



Comparison of the pharmacologic profiles of arginine vasopressin and oxytocin analogs at marmoset, titi monkey, macaque, and human oxytocin receptors



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ARTICLE INFO

Keywords:

Arginine vasopressin
Oxytocin
Oxytocin receptor
G-protein coupled receptor
Calcium-activated potassium channel

ABSTRACT

The oxytocin-arginine vasopressin (OT-AVP) ligand-receptor family influences a variety of physiological, behavioral, and social behavioral processes in the brain and periphery. The OT-AVP family is highly conserved in mammals, but recent discoveries have revealed remarkable diversity in OT ligands and receptors in New World Monkeys (NWMs) providing a unique opportunity to assess the effects of genetic variation on pharmacological signatures of peptide ligands. The consensus mammalian OT sequence has leucine in the 8th position (Leu⁸-OT), whereas a number of NWMs, including the marmoset, have proline in the 8th position (Pro⁸-OT) resulting in a more rigid tail structure. OT and AVP bind to OT's cognate G-protein coupled receptor (OTR), which couples to various G-proteins (G_{i/o}, G_q, G_s) to stimulate diverse signaling pathways. CHO cells expressing marmoset (mOTR), titi monkey (tOTR), macaque (qOTR), or human (hOTR) OT receptors were used to compare AVP and OT analog-induced signaling. Assessment of G_q-mediated increase in intracellular calcium (Ca²⁺) demonstrated that AVP was less potent than OT analogs at OTRs from species whose endogenous ligand is Leu⁸-OT (tOTR, qOTR, hOTR), relative to Pro⁸-OT. Likewise, AVP-induced membrane hyperpolarization was less potent at these same OTRs. Evaluation of (Ca²⁺)-activated potassium (K⁺) channels using the inhibitors apamin, paxilline, and TRAM-34 demonstrated that both intermediate and large conductance Ca²⁺-activated K⁺ channels contributed to membrane hyperpolarization, with different pharmacological profiles identified for distinct ligand-receptor combinations. Understanding more fully the contributions of structure activity relationships for these peptide ligands at vasopressin and OT receptors will help guide the development of OT-mediated therapeutics.

1. Introduction

AVP and OT are synthesized in the magnocellular neurons of the supraoptic and paraventricular nuclei in the hypothalamus and stored in the posterior pituitary, where they are secreted in the bloodstream and affect a number of physiological functions including maintaining water homeostasis (AVP) and stimulating parturition and lactation (OT)

[1,2]. In the CNS, AVP and OT are expressed in the social brain network [3] in distinct cell populations [4,5] and are associated with opposing roles in behavioral and physiological functions [6–9]. These central AVP and OT projections are involved in social perception, cognition and decision making, and perturbations are associated with psychopathologies including autism spectrum disorder, schizophrenia, anxiety, and depression [6].

Abbreviations: AVP, arginine vasopressin; BK_{Ca}, large conductance calcium-activated potassium channel; BLAST, basic local alignment search tool; BSA, bovine serum albumin; Ca²⁺, calcium; CHO, Chinese hamster ovary; CI, confidence interval; CNS, central nervous system; EC₅₀, half-maximal response; E_{MAX}, maximum response achievable; FMP, FLIPR membrane potential; GIRKs, G-protein-coupled inwardly-rectifying potassium channels; GPCR, G-protein-coupled receptor; hOTR, hOT receptor human oxytocin receptor; IC₅₀, concentration producing 50 % inhibitory response; IK_{Ca}, intermediate conductance calcium-activated potassium channel; Leu⁸-OT, consensus mammalian oxytocin sequence; mOTR, mOT receptor marmoset oxytocin receptor; NCBI, National Center for Biotechnology Information; NWM, new world monkeys; OT, oxytocin; OTR, oxytocin receptor; K⁺, potassium; OWM, old world monkeys; Pro⁸-OT, oxytocin sequence with proline in 8th position; PKC, protein kinase C; PTX, pertussis toxin; qOTR, qOT receptor macaque oxytocin receptor; SERCA, sarco-endoplasmic reticulum calcium ATPase; SK_{Ca}, small conductance calcium-activated potassium channel; Tg, thapsigargin; tOTR, tOT receptor, titi monkey oxytocin receptor

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<https://doi.org/10.1016/j.bioph.2020.109832>

Received 16 September 2019; Received in revised form 5 December 2019; Accepted 13 December 2019

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Evolutionary analysis reveals extreme conservation of AVP and OT in eutherian mammals which differ at amino acids 3 and 8 [10]. Vasopressin and OT are believed to have arisen from a gene duplication event prior to vertebrate divergence, whereas invertebrates generally have only one homolog [7,11]. In New World Monkeys (NWMs) AVP is conserved; however, an unusual level of OT variability has been observed with six distinct ligand variants identified to date [12–15]. OT is highly sensitive to structural modifications; small changes can affect pharmacological profiles [16–18]. Coevolution is observed for both OT and AVP ligands and their receptors [12,13,19] and subtle differences in receptor sequences between human and rat result in important alterations in the selectivity profiles of some ligands [9].

The mammalian OT-vasopressin receptor family is comprised of four G-protein coupled receptors: one canonical OT receptor and three vasopressin receptors (V1a, V1b, V2) [20]. Both OT and V1a receptors are robustly expressed in the brain [21]. Notably, there is ~85 % structural homology between OT and V1a receptors resulting in significant cross-reactivity [2,13,22–24].

The promiscuous activation of multiple G-proteins by OT receptors demonstrates some cell-specific biases [25–30]. Evolutionary factors lead to changes in ligand-receptor structure, which then may have consequences for physiological function and social behavior. Here, we assess natural variation in AVP and OT peptide ligands and OT receptors to assess ligand-receptor signaling cascades in four primate species representing ‘natural experiments’ in genetic variation in the OT receptor that are associated with phenotypic consequences: marmoset (*Callithrix jacchus*, Pro⁸-OT, NWM, socially monogamous), titi monkey (*Callicebus*, Leu⁸-OT, NWM, socially monogamous), macaque (*Macaca*, Leu⁸-OT, OWM, nonmanogamous), and human (*Homo sapiens*, Leu⁸-OT, apes, socially monogamous) [12,13]. Interestingly, binding analysis at these four OTRs demonstrated that Pro⁸-OT bound at modestly higher affinities than AVP or Leu⁸-OT, suggesting that rather than binding properties, downstream signaling pathways are primarily responsible for differences in physiological function and social behaviors [31]. We stably transfected CHO cells expressing marmoset (mOTR), titi monkey (tOTR), macaque (qOTR), and human oxytocin receptor (hOTR) and characterized AVP- and OT-analog G-protein signaling pathways by monitoring Ca²⁺ mobilization and membrane hyperpolarization.

2. Materials and methods

2.1. Chinese hamster ovary (CHO) cell cultures

Transcriptome profiling confirmed that neuronal cell lines ND7/23, F-11 and SH-SY5Y express OT receptor [32]. Likewise, RT-PCR showed neuroblastoma cell lines SK-N-SH, SH-SY5Y, IMR-32 and astrocytoma cell line MOG-G-UVW all express OT receptor [33]. We excluded cell lines that express OT receptors from our studies inasmuch as endogenous expression would contribute to observed ligand-induced signaling. Therefore, we selected CHO cells as a common transfection background since they do not express OT receptors [34]. Wild-type Chinese hamster ovarian-K1 (CHO-K1) cells were purchased from ATCC (CCL-61) and cultured in Ham's F12 (Hyclone SH30026.01), 10 % fetal bovine serum (FBS) (Atlanta Biologicals S11550), 1.5 % HEPES 1 M Solution (Hyclone SH30231.01), 1 % Penicillin-Streptomycin (10,000 U/mL; Life Technologies 15140-163). Human oxytocin receptor (hOTR) expressing CHO-K1 cell lines were purchased from Genscript (M00195). Marmoset oxytocin receptor (mOTR) plasmid was purchased from Genscript and stably-transfected into CHO-K1 cells as described [34] CHO-K1 cells stably transfected with tOTR and qOTR as described [31] and were received from Dr. Myron Toews lab (UNMC). mOTR, tOTR, qOTR and hOTR expressing cells were cultured in Ham's F12 (Hyclone SH30026.01), 10 % FBS (Atlanta Biologicals S11550), 1.5 % HEPES 1 M Solution (Hyclone SH30231.01), 1 % Penicillin-Streptomycin (10,000 U/mL; Life Technologies 15140-163) and 400 mg/mL G418 (RPI Corp. G64000-5.0). Human Kappa-opioid (κ O) receptor expressing CHO cells

(κ OR-CHO) were cultured in RPMI-1640 medium supplemented with 10 % FBS (Atlanta Biologicals S11550). Cells were cultured at 37 °C in 5 % CO₂ and 90 % humidity.

2.2. Drugs

AVP, Leu⁸-OT, and Pro⁸-OT Anaspec 58863 were reconstituted in DMSO Sigma-Aldrich D4540. NS-1619 Sigma-Aldrich N170, Paxilline Sigma-Aldrich P2928, SKA-31 Sigma-Aldrich S5573, thapsigargin Sigma-Aldrich T9033, and TRAM-34 Sigma-Aldrich T6700 were reconstituted in DMSO. Pertussis toxin Sigma-Aldrich P7208 was reconstituted in ultrapure water with 5 mg/mL bovine serum albumin Fisher Scientific BP1600-100. Dynorphin A (1–13) amide (American Peptide 26-4-51A) was dissolved in 25 mM Tris at pH 7.4. Apamin (Sigma-Aldrich A1289) was reconstituted in 0.05 M acetic acid.

2.3. Ca²⁺ mobilization assay

The effects of AVP and OT analog addition on Ca²⁺ mobilization were examined using Fluo3-AM fluorescence (Molecular Probes F1241) monitored with a FLIPR2 plate reader (Molecular Devices, Sunnyvale, CA). FLIPR operates by illuminating the bottom of a 96-well microplate with an air-cooled laser and measuring the fluorescence emissions from cell-permeant dyes in all 96 wells simultaneously using a cooled CCD camera. This instrument is equipped with an automated 96-well pipettor, which can be programmed to deliver precise quantities of solutions simultaneously to all 96 culture wells from two separate 96-well source plates.

Cells were plated at 0.3 million cells/mL in 96-well plates (MidSci P9803) and cultured overnight in culture media at 37 °C in 5 % CO₂ and 95 % humidity. On the day of assay, growth medium was aspirated and replaced with 100 μ l dye-loading medium per well containing 4 μ M Fluo-3 AM and 0.04 % pluronic acid (Molecular Probes P3000MP) in Locke's buffer (8.6 mM HEPES, 5.6 mM KCl, 154 mM NaCl, 5.6 mM glucose, 1.0 mM MgCl₂, 2.3 mM CaCl₂, 0.5 mM probenecid; pH 7.4). Cells were incubated for 1 h at 37 °C in 5 % CO₂ and 95 % humidity and then washed four times in 180 μ l fresh Locke's buffer using an automated microplate washer (Bio-Tek Instruments Inc., VT). Baseline fluorescence was recorded for 60 s, prior to a 20 μ l addition of various concentrations of Leu⁸-OT and Pro⁸-OT. Cells were excited at 488 nm and Ca²⁺-bound Fluo-3 emission was recorded at 538 nm at 2 s intervals for an additional 200 s.

