Regulation of the Human Delta-Opioid Receptor by Alkaloids: Different Roles of Arrestins

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Abstract

Study background: We previously revealed that the arrestin 2 was differentially involved in the regulation of the human delta opioid receptor (hDOR) by peptidic and alkaloid agonists. In the present study, we examined whether i) hDOR regulation by two related alkaloid agonists (etorphine and morphine) involved the same arrestins and ii) similar arrestin-dependent mechanisms occur upon short- and long-term agonist exposure.

Methods: The human neuroblastoma SK-N-BE cells, endogenously expressing hDOR, were transfected with wild-type or mutant arrestins. Using shRNA, we also generated a clonal cell line depleted in arrestin 2, the only arrestin isoform expressed in this cell line. Cells were then exposed to either morphine or etorphine for short-(1 h) or long-term (18 h), then receptor desensitization was examined on the cAMP pathway and internalization was visualized by confocal microscopy.

Results: In arrestin 2-depleted cells, we observed a strong reduction of desensitization after 18 h of morphine exposure but not with etorphine. Over-expression of wild-type or mutant arrestins produced an opposite modulation of receptor desensitization induced by morphine and etorphine both upon short- and long-term exposure. Confocal fluorescence microscopy experiments did not reveal any strong impact on receptor internalization when different arrestins were over-expressed.

Conclusion: Our results showed that i) two related opioid alkaloid agonists produce hDOR desensitization by different mechanisms ii) the involvement of arrestins in hDOR desensitization depends on the duration of agonist exposure.

Keywords: Arrestin; Camp pathway; Desensitization; Internalization; Morphine; Opioid receptor

Abbreviations: Arr 2: Arrestin 2 (or beta-arrestin 1); Arr 3: Arrestin 3 (or beta-arrestin 2); Deltorphin I: Tyr-D-Ala-Phe-Asp-Val-Val-Gly; NH2; DPDPE: D-Penicillamine(2,5)-enkephalin; GPCRs: G protein-coupled receptors; hDOR: Human delta-opioid receptor; KO: Knock-out; MOR: Mu opioid receptor; shRNA: Short hairpin RNA

Introduction

It is well admitted that the opioid system plays a central role in pain control but sustained exposure to exogenous opioids such as morphine that are used for their analgesic properties provokes tolerance in human and in animal (see for review) [1]. Tolerance that is defined by a decrease of the drug response following acute or chronic exposure, is closely related to opioid receptor desensitization observed in vitro. Mechanisms of both tolerance and desensitization have been extensively studied and appeared complex but arrestins were suggested to have a pivotal role as previously reviewed [2]. In arrestin 3-knock out (KO) mice, mu opioid receptor (MOR) desensitization and tolerance were decrease upon chronic morphine exposure compared to wild-type (WT) mice [3,4]. Those data are in good agreement with the canonical model of G protein-coupled receptors (GPCRs) regulation, including opioid receptors, indicating that arrestins are negative regulators of signaling by promoting uncoupling and endocytosis of cell surface receptors [5]. There is now accumulating evidence showing that opioid receptors are differentially regulated upon opioid agonist activation (see for review) [6]. This was demonstrated for interactions between DOR and MOR with G proteins and arrestins [7,8]. While different ligands could bind to opioid receptors, several laboratories, including ours, reported difference in their ability to regulate such receptors [9-13]. So, in the present study, we adressed the role of arrestins in hDOR regulation upon short- and long-term activation by two related alkaloid agonists morphine and etorphine. This study was conducted in the SK-N-BE cells, endogenously expressing hDOR [14,15]. To answer to those questions, we studied both desensitization, on the cAMP pathway, and receptor internalization in a cellular context over-expressing WT or mutant arrestins or in arrestin-depleted cells.

Material and Methods

Cell culture

SK-N-BE cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) (Sigma, Saint-Quentin Fallavier, France),...
supplemented with 10% fetal calf serum (FCS) (Biowest Nuaille, France), 1% antibiotic–antimycotic mixture (Sigma), and 2 mM L-glutamine at 37 °C in a water-saturated atmosphere containing 5% CO2.

