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Introduction

The endocannabinoid system (ECS) is a complex lipid-based signalling network involved in a wide variety of physiological and cognitive processes such as pain regulation, immune

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Visualization of membrane localization and the functional state of CB₂R pools using matched agonist and inverse agonist probe pairs†

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The diversity of physiological roles of the endocannabinoid system has turned it into an attractive yet elusive therapeutic target. However, chemical probes with various functionalities could pave the way for a better understanding of the endocannabinoid system at the cellular level. Notably, inverse agonists of CB_2R – a key receptor of the endocannabinoid system – lagged behind despite the evidence regarding the therapeutic potential of its antagonism. Herein, we report a matched fluorescent probe pair based on a common chemotype to address and visualize both the active and inactive states of CB_2R , selectively. Alongside extensive cross-validation by flow cytometry, time-lapse confocal microscopy, and superresolution microscopy, we successfully visualize the intracellular localization of CB_2R pools in live cells. The synthetic simplicity, together with the high CB_2R -selectivity and specificity of our probes, turns them into valuable tools in chemical biology and drug development that can benefit the clinical translatability of CB_2R -based drugs.

response, appetite control, learning and memory formation, cardiovascular regulation, and addictive-like behaviour.¹ The ECS consists of two cannabinoid receptor subtypes (CB₁R and CB₂R) that belong to the class A G protein-coupled receptor (GPCR) family. Arachidonoylethanolamide (AEA) and 2-arachidonoylglycerol (2-AG) are the endogenous ligands of both CB1R and CB₂R.² Regardless of their high homology, the key difference between the two receptors is their distribution.³ CB₁R is predominantly expressed in the central nervous system with the highest density in the cerebellum, hippocampus, and cerebral cortex,⁴⁻⁶ while CB₂R is more abundant in peripheral organs, such as the spleen and tonsils, and is mainly expressed in cells associated with the immune system.3 It has been shown that expression of CB₂R is strongly upregulated under pathological conditions such as cancer,7,8 immunological disorders,9 inflammation, neurodegenerative diseases,^{10,11} and drug abuse.¹² Therefore, modulating CB₂R activation will be a valuable therapeutic approach for several diseases including inflammation, autoimmune and metabolic disorders, chronic pain, multiple sclerosis and cancer. For example, agonistmediated activation of CB2R was previously shown to be beneficial for neuroprotection in chronic neurodegenerative disorders such as Huntington's and Alzheimer's diseases.13 Conversely, inactivation of CB2R via an inverse agonist/ antagonist was found to have therapeutic potential for treatment of various diseases associated with neuroinflammation



Fig. 1 Chemical structures of CB_2R agonist 1 and inverse agonist 2 sharing a 5- and 6-substituted picolinamide core.

and the immune system.^{14,15} Due to different expression patterns of CB_2R and CB_1R , as well as their distinct functions, the selective activation or deactivation of CB_2R does not involve undesired psychotropic responses which is considered a major therapeutic advantage and makes it a more attractive target compared to CB_1R . However, despite its great potential, no CB_2R -selective drug has made its way to the market to date, as clinical translatability from preclinical models deduced from different species is currently challenging.¹⁴ This is largely attributed to the highly inducible nature and complexity of CB_2R signalling pathways at the cellular level and the unclear understanding of its expression, localization and function.¹⁶ For



Fig. 2 (A) Modification of linker attachment from previous work (compound 3; sulfur linker) led to the discovery of a new versatile exit vector (compound 4; amide linker), and the diethylglycine moiety as the centerpiece hub in red. (B) Docking poses of compounds **10b** (magenta sticks) and **11b** (blue sticks) within active (green; PDB: 6KPF) and inactive (orange; PDB: 5ZTY) states of CB₂R, respectively. The red dashed line indicates the approximate boundary of the lipid bilayers. For a detailed description of the docking studies see ESI, S3.†

example, a number of studies indicate that the cellular responses associated with CB₂R activation are not only limited to plasmalemmal receptors but also to the intracellular pools.¹⁷⁻¹⁹ The absence of CB₂R-specific monoclonal antibodies, which are important tools for obtaining expression data at a cellular or tissue level, is further aggravating this situation.

Recent CB₂R-selective agonist fluorescent probes²⁰⁻²² could partially fill these gaps by addressing the activated state and providing information on CB₂R localization, expression, target engagement, pharmacokinetics and dynamics in real-time; the scarcity of labeled CB₂R-selective inverse agonist probes has resulted in a lack of information on the distribution of intraand extracellular CB₂R pools in the inactivated state. There are only a limited number of reports about inverse agonist fluoprobes labeling CB₂R. For example, rescent the chromenopyrazole-based inverse agonist probe was originally generated from an agonist but upon attachment of the Cv5 fluorescent dye, the functionality was altered.23 The surface receptors of CB₂R expressing HEK-293 cells were labeled via the aforementioned probe.23 Another example is NIR-mbc94, an analogue of selective inverse agonist SR144528, which has been shown to be an imaging agent for the unbiased highthroughput screening of compounds interacting with CB₂R as a therapeutic target.²⁴ Despite the wide range of applications and a high demand for CB₂R inverse agonist probes, no versatile probe platform with diverse fluorophores is available so far.