To assess the role of intracellular Ca²⁺ in OT-mediated mobilization of Ca²⁺, the sarco/endoplasmic reticulum calcium ATPase (SERCA) inhibitor thapsigargin was used to deplete intracellular Ca²⁺ stores. Cells were incubated in 100 μ l dye-loading medium per well containing 4 μ M Fluo-3 AM and 0.04 % pluronic acid in Locke's buffer (8.6 mM HEPES, 5.6 mM KCl, 154 mM NaCl, 5.6 mM glucose, 1.0 mM MgCl₂, 2.3 mM CaCl₂, 0.5 mM probenecid; pH 7.4). Cells were incubated at 37 °C in a 5 % CO₂ and 95 % humidity for 60 min prior to washing four times in 180 μ l Locke's buffer and 10 μ l addition of thapsigargin (1 μ M final concentration) and incubated for an additional five minutes. Ca²⁺ mobilization assays were performed as described above.

2.4. Membrane potential assay

To assess changes in membrane potential the FLIPR Membrane Potential Assay (FMP blue; Molecular Probes F1241) was used. Cells were plated at 0.3 million cells/ml in 96-well plates (MidSci P9803) and cultured overnight in culture media at 37 °C in 5 % CO₂ and 95 % humidity. Growth medium was removed and replaced with 190 μ l per well of FMP Blue in Locke's buffer (8.6 mM HEPES, 5.6 mM KCl, 154 mM NaCl, 5.6 mM glucose, 1.0 mM MgCl₂, 2.3 mM CaCl₂ pH 7.4). Cells were incubated at 37 °C in 5 % CO₂ and 95 % humidity for 45 min. Baseline fluorescence was recorded for 60 s, prior to a 10 μ l addition of log concentrations of AVP, Leu⁸-OT and Pro⁸-OT. Cells were excited at

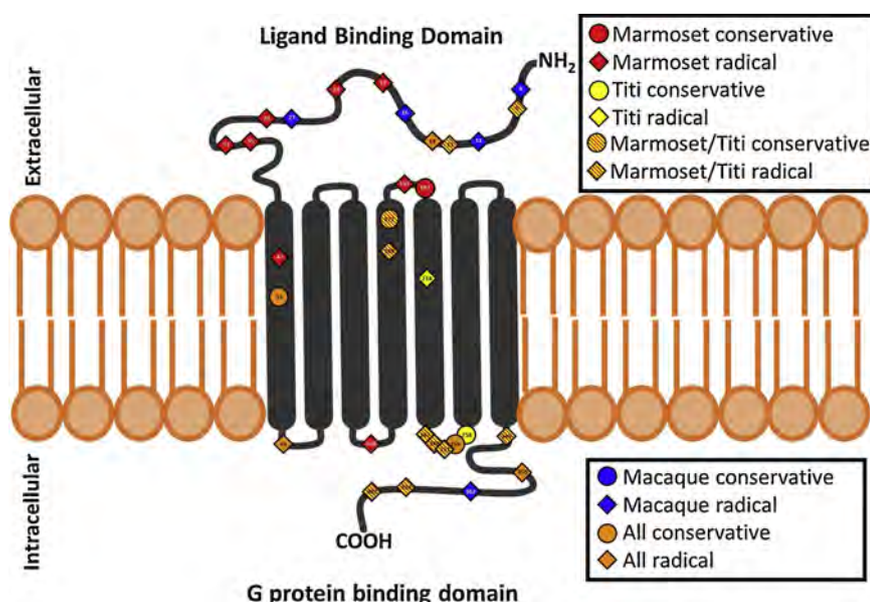


Fig. 1. Comparison of marmoset, titi monkey and macaque OTR amino acid sequence to human OTR. Identification of amino acid substitutions in marmoset (red), titi monkey (yellow), marmoset and titi monkey (red/yellow striped), macaque (blue), or all three (orange) OTRs relative to human OTR. Numbers represent the location of the amino acid substitution. Radical physiochemical substitutions that differ in size, polarity and/or charge are indicated by diamonds and amino acid substitutions that do not differ by these properties are conservative changes indicated by circles (Supplementary Table 1).

530 nm and emission was recorded at 565 nm at 2 s intervals for an additional 200 s.

To assess the role of $G_{i/o}$ in the OT ligand-induced membrane hyperpolarization, cells were incubated overnight with pertussis toxin (PTX) to inactivate $G_{i/o}$ [35]. Cells were plated at 125,000 cells/mL in 96-well plates. PTX (150 ng/ml) was added 24 h after plating and incubated for an additional 24 h. Membrane potential assay was performed as described above. To confirm the influence of PTX on a known $G_{i/o}$ mediated response, the effect of PTX on kappa-opioid receptor (κ OR) mediated hyperpolarization was used as a positive control [36]. κ OR-CHO were used for these experiments. The PTX assays were performed as described above for mOTR- and hOTR-expressing CHO cells, except for stimulation with dynorphin A 1–13 (DynA 1–13) rather than OT analogs.

To assess potential OT ligand-induced membrane hyperpolarization through Ca^{2+} -activated K^+ channels, we tested three inhibitors targeting Ca^{2+} -activated K^+ channel subtypes. G_q -mediated activation of protein kinase-C (PKC) causes an increase in cytosolic Ca^{2+} [37] with attendant activation of Ca^{2+} sensitive K^+ channels. Ca^{2+} -activated K^+ channels are separated into three subtypes of large (BK_{Ca}), intermediate (IK_{Ca}), and small conductance (SK_{Ca}) channels [38]. Paxilline is a selective inhibitor of the BK_{Ca} channel [39]. TRAM-34 is a selective inhibitor of the IK_{Ca} channel, $K_{Ca3.1}$ [40,41]. Apamin is a selective inhibitor of SK_{Ca} channels [42,43]. Cells were incubated with FMP in Locke's buffer at 37 °C in a 5 % CO_2 and 95 % humidity for 35 min prior to a 10 μ l addition of paxilline, TRAM-34, and/or apamin. Cells were incubated for an additional 10 min after drug addition. Membrane potential assays were performed as described above.

NS-1619 is a BK_{Ca} channel activator [44,45]. If changes in Ca^{2+} are responsible for activation of the BK_{Ca} , the response should be NS-1619 sensitive. Cells were incubated at 37 °C in a 5 % CO_2 and 95 % humidity for 35 min prior to a 10 μ l addition of paxilline. Cells were incubated for an additional 10 min after paxilline addition. Membrane potential assays were performed as described above, with the exception of stimulation with NS-1619 rather than OT analogs.

SKA-31 is an activator of IK_{Ca} channel $K_{Ca3.1}$ [46,47]. If changes in Ca^{2+} are responsible for the activation of $K_{Ca3.1}$, the response should be SKA-31 sensitive. Cells were incubated at 37 °C in a 5 % CO_2 and 95 % humidity for 35 min prior to a 10 μ l addition of TRAM-34. Cells were incubated for an additional 10 min after TRAM-34 addition. Membrane potential assays were performed as described above, with the exception of stimulation with SKA-31 rather than OT analogs.

To assess the role of intracellular Ca^{2+} in AVP and OT ligand-

induced changes in membrane potential, thapsigargin was used. Cells were incubated with FMP in Locke's buffer at 37 °C in a 5 % CO_2 and 95 % humidity for 40 min prior to a 10 μ l addition of thapsigargin. Cells were incubated for an additional 5 min after drug addition.

2.5. Data analysis

All concentration-response data were analyzed and graphs generated using GraphPad Prism (San Diego, CA, U.S.A.) software. Fluo-3 relative fluorescence changes were plotted as $F_{MAX} - F_0$. F_{MAX} is the maximum fluorescence achieved by the ligand. F_0 is the average of the baseline fluorescence reading for 30 images (60 s). Similarly, FMP relative fluorescence changes were plotted as $F_{MIN} - F_0$. F_{MIN} is the minimum fluorescence achieved by the ligand (indicative of hyperpolarization) and F_0 is the average of the baseline fluorescence reading for 30 images (60 s). EC_{50}/IC_{50} and E_{MAX} values for AVP- and OT peptide-stimulated increases in Fluo-3 fluorescence or decreases in FMP Blue fluorescence were determined by nonlinear least-squares fitting of a logistic equation to the peptide concentration versus fluorescence area under the curve data. The 95 % confidence intervals for all EC_{50}/IC_{50} and E_{MAX} were used to assess differences in potency/efficacy. R^2 was used to assess goodness of fit. Standard error bars have been included for all graphs; however, if the error bar is smaller than the size of the symbol, GraphPad Prism does not draw it.

3. Results

3.1. Comparison of marmoset, titi monkey and macaque OTR amino acid sequence to human OTR

Significant coevolution has been shown to exist between OT ligands and receptors in NWMs [12,13]. Marmoset, titi monkey, macaque and human OT receptor amino acid sequences were accessed from the National Center for Biotechnology Information (NCBI) [12]. Alignment using the NCBI basic local alignment search tool (BLAST) [48] indicates that human and macaque (*Macaca mulatta*) OT receptor are 98 % conserved; human and titi monkey (*Calicebus cupreus*) OT receptor 96 % conserved; and human and marmoset (*Callithrix jacchus*) 94 % conserved. Physiochemical changes to marmoset, titi monkey and macaque OT receptor sequences were classified as radical when the amino acid substitution differed by size, polarity and/or charge, and conservative if the substitution did not differ by these properties (Fig. 1; Supplementary Table 1). Radical amino acid substitutions were primarily observed

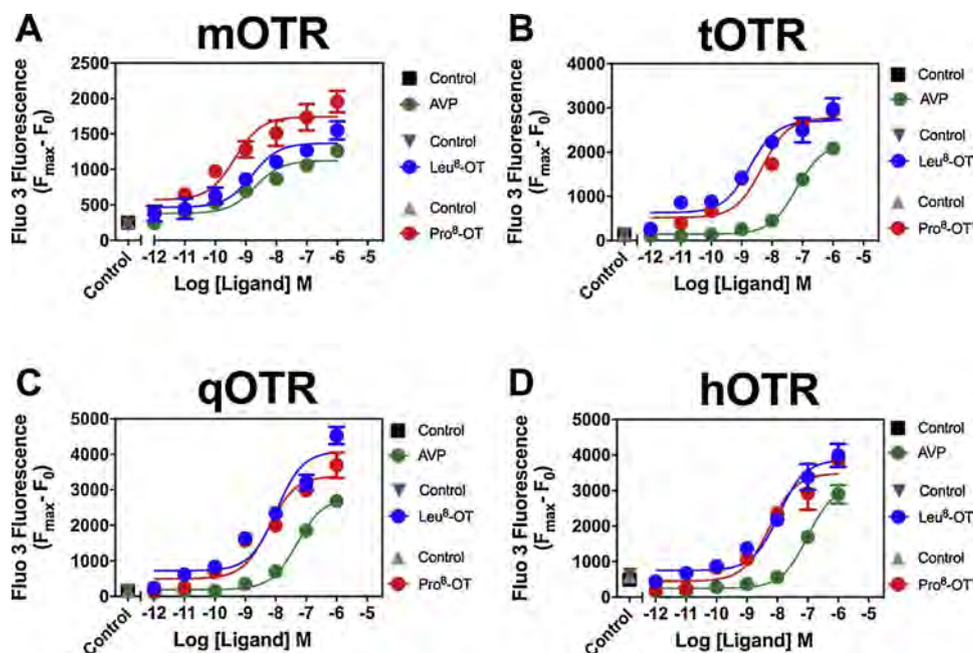


Fig. 2. AVP, Leu⁸-OT and Pro⁸-OT induced calcium mobilization in mOTR-, tOTR-, qOTR-, or hOTR- expressing CHO cells. AVP, Leu⁸-OT, and Pro⁸-OT concentration-response relationships in mOTR cells (A), in tOTR cells (B), in qOTR cells (C), and in hOTR cells (D). AVP, Leu⁸-OT and Pro⁸-OT analogs were run in parallel on the same plates, at the same time, with the same split of cells. N = 3 experiments (3–4 replicates per dose per experiment). Time-response (Supplementary Fig. 1). Statistical analysis (Table 1).

in the in the *N-terminus*, extracellular regions, and transmembrane regions that may play a role in ligand binding. In intracellular regions and *C-terminus* region, radical substitutions may affect G-protein coupling and downstream signaling. Radical changes were more numerous for mOTR with Pro⁸-OT as its endogenous ligand, than for tOT or qOT receptors. Amino acids 14, 51, 69, 255, and 355, in mOT, tOT and qOT receptors all differed from hOT receptor. Together, the expression of these receptors in CHO cells using a common cellular background allowed us to examine how this natural variation in ligands and receptors impacts receptor activation and cellular signaling.