Plasmids and transfection

Plasmids containing arrestin 2-GFP, arrestin 3-GFP, arrestin 2319-418, GFP, R169E arrestin 2-GFP, FLAG-tagged hDOR were kindly provided by Prof. S. Cotecchia (Université de Lausanne, Switzerland), Dr. A. Benmerah (Institut Cochin, INSERM U1016, UMR 8104, Université Paris-Descartes, France) Prof. N.W. Bunnett (University of California, San Francisco, USA), Prof. V.V. Gurevich (Vanderbilt University Medical Center, Nashville, USA), and Prof. M. Bouvier (Université de Montréal, QC, Canada), respectively. Transfection and the generation of the different clonal cell lines were previously described [16,17]. The different clonal cell lines were cultured under selection using 1 mg.ml⁻¹ geneticin (G418, Sigma Aldrich).

Western blotting

Expression of WT and mutant arrestins and the inhibition of the endogenous arrestin 2 expression by shRNA was evaluated by western blot as previously reported [17]. Whole cell lysates were prepared from wild-type SK-N-BE cells and the clonal cell lines. Cells were harvested by centrifugation (100 g, 5 min) and suspended in lysis buffer (10 mM Tris–HCl, 1 mM EDTA, 0.1% (v/v) Triton-X100, pH 7.4), then sonicated. After a centrifugation (15 min at 20,000 g and 4°C), protein concentration from the supernatant was determined by the Bradford assay and equal amounts were separated on 10% (w/v) acrylamide gels by SDS-PAGE. After electroblotting, nitrocellulose membranes were incubated with anti-arrestin 2 and 3 antibody (kindly provided by Prof. S. Cotecchia (Université de Lausanne, Switzerland), Dr. A. Benmerah (Institut Cochin, INSERM U1016, UMR 8104, Université Paris-Descartes, France) Prof. N.W. Bunnett (University of California, San Francisco, USA), Prof. V.V. Gurevich (Vanderbilt University Medical Center, Nashville, USA), and Prof. M. Bouvier (Université de Montréal, QC, Canada), respectively). Transfection and the generation of the different clonal cell lines were previously described [16,17]. The different clonal cell lines were cultured under selection using 1 mg.ml⁻¹ geneticin (G418, Sigma Aldrich).

Measurement of intracellular cAMP

Inhibition of adenyl cyclase was determined by measuring [¹H]cAMP accumulation as described previously [18]. This radioligand is a lipophilic antagonist that binds both cell surface and intracellular receptors. Cells were seeded in 24-well plates at a density of 100,000 cells/well and were allowed to grow for 48 h. Before binding experiments, cells were washed with DMEM/20 mM Heps for 5 min and then incubated for 30 min at 37°C with appropriate concentrations of [¹H]diprenorphine (0.05-5 nM) in a 0.3-ml final volume of 50 mM Tris–HCl/1% BSA (w/v), pH 7.4. Total and nonspecific binding were determined in the absence or in the presence of 20 µM levorphanol, respectively. The medium was rapidly removed and cells were harvested in 200 µl of 1 N NaOH and placed into vials in the presence of 3 ml of scintillation cocktail (PerkinElmer Life and Analytical Sciences). Each determination was carried out in triplicate. Scatchard analysis (supplemental data) was performed using SigmaPlot software to calculate the dissociation constant Kd (nM) and the receptor maximum binding capacity Bmax (fmol.mg⁻¹ of protein) values.

Inhibition of adenylyl cyclase was determined by measuring [¹H]cAMP accumulation as described previously [18]. Cells were seeded in 24-well plates at a density of 50,000 cells per well in a culture medium supplemented with 0.6 mCi [¹H]adenine and incubated overnight. cAMP accumulation was determined in the presence of isobutylmethylxanthine, forskolin and in the absence or in the presence of agonists. After 5 min at 37°C, the reaction was stopped by addition of 5% (w/v) trichloroacetic acid. The [¹H]cAMP content of each well was isolated by chromatography on acid alumina columns, mixed with 8 ml of scintillation mixture (Pico-Fluor-40, PerkinElmer), before assaying in a scintillation counter (PerkinElmer). Maximal inhibitory levels of opioid agonists were determined for each clonal cell line at 0.1X, 1X and 10X where X corresponds to the concentration producing the maximum response in the WT SK-N-BE cells SK-N-BE cells [11,19]. For desensitization experiments, cells were pretreated or not (naïve) either for 1 or 18 h with the concentration of etorphine or morphine producing the maximal inhibition of cAMP accumulation in each clonal cell line (Table 1). Then, the hDOR-induced adenyl cyclase inhibition was measured for 5 min at 37°C without removing the medium to avoid adenyl cyclase superactivation in the presence of 1 mM isobutylmethylxanthine alone (basal activity) or in combination with 40 µM forskolin (FSK) (stimulated activity). We checked that addition of freshly prepared agonist (etorphine or morphine) after 1 or 18 h exposure did not promote additional inhibition of cAMP accumulation demonstrating that the reduction of cAMP inhibition was due to hDOR desensitization and not agonist degradation. All experiments were carried out in triplicate and repeated at least three times with similar results.

