Differential β-arrestin2 requirements for constitutive and agonist-induced internalization of the CB₁ cannabinoid receptor

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Abstract

CB₁ cannabinoid receptor (CB₁R) undergoes both constitutive and agonist-induced internalization, but the underlying mechanisms of these processes and the role of β-arrestins in the regulation of CB₁R function are not completely understood. In this study, we followed CB₁R internalization using confocal microscopy and bioluminescence resonance energy transfer measurements in HeLa and Neuro-2a cells. We found that upon activation CB₁R binds β-arrestin2 (β-arr2), but not β-arrestin1. Furthermore, both the expression of dominant-negative β-arr2 (β-arr2-V54D) and siRNA-mediated knock-down of β-arr2 impaired the agonist-induced internalization of CB₁R. In contrast, neither β-arr2-V54D nor β-arr2-specific siRNA had a significant effect on the constitutive internalization of CB₁R. However, both constitutive and agonist-induced internalization of CB₁R were impaired by siRNA-mediated depletion of clathrin heavy chain. We conclude that although clathrin is required for both constitutive and agonist-stimulated internalization of CB₁R, β-arr2 binding is only required for agonist-induced internalization of the receptor suggesting that the molecular mechanisms underlying constitutive and agonist-induced internalization of CB₁R are different.

Keywords: β-arrestin, bioluminescence resonance energy transfer (BRET), CB₁ cannabinoid receptor, constitutive internalisation, G protein-coupled receptors (GPCR), receptor endocytosis

1. Introduction
Internalization of G protein-coupled receptors (GPCRs) is an important process in the regulation of receptor function. Although its main function is the modulation of receptor number on the cell surface, thereby adjusting the sensitivity of the cell to external stimuli, it also plays role in the resensitization and signaling of GPCRs (Ferguson, 2001; Hunyady and Catt, 2006; Shenoy and Lefkowitz, 2011). At the molecular level, β-arrestins are key regulatory proteins of receptor internalization, as they can bind to activated GPCRs, as well as to clathrin and the adaptor protein AP-2, thus directing the receptor towards clathrin-mediated endocytosis (Shenoy and Lefkowitz, 2011). β-arrestins also mediate receptor desensitization, as their binding to the activated GPCRs causes the uncoupling of the receptor from its cognate G protein (Shenoy and Lefkowitz, 2011). Furthermore, they play important roles in the activation of G protein-independent signal transduction pathways, e.g. the activation of MAP kinases, phosphatidylinositol 3 kinase, Akt or the small GTP-ase RhoA (DeWire et al., 2007; Wei et al., 2003).

β-arrestin1 (β-arr1) and β-arrestin2 (β-arr2) are two ubiquitously expressed isoforms of β-arrestins (Ferguson, 2001). Although β-arrestin binding is a general property of most activated GPCRs, the selectivity and stability of these binding shows receptor specific differences. Namely, class A receptors (e.g. the β2 adrenergic receptor) bind β-arr2 with a higher affinity than β-arr1, and this binding is transient, i.e. it can only be detected at or near the plasma membrane. In contrast, class B GPCRs, such as the AT1 angiotensin receptor, bind both β-arr1 and β-arr2 with relatively high affinity and form
stable complexes with β-arrestins, so that β-arrestins remain bound to the receptor after internalization and can be detected on intracellular vesicles (Oakley et al., 2000).

The CB₁ cannabinoid receptor (CB₁R) belongs to the superfamily of G protein-coupled receptors (GPCRs). The receptor plays role in many important physiological processes, such as learning, thinking, nociception or the regulation of food-intake (Pacher et al., 2006). Activation of presynaptic CB₁Rs by postsynaptic endocannabinoid release, which mediates retrograde transmission, is a key regulatory mechanism in the central nervous system, but paracrine activation of CB₁Rs with a similar mechanism can also occur in extraneural tissues (Freund et al., 2003; Gyombolai et al., 2012; Sanchez et al., 2001; Turu et al., 2009; Szekeres et al., 2012). The cellular signaling events following CB₁R activation are mainly associated with the activation of heterotrimeric Gᵢₒ/proteins and include inhibition of adenylyl cyclases, activation of Kᵩ channels, inhibition of Caᵥ channels and phosphorylation and activation of different subtypes of mitogen-activated protein kinases (MAP kinases) (Turu and Hunyady, 2010). G protein-independent signaling events following CB₁R stimulation have also been reported (Sanchez et al., 2001).

Similar to most GPCRs, CB₁R internalizes upon agonist stimulation. This has been demonstrated in many cell lines, including CHO (Rinaldi-Carmona et al., 1998), AtT20 (Hsieh et al., 1999; Jin et al., 1999; Roche et al., 1999), F11 (Coutts et al., 2001), neuroblastoma N18TG2 (Keren and Sarne, 2003) and HEK293 (Keren and Sarne, 2003; Leterrier et al., 2004) cells, as well as in hippocampal neurons, which naturally express CB₁R (Coutts et al., 2001; Leterrier et al., 2006). According to different studies, this agonist-induced CB₁R endocytosis occurs via clathrin- and/or caveolin-mediated
pathways in different cell types (Bari et al., 2008; Hsieh et al., 1999; Keren and Sarne, 2003; Wu et al., 2008).

Numerous studies suggest that β-arr2 is involved in the regulation of CB₁R. Co-expression of both GRK3 and β-arr2 was needed for the proper desensitization of the receptor in Xenopus oocyte (Jin et al., 1999), and dominant negative GRK2 and β-arrestin constructs reduced CB₁R desensitization in hippocampal neurons (Kouznetsova et al., 2002). In β-arr2 knockout mice desensitization and downregulation of CB₁R were impaired in certain regions of the central nervous system (Nguyen et al., 2012). Recruitment of β-arr2 to the activated CB₁R has also been demonstrated (Daigle et al., 2008a). Mutation of amino acids S426 and S430 was shown to inhibit receptor desensitization as well as late phase receptor endocytosis, but not β-arrestin binding (Daigle et al., 2008b; Jin et al., 1999). It has been demonstrated that serine and threonine residues at the C-terminus of CB₁R are involved in its β-arr2 binding and agonist-induced endocytosis (Daigle et al., 2008b; Hsieh et al., 1999). In contrast to the various studies that clearly point to an interaction between CB₁R and β-arr2 upon agonist stimulation, no direct data have been hitherto presented concerning the β-arr1 binding of CB₁R. In some structural studies, the association of β-arr1 with a synthesized CB₁R C-terminus has been shown (Bakshi et al., 2007; Singh et al., 2011), however, such binding has not been demonstrated with the intact CB₁R in living cells.

Constitutive internalization of CB₁R (i.e. spontaneous internalization in the absence of CB₁R agonists) has also been detected in hippocampal neurons, as well as in CHO and HEK cells (Leterrier et al., 2004, 2006; McDonald et al., 2007a; Turu et al., 2007). It has been suggested that constitutive CB₁R internalization is the consequence of
its basal activity, since inverse agonist treatment or inhibition of basal activity with a DAG lipase inhibitor (e.g. tetrahydrolipstatin) interfered with this process (Leterrier et al., 2004, 2006; Rinaldi-Carmona et al., 1998; Turu et al., 2007). However, other studies have concluded that constitutive internalization occurs independently of receptor activity (McDonald et al., 2007a, 2007b; Kleyer et al., 2012). The latter statement raises the possibility that constitutive and agonist-induced internalization of CB₁R may occur via distinct endocytic mechanisms (McDonald et al., 2007a). However, no evidence has been hitherto presented showing that these two processes are truly different in that they require distinct endocytic machinery to take place.

