Heteromerization of the \( \mu \)- and \( \delta \)-Opioid Receptors Produces Ligand-Biased Antagonism and Alters \( \mu \)-Receptor Trafficking

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Received January 7, 2011; accepted March 18, 2011

ABSTRACT

Heteromerization of opioid receptors has been shown to alter opioid receptor pharmacology. However, how receptor heteromerization affects the processes of endocytosis and postendocytic sorting has not been closely examined. This question is of particular relevance for heteromers of the \( \mu \)-opioid receptor (MOR) and \( \delta \)-opioid receptor (DOR), because the MOR is recycled primarily after endocytosis and the DOR is degraded in the lysosome. Here, we examined the endocytic and postendocytic fate of MORs, DORs, and DOR/MOR heteromers in human embryonic kidney 293 cells stably expressing each receptor. We show that DOR/MOR heteromers that are endocytosed in response to methadone are targeted for degradation, whereas MORs in the same cell are significantly more stable. It is noteworthy that we found that the DOR-selective antagonist naltriben mesylate could block both methadone- and \([\alpha\text{-Ala2,}\text{NMe-Phe4,}\text{Gly-ol5}]\text{-enkephalin}]\)-induced endocytosis of the DOR/MOR heteromers but did not block signaling from this heteromer. Together, our results suggest that the MOR adopts novel trafficking properties in the context of the DOR/MOR heteromer. In addition, they suggest that the heteromer shows “biased antagonism,” whereby DOR antagonist can inhibit trafficking but not signaling of the DOR/MOR heteromer.

Introduction

\( \delta \)-Opioid receptors (DORs) and \( \mu \)-opioid receptors (MORs) belong to the G protein-coupled receptor (GPCR) superfamily, and upon activation they regulate a variety of physiological functions including pain processing, anxiety, and reward (for review see Bodnar, 2010). After activation, opioid receptors, like most GPCRs, can be rapidly phosphorylated by GPCR kinases, bind arrestin proteins (Ferguson et al., 1998), and be endocytosed. After endocytosis receptors are then either targeted to degradation (for the DOR) (Whistler et al., 2002) or recycled back to the cell surface (for the MOR) (Law et al., 2000; Whistler et al., 2002).

This study was supported by a Schrödinger fellowship from the Austrian Science Fund [Fellowship J 2967-B09] (to L.M.-L.); the National Institutes of Health National Institute on Drug Abuse [Grant DA015232] (to J.L.W.); and funds provided by the State of California for medical research through the University of California, San Francisco (to J.L.W.). Article, publication date, and citation information can be found at http://jpet.aspetjournals.org. doi:10.1124/jpet.111.179093. [1] The online version of this article (available at http://jpet.aspetjournals.org) contains supplemental material.

Many GPCRs, including opioid receptors, are believed to function as dimers or higher-order oligomers (Rozenfeld and Devi, 2010). There is substantial evidence that the MOR and DOR form heteromers in vitro (Cvejic and Devi, 1997; George et al., 2000; Gomes et al., 2000, 2004; Fan et al., 2005; Hasbi et al., 2007) and mounting evidence that they form functional heteromers in vivo as well (Gupta et al., 2010; Wang et al., 2010; He et al., 2011). Coexpression of opioid receptors has been shown to alter opioid ligand properties and affect receptor signaling in cell culture model systems (Jordan and Devi, 1999; George et al., 2000; Gomes et al., 2004; Rozenfeld and Devi, 2007; Kabli et al., 2010), and these differences are hypothesized to occur as a consequence of receptor heteromerization. In addition, the DOR/MOR heteromer is reported to couple preferentially with the inhibitory pertussis toxin-insensitive \( G_{\text{ai}} \) subunit instead of pertussis toxin-sensitive \( G_{\text{ai}} \) (Fan et al., 2005; Hasbi et al., 2007). Furthermore, DOR/MOR heteromerization seems to also influence receptor maturation (Décaillot et al., 2008) and arrestin-mediated signaling (Rozenfeld and Devi, 2007). In addition, some MOR- and DOR-selective agonists have been shown to promote endocytosis when both receptors are coexpressed, although this

