The GnRH system in the European sea bass (Dicentrarchus labrax)

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Abstract

The cDNA sequences encoding three GnRH forms, sea bream GnRH (sbGnRH), salmon GnRH (sGnRH) and chicken GnRH II (cGnRH II), were cloned from the brain of European sea bass, Dicentrarchus labrax. Comparison of their deduced amino acid sequences to the same forms in the gilthead sea bream, Sparus aurata, and striped bass, Morone saxatilis, revealed high homology of the prepro-cGnRH II (94% and 98% respectively), and prepro-sGnRH (92% to both species). The sbGnRH exhibited dissimilar identities, with high homology to the striped bass (93%), and lower homology (59%) to the gilthead sea bream. Two transcript types were identified for the GnRH-associated peptide (GAP)-sGnRH as well as for the GAP-cGnRH II, which suggests a possible alternative splicing followed by the addition of an early stop codon. In order to obtain antibodies specific for the three GnRH precursors, recombinant GAP proteins were produced. The differential expression of the three GnRHs previously reported in the brain by means of in situ hybridization, using riboprobes corresponding to the GAP-coding regions, was fully confirmed by immunocytochemistry using antibodies raised against the recombinant GAP proteins, indicating that the transcripts are translated into functional proteins. Moreover, this approach allowed us to follow, for the first time, the specific projections of the different cell groups: sGAP fibers are distributed mainly in the forebrain with few projections reaching the pituitary, sbGAP fibers are mainly present in the preoptic area, mediobasal hypothalamus and predominantly project to the pars distalis of the pituitary, whereas cGnRH II fibers have a widespread distribution primarily in the posterior brain, and do not project to the pituitary. These new tools will be extremely useful to study further the development, regulation and functional significance of three independent GnRH systems in the brain of vertebrate species.


Introduction

Gonadotropin releasing hormone (GnRH) is a neuro-peptide known to regulate gonadotropin expression and secretion in vertebrates. To date, fourteen forms of GnRH decapetides have been found in vertebrates, in which at least one unique form has been determined from a representative species of all vertebrate classes. Ten GnRH forms out of the fourteen were found in fish and six of them in teleosts: chicken GnRH II (cGnRH II; Miyamoto et al. 1984), mammalian GnRH (mGnRH; Matsuo et al. 1971, Burgus et al. 1972), salmon GnRH (sGnRH; Sherwood et al. 1983), catfish GnRH (cfGnRH; Ngamvongchon et al. 1992), sea bream GnRH (sbGnRH; Powell et al. 1994), herring GnRH (hGnRH; Carosfeld et al. 2000) and Medaka GnRH (mdGnRH; Okubo et al. 2000). cGnRH II is present in all vertebrate classes examined to date, and the mGnRH form is present in primitive non-teleostean bony fish, primitive teleosts (eels and others), amphibians and mammals. sGnRH, cfGnRH and sbGnRH are found in higher teleosts. This taxonomic distribution indicates that cGnRH II and mGnRH are ancient vertebrate forms that gave rise to other variants.

The typical structure of the GnRH precursor protein consists of 1) a signal peptide at the N-terminal (about 23 amino acids (aa) in size), 2) the GnRH decapetide which is the bioactive peptide, followed by a 3 aa cleavage site (Gly-Lys-Arg) and 3) a GnRH-associated peptide (GAP) at the C-terminal (about 60 aa in size). The bioactive
decapptide is processed from the precursor by removal of the signal peptide and cleavage at the dibasic aa (Lys–Arg) (Klungland et al. 1992). The GnRH decapptide is highly conserved at positions 1–4, 6, 9–10 (Sower 1997). While the decapptide sequence of any GnRH form is highly conserved among species, the signal peptide and the GAP are more diverse both in their amino acid and nucleotide sequences. Although still controversial, the function of the conserved three peptides.

Table 1 Oligonucleotides used for PCR amplification of the three GnRH forms

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Oligonucleotides used for PCR amplification of the three GnRH forms

Materials and Methods

Isolation of European sea bass GnRH and GAP cDNAs: salmon, sea bream and chicken II forms

Total RNA was extracted from the brain of sexually mature female sea bass using a scaled-down guanidinium isothiocyanate method (Chomczynski & Sacchi 1987). cDNA was synthesized using 1 µg total RNA, AMV reverse transcriptase (Promega, Madison, WI, USA) and the universal oligo dT-adaptor primer (Frohman 1990) (no. 15, Table 1). Prior to use in 5′ RACE, the cDNA was tailed with terminal deoxynucleotidyl transferase (Promega) using dCTP.

