Detection of Heteromers Formed by Cannabinoid CB₁, Dopamine D₂, and Adenosine A₂A G-Protein-Coupled Receptors by Combining Bimolecular Fluorescence Complementation and Bioluminescence Energy Transfer

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Functional interactions in signaling occur between dopamine D₂ (D₂R) and cannabinoid CB₁ (CB₁R) receptors, between CB₁R and adenosine A₂A (A₂AR) receptors, and between D₂R and A₂AR. Furthermore, direct molecular interactions have been reported for the pairs CB₁R-D₂R, A₂AR-D₂R, and CB₁R-A₂AR. Here a combination of bimolecular fluorescence complementation and bioluminescence energy transfer techniques was used to identify the occurrence of D₂R-CB₁R-A₂AR hetero-oligomers in living cells.

KEYWORDS: G-protein-coupled receptors, adenosine A₂A receptor, cannabinoid CB₁ receptor, dopamine D₂ receptor, heterotrimers, oligomers of three different protomers, bimolecular fluorescence complementation, bioluminescence energy transfer, BRET

INTRODUCTION

The mechanism of action responsible for the motor-depressant effects of cannabinoids, which operate through centrally expressed cannabinoid CB₁ receptors (CB₁R), is still a matter of debate. The cellular and subcellular localization of CB₁R and D₂ receptors (D₂R) in the basal ganglia have been described in
The evidence suggests a colocalization of striatal CB₁R and D₂R predominantly in the soma and dendrites of the GABA enkephalinergic neurons and also in corticostriatal glutamate terminals where A₂A receptors (A₂AR) are also present[5,6,7,8,9,10]. Functional interactions between CB₁R and A₂A R or D₂R receptors that are relevant for striatal function have been reported. Thus, CB₁R and A₂AR form heteromers in cotransfected HEK-293T cells and in rat striatum[11]. In a human neuroblastoma cell line, CB₁R signaling was found to be completely dependent on A₂AR activation. Accordingly, blockade of A₂AR counteracted the motor-depressant effects produced by the intrastriatal administration of a CB₁R agonist[11]. The existence of CB₁R-D₂R heteromers has been demonstrated in transfected cell lines by coimmunoprecipitation[12] and by fluorescence resonance energy transfer (FRET) experiments[13]. Antagonistic CB₁R-D₂R interactions have been discovered at the behavioral level[13,14,15,16]. In rats, the CB₁R receptor agonist CP 55,940 at a dose that did not change basal locomotion was able to block quinpirole-induced increases in locomotor activity. In addition, not only the CB₁R antagonist rimonabant, but also the specific A₂AR antagonist MSX-3, blocked the inhibitory effect of CB₁R agonist on D₂-like receptor agonist-induced hyperlocomotion[13]. Taken together, these results give evidence for the existence of antagonistic CB₁-D₂ receptor-receptor interactions within CB₁R-D₂R heteromers in which A₂AR may also participate.

A₂AR-D₂R was one of the first reported heteromers[17]. A close physical interaction between both receptors has been shown using coimmunoprecipitation and colocalization assays[17], and FRET and bioluminescence resonance energy transfer (BRET) techniques[18,19,20]. At the biochemical level, two kinds of antagonistic A₂AR-D₂R interactions have been discovered that can explain the A₂AR-D₂R interaction observed at both the functional and behavioral levels[9,21,22,23]. First, by means of an intramembrane interaction, i.e., by intramolecular cross-talk within the heterodimer, stimulation of A₂AR decreases the affinity of D₂R for their agonists[24]. Second, the stimulation of D₂R, a Gi/o protein–coupled receptor, inhibits the cAMP accumulation induced by the stimulation of the Gs/olf protein–coupled A₂AR[17]. Therefore, it has been suggested that the A₂AR-D₂R interaction cross-talk in the central nervous system may provide new therapeutic approaches for Parkinson’s disease, schizophrenia, and drug addiction[23,25].

Since trimers formed by cannabinoid, adenosine, and dopamine receptors were suspected, strategies to detect them were developed in our laboratory. In one of them, trimers were detected by sequential application of BRET and FRET[26]. In this paper, the formation of hetero-oligomer complexes formed by cannabinoid CB₁, dopamine D₂, and adenosine A₂AR G-protein-coupled receptors (GPCRs) is confirmed by another technique[27], which consists of combining bimolecular fluorescence complementation and bioluminescence energy transfer.