Our main goal was, considering the important consequences of β-arrestin recruitment in the regulation of cell function, to characterize the β-arrestin binding properties of CB₁R and to reveal possible differences in constitutive and agonist-driven CB₁R endocytosis by investigating the role of β-arrestins in these processes.

2. Materials and Methods

2.1. Materials

The cDNAs of the rat vascular CB₁R and CB₁R-eYFP were provided by Zsolt Lenkei (Centre National de la Recherche Scientifique, Paris). β-arrestin1, β-arrestin2 and β-arrestin2-eGFP cDNAs were kindly provided by Dr. Marc G. Caron (Duke University, Durham, NC). Molecular biology enzymes were obtained from Fermentas (Vilnius, Lithuania) and Stratagene (La Jolla, CA). pcDNA3.1 vector, coelenterazine h, fetal bovine serum (FBS), OptiMEM, Lipofectamine 2000, and PBS-EDTA were from
Invitrogen (Carlsbad, CA). WIN55,212-2 and AM251 were from Tocris (Bristol, UK). Cell culture dishes and plates for BRET measurements were from Greiner (Kremsmunster, Austria). HaloTag® Alexa Fluor® 488 Ligand was from Promega (Madison, WI). Control siRNA, human β-arrestin2-specific siRNA, mouse β-arrestin2-specific siRNA and clathrin heavy chain-specific siRNA (with sequences 5'-UUCUCCGAACGUGUCACGU-3', 5'-GGACCGCAAAGUGUUUGUG-3', 5'-ACGUCCAUGUCAACCAACAA-3' and 5'-GAAAGAAUCUGUAGAGAAA-3', respectively) were from Eurofins MWG Operon (Ebersberg, Germany). HeLa and Neuro-2a cells were from ATCC (American Type Culture Collection, Manassas, VA). Anti-β-arrestin2 and HRP-conjugated anti-rabbit and anti-mouse antibodies were from Cell Signaling Technology Inc. (Beverly, MA). Anti-clathrin heavy chain antibody was from Transduction Laboratories (Lexington, KY). Unless otherwise stated, all other chemicals and reagents were from Sigma (St. Louis, MO).

2.2. Plasmid constructs and site-directed mutagenesis

The mVenus-tagged rat AT1a receptor (AT1R-mVenus), human β2 adrenergic receptor (β2AR-mVenus) and rat CB1R (CB1R-mVenus) were created by exchanging the sequence of fluorescent proteins in AT1R-YFP (Turu et al., 2006), β2AR-Sluc (Toth et al., 2012) or CB1R-YFP, respectively, to the sequence of mVenus using AgeI and NotI restriction enzymes. The mCherry-tagged AT1R (AT1R-mCherry) and Cerulean-tagged β2AR (β2AR-Cerulean) were created similarly. β-arrestin2-Rluc was constructed as described previously (Turu et al., 2006). β-arrestin1-Rluc and β-arrestin1-GFP were generated from β-arrestin2-Rluc and β-arrestin2-GFP, respectively, by replacing the cDNA of β-arrestin2
with that of β-arrestin1. CB$_1$R-mCherry and β-arrestin2-RFP were constructed by subcloning the cDNAs of CB$_1$R or β-arrestin2 into mCherry or RFP containing vectors (provided by Dr. R. Tsien, University of California, San Diego, CA), respectively. For the construction of Halo-CB$_1$R, the cDNA of HaloTag was first amplified from the HaloTag® pHT2 vector (Promega, Madison, WI) by PCR with the sense primer containing a cleavable signal sequence of influenza hemagglutinin (MKTIIALSYIFCLVFA) to achieve proper plasma membrane localization of the final construct (Guan et al., 1992). This product was inserted into a pEGFP-C1 vector (Clontech, Palo Alto, CA) in the place of eGFP sequence (pHalo-C1 vector). CB$_1$R cDNA was then inserted into pHalo-C1 after the HaloTag cDNA to yield Halo-CB$_1$R. CB$_1$R-Sluc was generated from CB$_1$R-YFP by replacing the eYFP coding sequence with the cDNA of super Renilla luciferase (Woo and von Arnim, 2008). EYFP-tagged ICAM-1 (ICAM-YFP) was constructed as described previously (Varnai and Balla, 2007). V54D mutation was inserted into the appropriate β-arrestin2 constructs by the QuikChange® site-directed mutagenesis kit (Stratagene, La Jolla, CA) according to manufacturer’s suggestions. Sequences of all constructs were verified using automated DNA sequencing.

2.3. Cell cultures and transfection

HeLa and Neuro-2a cells were maintained in DMEM supplemented with 10% FBS, (Invitrogen, Carlsbad, CA), 100 μg/ml streptomycin, and 100 IU/ml penicillin in 5% CO$_2$ at 37 °C. For confocal microscopy experiments, HeLa and Neuro-2a cells were grown on glass coverslips (coated with poly-L-lysine in the case of Neuro-2a cells) in 6-well plates and transfected with the indicated constructs using 1 μg/well of receptor constructs, 0.5
μg/well of β-arrrestin constructs (as indicated) and/or 25 pmol/well of siRNA (as indicated), with 2 μl/well Lipofectamine 2000 in OptiMEM following the manufacturer’s instructions.

For BRET time course experiments, HeLa cells were grown on 6-well plates and transfected with the indicated constructs using 1 μg/well of CB₁R constructs, 0.5 μg/well of β-arrestin constructs, 3 μg/well of ICAM-YFP and/or 25 pmol/well of siRNA (as indicated), with 2 μl/well Lipofectamine 2000 in OptiMEM following the manufacturer’s instructions. In the case of BRET titration experiments, HeLa cells were transfected with constant amounts of donor (β-ar1-Rluc or β-ar2-Rluc) and varying amounts of acceptor (CB₁R-mVenus, β₂AR-mVenus or AT₁R-mVenus) constructs with 2 μl/well Lipofectamine 2000 in OptiMEM following the manufacturer’s instructions. The amount of total transfected DNA was held constant (2.5 μg/well) with addition of varying amounts of empty pcDNA3.1 plasmid.

2.4. Bioluminescence resonance energy transfer (BRET) measurements

The BRET assay measuring the effects of different CB₁R variants on G protein activation was carried out as described previously (Turu et al., 2007). To measure BRET between β-arrestins and receptors, HeLa cells were transfected with Renilla luciferase-tagged β-arrestin isoforms (β-ar1-Rluc or β-ar2-Rluc) and mVenus-tagged AT₁R, β₂AR or CB₁R, and measurements were performed on the day after transfection. To measure BRET between ICAM and CB₁R, HeLa cells were transfected with eYFP-tagged ICAM-1, super Renilla luciferase-tagged CB₁R and the indicated β-ar2 construct or siRNA, and measurements were performed two days after transfection. Before the experiments cells
were detached with PBS-EDTA and centrifuged. Cells were resuspended in a modified Krebs-Ringer buffer containing 120 mM NaCl, 4.7 mM KCl, 1.2 mM CaCl$_2$, 0.7 mM MgSO$_4$, 10 mM glucose, 0.1% BSA and Na-HEPES 10 mM, pH 7.4; and transferred to white 96-well plates. Measurements were performed at 37 °C, after the addition of the cell permeable substrate, coelenterazine h at a final concentration of 5 μM. Counts were recorded using a Mithras LB 940 multilabel reader (Berthold Technologies, Bad Wildbad, Germany) with filters at 485 (for Rluc) and 530 nm (for eYFP) wavelengths. BRET ratios were calculated as a 530 nm/485 nm emission ratio. In the case of BRET titration experiments, mVenus/Rluc emission ratios were calculated by dividing average mVenus fluorescence counts (measured before the addition of coelenterazine h, with 485 nm excitation and 530 nm emission) by average Rluc bioluminescence counts (measured in non-stimulated cells after the addition of coelenterazine h). This was plotted against the average BRET ratio change measured 3 to 5 min after agonist stimulus. The measurements were done in triplicates, and the BRET records shown here are calculated from at least three independent experiments. BRET change was defined as the BRET ratio of a given time point minus the average BRET ratio of the initial (non-stimulated) time points. BRET change was baseline-corrected to the vehicle curve.

