Estradiol interacts with an opioidergic network to achieve rapid modulation of a vocal pattern generator

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Abstract Estrogens rapidly regulate neuronal activity within seconds-to-minutes, yet it is unclear how estrogens interact with neural circuits to rapidly coordinate behavior. This study examines whether 17-beta-estradiol interacts with an opioidergic network to achieve rapid modulation of a vocal control circuit. Adult plainfin midshipman fish emit vocalizations that mainly differ in duration, and rhythmic activity of a hindbrain–spinal vocal pattern generator (VPG) directly establishes the temporal features of midshipman vocalizations. VPG activity is therefore predictive of natural calls, and ‘fictive calls’ can be elicited by electrical microstimulation of the VPG. Prior studies show that intramuscular estradiol injection rapidly (within 5 min) increases fictive call duration in midshipman. Here, we delivered opioid antagonists near the VPG prior to estradiol injection. Rapid estradiol actions on fictive calling were completely suppressed by the broad-spectrum opioid antagonist naloxone and the mu-opioid antagonist beta-funaltrexamine, but were unaffected by the kappa-opioid antagonist nor-binaltorphimine. Unexpectedly, prior to estradiol administration, all three opioid antagonists caused immediate, transient reductions in fictive call duration. Together, our results indicate that: (1) vocal activity is modulated by opioidergic networks, confirming hypotheses from birds and mammals, and (2) the rapid actions of estradiol on vocal patterning depend on interactions with a mu-opioid modulatory network.

Keywords Nongenomic · Extranuclear · Membrane · Teleost · Opioid

Introduction

Estrogens rapidly regulate behavior within minutes in a variety of contexts, most notably the sexual behavior of male rats (Cross and Roselli 1999) and Japanese quail (Cornil et al. 2006b). Estrogens also act within seconds-to-minutes on the neurophysiological properties of striatal, hippocampal and hypothalamic neurons (Qiu et al. 2003; Woolley 2007). These actions on neuronal excitability occur via interactions with ion channels and/or neurotransmitter receptors, including those for opioids (reviews: Kelly and Ronnekleiv 2002; Cornil et al. 2006a). Taken together, these observations suggest that the mechanisms established for estrogen action on neuronal activity mediate the rapid behavioral effects of estrogens. However, it remains largely unknown how estrogens rapidly modulate discrete networks of neurons that pattern behavioral output. In principle, estrogen-dependent modulation could be via either direct effects on central pattern generators or indirect effects on afferent modulatory networks.

17-β-Estradiol (E₂) modulates the activity of a vocal central pattern generator in a sound-producing teleost fish, the plainfin midshipman (Porichthys notatus) (Remage-Healey and Bass 2004, 2007). Male midshipman establish and defend nests they build under rocky shelters, and emit long-duration (>1 min) advertisement calls known as “hums” to attract females to their nests. Adults of both

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W. K. L. and C. A. M. have been identified in many species as playing a role in the regulation of sexual behavior. An interesting finding is that both androgens and glucocorticoids can cause rapid changes in behavior, with the rapid effects of androgens being blocked by cyproterone acetate (antiandrogen) and RU486 (antiglucocorticoid), respectively (Remage-Healey and Bass 2004). A similar candidate receptor mechanism has been proposed to mediate rapid corticosteroid suppression of sexual behavior in mammals (Remage-Healey and Bass 2004, 2007), and these effects are both short-lived (lasting 30–45 min) and stereospecific. Androgens and glucocorticoids also cause rapid changes in the duration of active calls in adult male and female midshipman (Remage-Healey and Bass 2004, 2007), and these effects are both short-lived (lasting 30–45 min) and stereospecific. Androgens and glucocorticoids also cause rapid changes in the duration of active calls in adult male and female midshipman (Remage-Healey and Bass 2004, 2007), and these effects are both short-lived (lasting 30–45 min) and stereospecific. Androgens and glucocorticoids also cause rapid changes in the duration of active calls in adult male and female midshipman (Remage-Healey and Bass 2004, 2007), and these effects are both short-lived (lasting 30–45 min) and stereospecific.