ABBREVIATIONS: DOR, \( \delta \)-opioid receptor; MOR, \( \mu \)-opioid receptor; GPCR, G-protein coupled receptor; HEK, human embryonic kidney; PBS, phosphate-buffered saline; ANOVA, analysis of variance; RFU, relative fluorescence units; PAGE, polyacrylamide gel electrophoresis; DPDPE, \([\alpha\text{-Pen2,} \alpha\text{-Pen5}]\text{-enkephalin}]\); BNTX, 7-benzylidene naltrexone maleate; NTB, naltrexone mesylate; NTI, naltrindole hydrochloride; DAMGO, \([\alpha\text{-Ala2,}\text{NMe-Phe4,}\text{Gly-ol5}]\text{-enkephalin}]\); HA, hemagglutinin.
phenomenon seems to be ligand-dependent (Hasbi et al., 2007; Kabli et al., 2010), occurring with some but not all agonists. However, those prior studies did not examine endocytosis or postendocytic trafficking of DOR/MOR heteromers in response to many of the clinically relevant opioid drugs. In particular, there has been no exploration of the postendocytic fate of the DOR/MOR heteromer after activation by MOR agonists. This is particularly important for heteromers containing MOR and DOR because these two receptors have dramatically different postendocytic fates (Law et al., 2000; Tsaod and von Zastrow, 2000; Whistler et al., 2002). Specifically, after endocytosis MORs are reported to be recycled (Law et al., 2000; Whistler et al., 2002; Liang et al., 2008) and show rapid functional resensitization (Alvarez et al., 2002). In contrast, the DOR binds the GPCR-associated sorting protein and is targeted to the lysosomal degradation pathway after endocytosis (Tsaod and von Zastrow, 2000; Whistler et al., 2002), although the rate and extent of degradation are reported to be agonist-dependent (Zhang et al., 1999; Lecq et al., 2004; Binyaminy et al., 2008; Archer-Lahlu et al., 2009).

Here, we examined the endocytic and postendocytic trafficking properties of the DOR/MOR heteromers and examined whether the heteromer showed changes in “biased agonism” for trafficking compared with the receptor homomers.

Materials and Methods

Reagents. [α-Ala2,NMe-Phe4,Gly-ol5]-enkephalin (DAMGO), naltre- 
binen mesylate (NTB), [d-Pen2,d-Pen5]-enkephalin (DPDPE), and 7-benzylidene naltrexone maleate (BNTX) were purchased from Tocris 
Bioscience (Ellisville, MO). Naltrindole hydrochloride (NTB) and meth-
ando hydrochloride were purchased from Sigma-Aldrich (St. Louis, 
MO). All compounds were dissolved in water, with the exception of 
BNTX and NTB, which were dissolved in 5% dimethyl sulfoxide. Mouse 
M1 and M2 monoclonal antibody, anti-FLAG M2 affinity matrix, albu-
min from bovine serum, 1-glutathione, iodoacetamide, gelatin from 
bovine skin type B, Triton X-100, and Tween 20 were purchased 
from Sigma-Aldrich. Anti-HA.11 beads were from Covance Re-
search Products (Princeton, NJ).

Cell Culture. HEK293 cells (American Type Culture Collection, 
Manassas, VA) were grown in Dulbecco’s modified Eagle’s medium 
(Invitrogen, Carlsbad, CA) supplemented with 10% fetal bovine se-
rum (HyClone Laboratories, Logan, UT). N-terminal signal sequence 
and either HA- or FLAG-tagged c-DNA murine opioid receptor con-
structs were stably expressed in HEK293 cells. For generation of 
clonal stable cell lines, single colonies were chosen and propagated in the presence of selection-containing medium. Cell lines were care-
fully matched for expression (see Waldhoer et al., 2005).

Immunofluorescence Confocal Microscopy. HEK293 cells stably 
expressing N-terminal FLAG-MOR, HA-DOR alone, or 
FLAG-MOR and HA-DOR together were incubated with monoclonal 
anti-HA.11 antibody (Covance Research Products) and/or M1 anti-
FLAG antibody (Sigma-Aldrich) for 30 min to label surface receptors. 
Cells were treated as indicated. Subsequently, cells were fixed with 
3.7% formaldehyde in PBS for 20 min at room temperature and permeabilized with 0.1% Triton X-100, essentially as de-
scribed (Whistler and von Zastrow, 1998). Cells were then incu-
bated with subtype-selective fluorescent anti-mouse antibody di-
rected against M1 (Alexa488 IgG2b) and HA (Alexa594 IgG1, 
1:1000, 30 min) (Invitrogen) to label MOR and DORs with differ-
cent colors. After staining, cells were mounted in Vectashield 
mounting medium (Vector Laboratories, Burlingame, CA) and 
analyzed using a Zeiss LSM 510 META Axioplan 2 confocal mi-
croscope (Carl Zeiss Inc., Thornwood, NY).