Primers specific for each gap form were designed according to the striped bass GAP sequences (Genbank...
accession numbers: sbGnRH, AF056314; cGnRH II, AF056313; sGnRH, Y Zohar, personal communication). The first PCR amplification was carried out using primers located 5’ to the GAP sequence (Table 1: nos 3, 4, 5), followed by a nested PCR using a set of primers designed for the amplification of the full length GAP sequence (nos 6 and 9 for sbGAP, nos 7 and 10 for sGAP, nos 8 and 11 for cIIGAP). The nested primers contained specific restriction sites and a stop codon in the 3’ primer to facilitate introduction into an expression vector.

The full-length cDNA sequences of the three GnRH forms were isolated by 5’ RACE PCR amplification. 3’ gene specific primers (nos 9, 10, 11) and oligo-d-G adaptor 2 (no. 1) amplified oligo-dC tailed cDNA template. Nested PCR amplification was carried out using adaptor 2 (no. 2) and 3’ gene specific primers (nos 12, 13, 14). The total PCR products were cloned into pGem-T easy vector (Promega) and the clones containing the GnRH cDNA were identified by colony hybridization using the corresponding specific GAP sequence as a probe.

All PCR reactions were carried out in 25 μl using 1 unit Taq DNA polymerase (Promega), buffer (Promega), 1·5 mM MgCl2, nucleotides (0·2 mM final concentration of each nucleotide), 12·5 pmol of each primer and in a final dilution of 1:50 cDNA. Cycling parameters were: 3 min denaturation at 94°C followed by 30 cycles of 1 min denaturation at 94°C, 1 min annealing at 50°C and 1 min extension at 72°C (last extension step lasted 10 min).

The chosen PCR products were isolated by excision from agarose gel or direct clearing from PCR reaction using purification kits (gel extraction kit or PCR purification kit, Qiagen) and the clones containing the GnRH cDNA were ligated into pGEMT-Easy plasmid (Promega) and the clones containing the GnRH cDNA were identified by colony hybridization using the corresponds specific GAP sequence as a probe.

DNA sequencing
Double stranded plasmid DNA was sequenced either using the Sequenase version II kit (Amersham Pharmacia Biotech AB, Uppsala, Sweden) or using enzyme F S Taq dye deoxy terminator cycle sequencing at the DNA Sequencing Biological Services (Weizmann Institute, Rehovot, Israel). Universal primers, SP6 and T7, as well as gene-specific primers were used to obtain the complete sequence of both DNA strands.

Production of recombinant GAP proteins
The sequences encoding for the three GAPs were engineered into the PQE-31 expression vector (Qiagen) which contains a 6×HIS residue tag 5’ to the polylinker and produces N-terminal 6×His tagged recombinant proteins. The processes of ligation, transformation into M15 [pREP4] cells, expression of the protein and purification (using Ni-NTA agarose columns) were performed using the QIAexpress kit (Qiagen) under denaturing conditions, according to the manufacturer’s protocol.

The sequence encoding for the cIIGAP was engineered into pGEX-2 expression vector by using a 5’ gene-specific primer (5’GAATTCCAGGCAGGGAACTG3’) which was designed especially for insertion in the correct reading frame. This system expresses a recombinant protein which is fused in its N-terminal end to glutathione transferase. The construct was transformed into DH5α E. coli cells which were grown in the presence of ampicillin (50 μg/ml). Protein expression was induced by addition of 1 mM IPTG and further incubation for 3 h. The bacterial cells were then harvested by centrifugation (5000 x g, 15 min, 4°C), the pellet was frozen overnight at −20°C and resuspended in 10 ml ice-cold PBS/50 ml bacterial culture. Cells were lysed in the presence of lysozyme (20 μg/ml) for 45 min on ice, followed by 5–6 rounds of 15 s sonication (Microson cell disruptor model XL2005, microprobe 2·4 × 11·5 mm at level 18 of the scale). Triton X-100 was added to the supernatant to a final concentration of 1% followed by centrifugation at 10 000 × g for 5 min at 4°C.