3.2. AVP and OT analogs induce G_q-mediated intracellular Ca²⁺ mobilization

G_q activates the phospholipase Cβ and inositol 3-phosphate signaling pathway [27] resulting in an increase in intracellular Ca²⁺. To compare AVP and OT analog activation of OTR-coupled G_q, we performed functional assays using the Ca²⁺ indicator dye Fluo3-AM. At the mOT receptor, whose endogenous ligand is Pro⁸-OT, AVP demonstrated a similar potency (EC₅₀) to Pro⁸- and Leu⁸-OT, with the 95 % confidence intervals (95 % CI) overlapping (Fig. 2A; Table 1; Supplementary Fig. 1A-C). However, the endogenous Pro⁸-OT was more

Table 1
AVP and OT-analog induced intracellular calcium mobilization.

		Ligand (nM)		
Line	Parameter	AVP	Leu ⁸ -OT	Pro ⁸ -OT
mOTR	EC ₅₀	2.00	1.61	0.44
	95 % CI	0.63 to 6.32	0.61 to 5.13	0.14 to 1.40
	R ²	0.74	0.69	0.70
tOTR	EC ₅₀	63.26	1.87	4.68
	95 % CI	42.69 to 93.75	0.72 to 4.84	2.05 to 10.72
	R ²	0.98	0.89	0.91
qOTR	EC ₅₀	49.78	10.61	5.95
	95 % CI	36.78 to 67.38	4.57 to 24.19	2.41 to 14.67
	R ²	0.99	0.90	0.89
hOTR	EC ₅₀	98.63	11.02	5.58
	95 % CI	62.23 to 153.00	5.18 to 23.48	2.45 to 12.72
	R ²	0.97	0.92	0.91

Potency of AVP, Leu⁸-OT and Pro⁸-OT at inducing calcium mobilization in mOTR, tOTR, qOTR and hOTR CHO cells. N = 3 experiments (3–4 replicates per dose per experiment). Sigmoidal curves (Fig. 2).

efficacious at 1742 relative fluorescence units (RFUs) (95 % CI 1573–1912), whereas AVP was 1124 RFUs (95 % CI 1000–1248) and Leu⁸-OT 1372 RFUs (95 % CI 1220–1523). In the three primate receptors whose endogenous ligand is Leu⁸-OT (tOT, qOT, and hOT receptors), AVP was less potent and efficacious than either OT analog (Fig. 2B-D; Table 1; Supplementary Fig. 1D-L). In tOT receptor cell lines, AVP was 33X less potent than Leu⁸-OT and 13X less potent than Pro⁸-OT (Table 1). Leu⁸-OT 2710 RFUs (95 % CI 2424–2996) displayed equivalent efficacy with Pro⁸-OT 2779 RFUs (95 % CI 2485–3047), whereas AVP was less efficacious 2198 RFUs (95 % CI 2014–2382). In qOT receptor cell lines, AVP was 5X less potent than Leu⁸-OT and 8X less potent than Pro⁸-OT (Table 1). Likewise, Leu⁸-OT 4089 RFUs (95 % CI 3585–4594) displayed similar efficacy to Pro⁸-OT 3386 RFUs (95 % CI 2955–3816), but was more efficacious than AVP 2775 RFUs (95 % CI 2608 to 2942). In hOT receptor cell lines, AVP was 9X less potent than Leu⁸-OT and 16X less potent than Pro⁸-OT (Table 1). Leu⁸-OT 3867 RFUs (95 % CI 3473–4293) displayed similar efficacy to Pro⁸-OT 3483 RFUs (95 % CI 3075–3890), but was more efficacious than AVP 3150 RFUs (95 % CI 2787–3441). AVP, Leu⁸-OT, and Pro⁸-OT did not elevate intracellular Ca²⁺ in non-transfected CHO-K1 cells (Supplementary Fig. 2). These data demonstrate that the dose-dependent responses observed in OTR-expressing lines is attributable to OTR expression.

Sarco/endoplasmic reticulum calcium ATPase (SERCA) maintains the Ca²⁺ gradient between the cytosol and endoplasmic reticulum, and therefore SERCA pump inhibition results in depletion of intracellular Ca²⁺ stores [49,50]. To confirm the role of intracellular Ca²⁺ stores in AVP- and OT-analog mediated Ca²⁺ influx, cells were pretreated with thapsigargin, a potent SERCA inhibitor. We pretreated all four lines with thapsigargin and evaluated AVP, Leu⁸-OT and Pro⁸-OT induced Ca²⁺ influx. In control assays AVP and OT analogs produced concentration-dependent increases in Ca²⁺; however, pretreatment with thapsigargin blocked AVP and OT-analog evoked increases in Ca²⁺ (Fig. 3; Supplementary Fig. 3) [34] demonstrating the dependence on intracellular Ca²⁺ stores.

3.3. AVP and OT analog-induced changes in membrane potential are dependent on g_q mediated Ca²⁺ mobilization

To assess the role of AVP and OT-induced membrane hyperpolarization, we performed functional assays using FMP blue, a membrane potential-sensitive dye that displays increased fluorescence in response

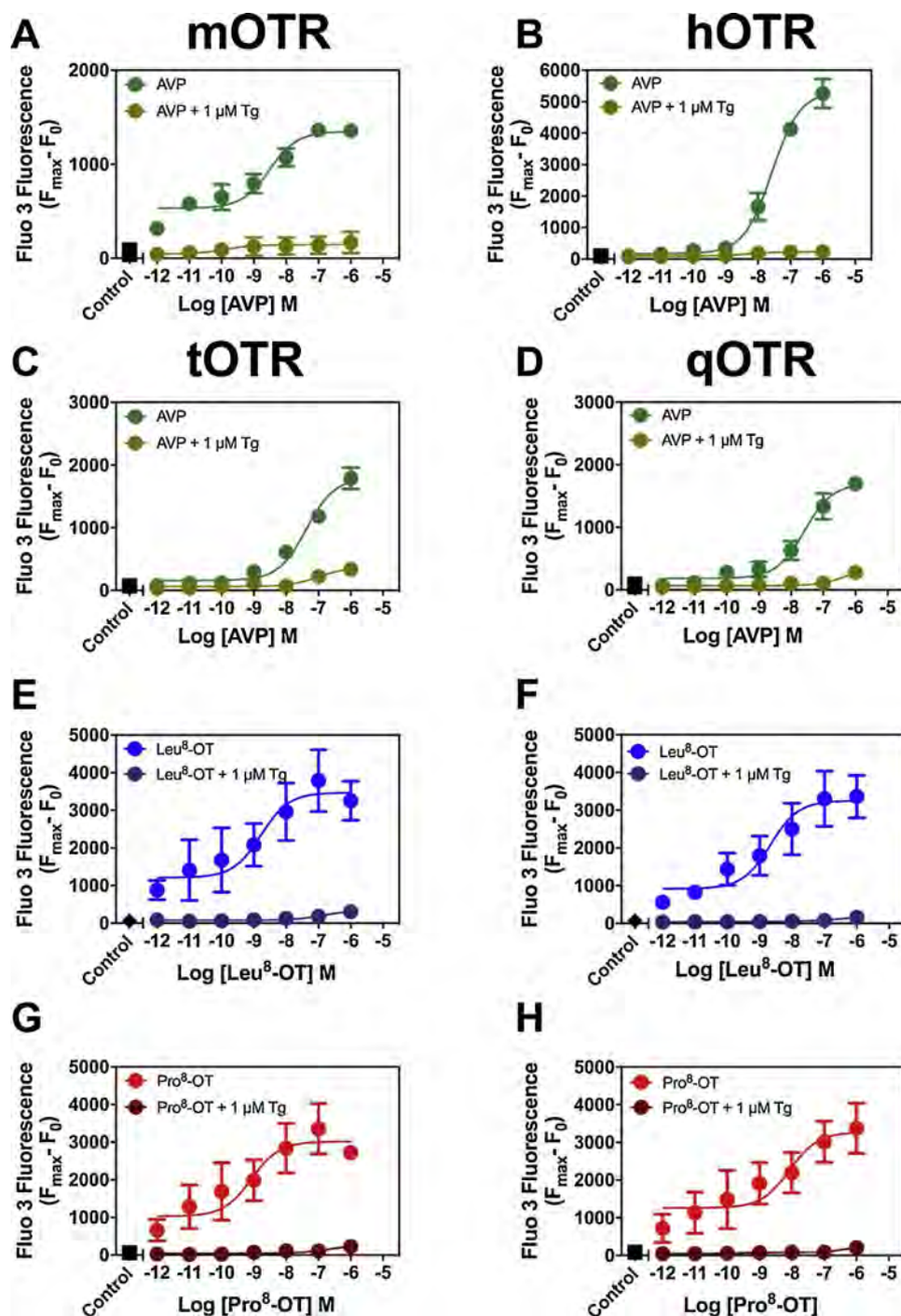


Fig. 3. Effects of pretreatment with thapsigargin (Tg) on AVP, Leu⁸-OT and Pro⁸-OT induced changes on intracellular Ca²⁺ mobilization in mOTR-, hOTR-, tOTR-, and qOTR-expressing CHO cells. Control AVP and Tg-pretreated concentration response relationships in mOTR-expressing cells (A), in hOTR-expressing cells (B), in tOTR-expressing cells (C), in qOTR-expressing cells (D). Leu⁸-OT and Tg-pretreated concentration response relationships in tOTR-expressing cells (E) and qOTR-expressing cells (F). Pro⁸-OT and Tg-pretreated concentration response relationships in tOTR-expressing cells (G) and qOTR-expressing cells (H). Control and Tg-pretreated replicates were run in parallel on the same plates, at the same time and with the same split of cells. N = 3 experiments (five replicates per dose per experiment). Time response (Supplementary Fig. 3).

