Localization and analysis of natriuretic peptide receptors in the gills of the toadfish, *Opsanus beta* (teleostei)

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This text continues with detailed analysis of the distribution and characterization of natriuretic peptide receptors in the gills of *Opsanus beta*. It discusses the types of receptors, their localization, and the methods used for their analysis. The text highlights the role of NPs in cardiovascular and osmoregulatory processes and their presence in the gills of teleost fish, including *Opsanus beta*. The study used autoradiography, competition assays, and sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) to characterize the receptors. The findings contribute to the understanding of NP physiology in different animal models, including teleost fish.
after netting or trapping in the Gulf of Mexico. Fifteen animals were used for the study. The fish were maintained at 23–25°C in seawater in glass aquariums with charcoal-cotton filtration and were fed weekly. Atlantic sea water (32 34 ppt) was collected from the running seawater system at Marineland, Florida. Fish were anesthetized in tricaine (MS-222; ethyl-m-aminobenzoate; 1:7,500) before use. In preliminary experiments, the rat kidney was used as a control tissue to verify the autoradiography protocols. Male Sprague-Dawley rats were kept at constant room temperature and controlled dark-light periodicity with ad libitum access to water and pellets chow. Rats were killed by a sharp blow to the head and exsanguination.

**Autoradiography**

For autoradiography, tissues were dissected free and freeze mounted in Tissue Tek (Miles; Elkhart, IN) in a microtome cryostat (Minotome, IEC; Needham Heights, MA), and 20–μm sections were cut and mounted on gelatin-chromium aluminum-coated slides and dried overnight under vacuum at 4°C. The sections were used immediately or were stored in sealed boxes at −20°C until used.

Precubation of the tissue sections was performed at room temperature (22–24°C) for 15 min in 50 mM tris(hydroxymethyl)aminomethane (Tris)-HCl buffer (pH 7.4), 50 mM NaCl, 5 mM MgCl₂, 0.1% bovine serum albumin (BSA), and 0.05% bacitracin. This was followed by incubation for 60 min in the same buffer supplemented with 4 μg/ml of leupeptin, 2 μg/ml chymostatin, 2 μg/ml pepstatin, 10–6 M phenylmethylsulfonyl fluoride (PMSF), and 10–5 M ethylenediaminetetraacetic acid (EDTA), or 125I-labeled rat ANP (called rANP-I), or 125I-labeled Tyr(o)-CNP (termed 125I-pCNP here). Displacement of specific binding for 125I-rANP was examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15). Displacement of specific binding for 125I-pCNP was similarly examined in consecutive sections in the presence of 1 μM rANP, 1 μM pCNP, and 1 μM rat C-ANF, a truncated ANP that binds only to NPR-C of mammals (15).

**In Vitro NP Binding Assays**

The gills were removed from anesthetized fish and placed in ice-cold phosphate-buffered saline (pH 7.4). The bony arch was dissected off and the gill filaments were placed in a 50-ml centrifuge tube in 10 volumes of 50 mM Tris and 1 mM NaHCO₃ (pH 7.4) and homogenized with a Tissue-Tearor (Biospec; Bartlesville, OK) at setting 7 for 1 min. The homogenate was diluted 1:1 with 50 mM Tris, 1 mM EDTA, and 1 mM MgCl₂ (pH 7.4) and centrifuged at 800 g for 10 min at 4°C. The supernatant was collected and centrifuged at 30,000 g for 20 min. The pellet was washed with 50 mM Tris (pH 7.4) and 250 mM sucrose and resuspended in the same buffer, and the protein concentration was determined with a BCA protein assay kit, calibrated against BSA standards. Membranes were stored at −70°C until use.

For competition binding assays, 25–50 μg protein of the gill membrane fraction were incubated in the same buffer used for sections and 10–25 pM of 125I-rANP and 20–30 pM of 125I-pCNP. Competition for the iodinated ligands was performed using the unlabeled peptides rANP, pCNP, or C-ANF in increasing concentrations (10–12 to 10–6 M). The incubation volume for the assays was either 0.125 or 0.250 ml. Binding reactions were performed for 2 h at 25°C; preliminary experiments showed that maximal binding was achieved after 60 min. Binding was terminated by dilution with 2 ml of ice-cold 50 mM Tris-HCl (pH 7.4), and bound ligand was separated from free ligand by filtration through 1% polyethyleneimine-treated Whatman GF/C filters. Filters were washed with 5 ml of ice-cold 50 mM Tris-HCl (pH 7.4), and the radioactivity on the filter was determined in a Beckman Gamma counter with 78% efficiency.