We have previously reported a high-affinity, cell-permeable fluorescent CB₂R probe **3** based on a reverse-design approach using a preclinically validated drug-derived CB₂R agonist **1** (Fig. 1 and 2).²⁰ The probe successfully detected CB₂R in several *in vitro* and *in vivo* settings across species. For example, **3** was also recently used to visualize the high expression levels of CB₂R in primary neonatal microglia isolated from wild-type and Tg2576 mice, the latter being used as an Alzheimer's Disease (AD) model.²⁵

However, for any chemical probe approach, it is desirable to have access to a matched molecular pair of agonist and inverse agonist with high structural similarity which are correspondingly labeled. Such chemical probes are most suitable to address distinct mechanisms of action, *e.g.* by distinguishing the activated or resting state of the receptor or allowing differential analysis of agonist-stimulated internalization of the receptor, while excluding the cellular phenotype.

With the goal of expanding the scope of our probe platform and addressing both active and inactive states of the receptor, we designed a matched molecular pair of CB_2R agonist and antagonist fluorescent probes derived from highly similar chemotypes of advanced preclinical CB_2R agonist 1 (ref. 26) and inverse agonist 2 (ref. 27) drug candidates (Fig. 1). In addition, we were able to attach a variety of fluorescent dyes leading to CB_2R probes that span a broad range of physicochemical properties. At last, varying combinations of agonist and antagonist with cell-permeable (*e.g.* TAMRA) or impermeable (*e.g.* Alexa488) fluorophores gave us access to a valuable toolbox suitable for detecting extra- and intracellular receptor pools. To explore the spatial-temporal dynamics of CB_2R , we employed these novel probes to investigate the expression and subcellular localization of the active and inactive states of CB_2R in living cells, utilizing super-resolution confocal imaging techniques.

Results and discussion

Probe design and molecular modelling

Our previous probes were derived from a drug-like CB_2R agonist bearing a 5,6-substituted picolinamide **1** (Fig. 1). Interestingly, it was shown that different substitutions at the 5- and 6-positions of picolinamide could alter the functionality of the ligand while maintaining high CB_2R affinity.^{27,28} For example, the replacement of the cyclopropyl moiety at position 5 of agonist **1** with 3-methoxy-azetidine alters the functionality from agonism to inverse agonism (2). This substitution causes a flip of the side chain of the toggle switch residue W258^{6.48} (Ballesteros–Weinstein numbering in superscript, Fig. 2B).²⁹ Besides the 5,6substituents, both ethyl side chains are involved in favorable van der Waals interactions with surrounding phenylalanine side chains F91, F94, and F106.

The drug-derived inverse agonist 2 possessing an exceptional selectivity profile (CB₂R $K_i = 0.3$ nM; CB₁R $K_i = 721$ nM; a selectivity factor of 1403 over CB₁R) was an ideal starting point for generating a matched agonist and inverse agonist-based probe pair with a 5,6-substituted picolinamide core in common.^{28,30,31}

The first and the most critical step in probe design is the identification of a suitable attachment point between the recognition element and the reporter unit, *i.e.* the ligand and fluorescent dye, respectively. In most cases, the recognition element and the fluorescent dye are distanced using a suitable linker, which allows the dye to access the extracellular space without compromising overall binding affinity.32 Previously, we introduced a hybrid of thio- and polyether chains to one of the ethyl groups of the diethylglycine moiety as the centerpiece hub (3, Fig. 2A). Even though our previous probes showed highly consistent interspecies affinity and potency for both human and mouse CB_2R , the presence of the sulfur atom in the linker posed a possible experimental imponderability in some of the advanced settings, as sulfur might be prone to oxidation.^{20,33} Therefore, with the goal of improving the physicochemical properties and simplifying the synthetic strategy, SAR studies were performed to investigate alternative sites for linker attachment at the diethylglycine centerpiece hub. For this, we used the ester functionality which after substitution by an amide moiety served as an attachment point. This design approach has the advantage that no chiral center is present and the synthesis route is greatly simplified compared to our previous probes (4, Fig. 2A).

Docking experiments were conducted to estimate the required linker length to reach out into the extracellular space and support linkerology studies. In Fig. 2B the best docking poses for agonist (**10b**) and inverse agonist (**11b**) pharmacophores, respectively, are depicted. For both ligands, the proper range of polyethylene glycol PEG chains to access the extracellular space for insertion of the fluorescent dye was estimated to be n = 2.