to depolarization and decreased fluorescence in response to membrane hyperpolarization [51,52]. Membrane hyperpolarization can be influenced by numerous ion channels, including Gi/o activation of GIRK channels and Gq activation of calcium-activated potassium channels [34]. Thus, potential ligand bias [53], and the potential for activation of multiple pathways suggest these data may be different from the Fluo-3 assays. In cell lines for all four OT receptors, AVP and both OT-analogs produced concentration-dependent decreases in FMP blue fluorescence consistent with a hyperpolarizing response (Fig. 4; Table 2; Supplementary Fig. 4). In mOT receptor cell lines, the 95 % confidence intervals for potency for all three ligands overlapped (Fig. 4A; Table 2; Supplementary Fig. 4A-C). However, Leu⁸-OT 6185 (95 % CI 5479–6891) and Pro⁸-OT 5207 RFUs (95 % CI 4862–5553), were more efficacious than AVP 4264 RFUs (95 % CI 3869–4659). In tOT and hOT

receptor cell lines, Leu⁸-OT was 4x more potent than Pro⁸-OT, and Pro⁸-OT was more potent than AVP by ~10X and ~4X, respectively (Fig. 4B,D; Table 2; Supplementary Fig. 4D-F, J-L). In tOT receptor expressing cells, Leu⁸-OT 5477 (95 % CI 5149–5805) and Pro⁸-OT 5347 RFUs (95 % CI 5094–5600), were more efficacious than AVP 4803 RFUs (95 % CI 4575–5031). In hOT receptor expressing cells, there was no substantial difference in efficacy with AVP 5049 RFU (95 % CI 4817–5281), Leu⁸-OT 5064 (95 % CI 4787–5341) and Pro⁸-OT 5303 RFUs (95 % CI 4970–5635). In qOT receptor cell lines, no significant difference was observed between Pro⁸-OT and Leu⁸-OT, but Leu⁸-OT was 30X more potent than AVP (Fig. 4C; Table 2; Supplementary Fig. 4G-I). However, the 95 % CI of efficacies for all three ligands overlapped, with AVP 2033 RFU (95 % CI 1310–2755), Leu⁸-OT 2268 (95 % CI 1827–2709) and Pro⁸-OT 1878 RFUs (95 % CI

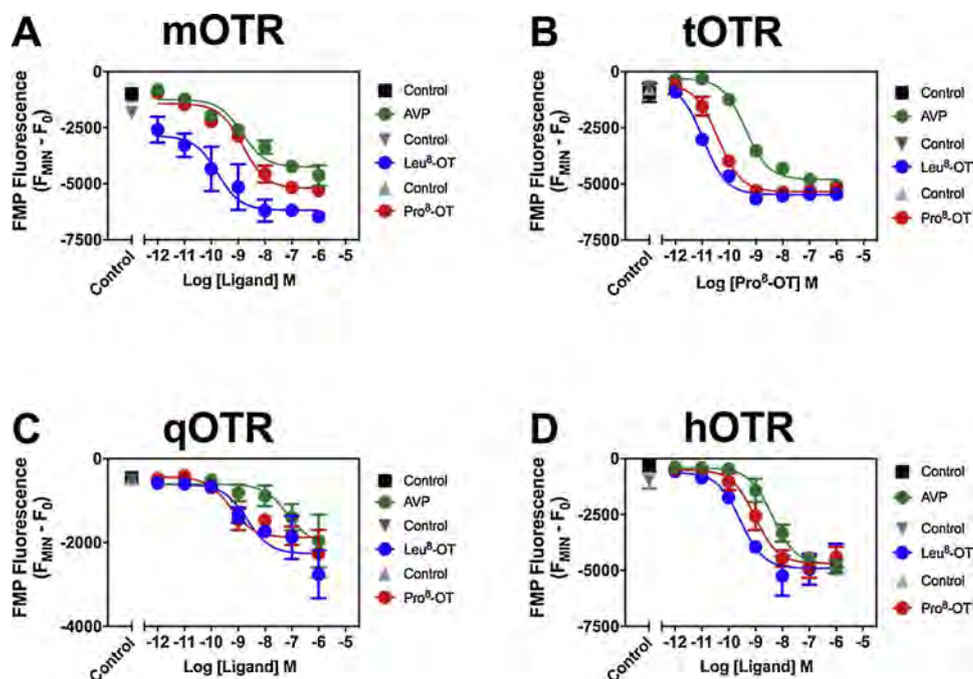


Fig. 4. AVP, Leu⁸-OT and Pro⁸-OT induced changes in membrane potential in mOTR-, tOTR-, qOTR-, or hOTR- expressing CHO cells. AVP, Leu⁸-OT, and Pro⁸-OT concentration-response relationships in mOTR cells (A), in tOTR cells (B), in qOTR cells (C), in hOTR cells (D). AVP, Leu⁸-OT and Pro⁸-OT analogs were run in parallel on the same plates, at the same time, with the same split of cells. N = 3 experiments (3–4 replicates per dose per experiment). Time response (Supplementary Fig. 4). Statistical analysis (Table 2).

Table 2
AVP and OT-analog induced membrane hyperpolarization.

		Ligand		
Line	Parameter	AVP	Leu ⁸ -OT	Pro ⁸ -OT
mOTR	IC ₅₀	1.41 nM	0.17 nM	1.49 nM
	95 % CI	0.57 to 3.50	0.03 to 1.03	0.79 to 2.81
	R ²	0.81	0.53	0.9
tOTR	IC ₅₀	0.4 nM	10.83 pM	38.60 pM
	95 % CI	0.27 to 0.61	5.08 to 23.09	22.56 to 66.05
	R ²	0.97	0.98	0.91
qOTR	IC ₅₀	59.46 nM	1.83 nM	0.53 nM
	95 % CI	8.17 to 432.60	0.36 to 9.27	0.14 to 2.09
	R ²	0.49	0.58	0.68
hOTR	IC ₅₀	4.20 nM	0.27 nM	0.95 nM
	95 % CI	2.24 to 8.03	0.09 to 0.89	0.40 to 2.20
	R ²	0.90	0.70	0.82

Potency of AVP, Leu⁸-OT and Pro⁸-OT at inducing membrane hyperpolarization in mOTR, tOTR, qOTR and hOTR CHO cells. N = 3 experiments (3–4 replicates per dose per experiment). Sigmoidal curves (Fig. 3).

1586–2170). The absence of effects on membrane potential in untransfected CHO-K1 cells demonstrated the requirement for OTR transfection in the observed hyperpolarizing response to AVP (Supplementary Fig. 5), Leu⁸-OT and Pro⁸-OT [34]. Together, AVP is more potent at OTRs from marmoset with Pro⁸-OT as its endogenous ligand, than species whose cognate ligand is Leu⁸-OT.

Pertussis toxin catalyzes ADP-ribosylation of the G $\alpha_{i/o}$ subunit [54], locking G $\alpha_{i/o}$ in its GDP-bound inactive state and preventing the pertussis toxin-sensitive G-protein from interacting with a GPCR. To assess the role of G $\alpha_{i/o}$ in AVP and OT-analog induced hyperpolarization, we pretreated cells with pertussis toxin. In mOT receptor CHO cells we previously demonstrated Leu⁸-OT mediated hyperpolarization was partially sensitive to pertussis toxin, suggesting both G $\alpha_{i/o}$ -mediated and pertussis toxin-insensitive pathways contribute to the hyperpolarizing response [34]. Pertussis toxin treatment did not inhibit AVP induced hyperpolarization in any of the four cell lines (Fig. 5A-D; Supplementary Fig. 6A-H; Supplementary Table 2), Leu⁸-OT induced hyperpolarization in tOT or qOT receptor cell lines (Fig. 5E-F; Supplementary Fig. 6I-L, Supplementary Table 2), or Pro⁸-OT induced hyperpolarization in tOT or qOT receptor cell lines (Fig. 5G-H; Supplementary Fig. 6M-P, Supplementary Table 2), suggesting that G $\alpha_{i/o}$ does not

contribute to the hyperpolarizing response for these ligand-receptor combinations [34]. Given that the kappa-opioid (κ O) receptor couples to G α_i , we used a κ OR expressing CHO cell line (κ OR-CHO) as a positive control to demonstrate pertussis toxin's ability to disrupt G α_i . DynorphinA1–13-NH₂, a κ O receptor agonist, produced a robust hyperpolarizing response in control κ O receptor cell lines, that was abolished in cells pretreated with pertussis toxin (Supplementary Figure 7). These data support our conclusion that AVP and OT analog-induced hyperpolarization in OTR-expressing cells do not involve the pertussis toxin-sensitive G-proteins, G $\alpha_{i/o}$.

OT analog induced hyperpolarization involves mOT and hOT receptors coupling to G α_q activating the G α_q /phosphoinositide-phospholipase C pathway and Ca²⁺-dependent K⁺ channel activation [34]. To explore the role of Ca²⁺-activated K⁺ channels in AVP-induced changes in membrane potential, as well as extending our data with OT analogs in tOT and qOT receptors, we used a pharmacological approach with inhibitors that discriminate between subtypes of Ca²⁺-activated K⁺ channels. There are three subtypes of Ca²⁺-activated K⁺ channels, including small conductance (SK_{Ca}), intermediate conductance (IK_{Ca}) and large conductance (BK_{Ca}) channels [38]. To assess the role of SK_{Ca} channels in AVP and OT-mediated membrane hyperpolarization cells were pretreated with the SK_{Ca}-selective blocker apamin, which blocks SK_{Ca} channels through an allosteric mechanism [43]. In mOT and hOT receptor CHO cells, we previously observed no major contribution ($\leq 15\%$) on Leu⁸-OT and Pro⁸-OT induced membrane hyperpolarization after pretreatment with apamin [34]. Similarly, in all four cell lines, apamin did not significantly affect AVP-induced changes in membrane potential (Fig. 6A-D; Supplementary Figures 8A–B, 9A, 10A; Supplementary Table 3). Likewise, in tOT and qOT receptor CHO cells, we observed no substantial effects ($\leq 15\%$) of pretreatment with apamin on Leu⁸-OT or Pro⁸-OT induced membrane hyperpolarization (Fig. 6E-H; Supplementary Figures 9B-C, 10B-C; Supplementary Table 3). Together, these data suggest that SK_{Ca} channels provide little or no contribution to AVP and OT mediated changes in membrane potential across these four primate OTRs.