One milliliter of a 50% slurry of glutathione–agarose beads (Sigma, St Louis, MO, USA) was added to the supernatant and the sample mixed for 2 min. The beads were then washed twice by adding 50 ml cold PBS and centrifuged for 10 s at 500 × g. Fused recombinant protein was eluted with 1 ml 50 mM Tris/Cl (pH 8·0)/5 mM reduced glutathione, mixed for 2 min and centrifuged for 10 s at 500 × g.

Generation of specific antibodies against recombinant GAPs
The purified recombinant GAPs were used for the immunization of guinea pigs. Nine female guinea pigs (5/6 weeks old; 345–365 g), three for each GAP, were acclimated for one week before the beginning of the immunization protocol. Each injection consisted of 25 μl recombinant GAP (1 μg/μl), 75 μl distilled water, and 100 μg incomplete Freund’s adjuvant. Preimmune sera were collected on the day of the first injection and stored at −20°C. The first injection was intradermal in the back (6–7 points) and the others, at one month intervals, were given intramuscularly in the hind legs. Eight days after the fourth injection, blood was collected under anesthesia by cardiac puncture, centrifuged and the serum was stored at −25°C.

Dot blot analysis for antibodies specificity
The specificity of antisera was assessed by dot blot analysis using the recombinant sGAP, sbGAP and cIIGAP proteins as antigens. Recombinant GAP proteins (6–12 μg) were
added to individual nitro-cellulose filters. The slices were blocked for 1 h in Tris-buffered saline buffer (TBS; 25 mM Tris, 140 mM NaCl, 2.7 µM KCl, pH 7.4) containing 5% non-fat dry milk. Samples were then incubated for 12 h with the primary antisera at a 1:2000 dilution followed by several washes with TBS containing 0·05% Tween 20. The slices were incubated for 1 h in peroxidase-conjugated anti-guinea pig IgG (Jackson ImmunoResearch Laboratories, West Grove, PA, USA) in 1:3000 dilution, and washed as before. The filter was developed in 0·024% 4-chloro-1-naphthol (Sigma) in the presence of 0·002% H₂O₂ in 50 mM Tris buffer.

In situ hybridization

The in situ hybridization procedure was according to the Anglade et al. (1993) protocol. Two vitellogenic females and two spermatizing males were used in the process, as described in detail by González-Martínez et al. (2001). It is important to note the use of riboprobes corresponding to the three GAP regions.

Immunohistochemistry

Fish (two males and two females) were anaesthetised with 2-phenoxyethanol (Sigma) and perfused via the aortic bulb with Bouin’s fixative (4% paraformaldehyde and 0·2% picric acid in 0·1 M phosphate buffer, pH 7·4). Brains, with the pituitary gland attached, were then removed and post-fixed in the same fixative for 5–7 h in the dark at 4 °C. After overnight cryoprotection in 0·1 M phosphate buffer containing 15% sucrose, brains were embedded in Tissue-Tek, frozen in cold isopentane and serial coronal and sagittal sections of 16 µm thick were obtained with a cryostat.

Immunohistochemical staining was performed using a streptavidin–biotin–peroxidase complex method. Endogenous peroxidase activity was blocked with 1% hydrogen peroxide in Coons buffer (CBT, 0·01 M Veronal, 0·15 M NaCl) containing 0·1–0·2% Triton X-100 for 30 min. Before immunostaining, sections were transferred for 5 min to CBT and then saturated in CBT containing 0·5% casein for 30 min. The sections were incubated overnight in a moist chamber at room temperature with anti-sGAP, anti-sbGAP and anti-cIGAP antisera (1:500, 1:1000 dilution in CBT/0·5% casein). The sections were then washed in CBT (2 × 15 min) and incubated for 1·5 h at room temperature with Biotin-sp–Conjugated-AffiniPure Goat Anti-GuineaPig–IgG (Jackson ImmunoResearch Laboratories) diluted 1:1000 in CBT. After washing in CBT (2 × 15 min), the sections were incubated for 1·5 h at room temperature with peroxidase–conjugated–streptavidin complex (Jackson ImmunoResearch Laboratories) diluted 1:1000 in CBT. Finally, the sections were washed in CBT followed by Tris–HCl (0·05 M, pH 7·4) and peroxidase activity was visualized either in 0·05 M Tris–HCl, pH 7·6 containing 0·025% 3,3 diaminobenzidine tetrahydrochloride (Sigma) and 0·01% hydrogen peroxide or in 0·04% 4-chloro-1-naphthol (Sigma) and 0·01% hydrogen peroxide. To confirm the specificity of the immunostaining, controls were performed by replacement of primary antisera with the corresponding preimmune sera and omission of primary or biotinylated antisera.

