Unusually persistent \( \text{G}_\alpha \text{i} \)-signaling of the neuropeptide \( \text{Y}_2 \) receptor depletes cellular \( \text{G}_i/o \) pools and leads to a \( \text{G}_i \)-refractory state

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Abstract

Background: A sensitive balance between receptor activation and desensitization is crucial for cellular homeostasis. Like many other GPCR, the human neuropeptide \( \text{Y}_2 \) receptor (h\( \text{Y}_2\)R) undergoes ligand dependent activation and internalization into intracellular compartments, followed by recycling to the plasma membrane. This receptor is involved in the pathophysiology of distinct diseases e.g. epilepsy and cancer progression and conveys anorexigenic signals which makes it an interesting and promising anti-obesity target. However, \( \text{Y}_2\)R desensitization was observed after daily treatment with a selective PYY\(_{1-36}\) analog in vivo by a yet unknown mechanism.

Materials: We studied the desensitization and activatability of recycled \( \text{Y}_2\)R in transiently transfected HEK293 cells as well as in endogenously \( \text{Y}_2\)R expressing SH-SYSY and SMS-KAN cells. Results were evaluated by one-way ANOVA and Tukey post test.

Results: We observed strong desensitization of the \( \text{Y}_2\)R in a second round of stimulation despite its reappearance at the membrane. Already the first activation of the \( \text{Y}_2\)R leads to depletion of the functional cellular \( \text{G}_\alpha \text{i/o} \) protein pool and consequently desensitizes the linked signal transduction pathways, independent of receptor internalization. This desensitization also extends to other \( \text{G}_\alpha \text{i/o} \)-coupled GPCR and can be detected in transfected HEK293 as well as in SH-SYSY and SMS-KAN cell lines, both expressing the \( \text{Y}_2\)R endogenously. By overexpression of chimeric \( \text{G}_\alpha \text{i} \) proteins in a model system, activation has been rescued, which identifies a critical role of the \( \text{G} \) protein status for cellular signaling. Furthermore, \( \text{Y}_2\)R displays strong allosteric coupling to inhibitory \( \text{G} \) proteins in radioligand binding assays, and loses 10-fold affinity in the \( \text{G} \) protein-depleted state observed after activation, which can be largely abrogated by overexpression of the \( \text{G}_\alpha \text{i} \)-subunit.

Conclusion: The unusually persistent \( \text{G}_\alpha \text{i} \)-signaling of the \( \text{Y}_2\)R leads to a state of cellular desensitization of the inhibitory \( \text{G}_\alpha \)-pathway. The strong allosteric effects of the \( \text{Y}_2\)R-\( \text{G}_\alpha \)-interaction might be a mechanism that contributes to the burst of \( \text{G}_\alpha \)-signaling, but also serves as a mechanism to limit the \( \text{Y}_2 \)-mediated signaling after recycling. Thus, the cell is left in a refractory state, preventing further \( \text{G}_\alpha \)-signaling of the \( \text{Y}_2\)R itself but also other \( \text{G}_\alpha \text{i/o} \)-coupled receptors by simply controlling the repertoire of downstream effectors.

Keywords: Neuropeptide Y, Neuropeptide Y receptors, GPCR, Desensitization, Internalization, G protein

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Background

G protein-coupled receptors represent a major family of cell surface receptors with approximately 800 different subtypes that share a common architecture of seven transmembrane helices connected by three intra- and extracellular loops. Members of the GPCR transduce a large spectrum of extracellular signals and consequently different regulatory mechanisms are fundamental to protect cells against overstimulation. One of the major processes limiting GPCR signaling is the reduction of receptor sensitivity towards a particular stimulus over time. This mechanism, defined as desensitization, includes a complex series of events e.g. receptor phosphorylation, arrestin-mediated internalization, receptor recycling, lysosomal degradation and decrease in mRNA levels [1–3]. However, the regulatory mechanisms beyond the sensitive equilibrium of receptor activation and desensitization are complex and likely distinct for individual receptors. These mechanisms are of great interest since many pharmacological agents targeting GPCR display diminished effectiveness over time [4–7]. Moreover, such knowledge is required for applications that use GPCR as a shuttle system for intracellular drug delivery [8, 9]. To date, approximately 35% of approved drugs address GPCR and the number of promising targets increases steadily [10]. This includes the neuropeptide Y hormone receptor family, consisting of four receptor subtypes — Y1R, Y2R, Y3R, and Y4R. Activated by the three endogenous ligands neuropeptide Y (NPY), peptide YY (PYY) and pancreatic polypeptide (PP), the Y receptors form a multi-ligand/multi-receptor system and contribute to a large variety of physiological processes within the human body. Beside the regulation of food intake [11] and gastrointestinal secretion, they also control blood pressure [12] and are involved in the pathophysiology of cancer progression as well as mood disorders [13, 14]. The Y2R, which is predominantly expressed in neuronal but also in peripheral tissue like liver, blood vessels and spleen, plays a central role in the development of new therapeutic drugs. Regarding its anorexigenic properties and its overexpression on specific tumor subtypes (glioblastoma; neuroblastoma) the Y2R represents a promising target for the treatment of obesity as well as for selective addressing of malignant tissue [15, 16]. However, intense Y2R desensitization was identified in mice and rats after daily treatment with a selective metabolically stable PYY13–36 analog [17]. This study indicates the importance of understanding the underlying mechanisms that controls and regulates receptor activity to improve the effect of administrated drugs. Previous investigations based on Y2R endocytosis and intracellular trafficking demonstrated an arrestin-dependent internalization, subsequent endosomal sorting and transport back to the cell membrane, which is highly regulated through distinct motifs within the C-terminus [18]. Moreover, Wanka et al. revealed different binding modes of arrestin 3 (arr-3) at the human Y1R and Y2R [19]. In consequence of the “tail” conformation, Y1R binds Gαq-protein as well as arr-3 simultaneously forming a supercomplex. In contrast, no supercomplex formation was observed for the Y2R. Owing to the “core” conformation, binding of arrestin to Y2R results in the dissociation of G protein, thereby terminating both the binding of the G protein to the receptor as well as the G protein-mediated signaling. Based on these findings we investigated the desensitization process of the human neuropeptide Y2R and identified a novel mechanism for signal suppression. We demonstrate here that activation of the Y2R results in an unusually persistent Gαq-mediated signaling, which is facilitated by strong allosteric coupling of the receptor to inhibitory G proteins and is terminated by the depletion of the functional cellular G protein pool. This leads to a state of cellular desensitization of the inhibitory Gαq-pathway for both the recovered Y2R as well as other Gαq-coupled receptors, protecting the cells against overstimulation by limiting the strong Y2R-mediated inhibitory G protein signaling.