**Affinity Cross-Linking and Electrophoresis of Gill Membranes**

Membranes were isolated as previously described and incubated in binding assay buffer with 125I-rANP or 125I-pCNP alone or in combination with either 1 μM rANP, 1 μM pCNP, or 1 μM C-ANF for 2 h. Affinity cross-linking was performed according to Ref. 16. After incubation, the covalent cross-linking agent disuccinimidyl carbonate in dimethyl sulfoxide was added to a final concentration of 1 mM and the incubation continued for 20 min at 4°C. The cross-linking reaction was stopped by addition of an equal volume of quench buffer containing 400 mM EDTA and 1 M Tris-HCl (pH 6.8). Membranes were centrifuged in an Eppendorf centrifuge at 13,000 g for 20 min to separate unbound hormone, and the resulting pellet was suspended for SDS-PAGE in 100 μl of sample buffer containing 62.5 mM Tris base, 2% SDS, 5% glycerol, 0.01% bromophenol blue, 2% β mercaptoethanol, pH 6.8, and then boiled for 3 min. The solubilized membranes were again centrifuged at 13,000 g to remove particulate matter, electrophoresis was performed on a 1.5% unidimensional polyacrylamide slab gel using a mini-gel apparatus, and proteins were stained with Coomassie brilliant blue (Bio-Rad; Richmond, CA). The gels were dried and exposed to Hyperfilm MP for 1–2 wk. Identification of the molecular weights of the labeled bands was done by use of reference standards (30,000–200,000) on the Coomassie blue-stained gel. A linear relationship between the relative mobility (Rf) and the log relative molecular mass of the standards was constructed and the unknown values calculated.

**Guanylate Cyclase Assays**

Gill membranes were isolated and the protein concentration was determined. Membranes were kept on ice until needed. For determination of guanylate cyclase activity, 50 μg of gill protein were added to 50 mM Tris-HCl, 2 mM isobutyl methylxanthine (IBMX), 10 mM creatinine phosphate, 1,000 U/ml creatine phosphokinase, 4 mM MnCl₂, 1 mM GTP, and increasing concentrations of rANP, pCNP, and C-ANF in a final volume of 100 μl. The basal rate of cGMP generation was determined in tubes without ligand. The incubations were performed for 15 min at 24°C and were terminated by the addition of 4 mM EDTA. The tubes were boiled for 3 min and centrifuged at 2,300 g for 15 min. The supernatant was collected and frozen until the cGMP content was determined with a cGMP radioimmunoassay kit.
Data analysis binding data were analyzed using the EBDA and LIGAND computer programs (13). Analysis of variance was performed using Statview SE on a Macintosh computer. Data are expressed as means ± SE.

**Materials**

Rat (3-[125I]iodotyrosine)-atrial natriuretic peptide (1,800–2,000 Ci/mmol), Hyperfilm-β-max, Hyperfilm MP, and the cGMP 3H assay system were purchased from Amersham (Arlington Heights, IL). Porcine 125I-Tyr4-CNP (1,400–1,600 Ci/mmol) and porcine Tyr4-CNP were obtained from Peninsula (Belmont, CA). rANP (3–28), pCNP, and C-ANF (rat des-Gln19, Ser19, Gly20, Leu21, Gly22)ANP-[4–23]-NH2 were obtained from Bachem (Torrance, CA). The BCA protein assay kit and dianacunimydil suberate were purchased from Pierce (Rockford, IL). GTP and IBMX were purchased from Sigma. All other chemicals were reagent grade.

**RESULTS**

**Autoradiographic Studies**

In preliminary experiments we verified the autoradiographic procedure by visualizing 125I-rANP binding sites on glomeruli of the rat kidney as previously reported (10). Specific binding was determined as the difference in binding between sections incubated with radiolabeled ligand alone and sections incubated with radiolabeled ligand and 1 nM unlabeled rANP (data not shown).