Biotin Protection Endocytosis and Endocytosis-Degradation 
Assays. HEK293 cells stably expressing N-terminal FLAG-
MOR alone or FLAG-MOR and HA-DOR together were grown to 90% 
confluence in 10-cm plates. Cells were washed twice in PBS and 
biotinylated with 0.3 mg/ml disulfide-cleavable biotin (Thermo 
Fisher Scientific, Waltham, MA) at 4°C for 30 min to selectively label 
a pool of receptors at the cell surface as described (Finn and Whistler, 
2001). For quantification of endocytosis, cells were washed in PBS 
and placed in prewarmed medium for 15 min before treatment with 
ligand or no treatment for 30 min. For quantification of stability/
degradation, cells were incubated with ligand for prolonged periods 
of time as indicated. Concurrent with ligand treatment total and 
strip plates remained at 4°C. After ligand treatment, plates were 
ashed in PBS, and the remaining cell surface-biotinylated recep-
tors were stripped in 50 mM glutathione, 75 mM NaCl, 75 mM 
NaOH, and 10% fetal bovine serum at 4°C for 60 min (twice 30 min; 
including strip but not total). Cells were quenched with PBS con-
taining 50 mM iodoacetamide and 10% bovine serum albumin for 30 
min (including total). Afterward, all cells were lysed in 0.1% Triton 
X-100, 150 mM NaCl, 25 mM KCl, and 10 mM Tris-HCl, pH 7.4 with 
protease inhibitors (Roche Diagnostics, Basel, Switzerland). Lysates 
were cleared by centrifugation at 10,600g (Eppendorf 5417R; Eppen-
dorf North America, New York, NY) for 10 min at 4°C. In cells 
expressing only one type of receptor they were immunoprecipitated 
overnight at 4°C with anti-FLAG M2 or HA.11 affinity matrix (de-
pending on the epitope tag), washed, and resolved by SDS-PAGE.
The “protected” pool of endocytosed receptors were visualized by streptavidin overlay. This protected pool shrinks across time for 
receptors that are degraded, because no new receptors are biotin-
ylated. This pool remains constant for receptors that are endocytosed, 
recycled, and re-endocytosed. For monitoring homomer versus het-
eromer trafficking in the same cells, cells were biotinylated, treated 
with agonist for the indicated time, stripped, quenched, and lysed as above. Lysates were then incubated with anti-FLAG M2 affinity 
matrix overnight at 4°C, which immunoprecipitated both FLAG-
MOR homomers and FLAG-MOR/HA-DOR heteromers. The lysate 
remaining was separated from the pellet and then immunoprecipi-
tated with HA.11 affinity matrix to isolate HA-DOR homomers. The 
Mt containing FLAG M2 affinity matrix, and therefore both MOR 
and DOR heteromers, was incubated with FLAG peptide to release all receptors to the lysate. This lysate was then incubated with HA.11 affinity matrix to selectively immunoprecipi-
tate HA-DOR/FLAG-MOR heteromers (that had already been immu-
noprecipitated with M2 matrix). The HA.11 affinity matrix con-
tained the MOR/DOR heteromers, whereas the lysate contained 
MOR homomers. Finally, the lysate remaining from the immunopre-
cipitation with HA.11 affinity matrix was incubated with anti-FLAG 
M2 affinity matrix to specifically isolate FLAG-MOR homomers. All 
matrix beads were washed and precipitates were deglycosylated with peptide N-glycosidase F (New England Biolabs, Ipswich, MA) in 
10 mM Tris, pH 7.5, for 1 h at 37°C, denatured with SDS sample buffer (no reducing agent), and resolved by SDS/PAGE. Blots were 
blotted in 5% milk, washed thoroughly, and incubated with Vec-
tastain ABC reagent (Vector Laboratories) for 30 min and washed 
throughout again. Blots were developed with enhanced chemilumines-
tence reagents (GE Healthcare, Chalfont St. Giles, Buckinghamshire, 
UK), scanned, and quantified using ImageJ software (National Insti-
tutes of Health, Bethesda, MD). No previous studies have used serial 
immunoprecipitation to specifically follow the postendocytic fate of ho-
mer and heteromer species in the same cell line.

