</i>		IHC		IHC		IHC	[127,130,131]
PROTOSTOMES								
NEMATODES	<i>Ascaridia galli</i>		IHC					[132]
	<i>Nematodirus battus</i>		IHC					[132]
	<i>Nippostrongylus brasiliensis</i>		IHC					[132]
ARTHROPODS	<i>Aeschna cyanea</i>		IHC					[133,134]
	<i>Eriocheir japonica</i>	N*						[9]
	<i>Eristalis aeneus</i>				IHC		IHC	[135]
	<i>Manduca sexta</i>		IHC		IHC	IHC	IHC	[136]
	<i>Periplaneta americana</i>	N*	IHC					[9,10,131,137,138]
ANNELIDS	<i>Hirudo medicinalis</i>		IHC					[131,139]

Table 1: Molecular and expression data available for the secretin members in metazoa. (Continued)

	<i>Lumbricus terrestris</i>	IHC	IHC		[131,140,141]
	<i>Nereis diversicolor</i>		IHC		[131,142]
	<i>Helix pomatia</i>		IHC		[143]
MOLLUSCS	<i>Mytilus galloprovincialis</i>		IHC		[144]
	<i>Planorbarius corneus</i>		IHC	IHC	IHC [145]
	<i>Sepioteuthis lessoniana</i>	N*			[9]
	<i>Viviparus ater</i>		IHC	IHC	IHC [146]
PLATYHELMINTHES	<i>Dugesia japonica</i>	N*			[9]
	<i>Schistosoma mansoni</i>		IHC		[147]
CNIDARIA	<i>Hydra magnipapillata</i>	N*			[9]

Nucleotide (N), Protein (P), Immunohistochemical (IHC) and *in situ* (IS)
 N* non-vertebrate nucleotide sequences deposited in NCBI

(*Danio rerio*) (PHI/VIP *a*, EU031789 and PHI/VIP *b*, EU031790) and of two PRP/PACAP (PRP/PACAP *a*, NM_152885 and PRP/PACAP *b*, AF329633) [46,47] and GCG/GLP precursors in zebrafish [48] suggests they are duplicates in fish and this has been confirmed by the identification of two PRP/PACAP and GCG/GLP genes in *Takifugu* (ENSTRUG00000003782 and ENSTRU G00000010059; ENSTRUG00000008721 and ENSTRU G00000004633, respectively) and *Tetraodon* (ENSTNI G00000017117 and ENSTNIG00000018649; ENSTNIG 00000013278 and ENSTNIG00000000614, respectively) genomes (Table 2) [8]. The greater number of secretin family genes identified in fish relative to tetrapods is most likely to be a result of the proposed teleost specific genome duplication and the absence of GIP and GHRH gene duplicates suggests they were probably deleted [49-51]. In common with *Xenopus*, no homologue of human SCT has been identified in fish genomes.

In the sea lamprey (*Petromyzon marinus*), a primitive vertebrate of the *Agnatha* clade, two homologues of human proglucagon (GCG/GLP) were characterised in the genome assembly Contig 31522 (GENSCAN 00000079364) and Contig 32128 which correspond to the previously reported proglucagon I (AF159707) and proglucagon II (AF159708) transcripts, respectively [52]. In the present study, searches of the partially sequenced lamprey genome retrieved putative PRP/PACAP (GENSCAN00000120210 on Contig3575.2) and PHI/VIP genes (shared between GENSCAN00000109335 and GENSCAN00000056150 localised on Contig20045.2 and Contig20045.3, respectively). However, GHRH, SCT and GIP were not identified possibly due to the present incomplete nature of its genome assembly.

Sequence and gene structure comparison

Sequence comparisons reveal that members of the secretin family are highly conserved and this also applies to

their secondary structure which consists of a random N-terminal structure and a C-terminal alpha-helix [53,54]. The lamprey, teleost, *Xenopus*, reptile and chicken members are in general 50% identical in amino acid sequence with the human homologues (Figure 2) and two main peptide subfamilies, which share in general a maximum of 60% sequence similarity between their members were identified; i) PACAP-like subfamily which includes 6 peptide groups (PACAP, PRP, VIP, PHI, GHRH and SCT) and ii) GCG-like subfamily which contains 4 peptide groups (GCG, GLP1, GLP2 and GIP). Highest sequence conservation from lamprey to human (>70% sequence identity) occurs within PACAP, VIP and GCG peptide groups which contrasts with PRP and GLP2 that are the most divergent and *Takifugu* PRP *b* and GLP2 predicted peptides are only 33% and 37% identical with the human homologues.

Consensus amino acids for peptide subfamilies were deduced and the overall conservation characterised. With few exceptions, 7 amino acid residues H¹, D³, F⁶, T⁷, Y¹⁰, Q¹⁶ and L²³ are the most abundant across the PACAP-like subfamily (Figure 2). The residues H¹, D³ and L²³ are present in all peptide groups with occasional exceptions in some taxa. However, the motif F⁶T⁷ is absent from the vertebrate PRP sequences and is only present in tunicate PRP and the residue Q¹⁶ is absent from the SCT mature peptide and Y¹⁰ is only present in the chicken SCT homologue. This suggests that specific modifications occurred within the conserved core domain of the PACAP-like subfamily members and their functional significance remains to be explored. A similar comparison of the vertebrate GCG-like subfamily indicates that 15 amino acid residues H¹, A², G⁴, T⁵, F⁶, S⁸, D⁹, S¹¹, L¹⁴, A¹⁹, K²⁰, F²², V²³, W²⁵, and L²⁶ are generally maintained across the 4 peptide groups. Even if taxa variability is taken into consideration, the residues G⁴, F⁶ and F²² and the motifs S⁸ D⁹

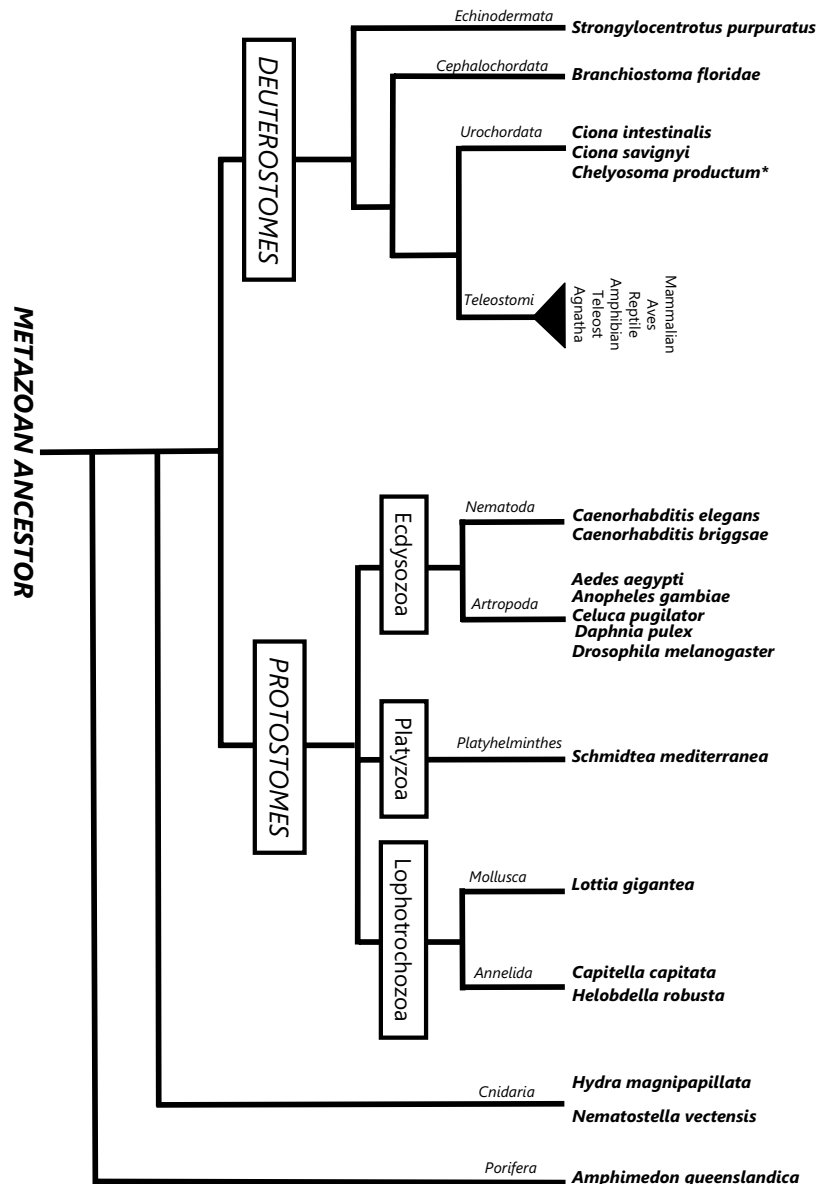


Figure 1 Phylogenetic position of the non-vertebrate genomes analysed. Simplified phylogeny of the metazoan evolution indicating the relative position of the early metazoa (Porifera and Cnidaria), protostome (Nematoda, Arthropoda, Platyhelminthes, Mollusca, Annelida) and early deuterostome (Echinodermata, Cephalochordata and Urochordata) genomes analysed (adapted from [96-98]). The tunicate *Chelyosoma productum* is also represented (*) since it is the only invertebrate in which secretin family members have been isolated [7].