The midshipman model system therefore presents an opportunity to identify mechanism(s) of acute estrogen action on a defined neural circuit (shared by all major vertebrate lineages, see Bass et al. 2008) that patterns vocalizations.

Here, we test the hypothesis that an opioidergic network provides a modulatory mechanism for the rapid actions of E_2 in midshipman. Our focus on opioidergic mechanisms is related to three previous observations. First, rapid E_2 actions on mammalian hypothalamic neurons occur via interactions with opioidergic systems (Kelly et al. 1992; Qiu et al. 2003). Second, a membrane-bound receptor that is proposed to mediate rapid corticosteroid suppression of clamping behavior in amphibians exhibits opioid-like binding properties (Evans et al. 2000). Lastly, longer-term (i.e., hours to days) interactions between steroids (estrogens and progesterins) and opioidergic neurons have emerged in studies examining the regulation of sexual receptivity in rodents (Eckersell et al. 1998; Sinchak and Micevych 2001; Acosta-Martinez and Etgen 2002). We report results for three opioid antagonists: naloxone, a broad-spectrum opioid blocker; beta-funaltrexamine, a blocker specific to mu-opioid neurons in other vertebrates; and nor-binaltorphimine, a blocker specific to kappa-opioid neurons in other vertebrates (Bodnar and Klein 2006). We note that the receptor specificity of these antagonists has not been fully characterized for opioid receptors in teleosts, but that opioid receptor subtypes are broadly conserved among vertebrates (Brooks et al. 1994; Ebbsesson et al. 1996; Khalap et al. 2005, see discussion).

Materials and methods

Animals

Midshipman fish have two adult male morphs that diverge in a large suite of traits including reproductive tactics and vocal behaviors (Brantley and Bass 1994). Type I males build and guard nests in the intertidal zone, whereas smaller type II males sneak and satellite spawn in attempt to steal fertilizations from type I’s. Only type I males are known to produce the long-duration advertisement hums as well as extended trains of brief agonistic grunts and growls that are intermediate in duration between hums and grunts (Bass and McKibben 2003). Type II males and females are only known to generate grunts in non-spawning contexts. This study included 40 type I males (13–20 cm standard length) collected from nest sites in northern California and Washington state. While females have the highest plasma E_2 levels, detectable levels of circulating E_2 are found in close to 70% of type I males during pre-nesting and nesting periods (Sisneros et al. 2004). Also, aromatase-containing glial cells surround the dorsolateral aspect of the vocal motor nucleus (VMN) and have processes that ramify throughout the VMN which likely provide a local source of E_2 via aromatization of circulating testosterone (Forlano et al. 2001; Sisneros et al. 2004).

Evoked activity of the vocal pattern generator (VPG)

Electrophysiology procedures closely followed those outlined in earlier studies with midshipman fish (Bass and Baker 1990; Goodson and Bass 2000; Remage-Healey and Bass 2004). The brain and rostral spinal cord were exposed by dorsal craniotomy under general anesthesia (0.025% benzocaine; Sigma) and long-lasting analgesia (0.25% bupivacaine; Abbott Laboratories, Chicago, IL; with 0.01 mg/mg epinephrine; International Medication Systems, El Monte, CA). Following surgery, fish were stabilized in a plexiglass tank during recovery and perfused through the mouth with fresh saltwater maintained at 17°C with a Peltier device. Exposed brain areas were covered with FluoroInert (3 M Corp., St. Paul, MN), and intramuscular injections of pancuronium bromide (0.5 mg/kg; Astra Pharmaceutical Products, Westborough MA) were used for immobilization. Brief electrical stimuli (pulse width = 0.1 ms, frequency = 250 Hz, duration = 25 ms) were delivered via an insulated tungsten electrode exposed at the tip (Fig. 1b) to a hindbrain–spinal vocal site VPP (Bass et al. 2008), formerly
designated as VM (Bass et al. 1994), that is vocally active (Remage-Healey and Bass 2004; 2006). Electrically evoked vocal motor volleys were recorded with an extracellular electrode (Teflon-coated silver wire with exposed ball tip) placed on a ventral occipital nerve root that carries vocal motor axons to the ipsilateral vocal muscle attached to the lateral walls of the swim bladder (Fig. 1b). Activity recorded from either left- or right-side occipital nerve root reflects bilateral output (Bass and Baker 1990). Rhythmic vocal motor volleys occurred as discrete bursts (fictive calls, Fig. 1a–c) similar to natural grunts (see Brantley and Bass 1994) and were recorded at 1-s intervals. Recordings were digitized on a Macintosh using IGOR Pro software (WaveMetrics, Inc., Lake Oswego, OR). The duration of each fictive call was measured from the recordings and the mean burst duration was computed for each time point and normalized to 100% of baseline output.