Autoradiograms of toadfish gill sections showed specific 125I-rANP binding sites along the efferent (relative to blood flow) and afferent sides of the gill filaments (Figs. 1a, 2a). Specific binding was also observed on the lateral margins of the secondary lamellae in the region of the marginal channels, but not over the entire lamellar body. No specific binding was observed on any gill arch structures, and little, if any, specific binding could be discerned on the epithelium of the filament body. The specific binding was displaced by incubation of sections with 1 µM rANP (Figs. 1b, 2b) and 1 µM pCNP. Compared with the rANP-displaced sections, a small amount of residual specific binding was present in sections incubated with 1 µM C-ANF (Fig. 2c).

The locations of binding sites within the complex gill microvasculature were determined by light-microscopic examination of sections coated with X-ray-sensitive emulsion. Specific binding was determined by comparing the silver grain densities in the presence of 125I-rANP alone (Figs. 3a, 4a) with the grain densities in sections incubated with 125I-rANP and 1 µM rANP (Figs. 3b, 4b). Specific binding sites were concentrated on the efferent filamental artery (Fig. 4a) and were very dense on efferent lamellar arterioles (Fig. 4a). Binding sites were also present on the afferent filamental artery and afferent lamellar arterioles (Fig. 3b). The binding sites on the lamellar arterioles extended along the marginal channel of the secondary lamellae, but little specific binding was present on the body of the lamellae. The binding on the blood vessels was mainly observed on the tunica intima and tunica media. The grain density on the filament body epithelium and the afferent and efferent branchial arteries did not appear to differ between experimental and control sections. In sections incubated with 1 µM C-ANF, binding sites had the same distribution but with reduced density.
The distribution of $^{125}$I-pCNP binding sites in the gills matched that observed for $^{125}$I-rANP. The binding was displaced by 1 nM rANP and 1 μM pCNP. Some residual binding was observed in sections incubated with 1 nM C-ANF. This binding had the same distribution as that in the sections incubated with $^{125}$I-rANP and 1 μM C-ANF.

**Characterization of NP Binding Sites**

The binding sites in the gills were analyzed by competitive radioligand assays, electrophoresis of membrane protein affinity cross-linked to radioligand, and determination of guanylate cyclase activity.

**Competition experiments.** The specificity and kinetics of the $^{125}$I-rANP and $^{125}$I-pCNP binding sites was examined by determining the ability of rANP, pCNP, and C-ANF to compete for binding sites in isolated membrane preparations.

$^{125}$I-rANP BINDING SITE. The concentration of each peptide that displaced 50% of the labeled ligand was 0.1 nM for rANP, 0.2 nM for pCNP, and 4 nM for C-ANF (Fig. 5, top). In the presence of 1 nM C-ANF, 7 ± 1% of specific $^{125}$I-rANP binding remained. Scatchard analysis of unlabeled rANP competing for the $^{125}$I-rANP binding data showed a nonlinear plot (EBDA), and subsequent curve-fitting analysis of the data by LIGAND gave a best
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Binding parameters were 1.28 ± 0.35 nM and 1.332 ± 588 fmol/mg protein, respectively (n = 4).

Affinity cross-linking. Under reducing conditions, electrophoresis of gill membranes cross-linked with ^125^I-rANP showed a major band at 75 kDa and a fainter band at 140 kDa (Fig. 7). Binding at both molecular masses was displaced when the membranes were incubated with ^125^I-rANP and 1 μM rANP, pCNP, and C-ANF (Fig. 7). In membranes incubated with ^125^I-pCNP, only the 75-kDa band was discerned, and this was also displaced with 1 μM rANP, pCNP, and C-ANF.

Stimulation of cGMP production by NPs. rANP and pCNP caused a dose-dependent increase in cGMP production (Fig. 8). Significant increases (P < 0.05) above basal were observed at 10^-7 and 10^-6 rANP and pCNP, respectively. C-ANF had no stimulatory effect (Fig. 8).