Both in situ hybridization and immunohistochemical sections were analyzed on a Leica photomicroscope and computer images were obtained with a Sony DKC-CM30 digital camera (Sony, Japan). The software used was Adobe Photoshop 5·0 and no subsequent alterations have been made. The determination of brain nuclei was made using a sea bass brain atlas (J M Cerdà-Reverter, S Zanuy & J A Muñoz-Cueto, unpublished results).

Results

Isolation of the cDNA sequences encoding three GnRH forms

PCR was used to amplify the cDNA sequences encoding the 5’ untranslated region (UTR), signal peptide, GnRH decapeptide, GAP and 3’ UTR of sGnRH, sbGnRH and cGnRH II (the latter is missing the 3’ UTR). 5’ and 3’ RACE were carried out on both hypothalamus and brain total RNA.

Salmon GnRH

The sGnRH cDNA was amplified using 5’ RACE and 3’ RACE. Two types of clones were obtained, one was 357 bp in length and contained 108 bp of the 5’ UTR. The second was 255 bp in length and contained only the last 6 bp of the UTR. The presence of the two 5’ RACE products was RNA dependent rather than RT/PCR dependent. This may indicate the presence of more than one potential transcription initiation. The full cDNA sequence consisting of 567 bp of the prepro sGnRH can be found in Genbank accession number AF224280. The cDNA sequence shows a high homology of 88–92% when compared with sGnRH precursors of other fish. The signal peptide identity is 78–83% and the GAP identity is 89–95% (Table 2).

Sea bream GnRH

The sbGnRH sequence was amplified using 5’ RACE and 3’ RACE. The full cDNA sequence of 394 bp, encoding for 99 aa, can be found in Genbank accession number AF224279. The cDNA sequence identity compared with other teleost sbGnRH precursors ranges from 50% in the case of the African cichlid to 92% in the striped bass. The leader peptide identity ranges from 32–86% and the GAP homology ranges from 35 to 94% (Table 2).

cGnRH II

The cGnRH II cDNA sequence of 462 bp in length that encodes for 85 aa of prepro-cGnRH II was amplified using 5’ and 3’ RACE. The nucleotide
sequence and the deduced amino acids sequence are available in Genbank accession number AF224281. The cGnRH II cDNA sequence shows high homology to other fish cGnRH II precursors, which ranges from 88%-99%. The cGnRH II leader peptide of the striped bass is completely homologous to that of the sea bass and the GAP sequences differ in only one amino acid. Comparison with the African cichlid and the sea bream preproGAP cGnRH II shows a slightly lower homology, which ranges from 88–96% (Table 2).

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<th>Fish species</th>
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<tr>
<td>Sea bream</td>
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### Salmon-, sea bream- and chicken II-GAPS

To raise anti-GAP specific antibodies, recombinant GAP proteins of each form were prepared in a bacterial expression system. The sequences encoding the three GAPs were amplified by PCR using GAP-specific primers (Table 1). Primers 7 and 10 for the sGAP, primers 6 and 9 for the sbGAP and primers 8 and 11 for the cIIGAP were used.

**Salmon GAP** Two transcripts of salmon GAP were amplified, one of 174 bp and the other of 681 bp. The larger product, which appeared to be more abundant, was an unprocessed mRNA GAP. Partial or full length introns 2 and 3 were recognized by the invariant GT and AG dinucleotides found in the 5’ and 3’ intron borders. Intron 2 containing 205 bp was located after amino acid 13, and intron 3 with 302 bp was located after amino acid 40 of the GAP. The short product encodes the predicted 56 amino acids of the GAP. However, a stop codon (tga) present in intron 2 may give rise to the translation of a shorter GAP peptide i.e. 10 amino acids (Fig. 1A).

**Chicken II GAP** RT-PCR amplification of the cIIGAP resulted in two transcripts, one of 450 bp and the other of 156 bp. Similar to the case of the sGAP, the 450 bp product of the cIIGAP appeared to contain introns 2 and 3. Again, an early stop codon (TAA) was found, which is located in exon 3 and may encode for a different peptide of 46 amino acids. Intron 2 spanned 77 bp, located after aa 12 of the GAP, and intron 3, 196 bp, located after aa 40. The small PCR product was the fully processed form (Fig. 1B).