and W²⁵L²⁶ are in general maintained (Figure 2). Peptide specific variations include for GIP, H¹ which is replaced by Y, L¹⁴ which is replaced by M or V (with exception of lizard) and A¹⁹ which is replaced by K or Q. In the GCG mature peptide sequence A² is replaced by S and in GLP2 with the exception of chicken and *Takifugu* T⁵ is replaced by S. Comparison of the metazoan PACAP-like and GCG-like subfamily consensus sequence revealed they

overlap for the residues H¹ and F⁶ which are key amino acids in secretin GPCR binding affinity [19,53,55,56]. This suggests that, after exon/chromosome duplication of their common ancestor exon, distinct evolutionary pressures within each subfamily occurred.

The chordate PACAP shares in general 92% of amino acid sequence similarity with VIP and these two peptides are the most conserved members (Figure 3). In contrast,

Table 2: Accession numbers (ENSEMBL) of non-mammalian secretin members.

	PRP/PACAP	PH/VIP	GHRH	GCG/GLP	GIP	SCT
Chicken (<i>Gallus gallus</i>)	ENSGALG0000 0014858	ENSGALG0000 0013604	ENSGALG0000 0003842	ENSGALG0000 0011104	ENSGALG0000 0001299	ENSGALG0000 0005081
Lizard (<i>Anolis carolinensis</i>)	ENSACAG0000 00008729	ENSACAG0000 00005619	ENSACAG0000 00011836	ENSACAG0000 00014182	ENSACAG0000 00006291	---
Xenopus (<i>Xenopus tropicalis</i>)	ENSXETG0000 0019179	ENSXETG0000 0027906	ENSXETESTG0 0000008409	ENSXETG0000 0013178	Scaffold_334	---
Takifugu (<i>Takifugu rubripes</i>)	ENSTRUG0000 0003782 ENSTRUG0000 0010059	ENSTRUG0000 0001139	---	ENSTRUG0000 0008721 ENSTRUG0000 0004633	---	---
Tetraodon (<i>Tetraodon nigroviridis</i>)	ENSTNIG0000 0017117 ENSTNIG0000 0018649	ENSTNIG0000 0007449	ENSTNIG0000 0007343	ENSTNIG0000 0013278 ENSTNIG0000 0000614	---	---
Zebrafish (<i>Danio rerio</i>)	ENSDARG0000 00004015 ENSDARG0000 00027740	ENSDARG0000 00079443 ENSDARG0000 00078247	ENSDARG0000 00069481	ENSDARG0000 00042999/ ENSDARG0000 00079296 ENSDARG0000 00040907	ENSDARG0000 00071306	---
Stickleback (<i>Gasterosteus aculeatus</i>)	ENSGACG0000 00017084 ENSGACG0000 00004163	ENSGACG0000 00001298	ENSGACG0000 00001298	ENSGACG0000 00013877 ENSGACG0000 00005606	---	---
Medaka (<i>Oryzias latipes</i>)	ENSORLG0000 0017872 ENSORLG0000 0011205	ENSORLG0000 0003905	---	ENSORLG0000 0002782 ENSORLG0000 0016891	---	---
Lamprey (<i>Petromyzon marinus</i>)	GENSCAN0000 00120210	GENSCAN0000 00109335/ GENSCAN0000 00056150	---	GENSCAN0000 00079364 Contig 32128	---	---

Contig/Scaffold numbers are indicated when ENSEMBL accession number are not available
Dashes indicate genes not identified

the vertebrate SCT demonstrates the lowest conservation (less than 62% within the PACAP-like subfamily) however this may be an artefact due to the restricted number of species in which it has been characterised. In addition, signature amino acid residues within or between peptide groups were also identified and they may reflect and support common evolutionary pathways and overlapping or specific functional roles. This includes, the PACAP Y¹³ and A²⁴A²⁵ motif, the V⁵, N⁹, F¹³ and N²⁴ for VIP, the motif N⁷K⁸A⁹ and residue H²⁴ within PRP, the L⁵, S⁸, E²⁴ and I²⁷ for PH, and the GHRH residues N⁸, I¹⁷, Q²⁴ and I²⁶ across the tetrapod and teleost members (Figure 2). Gene structure comparisons restricted to the mature peptide precursors revealed that PACAP and VIP exons are encoded in the same precursor as PRP and PH, respec-

tively [2,8] (Figure 3). Most common amino acids to the chordate PACAP and VIP members includes R¹⁴ and K¹⁵ and the motifs M¹⁷A¹⁸V¹⁹ and V²⁶L²⁷ and they share a maximum of 37% amino acid sequence identity with PRP and PH indicating that after exon duplication considerable changes occurred. The latter peptides have a similar sequence (81% similar) which is closely related to vertebrate GHRH (88% similar) with which they partition the residues A², K¹², L¹⁴, G¹⁵ and the motif S¹⁸A¹⁹R²⁰ and suggest a common evolutionary origin. The SCT gene has only been identified in tetrapods and the deduced peptide residues, E⁹ and Q²⁰ are maintained in the mammalian and chicken homologues. The evolutionary origin of SCT is still enigmatic and it is proposed to have been lost in the fish lineage [14,57].