Fig. 5. Top: competition study showing the ability of rANP, pCNP, and C-ANF to bind to toadfish gill membranes. Membranes were incubated to equilibrium with 25 pM ^125^I-rANP in the presence of increasing concentrations of cold peptide. Each point is the mean ± SE of 5 experiments. Bottom: representative Scatchard plot from 1 experiment of rANP competing for the ^125^I-rANP binding site. The plot is consistent with a 2-site model.

Fig. 6. Top: competition study showing the ability of rANP, pCNP, and C-ANF to displace ^125^I-pCNP. Membranes were incubated to equilibrium with 35 pM ^125^I-labeled pCNP in the presence of increasing concentrations of cold peptide. Each point is the mean ± SE of 5 experiments. Bottom: representative Scatchard plot from 1 experiment of pCNP competing for the ^125^I pCNP binding site. The plot is consistent with a 2-site model.
DISCUSSION

This study provides evidence for the presence of NP binding sites in the gills of the toadfish Opsanus beta. The autoradiograms of the gill sections showed that both 125I-rANP and 125I-pCNP binding sites were present on the afferent and efferent side of the filament; however, the binding on the filament body and the secondary lamellae did not appear to be above that attributed to nonspecific binding. Light microscopic examination of labeled gill sections dipped in nuclear-track emulsion revealed that the specific NP binding was predominantly localized on the afferent and efferent filamentary arteries and the afferent and efferent lamellar arterioles, with no significant specific binding on the afferent and efferent branchial arteries. Interestingly, extensive binding was also observed along the marginal blood channel of the secondary lamellae, proximal to the lamellar arterioles. The main conclusion that can be drawn from the anatomic studies is that the NP binding sites in the gills were located on vascular tissues rather than ionoregulatory tissues, e.g., chloride cells (see below). Previous studies have shown binding sites on epithelial (chloride?) cells in the gills of the European eel, Anguilla anguilla (2), and two antarctic species, Chionodraco hamatus and Pagothenia bernacchii (35). In addition, ANP stimulates the short-circuit current in the opercular epithelium of Fundulus heteroclitus (25), a response attributed to NPs acting directly on chloride cells. Thus it seems unusual that discernible binding sites on chloride cells were not observed in toadfish gills. The distribution of binding sites in toadfish was also different from that in the gill of the eel A. japonica, in which dense binding was found on the filamental cartilage ray (23).

The analysis of the NP binding sites in the gills shows that at least two populations of NPR are present. The predominant receptor in the gills exhibits many of the characteristics of the mammalian NPR-C, or clearance receptors, based on the following observations: 1) the majority of the 125I-rANP and 125I-pCNP binding is displaced by the truncated ligand C-ANF, which in mammals is a specific ligand of the NPR-C (17); 2) affinity cross-linking studies showed that a large population of receptors have a molecular mass of 75 kDa, which is in the vicinity of the monomeric mass of mammalian NPR-C under reducing conditions; 3) the ability of rANP and pCNP to compete against each radiolabeled ligand with similar affinity is characteristic of the binding of these ligands to NPR-C; and 4) C-ANF did not stimulate the production of cGMP, which is also characteristic of NPR-C, since they lack a guanylate cyclase domain (11). The results of this study, in combination with previous studies in shark (9), trout (5, 18), and Japanese eel (23), clearly show that a receptor characteristic of mammalian NPR-C is present in teleost and elasmobranch fish.

The second type of receptor in toadfish gills is a guanylate cyclase receptor as shown by the ability of NPs to stimulate cGMP production. It is likely that these receptors are the specific binding sites observed on the afferent and efferent arterial vasculature after incubation of sections with C-ANF (Fig. 2). This receptor may be the 140-kDa protein that bound 125I-rANP, since the guanylate cyclase receptors in mammals have similar molecular mass. However, the 140-kDa protein could not be visualized with 125I-pCNP even though pCNP stimulated guanylate cyclase activity. It is difficult to determine if the guanylate cyclase receptor is similar in nature to either the NPR-A or NPR-B of mammals, since both rANP and pCNP stimulated cGMP production in the gills with similar potency. Furthermore, rANP and pCNP competed for both the 125I-rANP and 125I-pCNP binding sites, which is not characteristic of NPR-A (high affinity for ANP) and NPR-B (high affinity for CNP). It is possible that the toadfish gill contains a single guanylate cyclase receptor that does not discriminate between rANP and pCNP. Indeed, isolated vascular
smooth muscle rings of toadfish respond equally to ANP and CNP, further suggesting that distinct NPR-A and NPR-B receptor proteins are not found in the branchial vasculature of this species (20). Molecular studies examining the homology of gill receptor protein genes with NPR-A and NPR-B genes may provide additional information on this point.