### Sea bream GAP

Only one transcript of the sbGAP was obtained by PCR amplification using primers 6 and 12. The transcript was 177 bp in length and seemed to be the fully processed form which encodes the predicted 59 amino acids of the GAP.

The RT-PCR product presented here represents mRNA and not genomic contamination for the following reasons: 1) the sGAP larger product was much more abundant than the smaller one, and 2) using the cDNA for sbGAP amplification resulted in only one transcript. Thus, it is possible that the larger products are a result of an alternative splicing of the original genes.

### Production of recombinant GAP peptides

All three GAP cDNAs were introduced into the bacterial His-tagged protein expression vector, pQE-31. The recombinant GAPs were expressed and purified. Analysis on 17% PAGE to detect the production of recombinant GAP peptides resulted in clear bands of the expected size of 6-5 kDa for the sea bream and the salmon GAPs. However, no recombinant cIIGAP could be detected. Following the purification process, which included binding to NI-NTA column and elution under denaturing conditions, clear clean bands for both sea bream and salmon GAPs were obtained, with some degradation seen in the late eluting fractions (Fig. 2A and B).

The cIIGAP was engineered into the pGEX2 vector and expressed as a fusion protein to glutathione S-transferase (GST). The purification procedure included binding to glutathione-agarose beads and elution with glutathione solution. SDS PAGE analysis of the eluted fractions revealed two bands: the cIIGAP/GST fused protein as a dominant band, 34 kDa in size, and a smaller faint band of 27 kDa which corresponds to the GST protein on its own (Fig. 2C).

### Production of antibodies

The three recombinant GAP proteins were injected into guinea pigs, three animals for each GAP, to obtain polyclonal antibodies. All nine antisera appeared to contain high levels of specific anti-GAP IgGs. As shown in Fig. 3,
A.

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Immunohistochemistry of GAP-expressing cells and fibers
To ensure that the GAP antibodies were suitable for the localization of preproGnRH expressing structures, the distribution of GAP-immunoreactive (ir) cells were compared with that of the corresponding GAP mRNAs, as detected by in situ hybridization. A very good correspondence between the distribution of GnRH-expressing perikarya and preproGnRH mRNA-containing cells was observed in all brain regions (Fig. 4). Cell bodies containing either sGAP mRNA or protein were mostly detected in the olfactory bulbs, either in the glomerular layer or in the ganglion cells of the terminal nerve (Fig. 4A and D).

B.

GAACCTGGACTCTTTTGGCAGTTCAGAGttgggatcagtttttctattttt
Cells expressing sbGAP mRNA or protein were detected in the caudal ventral telencephalon, the ventral preoptic region (Fig. 4B and E) and, more caudally, in the mediobasal hypothalamus. Finally, large cells expressing both cIIGAP mRNA and protein were only detected in the medial longitudinal fascicle of the synencephalon (Fig. 4C and F). The sGAP-ir fibers were more evident in the olfactory bulbs, ventral telencephalon, preoptic area, dorsal hypothalamus, ventral thalamus, pretectum and dorsal tegmentum. However, a conspicuous sGAP innervation was also observed in the dorsal telencephalon, optic tectum and ventral rhombencephalon. The pituitary gland only receives a relatively small number of sGAP-ir fibers. Sea bream GAP immunoreactive (sbGAP-ir) fibers were observed only in the ventral surface of the forebrain, associated with the ventral telencephalon, preoptic area and hypothalamus. These sbGAP-ir fibers strongly innervate the pituitary gland of sea bream. Finally, chicken-II GAP immunoreactive (cIIGAP-ir) fibers were abundant in the brain of sea bass, being especially evident in dorsal and ventral telencephalon, periventricular preoptic area and hypothalamus, dorsal and ventral thalamus, pretectum, posterior tuberculum, mesencephalic tectum and tegmentum, cerebellum, and rhombencephalon. No cIIGAP-ir fibers were detected in sea bream hypophysis.

Discussion

In this study, we have cloned three GnRH cDNAs encoding the precursors of three peptides from the European sea bass, providing further information on the presence of three variants in the brain of a perciform fish. Furthermore, we have produced recombinant GAP
proteins and raised specific antibodies suitable for the immunohistochemical detection of structures expressing preproGnRH.