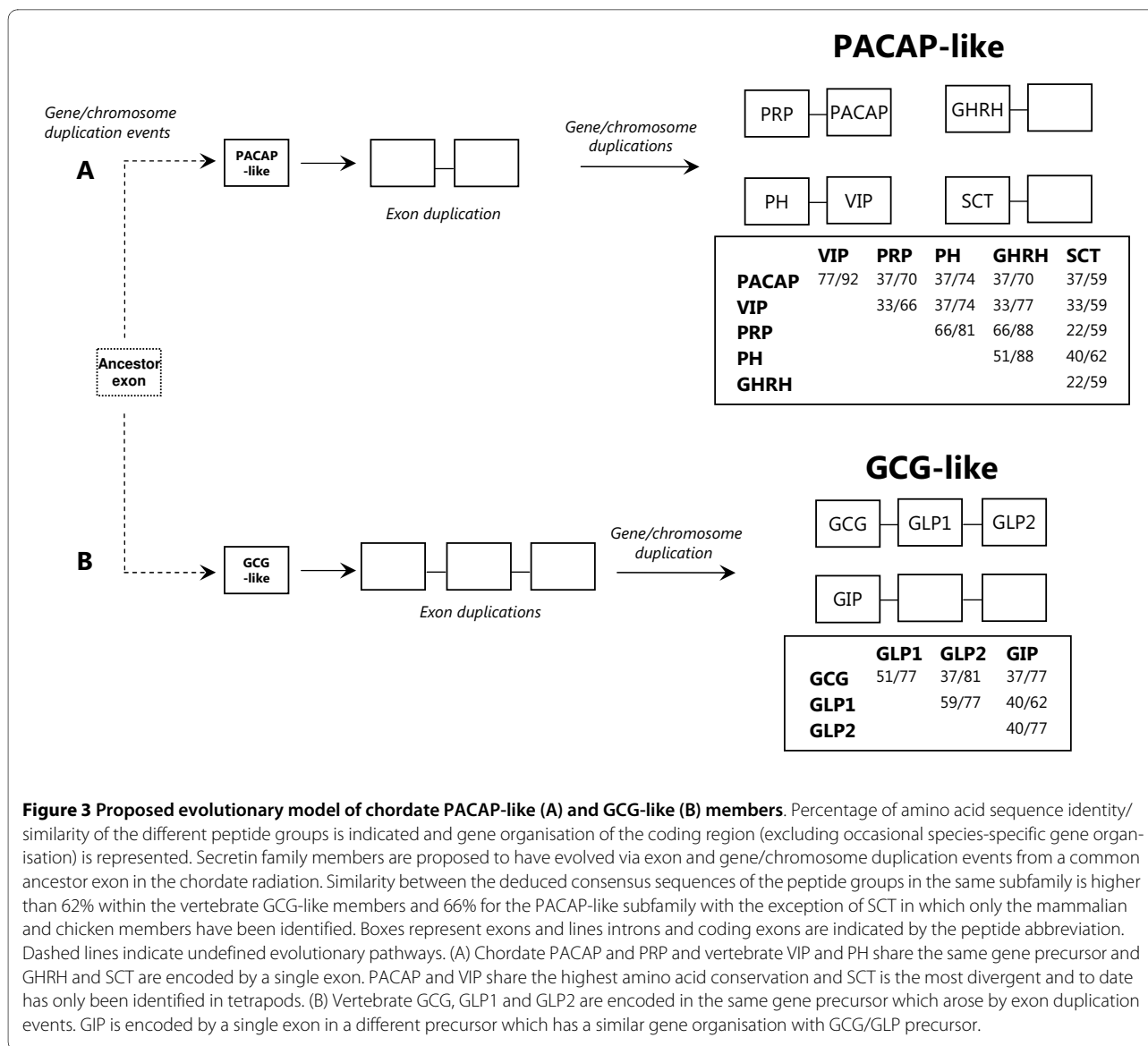
PACAP-like				GCG-like					
PACAP	1	10	20	%ID	GCG	1	10	20	%ID
Human	: HSDGIFTDSYSRYRQ MAVKKYLAAVL	100			Human	: HSQGTF TS SDYSKYLD SRRAQDFVQWLM	100		
Mouse	: HSDGIFTDSYSRYRQ MAVKKYLAAVL	100			Mouse	: HSQGTF TS SDYSKYLD SRRAQDFVQWLM	100		
Chicken	: HIDGIFTDSYSRYRQ MAVKKYLAAVL	96			Chicken	: HSQGTF TS SDYSKYLD SRRAQDFVQWLM	100		
Lizard	: HSDGIFTDSYSRYRQ MAVKKYLAAVL	100			Lizard	: HSQGTF TS SDYSKYLD TRRAQDFVQWLM	96		
Xenopus	: HSDGIFTDSYSRYRQ MAVKKYLAAVL	100			Xenopus	: HSQGTF TS SDYSKYLD SRRAQDFIOWLM	96		
Zebrafisha	: HSDGVFTDSYSRYRQ MAVKKYLAATVL	92			Zebrafisha	: HSEGTF SNDYSKYLETRRAQDFVQWLM	81		
Zebrafishb	: HSDGIFTDSYSRYRQ MAVQKYLAAVL	96			Zebrafishb	: HSEGTF SNDYSKYLETRRAQDFVQWLM	81		
Takifugua	: HSDGIFTDSYSRYRQ MAVKKYLAAVL	100			Takifugua	: HSEGTF SNDYSKYLEDRRAQDFVQWLM	74		
Takifugub	: HSDGIFTDSYSRYRQ MAVQKYLAAVL	96			Takifugub	: HSEGTF SNDYSKYLETRRAQDFVQWLM	77		
Sticklebackka	: HSDGIFTDSYSRYRQ MAVKKYLAAVL	100			Sticklebackka	: HSEGTF SNDYSKYLETRRAQDFVQWLM	77		
Sticklebackkb	: HSDGIFTDSYSRYRQ MAVQKYLAAVL	96			Sticklebackkb	: HSEGTF SNDYSKYLETRRAQDFVQWLM	70		
Lamprey	: HSDGLFTDLYSRYRQ MAVKKYLSVTL	85			LampreyI	: HSEGTF TS SDYSKYLD ENKQAKDFVQWLM	74		
<i>CheLyosoma1</i>	: HSDGIFTDSYSRYRQ MAVKKYLAAVL	96			LampreyII	: HSQGFT TS SDYSKYLD VKQAKDFVQWLM	70		
<i>CheLyosoma2</i>	: HSDGIFTDSYSRYRQ MAVKKYINALL	85			consensus	: HSEGTFSDYSKYLETRRAQDFVQWLM			
consensus	: HSDGIFTDSYSRYRQMAVKKYLAAVL								
VIP					GLP1				
Human	: HSDAVFTD NXTRLRQMAVKKYLSNL	100			Human	: HAEGTFT SDVSSYLEGQAAKEFIWLV	100		
Mouse	: HSDAVFTD NXTRLRQMAVKKYLSNL	88			Mouse	: HAEGTFT SDVSSYLEGQAAKEFIWLV	100		
Chicken	: HSDAVFTD NXSRFRQMAVKKYLSNL	88			Chicken	: HAEGTYT SDITSYLEGQAAKEFIWLV	88		
Lizard	: HSDAVFTD NXSRFRQMAVKKYLSNL	88			Lizard	: HADGTYT SDISSYLEGQAAKEFIWLV	88		
Xenopus	: HSDAVFTD NXSRFRQMAVKKYLSNL	88			Xenopusa	: HAEGTFT SDVTQQLDEKAAKEFIWLI	70		
Zebrafisha	: HSDAVFTD NXSRFRQMAAKKYLSNL	85			Xenopusb	: HAEGTYT NDVTEYLEKAAKEFIWLI	70		
Zebrafishb	: HSDAIFTD NXSRFRQMAVKKYLSNL	85			Xenopusc	: HAEGTFT NDMTNYLEKAAKEFIWLI	66		
Takifugua	: HSDAIFTD NXSRFRQMAVKKYLSNL	85			Zebrafisha	: HADGTYT SDVSSYLEQDQAAKEFVQWLM	74		
Takifugub	: HSDAIFTD NXSRFRQMAVKKYLSNL	85			Zebrafishb	: HAEGTYT SDVSSYLEQDQAAKEFVQWLM	74		
Sticklebackkb	: HSDAIFTD NXSRFRQMAVKKYLSNL	85			Takifugua	: HADGTYT SDVSSYLEKQAIKDFVQWLM	70		
Lamprey	: HSDAVFTD LFSRLRQQAEEKYAKSL	70			Takifugub	: HADGTYT SDVSTYLDQDQAAKEFVQWLM	70		
consensus	: HSDAVFTDNYSRFRQMAVKKYLSNL				Sticklebackka	: HADGTYT SDVSSYLEKQAIKDFVQWLM	70		
					Sticklebackkb	: HADGTYT SDVSSYLEQDQAAKEFVQWLM	66		
					LampreyI	: HADGTYT SDVSSYLEKQAIKDFVQWLM	55		
					consensus	: HADGTYTSDVSSYLEQDQAAKEFVQWLM			
PRP					GLP2				
Human	: VAHGILNEAYRKYVLDQLSAGKHLQSLV	100			Human	: HADGFSF DEMNTILDNLAARDFINWLI	100		
Mouse	: VAHEILNEAYRKYVLDQLSARKYLSQSV	85			Mouse	: HADGFSF DEMNTILDNLAARDFINWLI	92		
Chicken	: HADGIF SKAYRKLGLSARNYLHSLM	55			Chicken	: HADGTF TS SDINKL DDMAAKEFIWLI	55		
Lizard	: HADGIF NKAYRKYVLDQLSARKYLHSLM	66			Zebrafish	: HVDGTF TS SDVNVK LDLSIAAKEFIWLI	40		
Xenopus	: HADGLL NKAYRKYVLDQLSARKYLHSLM	66			Takifugu	: HVDGTF TS SDVNVK LDLSIAAKEFIWLI	37		
Zebrafisha	: HADGMFN KAYRKYVLDQLSARKYLHSLM	55			Stickleback	: HVDGTF TS SDVNVK LDLSIAAKEFIWLI	44		
Zebrafishb	: HADGLL DRALRDLILVQLSARKYLHSLM	51			LampreyII	: HSDGFT TS SDVNVK LDLSIAAKEFIWLI	48		
Takifugua	: HADGMFN KAYRKYVLDQLSARKYLHSLM	59			consensus	: HADGFSFSDMNKILDSMAAKEFIWLI			
Takifugub	: HAEEEL DRALREILGLTARHYRFLM	33							
Sticklebackka	: HADGMFN KAYRKYVLDQLSARKYLHSLM	59							
Sticklebackkb	: HAEEEL DRALREILGLTARHYRFLM	33							
<i>CheLyosoma1</i>	: HSDGIFTD KYRKYVLDQLSARKYLQWLM	48							
<i>CheLyosoma2</i>	: HSDGIFTD YRKYVLDQLSARKYLQWLM	48							
consensus	: HADGIFNKAYRKYVLDQLSARKYLHSLM								
PH					GIP				
Human	: HADGVFT SDFSKLLGQLSARKYLESML	100			Human	: YAEGTFT SDYSIAMDKIHQQDFVNWLL	100		
Mouse	: HADGVFT SDYSRLLGQLSARKYLESML	85			Mouse	: YAEGTFT SDYSIAMDKIHQQDFVNWLL	96		
Chicken	: HADGIFT SVHSHLLAKLSVRRYLSLI	62			Chicken	: YSEATL ASDYSRTMDNMLKKNFVWLL	51		
Lizard	: HADGLFT SGYSKLLGQLSARRYLESLE	77			Lizard	: YSEGLT ASDYSRTMDNMLKKNFVWLL	51		
Xenopus	: HADGLFT SGYSKLLGQLSARRYLESLE	77			Xenopus	: YSEAIL ASDYSRSDNMLKKNFVWLL	44		
Zebrafisha	: HADGLFT SGYSKLLGQLSARRYLESLE	77			Zebrafish	: YAESTI ASDISKIVDSMVQKNFVWLL	51		
Zebrafishb	: HADGLFT SGYSKLLGQLSARRYLESLE	81			consensus	: YAEGLTASDYSRTMDNMLKKNFVWLL			
Takifugub	: HADGLFT SGYSKLLGQLSARRYLESLE	77							
Sticklebackkb	: HADGLFT SGYSKLLGQLSARRYLESLE	77							
Lamprey	: HADALFT HNNYKLLGQMSARRYFESLL	55							
consensus	: HADGLFTSGYSKLLGQLSARRYLESLE								
GHRH									
Human	: YADAIFT NSYRKYVLDQLSARKYLQDIM	100							
Mouse	: HVDAIFT NTYRKYVLDQLSARKYLQDIM	66							
Chicken	: HADAIFT NTYRKYVLDQLSARKYLQDIM	70							
Lizard	: HADAIFT NTYRKYVLDQLSARKYLQDIM	74							
Xenopus	: HVDAIFT NTYRKYVLDQLSARKYLQDIM	62							
Zebrafish	: HADAIFT NTYRKYVLDQLSARKYLQDIM	81							
Takifugu	: HADAIFT NTYRKYVLDQLSARKYLQDIM	85							
Stickleback	: HADAIFT NTYRKYVLDQLSARKYLQDIM	85							
consensus	: HADAIFTNSYRKYVLDQLSARKYLQDIM								
SCT									
Human	: HSDGFT TSELSRRLREGARLQRLQLGLV	100							
Mouse	: HSDGMFT TSELSRRLQDSARLQRLQLGLV	85							
Chicken	: HSDGLFT TSEYSKMRGNAQVQKFIQNL	51							
consensus	: HSDGMFTSELSRRLQDSARLQRLQLGLV								

