Molecular characterization of eluxadoline as a potential ligand targeting mu-delta opioid receptor heteromers

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Abstract

Eluxadoline, an orally active mixed \(\mu\) opioid receptor (\(\mu\)OR) agonist \(\delta\) opioid receptor (\(\delta\)OR) antagonist developed for the treatment of diarrhea-predominant irritable bowel syndrome, normalizes gastrointestinal (GI) transit and defecation under conditions of novel environment stress or post-inflammatory altered GI function. Furthermore, compared to loperamide, which is used to treat non-specific diarrhea, the effects of eluxadoline on GI transit occur over a wider dosage range. However, the mechanisms of action of eluxadoline are unclear. In this study, we compared the ability of eluxadoline and loperamide to activate G-protein- and \(\beta\)-arrestin-mediated signaling at \(\mu\)OR homomers or \(\mu\)OR-\(\delta\)OR heteromers in heterologous cells. We also examined the ability of both compounds to reduce castor oil induced diarrhea in wild type (WT) and mice lacking \(\delta\)OR. We find that eluxadoline is more potent than loperamide in eliciting G-protein activity and \(\beta\)-arrestin recruitment in \(\mu\)OR expressing cells. However, in cells expressing \(\mu\)OR-\(\delta\)OR heteromers, the potency of eluxadoline is higher, but its maximal effect is lower than that of loperamide. Moreover, in these cells the signaling mediated by eluxadoline but not loperamide is reduced by \(\mu\)OR-\(\delta\)OR heteromer-selective antibodies. We find that in castor oil-induced diarrhea eluxadoline is more efficacious compared to loperamide in WT mice, and \(\delta\)OR appears to play a role in this process. Taken together these results indicate that eluxadoline behaves as a potent \(\mu\)OR agonist in the absence of \(\delta\)OR, while in the presence of \(\delta\)OR eluxadoline’s effects are mediated through the \(\mu\)OR-\(\delta\)OR heteromer.

Keywords

\(\mu\)OR-\(\delta\)OR heteromer; Anti-diarrhea; Eluxadoline; Loperamide; Irritable bowel syndrome

1. Introduction

Opioid receptors are therapeutic targets for the treatment of pain. Morphine, the prototypic opioid, targets the \(\mu\) opioid receptor (\(\mu\)OR) and is clinically preferred for the treatment of
chronic pain [1]. However, chronic morphine administration leads to a number of side-effects including development of analgesic tolerance and constipation. Studies seeking to decrease the side-effects associated with chronic morphine use found that delta opioid receptor (δOR) antagonists could enhance morphine-induced analgesia while preventing the development of tolerance to this drug [2-6] which suggested interactions between μOR and δOR. These interactions were examined using cells heterologously expressing either μOR or δOR or a combination of both receptors and showed that δOR selective antagonists, irrespective of their nature (peptidic or non-peptidic), could enhance μOR selective ligand binding and signaling only in cells co-expressing both receptors [7,8]. Moreover, these in vitro studies showed that the δOR antagonist decreased the dissociation rate of radioligand bound to μOR [9]. These data supported the idea that the δOR antagonist allosterically enhances μOR ligand binding leading to potentiation of μOR-mediated signaling and antinociception. One way in which allosteric modulation of μOR properties by δOR could occur is via the formation of μOR-δOR heteromers; μOR-δOR heteromerization is supported by studies using antibodies that selectively target the heteromer [10] or TAT peptides that can disrupt the formation of μOR-δOR heteromers [11]. Ligands targeting μOR-δOR heteromers either by having μOR agonist/δOR antagonist activity such as bivalent ligands or ligands possessing mixed μOR agonist and δOR antagonist activity have been generated [12-17]. Studies using a bivalent ligand comprising of a μOR agonistic pharmacophore separated by a 21-atom spacer arm from a δOR antagonistic pharmacophore (MDAN21) [15,17] showed that it exhibited 100-times higher antinociceptive potency compared to morphine without significant development of tolerance or dependence [15]. Similarly, studies using ligands possessing mixed μOR agonist/δOR antagonist activity show that their chronic administration leads to lesser side-effects compared to morphine [13]. Taken together these results suggest that targeting the μOR-δOR heteromer could lead to the development of drugs that are likely to have lower side effects than drugs targeting μOR alone.