The presence of NPR-C in the branchial vasculature of toadfish is consistent with the recent findings of a physiological study in the trout (18). These authors showed that the single-pass extraction of $^{125}I$-ANP by the gills was 60% and that this process was mediated by an NPR-C type of receptor, since it was markedly reduced when C-ANF (SC-46542) was injected into the circulation, which would occupy the NPR-C. Furthermore, the extraction of ANP occurred nearly exclusively in the arterioarterial (respiratory) pathway, which correlates well with the anatomic locations of NP binding sites in the gills of toadfish. The ability of C-ANF to compete for the binding sites in toadfish strongly indicates that the gills of this species have a similar clearance function. Because in fish all blood must pass through the gills, variable expression of clearance receptors in vascular smooth muscle cells of the intrafilamental circulation could be an important mechanism for regulating the levels of plasma NPs by differential clearance of the peptide from the circulation (18). Alternatively, it has recently been suggested that NPR-C may regulate the local concentration of NPs in the vicinity of the guanylate cyclase receptors (14).

The absence of discernible binding sites on the branchial epithelium does not exclude the possibility of the mitochondria-rich chloride cells, or any other epithelial cell type, being a target for NPs. There are several examples in mammals where potent cGMP stimulation has been shown in cells in which binding sites were not detected (12). This situation may apply to cells that do not express the low-molecular-weight NPR-C, which is often the predominant NPR in many tissues. In the present study, it is possible that a small number of binding sites would not be detected above the level attributed to nonspecific binding. However, since two studies have clearly shown NP binding sites on gill epithelial cells, including chloride cells, using techniques similar to those employed in this study, the absence of binding on the epithelium of toadfish is initial evidence that the epithelium is not targeted by circulating NPs. Interestingly, no detectable binding sites can be found in the kidney of O. beta (Donald, unpublished observations), further suggesting that direct regulation of ion-transporting tissues in toadfish is limited.

Perspectives

Although NPs have been shown to mediate a number of osmoregulatory effects in fish, the data are not substantial and are somewhat conflicting (e.g., effects in eel vs. other species; Ref. 7). Current data show that the plasma level of immunoreactive NP is higher in seawater-adapted rather than freshwater-adapted animals, suggesting that the peptide is more important in the regulation of salt load rather than volume load. In contrast, the vasodilatory actions of NPs have been observed in all species studied, suggesting that the NPs are involved in cardiovascular regulation in fish. The results of the present study provide evidence that the arterioarterial branchial vasculature of O. beta is regulated by circulating NPs. The presence of guanylate cyclase receptors on both the afferent and efferent lamellar arterioles and filamentary arteries indicates that NPs could be important in the control of pre- and postlamellar blood flow. Dilation of the branchial vasculature would facilitate changes in intrafilamental hemodynamics that could alter the functional surface area of the gills by increasing or decreasing the number of perfused secondary lamellae (see Refs. 1, 23). Adjustments in the functional surface area affect both osmoregulation and gas exchange, and therefore it is relevant to consider that the physiological effects of NPs could extend to gill functions other than the regulation of salt and water balance and blood flow. For example, if stretching of the atrial wall stimulates the release of NPs into the blood, then the peptide will be released during increases in cardiac output associated with activities such as exercise, since this is facilitated primarily by an increase in stroke volume rather than heart rate (24).

Many studies have shown that the branchial circulation is regulated by both neural and hormonal mechanisms (17). Clearly, NPs are an important new hormone that may interact with other neural and hormonal systems to control blood flow in the branchial vasculature.

This research was supported by National Science Foundation Grant DCR-8916413 to D. H. Evans.

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Received 4 November 1993; accepted in final form 16 April 1994.

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