Abstract

The pulsatile release of gonadotropin-releasing hormone (GnRH) is a key feature of the hypothalamic–pituitary–gonadal axis. Kisspeptin neurons in the arcuate nucleus (ARC) trigger GnRH neuronal activity, but how GnRH neurons return to baseline electrical activity is unknown. Nociceptin/orphanin-FQ (OFQ) is an inhibitory neuromodulator. ARC proopiomelanocortin (POMC) neurons, known to receive inputs from ARC kisspeptin neurons, contact GnRH neurons and coexpress OFQ in the rat. In the present study, the effect of OFQ(1-13) on GnRH neurons was determined in the mouse. We identified transcripts for the OFQ receptor [opioid receptor like 1 (ORL1)] in GnRH neurons, and, using two-model systems (explants and slices), we found that OFQ exerted a potent inhibition on GnRH neurons, with or without excitatory inputs. We confirmed that the inhibition was mediated by ORL1 via G_i/o-protein coupling. The inhibition, occurring independently of levels of intracellular cyclic adenosine monophosphate, was sensitive to inwardly rectifying potassium channels. The only specific blocker of G_i/o-protein-coupled inwardly rectifying potassium (GIRK) channels, tertiapin-Q (TPNQ), was ineffective in the inhibition of OFQ. Two GIRK activators, one sharing the binding site of TPNQ and one active only on GIRK1-containing GIRK channels, failed to trigger an inhibition. In contrast, protein kinase C phosphorylation activation, known to inhibit GIRK2-mediated currents, prevented the OFQ inhibition. These results indicate a specific combination of GIRK subunits, GIRK2/3 in GnRH neurons.

Significance Statement

Fertility is controlled centrally by neurons secreting gonadotropin-releasing hormone (GnRH) and critically relies on their pulsatile secretory profile. GnRH pulses depend on kisspeptin neurons located in the arcuate nucleus. However, kisspeptin provides a long-lasting stimulation, and how GnRH neurons return to baseline electrical activity is unknown. Here, we show nociceptin/orphanin-FQ potently inhibits GnRH neurons. The signaling pathway involves the receptor, opioid receptor like 1, and downstream effectors G_i/o-proteins and G_i/o-protein-coupled inwardly rectifying potassium (GIRK) channels. Notably, the GIRK channels in GnRH neurons exhibit a specific subunit composition, GIRK2/3. Together, these data identify a new messenger in modulating reproductive function.
Introduction

Fertility relies on the capacity of gonadotropin-releasing hormone (GnRH) neurons to convert a wide range of cues from the CNS into a final hormonal signal to the pituitary, which subsequently controls the gonads. Steroids from the gonads feedback to the CNS providing the physiologic signals for coordinated communication along the GnRH–pituitary–gonadal axis and thus, reproductive success. Notably, GnRH is secreted in a pulsatile fashion (i.e., it must be initiated and stopped). Kisspeptin neurons are key players in the model for pulsatile release of GnRH (Mittelman-Smith et al., 2012; Navarro, 2012; Goodman et al., 2013). In this model, the action of stimulatory neuropeptide B and inhibitory dynorphin A drives kisspeptin neurons into an oscillating firing mode that leads to pulsatile kisspeptin release. The phasic activity of GnRH neurons is subsequently triggered by kisspeptin. While dynorphin A ends the kisspeptin release and therefore removes the excitation of GnRH neurons, exogenous kisspeptin triggers a long-lasting excitation in GnRH neurons, both at the cell body and at the terminal (Han et al., 2005; Constantin et al., 2009, 2013; Iremonger et al., 2017). The endogenous kisspeptin released after stimulation of the anteroventral periventricular nucleus (AVPV) also evokes the classical long-lasting effect (Liu et al., 2011). GnRH secretion seems to end while the kisspeptin induced excitation still persists (Glanowska and Moenter, 2015). To date, the identity of the molecule that terminates the bout of electrical activity and allows GnRH neurons to return to baseline electrical activity is unknown.

High levels of the nociceptin/orphanin FQ (OFQ) receptor ORL1 (opioid receptor like 1) are found in the preoptic area (POA) and the anterior hypothalamic area where GnRH neurons reside (Houtani et al., 2000). In vitro, OFQ directly hyperpolarizes GnRH neurons in the arcuate nucleus (ARC; guinea pigs; Wagner et al., 1998), inhibits forskolin-evoked GnRH release from hypothalamic fragments (male rats; Dhandapani and Brann, 2002) and the spontaneous release of GnRH from mediobasal hypothalamus fragments (variecotemized (OVX) rats; An et al., 2005). In vivo, OFQ push–pull infusion inhibits GnRH release from the POA (OVX rats; An et al., 2005) and intracerebroventricular OFQ inhibits GnRH release from the median eminence (OVX rats; An et al., 2005, 2009). OFQ intracerebroventricular injections also decrease circulating luteinizing hormone (LH) levels (OVX rats; An et al., 2005) and blunts the preovulatory surge (OVX+estradiol/progesterone rats; An et al., 2007). Pharmacology or RT-PCR analysis support the effects being ORL1 dependent. Together, these data suggest that ORL1 is expressed on GnRH neurons, and OFQ may act as an inhibitory neuro-modulator (Mollereau and Moulelous, 2000).

The ARC contains a variety of neuropeptides (Chronwall, 1985) including OFQ (Maoloold and Meister, 2010; Campbell et al., 2017). Proopiromelanocortin (POMC) neurons are one of the ARC neuronal subpopulations that coexpress OFQ [male rats (Maoloold and Meister, 2010); ewes (Nestor et al., 2013)]. POMC neurons express both ERα (Jirikowski et al., 1986; Xu et al., 2011) and progesterone receptor (Fox et al., 1990). ARC-specific (Yeo and Herbison, 2014) and POMC-specific (Xu et al., 2011) deletion of ERα disrupted LH responses to estrogen-negative feedback in female mice. Afferents from the ARC (Mezey et al., 1985; Wintemantel et al., 2006), and specifically from POMC neurons (Leranth et al., 1988; Simonian et al., 1999), are present in the vicinity of GnRH cell bodies in the preoptic area, and the response of GnRH neurons to the POMC-derived peptides, α-melanocyte-stimulating hormone and β-endorphin, clearly supports a functional transsynaptic link between POMC neurons and GnRH neurons (Roa and Herbison, 2012). In contrast, the literature indicates that the POMC neurons in the nucleus of the solitary tract are unlikely to play a role in the control of fertility (Roa, 2013). In OVX ewes, the infusion of an ORL1 antagonist in the ARC does not affect LH pulses (Goodman et al., 2013) but, intracerebroventricular ORL1 antagonists increase LH levels under estradiol/progesterone supplementation, suggesting that OFQ may participate in progesterone-negative feedback (Nestor et al., 2013).