PACAP-like most conserved aa H.D..FT..Y.....Q.....L....

GCG-like most conserved aa HA.GTF.SD.S..L....AK.FV.WL.

Prototype sequence model **HADGTF**TS~~SDYS~~..L.Q...AK.F..WL.

Figure 2 Amino acid sequence conservation of vertebrate secretin family mature peptides. The mature peptide sequences were extracted by comparison with the human homologues and only the amino acid (aa) residues 1 to 27 are represented with the exception of the first 5 residues of human GLP1 (P01275). *Takifugu* GHRH was obtained from [93] (N000079, Assembly_4) and the lamprey GLP2 sequence for proglucagon II was not used since it was found to share very little similarity with other vertebrate members suggesting it has undergone species-specific evolution. Vertebrate peptides are grouped according to their potential origin from a PACAP-like exon or GCG-like exon. Percentage of identity (%ID) for the human homologue is given and the consensus sequences for each peptide group were deduced using the GeneDoc programme [90] and used to generate a PACAP-like and GCG-like subfamily peptide. The most frequent residues within the different peptide groups are annotated in bold and a prototype model sequence for the chordate secretin family members was derived by fusing the conserved PACAP-like and GCG-like subfamily amino acid sequence (overlapping residues are annotated in bold and italics). Accession numbers of the teleost and non-mammalian sequences used are indicated in Table 2. The human precursors are PHM/VIP, P01282; PRP/PACAP, P18509; GHRH, P01286; GCG/GLPs, P01275; GIP, P09681; and SCT, P09683 and mouse (*Mus musculus*) accession numbers are PHM/VIP, P32648; PRP/PACAP, O70176; GHRH, P16043; GCG/GLPs, P55095; GIP, P48756; and SCT, Q08535.



Members of the GCG-like subfamily share at least 62% amino acid similarity and GLP1 and GLP2 have the highest identity (59%, Figure 3). In addition to the general sequence conservation of the GCG-like subfamily members, conserved amino acid positions within the vertebrate peptide groups were also identified (Figure 2). This includes S², Q²⁴, M²⁷ and the motifs R¹⁷R¹⁸ and Q²⁰D²¹ for GCG; V¹⁰, S¹² and Q¹⁷ for GLP1; S⁵, N¹¹ and L²³ for GLP2; and Y¹ and L²⁷ in GIP. The GLP1 and GLP2 peptides are encoded in the same precursor (proglucagon precursor) as GCG with which they share 77% and 81% amino acid sequence similarity, respectively and the mature peptide coding exons are proposed to be a consequence of exon duplication events. Vertebrate GIP shares 37% and 40% sequence identity with GCG and GLPs, respectively and studies based upon gene structure comparisons suggest the latter emerged from the same exon as GCG, however

the results of sequence analysis are inconclusive and more data is required [2,27].

The secretin members in non-vertebrates

Database searches using the vertebrate nucleotide and peptide sequences of secretin family members and the duplicate urochordate *Chelyosoma productum* PRP/PACAP transcripts failed to identify conserved sequence and structure homologues in genome or EST databases from porifera, cnidaria, protostome and early deuterostomes (*Ciona*, amphioxus and sea urchin). Instead, short sequence matches were identified in unrelated genes or non-annotated genome segments. In depth analysis of the best matches, revealed homologies for the central region and C-terminal ends (outside the bioactive core) of the chordate mature peptides and, when the invertebrate fragments were used to interrogate vertebrate data-

bases, they failed to retrieve a secretin family homologue suggesting that members of this family are absent from non-vertebrate genomes.

The exception is PACAP for which a highly conserved partial sequence corresponding to the exon encoding the mature peptide (>89% amino acid identity, [8]) has been isolated in *Hydra magnipapillata* (AB083650), in the tunicate, *Halocynthia roretzi* (AB121759) and in several protostomes such as planarian (*Dugesia japonica*, AB083649), crab (*Eriocheir japonica*, AB121765), squid (*Sepioteuthis lessoniana*, AB083651) and cockroach (*Periplaneta americana*, AB083652) [9]. Database searches using the non-vertebrate PACAP nucleotide or deduced peptide sequences failed to retrieve homologues from protostomes with available genome data (*Helobdella robusta*, *Capitella sp. 1*, *Lottia gigantea*, *Daphnia pulex*, *Drosophila melanogaster*, *Aedes aegypti*, *Anopheles gambiae*, *Caenorhabditis elegans* and *Caenorhabditis briggsae*). Moreover, searches performed in *Hydra magnipapillata* and related species *Nematostella vectensis* genome assemblies with the *Hydra* PACAP nucleotide or deduced peptide sequence also failed to confirm the existence of a gene encoding PACAP. In early deuterostomes, searches using the *Chelyosoma productum* PRP/PACAP nucleotide or deduced peptide sequences in *Ciona intestinalis* and *Ciona savignyi* genomes or available tunicate ESTs failed to identify possible sequence homologues in urochordate. Taken together these results raise questions about the authenticity of the previously reported sequences.

An alternative strategy utilized a secretin family prototype sequence model based on the assumption that the chordate members arose from a common precursor gene which duplicated to give PACAP-like and GCG-like subfamily exons (Figure 2 and 3). The prototype sequence deduced *in silico* was H¹A²D³G⁴T⁵F⁶T⁷S⁸D⁹Y¹⁰S¹¹xxL¹⁴xQ¹⁶xxA¹⁹K²⁰xF²²xxW²⁵L²⁶ (x represents variable position) and contains 18 conserved amino acid positions and high conservation was found for the N-terminal region. Structural characterisation using Pfam analysis classified the generated prototype sequence as a Hormone_2 member (PF00123, which includes the vertebrate secretin family members) [58] and sequence similarity searches performed retrieved secretin family members in vertebrates but failed to identify potential members in non-vertebrates. A second approach using HMM models and searching the general NCBI non-redundant (nr) peptide and an invertebrate subset of the NCBI nucleotide and EST (est_others) databases corroborated the preceding results.

Phylogenetic analysis

The optimal maximum likelihood (ML) tree with bootstrap support values higher than 50% is presented in Fig-

ure 4. Despite the high level of sequence identity and short sequences utilized, phylogenetic analyses of the chordate mature peptides (1-27 aa) and their corresponding nucleotide sequences resulted in similar tree topologies and suggests that members of the secretin family share a common ancestry. In the optimal ML tree presented in Figure 4 two clades PACAP-like and GCG-like were obtained suggesting that after ancestral exon duplication two main peptide subgroups emerged and underwent distinct evolutionary trajectories. Similar tree topologies were obtained using Bayesian approaches (Supplementary table 2) and a PACAP-like derived clade includes the peptides PACAP, PRP, PH, VIP, GHRH and SCT and a GCG-like clade the peptides GCG, GLP1, GLP2 and GIP. The *Chelyosoma* PACAP deduced peptide sequences always grouped with the vertebrate peptide sequence homologues and the tunicate PRPs did not cluster with any particular peptide clade and in all the analysis performed tended to be more closely related to the tetrapod SCT.