**MATERIALS AND METHODS**

**Cell Culture**

HEK-293T cells were grown in Dulbecco’s modified Eagle’s medium (DMEM) (Gibco) supplemented with 2 mM L-glutamine, 100 U/ml penicillin/streptomycin, and 5% (v/v) heat inactivated fetal bovine serum (FBS) (all supplements were from Invitrogen, Paisley, Scotland, U.K.). Cells were maintained at 37°C in an atmosphere of 5% CO₂, and were passaged when they were 80–90% confluent, i.e., approximately twice a week.

**Fusion Proteins and Expression Vectors**

Full-length YFP was subcloned in the XhoI site of pcDNA3.1 vector (Invitrogen). The N-terminal truncated version of YFP, named nYFP (amino acids 1 to 155), was made by PCR amplification and cloning into the XhoI site of pcDNA3.1 using the following primers: FnYFP (5’-
CCGCTCGAGACCATTGGAGCAAGGGCGAGGAC-3') and RnYFP (5'-CCGTCTAGATCGCCATGATATAGCTTG-3'). Also, a C-terminal truncated version of YFP, named cYFP (amino acids 155 to 231), was made using the same strategy and the following primers:FcYFP (5'-CCGCTCGAGACCATGGACAAGCAGAAGAACGGC-3') and RcYFP (5'-CCGTCTAGATTACTTGTACAGCTCGTCCAT-3'). Gaα cloned in SFV1 vector (generously given by H. Vogel, Ecole Politecnique Federal de Laussane, Switzerland) or Gaγ and Gbβ cloned in pEYFP-C1 vector (generously provided by S. Cotecchia, Department of Pharmacology and Toxicology, University of Lausanne, Switzerland) were amplified to miss their stop codons using sense and antisense primers harboring unique NheI and BamHI sites to clone Gaα and Gbβ in pcDNA3.1-nYFP and pcDNA3.1-cYFP, respectively, and HindIII and BamHI sites to clone Gaγ in Rluc vector (pRluc-N1 PerkinElmer, Wellesley, MA). The amplified fragments were subcloned to be in-frame with the multiple cloning site of the vectors to give the plasmids Gaα-nYFP, Gbβ-cYFP, and Gaγ-Rluc, respectively.

All plasmids express luminescent or part of fluorescent proteins on the C-terminal ends of proteins. The human cDNAs for A2aR, CB1R, D2R, or D4.4R cloned in pcDNA3.1 were amplified without their stop codons using sense and antisense primers harboring unique NheI and BamHI sites to clone A2aR, CB1R, and D2R in pcDNA3.1-cYFP, pcDNA3.1-nYFP, and pRluc-N1, respectively, and XhoI and BamHI sites to clone D4.4R in pRluc-N1 vector. The amplified fragments were subcloned to be in-frame with the multiple cloning site of the vectors to give the plasmids A2a-cYFP, CB1-nYFP, D2-Rluc, or D4-Rluc.

Translational Transfection and Protein Determination

HEK-293T cells growing in six-well dishes were transiently transfected with the corresponding fusion protein cDNAs by PEI (PolyEthylenimine, Sigma, Steinheim, Germany) method. Cells were incubated with the corresponding cDNA, 5.47 mM (in nitrogen residues) PEI, and 150 mM NaCl in a serum-free medium. After 4 h, cells were placed in a fresh complete culture medium. Forty-eight hours after transfection, cells were rapidly washed twice in HBSS containing 10 mM glucose, detached, and resuspended in the same buffer. To control for cell number, sample protein concentration was determined using a Bradford assay kit (Bio-Rad, Munich, Germany) using bovine serum albumin as reference. Cell suspension (20 µg of protein) was distributed into 96-well black plates with a transparent bottom for fluorescence determinations or white plates with white bottom for BRET experiments.

Fluorescence Measurements

To quantify fluorescence, cells (20 µg protein) were distributed in 96-well microplates (black plates with a transparent bottom) and fluorescence was read in a Mithras LB 940 (Berthold Technologies, DLReady, Germany) using a 10-nm bandwidth excitation and emission filters at 485 and 530 nm, respectively. Protein fluorescence expression was determined as fluorescence of the sample minus the fluorescence of cells that were not transfected.