The European sea bass GnRH precursors exhibit the same general organization as all other known GnRH precursors: a signal peptide at the N terminus, the GnRH decapeptide, a conserved processing tripeptide sequence and the GAP region at the C-terminus. This typical organization suggests that all these preproGnRHs have a common ancestral origin.

The cDNA encoding for the precursors of the sGnRH and the cGnRH II have 87%-99% sequence identity with the corresponding sequences in other perciforms (Table 2, Fig. 5A and C). Both the signal peptide and the GAPs are quite conserved. However, the sbGnRH precursor sequence identity was approximately 90% with the striped bass and dropped to 40–60% with the sea bream and the African cichlid (Table 2, Fig. 5B). Interestingly, this suggests that the gene coding for the sbGnRH precursor has evolved faster than those coding for sGnRH and cGnRH II and also that sbGAP does not have strong physiological significance.

Two kinds of transcripts were found for the chicken II and salmon GAPs: a processed and an unprocessed form. The existence of introns in GnRH transcripts was previously reported in several studies: in the gilthead sea bream brain, the sGnRH form contains intron 1 as a 5’ UTR (Gothilf et al. 1996). Similarly, the sGnRH expressed in the gonads of the rainbow trout retains intron 1 to generate different sizes of 5’UTRs (Von Schalburg & Sherwood 1999). In the ovary of sea bream, sbGnRH and cGnRH II transcripts retained all three introns, and the possibility of an alternate stop codon that changes the GAP amino acid sequence is suggested (Nabissi et al. 2000). In the sea lamprey, three transcripts of the lamprey GnRH I form, differing in the length of GAP coding region due to alternative splicing in intron 2 were reported (Suzuki et al. 2000). In this study, the different transcripts of the cGnRH II and sGnRH were found to be expressed simultaneously in the brain of the European sea bass. Similar to the finding in the sea bream gonads, an alternative stop codon that truncates the amino acid sequences of the GAPs was found. The phenomenon of multiple transcripts of the same GnRH gene suggests that one GnRH form may have different functions.

The recent evidence that the brain of modern teleosts expresses three different GnRH genes encoding three different GnRH forms raises interesting questions in terms of distribution, functional significance, regulation and ontogenesis. This warrants thorough investigations with respect to the distribution of the cells expressing the different forms and their respective projections in adult animals and throughout development.

![Figure 4](http://www.endocrinology.org/journal/172/105-116/N ZMORA and others · Localization of GnRH in the European sea bass brain)
Until now, the organization of central GnRH systems has mainly been studied by means of immunohistochemistry using antibodies more or less specific for the endogenous decapeptides. Despite this lack of specificity, this technique has allowed the establishment of the organization of the GnRH systems in teleosts, and the presence of immunoreactive GnRH cells not only in the ventral forebrain, but also in the midbrain tegmentum (Münz et al. 1981, Dahmen et al. 1986). But specificity turned out to be a critical problem when it became clear that different GnRH forms were produced in the brain of a single species (Yu et al. 1988). Nevertheless, using well-characterized antibodies, several studies succeeded in showing the differential expression pattern of two different GnRHs, for example in the brain of the Masu salmon (Amano et al. 1991), the Siberian sturgeon (Leprêtre et al. 1993), the European eel (Montero et al. 1994) or the goldfish (Kim et al. 1995). The cDNA sequences encoding different GnRH forms allowed this problem to be approached by using in situ hybridization. This was first achieved in the African catfish to map the cell expressing cGnRH and cGnRH II by selecting probes corresponding to the highly divergent cGAP and cIIIGAP sequences (Zandbergen et al. 1995).