Gene environment comparisons

The immediate gene environment of vertebrate secretin members was compared and indicates that PRP/PACAP, GCG/GLP, and GHRH genome regions are syntenic and gene order is in general maintained (Figure 5). The GCG/GLP gene environment shares at least 3 genes (KCNH7, IFIH1 and SLC4A10), PRP/PACAP, 2 genes (YES1 and METTL4) and GHRH, 1 gene (RPN2) when equivalent genome regions are compared between *Takifugu*, *Xenopus*, chicken and human. The VIP and GIP genomic regions are poorly conserved and no gene synteny or gene order was identified (data not shown). Moreover, searches also failed to identify conservation of gene linkage between the different vertebrate secretin family members.

In *Takifugu*, the neighbouring genes of the paralogue GCG/GLP (SLC4A10 and KCNH7, N000046 and N000036) and PACAP (YES1, N000223 and N000080) genes were also duplicates supporting the teleost gene or genome duplication event. Searches were extended to the lamprey and lizard genomes but contiguous sequences were too small to confirm the existence of gene environment conservation. Comparison of the SCT genome region between chicken (chromosome 5) and human (chromosome 11) revealed they are highly conserved and the order of 5 genes (MUCDHL, DRD4, DEAF1, EPS8L2 and PDDC1) is maintained. In *Xenopus* genome, the chicken and human SCT flanking genes are localised in the amphibian scaffold_296 and in *Takifugu*, where a gene homologue is proposed to be absent, they are distributed in two distinct genome regions: MUCDHL and DRD4 are localised in scaffold N000002 and the remaining genes in scaffold N000328 suggesting the existence of

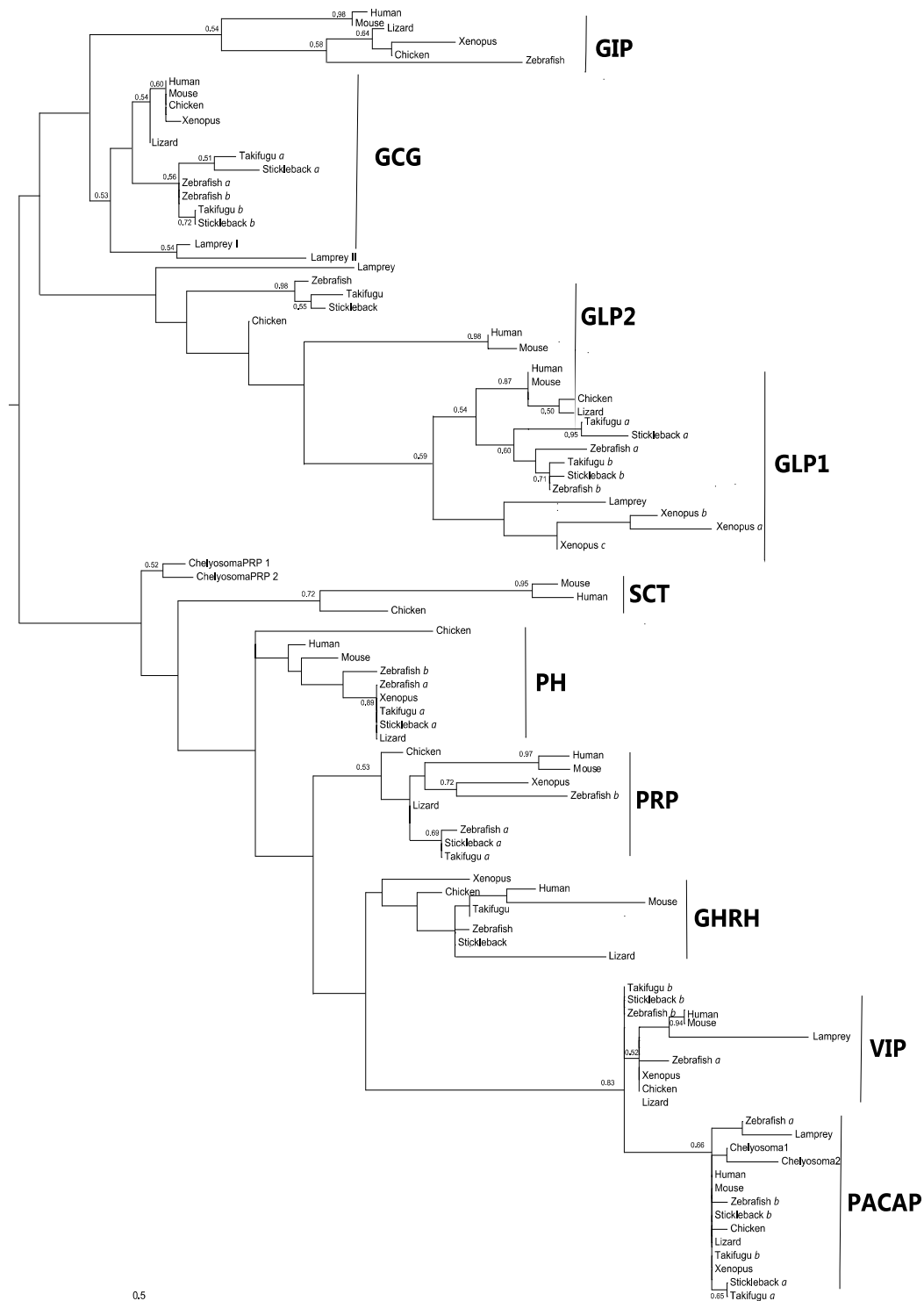


Figure 4 Evolutionary analysis of the chordate secretin family members. The maximum likelihood (ML) optimal tree topology is presented and was constructed with Phym1 3.0 [99]. ML bootstrap values higher than 50% are indicated at nodes and to facilitate interpretation a hypothetical root was added to the tree between the PACAP-like and GCG-like clades based upon gene structure evidence and proposed models for secretin family evolution. The different peptide groups are indicated and teleost duplicate genes are marked by *a* and *b*; *Xenopus* GLP1 exons by *a*, *b* and *c*. Accession numbers of the sequences used are described in Table 2 and for human and mouse members are: PHM/VIP (P01282 and P32648); PRP/PACAP (P18509 and O70176); GHRH (P01286 and P16043); GCG/GLPs (P01275 and P55095); GIP (P09681 and P48756); and SCT (P09683 and Q08535), respectively.