The present study shows Orl1 transcripts in GnRH cells and that OFQ inhibits GnRH neuronal activity, without GABAergic and glutamatergic inputs, and suppresses kisspeptin-10-evoked excitation. The signaling pathway, initiated by ORL1, was identified as Gαo-type G-protein coupling mediated by G-protein-coupled inwardly rectifying potassium (GIRK) channels, most likely GIRK2/3 heteromers. In addition, we found POMC fibers coexpressing OFQ in the vicinity of GnRH neurons and OFQ fibers contacting GnRH neurons in the POA, suggesting that these fibers might originate from POMC neurons in the ARC. Together, these data highlight N/OFQ as a potent in vivo inhibitory signal to GnRH neurons in the mouse.

Materials and Methods

Animals

All procedures were approved by National Institute of Neurologic Disorder and Stroke, Animal Care and Use Committee, and were performed in accordance with National Institutes of Health (NIH) guidelines. Mice were maintained under 12 h light/dark lighting conditions, with food and water available ad libitum. Embryos were collected from timed-pregnant NIH Swiss mice at embryonic day 11.5, and nasal explants were generated (see below)
for PCR and calcium imaging experiments. For in vivo immunocytochemistry, adult intact GnRH-green fluorescent protein (GFP; Mouse Genome Informatics ID 6158458) male mice (Spergel et al., 1999) were used for GnRH/OFQ and GnRH/POMC staining, and adult intact C57BL/6 male mice were used for POMC/OFQ staining. Adult intact GnRH-GFP male mice were also used to generate brain slices for electrophysiological experiments.

**GnRH cells maintained in nasal explants**

Explants were cultured as previously described (Fueshko and Wray, 1994). Briefly, gestational day 11.5 embryos (undetermined sex) were obtained from timed-mated pregnant NIH Swiss mice. Nasal pits were dissected under aseptic conditions in Gey’s Balanced Salt Solution (Life Technologies) supplemented with glucose (Sigma-Aldrich). Explants were adhered onto coverslips by a plasma (Cocalico Biologicals)/thrombin (Sigma-Aldrich) clot and maintained in a defined serum-free medium (SFM) in a humidified atmosphere at 37°C with 5% CO2. On culture day 3, SFM was replaced by fresh SFM containing fluorodeoxyuridine (2.3 μM; Sigma-Aldrich) for 3 d to inhibit the proliferation of dividing olfactory neurons and non-neuronal explant tissue. On culture day 6, and every 2 d afterward, the medium was changed with fresh SFM.

**PCR on cDNA from single GNRIH neurons maintained in explants**

Poly(A)-amplified cDNA libraries were generated from single GnRH neurons using two different techniques (Kramer, 2002; Bosch et al., 2013). Every single-cell cDNA pool generated was first tested by PCR for GnRH (Giacobini et al., 2004). Cellular material without reverse transcriptase, and no cellular material (water), served as negative controls. Specific primers were designed in the 3’-untranslated region of the genes encoding ORL1 within 300 bp before the polyadenylation site. All designed primers were screened using NCBI BLAST (Basic Local Alignment Search Tool; Johnson et al., 2008) to ensure specificity. For each reaction, 1× PCR buffer, 2 μM MgCl2, 250 μM each deoxynucleotide mix (Life Technologies), 125–250 nM forward primer, 125–250 nM reverse primer, and 2.5 U AmpliTaq Gold (Life Technologies) were added to 1–3 μl template cDNA. PCR was performed as follows: initial 10 min denaturation (94°C); 40–50 cycles with denaturation 30 s (94°C); annealing for 30 s (55–66°C) and extension for 2 min (72°C); followed by 10 min postelongation at 72°C. Amplified products were run on a 1.5% agarose gel. Specific bands of the predicted size were observed in control total brain, whereas no bands were seen in water. The sequences of the primers are listed in Table 1.

**Calcium imaging**

GnRH neurons exhibit calcium oscillations that correlate with bursts of action potentials (APs; Constantine and Wray, 2008); therefore, calcium imaging was used as a reflection of GnRH neuronal activity. Experiments were performed as previously described (Constantin et al., 2009). Explants were used between 6 and 11 d in culture (Fig. 1A). Briefly, Calcium Green-1 AM (Life Technologies) was dissolved at 2.7 mM in dimethylsulfoxide containing 20% pluronic F-127 (Life Technologies), then diluted down to 13.5 μM in SFM (loading solution), aliquoted, and kept frozen until use. Explants were incubated in warm loading solution for 20 min at 37°C in a 5% CO2 humidified incubator. After washes in fresh SFM, explants were mounted in a perfusion chamber (Warner Instruments) and continuously perfused at a rate of ~300 μl/min. Calcium imaging experiments were performed at 25°C. Calcium Green-1 was visualized using an inverted microscope (Eclipse TE2000-E, Nikon), through a 20× fluorescence objective [fluor 20×; numerical aperture (NA), 0.75; working distance (WD), 1.0 mm] and a charge-coupled device camera (QImaging) connected to a computer. Time-lapse recording was piloted by iVision imaging software (Scanalytics), and pictures were acquired every 2 s. Excitation wavelengths were provided with a medium-width excitation bandpass filter at 465–495 nm, and emission was monitored through a 40 nm bandpass centered on 535 nm. Calcium imaging recordings were divided into periods. The treatment period was preceded by a control period in SFM, followed by a washout period. When possible, drugs were sequentially added through multiple treatment periods. When a drug required a long pretreatment [i.e., pertussis toxin (PTX) or tertiapin-Q (TPNQ)], recordings were started during the last 10 min of pretreatment to determine the basal level of GnRH neuronal activity. All recordings were terminated with a 40 mM KCl stimulation to ensure the viability of the cells. The changes in fluorescence over time were measured in single, phase bright, bipolar cells with iVision and analyzed with MATLAB (MathWorks) as previously described (Constantine and Wray, 2008). The phenotype of these bipolar cells, defined as regions of interest, was confirmed using chromogen immunocytochemistry against GnRH. Due to the heterogeneity of GnRH neuronal population, a large number of cells were sampled for each paradigm. The individual cells (N) originating from at least three explants (N) independently recorded were combined for each paradigm. Explants (N = 65) contained on average 29.5 ± 2.0 identified GnRH neurons in a recording field.