As mentioned above, one of the severe side-effects associated with chronic morphine use is constipation; this suggests that opioid receptors in the gastrointestinal (GI) tract could be targeted for the treatment of GI tract disorders [18] such as diarrhea. This led to the development of loperamide, a peripherally active μOR agonist, as a therapeutic agent for the treatment of diarrhea [19,20]. However, one of the side-effects associated with the use of loperamide is the development of constipation [21,22]. The possibility that drugs having μOR agonist/δOR antagonist activity could have lesser side effects led to the synthesis of eluxadoline [14,16]. Recent studies show that eluxadoline is a locally acting μOR agonist/δOR antagonist that can normalize GI transit in stressed animals over a wide dose range [16]. Eluxadoline has limited systemic bioavailability which could potentially reduce its effects on the central nervous system and consequently prevent the development of side-effects associated with therapies currently used to treat irritable bowel syndrome with diarrhea (IBS-d). Currently, eluxadoline has completed Phase II [23] and is undergoing Phase III clinical trials for treatment of IBS-d. While in vivo preclinical studies indicate that eluxadoline modulates GI motility and decreases intestinal pain or visceral hyperalgesia without the constipation associated with drugs that activate μOR [16], its mechanism of action is not clear. Since eluxadoline is a mixed μOR agonist/δOR antagonist [14,16,23], it
is possible that it may mediate its effects by targeting \( \mu \text{OR}-\delta \text{OR} \) heteromers. Therefore, in this study we examined the mechanism of the \textit{in vitro} effects of eluxadoline by comparing its activity in cell lines (using an assay that specifically examines heteromer signaling) and in tissues from wild-type (WT) and knockout mice (\( \delta \text{OR}^{-/-} \) or \( \mu \text{OR}^{-/-} \)). Furthermore, we evaluated the extent to which eluxadoline affects GI transit in WT and \( \delta \text{OR}^{-/-} \) mice in a castor oil induced model of diarrhea. We find that eluxadoline-mediated signaling can be significantly, albeit partially, blocked by an \( \mu \text{OR}-\delta \text{OR} \) heteromer selective antibody in cells co-expressing both receptors. We also find that eluxadoline is more effective in blocking castor oil-induced diarrhea in WT mice as compared to \( \delta \text{OR}^{-/-} \) mice. These results suggest that eluxadoline, at least in part, mediates its effects by targeting \( \mu \text{OR}-\delta \text{OR} \) heteromers.

2. Methods

2.1. Cell culture

\( \mu \text{OR}^\beta \text{gal} \) and \( \mu \text{OR}^\beta \text{gal}-\delta \text{OR} \) expressing U2OS cells were a kind gift from DiscoveRx (Fremont, CA, USA). \( \mu \text{OR}^\beta \text{gal} \) cells expressing \( \mu \text{OR} \) tagged with a ProLink/\( \beta \)-galactosidase (\( \beta \text{gal} \)) donor (PK) fragment at the C-terminal region and \( \beta \)-arrestin tagged with a complementary \( \beta \text{gal} \) activator (EA) fragment were grown in MEM alpha (Life Technologies, Grand Island, NY, USA) containing 10% FBS (Biowest SAS, Nuaille, France), streptomycin-penicillin (Life Technologies), 500 \( \mu \text{g/ml} \) geneticin (Life Technologies) and 250 \( \mu \text{g/ml} \) hygromycin (Life Technologies). \( \mu \text{OR}^\beta \text{gal}-\delta \text{OR} \) cells expressing wild-type \( \delta \text{OR} \), \( \mu \text{OR} \) tagged with the PK fragment at the C-terminal region and \( \beta \)-arrestin tagged with the EA fragment were grown in MEM alpha containing 10% FBS, streptomycin-penicillin, 500 \( \mu \text{g/ml} \) geneticin, 250 \( \mu \text{g/ml} \) hygromycin and 0.25 \( \mu \text{g/ml} \) puromycin (Life Technologies).

2.2. \( [\text{35}S] \text{GTP}_\gamma \text{S} \) binding

Membranes were prepared from the spinal cord of either WT (Jackson Laboratories, Sacramento, CA, USA), \( \delta \text{OR}^{-/-} \) (Charles River Laboratories, Kingston, NY, USA), \( \mu \text{OR}^{-/-} \) (a gift from Dr. Charles Mobbs, Ichan School of Medicine at Mount Sinai, NY, USA) or from the ileal longitudinal muscle (containing myenteric plexus) of WT mice as described previously \[24,25\]. Membranes (10 or 20 \( \mu \text{g} \)) were subjected to a \([\text{35}S] \text{GTP}_\gamma \text{S} \) binding assay using DAMGO (R&D Systems, Minneapolis, USA), loperamide (Toronto Research Chemicals Inc., Ontario, Canada), eluxadoline (Furiex, Morrisville, NC, USA) (0–10 \( \mu \text{M} \) final concentration) in the presence or absence of TIPP (10 nM final concentration) (a gift from Dr. Peter Schiller, Institut de Reserches Cliniques de Montreal, Montreal, ON, Canada) as described previously \[25\]. \( \text{EC}_{50} \) and \( E_{\text{max}} \) were calculated using Prism 6 (GraphPad Software, Inc., La Jolla, CA, USA).

2.3. \( \beta \)-arrestin recruitment assay

U2OS cells expressing either \( \mu \text{OR}^\beta \text{gal} \) or \( \mu \text{OR}^\beta \text{gal}-\delta \text{OR} \) were plated in each well (5000 cells) of a 96-well white clear bottom plate in 100 \( \mu \text{l} \) of media. Next day, cells were treated with either DAMGO, loperamide, eluxadoline (0–10 \( \mu \text{M} \) final concentration) in the absence or presence of the \( \delta \text{OR} \) antagonist, TIPP (10 nM final concentration) (a gift from Dr. Peter Schiller) or in the absence or presence of antibodies (1 \( \mu \text{g/well} \)) to either \( \mu \text{OR} \), \( \mu \text{OR}-\delta \text{OR} \)
(generated as reported in [26]) or cannabinoid receptor type 1-angiotensin II receptor type 1 heteromer (CB1R-AT1R) (generated as reported in [27]) for 60 min at 37 °C. β-arrestin recruitment was measured using the PathHunter Chemiluminescence detection kit as described in the manufacturer’s protocol (DiscoveRx, Fremont, CA, USA). EC$_{50}$ and $E_{\text{max}}$ were calculated using Prism 6 (GraphPad Software, Inc., La Jolla, CA, USA).