Delivery of opioid antagonists

All opioid antagonists were obtained from Tocris Bioscience (Ellisville, MO) and dissolved in buffered teleost
Ringer’s (Forster–Taggart) solution (all solutions had pH between 6.81 and 8.19). Fictive calls evoked from the hindbrain–spinal VPP region were considered stable when a single fictive call was entrained to each stimulus. Once stability was established, a series of ten fictive calls was evoked at 1-s intervals to determine baseline (pre-drug) vocal motor output. Immediately following the baseline period, either saline vehicle (n = 4 animals) or one of three opioid receptor antagonists was pressure injected (see below) into the hindbrain (fourth) ventricle that directly borders the VMN (see Fig. 1b) (Bass and Baker 1990; Bass et al. 1994). Antagonists included the broad-spectrum opioid receptor antagonist naloxone (1, 10, 25 mM; n = 6, 4, 4 animals each, respectively), the mu-specific opioid antagonist beta-funaltrexamine (1, 10 mM; n = 4 each), and the kappa-specific opioid antagonist nor-binaltorphimine (0.66, 6.6 mM; n = 3 each). Drug doses were based on previous studies of opioid–steroid interactions (Evans et al. 2000; Acosta-Martinez and Etgen 2002; Ceccarelli et al. 2004; Schmitt and Kaufman 2005). The standard dose sequence (1, 10 mM) was lowered slightly for nor-binaltorphimine due to limited reagent availability; it is an effective kappa-opioid antagonist in both mammalian and non-mammalian vertebrates (Bradford et al. 2005).

Drugs and vehicle were delivered via a glass micro-pipette (microfilament 6010 1.0 mm OD, AM Systems, Carlsborg, WA) with broken tips (18–20 μm) using a picospritzer (Biomedical Engineering, Thornwood, NY). Only one dose of opioid antagonist or vehicle was used per experiment (i.e., in each animal). Injection parameters followed those well-established for this preparation (one pressure pulse per second at 30 psi and pulse duration of 10 ms for estimated infusion volume of approximately 0.16 nl; see Goodson and Bass 2000; Kittelberger et al. 2006). The picospritzer electrode was checked before and immediately following each experiment to verify that pressure delivery was functional. A simultaneous recording of ten fictive calls was obtained (as above) concurrent with pressure delivery of solutions to identify any instantaneous effects of either drug or saline delivery on vocal motor patterning (see also the experiment timeline in Fig. 1a). Two subsequent recordings of 15 fictive calls were obtained at 5 and 15 min following solution delivery (all treatments prior to E2 administration; see Fig. 1a) to determine whether drugs or saline altered vocal motor patterning. Five minutes after the 15-min post-drug recording, fish were administered an intramuscular injection (dorsal trunk muscle) of E2 (0.02 mg/kg) via a pre-inserted 23-G butterfly needle. This treatment dose rapidly increases fictive call duration (within 5 min and lasting up to 30 min) in type I male midshipman (Remage-Healey and Bass 2004, 2007). Immediately following the injection, a subsequent series of 15 fictive calls were obtained at 5, 15, 30, 45, and 60 min, and each time point was standardized to 100% of the baseline output. After the last recording, blood samples were obtained from the gill arch (Remage-Healey and Bass 2004, 2005; Remage-Healey and Bass 2006), and plasma was analyzed for E2 using enzyme immunoassay (Cayman Chemical, Ann Arbor, MI, techniques reported previously in Remage-Healey and Bass 2004, 2005, 2007). Plasma E2 levels were confirmed to be elevated (mean = 34.52 ng/ml ± 8.01 SEM; n = 16) for a subset of animals that received E2 injections (see also Remage-Healey and Bass 2004).