Methods

Peptides

All peptides were synthesized by automated solid-phase peptide synthesis using the 9-fluorenlymethoxy carbonyl/tert-butyl (Fmoc/tBu) strategy [20] and purified to > 95% homogeneity by preparative HPLC. Analyzation and identification were performed by MALDI-ToF mass spectrometry (Ultraflex III MALDI ToF/ToF, Bruker, Billerica, USA) and reversed-phase HPLC using linear gradients of solvent B (acetonitrile + 0.08% trifluoroacetic acid) in A (H2O + 0.01% trifluoroacetic acid).

Plasmids

The Y2R within the pEYFP-N1 expression vector (Clontech) was used for fluorescence microscopy, cAMP-assay, Ca2+-assay, IP-one assay and binding assays. An internalization deficient Y2 receptor mutant (S374T376T379D-Y2R) was generated by QwikChange site-directed mutagenesis (Stratagene) using appropriate primer pairs. The Y2R-mutant within C-terminally fused pEYFP-N1 was used for fluorescence microscopy, cAMP-assay, Ca2+-assay. For co-transfection experiments Y1R within the pEYFP-N1 and MCR1 within the pEYFP-pV2 expression vector were used for cAMP-assay and Ca2+-assay. For cAMP readout, pGL4.29[luc2P/ CRE/Hygro] encoding for a luciferase reporter gene (luc2P) was used (Promega). Overexpression of Gαq was performed by using Gαq-Venus-pcDNA3 vector. Additionally, for measuring receptor activation using the Gαq pathway, the chimeric GoαsiplexR protein was used (kindly provided by E. Kostenis, Rheinische Friedrich-Wilhelms-Universität, Bonn, Germany) [21]. The identity of all plasmid constructs was verified by Sanger dideoxy sequencing.
Cell culture

All cell lines were cultured in flasks to confluence prior to use in a humidified atmosphere at 37 °C and 5% CO₂. HEK293 cells (human embryonic kidney) were maintained in Dulbecco’s modified Eagle’s medium (DMEM) with 4.5 g/l glucose and L-glutamine and Ham’s F12 (1:1, Lonza) supplemented with 15% (v/v) heat-inactivated fetal bovine serum (FBS). Stably transfected HEK293-HA-Y2R-eYFP were generated by transfecting 13 μg of linearized HA-hY2R-eYFP-pVitro plasmid with 20 μl Lipofectamin® 2000 (Invitrogen™) according to the manufacturer’s protocol. For culturing the same conditions as previously described were used and 100 μg/ml hygromycin B gold (Invivogen) was supplied to the medium to ensure stable transfection. SHSY5Y, human bone marrow neuroblastoma cells, derived from a parental SK-N-SH human neuroblastoma cell line [22] were maintained in DMEM with 4.5 g/l glucose and L-glutamine and Ham’s F12 (1:1, Lonza) supplemented with 15% (v/v) heat-inactivated fetal bovine serum (FBS) and 0.1 mM NEAA (Lonza). SMS-KAN cells were cultivated in Eagle’s Minimal Medium (EMEM; Lonza; Lonza) supplemented with 15% FCS, 4 mM L-glutamine (Lonza) and 0.02 mM nonessential amino acids (NEA) (Lonza). All cells were kept in a humidified atmosphere at 37 °C and 5% CO₂.

Short tandem repeat analyses were performed to verify cell line identity, and all cell lines were tested negative for mycoplasma contamination.

Live cell microscopy

HEK293 or HEK293-HA-Y2R-eYFP were re-seeded (150.000/well) into sterile poly-D-lysine covered μ-slide 8 wells (Ibidi) and grown in a humidified atmosphere at 37 °C and 5% CO₂. For transfection, cells were cultured up to 70–80% confluency and subsequently transfected with 1.0 μg total DNA using Lipofectamin® 2000 transfection reagent (Invitrogen) according to the manufacturer’s protocol. For single transfection in empty HEK293 cells 1.0 μg of HA-Y2R-eYFP-N1 plasmid DNA was used. At the experimental day cells were starved in Opti-MEM® reduced serum medium for 30 min, followed by stimulation with 1 μM TAMRA-NPY for 60 min and subsequently washed three times with PBS and incubated in recovery medium for indicated time periods. Gradually preparation of all samples was performed in 10 min intervals of 2% paraformaldehyde (PFA)/PBS on ice, 2% PFA/PBS at RT and finally 4% PFA/PBS at RT. Samples were blocked and permeabilized with 10% BSA/0.1% TritonX-100/PBS for 1 h at RT. Incubation with anti-EEA1 (Santa Cruz, sc-33,585) (1:100) in 1% BSA/0.01% TritonX-100/PBS occurred overnight at 4 °C. The next day, samples were treated with anti-rabbit-AF350 (InvitrogenTM) (1:400) in 1% BSA/0.01% TritonX-100/PBS for 2 h at RT and mounted with Fluoromount-G®.

Immunostaining

HEK293-HA-hY2R-eYFP cells were seeded on poly-D-lysine coated coverslips (500.000/well). After 24 h incubation under humidified atmosphere at 37 °C and 5% CO₂, cells were starved in Opti-MEM® reduced serum medium for 30 min, followed by stimulation with 1 μM TAMRA-NPY for 60 min and subsequently washed three times with PBS and incubated in recovery medium for indicated time periods. Gradually preparation of all samples was performed in 10 min intervals of 2% paraformaldehyde (PFA)/PBS on ice, 2% PFA/PBS at RT and finally 4% PFA/PBS at RT. Samples were blocked and permeabilized with 10% BSA/0.1% TritonX-100/PBS for 1 h at RT. Incubation with anti-EEA1 (Santa Cruz, sc-33,585) (1:100) in 1% BSA/0.01% TritonX-100/PBS occurred overnight at 4 °C. The next day, samples were treated with anti-rabbit-AF350 (InvitrogenTM) (1:400) in 1% BSA/0.01% TritonX-100/PBS for 2 h at RT and mounted with Fluoromount-G®.

IP-one assay

HEK293 cells were grown in 25 cm² culture flask and co-transfected with 800 ng plasmid encoding the Gαs-qi4myr protein and 3200 ng plasmid encoding the Y2R fused C-terminally to eYFP applying Metafectene® Pro (Biontex Laboratories GmbH) according to the manufacturer’s protocol. One-day post transfection, cells were respectively. Second stimulation experiments using 5-carboxytetramethylrhodamine (TAMRA)-NPY were performed by stimulating cells in a first round with 1 μM NPY, subsequent acidic wash and either direct stimulation in a 2nd round with 100 nM TAMRA-NPY or incubation in ligand-free media supplied with CHX and with or without NH₄Cl for 60 min prior to the 2nd round stimulation. All microscopy images were obtained using the AxioObserver.Z1 microscope equipped with an Apo-Tome imaging system (Zeiss, Jena). Within one experimental setup, all images were taken using a fixed exposure time for the single fluorescence channels.