Internalization studies by confocal microscopy

Localization of both hDOR (red) and arrestins (green) were visualized as described previously [17]. For image analysis, the freeware ImageJ 1.47v was used to quantify the cell surface receptors (http://imagej.nih.gov/ij). We determined the integrated density of cell surface labeling using the TRITC channel both in GFP-positive and negative cells.

Docking studies

For each docked compound a preliminary calculation on its protonation state at pH 7.4 was carried out using standard tools of the ChemAxon Package (http://www.chemaxon.com/) and the majority of the molecules protonated on nitrogen (corresponding to among 98% for morphine as well as for etorphine) at this pH was used for docking studies. The crystallographic coordinates of hDOR used for docking studies were obtained from X-ray structure of the naltrindole/delta-opioid receptor complex (PDB ID 4N6H, a structure refined to 1.8 Å with an R Factor of 17.3%) [20]. Docking of morphine and etorphine into hDOR was carried out by means of the GOLD program with the default parameters [21]. This program applies a genetic algorithm to explore conformational spaces and ligand binding modes. To evaluate the proposed ligand poses, the ChemPLP, GoldScore and ChemScore fitness functions were applied (see the scores in supplemental data). The binding site in the hDOR model was defined as a 10 Å sphere from the co-crystalized ligand natrindole using the detection cavity algorithm.

Statistical analysis

All results are expressed as the mean ± standard error of the mean (S.E.M.) of n experiments. ANOVA (GraphPad Prism 4.0®) followed either by the Dunnett or the Bonferroni Test or Student t-test, when appropriate, were used to determine the statistical significance.

Results

Expression of WT or mutant arrestins in SK-N-BE cells

Stable expression of arrestins 2 or 3, the constitutively active mutant R169E arrestin 2 or a dominant negative (DN) mutant (arrestin 229-
Etorphine and morphine promote adenylyl cyclase inhibition in the different clonal cell lines. The concentration ([L]) and the maximum cAMP inhibition (Imax) produced were determined in BE-WT and in the different clonal cell lines. Data are the means ± S.E.M. performed in triplicate. Significance compared to BE-WT cells is indicated: *, P<0.05 or **, P<0.01, One-way ANOVA followed by Dunnett’s multiple comparison tests.

Short- and long-term hDOR desensitization promoted by etorphine or morphine

Then, we examined the impact of arrestin over-expression or KO of endogenous arrestin 2 on the endogenous hDOR level and the ability of both morphine and etorphine to inhibit adenylyl cyclase. We showed no significant modifications of Bmax values while over-expression of arrestin 3-GFP slightly but significantly increase Kd value by about 3-fold (Table 1). In functional studies, we observed that arrestin 3-GFP over-expression and KO of endogenous arrestin 2 significantly decreased the maximal inhibition produced by etorphine while the production of MM sequence increased morphine-induced cAMP inhibition and the dominant negative mutant of arrestin reduced morphine potency (Table 2). Then, we selected concentrations of etorphine and morphine producing almost similar and maximal inhibition of adenylyl cyclase to promote hDOR desensitization after short- and long-term exposure in the various clonal cell lines. The inhibition of cAMP accumulation produced by each agonist was normalized to 100% in control (Figure 2A and 2B). Only the over-expression of arrestin 2-GFP significantly increased hDOR desensitization upon long-term etorphine treatment which was evidenced by a superactivation of adenylyl cyclase (Figure 2C). In contrast, when WT and mutant of arrestins were over-expressed we rather observed a decrease in adenylyl cyclase superactivation after 18 h morphine exposure (Figure 2D). When endogenous arrestin 2 expression was impaired by shRNA, we observed a major reduction of desensitization by comparison with the MM clonal cell line but only upon morphine treatment (Figure 2D).