Analysis

As described above, fictive call durations were sampled and standardized to 100% of baseline. Statistical analyses of the effects of drugs on fictive call duration were performed using Statview 4.57. Repeated-measures ANOVA was performed for data from all treatments for analysis of treatment effects over time in a within- and between-subjects design. Tukey’s post hoc tests are reported for between-subject effects for comparisons of saline versus drug treatments for P < 0.05.

Results

Overview

We first show that each of the opioid antagonists used here—naloxone, beta-funaltrexamine, and nor-binaltorphimine—caused a rapid and brief (either immediate or within 5 min of infusion) suppression of fictive calling when delivered to the hindbrain VPP region. All antagonists had similarly brief effects when initially applied, but calling recovered to baseline quickly (within 5–10 min). In contrast, subsequent attenuation of rapid E2 effects on fictive calling was both antagonist-specific and prolonged, lasting up to 30–60 min. This encompasses the duration of E2’s actions on fictive calling in this species (Remage-Healey and Bass 2004, 2007). Only naloxone and the mu-opioid-specific antagonist beta-funaltrexamine, but not the kappa-opioid antagonist nor-binaltorphimine, inhibited E2’s effects on calling.

Antagonists alone

Naloxone

The broad-spectrum opioid antagonist naloxone (1, 10, and 25 mM) caused a transient reduction in fictive call duration, in a dose-dependent manner (Fig. 2a). There were significant effects of time-after-delivery (F = 8.74; df = 3,42; P = 0.0001), a dose × time interaction (F = 3.36; df = 9.42; P = 0.003), and a non-significant trend for the effect of dose...
The broad-spectrum opioid antagonist naloxone (a) the mu-specific-opioid antagonist beta-funaltrexamine (b) and the kappa-specific-opioid antagonist nor-binaltorphimine (c) each transiently suppress fictive call duration. All treatment effects recover to baseline at 15 min. *P<0.05 for Tukey’s post hoc test for between-subject comparisons of fictive calling during infusion of antagonists versus vehicle (saline).

(F = 3.12; df = 3.14; P = 0.06). Tukey’s post hoc test revealed that naloxone produced significant (P<0.05) reductions in fictive call duration at 5 min after delivery for all three doses (1, 10, 25 mM) when compared to saline alone. Fictive call duration was indistinguishable from baseline at 15 min following naloxone treatment. For the 5-min time point, the response to naloxone was linearly dose-dependent; i.e., there was a larger reduction in fictive call duration as the doses of naloxone increased.

**Beta-funaltrexamine**

The mu-specific opioid antagonist beta-funaltrexamine (1, 10 mM) also caused a transient reduction in fictive call duration (Fig. 2b). There were significant effects of time-after-delivery (F = 11.43; df = 3.21; P = 0.001; a dose × time interaction (F = 11.91; df = 6.21; P < 0.001), and a significant effect of dose (F = 29.7; df = 2.7; P = 0.0004). Tukey’s post hoc test revealed that beta-funaltrexamine produced significant (P<0.05) reductions in fictive call duration immediately upon delivery (‘spritz’; 10 mM dose) and at 5 min after delivery (1 mM dose) when compared to saline alone. While the time-dependence of the response to beta-funaltrexamine differed slightly for the two doses, they were similar in temporal pattern. That is, the larger dose (10 mM) produced only an immediate reduction in fictive calling and the lower dose (1 mM) produced a reduction only at 5 min after delivery of the drug; each recovered to baseline by the next time point tested, i.e., within at least 5 min (10 mM) or 10 min (1 mM). Fictive call duration was indistinguishable from baseline at 15 min following beta-funaltrexamine treatment.