BRET Assays with Bimolecular Fluorescence Complemented Proteins

HEK-293T cells were transiently cotransfected with a constant amount of cDNA encoding for the protein fused to Rluc and with increasingly equal amounts of cDNA corresponding to proteins fused to one of the two complementary parts of the YFP protein (nYFP and cYFP). Fluorescence was measured as indicated above. The equivalent to 20 µg of cell suspension was distributed in 96-well microplates (Corning 3600, white plates with white bottom) and 5 µM coelenterazine H (Molecular Probes, Eugene, OR) was added. After 1-min delay, collection of readings started using a Mithras LB 940, which allows the integration of the signals detected in the short-wavelength filter at 485 nm (440–500 nm) and the long-wavelength filter.
at 530 nm (510–590 nm). To quantify for protein-Rluc expression, luminescence readings were performed after 10 min of adding 5 µM coelenterazine H. The net BRET is defined as \([\text{long-wavelength emission}}/(\text{short-wavelength emission})\]-Cf, where Cf corresponds to \([\text{long-wavelength emission}}/(\text{short-wavelength emission})\] for the Rluc construct expressed alone in the same experiment.

**Immunostaining**

For immunocytochemistry, transiently transfected HEK-293T cells were fixed in 4% paraformaldehyde for 15 min and washed with PBS containing 20 mM glycine (buffer A) to quench the aldehyde groups. Then, after permeabilization with buffer A containing 0.2% Triton X-100 for 5 min, cells were treated with PBS containing 1% bovine serum albumin. After 1 h at room temperature, cells expressing D2R-Rluc were labeled with the primary mouse monoclonal anti-Rluc antibody (1/100, Chemicon) for 1 h, washed, and stained with the secondary antibody Cy3 Donkey antimouse (1/100, Jackson Immunoresearch Laboratories, Baltimore, PA). Heterodimers of receptors fused to complementary fragments of YFP were detected by their fluorescence properties. Samples were rinsed and observed in a Leica SP2 confocal microscope (Leica Microsystems, Mannheim, Germany).

**RESULTS AND DISCUSSION**

In the early 1980s, and based on indirect functional evidence, it was proposed that GPCRs could interact at the level of the neuronal plasma membrane. In the early 1990s, electrophoretic mobility and coimmunoprecipitation assays gave the first indication of GPCR homomerization. More recently, the development of the biophysical techniques BRET and FRET allowed the demonstration of GPCR homodimerization and heteromerization of two GPCRs in living cells\[23,28,29,30,31,32,33,34,35,36\]. Nevertheless, the lack of assays monitoring interactions between more than two proteins simultaneously makes it very difficult to draw a map of molecular networks involving protein-protein interactions. By the approach previously reported by Héroux et al.\[27\], we here show that the combination of bimolecular fluorescence complementation and bioluminescence energy transfer is useful to detect heteromerization of three different GPCRs.

For bimolecular fluorescence complementation assay, the reconstitution of a reporter fluorescence protein (YFP) from its two fragments attached to the potential interacting protein partners under study\[37,38\] is taken as evidence for the molecular interaction between the partners\[39\]. The usefulness of the bimolecular fluorescence complementation technique to detect protein heterodimers was proved using cells transfected with the following subunits of heterotrimeric G proteins: GαnYFP and GβcYFP. Positive complementation was detected by the increase of fluorescence at 530 nm upon increasing the amount of transfected proteins (Fig. 1). As negative control, no signal was detected when either GαnYFP and cYFP, or GβcYFP and nYFP, were cotransfected (Fig. 1). These results prove the ability of the bimolecular complementation technique to detect heterodimers. The bimolecular fluorescence complementation technique was used to detect CB1R-A2AR heterodimers in HEK-293 cells. Fusion of nYFP and cYFP fragments to CB1R or to A2AR did not prevent the receptor functionality determined as ERK1/2 phosphorylation (results not shown). Cells transfected with CB1RnYFP and A2ARcYFP (see Methods) showed fluorescence at the membrane level (Fig. 2A). No fluorescence was detected when cells were cotransfected with A2ARcYFP and nYFP (Fig. 2A) or with CB1RnYFP and cYFP (results not shown), showing the specificity of the signal. In agreement, positive complementation was detected by the increase of fluorescence at 530 nm upon increasing the amount of transfected receptors (Fig. 2B). As a negative control, no signal was detected when either A2ARcYFP and nYFP, or CB1RnYFP and cYFP, were cotransfected (Fig. 2B). Taken together, these results validate the usefulness of the bimolecular fluorescence complementation technique to monitor the formation of CB1R-A2AR heteromers.
FIGURE 1. Molecular interaction between $G_\alpha$ and $G_\beta$ subunits of heterotrimeric G proteins detected by bimolecular fluorescence complementation. HEK-293 cells were cotransfected with equal amounts of cDNAs corresponding to the fusion proteins $G_\alpha nYFP$ and $G_\beta cYFP$ (black), $G_\alpha nYFP$ and $cYFP$ (white), or $G_\beta cYFP$ and $nYFP$ (grey). Forty-eight hours post-transfection, fluorescence was determined at 530 nm. Values are mean ± SEM of four independent experiments. One-way ANOVA followed by Newman-Keuls test showed significant differences respective to both negative controls. ***$p < 0.001$.