Etorphine and morphine promote hDOR endocytosis upon short- and long-term exposure

To study the influence of arrestins on hDOR internalization upon short- and long-term exposure to etorphine or morphine, SK-N-BE cells stably expressing the FLAG-tagged hDOR were transiently transfected with WT, mutants of arrestins or shRNA directed against the endogenous arrestin 2. Then, we monitored the FLAG-tagged hDOR (red) and arrestins-GFP (green) localization by confocal microscopy and semi-quantitative analyses were performed by determining plasma

![Figure 1: Arrestins expression in the BE-WT and the clonal cell lines. (A) Whole cell lysates were prepared from wild type SK-N-BE cells (lane 1), arrestin 2-GFP (lane 2), arrestin 3-GFP (lane 3). R169E-arrestin2-GFP (lane 4) or the dominant negative mutant arrestin 2 319-418-GFP (lane 5) clonal cell lines. (B) Whole cell lysates were also prepared from clonal cell lines expressing shRNA directed against the endogenous arrestin 2 (lane 1) or a mismatch sequence (lane 2). Proteins were resolved by SDS/PAGE and expression of both endogenous and exogenous arrestins as well as actin were determined.](image)

![Figure 2A: Etorphine promotes hDOR desensitization.](image)

![Figure 2B: Morphine promotes hDOR desensitization.](image)

![Figure 2C: Short-term exposure to arrestins.](image)

![Figure 2D: Long-term exposure to arrestins.](image)

Table 1: Etorphine and morphine promote adenylyl cyclase inhibition in the different clonal cell lines. The concentration ([L]) and the maximum cAMP inhibition (Imax) produced by etorphine (Eto) and morphine (Mor) were determined in the BE-WT and in the different clonal cell lines. Data are the means ± S.E.M. of 3-5 independent experiments performed in triplicate. Significance compared to BE-WT cells is indicated: *, P<0.05 or **, P<0.01, One-way ANOVA followed by Dunnett’s multiple comparison tests.

Table 2: Endogenous hDOR level in the different clonal cell lines. Endogenous hDOR expression was determined in the different clonal cell lines. Data are means ± S.E.M. of 2-3 different experiments performed in triplicate. Significance compared to BE-GFP cell line is indicated: *, P<0.05. One-way ANOVA followed by Dunnett’s multiple comparison tests.
membrane labelling using Image J as previously reported [22]. In naïve cells transfected (GFP+) or not (GFP-) with the different plasmids containing the WT or the mutants of arrestins, the hDOR was mainly localized at the plasma membrane (Figure 3, naïves). When cells were pretreated either with etorphine or morphine for 1 h, we observed a strong internalization of the opioid receptor (Figure 3, Eto 1 h and Mor 1 h) without any significant effect of arrestin expression (Figure 4). In the shRNA clonal cell line, we showed that decrease of endogenous arrestin 2 expression significantly reduced etorphine-induced hDOR internalization while it increased receptor internalization in the case of morphine but very slightly (Figure 4, Eto 1 h and Mor 1 h). When the time of pretreatment was extended to 18 h, we observed a strong decrease of hDOR immunolabeling for both alkaloid agonists (Figures 3 and 4) without any significant difference between GFP+ and GFP except in the dominant negative mutant clonal cell line where morphine-induced hDOR endocytosis was significantly but slightly decreased (Figure 4, Mor 18 h).

Discussion
The present study showed that arrestins produced a different hDOR regulation upon activation by two related alkaloid agonists, etorphine and morphine. Upon short-term exposure, both agonists promote an arrestin-independent desensitization and internalization as demonstrated in shRNA cell line. While this result contrasts with the canonical model of GPCR regulation, previous reports also showed the lack of arrestin implication in morphine-induced MOR desensitization [23,24]. This is unlikely due to an absence of a total inhibition of arrestin 2 expression by shRNA since we previously reported the role of this protein in DPDPE- and deltorphin I-mediated hDOR desensitization using the same clonal cell line [17]. Those data clearly show that upon short-term exposure morphine and etorphine, on one hand, and DPDPE and deltorphin I, on the other hand, are biased agonists at the hDOR to recruit arrestin 2 for desensitization. Receptor desensitization could be due to sequestration as previously demonstrated for etorphine [18] but involvement of other actors such as RGS [25] or kinases [26] cannot be ruled out. Since expression of exogenous WT or mutant of arrestins were not equivalent between the different clonal cell lines (Figure 1) and over-expression of arrestins could affect the binding properties of hDOR (decrease of affinity by arrestin 3-GFP) (Table 2), their comparison would therefore be misleading. The increase of Kd value observed in the arrestin 3-GFP clonal cell line could explain the decrease of the maximal inhibitory effect of etorphine; but this was not detected upon morphine treatment. Furthermore, reduction of endogenous level of arrestin 2 also decreased the I₅₀ value for etorphine indicating that modulation of endogenous and exogenous arrestin levels had a complex impact on hDOR. However, in a given clonal cell line, comparison between the two alkaloid agonists is possible. When over-expressing arrestins, we observed their differential involvement in etorphine- and morphine-induced receptor desensitization. In a cellular context of arrestin 2 over-expression, we showed an increase of receptor desensitization both at short- and long-term exposure only for etorphine suggesting a poor interaction between those proteins that could be overcome by increasing the level of arrestin 2. This weak interaction is not related to the lack of receptor phosphorylation since etorphine enhanced Ser363 phosphorylation and promoted a GRK2-dependent desensitization [26]. In constrast, arrestin 2 was unable to promote receptor short-term desensitization upon morphine exposure even when its level was over-expressed. This indicates that the conformation of the receptor-morphine complex would be a poor substrate for arrestin 2 binding. When considering