2.4. Animals

Male C57BL/6 WT and δOR$^{-/-}$ mice (25–35 g; 6–12 weeks old) were obtained from either Jackson Laboratories (Sacramento, CA, USA; WT mice) or Charles River Laboratories (Kingston, NY, USA; δOR$^{-/-}$ mice). All mice were maintained on a 12-h light:12-h dark cycle with rodent chow and water available ad libitum, and housed in groups of five until testing. Animal studies were carried out according to protocols approved by the Icahn School of Medicine at Mount Sinai Animal Care and Use Committee.

2.5. Drug administration

Loperamide (Toronto Research Chemicals, Inc., Ontario, Canada) and eluxadoline (Furiex, Morrisville, NC, USA) were dissolved in 0.5% methylcellulose and 2% DMSO in water. Corresponding vehicle was used for control group. Mice were administered these drugs orally (p.o.). Naltrexone (R&D Systems, Minneapolis, USA) was dissolved in saline and administered intraperitoneally (i.p.). For chronic treatment with eluxadoline and loperamide, mice were treated with these drugs at the dose of 10 mg/kg (p.o.) once a day for 5 days. On the 6th day, mice were euthanized by cervical dislocation and ileum was collected (3–4 mice/sample) for ELISA assay.

2.6. Castor oil-induced diarrhea

Mice were placed in new absorbent lined bottomed cages with no access to food and water for 2 h before the test. Immediately before the test, the absorbent liner was discarded, and a fresh pre-weighed liner was placed in the cage. Diarrhea was induced by oral administration of castor oil (0.6 ml/mouse) (ACROS Organics, Geel, Belgium) to WT or δOR$^{-/-}$ mice. Stools were scored (diarrhea score of 0 = normal; 1 = wet and irregular shape; or 2 = shapeless) and weighed over a 4-h period as described previously [28,29]. After every hour, the absorbent liner was weighed, and another pre-weighed liner was placed in the cage. Diarrhea score represents the most marked change in feces for individual mice during a 4-h period. Loperamide and eluxadoline were administered orally 15 min before the castor oil administration. When naltrexone (10 mg/kg, i.p.) was used, it was administered 20 min before loperamide or eluxadoline administration. Body weight was measured before and 4 h after castor oil administration.

2.7. ELISA assay

Ileal longitudinal muscle (containing myenteric plexus) was prepared as described previously [24]. Membranes (10 μg) from mouse ileal longitudinal muscle were subjected to an ELISA assay using rat anti-μOR antibody (1:500), rat anti-δOR antibody (1:500) (generated as reported in [26]) or mouse anti-μOR-δOR heteromer selective antibody (1:100) as primary antibodies and anti-mouse IgG (1:1000) (Vector laboratories, Inc.,

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Burlingame, CA, USA) or anti-rat IgG (1:1000) (Jackson ImmunoResearch Laboratories, Inc., West Grove, PA, USA) coupled to horseradish peroxidase as secondary antibodies as described previously [30]. ELISA for each sample was performed in triplicate.

2.8. Statistical analysis

The data were expressed as means ± S.E.M. Student’s t-test or one-way ANOVA and multiple-comparison test (Student–Newman–Keuls test or Dunnett’s test) were used to analyze the data. A difference was considered to be significant at $p < 0.05$.

3. Results

3.1. Eluxadoline-mediated signaling

Although phase II clinical studies show that eluxadoline, a locally acting mixed μOR agonist and δOR antagonist, is effective in the treatment of IBS-d patients [23] very little is known about its cellular mechanism of action through heteromers. Therefore, we determined the signaling of eluxadoline and compared it to that of DAMGO, a selective μOR agonist, and of loperamide, a peripherally active μOR agonist used in the treatment of diarrhea [19,20].