FIGURE 2. $CB_1R$-$A_2AR$ heterodimers in HEK-293 cells. HEK-293 cells were cotransfected with (A) 2 µg of cDNA corresponding to the fusion proteins $CB_1RnYFP$ and $A_2ARcYFP$ (left panel), or $A_2ARcYFP$ and $nYFP$ (right panel) or (B) equal amounts of cDNAs corresponding to $CB_1RnYFP$ and $A_2ARcYFP$ (black), $A_2ARcYFP$ and $nYFP$ (grey), or $CB_1RnYFP$ and $cYFP$ (white). In (A), confocal microscopy images obtained 48 h post-transfection are shown. In (B), fluorescence at 530 nm was determined 48 h post-transfection. Values are mean ± SEM of four independent experiments. One-way ANOVA followed by Newman-Keuls test showed significant differences respective to both negative controls. ***$p < 0.001$. 
The positive results on the formation of CB₁R-A₂A R heteromers, obtained by the bimolecular fluorescence complementation technique, opened the possibility of combining this technology with BRET to investigate the existence of D₂R-CB₁R-A₂A R hetero-oligomers. The usefulness of the combination of the bimolecular fluorescence complementation technique and BRET to detect oligomers formed by three different proteins was first tested by transfecting the following fusion proteins of the three subunits of heterotrimeric G proteins: Gγ-Rluc, GαnYFP, and GβcYFP. Positive BRET was detected between Gγ-Rluc and complemented GαnYFP-GβcYFP. A hyperbolic BRET saturation curve was obtained upon increasing the GαnYFP-GβcYFP expression (Fig. 3A). This result proves the ability of the combination of the two techniques to detect trimolecular protein complexes as those detected for functional calcitonin gene-related peptide receptors, which are formed by the asymmetric assembly of a calcitonin receptor-like receptor homo-oligomer and a monomer of receptor activity-modifying protein-1[27].