![Figure 2: Role of arrestins in short- and long-term desensitization of hDOR. Wild type SK-N-BE cells and the different clonal cell lines were pretreated or not (naive) for 1 or 18 h with etorphine (A and C, Eto) or morphine (B and D, Mor). Agonist-induced inhibition of cAMP in naive cells was referred as 100%. Data are means ± S.E.M. of 3-9 different experiments performed in triplicate. *, P<0.05 and **, P<0.01, one-way ANOVA followed by Dunnett’s test compared to BE-WT cells, $$, P<0.01, t-test compared to shRNA MM.](image-url)
Figure 3: Role of arrestins in hDOR internalization. SK-N-BE cells stably over-expressing the FLAG-tagged hDOR were transiently transfected with arrestin 2-GFP (Arr 2), arrestin 3-GFP (Arr 3), the constitutively active mutant R169E-arrestin2-GFP (R169E), the dominant negative mutant arrestin 2319-418-GFP (319-418), a plasmid producing both GFP and shRNA directed against the endogenous arrestin 2 (shRNA) or a mismatch sequence (MM). Cells were treated or not (naïve) with etorphine (Eto) or morphine (Mor) for either 1 or 18 h. After treatment, cells were fixed and immunostained with the anti-FLAG M2 antibody. Immunoreactivity was revealed with a TRITC-conjugated secondary antibody. Localization of the hDOR (in red) and arrestins or shRNA (in green) was observed by confocal microscopy at a 60X lens. Images are representative of 3 to 4 independent experiments.