Calcium Mobilization Assay. HEK293 cells stably expressing 
N-terminal FLAG-MOR alone or FLAG-MOR and HA-DOR 
together were seeded onto 96-well black clear-bottom plates from Corning Life 
Sciences (Lowell, MA). Cells were then transiently transfected with chimeric G protein δ-Gq₄₅-myr (100 ng for every 70,000 cells) (Ko-
stenis, 2001). One day after transfection, cells were loaded for 60 min 
with a Ca²⁺ fluorophore (Molecular Devices, Sunnyvale, CA) and 
stimulated with ligand as indicated in the figure legends. Intracel-

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lular Ca\(^{2+}\) release was measured immediately after agonist application in a Flex apparatus (Molecular Devices) for 2 min. For experiments with antagonist, cells were preincubated with antagonist at the stated concentrations for 20 min before measurement of Ca\(^{2+}\) release in relative fluorescence units (RFU). Data are represented as percentage of the maximal effect given by the MOR agonist.

**Results**

**MOR Agonists Promote Endocytosis of DORs When Both Opioid Receptors Are Coexpressed.** Epitope-tagged versions of the murine DOR (HA-DOR) and MOR (FLAG-MOR) were stably expressed alone or together in a set of cell lines carefully matched for expression as described previously (Waldhoer et al., 2005). We then examined whether endocytosis of the MOR was affected by the presence of the DOR and vice versa. Cells expressing MOR or DOR alone or coexpressing MOR and DOR were treated with 1 \(\mu\)M of the MOR agonists methadone and DAMGO (Fig. 1). MORs were endocytosed upon activation with both DAMGO and methadone (Fig. 1A). The MOR agonist morphine did not promote endocytosis of either MOR or DOR (Supplemental Fig. 1). DORs were not endocytosed after application of the MOR agonists DAMGO or methadone when DOR was expressed alone (Fig. 1B), consistent with the low affinity of these drugs for the DOR (Raynor et al., 1994; Schmidt et al., 2002). However, DORs were readily endocytosed in response to the MOR agonist DPDP (Supplemental Fig. 1).

We next examined whether coexpression of MOR and DOR altered endocytosis in response to the small-molecule agonist methadone. Indeed, in the presence of MOR, DORs were endocytosed in response to methadone (Fig. 1C). Likewise, the peptide agonist DAMGO also promoted endocytosis of the MOR when it was coexpressed with MOR (Fig. 1C). Morphine did not promote endocytosis of MOR and DOR even when they were coexpressed (Supplemental Fig. 1).

The effects of methadone and DAMGO on endocytosis of MOR and DOR homomers and MOR/MOR heteromers were then quantified by biotin protection assay and serial immunoprecipitation as described under Materials and Methods. In brief, cells coexpressing DORs and MORs were biotinylated with thio-cleavable biotin to label all the receptors present at the surface. Cells were then pretreated with the MOR agonist methadone or DAMGO (1 \(\mu\)M, 30 min; Fig. 1, D and E). Residual surface biotin was stripped leaving only the protected endocytosed pool. We then selectively immunoprecipitated DOR/MOR heteromers and MOR homomers in the DOR/MOR coexpressing cell line by serial immunoprecipitation (see Materials and Methods). Both methadone and DAMGO promoted endocytosis of MOR homomers (Fig. 1, D and E middle) and DOR/MOR heteromers (Fig. 1, D and E top), but not DOR homomers in the same cells (Fig. 1, D and E bottom).

**MOR Endocytosis Is Inhibited in the Presence of DOR by MOR Antagonist.** Based on these results, we hypothesized that the DORs were being coendocytosed with the MORs in response to methadone (or DAMGO) as a heteromeric complex. If this were the case, we expected that occupying the DOR with an antagonist might selectively influence the trafficking of the DOR/MOR heteromer but not MOR homomers in response to MOR agonist. To examine this hypothesis, cells expressing MOR or DOR alone or coexpressing MOR and DOR were preincubated with the MOR-selective antagonist NTB for 20 min, then challenged with 1 \(\mu\)M of methadone or DAMGO for another 30 min still in the presence of antagonist. Under these conditions, NTB seemed to inhibit the endocytosis of MORs in the cells expressing MOR and