For this, we measured G protein activity by carrying out [35S]GTPγS binding assays using spinal cord membranes from WT, δOR−/− or μOR−/− mice. We find that DAMGO, the selective μOR agonist, dose dependently increases [35S]GTPγS binding to spinal cord membranes from WT (EC$_{50}$ ~ 192 nM; $E_{\text{max}}$ ~ 143%) and from δOR−/− (EC$_{50}$ ~ 188 nM; $E_{\text{max}}$ ~ 147%) but not from μOR−/− mice (Fig. 1A). Similarly, loperamide, the peripherally active μOR agonist, dose dependently increases [35S]GTPγS binding to spinal cord membranes from WT (EC$_{50}$ ~ 63 nM; $E_{\text{max}}$ ~ 139%), δOR−/− (EC$_{50}$ ~ 29 nM; $E_{\text{max}}$ ~ 144%) but not from μOR−/− mice (Fig. 1B). Interestingly, although eluxadoline like DAMGO and loperamide did not increase [35S]GTPγS binding to membranes from μOR−/− mice, it induced a dose dependent increase in [35S]GTPγS binding to spinal cord membranes from WT mice with a higher potency (EC$_{50}$ ~ 7 nM) and lower efficacy ($E_{\text{max}}$ ~ 120%) than seen with either DAMGO or loperamide (Fig. 1C). Moreover, in membranes from δOR−/− mice the efficacy of eluxadoline ($E_{\text{max}}$ of ~143%) was higher than in WT membranes ($E_{\text{max}}$ of ~120%) and similar to that seen with DAMGO in δOR−/− mice ($E_{\text{max}}$ of ~147%) (Fig. 1A and C). These results suggest that a portion of eluxadoline-mediated signaling in the presence of both μOR and δOR could be due to activation of μOR-δOR heteromers since in the absence of δOR it behaves like a pure μOR agonist.

Since eluxadoline is effective in the treatment of IBS-d [23] we also carried out [35S]GTPγS binding assays using ileal membranes from WT animals. We find that eluxadoline, DAMGO and loperamide cause dose-dependent increases in [35S]GTPγS binding (Fig. 1D–F). Interestingly, we find that in ileal membranes a combination of the δOR antagonist, TIPPψ, with the μOR agonist, DAMGO, or eluxadoline (a μOR agonist/δOR antagonist) give a higher increase in [35S]GTPγS binding compared to either DAMGO or loperamide (μOR agonists) (Fig. 1G). These results would suggest the presence of μOR-δOR heteromers in ileal tissue that could contribute to the effectiveness of eluxadoline in the treatment of IBS-d.
A common feature of GPCR signaling is that following receptor activation, the receptor is phosphorylated by GRKs and this leads to the recruitment of β-arrestin to the phosphorylated receptor, cessation of G-protein mediated signaling and induction of β-arrestin-mediated signaling (reviewed in [31]). Moreover, several studies in the last decade show that some agonists preferentially signal via one signaling pathway (for e.g., G protein-mediated) versus another (for e.g., β-arrestin-mediated) leading to biased agonism (reviewed in [31]). We therefore compared the ability of DAMGO, loperamide and eluxadoline to induce β-arrestin recruitment in cells expressing either μOR homomers (μORβgalOR cells) or μOR-δOR heteromers (μORβgalδOR cells) using an enzyme complementation assay that we recently used to identify and characterize compounds that preferentially activate the μOR-δOR heteromer [26]. Using this assay, we find that eluxadoline is more potent (i.e., lower EC50) at inducing β-arrestin recruitment than DAMGO or loperamide in μORβgalOR (Fig. 2A–C) or in μORβgalδOR cells (Fig. 2D–F). Interestingly, eluxadoline is less efficacious than DAMGO or loperamide in μORβgalδOR cells suggesting that it’s intrinsic δOR antagonistic activity could affect its ability to induce β-arrestin recruitment (Fig. 2D–F). This is supported by our previous observation that the δOR antagonist, TIPPψ, decreased DAMGO-mediated β-arrestin recruitment only in μORβgalδOR cells but not in μORβgalOR cells [26]. We therefore examined the effect of TIPPψ on DAMGO-, loperamide- or eluxadoline-mediated β-arrestin recruitment. We find that TIPPψ has no effect on β-arrestin recruitment induced by either DAMGO, loperamide or eluxadoline in μORβgalOR cells (Fig. 2A–C). However, in cells expressing μORβgalδOR, TIPPψ decreased DAMGO- or loperamide-induced β-arrestin recruitment but not that induced by eluxadoline (Fig. 2D–F). The lack of effect of TIPPψ on eluxadoline-mediated β-arrestin recruitment in μORβgalδOR cells could be because eluxadoline already exhibits antagonistic activity at δOR. In this context, it is interesting to note that in μORβgalδOR cells the efficacy of eluxadoline to elicit β-arrestin recruitment is 50% lower (Emax ~ 433%) than seen with DAMGO or loperamide (Emax ~ 863% and 956%, respectively). Together these results indicate that a component of eluxadoline’s effects is different from DAMGO or loperamide and is possibly through activation of μOR-δOR heteromers. We tested this by using μOR-δOR heteromer-selective antibodies that we have generated and previously shown that they can block heteromer-mediated binding and signaling [10,26]. Also, we had previously reported that activation of δOR can lead to β-arrestin recruitment to μOR in μORβgalδOR cells and this can be blocked by μOR-δOR heteromer-selective antibodies [26]. In agreement with these observations we find that deltorphin II (Delt II; R&D Systems, Minneapolis, USA), the δOR selective agonist, induces β-arrestin recruitment to μOR in μORβgalδOR cells (Fig. 3B) but not to μβgalOR cells (Fig. 3A) and this can be blocked by μOR-δOR heteromer-selective antibodies and not by μOR selective antibodies or by antibodies selective to an unrelated heteromer, i.e., the CB1R-AT1R heteromer (Fig. 3B). We find that loperamide-mediated β-arrestin recruitment both in μβgalOR (Fig. 3A) and μORβgalδOR cells (Fig. 3B) is significantly blocked by μOR-selective antibodies but not by antibodies selective for either the μOR-δOR or CB1R-AT1R heteromer. This supports that loperamide exerts its effects by activating μOR. In the case of eluxadoline, we find that its ability to induce β-arrestin recruitment is blocked by μOR-selective antibodies by ~40% in μβgalOR cells (Fig. 3A) and by ~17% in μORβgalδOR cells (Fig. 3B). In addition, we find that μOR-δOR heteromer-selective antibodies block 27% of eluxadoline-mediated β-arrestin recruitment in μORβgalδOR cells (Fig. 3B) but have no
effect in $\mu^\beta\text{gal}$OR cells (Fig. 3A); no effect of CB1R-AT1R heteromer-selective antibodies on eluxadoline-mediated $\beta$-arrestin recruitment was observed in either $\mu^\beta\text{gal}$OR (Fig. 3A) or $\mu\text{OR}^\beta\text{gal}$-OR cells (Fig. 3B). Taken together these results suggest that in cells co-expressing $\mu$OR and $\delta$OR a portion of eluxadoline-mediated signaling occurs via activation of $\mu$OR-$\delta$OR heteromers while in cells expressing only $\mu$OR eluxadoline functions as a $\mu$OR agonist.