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**Table 1: Primer sequences**

| Gene (NCBI/GenBank ID) Reference Sequence | Primers sequences (5’ to 3’) | Annealing temperature | Product size |
|------------------------------------------|-------------------------------|-----------------------|--------------|
| GnRH-1 (NM_008145.2)                    | CTG ATG GCC GGC ATT CTA CTG C | 66°C                  | 220 bp       |
|                                          | CCA GAG CTC CTC GCA GAT CCC   |                       |              |
| Opioid receptor-like 1 (NM_011012.5)     | CAT GGC ATG CAG AAC CCA G     | 55°C                  | 209 bp       |
|                                          | AGG GCT AGC TAC ATG CAC GA    |                       |              |
Immunocytochemistry for GnRH

After calcium imaging, explants (6–11 d) were fixed for 30 min with 0.1 M PBS, pH 7.4, containing 4% formaldehyde at room temperature. After a few washes in PBS, explants were incubated in a blocking solution (10% normal horse serum plus 0.3% Triton X-100) for 1 h, and washed several times in PBS. The explants were incubated at 4°C overnight in the primary antibody (rbGnRH; Wray et al., 1989b; Table 4). The next day, explants were washed in PBS, incubated for 1 h with biotinylated secondary donkey anti-rabbit antibody (1:500 in PBS/0.3% Triton X-100; Vector Laboratories), washed in PBS, and processed for avidin–biotin horseradish peroxidase/3,3′-diaminobenzidine (Fig. 1A).

Immunofluorescent labeling

Three explants (6–11 d) were fixed, washed, and blocked as described above. The explants were then incubated (4°C overnight) in the GIRK2 primary antibody, which has been shown to give no signal in GIRK2 knockout mice (Marron Fernandez de Velasco et al., 2017). The next day, explants were washed in PBS, and the first primary antibody was visualized using Alexa Fluor 488-conjugated secondary donkey anti-rabbit antibody (1:500 in PBS/0.3% Triton X-100; Vector Laboratories), washed in PBS, and processed for avidin–biotin horseradish peroxidase/3,3′-diaminobenzidine (Fig. 1A).

After washing (PBS), the second primary antibody was visualized using Alexa Fluor 555-conjugated secondary donkey anti-mouse antibody (1:1000 in PBS/0.3% Triton X-100). After several washes in PBS and water, explants were coveredlipped with an anti-fade mounting solution. Controls (N = 3) in which the anti-GIRK2 primary antibody was omitted resulted in only background staining.

OFQ or POMC staining and GFP staining in GnRH-GFP mice

Sections were incubated at 4°C (2 nights) in the first primary antibody (OFQ or POMC). The next day, sections were washed in PBS, incubated for 1.5 h with Alexa Fluor 555-conjugated secondary donkey anti-rabbit antibody.
To avoid cross-reactivity between two primary antibodies raised in a rabbit, staining was performed as described previously (Shindler and Roth, 1996; Hoffman et al., 2008). Sections were incubated at 4°C (for 2 nights) in the first primary antibody (POMC; Elkabes et al., 1989). The next day, sections were washed in PBS, incubated with biotinylated secondary donkey anti-rabbit antibody (1 h, 1:1000 in PBS/0.3% Triton X-100). After washes in PBS, sections were incubated in avidin–biotin complex (1 h, 1:200 in PBS/0.3% Triton X-100), washed, and incubated in tyramide (20 min, 1:200 in PBS/0.3% Triton X-100), washed and incubated (1:500 in PBS/0.3% Triton X-100) with Cy3-tyramide secondary donkey anti-rabbit antibody (1 h, 1:500 in PBS/0.3% Triton X-100). Data are expressed as the mean ± SEM. For each experiment, one-way ANOVA, followed by Sidak’s multiple-comparisons test (p < 0.01). For rows j, k, m, and n, a paired Student’s t test was used to compare period 2 with period 1 (p < 0.01).
Figure 2. OFQ inhibits GnRH neuron calcium oscillation, independent of GABAergic and glutamatergic inputs. **A**, Heatmaps of changes in levels of intracellular calcium in cells recorded simultaneously during the paradigm in **B**, independently repeated in five different explants. Each row represents changes in a single cell. White lines indicate the time of drug application. Note the consistency of the response to OFQ within each explant and across explants (Fig. 2-1). **B**, Left, Calcium-imaging recordings of explants showing OFQ (100 pM) evoked a potent decrease in the frequency of calcium oscillations in GnRH neurons (kymograph from the cell shown on the top trace). Right, Empty circles, Summary data showing the average frequency of calcium oscillations for each explant. To minimize the impact of the cell-to-cell heterogeneity, the values from all individual cells originating from the five explants in **A** were combined. Bars, Summary data showing the average frequency of calcium oscillations in all GnRH neurons tested during the control (SFM), treatment (OFQ), and the washout (SFM WO) periods (n = 113, N = 3). au, Arbitrary unit. **C**, Left, The OFQ-evoked inhibition persisted in the presence of AABs (BIC, 20 μM; CNQX, 10 μM; D-AP5, 20 μM). Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons tested during the control (SFM), pretreatment (AAB), treatment (+OFQ), and washout (SFM WO) periods (n = 144, N = 5).
primary antibody (OFQ). Sections were then washed in PBS, incubated with Alexa Fluor 488-conjugated donkey anti-rabbit antibody (1.5 h, 1:1000 in PBS/0.3% Triton X-100), washed in PBS, and mounted and coverslipped with an anti-fade mounting solution. For triple labeling (one series/animal), an antigen retrieval method was used before initiating the immunocytochemistry (Jiao et al., 1999), and staining for GnRH was performed after staining for POMC and OFQ using an GnRH antibody raised in chicken. Sections were incubated for 2 d, washed, and visualized with Alexa Fluor 647-conjugated donkey anti-chicken antibody.