The present study investigates the differential expression of three GnRH genes at both the mRNA and protein levels. Indeed, the distribution of the cells expressing prepro-sGnRH, -sbGnRH and -cGnRH-II mRNAs has already been studied in different perciforms (White et al. 1995, Gothilf et al. 1996, Okuzawa et al. 1997, Parhar et al. 1998). However, the respective projections of the different cell groups remained unknown. The results of our in situ hybridization study confirm the differential expression of the three forms by showing that sGnRH mRNA is mainly detected in the olfactory bulbs, sbGnRH mRNAs in the preoptic area and cGnRH II in the synencephalon, as shown in other perciforms (White et al. 1995, Gothilf et al. 1996, Okuzawa et al. 1997, Parhar et al. 1998, White & Fernald 1998). Because of the high similarity in the primary sequences of the three corresponding GnRH decapeptides, it was decided to generate antibodies against recombinant GAPs. This enabled us to circumvent the problem of specificity. It also allowed the use of a longer peptide to generate antibodies without any coupling to a carrier protein. The preliminary results obtained with all three antisera designed for each GAP fully confirmed the distribution of the corresponding mRNAs as well as the strict specificity of the antisera. The overall distribution of the GnRH-expressing cells also confirms what was previously reported in the same species using antibodies against sGnRH (Kah et al. 1991). The antibodies used in this latter study most likely recognized both sGnRH and sbGnRH, since the distribution of GnRH cell bodies corresponded exactly to the sum of sGAP- and sbGAP-immunoreactive cells detected in the present work. These results not only demonstrate that translation occurs in the different mRNA-expressing cell groups, but also that these antisera against GAPs are suitable to map the different GnRH systems.

In addition to confirming the distribution of the different preproGnRH mRNAs, these antibodies demonstrated that sGAP immunoreactive fibers project mainly to the forebrain, but that a few projections also reach the pituitary. Fibers immunoreactive for sbGAP were mainly observed in the preoptic area, the mediobasal hypothalamus and predominantly project to the pars distalis of the pituitary. The cIIIGAP-positive projections are widespread in the brain of sea bass but mainly concerned the posterior brain and did not reach the pituitary.

Figure 5

Comparison of GnRH precursor sequences of perciform representatives with the sea bass precursors. The GnRH decapeptide is underlined. Homologous amino acids are shown as dashes. (A) sGnRH precursor. (B) sbGnRH precursor. (C) cGnRH II precursor.
sbGnRH innervation. This assumption is consistent with previous physiological studies performed in another periconform species, the gilthead sea bream (Powell et al. 1994, Gothilf et al. 1996, Holland et al. 1998) as well as in the European sea bass (Rodriguez et al. 2000). Moreover, sbGnRH mRNA levels in the brain of both species were shown to fluctuate in correlation with ovarian development, suggesting their relevance in this process (Gothilf et al. 1997, Rodriguez et al. 2000). However, the present results also indicate a minor contribution of sGnRH to the pituitary, although the precise origin of these projections remains to be established. In the dwarf gourami, the terminal nerve ganglion cells have been shown to project widely in the brain, but not in the pituitary (Oka & Matsushima 1993). It is generally believed that GnRH-immunoreactive cells of the terminal nerve are not essential for the regulation of gonadotropin release, an assumption strongly reinforced by the fact that sectioning the olfactory tract in female goldfish does not impair gonadal development and ovulation (Kobayashi et al. 1994). Thus, sGAP-ir fibers entering the pituitary of sea bass could also be involved in the regulation of the functions of other adenohypophyseal cells.

At the present stage, it is not known whether the existence of three GnRH genes in periconform is the result of recent gene duplication in modern teleosts or if the situation is similar in other fish families or even in other classes of vertebrates. However, recent data obtained in herring also indicate the existence of three GnRH variants in the brain of this very primitive teleost (Carosfeld et al. 2000), which suggests that this feature could be common to most teleosts. Phylogenetic analyses indicate that sbGnRH-coding sequences form a branch different from that of sGnRH and cGnRH II-coding sequences (J A Muñoz-Cueto & O Kah unpublished data, Gothilf et al. 1995, White et al. 1998). In addition, there is recent evidence in frog that the brain of amphibians also expresses sGnRH and cGnRH II-coding sequences (J Año-Cueto JA & Kah O 1996). It is generally believed that a third GnRH peptide, possibly sGnRH is expressed in higher vertebrates, including humans (Yahalom et al. 1999). Furthermore, it has been shown in the rhesus macaque that two different populations of GnRH-immunoreactive neurons differentiate during development with different morphological features and a different final distribution in the brain (Quanbeck et al. 1997). On the other hand, a recent study based on promoter transgensics in mice suggests the existence of different GnRH neuronal populations of different embryological origins during development (Skenner et al. 1999).

Altogether, our results indicate that the molecular and immunological tools generated in the sea bass are highly specific and are suitable for future studies aimed at investigating the development, organization, regulation and functional significance of three GnRH systems in the brain of a single species.

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