### 3.2. Castor oil-induced diarrhea

Previous studies showed that eluxadoline can normalize GI transit over a wider dose range compared to loperamide in WT mice [16]. Although WT mice express both $\mu$OR and $\delta$OR in myenteric plexus [32], not much is known about the contribution of each receptor type or of $\mu$OR-$\delta$OR heteromers to the effects of eluxadoline on GI transit. To investigate this we used WT and $\delta$OR$^{-/-}$ mice and compared the effects of eluxadoline and loperamide on castor oil induced diarrhea. We find that oral administration of castor oil induces severe diarrhea (i.e., higher score) both in WT (Fig. 4A) and $\delta$OR$^{-/-}$ mice (Fig. 4B). The castor oil-induced diarrhea in WT mice was reduced by administration of 5 or 10mg/kg of either loperamide or eluxadoline (Fig. 4A). Moreover, this reduction in castor oil-induced diarrhea by either loperamide or eluxadoline at 10 mg/kg were not observed if the animals were pre-treated with the $\mu$OR antagonist, naltrexone (Fig. 4A); studies have reported that antagonizing the activity of opioid receptors, including $\mu$OR, has no effect on castor oil induced diarrhea [33, 34]. Interestingly, in $\delta$OR$^{-/-}$ mice a 5 mg/kg dose of either loperamide or eluxadoline did not significantly reduce castor oil-induced diarrhea (Fig. 4B); a higher dose of 10 mg/kg was required to completely eliminate diarrhea in the case of loperamide and partially eliminate in the case of eluxadoline (Fig. 4B).

Next, we correlated in the same animals the diarrheal scores with fecal output by measuring fecal weight. We find that castor oil administration significantly increases the fecal output in WT (Fig. 5A) and $\delta$OR$^{-/-}$ mice (Fig. 5B). In addition, we find that in WT mice, treatment with either loperamide or eluxadoline significantly reduces the castor oil induced fecal output and the values were below that seen with controls administered with vehicle instead of castor oil (Fig. 5A). These effects were blocked by coadministration of naltrexone (Fig. 5A). Similar results were obtained in $\delta$OR$^{-/-}$ mice (Fig. 5B) although a 5 mg/kg dose caused a less pronounced decrease in fecal output compared to that seen in WT animals for both loperamide and eluxadoline (Fig. 5A and B). The fact that a 5 mg/kg dose of either loperamide or eluxadoline significantly decreased fecal output in $\delta$OR$^{-/-}$ mice but had no significant effect on the diarrhea scores (Fig. 4B) in the same animals could be due to the qualitative nature of the diarrhea score. Taken together these results showing that eluxadoline (at 5 mg/kg) produced a lower blockade of diarrhea and fecal output in $\delta$OR$^{-/-}$ mice compared to WT mice suggests that $\delta$OR and/or $\mu$OR-$\delta$OR heteromers could contribute to its anti-diarrheal effect in WT mice. Moreover, a dose-response effect with eluxadoline is observed in $\delta$OR$^{-/-}$ but not in WT mice which would again indicate that $\delta$OR activity may modulate eluxadoline’s effects on $\mu$OR in WT mice and that eluxadoline behaves as a $\mu$OR agonist in mice that lack $\delta$OR.

Next we examined the changes in body weight in WT and $\delta$OR$^{-/-}$mice and found that in both groups castor oil-induced diarrhea led to decreases in body weight (Fig. 6A and B) and
this was blocked by loperamide and eluxadoline. Both loperamide and eluxadoline effects could be partly blocked by naltrexone. Although not statistically significant, the amount of body weight change in δOR−/− appeared more robust compared to WT mice. Interestingly, the effect of naltrexone was more pronounced in δOR−/− mice (Fig. 6). The fact that 5 mg/kg eluxadoline was less effective in δOR−/− mice as compared to the WT mice is consistent with the results with diarrhea score and fecal output and further support a role for δOR (or μOR-δOR heteromer) in this effect in WT mice.