Microscopy pictures were processed with the standard software Axio vision 4.8 and exported as an 8 Bit TIFF file (Tagged Image File Format). The open access software Image J was applied for the analysis of the pictures. For determination of mean cell surface fluorescence (MCSF) ten nonadjacent cells per image were measured using the segmented line function. Calculation of the relative TAMRA fluorescence, the Raw Intensity Density was measured under distinct conditions. The particular background fluorescence of each image was subtracted after every evaluation. Since this program uses the single gray levels for evaluation, the calculated values were evaluated retroactively as relative fluorescence intensity and analyzed with the statistical program GraphPad Prism.
seeded (75,000 cells/well) into white poly-D-lysine covert 96-well plates (Greiner Bio-one) and incubated overnight at 37°C. Receptor activation studies were performed by using the IP-one Gq assay kit (Cis-Bio). Prior to detection of IP- species; cells were either stimulated with buffer, 100 nM or 1 μM NPY for 60 min and washed with acidic wash buffer and HBSS. For recycling, cells were incubated for 60 min with assay buffer, supplemented with 100 μg/ml CHX (Merck/Calbiochem®). After recovery, the recycling medium was removed and 30 μl stimulation solution containing NPY in the concentration range of 10^{-13} M to 10^{-7} M and LiCl (inhibition of IP-species degradation) was added and cells were incubated for 1.5 h at 37°C. Stimulation was stopped by adding lysis buffer supplied with antibody 1 and antibody 2 according to the manufacturer’s protocol. After 60 min of incubation at room temperature, the emission at 620 nm and 665 nm was measured and the ratio (acceptor 665 nm/donor 620 nm) was calculated.

**Ca^{2+}- assay**

HEK293 cells were grown in 25 cm² culture flask up to 70–80% confluence in a humidified atmosphere at 37°C and 5% CO₂. Co-transfection of 800 ng plasmid encoding the GαΔg4myr protein and 3200 ng plasmid encoding for the Y2R fused C-terminally to eYFP was performed by using Metafectene® Pro (Biontex Laboratories GmbH) according to the manufacturer’s protocol. One-day post-transfection cells were re-seeded (100,000 cells/well) into a black poly-D-lysine covert 96-well plates (Greiner Bio-one) and incubated overnight at 37°C. Two-days post transfection Ca^{2+} experiments were performed with the help of the FLIPR® Calcium Assay Kit (Molecular Devices). Cells were stimulated in a first period either with buffer, 10 nM, 100 nM or 1 μM NPY for 60 min, followed by the removal of the peptide solution and subsequent acidic wash two times with acidic wash buffer (50 mM glycine, 100 mM NaCl, pH 3.0) and one neutralization step with Hank’s balanced salt solution (HBSS; PAA). For recycling, cells were incubated for 60 min with assay buffer (20 mM HEPES, 2.5 mM Probenecid in HBSS, pH 7.5) supplemented with 100 μg/ml CHX (Merck/Calbiochem®) and 1 μM Fluor-2-AM staining dye. For measurement, a subset of respectively stimulated cells was treated with a mixture of NPY in a concentration range 10^{-12} M to 10^{-6} M and 5 μM forskolin and incubated for 4 h at 37°C. As a positive control only 5 μM forskolin were added to the wells. After incubation, medium was exchanged by 30 μl fresh DMEM (RT). 30 μl ONE-Glo substrate was added and after 5 min of incubation the cAMP level was measured by detecting the luminescence. For co-transfection experiments the transfection of two receptors simultaneously and pGL4.29[luc2P/CRE/Hygro] plasmid were performed in a ratio 1:1:1, keeping the total amount of 6000 ng. For Y1/Y2 co-transfection, cells were treated with either 1 μM F^{34}NPY or Ahx^{5–24}NPY first, followed by the washing step and the recovery period. The second stimulation was performed in a peptide concentration range, addressing the respective receptor that was missed in the first round. Procedure was analogous concerning co-transfection with melanocortin 1 receptor, first stimulation was performed either with 1 μM NPY or 1 μM NAPamide.

Endogenous Y2 receptor activation was performed in SMS-KAN cells by using the Gs dynamic assay kit (Cis-Bio). Prior to detection of cAMP concentration, SMS-KAN cells were grown in 12 well plates in a humidified atmosphere at 37°C and 5% CO₂. At the experimental day, cells were stimulated first either with buffer (control cells) or 1 μM NPY for 60 min, followed by the removal of the
peptide solution and subsequent washing steps with acidic wash buffer twice (50 mM glycine, 100 mM NaCl, pH 3.0) and one neutralization step with HBSS (PAA). For recycling, cells were incubated for 60 min with DMEM supplemented with 100 μg/ml CHX. For cAMP measurement, the cells were detached from the 12-well plate by adding 150 μl stimulation buffer and gently using a cell scraper. Finally, a subset of 50,000 cells in an end volume of 5 μl was reseeded in a 384 well plate. Cells were treated with a mixture of NPY in a concentration range of 2 × 10⁻¹² M to 2 × 10⁻⁵ M (final concentration 10⁻¹² M to 10⁻⁵ M) and 10 μM forskolin (final concentration 5 μM) simultaneously and incubated for 1 h at 37 °C. As a positive control only 5 μM forskolin (final concentration) was added to the wells. Stimulation was stopped by adding 5 μl of cAMP-d2 and cAMP-cryptate working solution diluted in lysis- and detection buffer to each well and incubated at room temperature for another 60 min according to manufacturer’s protocol. After 60 min of incubation at room temperature, the emission at 620 nm and 665 nm was measured and the ratio (acceptor 665 nm/donor 620 nm) was calculated.

**Specific radioligand binding assay**

HEK293 cells were maintained in 25 cm² culture flask up to 70–80% confluency in a humidified atmosphere at 37 °C and 5% CO₂. Cells were transfected with 4000 ng plasmid encoding for the Y₂R using Metafectene® Pro (Biontex Laboratories GmbH) according to the manufacturer’s protocol. For Gαi-overexpression experiments cells were transfected with 3000 ng plasmid encoding for the Y₂R and 1000 ng plasmid encoding for the Gαi₂-subunit. One-day post-transfection cells were seeded into poly-D-lysine covert 48-well plates (Greiner Bio-one) and incubated overnight at 37 °C. Two-day post-transfection cells were treated either with buffer or 1 μM NPY for 60 min subsequently washed twice with acidic wash buffer and once with HBSS, followed by a recovery period in ligand free medium containing 100 μg/ml CHX (Merck/Calbiochem). After treatment, cells were immediately cooled down on ice, washed once with PBS and incubated with 6 × 10⁻¹¹ M human [¹²⁵I]-PYY in binding buffer for 4 h. Binding buffer consisted of OptimEM, 50 mM Pefabloc® SC, 1% BSA and for replacement NPY in a concentration range 10⁻¹¹ M to 10⁻⁹ M. After incubation, cells were washed twice with ice-cold PBS and lysed with 0.2 M NaOH. Lysates were transferred into scintillation cocktail and radioactivity was detected with the help of Microbeta2” counter.

**Statistical analysis**

Calculations of means, S.E.M. and statistical analysis were performed using PRISM 5.0 program (GraphPad Software, San Diego, USA). Significances were calculated according to one-way ANOVA and Tukey’s or paired, two-tailed t-test.