Fig. 1. Agonist-occupied MORs promote coendocytosis of DORs. A to C, HEK293 cells stably expressing FLAG-MOR (A), HA-DOR (B), or FLAG-MOR and HA-DOR (C) were incubated with antibody recognizing the N-terminal epitope tags for 30 min to label surface receptors. Cells were then treated with 1 \(\mu\)M MOR agonist methadone (MD) or MOR agonist DAMGO (DG) for 30 min at 37°C. Cells were fixed and stained as in Materials and Methods. In cells coexpressing DOR and MOR, images were captured consecutively from dual color channels (green and red fluorescence). Images are representative examples of multiple independent experiments. D and E, endocytosis of DOR/MOR heteromers, MOR homomers, and DOR homomers was analyzed by a biotin protection endocytosis assay in cells coexpressing FLAG-MOR and HA-DOR. Cells were biotinylated then treated with 1 \(\mu\)M methadone (MD) or DAMGO (DG) for 30 min. Endocytosed-protected homomeric and heteromeric receptors were separated by serial immunoprecipitation and resolved by SDS-PAGE as three populations: DOR homomers, MOR homomers, and DOR/MOR heteromers (see Materials and Methods). Total refers to the biotinylated receptor signal present in cells after initial labeling and without further manipulation; strip refers to biotinylated cells that were reacted with glutathione without other manipulations, demonstrating the efficiency with which biotin was cleaved from receptors and represents the background. Both total and strip serve as internal controls within each experiment. A representative immunoblot is shown for DOR/MOR heteromers (top), MOR homomers (middle), and DOR homomers (bottom).

Fig. 2. MOR antagonist NTB inhibits endocytosis in the DOR/MOR cell line. HEK293 cells stably expressing FLAG-MOR (A), HA-DOR (B), or FLAG-MOR and HA-DOR (C) were incubated with antibody recognizing the N-terminal epitope tags for 30 min at 37°C to label surface receptors. The DOR antagonist NTB (1 \(\mu\)M) was then added, followed 20 min later by the MOR agonist methadone (MD; 1 \(\mu\)M) or DAMGO (DG; 1 \(\mu\)M). After 30 min at 37°C, cells were fixed and stained as in Materials and Methods. In cells coexpressing DOR and MOR, images were captured consecutively from dual color channels (green and red fluorescence). Images are representative examples of multiple independent experiments.
DOR (Fig. 2C). This was unlikely to be a nonspecific effect of NTB on the MOR homomers, because NTB did not block endocytosis of the MOR in cells expressing only MOR (Fig. 2A), whereas it did block endocytosis of the DOR in response to the DOR agonist DPDP in cells expressing DOR alone (Supplemental Fig. 1C). As expected, pretreatment with a MOR-specific antagonist, CTAP (D-Phe-Cys-Trp-Arg-Thr-Pen-Thr-NH2), blocked endocytosis of MOR homomers in response to methadone in a dose-dependent manner (Supplemental Fig. 2A), as well as DOR/MOR heteromers (Supplemental Fig. 2B).

To further examine and quantify whether the effects of the DOR antagonist on MOR endocytosis were selective to the heteromer, we used serial immunoprecipitation and the modified version of the biotin protection assay to independently monitor the extent of endocytosis of DOR/MOR heteromers and MOR homomers. Cells coexpressing DORs and MORs were biotinylated with thio-cleavable biotin then pretreated with the DOR antagonist NTB (from 0.05 to 1 μM, 20 min) or vehicle, followed by methadone or DAMGO (1 μM, 30 min; Fig. 3). Residual surface biotin was then stripped, leaving only the protected endocytosed pool. We then selectively immunoprecipitated DOR/MOR heteromers and MOR homomers in the DOR/MOR-coexpressing cell line by serial immunoprecipitation. We found that both MOR homomers and DOR/MOR heteromers were endocytosed by both methadone and DAMGO (Fig. 3A, compare with untreated). The DOR antagonist NTB significantly inhibited DOR/MOR endocytosis in response to both methadone and DAMGO in a concentration-dependent manner compared with its effect on MOR endocytosis in the same cell line (Figs. 2A, top compared with bottom and 3B). NTB has a >100-fold higher affinity for DOR over MOR (Kᵢ = 0.013 nM for DOR and Kᵢ = 12 nM for MOR) (Raynor et al., 1994; Kim et al., 2001). Nevertheless, at high concentrations, selectivity can be lost (Kim et al., 2001). Consistent with this, endocytosis of MOR homomers was also inhibited by the highest concentration of the DOR antagonist NTB, albeit to a significantly smaller degree than DOR/MOR heteromer endocytosis (Fig. 3, A, top compared with bottom, and B). These effects were not unique to NTB. Pretreatment of cells coexpressing MOR and DOR with two other DOR antagonists, either BNTX (Kᵢ = 0.66 nM for DOR and Kᵢ = 18 nM for MOR) or NTI (Kᵢ = 0.02 nM for DOR and Kᵢ = 64 nM for MOR) (1 μM, 20 min), also significantly and selectively inhibited methadone- and DAMGO-induced endocytosis of the DOR/MOR heteromer compared with the MOR homomers present in the same cells (Fig. 4).