3.3. Chronic eluxadoline treatment

We have previously reported/noted that chronic treatment with morphine under a paradigm that leads to the development of tolerance causes an increase in μOR-δOR heteromer expression in different brain regions [10]. Therefore, in this study we examined whether μOR-δOR heteromers are present in myenteric neurons from GI tissue and ascertained the effect of long-term treatment with loperamide or eluxadoline on receptor expression and on body weight. We find that administration of these compounds at the dose of 10 mg/kg/day for 5 days did not lead to significant changes in body weight (Fig. 7A), indicating that repeated oral treatment with either drug would not affect the nutrient absorptive function in intestine. In addition, using antibodies selective for either μOR, δOR or μOR-δOR heteromers we can detect the presence of μOR, δOR or μOR-δOR heteromers in mouse ileal longitudinal muscle (containing the myenteric plexus) (Fig. 7B). In vehicle treated animals levels of δOR appear to be the highest, followed by levels of μOR and μOR-δOR heteromers. These results support findings by a study reporting co-localization of μOR and δOR in myenteric neurons [32]. In addition, we find that chronic treatment with loperamide or eluxadoline does not induce significant changes in the levels of μOR, δOR or μOR-δOR heteromer levels in ileal longitudinal muscle preparations containing myenteric plexus from WT mice.

4. Discussion

Several studies have examined the pharmacological profiles of μOR-δOR heteromers (reviewed in [35,36]). Among them a few have demonstrated that occupancy of δOR enhances μOR activity. Thus δOR selective antagonists were shown to enhance μOR ligand binding, μOR ligand-mediated signaling and μOR-mediated (i.e., morphine-mediated) analgesia [7-9,37]. Furthermore, studies showed that mice with reduced δOR levels (through the use of antisense oligonucleotides) or δOR−/− mice did not develop tolerance or dependence to morphine [38,39]. Together these studies indicate that the use of δOR antagonists could lead to a decrease in the adverse effects associated with in vivo administration of μOR agonists. Based on this possibility a number of ligands were synthesized that have dual μOR agonist and δOR antagonist activity [13-15,17,26,40,41]. The earliest studies with a single compound possessing mixed μOR agonistic and δOR antagonistic activities involved peptide ligands [42]. These peptides were found to produce potent antinociception with reduced tolerance compared to morphine and no physical dependence was observed upon chronic administration [42]. Among the non-peptide ligands possessing mixed μOR agonistic and δOR antagonistic activities, 14-alkoxypyridomorphinans have been reported to induce potent antinociception but
diminished tolerance development as compared to morphine [13]. However, to date it is not clear as to whether these compounds exert their effects by binding to individual μOR or δOR or by targeting the μOR-δOR heteromer.

In this study, we find that loperamide, a μOR agonist, is more potent but as efficacious as DAMGO (a peptidic μOR agonist) in promoting [35S]GTPγS binding while being more potent but less efficacious at recruiting β-arrestin in μβgalOR cells, suggesting that it exhibits bias towards G-protein mediated signaling. Biased signaling, i.e., the ability of some agonists to preferentially signal via one signaling pathway (for e.g., G protein-mediated) versus another (for e.g., β-arrestin-mediated) is being increasingly reported for G-protein coupled receptors (reviewed in [31]). Interestingly, eluxadoline, a mixed μOR agonist/δOR antagonist acts as a μOR agonist in the absence of δOR, but exerts some of its effects via the μOR-δOR heteromer in the presence of δOR. Importantly, blockade of eluxadoline-mediated signaling by μOR-δOR heteromer selective antibody supports the idea that this compound targets μOR-δOR heteromers. Furthermore, the in vivo findings of the differences in anti-diarrheal effect of lower dose of eluxadoline between WT and δOR−/− mice is consistent with the notion that, eluxadoline, at least in part, targets μOR-δOR heteromers.

Previous studies that tested the anti-diarrheal effects of eluxadoline in either novel-environment stressed mice or in intracolonic mustard oil-induced intestinal inflammatory model found that it reduces GI transit and fecal output [16]. Consistent with this we find that eluxadoline exhibits similar effects on castor oil induced diarrhea (Figs. 4 and 5). However, in the present study we find that the lower dose (5 mg/kg) of eluxadoline slightly reduced fecal output to below the vehicle control in WT mice (i.e., induced constipation). This is in contrast to previous reports showing that eluxadoline even at doses of 5–25 mg/kg did not cause constipation in the novel-environment stressed mice model of diarrhea [16]. These differences could be due to the differences in the stressor used in the studies (novel environment vs. castor oil) and the time period for measuring fecal output since Wade et al. (2012) measured for only 1 h after the novel-environment stress, while we measured for 4 h after castor oil injection. Castor oil is known to release ricinoleic acid followed by alterations in ion transport and water flux in the intestine [43-46] leading to increases in fecal output or diarrhea. In contrast, novel environment stress is a form of psychological stress that is accompanied by behavioral changes like grooming, rearing and sniffing [47-49]. It is known that the novel-environment-induced increase in fecal output is mediated by an increase in corticotropin-releasing hormone and thyrotropin-releasing hormone and by activation of cholinergic and serotoninergic neurons [50]. It is possible that these differences as well as changes in the levels and/or activity of intestinal μOR-δOR heteromers under these two assay conditions are responsible for the observed differences.