2.5. Confocal laser-scanning microscopy

Cells were grown on glass coverslips and transfected with the indicated constructs, as described above, 48 hours prior to measurement. EGFP (or Halo-Alexa488) and mCherry (or RFP) were excited with 488 nm argon, and 543 nm helium/neon lasers, respectively, and emitted fluorescence was detected with 500-550 bandpass and 560 longpass filters,
respectively, using a Zeiss LSM 510 confocal laser scanning microscope. For quantification of confocal measurements, 20 images of individual cells were taken from each sample, and intracellular and total cell fluorescence intensities of the images were determined using the ImageJ software (W. S. Rasband, ImageJ, United States National Institutes of Health, Bethesda, MD (rsb.info.nih.gov/ij/)). Intracellular fluorescence was divided by total cell fluorescence to yield IC/total cell fluorescence ratio. Blind selection and analysis of the cells were performed to avoid any bias during the evaluation of the internalization data.

2.6. Halo-labeling protocols

To measure agonist-induced internalization of the receptor, Halo-CB₁R-transfected cells were stained with Halo-Alexa488 ligand diluted 1:20 000 in DMEM, and incubated at 37°C, 5% CO₂ for 15 minutes. Cells were then washed twice with DMEM to remove unbound Halo-Alexa488, and treated with DMSO, WIN55,212-2 (10 μM) or WIN55,212-2+AM251 (10 μM+30 μM, respectively) at 37°C, 5% CO₂ for 30 minutes. At the end of stimulation period, cells were washed twice with PBS and fixed with 4% formaldehyde at 4°C for 15 minutes. Fixed cells were analyzed under confocal microscope as described above.

To measure constitutive internalization of the receptor, Halo-CB₁R-transfected cells were stained with Halo-Alexa488 ligand diluted 1:20 000 in DMEM, and incubated at 37°C, 5% CO₂ for 15 minutes. Cells were then washed twice with DMEM to remove unbound Halo-Alexa488 and incubated in DMEM at 37°C, 5% CO₂ for 5 hours and 45 minutes without any further treatment or in the presence of DMSO, WIN55,212-2 (10
μM) or AM251 (30 μM). To control the effectiveness of siRNA transfection during the 6 hours of incubation, parallel samples were subjected to the same washing procedures, however, without the staining period at the beginning. For these cells, an ‘agonist-induced’ protocol (15 min staining + 30 min WIN55 stimulus, see above) was applied 45 minutes before the end of the experiment. At the end of the 6 hours, all cells were washed twice with PBS and fixed with 4% formaldehyde for 15 minutes at 4°C. Fixed cells were analyzed under confocal microscope as described above.

2.7. Western blot analysis

Cells for Western blot analysis were transfected on 6-well plates parallel to the cells transfected for confocal analysis. On the day of experiment, cells were placed on ice and washed with ice-cold PBS. Cells were scraped into SDS sample buffer, briefly sonicated, boiled at 95°C for 5 min, centrifuged at 4°C for 10 min and separated on SDS polyacrilamide gels. The proteins were transferred to PVDF membranes, blocked (30 min, 5% fat-free milk powder in PBS with 0.05% Tween 20 (PBST)) and incubated with rabbit anti-β-arrestin2 primary antibody or mouse anti-clathrin heavy chain primary antibody (1 h, diluted 1:1000 or 1:4000, respectively, in PBST containing 5% fat-free milk powder) and HRP-linked goat anti-rabbit or anti-mouse (respectively) secondary antibody (30 min, diluted 1:2000 in PBST containing 5% fat-free milk powder). To control the total protein amount of cell samples, antibodies were removed with standard stripping buffer and a second immunostaining procedure was carried out with mouse anti-β-actin primary antibody (1 h, 1:10 000) and HRP-linked goat anti-mouse secondary antibody (30 min, 1:10 000). The antibodies were visualized using SuperSignal West
Pico reagen (Pierce Biotechnology Inc., Rockford, IL), according to manufacturer’s instructions. Western blot images were scanned and quantified using the ImageJ software. Changes in β-actin levels were used to normalize changes in β-arr2 or clathrin heavy chain levels.

2.8. Statistical analysis

Statistical analysis was made using the software SigmaStat for Windows 3.5 (Systat Software Inc., Richmond, CA). The effects of co-transfected siRNAs were evaluated using Student’s t-test or two-way ANOVA with Holm-Sidak’s post-hoc test, as appropriate. Data shown are mean ± SEM values.

3. Results

3.1. β-arrestin binding properties of CB₁R

β-arrestin binding properties of CB₁R were first analyzed using a bioluminescence resonance energy transfer (BRET) based approach. BRET was measured between Renilla luciferase tagged β-arr1 or β-arr2 (β-arr1-Rluc and β-arr2-Rluc) and the mVenus-tagged CB₁R (CB₁R-mVenus) in transiently transfected HeLa cells. Stimulation using WIN55,212-2 (WIN55, 10 µM), a potent synthetic CB₁R agonist, led to an increase in the BRET signal when β-arr2-Rluc was co-expressed with CB₁R-mVenus (Fig. 1A) reflecting the recruitment of the β-arr2-Rluc molecules to CB₁R-mVenus following receptor activation. In contrast, the same stimulus did not cause any detectable change in
the BRET signal between β-arr1-Rluc and CB₁R-mVenus (Fig. 1A), indicating that this subtype of β-arrestin does not bind to CB₁R following receptor activation.