**Results**

Y₂R and NPY co-internalize but underlie different trafficking routes

Understanding regulatory mechanisms such as desensitization is fundamental to understand the endocytic and post-endocytic fate of a receptor. Previous investigation clearly demonstrated that human Y₂R internalizes by arrestin-dependent and independent mechanisms following high agonist concentration. Arrestin 3 is fully engaged by the C-terminus of the receptor and the transmembrane core, thus physically blocking G protein interactions and terminating signaling. Once internalized, the receptor re-appears at the cell membrane after agonist washout. To get more detailed information about the cellular transport mechanism, we first examined the post-endocytic sorting of the receptor and its agonist NPY. We used a stably transfected HEK293 cell line expressing the Y₂R N-terminally fused to hemagglutinin (HA), and carrying enhanced yellow fluorescent protein (eYFP) at its C-terminus. Enabled by the YFP-labeling of the receptor and an additional TAMRA-NPY variant, live cell fluorescence microscopy was applied to study the intracellular fate of the receptor after agonist stimulation as graphically illustrated in Fig. 1a. As shown in Fig. 1b, the receptors are expressed predominantly in the plasma membrane prior to agonist stimulation. For quantification, the mean cell surface fluorescence (MCSF) was determined and set to 100% (Fig. 1c, w/o). Stimulation of HEK293-HA-Y₂R-eYFP with 1 μM NPY led to strong internalization and accumulation into intracellular compartments (33 ± 2%, MCSF, white bar). Removing the stimulation solution and incubation in ligand free medium for 60 min resulted in a reappearance of receptor at the plasma membrane up to 69% (SEM ± 4%, light grey bar), which was inhibited by adding 20 mM NH₄Cl as recycling inhibitor. Co-internalization of the ligand-receptor-complex was observed by incubation with TAMRA-NPY (Fig. 1d/e). Immunostaining of the early endosome antigen EE1A displayed an intracellular accumulation of TAMRA-NPY in early and sorting endosomes, prior to further distribution (Fig. 1e). After ligand removal and during the recycling period, the amount of co-localized NPY-receptor-complexes decreased in early endosomes. While the receptor fluorescence at the cell membrane increased, the total amount of red peptide fluorescence decreased. Using a lysosomal stain, the reduction of peptide fluorescence was identified as a result of degradation in lysosomes (Fig. 1d). Accordingly, the Y₂R and its ligand pass through different intracellular trafficking routes. While the receptor recycles back to the plasma membrane, and should be available for a further activation, the ligand is degraded in lysosomes.
Reduced Y2R internalization in second stimulation experiments despite recycling

Next, we aimed to confirm the functionality of the recycled receptors by testing whether recycling represents a quick resensitization and qualifies the system as a suitable drug-shuttle. Thus, we examined the internalization properties in a second period of stimulation, which is summarized in a graphical scheme in Fig. 2a. Ligand uptake of HEK293-HA-Y2R-eYFP was measured by Raw Intensity Density using Image J. First stimulation of cells with 100 nM TAMRA-NPY resulted in high yields of internalized ligand after 60 min (Fig. 2b), which was set to 100% as control (Fig. 2c). Stimulation of the cells with 1 μM non-fluorescent NPY, intensive acidic wash and subsequent stimulation with 100 nM TAMRA-NPY led to an expected significantly reduced TAMRA-NPY uptake (45 ± 5%, 0 RE), due to receptor internalization and reduced cell surface receptor amounts. However, the peptide uptake was similar and not increased when the receptor was allowed to recycle after extending the recovery period up to 60 min (49 ± 4%, 60 RE). Next, the uptake of TAMRA-NPY was measured in presence of the recycling inhibitor NH4Cl during recovery period. No further reduction of internalized peptide was detectable (48 ± 46%, 60 RE + NH4Cl), confirming that the recycled receptors do not contribute to peptide uptake into the cell. This indicates a significantly reduced functionality of the recycled receptors with respect to internalization.
Y2R stimulation but not internalization leads to sustained reduction of cAMP response

To clarify whether the diminished endocytosis of recycled receptors is due to impaired arrestin recruitment and thus specific to endocytosis or rather a consequence of reduced receptor functionality in general, we investigated receptor activation in a next step by analyzing cellular cAMP levels (Fig. 3). Without NPY-stimulation, 5 μM of the adenylyl cyclase activator forskolin typically increases the baseline of intracellular cAMP concentration in Y2R-expressing HEK293 cells by ~ 10 fold (set to 100%), which is completely reversed by stimulation with NPY with an EC50 of 0.04 nM. Surprisingly, the maximal inducible cAMP response, induced by forskolin was dramatically reduced in cells with recycled Y2R (Fig. 3a). Stimulation with 100 nM or 1 μM NPY followed by agonist washout and 60 min recycling period (same procedure as used in microscopy experiments) led to a drop of cellular cAMP in response to 5 μM forskolin (18 ± 5%, 14 ± 9% respectively, Fig. 3a). Even stimulation with 10 nM NPY already resulted in ~ 60% reduction of the susceptibility of AC to Forskolin (41 ± 7%, Fig. 3a, brown curve). Moreover, we have confirmed our findings in a biologically relevant system by investigating the cellular cAMP concentration in SMS-KAN cells that have been isolated from a primary human brain tumor and endogenously express the Y2 receptor [23]. Interestingly, we observed the same pattern of sustained Gαi-signaling and diminished cellular activatability after stimulation with 1 μM NPY in this transfection-free cellular model (Fig. 3b). The maximally inducible cAMP response with 5 μM forskolin was also significantly reduced in cells with recycled Y2R and did not reach the bottom-line of control cells.