**Nor-binaltorphimine**

The kappa-specific opioid antagonist nor-binaltorphimine (0.66, 6.6 mM) also caused a transient reduction in fictive call duration that was, like naloxone, dose-dependent (Fig. 2c). There were significant effects of time-after-delivery (F = 11.43; df = 3.21; P = 0.001), a dose × time interaction (F = 11.91; df = 6.21; P < 0.001), and a significant effect of dose (F = 29.7; df = 2.7; P = 0.0004). Tukey’s post hoc test revealed that nor-binaltorphimine produced significant (P<0.05) reductions in fictive call duration immediately upon delivery (0 min; 6.6 mM dose) and at 5 min after delivery (0.66 and 6.6 mM doses) when compared to saline alone. Fictive call duration was indistinguishable from baseline at 15 min following nor-binaltorphimine treatment. Similar to beta-funaltrexamine, the dose-dependence of the response to nor-binaltorphimine was apparent in that only the highest dose (6.6 mM) produced an immediate reduction in fictive calling. Unlike beta-funaltrexamine, both low and high doses produced reductions in fictive calling at the 5-min time point. Similar to both naloxone and beta-funaltrexamine, fictive call suppression that was present at 5 min recovered to baseline at the next time point tested, i.e., 15 min after picospritzing the drug.

**Antagonists together with E2**

**Naloxone**

The broad-spectrum opioid antagonist naloxone (1, 10, and 25 mM) caused suppression of E2’s rapid effects on fictive call duration that was dose-dependent (Fig. 3a). There were significant effects of time-after-injection (F = 2.43; df = 5.60; P = 0.04), a dose × time interaction (F = 2.23; df = 15.60; P = 0.01), and a significant effect of naloxone dose (F = 5.33; df = 3.12; P = 0.01) when comparing naloxone + E2 treatments versus saline + E2 treatment. Tukey’s post hoc test revealed that naloxone + E2 treatments produced significantly shorter fictive call duration from saline + E2 treatment at 5 min (10 and 25 mM),
15 min (25 mM), and 30 min (10 and 25 mM). Therefore, only the intermediate (10 mM) and high (25 mM) doses of naloxone were effective in suppressing the rapid actions of E2.

**Beta-funaltrexamine**

The mu-specific opioid antagonist beta-funaltrexamine (1, 10 mM) also suppressed E2’s rapid effects on fictive call duration (Fig. 3b). There was no significant effect of time-after-injection ($F = 0.68$; $df = 5.45$; $P = 0.64$), but there was a significant dose × time interaction ($F = 3.44$; $df = 10.45$; $P = 0.002$) and a significant effect of beta-funaltrexamine dose ($F = 4.26$; $df = 2.9$; $P = 0.05$) when comparing beta-funaltrexamine + E2 treatments versus saline + E2 treatment. Like naloxone, Tukey’s post hoc test revealed that beta-funaltrexamine + E2 treatments produced significantly shorter fictive call duration from saline + E2 treatment at 5 min (1 and 10 mM), 15 min (1 mM), and 30 min (1 mM). Thus, unlike naloxone, only the low (1 mM) dose of beta-funaltrexamine was effective in suppressing the entire time-course of rapid E2 actions, while the response to the high (10 mM) dose was no longer significantly different from the saline + E2 treatment after 15 min.