To assess the roles of BK_{Ca} channels in AVP and OT-analog mediated changes in membrane potential, cells were pretreated with the BK_{Ca} blocker paxilline, which stabilizes the channel in its closed conformation [55]. In mOT receptor CHO cells, paxilline produced a moderate inhibition of AVP induced hyperpolarization, resulting in a 20 %

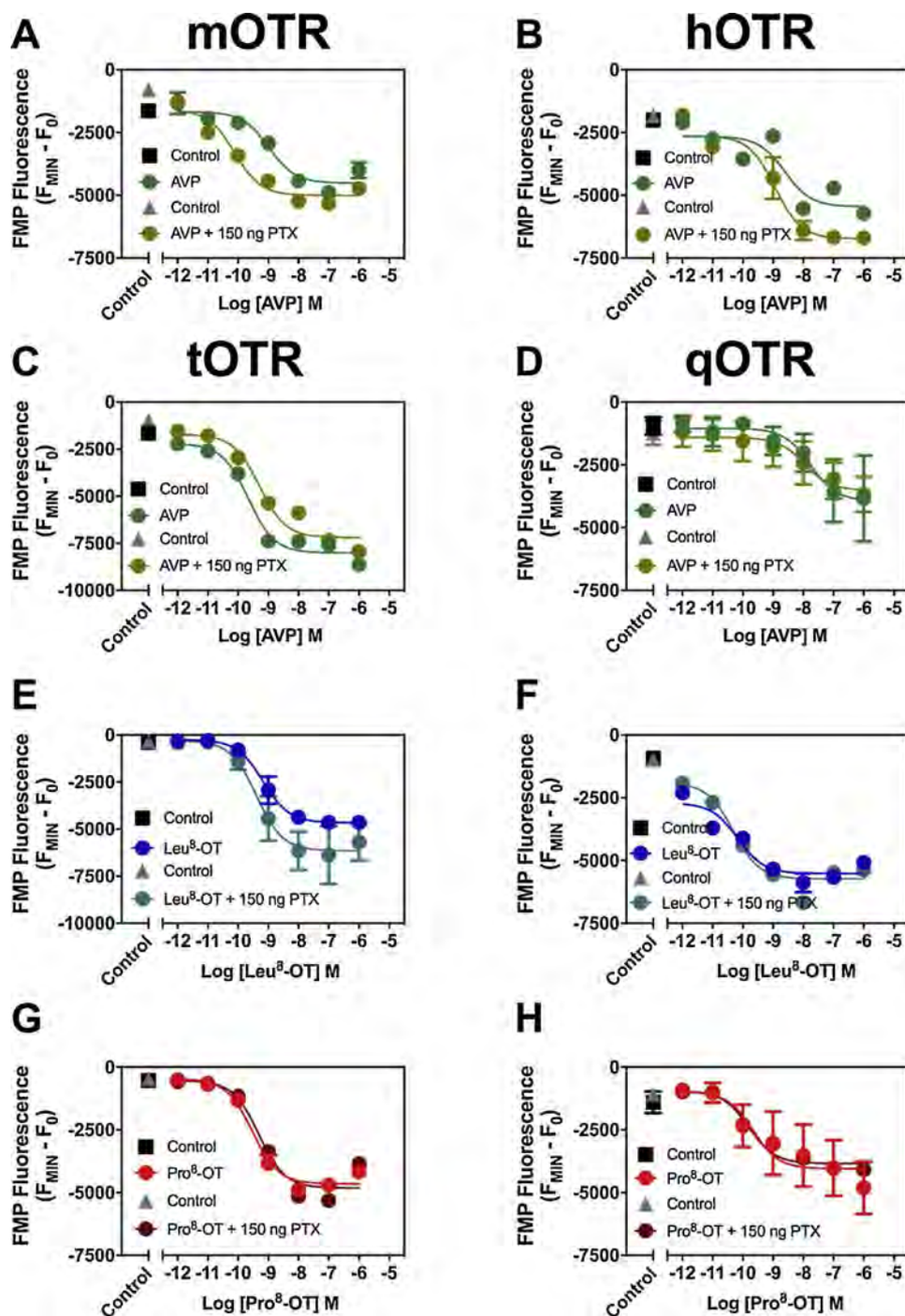


Fig. 5. Lack of effect for pretreatment with PTX on AVP, Leu⁸-OT and Pro⁸-OT induced changes in membrane potential in mOTR-, tOTR-, qOTR- and hOTR-expressing CHO cells. Control AVP and PTX-pretreated concentration response relationships in mOTR-expressing cells (A), in hOTR-expressing cells (B), in tOTR-expressing cells (C), in qOTR-expressing cells (D). Leu⁸-OT and PTX-pretreated concentration response relationships in tOTR-expressing cells (E) and qOTR-expressing cells (F). Pro⁸-OT and PTX-pretreated concentration response relationships in tOTR-expressing cells (G) and qOTR-expressing cells (H). Control and PTX-pretreated replicates were run in parallel on the same plates, at the same time and with the same split of cells. N = 3 experiments (five replicates per dose per experiment). Time-response (Supplementary Fig. 6). Statistical analysis (Supplementary Table 2).

inhibition (Fig. 6A; Supplementary Figure 8C, Supplementary Table 3) and produced greater inhibition (52 %) in hOT receptor cells (Fig. 6B; Supplementary Figure 8D, Supplementary Table 3). In tOT receptor CHO cells, paxilline produced minimal inhibition in AVP and moderate inhibition in Leu⁸-OT and Pro⁸-OT induced changes in membrane potential, resulting in a 13 %, 38 % and 39 % reduction of the hyperpolarizing response, respectively (Fig. 6C,E,G; Supplementary Figure 9D-F; Supplementary Table 3). In qOT receptor CHO cells, paxilline produced a moderate inhibition in AVP and Pro⁸-OT induced changes in membrane potential and substantial inhibition in Leu⁸-OT, resulting in a 20 %, 35 % and 46 % inhibition, respectively (Fig. 6D,F,H; Supplementary Figure 10D-F; Supplementary Table 3). Together, these data suggest that BK_{Ca} channels contribute to a fraction of the AVP and OT-analog induced changes in membrane potential. In mOT and hOT

receptor CHO cells, the BK_{Ca} activator NS-1619 is inhibited with paxilline [34]. Here we challenged tOT and qOT receptor CHO cells with NS-1619, and the evoked hyperpolarization was blocked with 30 μ M paxilline by 80 % and 72 %, respectively (Supplementary Figure 11 A–B,E–F; Supplementary Table 4) supporting a role for BK_{Ca} channels in the regulation of CHO cell membrane potential.

To assess the role of IK_{Ca} channels in AVP and OT-mediated membrane hyperpolarization, cells were pretreated with TRAM-34, an IK_{Ca} channel blocker that specifically blocks K_{Ca}3.1 by occupying the K⁺ binding site [40]. TRAM-34 produced the most effective inhibition in AVP and OT-analog mediated changes in membrane potential, with AVP less impacted than OT analogs in three OT receptor species (mOT, tOT, and hOT receptor). In mOT receptor CHO cells, TRAM-34 produced a moderate inhibition in AVP-induced membrane potential (33

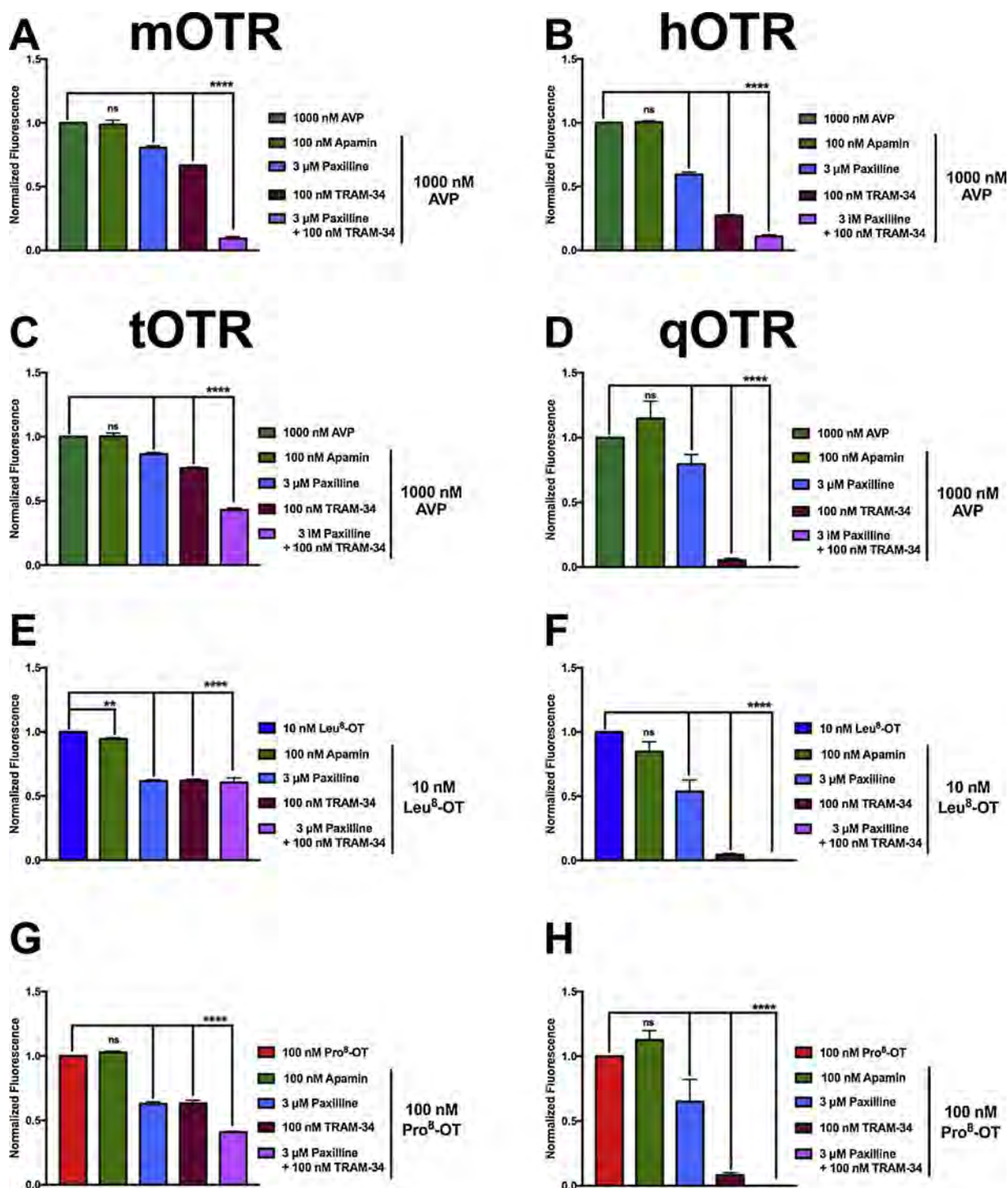


Fig. 6. Effects of pretreatment with Ca^{2+} -activated K^{+} inhibitors on AVP and/or $\text{Leu}^8\text{-OT}$ or $\text{Pro}^8\text{-OT}$ induced changes in membrane potential in mOTR-, hOTR-, tOTR-, and qOTR-expressing CHO cells. Inhibitor fluorescence was normalized to AVP (A–D), $\text{Leu}^8\text{-OT}$ (E–F), or $\text{Pro}^8\text{-OT}$ (G–H) membrane hyperpolarization. Time response (Supplementary Figures 8–10). Area under the curve (negative peaks only) were assessed and a one-way Anova was performed with Sidak's multiple comparisons to determine statistical significance. $N = 3$ experiments for each inhibitor/analog/cell line combination (10 replicates per dose per experiment). Adjusted p-values (Supplementary Table 3).