Next, we were interested whether this behavior is internalization dependent. Fluorescence microscopy experiments of Y2R cells stimulated with 10 nM NPY displayed no internalization at this concentration (Figure S1), indicating an internalization independent mechanism. To confirm this hypothesis, we generated an internalization deficient mutant, exchanging the important phosphorylation residues [S374,T376,T379] to aspartate within the C-terminus [18]. This Y2R variant exhibited the same loss of activity as the wild type (Fig. 3c), excluding cellular processes accompanying receptor endocytosis as a cause of these findings. Next, we co-transfected cells with Y1R and Y2R simultaneously and compared the effects of receptor stimulation. The use of selective peptide agonists allows the addressing of the distinct receptors. Stimulation with either the receptor selective agonist for Y1R F7, P34-NPY or for Y2R Ahx5-24-NPY resulted in a robust receptor...
Fig. 3 (See legend on next page.)
activity with EC_{50}-values in the nanomolar range (F^{24}, p^{34}-NPy: 4.4 nM, Ahx^{24}-NPy: 14 nM, orange and dark blue line respectively), demonstrating the functionality of the assay setup. By stimulation with 1 μM F^{24}, P^{34}-NPy to address Y2R, followed by stimulation with the Y2R selective Ahx^{24}-NPy after 60 min recovery period, no significant changes in the forskolin-inducible cAMP signal or receptor potency (Fig. 3d, Emax = 74%, EC_{50} = 28 nM, light blue dashed line) was observed. However, addressing the Y2R first by using 1 μM Ahx^{24}-NPy and second stimulation with the Y2R selective agonist resulted in a significant loss of forskolin-inducible cAMP (Fig. 3d, Emax = 13% orange dashed line). As an additional control, we used melanocortin 1 receptors (MCRI), which endogenously couple to Gαi-protein, to test the influences on other downstream signaling pathways. Melanocortin receptors are robustly activated by NAPamide, a modified MCRI agonist, also after stimulation of co-transfected Y2R with 1 μM NPY. The maximal cellular cAMP levels however, were slightly reduced by ~17% (Emax = 83 ± 10%, green dashed line) compared to the control without Y2R stimulation, indicating that the inhibitory effects of Y2R on cellular cAMP are still present (Fig. 3e). These data suggest a Y2R-specific Gαi-mediated mechanism, independent of other Ga-signaling pathways. To confirm this, pertussis toxin (PTX) was used, a Gαi sensitive exotoxin that prevents interaction with the G protein and its receptor [24, 25]. Different concentrations were tested first probing the optimal in vitro condition (Fig. 3f). All further experiments were performed using 250 ng/ml PTX. Cell toxicity was examined by microscopy studies and excluded, since no differences in cell morphology were observed (Fig. 3g, images). The presence of PTX clearly reduced the maximal cAMP response induced by forskolin after treatment of cells with NPY (Fig. 3g). Furthermore, applying PTX also on MCR1/ Y2R co-transfected cells supported Gαi-specific effects by abolishing the slight inhibitory effect, seen in the experimental setup, when Y2R was stimulated with NPY first (Fig. 3e, right panel). These data demonstrate that Y2R stimulation and activation leads to a long-lasting inhibition of Gαi-activated effector proteins and consequently downstream in the Gαi-pathway.

High G protein turnover of Y2R is terminated by depleted intracellular G protein pool

These data raise the question, whether this sustained inhibition of Gαi-pathway is due to a specific and exceptionally tight interaction between Y2R-activated Gαi-protein and the adenylyl cyclase (AC) or whether a very high G protein turnover and thus high number of active Gαi-GTP is responsible for the phenomenon. We took advantage of the chimeric Gα_{D_{f}i4myr} protein (Gα_{D_{f}i}) as a tool system. This chimeric Gα-protein couples to G_{i}-preferring receptors, but addresses cellular effectors of the Gαi-pathway and activates phospholipase C, which is directly measurable by the increase of cellular inositol phosphate (IP) and Ca^{2+}-influx [21]. Moreover, the chimeric G protein is co-transfected with the receptor, whereby modulation of both receptor and G protein expression levels and ratio by transfection provides an excellent assay setup. First, we tested receptor activity using both IP accumulation assay and Ca^{2+}-assay, as a live imaging assay, since the Gα_{D_{f}i} re-routes the native Gαi-pathway to the phospholipase pathway. Both assay setups were suitable for measuring receptor activity and showed similar EC_{50} -values compared to the endogenous signaling pathway detected with cAMP accumulation assay. In accordance with our previous results, the measured activity of stimulated and recycled receptors displayed an impaired functionality comparable to cAMP accumulation assay (Fig. 4a/b).

For the IP accumulation assay, we first stimulated the Y2R with either 100 nM or 1 μM NPY, followed by an extensive washing step and recovery period in ligand free medium. All pre-treatment and recovery steps were...
conducted in the absence of lithium chloride to ensure normal cellular IP turnover. Nonetheless, we found a sustained IP-response, since the initial IP-level for NPY stimulation is still at 60% and does not reach the basal IP-level after recovery period again (Fig. 4a, red curve/bar). This effect is even higher for stimulation with 1μM NPY (Fig. 4a, olive curve/bar).

These results support our findings of a sustained Gαi signal transduction is reduced but G protein overexpression counteracts the effect in HEK293 cells. A Concentration-response curves of transiently co-transfected HEK293-hY2R+Gαqi were measured by IP-accumulation assay, using the chimeric G protein as a tool. Prolonged IP signal was observed after first stimulation with 100 nM (red) or 1μM (olive-green), subsequent washing and 60 min recover of receptor in ligand free media compared to control curve without stimulation (w/o, black). B Decreased Ca2+ response from transiently transfected HEK293-hY2R-eYFP cells was obtained after stimulation with either 10 nM (brown) or 1μM (olive-green) for 60 min at 37°C followed by subsequent washing and 60 min recycling. C Overexpression of G protein and modifying the ratio between receptor and G protein led to a partial gain of Ca2+ signal after stimulation with 1μM NPY, followed by washing step and 60 min recovery. Fluorescence intensity was investigated to quantify the receptor-eYFP fusion protein. Data were confirmed by testing Gαqi-signaling with SH-SY5Y cells that endogenously expressing Y2R, and were transiently transfected with the chimeric Gαqi protein. We observed an attenuated Ca2+-response for receptors stimulated either with 1μM NPY (olive-green) or 1μM Ahx24-24-NPY (light grey), a selective Y2R agonist, and stimulated with NPY in a second experiment. Treatment with F7-P34-NPY, a selective Y1R agonist (berry) proved Y2R specificity. Experiments represent data n ≥ 3; significance was determined by one-way ANOVA, Tukey post test, ns: not significant, ***: P < 0.0001

In the next step, we performed Ca2+-measurements downstream of the chimeric Gαqi protein. Beside the fast rise of intracellular Ca2+-levels, an advantage of measuring Ca2+-influx after receptor stimulation is the rapid decrease and resetting to the basal Ca2+-levels. Thus, the increased cellular Ca2+ levels after the first stimulation remain “invisible” and the measurements give access to the actual amount of G protein activation after the receptor recovery period. The baselines of control and after the first round of stimulation start at a comparable intensity contrary to IP-one accumulation assay and thereby the assay window is magnified.
Stimulating cells in a concentration-response range resulted in a robust receptor activity response curve with EC_{50}-values in the nanomolar range (Fig. 4b). Interestingly, also in this transient measurement, we found a loss of activity after the first stimulation. Stimulation first with 10 nM NPY resulted in 40% reduced Ca^{2+}-influx (Fig. 4b, 62% ± 6), which is even more reduced after stimulation with 1 μM (Fig. 4b, 21% ± 1%). Taken together, our findings suggest that a very high G protein turnover of the NPY-activated Y_{2}R depletes the functional cellular G protein pools, limiting signaling responses even after agonist washout and a recovery period of 1 h. To clarify whether a longer recovery period might improve Y_{2}R functionality after the first stimulation, we also modified and prolonged the recycling period up to 6 h. However, even under these conditions the signal did not exceed 50% of control cells, which further underlines the strong and persistent downregulation of receptor activity (Figure S2).