**Nor-binaltorphimine**

Unlike naloxone and beta-funaltrexamine, the kappa-specific antagonist nor-binaltorphimine (0.66, 6.6 mM) had no significant effect on the rapid increase in fictive call duration induced by E2 (Fig. 3c). There was a significant effect of time-after-injection (response to E2; $F = 10.70$; $df = 5.35$; $P < 0.0001$), but there was no time × dose interaction ($F = 0.57$; $df = 10.35$; $P = 0.83$), and no significant effect of binaltorphimine dose ($F = 0.42$; $df = 2.7$; $P = 0.67$) when comparing nor-binaltorphimine + E2 treatments versus saline + E2 treatment. Tukey’s post hoc test did not detect significant differences between nor-binaltorphimine + E2 and saline + E2 treatment for any time point at any dose. It is possible that a higher dose of nor-binaltorphimine would have caused suppression of fictive calling. However, the current dose regimen matched the order of magnitude for the other opioid antagonists used in this study (micro- to milli-molar range) and is within the high-range of doses for effective kappa-opioid blockade as reported in the literature (Gogas et al. 1996; Mitchell et al. 2005; Bodnar and Klein 2006; Pascoe et al. 2008).

**Discussion**

This study reports that intracerebroventricular microinjection of opioid antagonists can acutely inhibit fictive calling in midshipman fish. We also report that the rapid neuromodulation of fictive calling by E2 is suppressed during specific inhibition of a central mu-opioidergic network. Our findings are therefore consistent with the hypothesis that rapid E2 actions on hindbrain–spinal VPG activity occur via interactions with a central mu-opioidergic network whose activation, in turn, modulates the VPG (as summarized in Fig. 1).

**Opioidergic network and vocal mechanisms**

This study provides evidence that opioids are neuromodulatory in contexts outside of their well-established role in regulating nociception and reinforcement/reward via actions in the forebrain (Bodnar and Klein 2006). The current observation that opioidergic neurons are involved in modulating vocal motor mechanisms may not be unique to midshipman fish.
fish, since opioids and their receptors are expressed throughout the song-control pathway in oscine (Deviche et al. 1993) and non-oscine (Durand et al. 1998) songbirds. Furthermore, opioids regulate vocal behaviors in both songbirds and mammals (Riters et al. 2005; Bodnar and Klein 2006), and, consistent with these earlier observations, the current study provides neurophysiological evidence that opioidergic neurons participate in the patterning of vocalizations.

Both the mu-specific (beta-funaltrexamine) and the kappa-specific (nor-binaltorphimine) opioid antagonists caused immediate suppression of fictive call duration during infusion (‘spritz’; latency < 1 min), whereas the broad-spectrum opioid antagonist naloxone caused vocalization suppression after a latency of 5 min. Therefore, suppression was common to all three antagonists as was a rapid recovery to baseline. Relative to the other antagonists, the naloxone effect was delayed but similarly short-lived, which may relate to differences in the ability of each drug to penetrate neural tissue due to chemical structure or potency (see also Martin et al. 2006; Chatjale et al. 2008). It is noteworthy that the binding affinity of opioid antagonists to mu- and kappa-opioid receptors have been well-characterized for mammals, but not yet for teleost fishes. A lack of strict dose-dependence for these opioid antagonist treatments on fictive calling makes non-specific effects of microinjection (such as concentration-dependent changes in pH or other interfering substances) unlikely. However, we cannot formally rule out the possibility of immediate and non-specific artifacts of microinjection on vocal motor network properties given that all of the antagonists, in general, suppress fictive call duration.

Opioid peptide and receptor expression in the midshipman central nervous system is currently un-described. Opioids and their receptors are highly conserved among vertebrates (Khalap et al. 2005), and neuroanatomical evidence for opioid networks exist for other teleost fishes (Brooks et al. 1994; Ebbesson et al. 1996). In coho salmon, mu-opioid binding sites are reported throughout the telencephalon, diencephalon and mesencephalon (Ebbesson et al. 1996), but the distribution of mu-opioid binding sites in the hindbrain are currently unknown in any teleost (delta-opioid receptors are found in the hindbrain of zebrafish, specifically in the periventricular zone and abutting the nucleus of the medial longitudinal fasciculus, Porteros et al. 1999). The indication that a mu-opioidergic network is critical for rapid actions of estrogens in midshipman fish is intriguing in light of the observation that brain mu-opioid systems are the most highly conserved among vertebrates (Li et al. 1996; Darlison et al. 1997).