To investigate the β-arrestin binding of CB₁R more thoroughly, BRET titration experiments were performed. In these experiments, BRET signal is measured and plotted as a function of the BRET acceptor/donor (mVenus/Rluc) expression ratio. In case of a specific interaction, such as the binding between a GPCR and β-arrestins upon agonist stimulus, BRET titration follows a saturation curve, the slope of which corresponds to the affinity between the two BRET partners (Marullo and Bouvier, 2007). Thus, BRET titration experiments were carried out between CB₁R-mVenus and β-arr1-Rluc or β-arr2-Rluc. Furthermore, the mVenus-tagged β₂-adrenoceptor and AT₁ angiotensin receptor (β₂AR-mVenus and AT₁R-mVenus, respectively) were also tested for their β-arrestin affinities in the same system, as these two receptors are prototypical class A and class B GPCRs, respectively (Oakley et al., 2000), and therefore their β-arrestin binding properties could be used as a reference in our experiments. As expected, BRET titration curves of β₂AR-mVenus and AT₁R-mVenus followed saturation kinetics with both β-arrestin isoforms after agonist stimulus. While AT₁R-mVenus binds both isoforms with high affinity (Fig. 1B), in case of β₂AR-mVenus, substantial difference was observed between β-arr1-Rluc and β-arr2-Rluc binding, showing that this GPCR has substantially higher affinity for β-arr2 (class A GPCR, Fig. 1C). BRET titration curve of CB₁R-mVenus and β-arr2-Rluc also followed saturation kinetics, however, with a very low steepness. In contrast, no detectable BRET signal change occurred with β-arr1-Rluc even at high acceptor/donor ratios (Fig. 1D).
β-arrestin binding of CB₁R was also visualized using confocal microscopy. In HeLa cells transiently transfected with mCherry-tagged CB₁R (CB₁R-mCherry) and GFP-tagged β-arr1 or β-arr2 (β-arr1-GFP or β-arr2-GFP, respectively), CB₁R-mCherry was present at the plasma membrane as well as in intracellular vesicles in non-stimulated cells. This distribution corresponds to constitutive internalization of CB₁R, a well-known property of this receptor (Fig. 2B and F). On the other hand, both β-arr1-GFP and β-arr2-GFP showed diffuse cytoplasmic (and, in the case of β-arr1-GFP, also nuclear) localization in non-stimulated cells, without any detectable β-arrestin either at the plasma membrane or in intracellular vesicles (Fig. 2A and E). After stimulation with WIN55, marked redistribution of β-arr2-GFP could be observed within a few minutes, namely it appeared in punctuate structures at the plasma membrane. However, the inner regions of the cytoplasm showed no accumulation of β-arr2-GFP containing vesicles even after longer periods of stimulation (Fig. 2G). In contrast to β-arr2-GFP, β-arr1-GFP showed no change in its localization after WIN55 stimulus (Fig. 2C), indicating that this subtype of β-arrestin does not bind to CB₁R. In the same system, fluorescently labeled β₂AR and AT₁R showed β-arrestin binding pattern, which are in good agreement with their class A and class B behavior, respectively, i.e. β₂AR bound β-arr1 and β-arr2 only at the plasma membrane (Suppl. Fig. 1A-D), whereas AT₁R remained bound with both isoforms in endosomal vesicles (Suppl. Fig. 1E-H).

The above data indicate that, based on its β-arrestin binding properties, CB₁R belongs to the class A group of GPCRs, i.e. upon activation, β-arr2 binds to the receptor, but remains bound to it only in the proximity of the plasma membrane, whereas CB₁R does not recruit β-arr1.
3.2. Dominant-negative β-arr2 impairs agonist-induced CB₁R internalization

Next, the agonist-induced internalization of CB₁R was investigated. We utilized the Halo labeling technique, as the analysis of the internalization of C-terminally tagged CB₁R variants using confocal microscopy is difficult because of the presence of the constitutively internalized intracellular receptor population. The HaloTag protein was fused to the N-terminus of the receptor (Halo-CB₁R). When expressed by the cells, HaloTag is able to rapidly and covalently bind Halo-Alexa488, its membrane-impermeant fluorescent ligand. Using this method, the receptor population residing in the plasma membrane can be selectively stained, and their internalization can be followed. The functionality of Halo-CB₁R was tested using BRET assays measuring the G protein activation and the β-arr2 binding of the receptors upon WIN55 stimulus, and no significant difference was detected between the EC₅₀ (mol/l) values of wild-type and HaloTag-labeled CB₁R dose-response curves (pEC₅₀ 8.98±0.22 versus 9.02±0.12 for G protein activation and 6.48±0.14 versus 6.49±0.08 for β-arr2 binding, respectively, n=3).

To study agonist-induced internalization, cells were stained for 15 minutes, treated with vehicle or CB₁R ligand for 30 minutes, and then fixed and analyzed using confocal microscopy. In vehicle-treated cells, plasma membrane staining was clearly detectable, and labeling of intracellular vesicles was minimal (Fig. 3A). WIN55 treatment resulted in the appearance of numerous intracellular vesicles, representing intensive agonist-induced internalization, which was inhibited by the CB₁R inverse agonist AM251 (Fig. 3B-C). To study the role of β-arr2 in this process, first, a dominant-negative β-arr2 mutant, β-arr2-V54D (β-arr2-V54D) was applied. This mutant has reduced ability to bind to GPCRs
(Ferguson et al., 1996; Krupnick et al., 1997), and has been widely used to study the role of β-arr2 in receptor internalization. We co-expressed the RFP-tagged form of wild-type or V54D mutant β-arr2 (β-arr2-RFP and β-arr2-V54D-RFP, respectively) with Halo-CB1R. Co-expression of β-arr2-RFP did not impair the WIN55-induced CB1R internalization (Fig. 3D-G). However, in cells expressing β-arr2-V54D-RFP, agonist-induced internalization of Halo-CB1R was markedly reduced (Fig. 3H-K), suggesting a role for β-arr2 in agonist-induced CB1R internalization.

3.3. siRNA-mediated knock-down of β-arr2 impairs agonist-induced internalization of CB1R

siRNA-mediated knock-down of a protein provides an alternative approach to study its role in a given mechanism. Therefore, the effects of transfection of cells with β-arr2-specific siRNA (β-arr2 siRNA) were also investigated. Western blot analysis showed approximately 50% reduction in β-arr2 protein levels of these cells (Fig. 4A). Internalization of Halo-CB1R in these experiments was analyzed as described above. CB1R internalization upon WIN55 treatment was intact in cells transfected with control siRNA, but was markedly reduced in β-arr2 siRNA-transfected cells (Fig. 4B-E). As the cells that were transfected with siRNA could not be individually identified in these experiments, quantification of the samples was carried out to properly evaluate the effects of siRNA transfection. This showed a significant inhibitory effect of β-arr2 siRNA on agonist-induced CB1R internalization (Fig. 4F).

3.4. Agonist-induced internalization of CB1R
To investigate the agonist-induced internalization of CB₁R in a more quantitative way, we monitored the movement of the receptor between the plasma membrane and the cytoplasm, by measuring BRET between the YFP-tagged form of ICAM-1, a plasma membrane resident protein (ICAM-YFP), and the SuperRenilla luciferase-tagged CB₁R (CB₁R-Sluc) in transiently transfected HeLa cells. In cells expressing CB₁R-Sluc and ICAM-YFP, WIN55 stimulus led to a decrease in the BRET signal, indicating the removal of CB₁R-Sluc from the plasma membrane (Fig. 5A). This decrease reached its maximum 10 to 20 minutes after stimulation, and showed then no considerable change until the end of the measurement (at approximately 40 minutes after stimulation). Overexpression of wild-type β-arr2 had no significant effect on CB₁R internalization. However, when β-arr2-V54D was co-expressed in the cells, the extent of the BRET-decrease upon WIN55 stimulus was significantly lower than that observed in control cells (Fig. 5A). The effect of siRNA mediated β-arr2 knock-down on CB₁R internalization was also studied using the same BRET assay. We found that the knock-down of β-arr2 with β-arr2 siRNA resulted in a marked reduction of BRET signal decrease after CB₁R activation, reflecting the impaired agonist-induced internalization of the receptor (Fig. 5B).