**NTB Inhibits Endocytosis of DOR/MOR Heteromers Without Inhibiting Signaling.** We next examined whether the DOR antagonist NTB affected signaling of DOR/MOR heteromers in response to MOR agonist. Cells expressing MOR or DOR alone or coexpressing DOR and MOR were pretreated with increasing concentrations of NTB followed by treatment with methadone (1 μM; Fig. 5A) or DAMGO (1 μM; Fig. 5B). Drug-mediated signaling was assessed by measuring Ca²⁺ release from intracellular stores (see Materials and Methods and Kostenis, 2001). Both methadone and DAMGO showed equivalent potency and efficacy in MOR and DOR/MOR cell lines and significantly reduced potency and efficacy in cells with only DOR (see Table 1). Although NTB inhibited endocytosis of the DOR/MOR heteromer (Fig. 3), we found that, even at the highest doses, NTB inhibited only...
pressing only MOR (Fig. 5), where NTB inhibited 63 ± 10% of the methadone- and 52 ± 4% of the DAMGO-mediated signaling. As expected, NTB was very effective at inhibiting both methadone-mediated (87 ± 2%) (Fig. 5A) and DAMGO-mediated (87 ± 12%) (Fig. 5B) signaling in cells expressing only DOR (Fig. 5A). NTB also effectively blocked signaling from the DOR-selective agonist DPDPE in cells expressing DOR and cells expressing DOR/MOR (Supplemental Fig. 3).

Therefore, NTB shows biased antagonism on the DOR/MOR heteromer by antagonizing only endocytosis but not DAMGO- or methadone-mediated signaling of the heteromeric complex (Figs. 3 and 5). It is noteworthy that this biased antagonism seems to be ligand-selective. Specifically, both BNTX and NTI not only blocked endocytosis of the DOR/MOR heteromer (Fig. 3), but also blocked methadone-mediated signaling on DOR/MOR cells (Fig. 6A; 81 ± 6 and 79 ± 6% of inhibition for BNTX and NTI, respectively). BNTX and NTI both also substantially blocked DAMGO-mediated signaling on DOR/MOR cells (Fig. 6B; 62 ± 8 and 48 ± 6% of inhibition for BNTX and NTI, respectively). As expected, all three DOR antagonists antagonized methadone- and DAMGO-mediated signaling in cells expressing only DOR (Fig. 6, C and D) (see Table 2).

**Fig. 5.** NTB shows significantly reduced antagonism of signaling on DOR/MOR heteromers. Cells coexpressing DOR/MOR (A), DOR only (●), or MOR only (▲) were pretreated with increasing concentrations of the DOR antagonist NTB for 20 min. Calcium release caused by chimeric G protein Δ6-Gαs-myristate was measured in a Flex apparatus upon stimulation with methadone (A) or DAMGO (B) (1 μM). Maximal effects for methadone (RFU) were: MOR (735 ± 51), DOR (562 ± 55), and DOR/MOR (956 ± 56). Data represent means ± S.E.M. of n = three to five experiments carried out in triplicate (two-way ANOVA and Bonferroni post-test; *p < 0.01; ***p < 0.001 DOR-only cells compared with DOR-only cells).

**Fig. 6.** Reduced antagonism of DOR/MOR heteromer signaling is NTB specific. A and B, cells coexpressing DOR/MOR were pretreated with increasing concentrations of NTB (●), BNTX (▲), or NTI (◇) for 20 min and signaling in response to methadone (A) or DAMGO (B) (both 1 μM) was measured as in Fig. 5. C and D, cells expressing only DOR are shown as a control for each antagonist upon methadone (C) or DAMGO (D) (both 1 μM) stimulation. Data represent means ± S.E.M. of n = four to five experiments carried out in triplicate (two-way ANOVA and Bonferroni post-test; *, p < 0.05; **, p < 0.01; ***p < 0.001 NTB compared with BNTX; ###, p < 0.001 NTB compared with NTI). See Table 2.