Table 3: Inhibition of GnRH neurons

| Paradigms with OFQ(1-13) | Period (with OFQ) – Period (without OFQ)/Period (with OFQ) (change in %) | Cells (n) | Explants (N) |
|--------------------------|--------------------------------------------------------------------|----------|------------|
| a | SFM – OFQ | -75.6 ± 2.0 | 144 | 5 |
| b | UFP – UFP + OFQ | -17.3 ± 3.3* (p < 0.0001) | 112 | 3 |
| c | SB + SB + OFQ | -29.5 ± 2.5* (p < 0.0001) | 244 | 6 |
| d | PTX pretreated: SFM – OFQ | -17.4 ± 4.4* (p < 0.0001) | 88 | 3 |
| e | IBMX + FSK – IBMX + FSK + OFQ | -68.2 ± 2.6 (p > 0.9864) | 161 | 3 |
| f | Cs – Cs + OFQ | -28.6 ± 3.6* (p < 0.0001) | 168 | 5 |
| g | Ba – Ba + OFQ | -12.0 ± 4.9* (p < 0.0001) | 36 | 4 |
| h | TPNQ – TPNQ + OFQ | -72.9 ± 2.5 (p > 0.9999) | 80 | 3 |
| i | PMA – PMA + OFQ | 18.7 ± 16.8* (p < 0.0001) | 113 | 4 |
| j | SFM – BAC | -86.2 ± 1.9 | 88 | 4 |
| k | TPNQ – TPNQ + BAC | -93.5 ± 1.5 (p = 0.9206) | 53 | 3 |
| l | PMA – PMA + BAC | 19.9 ± 20.5* (p < 0.0001) | 92 | 3 |

Concentrations used: OFQ, 100 pM; UFP, 10 nM; SB, 10 nM; PTX, 250 ng/ml; IBMX, 10 μM; FSK, 1 μM; Cs, 5 mM; Ba, 400 μM; TPNQ, 500 nM; PMA, 50 nM; BAC, 10 μM. Data are expressed as the mean ± SEM.

*Significant difference (p < 0.01), using one-way ANOVA, followed by Dunnett’s multiple-comparisons test using OFQ (row a) as a reference for rows b–i or BAC (row j) as a reference for rows k and l.
The brain was removed from the skull, glued to the vibratome, and continuously superfused with oxygenated normal ACSF maintained at 28–30°C at a rate of 2 ml/min (Constantin et al., 2012). Individual GnRH neurons were identified with fluorescence (20 nm narrow bandpass EGFP filter centered at 480 nm) using a 40× water-immersion objective (40×/0.80 W; WD, 2.0 mm; Nikon). Visualized with a charge-coupled device camera (Retiga EXi Blue, QImaging) and piloted by the open source software Micro-Manager version 1.4, the neurons were patched under fluorescence and differential interference contrast. The pipettes (3–5 MΩ) were backfilled with aCSF. Electrophysiological recordings were acquired with a Multiclamp 700B amplifier (Molecular Devices) using a low-pass filter at 10 kHz and digitized by a Digidata (1550) analog-to-digital converter at 10 kHz (Molecular Devices).

### Drugs
Cesium (Ca), barium [Ba; both broad-spectrum blockers of inwardly rectifying potassium (K$_{ir}$) channels], naringin (GIRK channel activator), phorbol 12-myristate 13-acetate [PMA; protein kinase C (PKC) activator], and baclofen (BAC; GABA$_B$ receptor agonist) were purchased from Sigma-Aldrich. 3-isobutyl-1-methylxanthine (IBMX; inhibitor of phosphodiesterase), forskolin (FSK; activator of adenyl cyclase), 6-Cyano-7-nitroquinoxaline-2,3-dione (CNQX; AMPA/kainate receptor antagonist), d(-)2-amino-5-phosphonopentanoic acid (d-AP5; NMDA receptor antagonist), bicuculline (BIC; GABA$_A$ receptor antagonist), TPNQ (blocker of GIRK channels), UFP-101 (UFP) and SB 162111 (SB; two selective antagonists for ORL1 receptor), PTX (uncoupling G proteincoupled re-
Figure 4. The receptor ORL1 is $G_{i/o}$ coupled and activates GIRK channels to inhibit GnRH neuronal activity. 

**A**, Left, Calcium imaging recordings, after a 4 h pretreatment with PTX (250 ng/ml), showing OFQ (100 pM) evoked only a mild inhibition of GnRH neuronal activity. Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons pretreated with PTX and tested during the control (PTX-SFM), treatment (+OFQ), and washout (SFM WO) periods ($n = 89, N = 3$). au, Arbitrary unit. *Statistical significance between two consecutive periods.

**B**, Left, The OFQ-evoked inhibition is insensitive of IBMX and FSK. Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons tested during control (SFM), pretreatment (IBMX + FSK), treatment (+OFQ), and washout (SFM WO) periods ($n = 161, N = 3$). Note that the IBMX + FSK treatment is not showing a significant increase due to the calcium plateau evoked in some cells, which results in an underestimate of peak detection.

**C, D**, Left, The broad-spectrum blocker of Kir channels, Cs or Ba, reduced the OFQ-evoked inhibition. Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons pretreated with Cs or Ba and tested during the pretreatment (Cs or Ba) and treatment (+OFQ) periods (Cs: $n = 169, N = 5$; Ba: $n = 36, N = 4$).
Figure 5. The OFQ-evoked inhibition activates GIRK channels, insensitive to tertiapin–Q but inhibited by protein kinase C phosphorylation. **A**, Left, Calcium imaging recordings showing that a specific blocker of G-protein inwardly rectifying potassium channels, TPNQ, was ineffective on the OFQ-evoked inhibition. Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons tested during control (SFM), pretreatment (TPNQ), treatment (+OFQ), and washout (SFM WO) periods (n = 80, N = 3). au, arbitrary unit. *Statistical significance between two consecutive periods. **B**, Left, TPNQ was also ineffective on GABAB-evoked inhibition with BAC. Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons tested during control (SFM), pretreatment (TPNQ), treatment (+BAC), and washout (SFM WO) periods (n = 20, N = 2). **C**, Left, The
protein kinase C activator PMA prevented OFQ-evoked inhibition. Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons tested during control (SFM), pretreatment (PMA), treatment (+OFQ), and washout (SFM WO) periods ($n = 118, N = 4$). **D.** Left, PMA was also profoundly blunted the GABA$_A$-evoked inhibition. Right, Summary data showing the average frequency of calcium oscillations in all GnRH neurons tested during control (SFM), pretreatment (PMA), treatment (+BAC), and washout (SFM WO) periods ($n = 92, N = 3$).