These data indicate that β-arr2 binding is required for the agonist-induced internalization of CB₁R, since interference with this binding following receptor activation, either caused by dominant-negative effect or through the genetic knock-down of β-arr2, results in impairment of agonist-induced CB₁R internalization.

3.5. Constitutive CB₁R internalization is not affected by β-arr2 impairment
Recent data suggest that constitutive internalization may occur independently of CB₁R activity (McDonald et al., 2007a, 2007b; Kleyer et al., 2012), and since β-arr2 coupling is an activity-dependent process, it raises the possibility that it is a second, differently regulated endocytic process. Therefore we monitored receptor constitutive endocytosis using similar approaches with Halo-tagged CB₁R to address possible differences in its regulation compared to agonist-induced internalization. In these experiments, cells were stained for 15 minutes with the Alexa488-conjugated Halo-ligand, and incubated for a total of 6 hours in medium to allow spontaneous CB₁R endocytosis to occur. In control cells, a large number of vesicles could be seen intracellularly, demonstrating constitutive endocytosis of CB₁R (Fig. 6A). This could be further enhanced with WIN55 treatment, whereby hardly any detectable receptor was present on the plasma membrane by the end of the 6 hours incubation period (Fig. 6B). AM251 treatment along the 6 hours had no inhibitory effect on constitutive CB₁R internalization (Fig. 6C), showing that this form of internalization is not dependent on receptor activity. When studying the role of β-arr2 in this process, cells co-expressing β-arr2-RFP showed no detectable change in the extent of constitutive internalization (Fig. 6D-E). Moreover, in contrast to agonist-induced endocytosis, in cells co-expressing the dominant-negative mutant β-arr2 (β-arr2-V54D-RFP), intracellular Halo-CB₁R-containing vesicles were still present, in a similar extent as in control or β-arr2-RFP expressing cells (Fig. 6F-G). When cells were co-transfected with either control or β-arr2 siRNA, no significant difference in the intracellular vs. total cell fluorescence ratio could be detected between these cells after 6 hours of incubation, showing that constitutive internalization of CB₁R was not affected (Fig. 7A). The effect of siRNA was not attenuated throughout the incubation period, as in the same
experiments, agonist-induced CB\textsubscript{1}R internalization was still impaired when tested at the end of the incubation period (Fig. 7B). These data suggest that β-arr2 is not required for the spontaneous internalization of CB\textsubscript{1}R. Similar results were obtained when Halo-CB\textsubscript{1}R internalization was investigated in a Neuro-2a mouse neuroblastoma cell line, which is known to endogenously express CB\textsubscript{1}R (Graham et al., 2006). siRNA mediated knock-down of β-arr2 (resulting in an approximately 80% reduction of β-arr2 protein levels, Suppl. Fig. 1A) impaired agonist-induced internalization of Halo-CB\textsubscript{1}R significantly in these cells, but had no effect on the constitutive internalization of the receptor (Fig. 7C and D). These data indicate that the β-arr2 requirements of CB\textsubscript{1}R internalization are not specific to HeLa cells and are regulated similarly in Neuro-2a cells, which express endogenous CB\textsubscript{1}Rs.

3.6. siRNA-mediated knock-down of clathrin heavy chain impairs both agonist-induced and constitutive internalization of CB\textsubscript{1}R

Our results showed that constitutive CB\textsubscript{1}R internalization is independent of β-arr2 binding, which raises the question whether this form of CB\textsubscript{1}R endocytosis is clathrin-mediated. Therefore we applied siRNA-mediated knock-down of clathrin heavy chain in HeLa cells, and monitored constitutive endocytosis of Halo-CB\textsubscript{1}R under these conditions, as described above. Western blot analysis showed approximately 50% reduction in clathrin heavy chain levels of these cells (Suppl. Fig. 2B). We found that in cells transfected with clathrin heavy chain siRNA the spontaneous endocytosis of Halo-CB\textsubscript{1}R over 6 hours was significantly impaired compared to control siRNA-transfected cells (Fig. 8A). In the same experiments, agonist-induced CB\textsubscript{1}R internalization was also
inhibited in clathrin heavy chain siRNA-transfected cells (Fig. 8B). These results show that, although the β-arr2 requirements for agonist-induced and constitutive CB₁R internalization are different, both of them occur at least partly by a clathrin-mediated process.

4. Discussion

Our data show that, based on its β-arrestin binding characteristics, CB₁R can be classified as a class A GPCR. The relatively low affinity of the binding between CB₁R and β-arr2 is indicated by the low steepness of slope of the BRET saturation curve between the two molecules upon CB₁R stimulation, and by the lack of detectable β-arrestin2 in late endosomes after CB₁R internalization in confocal experiments. Furthermore, we found in our BRET and confocal measurements that β-arr1 recruitment to the activated CB₁R is essentially absent. This was not caused by the presence of the fluorescent tag on the C-terminus of CB₁R because the binding of β-arr2 could be detected using the same receptor construct. The C-terminal tagging of β-arr1 is also not likely to cause the lack of the signal, as both β-arr1-Rluc and β-arr1-GFP constructs could be successfully used to detect β-arr1 recruitment to AT₁R and β2AR. In a previous study, Bakshi et al. have shown using nuclear magnetic resonance (NMR) that a C-terminal fragment of CB₁R is able to bind β-arr1 (Bakshi et al., 2007). Our data suggest that such binding between the two molecules, even if present, must be very weak compared to β-arr2, when tested with native proteins in living cells.

According to Oakley and colleagues, the class A or class B behavior of a GPCR is basically determined by the absence or presence (respectively) of multiple
serine/threonine-containing clusters at the receptor C-terminus (Oakley et al., 2001). Our findings demonstrating a class A pattern for CB₁R are in good agreement with this theory, as the C-terminus of CB₁R contains only one such cluster (463-SVSTDTS-469) (Oakley et al., 2001).