To corroborate our hypothesis, we altered the ratio of receptor to G protein in the cell. Relative overexpression of G_{αq6q4myr} leads to a partial rescue of the receptor activation after receptor recycling (Fig. 4c, olive green lines). Expression levels of the receptors were quantified by fluorescence measurements using the C-terminally fused YFP and are displayed in Fig. 4c (right panel). Only G_{αq} transfection served as negative control, while the commonly used condition 4:1 (3200 ng receptor: 800 ng G protein) was set to 100%. Even under conditions with 17-fold excess of G protein at DNA level (0.06:1) and a receptor-eYFP fluorescence minimally over untransfected control, the maximal signal in the second stimulation with NPY (olive green lines) did not exceed 50% of control cells (black lines). However, an EC_{50}-shift was also observable, comparing to control receptors (black line) in different transfection conditions, suggesting a sensitive assay-system with a high number of reserve receptors. By reducing the amounts of the receptors by decreased plasmid transfection the number of possible cell surface signaling receptors decline, which is reflected in the rightward shift of the concentration-response curve. Thus, this experiment clearly demonstrates that the availability of the G protein becomes the limiting factor after Y_{2}R stimulation in the second receptor activation. We further excluded that cellular Ca^{2+}-stores are the limiting factor, as i) Y_{2}R activation mainly triggers Ca^{2+}-influx from extracellular compartments rather than from cellular compartments (Figure S3A) and ii) stimulation in Y_{2}R/Y_{4} co-expression systems displays no signal reduction of PP (endogenous Y_{2}R ligand), but a strong loss for NPY by addressing the Y_{2}R (Figure S3B).

To confirm this mechanism in a more biologically relevant setting, we transferred the Ca^{2+}-influx experiments to human SH-SY5Y cells, which are derived from a parental SK-N-SH human neuroblastoma cell line and endogenously express the neuropeptide Y_{2}R [22, 26]. In a first step, we aimed to verify selective Y_{2}R signaling by treating the cells either with Y_{2}R specific agonist Ahx^{5–24}-NPY or Y_{1}R specific agonist F^{7}, P^{34}-NPY. Stimulation of the cells with the selective Y_{2}R agonist Ahx^{5–24}-NPY resulted in a robust receptor activation (Fig. 4d, dark grey curve, EC_{50} 41.9 nM), whereas no Ca^{2+}-signal was detectable when the cells were treated with selective Y_{1}R agonist F^{7}, P^{34}-NPY (Fig. 4d, berry curve, EC_{50} > 1000). In addition to Y_{1} and Y_{2} receptors, NPY can also signal through the related Y_{4} and Y_{5} receptors. To confirm exclusive Y_{2}R expression in SH-SY5Y cells we used PP as Y_{4} receptor subtype and [Ala^{31}, Aib^{32}]-NPY as Y_{5} receptor subtype selective ligands, and investigated the activity of these peptides in Ca^{2+}-flux assays [27, 28]. In agreement with literature [22, 26], these peptides displayed no activity in the endogenous background of SH-SY5Y cells (Figure S4).

As already observed in HEK293 cells, we detected a similar loss in activity of endogenous Y_{2}R after the first stimulation with both 1 μM NPY (olive green line/bar, E_{max}: 40 ± 7%, EC_{50}: 34 nM) and 1 μM Ahx^{5–24}-NPY (light grey line/bar, E_{max}: 56 ± 6%, EC_{50}: 10 nM) respectively. Thus, we demonstrate that the Y_{2}R specific effects on intracellular signaling after a first stimulation are not due to the overexpression in commonly used cell lines, but similarly can be found in a biological context.

**Different ligand affinity states serve as feedback mechanism and control Y_{2}R signaling**

Allosteric modulation and regulation is a natural principle to modify the activity of molecules and enzymes in cell signaling. The strength of attraction between a receptor and its ligand – defined as affinity – is crucial for transducing signals. Strong allosteric effects of G protein binding for ligand affinity have been suggested before [29]. Hence, we investigated whether changes in the cellular G protein pool are reflected in the binding properties. To assess ligand-receptor affinity and compare Y_{2}R binding prior (w/o, black line) and after stimulation with 1 μM NPY (olive green line), we performed specific [^{125}I]-PYY competition radioligand binding assay using transiently transfected Y_{2}R in HEK293 cells. The following experiments were performed on ice to prevent receptor internalization and thus allow for equilibrium binding. Specific binding of control receptors that were incubated with buffer represents 100% of possible binding sites (Fig. 5a, black bar). Interestingly, ligand binding of recycled receptors, treated with 1 μM NPY for 60 min, followed by acidic wash and recovery in ligand free medium for 60 min still showed a significant reduction in B_{max} (Fig. 5a, olive green bar, 32 ± 8%), contrary to our observation in microscopy studies that revealed a recycling rate approximately up to 70%. We attribute this to experimental limitations due to the low radioligand concentration compared to the
micromolar concentration of peptide used for stimulation. For normalization, total binding and unspecific binding of each condition were constrained to 100 and 0% respectively. NPY displacement experiments show a biphasic binding mode with a high affinity and a low affinity state (Fig. 5b, left panel, w/o black line), which is slightly more pronounced compared to the experiments with isolated membrane preparations [29]. Strikingly, the high affinity state was lost after first stimulation with 1 μM NPY (Fig. 5b, left panel, olive green line). While evaluating the two site fit calculations, control receptors clearly distribute into a high affinity state with a percentage of 45% and an IC50[high] of 0.3 nM and a low affinity state with an IC50[low] of 40 nM. Analyzing recycled receptors, the calculated IC50[high] and IC50[low] is 30 nM and comparable when applying one site fit (IC50 35 nM, data not shown), indicating an impaired ligand affinity due to the loss of a high affinity receptor conformation. To address the question, whether the affinity shift is a result of G protein depletion, we co-transfected Gαi2 with the Y2R (Fig. 5b, right panel). In consequence of the overexpression, an increase in the proportion of the high affinity state already for the control receptors were observable (Fig. 5b, right plot, w/o, black line, fraction [high]: 65 ± 7%, IC50[high]: 0.4 nM, IC50[low]: 59 nM). Moreover, the binding properties of recycled receptors were altered positively (Fig. 5b, olive green line). Applying the two site fit model under Gαi overexpressed condition subdivided also recycled receptors in a high-affinity state [IC50[high]: 5 nM] with a considerable share of 75 ± 19% (fraction[high]) and only a minor fraction of receptors in a low affinity state with a calculated IC50[low] of 270 nM. Taken together, overexpression of Gαi shifts the percentage of the receptor towards the active state as shown in Fig. 5c. Thus, the binding experiments support our finding that the G protein pool is depleted after Y2R stimulation, which limits further receptor-ligand binding and downstream cellular signaling.