Opioidergic neurons that modulate midshipman vocal patterning could have multiple neuroanatomical origins. One of the major descending opioid projections in vertebrates is the midbrain periaqueductal gray (PAG, Behbehani 1995), a critical node linking forebrain to hindbrain vocal neurons in midshipman fish (see Kittelberger et al. 2006, PAG is part of the mVAC in Fig 1 top) as well as in mammals (Jurgens 2008). The current findings are consistent with the hypothesis that descending opioidergic neurons from the PAG have terminals in the hindbrain–spinal VPG, and that their activity contributes to the rapid modulation of vocalizations in midshipman. This hypothesis predicts that a subset of the projections from the PAG to the VPG consist of opioidergic processes that are rapid modulated by estrogens (directly or indirectly, see below).

Opioids and rapid estrogen mechanisms

We report that mu-opioidergic antagonists block the rapid actions of $E_2$ during a 15–60 min period after microinjection into the hindbrain ventricle adjacent to the VPG. This indicates that opioid blockers exert long-lasting effects on the VPG comprised of mechanisms that could be substantially different from the immediate (0–5 min) inhibitory effects of opioid blockers on fictive calls. Opioid agonists and antagonists are known to have delayed (minutes-to-hours) modulatory actions on network function (Brown et al. 1998), which consist of mechanisms distinct from rapid opioidergic (seconds-to-minutes) actions on neural activity (Heinricher 2003; Nugent et al. 2007). The current results suggest that similarly divergent acute versus delayed opioid mechanisms can each exert modulatory influences on the VPG circuit in midshipman.

One mechanism to account for the current findings is that opioid antagonists broadly suppress neuronal activity, including activity within the VPG, and that this suppression precludes the rapid actions of $E_2$ on the VPG. Consistent with this hypothesis, opioid agonists and antagonists exert long-lasting suppression of activity within neural circuits in a variety of contexts (Bennett and Mayer 1979; Cheng et al. 1986; Charpak et al. 1988). However, in the current experiments, the midshipman VPG recovered from suppression (i.e., returned to 100% of baseline fictive calling) within 5–10 min following treatment with all three opioid receptor antagonists. Secondly, mu-opioid blockade suppressed baseline VPG activity (Fig. 2b) and subsequent rapid $E_2$ actions (Fig. 3b), yet kappa-opioid blockade only rapidly suppressed VPG activity (Fig. 2c), but did not subsequently influence $E_2$’s rapid actions (Fig. 3c). These findings together indicate that there are specific modulatory interactions between $E_2$ and a candidate mu-opioidergic network in the midshipman VPG region.

The candidate cellular mechanism(s) for combined $E_2$/mu-opioid regulation of the VPG include either $E_2$ binding directly to opioid receptors (Gutierrez et al. 1998), or to nonclassical binding sites on opioidergic neurons (Lagrange
These nonclassical actions of E2 could occur either directly on modulatory mu-opioidergic neurons, or indirectly via an as yet unidentified modulatory input afferent to mu-opioidergic neurons. The acute nature of our findings indicate that the site of rapid E2 action within the VPG region is a nonclassical estrogen-binding site. Although the selective estrogen-receptor modulator ICI 182,780 potently blocks long-term estrogen-opioid-receptor mechanisms in other teleosts (e.g., Muenet et al. 2005; Pinto et al. 2006a, 2006b), pilot studies indicate that ICI is ineffective at modulating E2’s rapid actions on vocal patterning in midshipman fish (Remage-Healey and Bass, unpublished observations). The binding site which mediates rapid E2 effects in midshipman may therefore be structurally dissimilar to the nuclear estrogen receptor (i.e., a membrane receptor/ion channel); alternatively, the rapid actions of E2 could be mediated by nonclassical mechanisms associated with the canonical cytosolic estrogen receptor(s) (e.g., Vasudevan et al. 2005). Previous evidence from rats indicates that E2 increases the availability of mu-opioid receptors in hypothalamic inhibitory neurons (Torres-Reveron et al. 2009), suggesting that the rapid effects of E2 observed here could be via similarly nonclassical interactions with hindbrain/spinal GABAergic neurons that are mu-opioid responsive. Although more neuroanatomical evidence is needed for teleosts, E2 can modulate opioidergic function in the brainstem of rats (Tashiro et al. 2008) and opioid receptor mRNA is co-localized with estrogen-receptor positive cells in brainstem circuits (Flores et al. 2003).