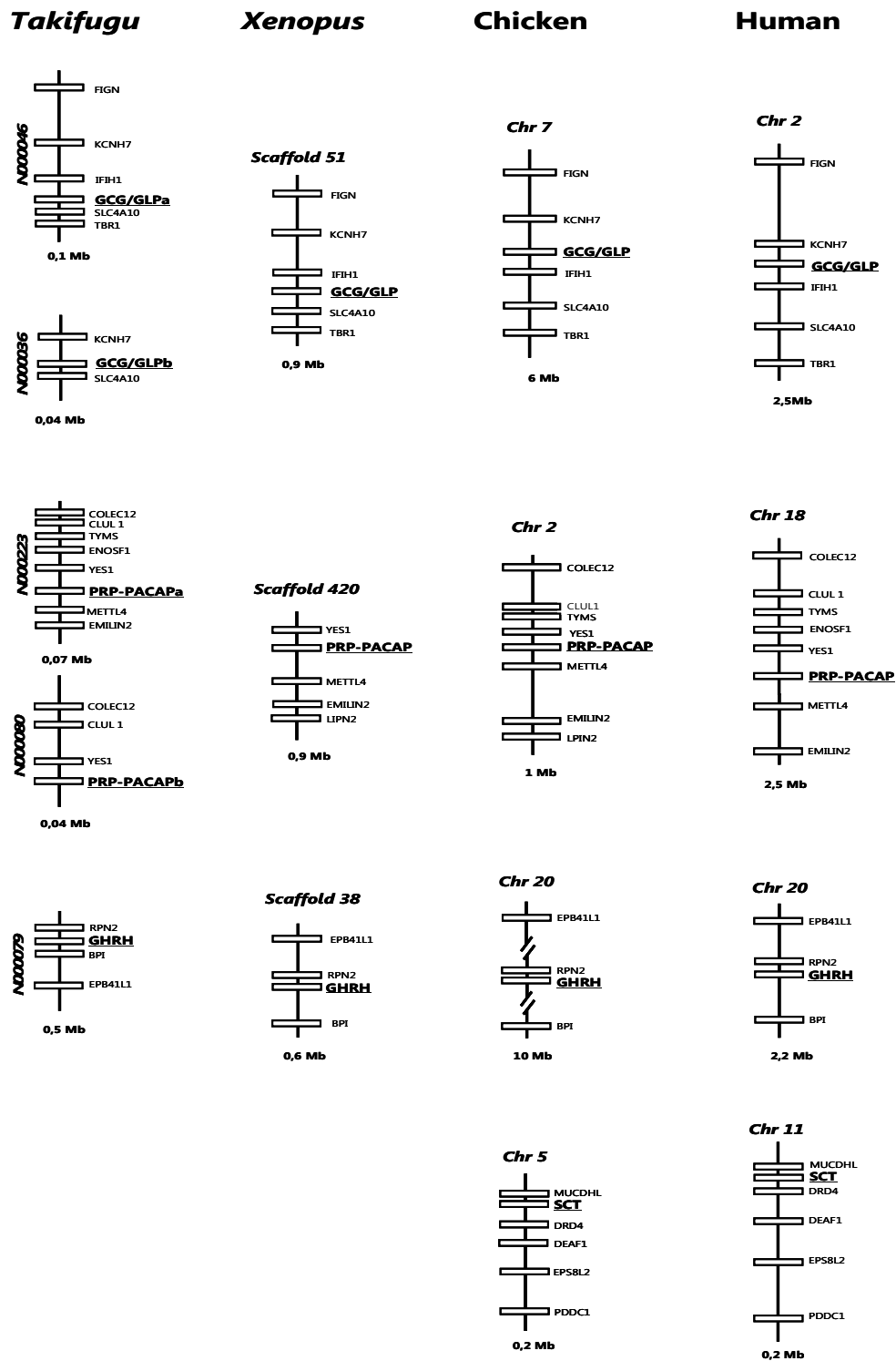


Figure 5 Gene environment comparisons of the GCG/GLPs, PRP/PACAP, GHRH and SCT genes in *Takifugu*, *Xenopus*, chicken and human. Homologue genes were identified using sequence similarity approaches with the *Takifugu* genes. *Takifugu* scaffolds are named according to the Assembly 4 available at [93] and have a direct correspondence with ENSEMBL (eg: N000046 corresponds to *Takifugu* Ensembl scaffolds_46). Genes were named based on HUGO annotation and the size of the genome regions analysed indicated within brackets. Genes are represented by boxes and genomic regions are indicated by lines. The figure is not drawn to scale and genes are positioned according to their relative distance in the genome assembly. For simplicity, only homologue genes are represented and GCG/GLP, PRP/PACAP, GHRH and SCT genes are edited in bold and underlined.

a conserved gene block prior to the emergence of the tetrapod gene (data not shown).

In order to identify a potential secretin family genome region in non-vertebrates, the conserved vertebrate gene environment was used to retrieve homologues in the *C. elegans*, *Drosophila* and *Ciona* genomes. Genes sharing similarity in sequence to those flanking the vertebrate GCG/GLP, PRP/PACAP and GHRH loci were identified although the genes encoding secretin family members were absent (Figure 6). In *C. elegans* the genes *drh-3*, *abts-1*, *src-1* and *M01A10.3* map to chromosome I and are respectively homologues of the human *IFIH1* and *SLC4A10* on chromosome 2, *YES1* on chromosome 18 and *RPN2* on chromosome 20. In *Drosophila*, *sei* and *CG6370* are localised in chromosome 2R and are the homologues of human *KCNH7* and *RPN2* and the fruit-fly *Dcr-2* and *CG14906* genes in chromosome 3R the correspondent in sequence of the human *IFIH1* and *METTL4*, respectively.

In the tunicate genome, homologues of the vertebrate secretin family flanking genes were identified scattered in the *Ciona* genome assembly. Homologues of human *KCNH7* (ENSCING0000000802, scaffold_167), *IFIH1* (ENSCING00000009744, scaffold_44) and *SLC4A10* (ENSCING00000006669, chromosome 9p) genes which flank GCG/GLP in vertebrates were identified. The conserved gene environment of the vertebrate PRP/PACAP genes: *YES1* (ENSCING00000008550 on scaffold_65) and *METTL4* (ENSCING00000013236 on chromosome 8q) genes are also present as well the putative urochordate homologue of the *RPN2* gene (ENSCING00000009122) conserved in the vertebrate GHRH genome region.

In the *C. elegans* genome chromosome 1 contained homologues of the genes which flank PRP/PACAP, GCG/GLP and GHRH in vertebrates. This suggests that this chromosome may be the protostome genome fragment that most resembles the potential metazoan ancestral secretin genome region from which the vertebrate members emerged. However, detailed analysis of chromosome positions (*src-1*, 1566932 to 1580204bp; *M01A10.3*, 5550508 to 5549145 bp; *drh-3*, 7820837 to 7826373 bp; and *abst-1*, 8307558 to 8296909 bp) revealed they are not mapped in close proximity and do not form a gene cluster. Moreover, no conserved linkage between the putative *Drosophila* and the *C. elegans* gene homologues exists.

Discussion

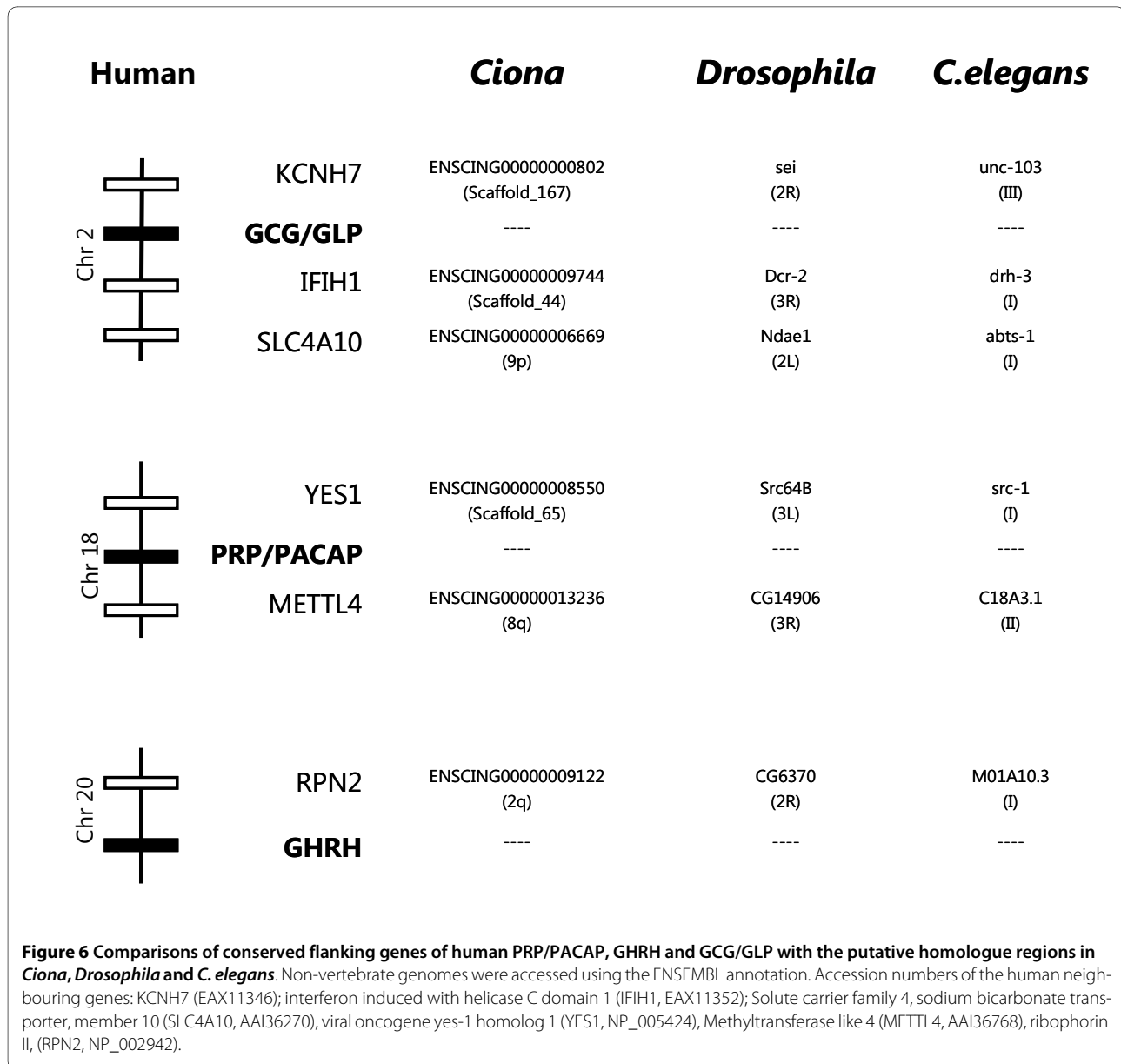
Comparative analysis of data from phylogenetically distant organisms is a major contributor for understanding gene and gene family evolution and the role of function and regulation in this process. The identification of gene homologues in vertebrates and early metazoan genomes provides a unique opportunity to perform comparative studies and to investigate gene family ancestries. The

secretin family is a well-studied group of peptides which activate specific receptors of family 2 GPCRs to bring about their pleiotropic actions in vertebrates. Secretin-like family GPCRs have been identified and cloned from non-vertebrate genomes [6,14,59,60] and their putative peptide ligands identified by immunohistochemistry (IHC) using antisera raised against the mammalian peptide homologues (Table 1). Both peptides and their corresponding receptors are proposed to have arisen by gene duplication events prior to the vertebrate radiation [2,6,8,10,11,18] and they represent an interesting model for studies of receptor-ligand evolution.