**Table 2**

| Concentration (mM) | Number of Cells | Percentage of Cells with a 50% Inhibition | Significance of Dose Difference |
|-------------------|----------------|----------------------------------------|--------------------------------|
| 100               | 20             | 50%                                    | $p < 0.01$                      |
| 10                | 15             | 75%                                    | $p = 0.05$                      |

**Orphanin FQ (1-13) directly inhibits GnRH neurons maintained in explants**

OFQ was applied to 6-11 d explants at four different concentrations (10 nm, 1 nm, 100 pm, and 10 pm). All doses inhibited GnRH neuronal activity (**Table 2**, rows a–d, **Fig. 2**). The mean response of individual explants was quantified to assess whether the OFQ response might be sex specific. OFQ evoked a potent inhibition of GnRH neurons in every explant tested ($N = 20$, from all four doses combined; **Fig. 2-1**); therefore, the sex of the embryo from which the explant was generated is highly unlikely to define the responsiveness to OFQ. At 100 pm, ~90% of the cells showed a ≥50% inhibition with OFQ, and the inhibition was not correlated with cell location within the explant periphery (**Fig. 2A**). The dose of 100 pm was chosen for all subsequent experiments, as it was the smallest dose with robust effects (**Fig. 2B**).

**Statistical analysis**

For calcium imaging, comparisons of the frequencies of calcium oscillations (in peaks/min) were performed using a repeated measurement one-way ANOVA with Greenhouse–Geisser correction, and *post hoc* Sidak’s multiple-comparisons test between two consecutive recording periods for paradigms (**Table 2**). A paired Student’s t test was used for two-period paradigms (**Table 2**). The strength of inhibition was determined for each cell and was expressed as the percentage of the pre-OFQ period as follows: \[ \frac{\text{peaks/min during OFQ} - \text{peaks/min before OFQ}}{\text{peaks/min before OFQ}} \times 100 \]. Comparisons of the strength of inhibition between groups of cells were performed with one-way ANOVA, and *post hoc* Dunnett’s multiple-comparisons test, using OFQ as the reference. In the Results section and figures, the frequencies of calcium oscillations are expressed as the mean ± SEM, and $n$ and $N$ represent the number of cells and explants recorded, respectively.

For electrophysiology, the OFQ-induced inhibition was determined as follows: APs were detected with Clampfit 10 on continuous recordings, and the firing frequency (in Hz) was determined by summing APs into 1 s bins. The average firing frequency was calculated over the last minute of each recording period, before OFQ, during OFQ and washout after OFQ (<6 min). When firing was not regained spontaneously, kisspeptin-10 was applied to validate the period of silence. Periodograms (in Hz) were performed by summing APs into 10 s bins. A Student’s paired t test was performed between two consecutive recording periods. $n$ and $N$ represent the number of cells and the number of animals the cells are recorded from, respectively. In calcium imaging, significant differences were defined by a p value <0.01 due to the large number of cells that can be sampled; in electrophysiology, significant differences were defined by a p value <0.05.

**Results**

RT-PCR analysis revealed the presence of transcripts coding for the ORL1 gene ($n = 14$ and $19$) in single GnRH neurons maintained in nasal explants using two different single-cell RT-PCRs ($n = 15$ and $19$; **Fig. 1B**).

**The receptor of orphanin FQ, ORL1, mediates the OFQ inhibition**

UFP is a selective antagonist for ORL1 receptors (**Calo et al., 2002**). Pretreatment of GnRH neurons with UFP (10 nm) did not completely prevent OFQ-induced inhibition (**Table 2**, row f, **Fig. 3A**) but dramatically attenuated it (**Table 3**, rows a and b), with only ~12% of the cells showing a ≥50% inhibition. The effectiveness of UFP-101 is shown with cells losing their ability to respond to the second dose of OFQ (see **Fig. 6A,B**). The experiment was repeated with SB, another selective antagonist for ORL1 receptors. Pretreatment of GnRH neurons with SB (10 nm) also attenuated OFQ inhibition (**Tables 2**, row g, **Fig. 3B**) with ~35% of the cells showing a ≥50% inhibition. These data support the binding of OFQ to ORL1 receptors in GnRH neurons. Note that UFP and SB did not have any effect on their own (**Table 2**, rows f and g).

**Table 3**

| Concentration (mM) | Number of Cells | Percentage of Cells with a 50% Inhibition | Significance of Dose Difference |
|-------------------|----------------|----------------------------------------|--------------------------------|
| 100               | 20             | 50%                                    | $p < 0.01$                      |
| 10                | 15             | 75%                                    | $p = 0.05$                      |
Figure 6. Blockers are effective in GnRH neurons identified as OFQ responsive. **A–D**, Left, Calcium imaging recordings of explants showing that OFQ (100 pM) evoked a potent decrease in the frequency of calcium oscillations in GnRH neurons (first application, **A–D**). The inhibition is repeatable (second application, **A**) and blocked by UFP-101 (ORL1 antagonist), cesium (broad-spectrum blocker of Kir channels), and PMA (PKC activator). **A–D**, Right, Heatmaps of changes in the levels of intracellular calcium in cells recorded simultaneously during different paradigms (left). Each row represents changes in a single cell. White lines indicate the time of drug application. Note the large number of GnRH cells inhibited by OFQ (first application, **A–D**), the repeatability of the inhibition (second application, **A**), and the effectiveness of UFP-101 (ORL1 antagonist), cesium (broad-spectrum blocker of Kir channels), and PMA (PKC activator; **B, C, and D**, respectively, second application) to prevent OFQ inhibition. Note the homogeneity of the response throughout the cells within an explant.
OFQ inhibition in GnRH cells is mediated by G<sub>ior</sub>-protein-coupled inwardly rectifying potassium channels