%; Fig. 6A; Supplementary Figure 8E; Supplementary Table 3). In hOT receptor CHO cells, TRAM-34 produced a more robust inhibition of AVP-induced membrane potential (61 %; Fig. 6B; Supplementary Figure 8 F; Supplementary Table 3). In tOT receptor CHO cells, TRAM-34 produced a moderate inhibition in AVP-induced membrane potential (25 %), whereas $\text{Leu}^8\text{-OT}$ and $\text{Pro}^8\text{-OT}$ were inhibited somewhat greater, 38 % and 37 %, respectively (Fig. 6C,E,G; Supplementary Figure 9G-I; Supplementary Table 3). In qOT receptor CHO cells, AVP,

$\text{Leu}^8\text{-OT}$ and $\text{Pro}^8\text{-OT}$ were robustly inhibited, ~98 % each (Fig. 6D,F,H; Supplementary Figure 10G-I; Supplementary Table 3). Involvement of $\text{K}_{\text{Ca}3.1}$ in hyperpolarization can be tested with the activator SKA-31. We challenged tOT and qOT receptor CHO cells with SKA-31 and showed that the hyperpolarization was inhibited by 300 nM TRAM-34 pretreatment by 78 % and 92 %, respectively (Supplementary Figure 11C-D, G-H; Supplementary Table 4). These results support the role for the $\text{K}_{\text{Ca}3.1}$ channel in the regulation of membrane

potential by AVP and OT-ligands.

To determine the combined contribution of BK_{Ca} and K_{Ca}3.1 channels in AVP and OT-mediated changes in membrane potential, cells were pretreated with both paxilline and TRAM-34. In mOT receptor CHO cells, the combined exposure of paxilline and TRAM-34 inhibited AVP-induced membrane hyperpolarization by 91 % (Fig. 6A; Supplementary Figure 8 G; Supplementary Table 3). In hOT receptor CHO cells, the combined exposure of paxilline and TRAM-34 inhibited AVP-induced hyperpolarization by 87 % (Fig. 6B; Supplementary Figure 8 H; Supplementary Table 3). In tOT receptor CHO cells, paxilline and TRAM-34 combined inhibited AVP, Leu⁸-OT and Pro⁸-OT by 57 %, 40 % and 60 % (Fig. 6C,E,G; Supplementary Figure 9J-L; Supplementary Table 3). In qOT receptor CHO cells, the combination completely inhibited AVP, Leu⁸-OT and Pro⁸-OT induced hyperpolarization (Fig. 6D,F,H; Supplementary Figure 10J-L; Supplementary Table 3). Together, these data indicated an additive effect of inhibition of BK_{Ca} and K_{Ca}3.1 channels demonstrating that they are primarily responsible for AVP and OT-analog induced membrane hyperpolarization in CHO cells expressing these OTRs.

To confirm the role of intracellular Ca²⁺ stores in AVP- and OT analog mediated membrane hyperpolarization, cells were pretreated with the SERCA inhibitor thapsigargin to [49,50]. As expected, pretreatment with thapsigargin eliminated the hyperpolarization produced by AVP, Leu⁸-OT, and Pro⁸-OT in at all four OT receptor cell lines (Supplementary Figure 12) [34]. Together, these data demonstrated that intracellular Ca²⁺ stores are the source of Ca²⁺ responsible for AVP and OT-analog engagement of K⁺ channels. Interestingly, in thapsigargin pretreated tOT and hO receptor CHO cells challenged with Leu⁸-OT, and in hOT receptor cells challenged with Pro⁸-OT a depolarizing response was observed (Supplementary Figure 12E) [34] indicating a possible dual modulation of K⁺ channel currents by OT analogs through these OT receptors.

4. Discussion

The molecular structure of OT-AVP family peptides and receptors, as well as a facilitation of physiological responses and social behaviors related to stress and anxiety are highly conserved in evolution [2,7,8,11,13]. One mechanism to assess genetic conservation is quantification via the ratios of nonsynonymous nucleotide substitutions relative to synonymous substitutions (dN/dS), where ratios less than 1.0 imply stabilizing selection and greater than 1.0 selection for diverse sequences. Analysis of dN/dS ratios for AVP is 0.005 and OT is 0.009, suggesting extreme conservation for both peptides [10]. Comparison of OT analogs across NWM suggests positions 2 and 8 are most variable [56], with a single amino acid substitution at the 8th position causing in a Leu > Pro change resulting in a rigid turn in the peptide backbone [12,15]. Moreover, genetic changes in ligand structure resulted in a significantly higher proportion of corresponding changes in the OT receptor sequences of NWMs, particularly at the N-terminus which is involved in ligand binding [12], as is evident in the greater number of mOT receptor substitutions (red) with Pro⁸-OT as its endogenous ligand as compared to tOT receptor (yellow) or qOT receptor (blue) substitutions that have the consensus mammalian Leu⁸-OT as their endogenous neuropeptide (Fig. 1). These changes likely affect the overall three dimensional architecture of the receptor, each of which may contribute to differences in cellular signaling. The natural variability within OT-AVP family in NWMs provides a unique opportunity to assess these signaling cascades in relation to ligand-receptor coevolution [12,57], particularly in relation to social behaviors. For example, social monogamy is exceptionally high in NWMs and phylogeny correlated statistical comparisons demonstrate it corresponds with ligand-receptor coevolution [12], whereas social monogamy is rare among mammals in general [13]. Both OT analogs have been shown to modulate social behavior in marmosets, with the cognate ligand Pro⁸-OT more effective than Leu⁸-OT in relation to sexual fidelity and reduced prosocial behaviors

towards strangers [58,59].

At the cellular level, ligand and receptor structure, as well as cellular context, appear to influence how GPCRs activate downstream signaling cascades. These signaling cascades result in changes in cell function, influencing physiological and behavioral effects at the organismal level [60]. A recent study assessed the binding properties for these ligands at the OTRs and found that the rank order affinity was Pro⁸-OT > Leu⁸-OT > AVP for receptors from all four species [31]. We recognize the importance of cellular context in GPCR signaling and understand that signaling profiles in the heterologous expression system (CHO cells) that we selected may differ from those of intact neurons. However, primary neuronal cultures from expressing the OT receptors from the species we are evaluating are not currently available. Established human neuroblastoma and glioma cell lines reportedly express the endogenous hOT receptor [33], although these cells do not successfully recapitulate the sensitivity of primary neurons in culture [61] and therefore signaling profiles may also differ from intact primary neuronal cultures. Thus, we believe that these initial *in vitro* experiments to define the signaling profiles of AVP and OT variants are a fundamental first step to understanding ligand-receptor function.

In this study we compared the pharmacological signatures of AVP, Leu⁸-OT, and Pro⁸-OT in CHO cells expressing marmoset, titi monkey, macaque, or human OT receptors. Our results confirm that all three ligands activated G_q signaling in concentration-dependent increases in intracellular Ca²⁺ through all four OT receptors. Interestingly, at OT receptors with Leu⁸-OT as their endogenous ligand (titi monkey, macaque, human) AVP was less potent and efficacious than OT analogs. In contrast, at the mOT receptor, with Pro⁸-OT is the cognate ligand, AVP potency comparable to OT analogs. Notably, Pro⁸-OT was more efficacious than AVP or Leu⁸-OT, suggesting evolutionary differences in receptor structure may play a role in ligand potency and efficacy at the OTR [12,13]. We confirmed the role of intracellular Ca²⁺ in AVP and OT analog-induced Ca²⁺ mobilization at all four OT receptors by depleting intracellular Ca²⁺ with thapsigargin, which inhibited the response to all three ligands [34].

Given the promiscuous coupling of OTRs to various G-proteins [28,29], we next assessed the ability of AVP, Leu⁸-OT, and Pro⁸-OT to trigger a hyperpolarizing response at the four OT receptors. AVP was less potent than either OT analog at the three OT receptors whose cognate ligand is Leu⁸-OT. However, at the mOT receptor, no significant differences were observed in the potency of AVP and Pro⁸-OT. Leu⁸-OT was more potent at the mOT, tOT and hOT receptors, whereas Pro⁸-OT was appeared slightly more potent at the qOT receptor. Notably, AVP was less efficacious at mOT and tOT receptors (both NWMs), whereas no differences in efficacy were observed for the three ligands at qOT receptor (an OWM) or hOT receptor, suggesting common nucleotide substitutions in the in mOT and tOT receptor may contribute to observed differences in ligand efficacy.

GPCR coupling to G_{i/o} stimulates G-protein-gated inward rectifying K⁺ channels via the Gβγ subunit [62]. Pertussis toxin inhibits G_{i/o} from coupling to the GPCR [63] thus blocking G_{i/o} protein mediated contribution to the hyperpolarizing response. Our previous data demonstrated that G_{i/o} is responsible for a minor portion of the Leu⁸-OT induced hyperpolarizing response at the mOT receptor, suggesting a dual modulation by a pertussis toxin-sensitive and pertussis toxin-insensitive G-proteins (Pierce et al. 2019). The Leu⁸-OT induced hyperpolarizing response was pertussis toxin-insensitive in OTRs from all three species whose cognate ligand is Leu⁸-OT. Both AVP and Pro⁸-OT are pertussis toxin-insensitive at all four OT receptors, suggesting G_{i/o} coupling does not contribute to their hyperpolarizing response. Since these *in vitro* analyses all utilize the same CHO-K1 cell line, cellular context is unlikely to be a contributing factor.

Since AVP and OT-ligand induced membrane hyperpolarization are largely pertussis toxin-insensitive, we assessed the contribution of Ca²⁺-activated K⁺ channels in AVP and OT-analog mediated hyperpolarization using channel blockers specific for SK_{Ca}, BK_{Ca} (K_{Ca}1.1),

and IK_{Ca} ($K_{Ca3.1}$). Apamin selectively blocks SK_{Ca} channels, and pretreatment with this inhibitor resulted in minimal or no inhibition ($\leq 15\%$) of AVP, Leu^8 -OT and Pro^8 -OT at the mOT, tOT, qOT and hOT receptors. These data demonstrate that SK_{Ca} channels provide minimal contribution to AVP and OT-analog induced membrane hyperpolarization. Paxilline is a selective blocker of BK_{Ca} channels, which contributes to neuronal excitability [64]. Pretreatment with paxilline resulted in partial inhibition for AVP and OT analogs in cells expressing each of the four OT receptors. This is consistent with previous reports that Leu^8 -OT mediated hyperpolarization through OT receptor G_q /phosphoinositide-phospholipase C pathway activation of BK_{Ca} channels [34,65]. Paxilline also inhibited the hyperpolarizing response to the BK_{Ca} channel opener NS-1619 in mOT, hOT [34], tOT and qOT receptor CHO cells, further supporting the role for BK_{Ca} channels in membrane hyperpolarization. TRAM-34 selectively inhibits a IK_{Ca} channel ($K_{Ca3.1}$) [40], and also regulates neuronal excitability [66]. Pretreatment with TRAM-34 produced the most robust inhibition of the hyperpolarizing response to all three ligands in cell lines expressing each of the four OT receptors. TRAM-34 also inhibited membrane the hyperpolarizing response to $K_{Ca3.1}$ channel opener SKA31 in mOT, hOT [34], tOT and qOT receptor CHO cells, demonstrating the involvement of $K_{Ca3.1}$ in membrane hyperpolarization. Combined pretreatment with paxilline and TRAM-34 suggested that OT receptor G_q /phosphoinositide-phospholipase C pathway activation of BK_{Ca} and the IK_{Ca} channel $K_{Ca3.1}$ were additive. Depletion of intracellular Ca^{2+} with SERCA inhibitor thapsigargin confirmed the role of intracellular Ca^{2+} stores in AVP and OT-ligand induced membrane hyperpolarization in the OT receptor- G_q /phosphoinositide-phospholipase C pathway activation of Ca^{2+} dependent K^+ channels.