**Discussion**

Understanding the regulatory mechanisms that control the interplay of receptor activation, receptor trafficking and signal transduction are key processes for the development of new therapeutic approaches. Interestingly, the underlying regulatory processes reveal no consistent scheme and can affect receptor sensitivity in a variety of ways [6, 30, 31]. The classical ligand-induced activation of GPCR is divided into three major steps and can shortly be summarized as i) ligand binding and stabilization of active receptor conformations, ii) binding and activation of the G protein and iii) phosphorylation of the activated receptor targeting for desensitization [32]. The latter protects the cells against chronic and acute overstimulation by fine tuning receptor sensitivity upon external stimuli [33, 34]. Since the neuropeptide Y2 receptor is involved in multiple physiological and pathophysiological processes, unraveling of this mechanism leads to a better comprehension of GPCR signaling and might improve the development of therapeutic drugs by limiting side effects [11–14]. For the Y2R, a typical pattern has been reported, which involves arrestin-depended internalization after stimulation with its endogenous ligand NPY [19, 35, 36]. Surprisingly, we identified in this study that despite the ligand-dependent internalization, the downstream Gαi pathway remains "turned on", which is reflected in severely depressed cellular cAMP levels or vice versa elevated inositol phosphate levels when using chimeric Gαi protein. Moreover, ineffectiveness of adenyl cyclase activator forskolin to counteract low cAMP levels and re-elevate cellular cAMP amount was observed as well, raising the question of an additional regulatory mechanism.
The concept that GPCRs internalize and continuously activate G proteins was demonstrated first by the experimental data of Shukla et al. They confirmed a large complex consisting of G protein, arrestin and receptor simultaneously [37]. Here, the G protein remains bound to the transmembrane core of the receptor while the arrestin interacts with the C-terminal part of the receptor without terminating signaling due to internalization [38]. However, based on this data, Wanka et al. recently demonstrated for the Y2R that both the receptor C-terminus and the transmembrane core are engaged in arrestin binding [19]. This so-called core-conformation mutually excludes G protein binding to the same receptor.

Nonetheless, our observations of a long-lasting G protein activation, which have been shown for inositol phosphate accumulation as well as for cAMP measurements are in contrast with these findings and doubt the classical role of arrestin-dependent internalization to hamper G protein signaling. Indeed, persistent G protein signaling was already reported for different members of the GPCR-family, e.g. thyrotropin receptor, parathyroid hormone receptors as well as sphingosine 1-phosphate receptor 1, but the prolonged activation was primarily found from intracellular compartments after internalization [39–41]. Since an internalization deficient Y2R mutant, lacking the important Ser374 and Thr376/379 residues within the internalization motif 373DSFTEATNV381 at the proximal C-terminus [18] displayed a similar activation pattern, we exclude intracellular signaling and rather suggest an additional regulatory mechanism concerning activation and desensitization that is delimited to the cell membrane and primarily independent of arrestin recruitment. Additionally, our hypothesis is strengthened because stimulation of the wild type Y2R with very low NPY concentration (10 nM) in a first round already resulted in a prolonged G protein activation. Comparable to the internalization-deficient mutant, the low NPY concentrations are insufficient for the recruitment of arrestin. These results are in accordance with previous investigation of Walter et al and Lindell et al, who postulated an arrestin-dependent internalization only at higher NPY concentration [18, 42]. The low affinity binding of arrestin towards the Y2R might provide the basis for a persistent G protein activation as a consequence of delayed arrestin recruitment and hence impaired termination of G protein signaling.

Moreover, another unexpected finding in our study is the non-responsiveness of the adenyl cyclase that was observed after NPY stimulation of the cells, since the global forskolin-dependent activation of AC was impaired. We speculate that the Ga12/13-inhibited AC5/6 subtypes are targeted by Ga1 signaling in HEK293 cells, as the cAMP signal was previously shown to be reduced by addition of CaCO3, and the cellular cAMP levels are elevated in the presence of a protein kinase C inhibitor, both of which is characteristic for these AC subtypes [43–45] (Figure S5).

Blocking Gi-signaling by treating the cells with PTX, the adenyl cyclase was still activatable by forskolin, indicating a receptor mediated regulation. We presume that long-lasting G protein activation of the Y2R and therefore the sustained inhibitory effect leads to controlled desensitization of the AC5/6 subtypes preventing overstimulation. However, signaling of co-transfected Ga12/13-coupled MCR1 was only very slightly affected after NPY treatment and stimulation with NAPamide still leads to a robust cAMP response. We suggest that additionally the forskolin-insensitive AC9 subtype contributes to the cAMP signal of this receptor. The AC9 subtype is characterized by its responsiveness to Ga12 but not to forskolin [43, 45]. Mullerhausen et al reported similar findings of AC desensitization of the sphingosine 1-phosphate receptor 1 and postulated that this phenomenon has been detected with several members of Gi protein-coupled GPCR and seems to be dependent on the amount of adenyl cyclase and Gs proteins [41, 46, 47]. Moreover Watts and Neve et al suggested that this so called heterologous sensitization of adenyl cyclase following receptor activation is independent of receptor desensitization, internalization and down-regulation and contributes to fundamental physiological processes within the neurotransmitter crosstalk [47]. However, at present, the biological role of Y2R-mediated inhibition of the cAMP system remains unknown.

Additionally to this desensitization of the adenyl cyclase/cAMP system, we found that after intracellular sorting and subsequent reappearance of the Y2R at the cell membrane, ligand binding, G protein activation and receptor internalization was diminished in the second stimulation as well, suggesting that receptor desensitization is regulated independently and controlled by additional factors.

To exclude receptor processing during the endocytosis and recycling route, we used the internalization deficient variant to further characterize desensitization of the receptors. Interestingly, our data clearly demonstrate a significant loss in receptor activation after repeated stimulation, although no endocytosis of the receptors occurred. Moreover, stimulation with 10 nM NPY (insufficient for receptor endocytosis) in the first experiment already resulted in a significantly impaired receptor activation in the followed set-up (Fig. 4c). Based on these findings, we hypothesize an additional desensitization mechanism that controls G protein signaling independently of receptor internalization. Indeed, several studies have already shown desensitization of receptors is not limited to receptor internalization. For different members of the GPCR family like m3 muscarinic-, endothelin- as well as adenosine A2a receptors an arrestin- and internalization-independent desensitization mechanism were already described [34, 46, 48, 49]. Moreover, specificity and selectivity of these data were confirmed by co-transfection of Y1 and Y2 receptors simultaneously. Selective stimulation of Y2R with Ahx5−25-NPY and subsequent activation of Y1R with F1P34-NPY resulted in a
significantly decreased Y1 receptor response, whereas vice versa selective first stimulation of Y1R and subsequent Y2R stimulation did neither affect receptor activation nor AC desensitization, indicating distinct “consumption” of effector proteins within one receptor family. Co-transfection with MCR1- a member of Goαi-coupled receptors – and subsequent stimulation of the Y2R did not lead to changes in the following cAMP signaling response of MCR1, indicating a mechanism specific for Goαi-signaling.