The class A behavior of CB₁R may have an important impact on the pattern of receptor internalization, de- and resensitization. Furthermore, one should be aware of this property of CB₁R when studying the signal transduction of the receptor. As mentioned in the introduction, β-arrestin can by itself moderate different intracellular signaling pathways. It was also shown that the localization of β-arrestin-dependent MAPK activation is dependent on the class A/class B characteristics of the involved receptor. After stimulation of class B receptors, β-arrestin activates a MAPK pool that remains associated with endosomes, whereas in the case of class A GPCRs, the MAPK activated by β-arrestin is able to reach the nucleus (Tohgo et al., 2003). CB₁R is also known to mediate ERK phosphorylation, and early gene expression changes are associated with this pathway (Derkinderen et al., 2003; Graham et al., 2006; Valjent et al., 2001). However, G protein-dependent and β-arrestin-dependent MAPK activation of GPCRs are known to be spatially and temporally different (Ahn et al., 2004). Although the role of β-arrestin molecules in CB₁R-mediated ERK activation has been suggested (Daigle et al., 2008a), the exact role of this process in CB₁R-mediated ERK activation requires additional studies. It is very likely that the class A behavior of CB₁R is associated with a characteristic distribution of MAPK activation, nevertheless, direct studies that address these questions are needed to confirm them.
Agonist-induced internalization of G protein-coupled receptors is usually the direct consequence of receptor activation. Similarly, constitutive receptor internalization is also generally considered to be a result of constitutive receptor activity. This scenario also implies that the mechanisms that control the endocytosis of a given receptor after agonist activation are similarly engaged in its spontaneous endocytosis. This has also been the prevailing idea in the case of CB$_1$R trafficking. In our study, the Halo labeling technique was utilized to analyze the mechanism of internalization of CB$_1$R. The cellular trafficking of GPCRs is often followed by the use of fluorescent proteins fused mainly to the C-terminus of the receptor. However, the detailed analysis of the internalization of such receptor variants under microscope is often hindered by the continuously visible intracellular receptor populations (representing either maturing or endosomal receptors), and this is especially true for constitutively internalizing receptors, such as CB$_1$R. Receptor distribution on cell surface and in intracellular vesicles is a function of internalization and recycling rates, representing an actual equilibrium of these processes. When cells are treated with a receptor antagonist, theoretically both processes can be altered. Receptor internalization can be inhibited by decreasing constitutive activity, or recycling may be altered by targeting the receptor to different intracellular trafficking routes. Thus, measuring C-terminally tagged receptor distribution in cells can be misleading when trafficking of constitutively internalizing receptors is measured. Furthermore, alterations in the trafficking of C-terminally tagged CB$_1$R have been also suggested (Rozenfeld, 2011). All these difficulties can be overcome by the use of N-terminal tags that can be covalently labeled with small, membrane-impermeable fluorescent ligands. Indeed, such receptor variants (i.e. SNAP and CLIP-tagged
receptors) have been utilized in the studying of CB₁R internalization (Ward et al., 2011a, 2011b). Here we used another variant, the HaloTag labeled form of CB₁R. As indicated in our study, G protein activation and β-arrestin binding of Halo-CB₁R have not been altered due to the N-terminal labeling, and therefore this receptor can be used to study receptor trafficking with the same advantages as the above mentioned ones.

The main finding of our study is that constitutive CB₁R internalization is not only independent of the active/inactive state of the receptor, but it occurs via a mechanism which is at least partly different from activity-driven endocytosis.

This finding is based on our observations that agonist-induced CB₁R internalization requires β-arr2 binding, whereas constitutive internalization of the receptor does not. As described in the introduction, the binding between β-arr2 and CB₁R as well as a role for β-arrestins in the desensitization of CB₁R have been demonstrated by former studies (Bakshi et al., 2007; Daigle et al., 2008a, 2008b; Jin et al., 1999; Kouznetsova et al., 2002; Nguyen et al., 2012; Singh et al., 2011). Therefore, the role of β-arr2 in CB₁R endocytosis may seem to be a logical consequence of its recruitment to the receptor. However, in fact, none of these studies provided direct evidence at the cellular level that β-arrestin binding is required for CB₁R internalization. This is important because various examples demonstrate that β-arrestin2 recruitment and receptor internalization can be independent processes, as it is the case e.g. with the PAR1 thrombin receptor (Paing et al., 2002), the CXCR2 chemokine receptor (Fan et al., 2001; Zhao et al., 2004), the N-formyl peptide receptor (FPR) (Vines et al., 2003), or at high agonist concentration the AT₁ angiotensin receptor (Gaborik et al., 2001; Hunyady and Catt, 2006; Zhang et al., 1996). Therefore, it is important to study directly the β-arrestin
dependence of CB₁R internalization. In the present study we performed these experiments using a dominant-negative β-arrestin2 mutant and siRNA-mediated knock-down of β-arr2. Our data demonstrate that β-arr2 binding is required for the agonist-induced internalization of CB₁R.

In contrast, using similar approaches, we have demonstrated in our study that, although both agonist-induced and constitutive CB₁R endocytosis are mediated at least partly by clathrin, constitutive internalization of CB₁R is a β-arr2-independent process. Earlier, McDonald et al. raised the idea that these two forms of CB₁R internalization may occur via different, or at least partly different, pathways (McDonald et al., 2007a). Our work provides experimental evidence to support their proposal. Such regulatory phenomenon is however not a unique property of CB₁R, as similar differences in the mechanisms of tonic versus agonist-induced internalization have been demonstrated in the case of other GPCRs, such as the TP thromboxane A2 receptor (Parent et al., 2001), the Y₁ neuropeptide Y receptor (Holliday et al., 2005) or the mGlu₁ metabotropic glutamate receptor (Dale et al., 2001). Furthermore, we also show here that the spontaneous internalization of CB₁R is not simply a consequence of its constitutive activity, since it also occurs in the presence of the CB₁R inverse agonist AM251, i.e. when the receptors are stabilized in their inactive state. This is in line with the data published by McDonald et al., which showed no alterations in the cellular distribution of CB₁R upon inverse agonist treatment in neurons (McDonald et al., 2007a). In contrast, a previous study using inverse agonist treatment (Leterrier et al., 2006), and our study using DAGL-inhibitor treatment (Turu et al., 2007) have shown an inhibition of basal endocytosis of the receptor upon these treatments, suggesting a role for basal CB₁R
activity in this process. Since active receptors internalize by β-arr2-dependent mechanism, those results may seem to be contradictory with our present results, where we now find that interfering with β-arr2 expression or function does not have effect on constitutive endocytosis. One explanation for these differences is our previous observation that CB₁R has different basal activity in different cell types used in these studies (data not shown). The other possibility is that inhibition of receptor activity may interfere with intracellular trafficking, so it might also accelerate recycling, thus the rate of activity-dependent constitutive internalization may be well overestimated in those experiments. This is supported by the observation, that in resting cells, β-arr2 is not translocated to the plasma membrane, suggesting a low percent of active receptors there (Fig. 2E). The advantage of using the Halo technique is that we can now selectively follow trafficking of surface receptors, thus recycling becomes less of an issue. Therefore, combined with previous results, the data presented here are consistent with a model in which continuous CB₁R internalization in resting cells is caused to a certain extent by basal release of endocannabinoids, the rate of which may vary in different cell types, while another part of basal CB₁R internalization is ‘truly’ constitutive, i.e. it is independent of CB₁R activation and requires a mechanism that is different from agonist-induced, β-arrestin2-dependent internalization.

Recent work by Kleyer and colleagues demonstrated that in immune cells expressing endogenous CB₁Rs, internalization of the receptor is independent of its activation (Kleyer et al., 2012), suggesting that agonist-dependent internalization may not have physiological role in primary cells. However, in hippocampal neurons, it has been demonstrated that agonist treatment leads to internalization (Coutts et al., 2001). These
data, together with differences detected in the regulation of constitutive internalization (Leterrier et al., 2006; McDonalds et al., 2007a) are in good agreement with our results showing that CB₁R internalizes through at least two different mechanisms. Depending on the environment in a particular cell type, or on the methods used by different groups, one or the other mechanism might be more prevalent.

The physiological role of spontaneous CB₁R endocytosis is not clear at the moment. It has been suggested that constitutive CB₁R internalization in neurons have a role in the axonal distribution of the receptor (Leterrier et al., 2006; McDonald et al., 2007a). This underlines the importance of constitutive CB₁R endocytosis in the central nervous system. However, CB₁Rs are also expressed in peripheral cannabinoid target tissues, and we have practically no data what is the importance of such trafficking feature of the receptor in these cells. On the other hand, maintained functionality of intracellular CB₁R populations has been demonstrated (Brailoiu et al., 2011; Rozenfeld and Devi, 2008), and this raises the idea that the activation of intracellular versus plasma membrane CB₁R populations may differentially regulate cell functions (Rozenfeld, 2011). Nevertheless, additional studies are needed to further clarify the importance of constitutive CB₁R endocytosis in different body tissues.