Figure 4: Quantification of hDOR internalization. Integrated density corresponding to cell surface hDOR labelling was determined as described in Material and Methods both in GFP-positive (expressing WT or mutant of arrestins or shRNA) and negative cells (non-transfected cells). Data are means ± S.E.M. of 3 independent experiments. **, P<0.01, ***, P<0.001, two-way ANOVA followed by Bonferroni’s test compared to GFP-negative cells.
short-term activation, we showed that arrestin 3 and the constitutively active mutant inhibit receptor desensitization upon morphine but not etorphine exposure probably by promoting a significant hDOR internalization and recycling as previously demonstrated [27]. Morphine, but not etorphine, would promote conformational receptor changes that could enable efficient interactions between the hDOR and the arrestin 3 or the constitutively active mutant. Such a preferential interaction between hDOR and arrestin 3 under morphine activation compared to etorphine was not previously observed [8]. However, it is highly difficult to compare our data with this study since those authors over-expressed hDOR to a level about 100 to 200-fold higher than in our cellular model and failed to detect any significant interaction between the hDOR and arrestin 2-GFP. Furthermore, it is well admitted that receptor over-expression increases spare receptors which consequently reduced desensitization [28]. The great difference in receptor expression (and in spare receptors) between our study and others could be a major factor to explain discrepancies about the role of arrestin in the desensitization process. While our data do not support any role of arrestin 3 in hDOR regulation, others showed that this protein preferentially interacts with nDOR [29] and promotes the mouse receptor desensitization [30]. Such discrepancy would be related to the substantial differences in the carboxy-terminal tail between the mouse and the human DOR, a critical region for interactions with arrestins [31]. After long-term agonist exposure, we observed a complete desensitization or a slight increase in adenylyl cyclase activity for morphine pretreatment. This was correlated with a global decrease of DOR immuno-reactivity. Prolonged activation of DOR by morphine or etorphine was reported to cause receptor down-regulation in different brain regions [32,33] and also in the SK-N-BE cell line in the case of etorphine [18,34]. When examining the potential role of arrestins in long-term receptor regulation, we found that decrease of endogenous arrestin 2 expression greatly reduced desensitization induced by morphine but not by etorphine. This indicates that i) hDOR desensitization would involve different molecular mechanisms upon short- and long-term treatment by morphine ii) etorphine and morphine, two related alkaloid agonists, induce desensitization but via different mechanisms implicating or not the arrestin 2. Interestingly, morphine (see for review) [35] and etorphine [26] were shown to promote acute desensitization of opioid receptors in a PKC-dependent manner. So, we can hypothesize that in such conditions arrestins would not be required. However, under long-term activation, morphine would promote a significant hDOR phosphorylation to enable the recruitment of arrestin 2. The reduction of hDOR desensitization observed after 18 h morphine exposure in arrestin 2-depleted cells is not related to a decrease of receptor internalization. This is not surprising since previous studies reported no correlation between opioid receptor internalization and desensitization [19,36]. This also suggests that very few cell surface receptors would enable an almost complete inhibition on the cAMP pathway and/or confocal microscopy is not an accurate tool to detect few receptors in such conditions. Furthermore, evaluation of hDOR level by binding experiments and immuno-labeling showed discrepancies indicating that the two methods have their own limitations and require a careful interpretation. Rather than a classical role in uncoupling and internalization, our data suggest that arrestin 2 would inhibit recycling and resensitization upon long-term morphine treatment as recently demonstrated for the MOR and arrestin 3 [37]. As observed for short-term pretreatment, arrestin 2 over-expression potentiates DOR desensitization upon 18 h etorphine which was characterized by a superactivation of adenylyl cyclase; this confirms that arrestin 2 is the preferred partner of the hDOR when activated by etorphine but not morphine. Concerning long-term morphine exposure, over-expression of either WT or mutant arrestins was shown to decrease adenylyl cyclase superactivation. Recent data

Figure 5: Modeling of naltrindole, etorphine and morphine binding on hDOR. X-ray structure of naltrindole co-crystallized with the hDOR (A) in comparison with docking configurations of morphine (B) and etorphine (C). The ligands and selected side chains are shown in the stick representation and the sodium ion (purple) and selected water molecules (red) are shown as balls. Polar interactions are presented as dash-lines. This figure was made with PYMOL v1.3 (DeLano Scientific, 2002, San Carlo, USA).
indicate that Src mediated-MOR phosphorylation would recruit Ras/Raf-1 proteins which in turn increase adenyly cyclase activity [38,39]. Over-expression of arrestins, which were shown to interact with Src [40], could interfere with such tyrosine phosphorylation and inhibit adenyly cyclase superactivation.

Recently, Fenali et al. identified in the crystal structure of the hDOR that the sodium ion and Asn131 played a major role in the constitutive activation of arrestin [20]. So, we hypothesized that morphine or etorphine could differentially modify the interaction network around sodium and/or Asn131 favouring arrestin interactions. However, based on the high resolution hDOR structure bound to naltrindole, our in silico analysis failed to reveal any significant modification at Asn131 upon either morphine or etorphine binding (Figure 5). This latter result does not challenge the biased agonism theory but could rather suggest that other regions of the receptor would be involved in the arrestin interactions. Indeed, several residues of the complex etorphine-mouse DOR were identified by 3D modeling that could modulate auxiliary sites for receptor partners such as arrestins [41]. Furthermore, the carboxy-terminal tail of the DOR, which was not included in the crystal structure, represents a putative region involved in the differential interaction between receptor and arrestins upon etorphine or morphine exposure.

Conclusions

It is now well admitted that opioid receptor desensitization plays a major role in tolerance. Our study showed that different molecular mechanisms occurred in short- and long-term hDOR desensitization upon morphine treatment. While both alkaloid agonists, etorphine and morphine, produced an arrestin 2-independent short-term hDOR desensitization, the endogenous arrestine 2 would rather inhibit recycling and potentiate desensitization after 18 h morphine exposure but not for etorphine. Our data also suggest that hDOR would poorly interact with endogenous arrestin 2 but when over-expressing arrestin, we could reveal a different role of WT or mutant arrestin under alkaloid agonists exposure. Our study shows that two closely related opioid agonists promote a complex receptor regulation and supports the notion of arrestin-biased agonism at hDOR.

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References


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