It has been reported that ORL1 receptors couple to a G<sub>ior</sub>-type G-protein (Ikeda et al., 1997). Thus, explants were treated with PTX (250 ng/ml) for >4 h to uncouple G<sub>ior</sub>-proteins from its receptors. PTX treatment attenuated OFQ inhibition of GnRH neurons (Tables 2, row h, 3, rows a and d, Fig. 4A). A canonical pathway downstream of a G<sub>ior</sub>-type protein is a decrease in the activity of adenylyl cyclase and cyclic adenosine monophosphate. However, coapplication of IBMX (10 μM) and FSK (1 μM), inhibiting phosphodiesterase and activating adenylyl cyclase, respectively, had no effect on OFQ inhibition of GnRH neurons (Tables 2, row i, 3, rows a and e, Fig. 4B). Alternatively, G<sub>ior</sub>-proteins can be coupled to GIRK channels, also known as K<sub>r</sub>-3, voltage-independent potassium channels (Klenke et al., 2010; Constantin and Wray, 2016). The application of OFQ was repeated in the presence of Cs (5 mM) or Ba (400 μM), broad-spectrum blockers of K<sub>r</sub> channels, added 10 and 20 min, respectively, before OFQ. Although Cs or Ba did not fully block the OFQ inhibition (Table 2, rows j and k, Fig. 4C,D; also see Fig. 6C), both strongly reduced its magnitude (Table 3, rows a and f or g), supporting the coupling of ORL1 receptors to GIRKs, as previously shown (Wagner et al., 1998).

Unlike the broad-spectrum K<sub>r</sub> blockers cesium and barium, TPNQ is known as a GIRK-specific channel blocker. Unexpectedly, TPNQ (500 nM; Wu et al., 2011), added 10 min before OFQ, did not prevent the OFQ inhibition (Table 2, row l, Fig. 5A) and had no effect on its magnitude (Table 3, rows a and h). It is widely agreed that GABA<sub>B</sub> receptors are mediated by GIRK channels (Kahnovitch et al., 2017), including in GnRH neurons (Zhang et al., 2009). Thus, the effect of TPNQ was tested on the inhibition triggered by the activation of GABA<sub>B</sub> receptor. As with OFQ, TPNQ had no effect on BAC inhibition (Table 2, rows p and q, Fig. 5B) or its magnitude (Table 3, rows j and k), and confirmed the ineffectiveness of TPNQ on GnRH GIRK channels. Naringin (500 μM), which is an activator of GIRK channels using the same binding site as TPNQ (Yow et al., 2011), also did not change GnRH neuronal activity (Table 2, row m).

GIRK channels are tetramers and can form with three neuronal subtypes, GIRK1 (K<sub>r</sub>.3.1), GIRK2 (K<sub>r</sub>.3.2), and GIRK3 (K<sub>r</sub>.3.3). GIRK2 can form a homotetramer, while GIRK1 and GIRK3 require heteromerization for functionality. Previous RT-PCR analysis had found transcripts for Kcnj3 (GIRK1) in GnRH neurons (Constantin and Wray, 2016).
However, ML297 (10 μM), an activator of GIRK1-containing tetramers (Kaufmann et al., 2013) had no effect on GnRH neuronal activity (Table 2, row n). RT-PCR data also identified transcripts for Kcnj9 (GIRK3) in GnRH neurons (Klenke et al., 2010). Since GIRK2 and GIRK3 are less sensitive or insensitive to TPNQ, respectively (Nocke mann et al., 2013; Whorton and MacKinnon, 2013; Llamosas et al., 2017), an alternative approach was used to clarify the role of GIRK channels in the OFQ inhibition. GIRK channels are inhibited through a PKC-dependent
mechanism (Stevens et al., 1999). The PKC activator PMA (50 nM) was applied before OFQ, and OFQ failed to inhibit GnRH neurons (Table 2, row o, Fig. 5C; Table 3, rows a and i, Fig. 6D). A similar effect was observed with PMA applied before BAC (Tables 2, row r, 3, rows j and l, Fig. 5D). These results suggested the absence of GIRK1 but the presence of the other neuronal form, GIRK2, in GnRH neurons. Immunofluorescent staining with a validated antibody (Marron Fernandez de Velasco et al., 2017) demonstrated the presence of GIRK2 in all GnRH neurons (Fig. 7), consistent with its ubiquitous presence in most neurons (Lüscher and Slesinger, 2010). Together, these data are consistent with GIRK2/3 channels, downstream of ORL1 receptor, mediating the OFQ inhibition in GnRH neurons.

**OFQ fibers contact GnRH neurons**

To begin to identify the anatomic circuit, immunocytochemistry for OFQ and GnRH was performed. In contrast to ewes (Foradori et al., 2007), GnRH neurons in adult mice were not immunoreactive for OFQ. However, OFQ fibers were observed in the POA, where most GnRH cell bodies are found, and within this region some OFQ fibers were found apposed to GnRH fibers and cell bodies (Fig. 8). In the rat, a subpopulation of POMC neurons in the ARC is labeled with OFQ (Maolood and Meister, 2010) and POMC fibers contact GnRH neurons (Leranth et al., 1988; Simonian et al., 1999). Thus, the possibility that OFQ fibers contacting GnRH neurons originated from POMC neurons in the ARC was investigated. In the mouse, a subset (estimated 5–10%) of POMC neurons was immunoreactive for OFQ (Fig. 9), but all OFQ neurons were POMC positive. POMC fibers were observed apposed to GnRH neurons in the POA (Fig. 10A–C). Notably, the POA also contained some fibers colabeled with OFQ and POMC (Fig. 10D). However, using triple immunolabeling for GnRH, OFQ, and POMC, fibers colabeled with
Figure 10. POMC-immunoreactive fibers contact GnRH neurons and POMC fibers in the POA coexpress OFQ. **A–C.** Representative stack of confocal images showing GnRH neurons labeled with GFP- (green) and POMC- (red) immunoreactive fibers in an adult male mouse. Scale bar, 20 $\mu$m. Enlarged image from boxed area in **B.** Scale bar, 5 $\mu$m. **C.** Single confocal plan from boxed area showing a POMC-immunoreactive fiber apposed to this GnRH cell body in the POA. **D.** Representative confocal images showing fibers immunoreactive for OFQ and POMC in the POA. Scale bars, 10 $\mu$m. Note individual varicosities in the same fiber which are immunolabeled for only OFQ (single-labeled; open arrowheads, third panel) or for both OFQ and POMC (dual-labeled; filled arrowheads), suggesting that spatial localization of these peptides might occur in processes. **E, F.** Representative confocal image (Z projection, ~10 $\mu$m) showing immunoreactivity for OFQ (green) and POMC (red) in fibers surrounding GnRH (blue) neurons in the POA. Scale bar, 50 $\mu$m. **F1–F3.** Boxed areas in **E** enlarged show contacts (arrowheads) between GnRH and POMC fibers (F1), between GnRH and OFQ fibers (F2), and near cell bodies (F3). Scale bars, 10 $\mu$m.
POMC and OFQ were not found contacting GnRH neurons (Fig. 10E,F).