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Figure legends

Fig.1 BRET measurements showing the recruitment of β-arr1 and β-arr2 to CB₁R, AT₁R and β₂AR upon agonist stimulus

A, CB₁R-mVenus was co-expressed with β-arr1-Rluc or β-arr2-Rluc in HeLa cells, and BRET was measured upon WIN55 (10 μM) stimulus. Measurements were baseline-corrected to vehicle curves (indicated by horizontal dashed line). Arrow indicates the time point of stimulation. Data are mean±SEM. B-D, BRET titration curves showing the relative affinities of AT₁R (B), β₂AR (C) and CB₁R (D) to β-arrestin isoforms. HeLa cells were transfected with constant amounts of β-arr1-Rluc or β-arr2-Rluc, and varying amounts of the mVenus-tagged receptor, yielding different acceptor/donor ratios. Average BRET change between 3 to 5 minutes after agonist stimulus (100 nM angiotensin II (B), 1 μM isoproterenol (C) or 10 μM WIN55 (D)) was plotted against mVenus/Rluc intensity ratios measured at the beginning of each experiment. Data resulting from at least 3 independent experiments were fitted using non-linear regression with a one-site binding equation.

Fig.2 Confocal microscopy analysis indicates class A β-arrestin binding pattern of CB₁R

β-arr1-GFP (A-D) or β-arr2-GFP (E-H) and CB₁R-mCherry were co-expressed in HeLa cells and analyzed by confocal microscopy. Under control conditions, β-arr1-GFP shows diffuse cytoplasmic and nuclear localization (A). After 20 minutes of WIN55 (10 μM)
stimulus, no change in β-arr1-GFP distribution can be detected (C). β-arr2-GFP shows diffuse cytoplasmic localization in control cells (E, inset). After 20 minutes of WIN55 (10 μM) stimulus, β-arr2-GFP can be detected in punctuate structures, however only in the close proximity of the plasma membrane (G, inset, arrows indicate β-arr2-GFP puncta). A large proportion of CB₁R is constitutively intracellular, reflecting to spontaneous endocytosis of the receptor (B,D,F,H). Images are representative from 3 independent experiments. Scale bar 10 μm.

**Fig.3 Agonist-induced internalization of CB₁R is impaired by dominant-negative β-arr2**

Halo-CB₁R was expressed in HeLa cells alone (A,B,C) or together with wild-type β-arr2-RFP (D-G) or β-arr2-V54D-RFP (H-K), and analyzed with confocal microscopy after 15 min Halo-Alexa488 staining and 30 min vehicle (A,D,E,H,I) WIN55 (10 μM, B,F,G,J,K) or WIN55+AM251 (10 μM+30 μM, respectively, C) treatment. Vehicle treatment causes no substantial Halo-CB₁R internalization (A,D,H) in either cell population, whereas WIN55 treatment leads to massive endocytosis of the receptor in control (B) or β-arr2-RFP- (F,G), but not in β-arr2-V54D-RFP- (J,K) expressing cells. WIN55-induced internalization is blocked by co-treatment with AM251 (C). Images are representative from 3 independent experiments. Scale bar 10 μm.

**Fig.4 Agonist-induced internalization of CB₁R is impaired by β-arr2-specific siRNA**

A, Western blot analysis shows an approximately 50% reduction in the β-arr2 protein levels of β-arr2 siRNA-transfected cells compared to control siRNA-transfected cells.
n=3 B-E, HeLa cells were transfected with Halo-CB₁R and control (B,D) or β-arr2-specific (C,E) siRNA, and analyzed with confocal microscopy after 15 min Halo-Alexa488 staining and 30 min vehicle (B,C) or WIN55 (10 μM, D,E) treatment. Vehicle treatment causes no substantial Halo-CB₁R internalization in either cell population (B,C), whereas WIN55 treatment leads to massive endocytosis of the receptor in control (D), but not in β-arr2 (E) siRNA-transfected cells. Scale bar 10 μm. F, Quantification of the data using an intracellular to total cell fluorescence ratio shows a significant increase in intracellular receptor number upon WIN55 stimulus in control siRNA-transfected cells, and this is significantly reduced in cells transfected with β-arr2 siRNA. Data are mean±SEM, n=6, *p<0.05, ns - not significant.

**Fig.5 BRET measurements showing the β-arr2 dependence of agonist-induced CB₁R internalization**

CB₁R-Sluc was co-expressed with ICAM-YFP in HeLa cells, and BRET was measured to follow the agonist-induced removal of the receptor from the plasma membrane. A, BRET signal decrease upon WIN55 (10 μM) stimulus can be detected in cells co-transfected with pcDNA3.1 (open circles). Co-expression of wild-type β-arr2 has no significant impact on BRET change (closed triangles). Co-expression of β-arr2-V54D substantially reduces BRET signal decrease (open triangles). Data are all mean±SEM, n=8. B, BRET signal decrease upon WIN55 (10 μM) stimulus can be detected in control siRNA-transfected cells (closed triangles). The BRET signal decrease is substantially reduced in cells transfected with β-arr2 siRNA (open triangles). Data are all mean±SEM, n=4.
Measurements were baseline-corrected to vehicle curves (indicated by horizontal dashed lines). Arrows indicate the time point of stimulation.

**Fig.6 Constitutive internalization of CB₁R is not affected by inverse agonist treatment or dominant-negative β-arr2**

Halo-CB₁R was expressed in HeLa cells alone (A,B,C) or together with wild-type β-arr2-RFP (D,E) or β-arr2-V54D-RFP (F,G), and analyzed by confocal microscopy after 15 min Halo-Alexa488 staining and 5 h 45 min incubation at 37°C, 5% CO₂. In control cells (A), a substantial amount of Halo-CB₁R can be detected intracellularly. Internalization is enhanced in the presence of WIN55 (10 μM, B) but not affected by AM251 (30 μM, C). Constitutive internalization of the receptor can be detected in cells expressing wild-type β-arr2-RFP (D,E) and also in cells expressing β-arr2-V54D-RFP (F,G, see cell indicated with arrow). Images are representative from 3 independent experiments. Scale bar 10 μm.

**Fig.7 Constitutive internalization of CB₁R is not affected by β-arr2-specific siRNA**

A and C, HeLa (A) or Neuro-2a (C) cells were transfected with Halo-CB₁R and control or β-arr2-specific siRNA, and analyzed with confocal microscopy after 15 min Halo-Alexa488 staining and 5 h 45 min incubation at 37°C, 5% CO₂. In both control and β-arr2 siRNA-transfected cells, substantial amounts of intracellular receptors can be detected. Quantification of the data using an intracellular to total cell fluorescence ratio shows no significant difference between control or β-arr2 siRNA-transfected cells after the 6 hours of incubation. B and D, In the same experiments, control or β-arr2 siRNA-transfected HeLa (B) or Neuro-2a (D) cells were incubated at 37°C, 5% CO₂ for 5 h 15 min followed
by 15 min Halo-Alexa488 staining and 30 min WIN55 (10 μM) treatment. A substantial amount of internalized receptors was detected in control, but not in β-arr2 siRNA-transfected cells. Data quantification shows that internalization upon WIN55 stimulus after 5 h 30 min is significantly reduced in β-arr2 siRNA-transfected cells. Scale bar 10 μm. Data are mean+SEM, n=3, *p<0.05.