In the present study, despite extensive *in silico* database searches it was not possible to identify members of the secretin peptide family in non-vertebrates regardless of the report of a full-length PACAP cDNA in the tunicate, *Chelyosoma productum* [7] and cDNA of the PACAP coding exon in crab, cockroach, squid, planarian and *Hydra magnipapillata* [7,9]. Moreover, using the partial PACAP cDNA previously identified in *Hydra magnipapillata* (a cnidaria) to interrogate its sequenced genome [61] failed to identify the gene homologue.

It seems unlikely that the failure to identify PACAP in non-vertebrates was related to the methodology used as a similar approach has been successfully utilized to identify the gene encoding the active nonapeptide hormone, vasotocin/vasopressin in the amphioxus genome assembly and also the gene loci in several chordates including teleosts [62]. In fact, the existence of neurohypophysial hormones (eg. vasopressin and oxytocin) in deuterostomes and protostomes has been amply confirmed by the isolation of both genes and peptides from representatives of a number of different phyla (reviewed in [62]). Similarly, conserved sequence homologues of vertebrate neuropeptide Y family members have been isolated from molluscs and also from fruit-fly and mosquito and GPCR ligand-receptor pairs similar to the vertebrate NPY system have been characterised [63-65].

A comprehensive *in silico* analysis of the fully sequenced *Drosophila* genome identified and classified GPCRs and compared their number to putative neuropeptide ligands. Although 5 secretin family GPCR members were identified in the arthropod genome, only two potential ligands, corticotrophin releasing factor (CRF)-related peptides and *amnesiac* genes, unrelated to chordate secretin peptide family members were predicted [66]. In contrast, recent studies performed in molluscs (*Helix pomatia*) using IHC and MALDI-TOF/TOF identified partial peptides with a similar mass to vertebrate PACAP in the snail whole hemolymph and CNS extracts [67]. Should the identity of the peptide be confirmed by sequencing taking in consideration the results of the present study a new paradigm will be required to explain secretin family evolution.



There is evidence that genes for *amnesiac* in *Drosophila* and maxadilan in sand-fly might encode functional homologues of the vertebrate PACAP despite their lack of sequence similarity [2,68-70] and the maxadilan peptide is able to activate mammalian PAC₁ receptors *in vitro* [71,72]. The activation of family 2 GPCRs members by the secretin family of ligands has been linked to their well conserved structure [53,54] which comprises an alpha helix in the mid and C-terminal region and an N-terminal loop. The region of maxadilan implicated in PAC₁ receptor activation has an identical structure to the N-terminal region of secretin family ligands and contains key amino acids involved in receptor activation (reviewed by [69]). In fact, mutation analysis with maxadilan demonstrated

that despite its greater size compared to vertebrate secretin family peptides, the disruption of four conserved cysteine residues (1 - 5 and 14 - 51) responsible for the formation of two disulfide bonds led to loss of activity [69]. Taking into consideration the degree of conservation of the N-terminal ligand binding domain of the secretin family GPCRs [14] and their relative promiscuity [73] it is unsurprising that protostome peptides activate vertebrate receptors. In nematode and *Drosophila*, PDF (Pigment Dispersing Factor) stimulated the homologues of vertebrate secretin GPCRs but they were not stimulated by secretin family members [14,59,60]. These observations suggest that specificity of the receptor members has changed during evolution and may explain the failure to identify conserved ligands.

A general model to explain peptide ligand binding and receptor activation has emerged recently for class B GPCRs (secretin family) [74]. The proposed mechanism, known as the "two-domain model" suggests that initial ligand-receptor interactions are mediated by the central and C-terminal peptide segments with the extracellular N-terminal receptor region and that activation occurs subsequently when the bioactive N-terminus of the ligand binds the receptor juxtamembrane domain [75]. In this context, the identification in the present study of putative "signature" amino acids conserved across taxa for each peptide groups may explain differences in selection and affinity for receptors. For example, mutation of conserved residues within the C-terminal region of VIP revealed that substitution of L²³ (common to PACAP-like subfamily members) decreased peptide biological activity without altering the predicted structure [76]. D³ is conserved across the PACAP-like peptides and this residue has a role in adenylyl cyclase (AC) stimulation and interacts with basic residues (R¹⁸⁸ and K¹⁹⁵) in the second transmembrane helix of VPAC [77]. Similarly, D⁹ which is conserved across the GCG-like members is essential in the activation of mammalian GCGR [78]. It will be of interest in the future to study the role of unique amino acid residues/motifs identified in the present study within the N-terminal and C-terminal regions of each peptide group.

Recently, a Darwinian evolutionary model was proposed to explain the origin of steroid hormones and their receptors and may also explain the emergence of metazoan secretin peptide-receptor pairs. The steroid hormones and receptors were proposed to have evolved through a molecular exploitation process in which structurally adapted receptors evolved prior to ligand emergence [79,80]. By reconstructing the sequence of the ancestral steroid receptors (eg. corticoid receptor) the authors verified that they are activated by hormones (eg. aldosterone) that only emerged in the tetrapod lineage. At present, evolutionary comparisons of the metazoan receptor members and secretin peptide family suggests that, receptors emerged prior to the ligands which were subsequently acquired as a consequence of genome evolution in the chordate radiation. Generally, two major gene or genome duplication events are proposed to have occurred at the origin of vertebrates and have accompanied increased organismal complexity and emergence of gene novelties [81,82]. Whilst the majority of gene duplicates were probably lost as a consequence of their functional redundancy, some were fixed in the genome by the gain of new biological functions or partitioning the function of the ancestral counterpart and this may be the case for the secretin family GPCRs and their ligands [8,17,83].

The origin of the chordate secretin peptide family has been previously associated with the insect adipokinetic hormone (AKH) and AKH-Precursor Related Peptides

(APRP) precursor evolution [84]. Despite their low sequence similarity, the arthropod AKH and APRP precursor was found to share a similar gene organisation and comparable functions with vertebrate GCG and GHRH and they were proposed to have shared common origin prior to protostome-deuterostome divergence (approximately 600 million years ago). However, against this hypothesis is the recent demonstration that insect ADK signals through a gonadotropin-releasing hormone (GnRH) like receptor (members of family 1 GPCRs) and also the isolation of putative nematode AKH-GnRH related precursors suggesting that the invertebrate ADK members may share common evolution with the metazoan GnRH system [85,86].

Conclusions

The present study does not confirm the results of IHC studies in the early 70's and 80's which identified putative invertebrate secretin family members using antisera against mammalian peptides. It was not possible to identify sequence homologues of the *Chelyosoma productum* PRP/PACAP peptides in the sequenced *Ciona* genomes, although at least 8 putative secretin-like family GPCRs have been reported [14]. The previous facts taken with i) the identification in snail of a putative PACAP peptide; and ii) the activation of a secretin-like family GPCRs in nematode and *Drosophila* by PDF but not by vertebrate secretin family members makes it difficult to establish when the peptide members emerged in the deuterostome lineage. It is hypothesised that the emergence of the full suite of receptors and their ligands accompanied the rapid genome changes during chordate evolution. The ancestral secretin family gene probably arose as part of an existing gene or gene fragment and via exon and gene duplication events generated the existing suite of family members (Figure 3). This occurred after the emergence of the secretin family GPCRs and led to the establishment of novel and specific receptor ligand interactions that contributed to the generation of novel physiological functions. In contrast, to other peptide families, such as NPY/PYY and Oxytocin/vasopressin which stimulate receptors of family 1 GPCRs and are highly conserved from protostomes to deuterostomes, members of the secretin family GPCRs appear to have adopted new ligands during evolution.