**TABLE 1**

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N.D., not determined.
* P < 0.05 and *** P < 0.001 DOR compared with DOR/MOR or MOR; # P < 0.05 MOR compared with DOR or DOR/MOR.
Endocytosis of DOR/MOR Heteromers Leads to Degradation. Several groups have reported that the MOR and DOR have different fates after endocytosis. However, the postendocytic fate of the DOR/MOR heteromer after endocytosis in response to MOR agonist is unknown. Consequently, we next assessed what postendocytic fate these heteromeric complexes followed, that of the MOR (recycling) or that of the DOR (degradation). Using a modified version of the biotin protection assay and serial immunoprecipitation, we assessed the postendocytic fate of MOR homomers and DOR/MOR heteromers in the DOR/MOR cell line (see Materials and Methods). As reported previously, endocytosed homomeric MORs were relatively stable even under constant agonist pressure for 2 h (Fig. 7, A bottom and B). In contrast, the endocytosed pool of DOR/MOR heteromers in the same cells were significantly more degraded after 2 h of agonist treatment (Fig. 7, A top and B). Thus, it seems that the DOR/MOR heteromer adopts the fate of the DOR after endocytosis in response to MOR agonist.

Discussion

Here, we show that the DOR/MOR heteromer is unique in its trafficking properties in several ways. First, the MOR agonists methadone and DAMGO can promote endocytosis of DORs when the receptors are coexpressed. Second, DOR-selective antagonists can selectively inhibit endocytosis of the DOR/MOR heteromer but not the MOR homomer in the same cell in response to activation by MOR agonists. Third, and especially intriguing, is our observation that the DOR-selective antagonist NTB inhibits DOR/MOR heteromer endocytosis in a dose-dependent manner, without antagonizing G protein-mediated signaling from this receptor. Fourth, we found that the DOR/MOR heteromer adopts the postendocytic fate of the DOR receptor after endocytosis in response to MOR agonist, and is, therefore, degraded rather than recycled. Consequently, NTB can selectively block down-regulation of DOR/MOR heteromers by inhibiting their endocytosis without blocking signaling from the heteromer.

Although opioid receptor heteromers are widely accepted to exist in heterologous expression systems, the existence, and functional significance, of receptor heteromers in native tissues is still a matter of controversy. It has been reported that MORs and DORs are not colocalized in the spinal cord of mice expressing a green fluorescent protein-tagged DOR (Scherrer et al., 2009). In opposition to these findings, two studies demonstrated that endogenous MORs and DORs can indeed colocalize in small dorsal root ganglia neurons of mice (Wang et al., 2010; He et al., 2011). It is noteworthy that several groups have reported increased expression of functional DORs after varying physiological stimuli, including chronic morphine treatment, stress, chronic inflammatory pain, and ethanol consumption (for review see Bie and Pan, 2007; Cahill et al., 2007), suggesting that the prevalence of DOR/MOR heteromers could change under these conditions. Indeed, chronic morphine treatment has been shown to promote up-regulation of an opioid receptor complex that is recognized by an antibody selective for DOR/MOR heteromers (Gupta et al., 2010). Taken together, we believe there is

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<th></th>
<th>Log IC50</th>
<th>Maximal Inhibition %</th>
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<td></td>
<td>Methadone</td>
<td>DAMGO</td>
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<tr>
<td>NTB</td>
<td>−6.8 ± 0.2</td>
<td>−6.9 ± 0.3</td>
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<tr>
<td>BNTX</td>
<td>−6.9 ± 0.2</td>
<td>−6.5 ± 0.3</td>
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<tr>
<td>NTI</td>
<td>−7.2 ± 0.3</td>
<td>−6.8 ± 0.2</td>
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**P < 0.01 and ***P < 0.001 NTB compared with BNTX; ### P < 0.001 NTB compared with NTI.