Fig.8 Constitutive internalization of CB₁R is impaired by clathrin heavy chain-specific siRNA

A, HeLa cells were transfected with Halo-CB₁R and control or clathrin heavy chain-specific siRNA (clathrin siRNA), and analyzed with confocal microscopy after 15 min Halo-Alexa488 staining and 5 h 45 min incubation at 37°C, 5% CO₂. A substantial amount of internalized receptors was detected in control, but not in clathrin siRNA-transfected cells. Quantification of the data using an intracellular to total cell fluorescence ratio shows significant difference between control or clathrin siRNA-transfected cells after the 6 hours of incubation. B, In the same experiments, control or clathrin siRNA-transfected HeLa cells were incubated at 37°C, 5% CO₂ for 5 h 15 min followed by 15 min Halo-Alexa488 staining and 30 min WIN55 (10 μM) treatment. A substantial amount of internalized receptors was detected in control, but not in β-arr2 siRNA-transfected cells. Data quantification shows that internalization upon WIN55 stimulus after 5 h 30 min is significantly reduced in clathrin siRNA-transfected cells. Scale bar 10 μm. Data are mean+SEM, n=3, *p<0.05.
Suppl. Fig.1 Confocal microscopy analysis showing the β-arrestin binding of AT₁R and β₂AR

AT₁R-mCherry (A-D) or β₂AR-Cerulean (E-H) were co-expressed with β.arr1-GFP (A,B,E,F) or β.arr2-GFP (C,D,G,H) in HeLa cells and analyzed by confocal microscopy. In AT₁R-mCherry expressing cells, both β.arr1-GFP and β.arr2-GFP could be detected in intracellular vesicles after 20 minutes of angiotensin II (Ang II, 100 nM) stimulus (A,C). In β₂AR-Cerulean expressing cells, β.arr1-GFP and β.arr2-GFP could be detected in puncta at the plasma membrane but not in intracellular vesicles after 20 minutes of isoproterenol (ISO, 1 μM) stimulus (E,G). Images are representative from 3 independent experiments. Scale bar 10 μm

Suppl. Fig.2 Western blot images showing siRNA induced protein knock-down

Neuro-2a (A) or HeLa (B) were transfected with Halo-CB₁R and control or mouse β-arrestin2-specific (A) or clathrin heavy chain specific (B) siRNA (clathrin siRNA) as described in Chapter 2.3. Western blot experiments were carried out as described in Chapter 2.7. A, Data quantification shows an approximately 80% reduction in the β.arr2 protein levels of β.arr2 siRNA-transfected Neuro-2a cells compared to control siRNA-transfected cells. B, Data quantification shows an approximately 50% reduction in the clathrin heavy chain (clathrin HC) protein levels of clathrin siRNA-transfected HeLa cells compared to control siRNA-transfected cells. Data are mean+SEM, n=3
• CB₁R binds β-arrestin2 but not β-arrestin1 upon activation.
• Agonist-induced internalization of CB₁R is β-arrestin2 dependent.
• Constitutive CB₁R internalization does not require β-arrestin2.
• Agonist-induced and constitutive CB₁R endocytosis are distinctly regulated processes.
Figure 1

A. CB₁R-mVenus

B. AT₁R-mVenus

C. β₂AR-mVenus

D. CB₁R-mVenus

BRET change upon stimulus

mVenus/Rluc

β-arr1-Rluc
β-arr2-Rluc

β-arr1-Rluc
β-arr2-Rluc

β-arr1-Rluc
β-arr2-Rluc

β-arr1-Rluc
β-arr2-Rluc
Figure 2

**β-arr1-GFP**

(A) Control

(C) WIN55 20 min

**CB₁R-mCherry**

(B) Control

(D) WIN55 20 min

**β-arr2-GFP**

(E) Control

(G) WIN55 20 min

**CB₁R-mCherry**

(F) Control

(H) WIN55 20 min
Figure 3

Halo-CB₁R

DMSO 30 min

WIN55 30 min

WIN55 + AM251 30 min

A

B

C

D

E

F

G

H

I

J

K

β-arr2-V54D-RFP

β-arr2-RFP

Halo-CB₁R

Halo-CB₁R
Figure 4

A

![Image of Western blot](image1)

- **β-arr2 siRNA** vs **control siRNA**
- Normalized protein level

B-C

- **DMSO 30 min** and **WIN55 30 min**
- **control siRNA** vs **β-arr2 siRNA**

D-E

- **DMSO 30 min** and **WIN55 30 min**
- **control siRNA** vs **β-arr2 siRNA**

F

- **IC/Total fluorescence**
- **DMSO** vs **WIN55**
- **control siRNA** vs **β-arr2 siRNA**

* p < 0.05

ns: not significant
Figure 5

A

BRET ratio change vs. Time (min)

- + pcDNA3.1
- + β-arr2-WT
- + β-arr2-V54D

B

BRET ratio change vs. Time (min)

- + control siRNA
- + β-arr2 siRNA
Figure 6

Halo-CB₁R

DMSO 6 h | WIN55 6 h | AM251 6 h

A B C

Halo-CB₁R β-arr2-V54D-RFP

D E

6 h incubation

Halo-CB₁R β-arr2-RFP

D E

6 h incubation

Halo-CB₁R β-arr2-V54D-RFP

F G

6 h incubation
Figure 7

A. HeLa cells

6 h incubation

control siRNA  |  β-arr2 siRNA

5 h 30 min incubation + 30 min WIN55

control siRNA  |  β-arr2 siRNA

B. HeLa cells

control siRNA  |  β-arr2 siRNA

5 h 30 min incubation + 30 min WIN55

C. Neuro-2a cells

6 h incubation

control siRNA  |  β-arr2 siRNA

D. Neuro-2a cells

5 h 30 min incubation + 30 min WIN55

control siRNA  |  β-arr2 siRNA
Figure 8

A

control siRNA  clathrin siRNA

6 h incubation

B

control siRNA  clathrin siRNA

5 h 30 min incubation + 30 min WIN55

IC/Total fluorescence
Supplementary Figure 1

(A) β-arr1-GFP

(B) AT₁R-mCherry

(C) β-arr2-GFP

(D) AT₁R-mCherry

(E) β-arr1-GFP

(F) β₂AR-Cerulean

(G) β-arr2-GFP

(H) β₂AR-Cerulean
Supplementary Figure 2

A

![Western blot images for β-arr2 and β-actin in control siRNA and β-arr2 siRNA conditions.](image)

- **β-arr2**
  - Control siRNA
  - β-arr2 siRNA
  - Molecular weight: 46 kDa

- **β-actin**
  - Control siRNA
  - β-arr2 siRNA
  - Molecular weight: 42 kDa

B

![Western blot images for clathrin HC and β-actin in control siRNA and clathrin siRNA conditions.](image)

- **clathrin HC**
  - Control siRNA
  - Clathrin siRNA
  - Molecular weight: 190 kDa

- **β-actin**
  - Control siRNA
  - Clathrin siRNA
  - Molecular weight: 42 kDa

**Graphs**

- **Normalized protein level**
  - X-axis: control siRNA, β-arr2 siRNA
  - Y-axis: normalized protein level

- **Normalized protein level**
  - X-axis: control siRNA, clathrin siRNA
  - Y-axis: normalized protein level

**Legend**

- control siRNA
- β-arr2 siRNA
- clathrin siRNA