Methods

Data mining

Using comparative sequence approaches the existence of putative non-vertebrate secretin family members were investigated in publicly available protostome and early deuterostome genome, EST and protein databases. The complete nucleotide and amino acid sequences of human secretin family members (PHM/VIP, P01282; PRP/

PACAP, P18509; SCT, P09683; GHRH, P01286; GIP, P09681; and GCG/GLP, P01275) mature peptides and their homologues in zebrafish and tunicate *Chelyosoma productum* PRP/PACAP precursors were used to interrogate databases (Figure 1, Additional file 1). Searches were performed in the metazoan genomes of porifera (*Amphimedon queenslandica*), cnidarians (*Nematostella vectensis* and *Hydra magnipapilata*), planarian (*Schmidtea mediterranea*), annelids (*Helobdella robusta* and *Capitella capitata*), mollusc (*Lottia gigantea*), crustacean (*Daphnia pulex*), insects (*Drosophila melanogaster*, *Aedes aegypti* and *Anopheles gambiae*); nematodes (*Caenorhabditis elegans* and *Caenorhabditis briggsae*); and in the early deuterostomes, sea urchin (*Strongylocentrotus purpuratus* Build 2), cephalochordate (*Branchiostoma floridae*) and urochordates (*Ciona savignyi* and *Ciona intestinalis*); and also in the vertebrate sea lamprey (*Petromyzon marinus*), teleosts (zebrafish, *Danio rerio*; *Takifugu rubripes*; *Tetraodon nigroviridis*; medaka, *Oryzias latipes*; stickleback, *Gasterosteus aculeatus*) and tetrapods, frog (*Xenopus tropicalis*), lizard (*Anolis carolinensis*) and chicken (*Gallus gallus*). Searches of the *Takifugu* genome were also performed in <http://fugu.nimr.mrc.ac.uk/blast>. To substantiate the results, further searches for secretin family members were also carried out in the NCBI EST data sets for porifera (Porifera (taxid:6040), cnidaria (Cnidaria (taxid:6073), protostomes (Protostomia (taxid:33317)) and early deuterostome (Echinoderms (taxid:7586); Cephalochordata (taxid:7735); Urochordata (taxid:7712) and also in species specific EST databases for the planarian (*Schmidtea mediterranea*), crab (*Celuca pugilator*), pacific oyster (*Crassostrea gigas*) and mussel (*Mytilus edulis*) (Additional file 1). In addition, the complete nucleotide precursor or the sequence corresponding to the deduced mature peptide of secretin family members in deuterostome were used to interrogate the general nucleotide (nr/nt) and protein databases (nr) available at NCBI [9] and UniProt [87] using the BLAST programme.

For small mature peptide sequences the BLAST algorithm was adjusted (scoring matrix PAM30, word size 2, highest expected value parameters, low complexity filter off, no adjustment) to permit identification of short peptide hits with strong similarities. Searches using short nucleotide sequences were also performed with word size 7; expected value 1000 and low complexity filter off. Best matches with significant scores or low E values <0.01 were retrieved and analysed.

Searches using a hidden Markov model were performed with the HMMER3 (3.03b) [88] suite of software on the NCBI non-redundant (nr) peptide, and custom-made invertebrate nucleotide (nt) (1614126 records) and est_others (ests minus human and mouse) (11209486 records) databases using hmmsearch. Subset databases

were constructed to reduce the computational burden of performing a HMMER3 search against the complete NCBI nucleotide and EST databases. The invertebrate subset databases were constructed by querying the NCBI databases using Entrez for all invertebrate GI numbers (Metazoa NOT Vertebrata) for both nt and est_others and filtered using fastacmd (part of the NCBI BLAST package). Peptide searches were performed with the Pfam model for the secretin peptide family members (Hormone_2 member, PF00123) and nucleotide queries with a model constructed in HMMER3 (3.03b) using the nucleotide aligned sequences of the 1-27 aa mature peptide regions of the secretin family members represented in Figure 2.

Sequence comparisons

The potential secretin family members identified were compared with existing vertebrate members. The non-vertebrate sequences that shared similarity with previously annotated genes or gene intron regions were discarded and the remaining candidates used to interrogate the NCBI database to confirm identity and failed to retrieve a homologue of the vertebrate secretin family. In contrast, homologues in vertebrate datasets were identified and the deduced amino acid sequences of the retrieved transcripts or predicted exon coding regions were compared using ClustalX 2.0 [89] with the conserved mature peptide region 1-27 aa of representatives of the secretin family with the exception of human GLP1 in which 5-32 aa were used that correspond to a unique coding exon. Peptide similarities/identities were determined using the GeneDoc programme [90] and amino acid consensus sequences within each peptide group were deduced and compared to demonstrate general levels of conservation for each subfamily. A prototype peptide representative of the chordate secretin family was constructed by fusing the most abundant amino acid residues within the PACAP-like and GCG-like peptides members. The *in silico* deduced sequence was submitted to Pfam [58] analysis to confirm identity as a secretin member and used to search the vertebrate and non-vertebrate NCBI databases with BLAST and adjusted parameters to identify homologues.

Phylogenetic analysis

Phylogenetic analyses were performed using the ClustalX 2.0 alignment of the 1-27 aa mature peptide region of secretin members. The amino acid sequence alignment produced was analysed with PROTTEST to select the model of protein evolution that best fits dataset [83] and phylogenetic analyses were conducted using 95 taxa with the maximum likelihood and Bayesian estimation methods (MrBayes and PhyloBayes, Additional file 2). The maximum likelihood analysis was carried out using

Phyml 3.0 [85] with 100 bootstrap replicates with a JTT substitution model with a discrete gamma distribution of rates among sites with 4 categories (Γ). A search for the optimal ML tree was also performed. Bayesian estimation using MrBayes [91] was performed with the Dayhoff model with Γ and PhyloBayes [92] with the CAT model plus Γ . The MrBayes analysis was conducted with two MCMC runs (each with 4 chains) for 200,000 generations with 20,000 samples. The PhyloBayes CAT analysis was performed using 2 independent run replicates (40727 and 40382 generations, respectively). Likelihoods were plotted against generation time and the MCMC chains were assumed to have reached stationarity when the curve plateaued. Phylogenetic sequence analysis was also performed using the nucleotide sequences of the 1-27 mature peptide domains with the ML method as previously described and the GTR model plus Γ and the individual peptide clades present in both nucleotide and amino acid trees obtained were similar (data not shown).

Gene environment comparisons

The gene environment of vertebrate secretin family members was determined to identify potentially conserved gene blocks; these were then used to search for putative ancestral secretin genome-like regions in *C. elegans*, *Drosophila* and *Ciona* assemblies. The *Takifugu* NIX annotated scaffolds [93] were used as a guide to characterise the *Xenopus* (*Xenopus tropicalis*) [94], the chicken and human homologue regions [95]. The NIX annotated scaffolds from *Takifugu* were used as they had greater information content than the homologue data deposited in ENSEMBL. The conserved vertebrate gene environment identified the genes YES1 and METTL4 for PRP/PACAP; the gene RPN2 for GHRH; and the KCNH7, IFIH1 and SLC4A10 genes within the vertebrate GCG/GLP genomic region. The, *C. elegans*, *Drosophila* and *Ciona* genomes were assessed using the ENSEMBL assembly annotation and homologues identified and compared with the vertebrate homologue region.

Additional material

Additional file 1 Publicly available genome and EST databases searched using the nucleotide and mature peptide sequence of the tunicate (*Chelysoma productum*) and human secretin family members.

Additional file 2 Phylogenetic support values (bootstrap proportions, bp; and posterior probabilities, pp) of the secretin family peptide clusters constructed with maximum likelihood (ml) (Figure 4), MrBayes and PhyloBayes methods.

Abbreviations

VIP: (Vasoactive Intestinal Peptide); PACAP: (Pituitary Adenylate Cyclase-Activating Polypeptide); SCT: (Secretin); GHRH: (Growth Hormone Releasing Hormone); PRP: (PACAP-Related Peptide); PH: (Peptide Histidine); PHI: (PH-Isoleucine); PHM: (PH-Methionine); GCG: (Glucagon); GLP: (GCG-like); GIP: (Glucose-dependent Insulinotropic Peptide); GPCRs: (G-Protein Coupled Receptors).

Authors' contributions

The majority of the practical work was carried out by JCRC in collaboration with FAV and ASG. JCRC and DMP planned the study, wrote, analysed the data and critically revised the manuscript for important intellectual content. All authors read and approved the final manuscript.

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Author Details

Centre of Marine Sciences (CCMAR), Universidade do Algarve, Campus de Gambelas, 8005-139 Faro, Portugal

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