The rapid E2/opioidergic modulation reported here is a part of a growing number of reports indicating that steroids interact with opioidergic systems to achieve rapid actions on neuronal activity. Steroids and steroidal drugs interact directly with opioid receptors, including mu-opioid receptors, to alter binding affinities for endogenous opioids and subsequent neuronal activity (Maggi et al. 1996; SanMartin et al. 1999; Sajadi et al. 2007; Brown et al. 2008). A membrane glucocorticoid receptor recently isolated in newts exhibits kappa-opioid binding properties, and has been proposed to mediate the rapid actions of corticosterone on reproductive clasing behavior (Evans et al. 2000). In guinea pig hypothalamus, E2 rapidly inhibits the mu-opioid effector system (Kelly et al. 1992), although not via a direct interaction with opioid receptors per se (Qiu et al. 2003). In contrast, in rat hypothalamus, E2 causes mu-opioid receptor trafficking inward from the membrane (internalization) within 30 min, and this effect can be blocked by treatment with the broad-spectrum opioid blocker naltrexone (Eckersell et al. 1998; Sinchak and Micevych 2001). Micevych and colleagues have proposed that E2 causes rapid release of opioid peptides, which then bind to mu-opioid receptors, leading to their internalization. The results of the current study are consistent with this model from rat hypothalamus, and support the hypothesis that the rapid effects of E2 cause the release of mu-opioids within the VPG and thereby increase fictive call duration. This model for the midshipman VPG generates two specific predictions: (1) opioidergic modulatory neurons express estrogen-binding sites, most likely presynaptically within the VPG itself, and (2) estrogen treatment will lead to rapid increases in local mu-opioid concentrations accompanied by rapid internalization of mu-opioid receptors in the VPG region of midshipman.

This study demonstrates that steroids and neuropeptides can interact in the minute-by-minute control of motor network patterning, suggesting that acute shifts in midshipman calling behavior in nature involve combined rapid actions of estrogens and opioids (Fig. 1). Our findings are consistent with a mechanism of second-order, or “meta”-modulation, in which the actions of one neuromodulator (in this case E2) influence the actions of an existing neuromodulatory circuit (in this case mu-opioid neurons). Metamodulation provides an important mechanism for generating a broad range of behavioral states from a discrete network of neurons (Katz and Edwards 1999; Mesce 2002; McLean and Sillar 2004). Metamodulation is likely an important mechanism for shaping vocal output in midshipman, since the neuropeptides arginine vasotocin and isotocin also exert rapid actions on fictive calling in this species (Goodson and Bass 2000).

The current results suggest that the steroid-sensitive neural circuitry that patterns vocal production in other vertebrates (Schlinger and Brenowitz 2002; Bass and Remage-Healey 2008; Jurgens 2008), could be regulated by acute steroid/opioid interactions similar to those in this study. Thus far, evidence has been suggestive in favor of this hypothesis. Masculinization of “crown” vocalizations in quail involves a testosterone-dependent upregulation of opioid binding sites in the brainstem (Bernroider et al. 1996). In addition, peripheral injections of opioid receptor antagonists increase male song production in starlings (Riters et al. 2005). However, to our knowledge neurophysiological evidence that opioids and steroids interact to regulate vocal patterning does not exist for birds and mammals. The mammalian hypothalamus exhibits sexually dimorphic expression of opioid peptides, and some neurons co-express mRNA for steroid receptors and opioids (Gu and Simerly 1994; Roepke et al. 2007). Therefore, rapid interactions between opioid- and steroid-dependent mechanisms, such as those outlined in this study, could be especially important for sex differences in brain function and behavioral plasticity in vertebrates.

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