Fig. 7. Endocytosis of the DOR/MOR heteromer leads to the degradation of the receptor complex. Postendocytic stability of DOR/MOR heteromers and MOR homomers from the same cell line were analyzed by biotin protection-degradation assay. Cells coexpressing FLAG-MOR and HA-DOR were biotinylated, then left untreated or treated with 1 μM methadone for 30, 60, 90, and 120 min before stripping. Total refers to the biotinylated receptor signal present in cells after initial labeling and without further manipulation; strip refers to biotinylated cells that were reacted with glutathione without other manipulations, demonstrating the efficiency with which biotin can be cleaved from surface receptors and represents the background. The stability of the protected endocytosed DOR/MOR heteromers (top) and MOR homomers (bottom) was assessed by serial immunoprecipitation followed by SDS-PAGE and streptavidin overlay (see Materials and Methods) at the time points stated. B, quantification of experiments in A is shown for DOR/MOR heteromers versus MOR homomers. Histogram shows the mean stability of the biotinylated endocytosed receptors relative to the endocytosed pool seen after 30 min of stimulation. Shown are the mean ± S.E.M. n = 4 to 10 independent experiments (two-way ANOVA, Bonferroni post-test: *, p < 0.05; **, p < 0.01; ***, p < 0.001).
increasing evidence that the DOR/MOR heteromer is a functional receptor unit in vivo. Thus, understanding the unique properties of this receptor heteromer could help reveal its functional role.

Once endocytosed, GPCRs can take distinct trafficking routes that further shape the signaling response. GPCRs are either 1) rapidly targeted to the lysosomes for its degradation, resulting in complete termination of receptor signal activity, 2) rapidly recycled back to the plasma membrane, resulting in resensitization and signal recovery, or 3) are retained in endosomes, traversing the degradative and/or recycling pathways at a much slower rate (for review see Marchese et al., 2008). Several groups have demonstrated that the MOR and DOR have different postendocytic fates. For example, we have seen that DORs heterologously expressed in HEK293 cells exhibitpronounced down-regulation within 2 h of exposure to agonist, whereas MORs expressed at similar levels are primarily recycled and significantly more stable (Whistler et al., 2002). The MOR has also been shown to resensitize in rat brain slices (Alvarez et al., 2002) and apparently in humans as well (Szeto et al., 2001). Thus, there remained the intriguing question as to the postendocytic fate of the DOR/MOR heteromer. We envisioned at least three possible outcomes for the DOR/MOR heteromers: 1) MORs within the heteromeric complex would take DORs back to the plasma membrane, converting DOR into a recycling receptor, 2) DORs would drag MORs in the heteromer to the lysosome, converting MOR into a degrading receptor, or 3) the receptors would separate after endocytosis and travel on their own. Here, we found that MORs that were heteromerized and coendocytosed with DORs were degraded more rapidly and to a further and greater extent than homomeric MORs in the same cells after endocytosis in response to MOR agonist. Thus, our data suggest either that the heteromer is a stable unit after endocytosis and that degradation “wins” in the battle for the fate of the heteromer, or coendocytosis of the MOR and DOR as a heteromer somehow marks the MORs in this complex for degradation. This codegradation of MOR and DOR would not be expected to occur under conditions where the MOR (and/or DOR) is not significantly endocytosed, such as after activation by morphine (Keith et al., 1998; Arttamangkul et al., 2008) (Supplemental Fig. 1). Thus, it is unlikely that the codegradation of MOR together with DOR is responsible for tolerance to the antinociceptive effects of chronic morphine as has been proposed (He et al., 2011).

Several groups have shown that heteromerization of MOR and DOR changes binding and signaling properties (Gomes et al., 2000, 2004; Kabli et al., 2010; Yekkirala et al., 2010). Therefore, we cannot rule out the possibility that this may be the case for NTB used in this study as well. Nevertheless, our observation that endocytosis of the DOR/MOR is blocked by NTB indicates that the ligand is engaging the heteromeric target and displaying biased antagonism. This biased antagonism is ligand-selective [BNTX and NTI, for example, block not only trafficking but also signaling (see Figs. 4 and 6)].

DOR antagonists are reported to be more effective for the reduction of tolerance to morphine in vivo (for review see Ananthan, 2006). Consequently, there has been much interest in combining DOR antagonists with MOR agonists to delay or reduce the development of opioid tolerance and dependence (Abdelhamid et al., 1991; Fundytus et al., 1995; Hepburn et al., 1997; Gomes et al., 2004; Daniels et al., 2005). It is noteworthy that our results suggest that the efficacy of these combination therapies probably will show ligand bias, and any interpretation of these data must include a consideration of the effects of these drug combinations not only on the signaling, but also on the trafficking of the MOR, DOR, and DOR/MOR heteromers.

Authorship Contributions

Participated in research design: Milan-Lobo and Whistler.

Conducted experiments: Milan-Lobo.

Wrote or contributed to the writing of the manuscript: Milan-Lobo and Whistler.

Other: Milan-Lobo and Whistler acquired funding for the research.

References


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