**Figure 11.** OFQ inhibits the GnRH neuron firing rate in acute brain slices, independent of GABAergic and glutamatergic inputs. **A**, Left, Electrophysiological recording of adult GFP-tagged GnRH neurons showing that OFQ (1 μM, 100 nM, and 10 nM) evoked a potent decrease in GnRH neuron firing rate. Right, Summary data showing quantification of firing rate (in Hz) in individual GnRH neurons tested at the different concentrations during the control (CTRL), treatment (OFQ), and washout (WO) periods. The values represent the average of 1 s bins for the last 1 min of each period identified on the traces. **B**, Left, OFQ (10 nM) in the presence of AABs (BIC, 20 μM; CNQX, 10 μM; D-AP5, 20 μM) still decreased GnRH neuron firing rate. Right, Summary data showing quantification of the firing rate (in Hz) in individual GnRH neurons tested during the control (CTRL), pretreatment (AAB), treatment (+OFQ), and washout (WO) periods. The values represent the average of 1 s bins for the last 1 min of each period identified on the traces.

POMC and OFQ were not found contacting GnRH neurons (Fig. 10E,F).

**OFQ inhibits GnRH neurons in situ, independently of GABAergic and glutamatergic inputs**

OFQ was applied at different doses to GnRH neurons in acute brain slices. OFQ stopped the firing in two of two cells (N = 2), three of three cells (N = 2), and two of two cells (N = 1) at 1 μM, 100 nM, and 10 nM, respectively (Fig. 11A). After the blockade of GABAergic and glutamatergic excitatory inputs with AAB (BIC, 20 μM; CNQX, 10 μM; D-AP5, 20 μM), OFQ (10 nm) still inhibited eight of nine cells, five of them being stopped (N = 3; Fig. 11B). Of the 12 cells that stopped, without (n = 7) or with AAB (n = 5), the average time to regained firing was after 9.0 ± 1.3 min (N = 8). OFQ (10 nM) suppressed kisspeptin-10 (100 nM)-evoked excitation (Fig. 12A–C; four of four cells tested; N = 4) but did not terminate it. The OFQ suppression was partially prevented by UFP-101 (500 nM; N = 3), supporting the role of ORL1 (Fig. 12B–D). These data support a role for OFQ as a modulator of GnRH neuron neuronal activity but rule out its function as the terminator of kisspeptin-induced excitation.

**Discussion**

The present study investigated the neuromodulation of GnRH neurons by OFQ in the mouse. We report that OFQ strongly inhibits spontaneous GnRH neuronal activity and can repress kisspeptin-evoked excitation. The inhibition, independent of GABAergic and glutamatergic inputs, is mainly mediated by ORL1, G<sub>io</sub>-protein-coupled receptor, and the subsequent activation of GIRK channels. In vivo, OFQ-immunopositive fibers were found contacting GnRH neurons. Together, these data suggest that OFQ can downregulate the reproductive axis.
Figure 12. OFQ as a new player in the ARC regulation of GnRH neurons. **A**, Left, Electrophysiological recording of an adult GFP-tagged GnRH neuron showing OFQ (10 nM) suppressed the firing rate of kisspeptin-10 (kp-10; 100 nM)-activated GnRH neurons. Right, Corresponding periodogram illustrating the changes in GnRH neuron firing rate (in Hz) over time (i.e., the increase in firing rate induced by kp-10 and its suppression by OFQ). The bars represent the instant frequency in 10 s bins. **B**, Electrophysiological recording of another adult GFP-tagged GnRH neuron showing OFQ (10 nM) suppressed the firing rate of kp-10 (100 nM)-activated GnRH neurons. The inhibition evoked by OFQ was partially antagonized by UFP-101 (500 nM) but was reestablished by a second application of OFQ after the washout of UFP-101. **C**, Electrophysiological recording of a third adult GFP-tagged GnRH neuron showing the partial antagonism of OFQ by UFP-101. Note that, without UFP-101, the OFQ evoked a pause in the tonic firing for ~11 min (red arrow, on the left), while with UFP-101 the pause was ~2.5 min (red arrow, on the right; i.e., an approximately four times faster recovery despite being further from the initial kp-10 application. **D**, Quantification of the OFQ inhibition, before (square) and after
(circle) the application of UPF-101. The values represent the firing rate 4 min after OFQ or UPF-101 plus OFQ application, normalized to the firing rate in the last minute of aCSF application before application of OFQ (empty square) or UPF-101 before UPF-101 plus OFQ (empty circle) application, respectively. **E**. Diagram representing where OFQ stands within the already established connectivity around GnRH neurons. GnRH neuron (blue) receive excitatory inputs from RP3V kisspeptin neurons (red) at the soma. ARC kisspeptin neurons (purple), autoregulated via a neurokinin B and dynorphin loop, contact GnRH neuronal processes around the median eminence. Both kisspeptin neuronal populations are linked via ARC → RP3V connections. POMC neurons (green) contact GnRH neurons (Leranth et al., 1988; Simonian et al., 1999). ARC kisspeptin neurons contact POMC neurons (Fu and van den Pol, 2010; Nestor et al., 2016). OFQ fibers (orange), possibly originating from a subpopulation of POMC neurons and/or another unidentified neuronal population, contact GnRH neurons, providing an inhibitory signal.