The OT-AVP family of neuropeptides and receptors plays a fundamental role in mediating physiological processes and social behaviors [67–70], with perturbations associated with various social and behavioral deficits [6]. These sociobehavioral functions prompt interest in OT as a potential therapeutic mediator in conditions such as autism spectrum disorder [71], depression and anxiety [6,72], schizophrenia [73,74], and post-traumatic stress disorder [75]. Clinically, OT is used peripherally to induce labor [76] and prevent postpartum hemorrhage [77], which provide further therapeutic potential for OT analogs. A major challenge is connecting ligand-induced receptor activation at the cellular level to changes in organismal physiological and behavioral functions [60]. The present results show that AVP, Leu^8 -OT and Pro^8 -OT display functionally distinct responses at the various OTRs examined. Some of these differences correspond to cognate ligand and receptor structure, such as OT-analogs increased potency at inducing membrane hyperpolarization and intracellular Ca^{2+} mobilization compared to AVP at the OTRs from species whose cognate ligand is Leu^8 -OT. The unique diversity of OT ligands and receptors in NWMs provide a natural experiment whereby we can begin to parse out how pharmacological profiles are affected by ligand and receptor structure, and how these correlate with specific behavioral outcomes, providing insight for the advancement of OT-mediated therapeutics.

5. Author contributions

Participated in research design: Murray, Pierce

Conducted experiments: Pierce, Mustoe

Performed data analysis: Pierce

Wrote or contributed to the writing of the manuscript: Pierce, French, Murray

Declaration of Competing Interest

None.

Aknowledgments

This work was supported by the National Institutes of Health Eunice Kennedy Shriver National Institute of Child Health and Human Development [Grant R01HD089147]. We thank Dr. Myron Toews, Nancy Schulte and Dr. Jack Taylor for providing the stably transfected qOTR and tOTR cell lines. We thank Dr. Jeffrey French and Dr. Aaryn Mustoe for providing the AVP, Leu^8 -OT and Pro^8 -OT peptides. We thank Dr. Aaryn Mustoe for helping with some of the experiments. We thank Dr. Suneet Mehrotra for his thoughtful discussions and Ms. Bridget Sefranek for her careful reading of the manuscript.

Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:<https://doi.org/10.1016/j.biopha.2020.109832>.

References

- [1] M. Ludwig, G. Leng, Dendritic peptide release and peptide-dependent behaviours, *Nat. Rev. Neurosci.* 7 (2) (2006) 126–136.
- [2] B. Jurek, I.D. Neumann, The oxytocin receptor: from intracellular signaling to behavior, *Physiol. Rev.* 98 (3) (2018) 1805–1908.
- [3] S.W. Newman, The medial extended amygdala in male reproductive behavior. A node in the mammalian social behavior network, *Ann. N. Y. Acad. Sci.* 877 (1999) 242–257.
- [4] D. Huber, P. Veinante, R. Stoop, Vasopressin and oxytocin excite distinct neuronal populations in the central amygdala, *Science* 308 (5719) (2005) 245–258.
- [5] H. Gainer, Cell-type specific expression of oxytocin and vasopressin genes: an experimental odyssey, *J. Neuroendocrinol.* 24 (4) (2012) 528–538.
- [6] I.D. Neumann, R. Landgraf, Balance of brain oxytocin and vasopressin: implications for anxiety, depression, and social behaviors, *Trends Neurosci.* 35 (11) (2012) 649–659.
- [7] R. Stoop, Neuromodulation by oxytocin and vasopressin, *Neuron* 76 (1) (2012) 142–159.
- [8] R. Stoop, Neuromodulation by oxytocin and vasopressin in the central nervous system as a basis for their rapid behavioral effects, *Curr. Opin. Neurobiol.* 29 (2014) 187–193.
- [9] B. Chini, M. Manning, Agonist selectivity in the oxytocin/vasopressin receptor family: new insights and challenges, *Biochem. Soc. Trans.* 35 (Pt 4) (2007) 737–741.
- [10] M. Wallis, Molecular evolution of the neurohypophysial hormone precursors in mammals: comparative genomics reveals novel mammalian oxytocin and vasopressin analogues, *Gen. Comp. Endocrinol.* 179 (2) (2012) 313–318.
- [11] Z.R. Donaldson, L.J. Young, Oxytocin, vasopressin, and the neurogenetics of sociality, *Science* 322 (5903) (2008) 900–904.
- [12] D. Ren, G. Lu, H. Moriyama, A.C. Mustoe, E.B. Harrison, J.A. French, Genetic diversity in oxytocin ligands and receptors in New World monkeys, *PLoS One* 10 (5) (2015) e0125775.
- [13] J.A. French, J.H. Taylor, A.C. Mustoe, J. Cavanaugh, Neuropeptide diversity and the regulation of social behavior in New World primates, *Front. Neuroendocrinol.* 42 (2016) 18–39.
- [14] P. Vargas-Pinilla, V.R. Paixao-Cortes, P. Pare, L. Tovo-Rodrigues, C.M. Vieira, A. Xavier, D. Comas, A. Pissinatti, M. Sinigaglia, M.M. Rigo, G.F. Vieira, A.B. Lucion, F.M. Salzano, M.C. Bortolini, Evolutionary pattern in the OXT-OXTR system in primates: coevolution and positive selection footprints, *Proc. Natl. Acad. Sci. U. S. A.* 112 (1) (2015) 88–93.
- [15] A.G. Lee, D.R. Cool, W.C. Grunwald Jr., D.E. Neal, C.L. Buckmaster, M.Y. Cheng, S.A. Hyde, D.M. Lyons, K.J. Parker, A novel form of oxytocin in New World monkeys, *Biol. Lett.* 7 (4) (2011) 584–587.
- [16] R. Beard, A. Stucki, M. Schmitt, G. Py, C. Grundschober, A.D. Gee, E.W. Tate, Building bridges for highly selective, potent and stable oxytocin and vasopressin analogs, *Bioorg. Med. Chem.* 26 (11) (2018) 3039–3045.
- [17] S. Scibola, G.H. Goetz, G. Bai, B.N. Rogers, D.L. Gray, A. Duplantier, K.R. Fonseca, M.A. Vanase-Frawley, N.M. Kablaoui, Systematic N-methylation of oxytocin: impact on pharmacology and intramolecular hydrogen bonding network, *Bioorg. Med. Chem.* 24 (16) (2016) 3513–3520.
- [18] M. Muttenthaler, A. Andersson, A.D. de Araujo, Z. Dekan, R.J. Lewis, P.F. Alewood, Modulating oxytocin activity and plasma stability by disulfide bond engineering, *J. Med. Chem.* 53 (24) (2010) 8585–8596.
- [19] J. Koehbach, C.W. Gruber, From ethnopharmacology to drug design, *Commun. Integr. Biol.* 6 (6) (2013) e27583.
- [20] H.H. Zingg, S.A. Laporte, The oxytocin receptor, *Trends Endocrinol. Metab.* 14 (5) (2003) 222–227.
- [21] Q.J. Pittman, S.J. Spencer, Neurohypophysial peptides: gatekeepers in the amygdala, *Trends Endocrinol. Metab.* 16 (8) (2005) 343–344.
- [22] Z. Song, H.E. Albers, Cross-talk among oxytocin and arginine-vasopressin receptors: relevance for basic and clinical studies of the brain and periphery, *Front. Neuroendocrinol.* (2017).
- [23] G. Gimpl, F. Fahrenholz, The oxytocin receptor system: structure, function, and regulation, *Physiol. Rev.* 81 (2) (2001) 629–683.