Contrary to the obviously low arrestin binding affinity, the unoccupied receptor apparently exists in a conformation that strongly favors G protein coupling. Kaiser et al. recently reported two different affinity states concerning Y2R binding properties [29]. Based on our radioligand binding experiments and in agreement with these findings, 125I-PYY ligand displacement experiments display a bi-phasic binding, including a high affinity state of ~ 40% receptors and a low affinity state. Notably, the high affinity state was apparently lost after stimulation with 1 μM NPY, which obviously correlates with the amount of available G protein. High affinity binding is a result of allosteric interactions between the G protein and ligand binding site of the receptor, and thus regulated by the association of G protein. Accordingly, decoupling of G protein-receptor-complex through desensitization is expected to lead to a loss in high affinity binding. Preassembly of the G protein and Y2R prior to agonist stimulation and disruption of the high affinity state by GTPγS was already suggested by Kaisser et al [29] and is consistent with our findings that demonstrate the shift to low affinity binding sites after first agonist stimulation. Similar results have been obtained for other Gi-coupled GPCR such as dopamine receptors and opioid receptors, indication a tight correlation between desensitization and changes in agonist-receptor-affinity states [5, 50–53]. Here, we further observed a dramatic decrease in total binding of pretreated receptors (Bmax 32%). Besides competing with remaining NPY due to inefficient washing, the loss in Bmax might be attributed to an insufficient ligand affinity in absence of preassembled G protein and would align well with the reported loss in total binding by addition of GTPγS [29]. As receptors that are exposed to agonist for the first time show a high affinity state, we propose that agonist binding results in a strong effector signaling leading to depletion of the intracellular G protein pool. This prevents reassembly of the R-G/ R*-G-complexes after agonist washout, resulting in prolonged desensitization, which is reflected in the loss of the high affinity state. Moreover, this hypothesis is supported by the previously described pattern of prolonged G protein signaling and is further corroborated by the over-expression of chimeric G protein. By varying G protein-receptor ratio towards an oversupply of G protein, we have confirmed a positive correlation with a higher receptor response and an increased Fmax after pretreatment with 1 μM NPY. Thus, very tight binding of the agonist leads to a long-lasting agonist-receptor-complex, which further activates all the nearby functional G proteins until the G protein pool is depleted. Hence, we postulate a new regulatory mechanism by which the Y2R functions as G protein magnet and captures all freely available G proteins. The high intrinsic affinity of the receptor to inhibitory G protein and strong allosteric connections between G protein and the ligand-binding site of the receptor contributes to very efficient activation and turnover of cellular G proteins, resulting in strong and persistent activation of the Goαi-pathway. This depletes the functional intracellular G protein repertoire before arrestin-mediated internalization can terminate signaling. Thus, the cell is left in a refractory state, preventing further Gi-signaling of both the Y2R itself but also other Goαi, -coupled receptors, suggesting that Y2R expression dominates Gi-signaling within the cell. Furthermore, our studies highlight that the availability of effector proteins critically affects the cellular signaling status, and simple depletion of a downstream effector adds to the stock of cellular control mechanisms. Up to now, the biological background remains unclear but we postulate an additional controlling mechanism within the presynaptic and postsynaptic transmitter crosstalk as these findings are not only observed in transfected HEK293 cells but also present in SH-SY5Y (bone marrow cells from metastatic neuroblastoma) and SMS-KAN (cells from primary brain tumor), both endogenously expressing the Y2 receptor. However, further investigations are necessary to completely unravel the pathway leading to a controlled receptor desensitization versus resensitization.

Conclusion

Our data demonstrate that activation of the Y2R results in a strong and persistent activation of the Goαi-pathway. A high intrinsic affinity of the receptor to inhibitory G protein and the strong allosteric interaction between G protein- and the ligand binding site of the receptor contributes to very efficient activation and turnover of cellular G proteins, which furthermore depletes the intracellular G protein repertoire before arrestin-mediated internalization can terminate signaling. Thus, the cell is left in a refractory state, preventing further Gi-signaling of both the Y2R itself but also of other Goαi,coupled receptors, suggesting that Y2R expression dominates Gi-signaling within the cell. Furthermore, our studies highlight that the availability of effector proteins critically affects the cellular signaling status, and simply depletion of a downstream effector adds to the stock of cellular control mechanism.

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10.1186/s12964-020-00537-6.

Additional file 1: Supporting Results. Figure S1. Internalization behavior of hY2 receptor wild type. Figure S2. Prolongation of the
Y2R activation mainly triggers Ca2+-influx from extracellular compartments and excludes Ca2+ as a limiting factor. **Figure S3.**

**Acknowledgments**

The authors gratefully thank for excellent assistance of Kristin Löbner, Janet Dammann and Ronny Müller.

**Authors’ contributions**

I.Z. and A.G.B-S. designed the study, with contributions from A.K., S.B. and K.M. on the experimental design. S.B. performed immunostaining assay and I.Z. and A.G.B-S. designed the study, with contributions from A.K., S.B. and K.M. All authors contributed to discussion and manuscript editing. I.Z. wrote the manuscript with support of A.G.B-S and with contributions from all authors.

All authors read and approved the final manuscript.

**Funding**

This work was supported by the German Science Foundation Deutsche Forschungsgemeinschaft (German Research Foundation) through CRC 1052, project number 209939838, subproject A03 and Z05, the European Union (EU) and the Federal State of Saxony.

**Availability of data and materials**

All other data needed to evaluate the conclusions in the paper are present in the paper or in the supporting Information. This manuscript is accompanied by supporting information including additional pharmacological analysis and characterization of Y2R. Even if these results are not essential to understand the manuscript, we think, it is adjuvant to underline the new findings.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**Received** 19 October 2019 **Accepted** 19 February 2020

**Published online:** 30 March 2020

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**Abbreviations**

ACN: Acetonitrile; AC: Adenylyl cyclase; ANOVA: Analysis of variance; Arr: Arrestin; cAMP: Cyclic adenosine monophosphate; CHX: Cycloheximide; DMEM: Dulbecco’s modified Eagle’s medium; eYFP: Enhanced yellow fluorescent protein; Fmoc: Fluorenylmethyloxycarbonyl; Go: Inhibitory G-protein; GPCR: G protein-coupled receptor; GTPγS: Guanosine 5′-O-(gamma-thio)-triphosphate; HA: Hemagglutinin; HBSS: Hank’s buffered salt solution; HEK: Human embryonic kidney; IP: Insolito phosphate; LIC: Lithium chloride; MALDI-ToF: Matrix-assisted laser desorption/time of flight; MCR1: Melanocortin 1 receptor; NH4Cl: Ammoniumchloride; NPY: Neuropeptide Y; PLC: Phospholipase C; PP: Pancreatic polypeptide; PTX: Pertussis toxin; PYY: Peptide YY; RP-HPLC: Reversed phase high performance liquid chromatography; TAMRA: 6-(6-carboxytetramethylrhodamine)Y2R; NPY receptor subtype i: i/o without. Without

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