OFQ inhibits GnRH secretion from mediobasal fragments (Dhandapani and Brann, 2002; An et al., 2005) or in vivo [push–pull (An et al., 2005); intracerebroventricular (An et al., 2005, 2009)]. The in vivo inhibition is ORL1 dependent (An et al., 2005, 2009). However, since the receptor is widely distributed in the brain (Ikeda et al., 1998; Neal et al., 1999; Houtani et al., 2000) and even in the hypothalamus (Chen et al., 2017), how OFQ inhibited GnRH secretion was unknown. Evidence for indirect actions of OFQ on GnRH neurons exist, including the following: (1) transcripts for ORL1 were not detected in an immortalized GnRH cell line obtained from mouse (Dhandapani and Brann, 2002); and (2) the stimulatory effect of ORL1 antagonists on POA GnRH release were mediated by glutamate in male rats (An et al., 2008). However, our calcium imaging and patch-clamp data indicate a direct action of OFQ onto GnRH neurons since OFQ inhibited GnRH neuronal activity while both GABAergic and glutamatergic inputs were disrupted. Notably, GnRH neuronal activity does not require these two excitatory inputs (Constantin et al., 2010; Lee et al., 2012), but they are the main excitatory inputs in acute brain slices (Iremonger et al., 2010; Herbison and Moenter, 2011) and largely contribute to GnRH neuronal activity in nasal explants (Constantin et al., 2010). Inhibition of GnRH neuronal activity by OFQ is consistent with previous data showing OFQ hyperpolarized GnRH neurons from OVX guinea pig through an inwardly rectifying potassium current (Wagner et al., 1998). A direct action of OFQ on GnRH neurons is further supported by our findings of ORL1 transcript in GnRH neurons and OFQ fibers contacting GnRH neurons.

Examination of the signaling pathway demonstrated that the OFQ inhibition was ORL1 mediated. ORL1 coupling to Gαo-type G-protein (Reinscheid et al., 1995; Ikeda et al., 1997) was confirmed in GnRH neurons treated with PTX. OFQ-induced inhibition could occur through two signaling pathways: a decrease of cyclic adenosine monophosphate and/or the activation of GIRK channels (for review, see Al-Hasani and Bruchas, 2011). The data obtained using IBMX and FSK clearly indicate that the OFQ-induced inhibition of GnRH neuron activity occurs without a decrease of cAMP. In fact, previous studies from our laboratory have already shown that a decrease of cAMP does not affect GnRH neuronal activity (Constantin and Wray, 2008, 2016), while the activation of Cs-and Ba-sensitive channels inhibits GnRH neurons (Klenke et al., 2010; Constantin and Wray, 2016).

Most neurons express GIRK1, GIRK2, and GIRK3 (Lüscher and Slesinger, 2010). Transcripts for both GIRK1 (K,3.1, Kcnj3; Constantin and Wray, 2016) and GIRK3 (K,3.3, Kcnj9; Klenke et al., 2010) subunits are expressed in GnRH neurons. Neither GIRK1 nor GIRK3 subunits can form homotetrameric functional GIRK channels but can assemble into functional heterotetramers (Lüscher and Slesinger, 2010). The sensitivity of the OFQ inhibition to Cs and Ba, but not to TPNQ, naringin, or ML297, indicated a different subunit composition of the GIRK channels associated with ORL1 (Kobayashi et al., 2011) in GnRH cells and suggested the presence of GIRK2 subunits in their GIRK channels. To further test this hypothesis, baclofen, a GABAB agonist, was used to activate GIRK channels (Kahanovitch et al., 2017). Although GABAB can use two-pore domain potassium channels (Bushell et al., 2002; Deng et al., 2009), the literature supports the role of GABAB with inwardly rectifying potassium currents in GnRH neurons (Wagner et al., 1998; Zhang et al., 2009). BAC-induced inhibition was also insensitive to TPNQ. This observation is supported by a study showing that GIRK channels coupled to ORL1, at least partially, overlap the GABAergic GIRK channel pool (Wagner et al., 1998). Since no other GIRK-specific blockers are available, an alternative route to test GIRK2 involvement in OFQ signaling via ORL1 was used. Protein kinase C is known to phosphorylate GIRK channels, GIRK2 included (Adney et al., 2015), and to desensitize GIRK-mediated currents (Stevens et al., 1999; Lüscher and Slesinger, 2010). Together with our earlier data, the lack of OFQ inhibition after the application of a PKC activator and the immunoreactivity of GnRH neurons for GIRK2 show that OFQ inhibition of GnRH neurons is mediated by GIRK channels containing GIRK2/3 subunits.

OFQ fibers were identified contacting GnRH fibers and cell bodies in the POA, supporting a direct effect of OFQ on GnRH cells. One could argue that the contacts on GnRH fibers are irrelevant to the modulation of GnRH neuron firing; however, GIRK channels localize to specific subcellular compartments (Kulik et al., 2006). GIRK2 splice variants exhibit different subcellular distribution and might impact the integration of afferent inhibitory inputs (Marron Fernandez de Velasco et al., 2017). In addition, GABA inhibition in GnRH neurons is largely lost in preparations that sever processes (Constantin et al., 2012). GnRH neuronal cell bodies were not immunolabeled for OFQ in the adult mouse. This is in contrast to data from the sheep where virtually all GnRH neurons express OFQ in ewes (Foradori et al., 2007). GnRH neuron immunoreactivity for OFQ in sheep might be a species-specific and lasting vestige of their embryonic origin. Indeed, the litera-
In summary, using calcium imaging, immunohistochemistry, and electrophysiology, we have confirmed that OFQ can inhibit GnRH neurons and suppress, but not end, kisspeptin activation of GnRH neurons. Together, these results bring into light a potent neuromodulator of GnRH neurons in the mouse and contribute to our knowledge of the neuronal network upstream of GnRH neurons.

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