Relatively few studies have examined the levels of opioid receptor proteins in the GI tract. Using enhanced green fluorescent protein (eGFP)-tagged δOR (δOReGFP) expressing mice, the distribution of δOR was found to be confined to enteric neurons and fibers within the muscularis externa; submucosal plexus and myenteric plexus [32]. This study also showed that, in the myenteric ganglia, over 80% of δOReGFP positive myenteric neurons co-expressed μOR, and 60% of μOR positive neurons co-expressed δOReGFP [32]. This is consistent with a previous study that showed co-localization of μOR and δOR in myenteric
neurons by immunohistochemistry [51]. These results suggest that μOR-δOR heteromers are present in the myenteric neurons. In this study, we detected the presence of μOR-δOR heteromers in ileal longitudinal muscle of mice that includes myenteric neurons [24] using μOR-δOR heteromer-selective antibodies. Together these results support the presence of μOR-δOR heteromers in ileal tissue.

Studies have shown that chronic morphine treatment leads to increase in μOR-δOR heteromer levels in select brain regions [10]. Moreover, μOR-δOR heteromerization changes morphine-mediated signaling from G-protein-into β-arrestin mediated which could contribute to side-effects such as the development of analgesic tolerance [11,37,52]. Interestingly, β-arrestin2 knockout mice exhibit less morphine-induced constipation than their WT counterparts [53], indicating an involvement of β-arrestin mediated signaling (potentially via μOR-δOR heteromers) in the constipating effects of morphine [54]. However, chronic morphine administration does not lead to development of tolerance to the constipating side-effect [55-58]. This led us to wonder if intestinal μOR, δOR, and μOR-δOR heteromer levels are altered following chronic treatment with drugs. In this study we did not detect significant changes in μOR, δOR, and μOR-δOR heteromer levels in the intestine following chronic treatment with either loperamide or eluxadoline compared to controls treated with vehicle (Fig. 7). However, preliminary studies detect a slight increase albeit not significant in μOR-δOR heteromer levels in ileum following chronic morphine administration (data not shown). This would suggest a differential regulation of μOR-δOR heteromer function in brain and gut, which is consistent with what has been previously reported [54].

The detection of μOR-δOR heteromers in mouse ileal tissue together with the anti-diarrheal effect of eluxadoline and in vitro data showing that eluxadoline-mediated signaling is reduced by μOR-δOR heteromer-selective antibodies indicates that eluxadoline, at least in part, mediates its effects by targeting μOR-δOR heteromers in the intestine. This would suggest that intestinal μOR-δOR heteromers could be a potential therapeutic target for the treatment of GI tract disorders including IBS-d. However, additional studies examining the level and changes in the localization of μOR-δOR heteromers in the human GI tract following IBS-d are required to demonstrate that the μOR-δOR heteromer is a novel therapeutic target for the treatment of this disorder.

In this study, we find that in the absence of δOR, eluxadoline behaves as a potent μOR agonist. However, co-expression of μOR and δOR alters the signaling profile of eluxadoline that can be partly blocked by μOR-δOR heteromer-selective antibodies. Thus, the actions of eluxadoline could, at least in part, be due to targeting of μOR-δOR heteromers in the gut. In addition, we find that eluxadoline can block castor oil-induced diarrhea in WT mice and this is attenuated in δOR−/− mice indicating the involvement of δOR probably through μOR-δOR heteromerization in the in vivo effects of eluxadoline.

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Abbreviations

GI  gastrointestinal
IBS-d  irritable bowel syndrome with diarrhea
μOR  mu opioid receptor
δOR  delta opioid receptor
βgal  beta-galactosidase
GTPγS  guanosine 5′-O-(3-thiotriphosphate)
DAMGO  [D-Ala², N-MePhe⁴, Gly-ol]-enkephalin
CB1R  cannabinoid receptor type1
AT1R  angiotensin II receptor type 1
ELISA  enzyme-linked immunosorbent assay
EC₅₀  50% effective concentration
E_max  maximum effective concentration
WT  wild-type
−/−  knockout
eGFP  enhanced green fluorescent protein