- [24] M. Manning, A. Misicka, A. Olma, K. Bankowski, S. Stoev, B. Chini, T. Durroux, B. Mouillac, M. Corbani, G. Guillon, Oxytocin and vasopressin agonists and antagonists as research tools and potential therapeutics, *J. Neuroendocrinol.* 24 (4) (2012) 609–628.
- [25] M. Ragenbass, Overview of cellular electrophysiological actions of vasopressin, *Eur. J. Pharmacol.* 583 (2–3) (2008) 243–254.
- [26] A. Reversi, V. Rimoldi, T. Marrocco, P. Cassoni, G. Bussolati, M. Parenti, B. Chini, The oxytocin receptor antagonist atosiban inhibits cell growth via a “biased agonist” mechanism, *J. Biol. Chem.* 280 (16) (2005) 16311–16318.
- [27] A. Reversi, P. Cassoni, B. Chini, Oxytocin receptor signaling in myoepithelial and cancer cells, *J. Mammary Gland Biol. Neoplasia* 10 (3) (2005) 221–229.
- [28] M. Busnelli, A. Sauliere, M. Manning, M. Bouvier, C. Gales, B. Chini, Functional selective oxytocin-derived agonists discriminate between individual G protein family subtypes, *J. Biol. Chem.* 287 (6) (2012) 3617–3629.
- [29] M. Gravati, M. Busnelli, E. Bulgheroni, A. Reversi, P. Spaiardi, M. Parenti, M. Toselli, B. Chini, Dual modulation of inward rectifier potassium currents in olfactory neuronal cells by promiscuous G protein coupling of the oxytocin receptor, *J. Neurochem.* 114 (5) (2010) 1424–1435.
- [30] M. Kolaj, L.P. Renaud, Vasopressin acting at V1-type receptors produces membrane depolarization in neonatal rat spinal lateral column neurons, *Prog. Brain Res.* 119 (1998) 275–284.
- [31] J.H. Taylor, N.A. Schulte, J.A. French, M.L. Toews, Binding characteristics of two oxytocin variants and vasopressin at oxytocin receptors from four primate species with different social behavior patterns, *J. Pharmacol. Exp. Ther.* (2018).
- [32] K. Yin, G.J. Baillie, I. Vetter, Neuronal cell lines as model dorsal root ganglion neurons: a transcriptomic comparison, *Mol. Pain* 12 (2016).
- [33] P. Cassoni, A. Sapino, A. Stella, N. Fortunati, G. Bussolati, Presence and significance of oxytocin receptors in human neuroblastomas and glial tumors, *Int. J. Cancer* 77 (5) (1998) 695–700.
- [34] M.L. Pierce, S. Mehrotra, A.C. Mustoe, J.A. French, T.F. Murray, A comparison of the ability of leu(8)- and pro(8)-Oxytocin to regulate intracellular Ca(2+) and Ca(2+)-Activated K(+) channels at human and marmoset oxytocin receptors, *Mol. Pharmacol.* 95 (4) (2019) 376–385.
- [35] X.B. Zhou, S. Lutz, F. Steffens, M. Korth, T. Wieland, Oxytocin receptors differentially signal via Gq and Gi proteins in pregnant and nonpregnant rat uterine myocytes: implications for myometrial contractility, *Mol. Endocrinol.* 21 (3) (2007) 740–752.
- [36] K.S. Murthy, G.M. Makhlof, Opioid mu, delta, and kappa receptor-induced activation of phospholipase C-beta 3 and inhibition of adenylyl cyclase is mediated by Gi2 and G(o) in smooth muscle, *Mol. Pharmacol.* 50 (4) (1996) 870–877.
- [37] S.L. Ritter, R.A. Hall, Fine-tuning of GPCR activity by receptor-interacting proteins, *Nat. Rev. Mol. Cell Biol.* 10 (12) (2009) 819–830.
- [38] C. Vergara, R. Latorre, N.V. Marrion, J.P. Adelman, Calcium-activated potassium channels, *Curr. Opin. Neurobiol.* 8 (3) (1998) 321–329.
- [39] M. Sanchez, O.B. McManus, Paxilline inhibition of the alpha-subunit of the high-conductance calcium-activated potassium channel, *Neuropharmacology* 35 (7) (1996) 963–968.
- [40] H.M. Nguyen, V. Singh, B. Pressly, D.P. Jenkins, H. Wulff, V. Yarov-Yarovsky, Structural insights into the atomistic mechanisms of action of small molecule inhibitors targeting the KCa3.1 channel pore, *Mol. Pharmacol.* 91 (4) (2017) 392–402.
- [41] R.G. Staal, T. Khayrullina, H. Zhang, S. Davis, S.M. Fallon, M. Cajina, M.E. Nattini, A. Hu, H. Zhou, S.B. Poda, S. Zorn, G. Chandrasena, E. Dale, B. Campbell, L.C. Biilmann Ronn, G. Munro, T. Mller, Inhibition of the potassium channel KCa3.1 by senicapoc reverses tactile allodynia in rats with peripheral nerve injury, *Eur. J. Pharmacol.* 795 (2017) 1–7.
- [42] A.L. Blatz, K.L. Magleby, Single apamin-blocked Ca-activated K+ channels of small conductance in cultured rat skeletal muscle, *Nature* 323 (6090) (1986) 718–720.
- [43] C. Lamy, S.J. Goodchild, K.L. Weatherall, D.E. Jane, J.F. Liegeois, V. Seutin, N.V. Marrion, Allosteric block of KCa2 channels by apamin, *J. Biol. Chem.* 285 (35) (2010) 27067–27077.
- [44] K. Lee, I.C. Rowe, M.L. Ashford, NS 1619 activates BKCa channel activity in rat cortical neurones, *Eur. J. Pharmacol.* 280 (2) (1995) 215–219.
- [45] G. Edwards, A. Niederste-Hollenberg, J. Schneider, T. Noack, A.H. Weston, Ion channel modulation by NS 1619, the putative BKCa channel opener, in vascular smooth muscle, *Br. J. Pharmacol.* 113 (4) (1994) 1538–1547.
- [46] A. Sankaranarayanan, G. Raman, C. Busch, T. Schultz, P.I. Zimin, J. Hoyer, R. Kohler, H. Wulff, Naphtho[1,2-d]thiazol-2-ylamine (SKA-31), a new activator of KCa2 and KCa3.1 potassium channels, potentiates the endothelium-derived hyperpolarizing factor response and lowers blood pressure, *Mol. Pharmacol.* 75 (2) (2009) 281–295.
- [47] P. Christophersen, H. Wulff, Pharmacological gating modulation of small- and intermediate-conductance Ca(2+)-activated K(+) channels (KCa2.x and KCa3.1), *Channels Austin (Austin)* 9 (6) (2015) 336–343.
- [48] G.M. Boratyn, A.A. Schaffer, R. Agarwala, S.F. Altschul, D.J. Lipman, T.L. Madden, Domain enhanced lookup time accelerated BLAST, *Biol. Direct* 7 (2012) 12.
- [49] S.M. Dravid, T.F. Murray, Spontaneous synchronized calcium oscillations in neocortical neurons in the presence of physiological [Mg(2+)]_i: involvement of AMPA/kainate and metabotropic glutamate receptors, *Brain Res.* 1006 (1) (2004) 8–17.
- [50] N.T. Quynh Doan, S.B. Christensen, Thapsigargin, origin, chemistry, structure-activity relationships and prodrug development, *Curr. Pharm. Des.* 21 (38) (2015) 5501–5517.
- [51] D.F. Baxter, M. Kirk, A.F. Garcia, A. Raimondi, M.H. Holmqvist, K.K. Flint, D. Bojanic, P.S. Distefano, R. Curtis, Y. Xie, A novel membrane potential-sensitive fluorescent dye improves cell-based assays for ion channels, *J. Biomol. Screen.* 7 (1) (2002) 79–85.
- [52] K.L. Whiteaker, S.M. Gopalakrishnan, D. Groebe, C.C. Shieh, U. Warrior, D.J. Burns, M.J. Coghlan, V.E. Scott, M. Gopalakrishnan, Validation of FLIPR membrane potential dye for high throughput screening of potassium channel modulators, *J. Biomol. Screen.* 6 (5) (2001) 305–312.
- [53] L.M. Luttrell, S. Maudsley, L.M. Bohn, Fulfilling the promise of “Biased” g protein-coupled receptor agonism, *Mol. Pharmacol.* 88 (3) (2015) 579–588.
- [54] T.F. Murray, J.F. Siebenaller, Differential susceptibility of guanine nucleotide-binding proteins to pertussis toxin-catalyzed ADP-ribosylation in brain membranes of two congeneric marine fishes, *Biol. Bull.* 185 (3) (1993) 346–354.
- [55] Y. Zhou, C.J. Lingle, Paxilline inhibits BK channels by an almost exclusively closed-channel block mechanism, *J. Gen. Physiol.* 144 (5) (2014) 415–440.
- [56] A. Mustoe, J.H. Taylor, J.A. French, Oxytocin structure and function in new world monkeys: from pharmacology to behavior, *Integr. Zool.* (2018).
- [57] D. Ren, K.R. Chin, J.A. French, Molecular variation in AVP and AVPR1a in New World monkeys (Primates, platyrrhini): evolution and implications for social monogamy, *PLoS One* 9 (10) (2014) e111638.
- [58] J. Cavanaugh, A.C. Mustoe, J.H. Taylor, J.A. French, Oxytocin facilitates fidelity in well-established marmoset pairs by reducing sociosexual behavior toward opposite-sex strangers, *Psychoneuroendocrinology* 49 (2014) 1–10.
- [59] A.C. Mustoe, J. Cavanaugh, A.M. Harnisch, B.E. Thompson, J.A. French, Do marmosets care to share? Oxytocin treatment reduces prosocial behavior toward strangers, *Horm. Behav.* 71 (2015) 83–90.
- [60] L.M. Luttrell, S. Maudsley, D. Gesty-Palmer, Translating in vitro ligand bias into in vivo efficacy, *Cell. Signal.* 41 (2018) 46–55.
- [61] K.T. LePage, R.W. Dickey, W.H. Gerwick, E.L. Jester, T.F. Murray, On the use of neuro-2a neuroblastoma cells versus intact neurons in primary culture for neurotoxicity studies, *Crit. Rev. Neurobiol.* 17 (1) (2005) 27–50.
- [62] R.A. Rifkin, S.J. Moss, P.A. Slesinger, G protein-gated potassium channels: a link to drug addiction, *Trends Pharmacol. Sci.* 38 (4) (2017) 378–392.
- [63] T.F.S. Murray, J.F. Siebenaller, Differential Susceptibility of Guanine Nucleotide-binding Proteins to Pertussis Toxin-catalyzed ADP-ribosylation in Brain Membranes of Two Congeneric Marine Fishes, *Bio. Bull.* 185 (1993) 346–354.
- [64] J. Kang, J.R. Huguenard, D.A. Prince, Development of BK channels in neocortical pyramidal neurons, *J. Neurophysiol.* 76 (1) (1996) 188–198.
- [65] T. Che, H. Sun, J. Li, X. Yu, D. Zhu, B. Xue, K. Liu, M. Zhang, W. Kunze, C. Liu, Oxytocin hyperpolarizes cultured duodenum myenteric intrinsic primary afferent neurons by opening BK(Ca) channels through IP(3) pathway, *J. Neurochem.* 121 (4) (2012) 516–525.
- [66] R.W. Turner, M. Kruskic, M. Teves, T. Scheidl-Yee, S. Hameed, G.W. Zamponi, Neuronal expression of the intermediate conductance calcium-activated potassium channel KCa3.1 in the mammalian central nervous system, *Pflugers Arch.* 467 (2) (2015) 311–328.
- [67] B.J. Crespi, Oxytocin, testosterone, and human social cognition, *Biol. Rev. Camb. Philos. Soc.* 91 (2) (2016) 390–408.
- [68] G. Missig, L.W. Ayers, J. Schulkin, J.B. Rosen, Oxytocin reduces background anxiety in a fear-potentiated startle paradigm, *Neuropsychopharmacology* 35 (13) (2010) 2607–2616.
- [69] T. Baumgartner, M. Heinrichs, A. Vonlanthen, U. Fischbacher, E. Fehr, Oxytocin shapes the neural circuitry of trust and trust adaptation in humans, *Neuron* 58 (4) (2008) 639–650.
- [70] J. Cavanaugh, S.B. Carp, C.M. Rock, J.A. French, Oxytocin modulates behavioral and physiological responses to a stressor in marmoset monkeys, *Psychoneuroendocrinology* 66 (2016) 22–30.
- [71] E. Anagnostou, L. Soorya, W. Chaplin, J. Bartz, D. Halpern, S. Wasserman, A.T. Wang, L. Pepa, N. Tanel, A. Kushki, E. Hollander, Intranasal oxytocin versus placebo in the treatment of adults with autism spectrum disorders: a randomized controlled trial, *Mol. Autism* 3 (1) (2012) 16.
- [72] H. Ji, W. Su, R. Zhou, J. Feng, Y. Lin, Y. Zhang, X. Wang, X. Chen, J. Li, Intranasal oxytocin administration improves depression-like behaviors in adult rats that experienced neonatal maternal deprivation, *Behav. Pharmacol.* 27 (8) (2016) 689–696.
- [73] M. Brambilla, M. Cotelli, R. Manenti, J. Dagan, D. Sisti, M. Rocchi, M. Balestrieri, S. Pini, S. Raimondi, F.M. Saviotti, P. Scocco, G. de Girolamo, Oxytocin to modulate emotional processing in schizophrenia: a randomized, double-blind, cross-over clinical trial, *Eur. Neuropsychopharmacol.* 26 (10) (2016) 1619–1628.
- [74] C.A. Pedersen, C.M. Gibson, S.W. Rau, K. Salimi, K.L. Smedley, R.L. Casey, J. Leserman, L.F. Jarskog, D.L. Penn, Intranasal oxytocin reduces psychotic symptoms and improves Theory of Mind and social perception in schizophrenia, *Schizophr. Res.* 132 (1) (2011) 50–53.
- [75] J.L. Frijling, Preventing PTSD with oxytocin: effects of oxytocin administration on fear neurocircuitry and PTSD symptom development in recently trauma-exposed individuals, *Eur. J. Psychotraumatol.* 8 (1) (2017) 1302652.
- [76] O.A. Viteri, B.M. Sibai, Challenges and limitations of clinical trials on labor induction: a review of the literature, *AJP Rep.* 8 (4) (2018) e365–e378.
- [77] F.J. Theunissen, L. Chinery, Y.V. Pujar, Current research on carbocetin and implications for prevention of postpartum haemorrhage, *Reprod. Health* 15 (Suppl 1) (2018) 94.