In order to detect possible formation of hetero-oligomers composed of CB₁, D₂, and A₂A receptors, HEK-293 cells were transfected with D₂Rluc, A₂ArCYPFP, and CB₁RnYFP. Fusion of Rluc to D₂R did not prevent the receptor functionality determined as ERK1/2 phosphorylation (results not shown). Fusion proteins did not affect the normal subcellular distribution of receptors (Fig. 3B). In fact, these receptors are predominantly colocalized in the plasma membrane of cotransfected cells. In conditions to give a BRET₅₀ fusion protein expression levels, measured as described previously[26] by radioligand binding in different experimental sessions, were between 0.5 and 0.7 pmols/mg protein for A₂ArCYPFP, between 0.9 and 1.1 pmols/mg protein for D₂RRluc, and between 0.6 and 0.8 pmols/mg protein for CB₁RnYFP. Triggering with coelenterazin H, these transfected cells gave a significant BRET signal. The BRET signal was specific as assessed by the saturation hyperbola obtained upon increasing the complemented A₂ArCYPFP-CB₁RnYFP expression and by the lack of the signal using D₂RRluc instead of D₂Rluc as a negative control (Fig. 3C). These data indicate that D₂R, CB₁R, and A₂A R form, at least, trimolecular oligomers in cotransfected living cells. This technique is validated by the identification of D₂-CB₁- and A₂A receptor heteromers by sequential resonance energy transfer (SRET)[26]. Apart from transmembrane regions, basic and acidic residues are involved in the epitope-epitope electrostatic interactions existing in D₂-A₂A receptor heteromers. In fact, mass spectometry and pull-down assays have been instrumental to show that the Arg-rich D₂R epitope may bind to two different epitopes in the C-terminal part of the A₂A R, one containing two adjacent Asp residues and another containing a phosphorylated Ser residue[18,20]. Then, it might be possible that one of the epitopes is involved in the interaction with D₂ and another in the interaction with CB₁. Further experimental work is necessary, however, to elucidate the amino acids constituting the interfaces in the D₂-CB₁-A₂A hetero-oligomer.
FIGURE 3. D₂R-CB₁R-A₂AR heteromers detected by a combination of bimolecular fluorescence complementation and BRET. In (A), as a positive control, BRET saturation curve was performed using HEK-293 cells cotransfected with 0.75 µg of cDNA corresponding to the fusion protein G₃Rluc (100,000 bioluminescence units) and increasing equal amounts of cDNAs corresponding to G₃αnYFP and G₃βcYFP (1000–6000 fluorescence units). In (B), confocal microscopy image of a cell after 48 h of transfection with 1 µg of cDNA corresponding to D₂Rluc, 2 µg of cDNA corresponding to CB₁RnYFP, and 2 µg of cDNA corresponding to A₂ARcYFP. Proteins were identified by fluorescence (green image) or by immunocytochemistry (red image) using a monoclonal anti-Rluc primary antibody and a cyanine-3-conjugated secondary antibody. Colocalization is shown in yellow in the right image. In (C), BRET saturation curve (red) was obtained using HEK-293 cells cotransfected with 1.5 µg of cDNA corresponding to D₂Rluc (100,000 bioluminescence units) and increasing equal amounts of cDNAs corresponding to CB₁RnYFP and A₂ARcYFP (1000–10,000 fluorescence units). As negative controls, cells were transfected with 1.5 µg of either the cDNA for D₄Rluc (black line) or for GABAB₂Rluc (green line) (100,000 bioluminescence units in each case).
There has been reported a coexpression of D_2R and A_2A_R in GABAergic striatal neurons (see [9]) and of CB_1R and A_2A_R in rat striatal fibrillar structures[11,13]. The demonstration of D_2R-CB_1R-A_2A_R heteromers in transfected cells, together with such striatal codistribution of the three receptors in the plasma membrane of striatal neurons, strongly suggests that these three receptors are forming part of a molecular network. The function of these neurons is particularly compromised in Parkinson’s disease and in the early stages of Huntington’s disease[21]. Furthermore, neuroadaptations of glutamatergic synapses of GABAergic enkephalinergic neurons localized in the nucleus accumbens (the ventral part of the striatum) seem to be involved in compulsive drug seeking and relapse[40]. Based on the existence of antagonistic interactions between A_2A_R and D_2R in the A_2A_R-D_2R heteromer[21,25], A_2A_R antagonists are giving successful results in clinical trials in patients with Parkinson’s disease[41]. Furthermore, A_2A_R antagonists are being considered as possible therapeutic agents in end-stage drug addiction[42]. Their clinical efficacy might be related to the recently demonstrated dependence of A_2A_R activation for CB_1R receptor signaling within the striatal A_2A_R-CB_1R heteromers[11]. Thus, A_2A_R antagonists behave as CB_1R antagonists, known to counteract cue-induced reinstatement of different addictive drugs in the experimental animal, a model for human relapse[43]. Although the intramembrane and intracellular cross-talk established by complexes formed by receptor heterodimers is already important to understand better the function of striatal enkephalinergic neurons, the occurrence of oligomers formed by three different receptors indicate a more diverse interplay between receptors for neurotransmitters. Taken together, the results already reported in the literature suggest that the A_2A_R-CB_1R-D_2R receptor heteromer may act as a processor mediating the neuronal computation needed to modulate striatal dopamine neurotransmission.

The demonstration of D_2R-CB_1R-A_2A_R heteromers in transfected cells, together with their striatal codistribution, opens new perspectives to understand the interplay between different neurotransmitter-neuromodulator systems. Pharmacological and functional diversification expand in a macromolecular complex containing three receptors by the same simple events as described for dimers, i.e., by (1) a change in the pharmacological profile of a receptor when another receptor in the complex is activated and (2) a change in the associated signaling response-pathways depending on the receptors present in the complex, their degree of activation, and the nature of the G proteins expressed in the horizontal molecular network involved[44].

The combination of bimolecular fluorescence complementation and bioluminescence energy transfer techniques constitutes a powerful approach to detect the protein-protein interactions localized in the plane of the membrane, and thus allows identification of the horizontal molecular networks like the receptor networks in local circuits. This new knowledge will hopefully provide novel therapeutic approaches for neurodegenerative diseases, mental disorders, and drug addiction.

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