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Fig. 1.
Effect of DAMGO, loperamide and eluxadoline on G-protein activation. (A–C) Membranes (10 μg) from spinal cords of WT, μOR−/− and δOR−/− mice were subjected to a [35S]GTPγS binding assay using DAMGO (A), loperamide (B), and eluxadoline (C) (0–10 μM final concentration) as described in Section 2. (D–G) Membranes (20 μg) from the ileum of WT mice were subjected to a [35S]GTPγS binding assay using DAMGO (D and G), loperamide (E and G), and eluxadoline (F and G) (0–10 μM final concentration) in the presence or absence of TIPPψ (10 nM final concentration) as described in Section 2. (G) Represents E_{max} (% of basal) obtained with 10 μM final concentration of DAMGO (±10 nM final concentration of TIPPψ), eluxadoline or loperamide. Basal values determined in the absence of the agonist were taken as 100%. Results are the mean ± S.E.M. n= 3–9. n.d., Not determined. *p < 0.05; **p < 0.01, Dunnett’s test.
Fig. 2. Effect of DAMGO, loperamide and eluxadoline on β-arrestin recruitment. Cells (5000/well) expressing either μβgalOR (A–C) or μβgalOR-δOR (D–F) were treated with either DAMGO (A and D), loperamide (B and E), eluxadoline (C and F) (0–10 μM final concentration) in the absence or presence of the δOR antagonist, TIPPψ (10 nM final concentration) for 60 min at 37 °C and β-arrestin recruitment was measured as described in Section 2. Results are the mean ± S.E.M. n = 4–12. *p < 0.05, **p < 0.01, vs. absence of TIPPψ, t-test.
Fig. 3.
Effect of μOR-δOR heteromer-selective antibody on eluxadoline-mediated signaling. Cells (5000 cells) expressing either μgalOR (A) or μgalOR-δOR (B) were treated with either deltorphin II (Delt II), loperamide or eluxadoline (1 μM final concentration) in the absence or presence of antibodies (Ab, 1 μg/well) to either μOR, μOR-δOR heteromer or CB1R-AT1R heteromer for 60 min at 37 °C and β-arrestin recruitment was measured as described in Section 2. Results are the mean ± S.E.M. n = 4. *p < 0.05, **p < 0.01, vs. no Ab treatment for each group, Dunnett’s test.
Fig. 4.
Effect of loperamide and eluxadoline on castor oil-induced diarrhea. Diarrhea was induced by oral administration of castor oil (0.6 ml/mouse) in WT (A) or δOR−/− (B) mice. Stools were scored for diarrhea (0 = normal; 1 = wet and irregular shape; or 2 = shapeless) for 4 hours described in Section 2. Loperamide and eluxadoline (5 or 10 mg/kg) were administered orally 15 min before the castor oil administration. Naltrexone (10 mg/kg, i.p.) was administered 20 min before loperamide or eluxadoline administration. Results are the mean ± S.E.M. n = 3–6. **p < 0.01, vs. vehicle control; ###p < 0.01, vs. castor oil alone; $$p < 0.01, vs. castor oil + loperamide (10 mg/kg, p.o.); ††p < 0.01, vs. castor oil + eluxadoline (10 mg/kg, p.o), Student–Newman–Keuls test.
Fig. 5.
Effect of loperamide and eluxadoline on castor oil-induced diarrhea (fecal output). Diarrhea was induced by oral administration of castor oil (0.6 ml/mouse) in WT (A) or δOR−/− (B) mice. Stools were collected and weighed during 4 h as described in Section 2. Loperamide and eluxadoline (5 or 10 mg/kg) were administered orally 15 min before the castor oil administration. Naltrexone (10 mg/kg, i.p.) was administered 20 min before loperamide or eluxadoline administration. Results are the mean ± S.E.M. n = 3–6. **p < 0.01, vs. vehicle control; ##p < 0.01, vs. castor oil alone; $$$$$p < 0.01, vs. castor oil + loperamide (10 mg/kg, p.o.); ††p < 0.01, vs. castor oil + eluxadoline (10 mg/kg, p.o), Student–Newman–Keuls test.
Fig. 6.
Effect of loperamide and eluxadoline on castor oil-induced body weight change. Diarrhea was induced by oral administration of castor oil (0.6 ml/mouse) in WT (A) or δOR−/− (B) mice. Body weight was measured before and 4 h after castor oil administration. Loperamide and eluxadoline (5 or 10 mg/kg) were administered orally 15 min before the castor oil administration. Naltrexone (10 mg/kg, i.p.) was administered 20 min before loperamide or eluxadoline administration. Results are the mean ± S.E.M. n = 3–6. **p < 0.01, vs. vehicle control; ##p < 0.01, vs. castor oil alone; $$p < 0.01, vs. castor-oil + loperamide (10 mg/kg, p.o.); †p < 0.05, ††p < 0.01, vs. castor oil + eluxadoline (10 mg/kg, p.o), Student–Newman–Keuls test.
Fig. 7.
Effect of chronic treatment of loperamide and eluxadoline on body weight (A) and on receptor expression levels in ileal longitudinal muscle (B). Mice were treated with loperamide or eluxadoline (10 mg/kg, p.o., once a day for 5 days), or with 0.5% methylcellulose (0.1 ml/10 g; vehicle). (A) Body weight was measured immediately before the daily administration. Results are the mean ± S.E.M. n = 6–7. (B) On 6th day, ileum was collected (3–4 mice/sample). Membranes (10 μg) from mouse ileal longitudinal muscle (containing the myenteric plexus) were subjected to an ELISA assay in as described in Section 2. Tissues from 3 to 4 individual animals were pooled and collected as one sample. ELISA was performed in triplicate. Results are the mean ± S.E.M. n = 5.