Chemistry

The synthetic pathway to access the Boc-protected intermediates and target fluorescent probes bearing various dyes is outlined in Scheme 1. Fmoc-protected diethylglycine 5 was used as the centerpiece unit to connect the 5,6-substituted picolinamide recognition elements to the desired linker. In order to elaborate the optimal linker length for dye attachment, compound 5 was functionalized with a series of linkers 6ad with different lengths under HATU-mediated amide coupling conditions. Fmoc-protecting group removal of compounds 7ad using DBU was followed by coupling to agonist 8 or inverse agonist 9 precursors in situ to afford Boc-protected congeners with matched chemotypes (10a-d and 11a-d). Compounds 8 (ref. 20) and 9 (ref. 31) were synthesized according to literature protocols. The final step was to conjugate a variety of broadly used fluorophores such as 5/6-TAMRA, SiR, 5/6-Alexa488 and Alexa647 to the selected intermediates (10b and 11b) with a linker length of n = 2, which turned out to be optimal in subsequent SAR studies. For this purpose, the Boc-protecting group of 10b or 11b was first cleaved using TFA. Subsequently, the resulting free amines were coupled with the desired fluorescent dyes either under suitable amide coupling conditions to furnish probes **12–15** and **17–18**. Compound **16** was synthesized *via* a variation of the aforementioned synthetic route starting with coupling of 5/6-TAMRA-COOH to Bocdeprotected **7b** followed by another amide coupling reaction to **9** using HATU as the coupling reagent (ESI, S32†).

Evaluation of the appropriate linker length

To identify the optimal linker length, binding affinities of unlabeled precursors **10a–d** and **11a–d** were measured *via* a competitive radioligand binding assay on CHO membranes stably expressing hCB_1R or hCB_2R (Table 1).

For both agonist and inverse agonist chemotypes, the PEG chain with n = 2 (**10b** and **11b**, respectively) showed the highest affinity and selectivity for CB₂R and was therefore chosen as the optimal linker length. The observed trend confirmed the predictions based on our docking studies (Fig. 2B). To assess whether the attachment of the linker affects receptor function, the efficacy of selected precursors was determined in a cAMP assay (Table 1). Notably, **10b** and **11b** preserved partial agonist and inverse agonist activity, respectively, with high potency



Scheme 1 Synthesis of fluorescent probes. Reagents and conditions: (i) HATU, DIPEA, DMF, rt, 1 h; (ii) (1) DBU, HOAt, DMF, rt, 20 min; (2) 8 or 9, HATU, DIPEA, DMF, rt, 10 h; (iii) TFA (9 equiv.), CH₂Cl₂, 0 °C to rt, 3 h; (iv) for compounds **13**, **15**, **17** and **18**: HATU, dye, DIPEA, DMF, rt, 10 h; for compound **12**: EDC·HCl, HOAT, dye, DIPEA, DMF, rt, 10 h; for compound **14**: dye, DIPEA, DMF, rt, 10 h; for compound **16**: see the ESI.†

Cmpd	Linker length	K_{i}^{a} (nM)						
		hCB ₁ R	hCB ₂ R	Selectivity ^b	EC_{50} or IC_{50}^{c} (nM)	E_{\max}^{d} (%)	Function	Parent cmpd
1	n.a.	4.2	0.2	21	6.5	102	Agonist	n.a.
2	n.a.	721	0.3	1403	4.4	-130	Inverse agonist	n.a.
10a	n = 1	836	28	30	n.d.	n.d.	n.d.	1
10b	n = 2	466	6	78	49	83	Agonist	1
10c	n = 3	748	63	12	n.d.	n.d.	n.d.	1
10d	n = 4	1472	144	10	n.d.	n.d.	n.d.	1
11a	n = 1	>10 000	2061	>4	n.d.	n.d.	n.d.	2
11b	n = 2	>10 000	106	>94	88	-63	Inverse agonist	2
11c	n = 3	>10 000	>10 000	n.a.	n.d.	n.d.	n.d.	2

^{*a*} K_i (nM) values obtained from [³H]CP55940 displacement assays on CHO membranes stably expressing hCB₁R or hCB₂R. Values are means of at least three independent experiments performed in duplicate. For details, see the ESI. ^{*b*} Selectivity was determined by calculating the ratio of K_i (CB₁R)/ K_i (CB₂R). ^{*c*} The potency (EC₅₀ or IC₅₀) of the selected compounds was measured using cells stably expressing hCB₂R in homogeneous time-resolved fluorescence (HTRF) cAMP assay. The data are the means of four independent experiments performed in technical replicates. ^{*d*} Maximum effect (E_{max} in %) was normalized to reference full agonist **APD371**. n.a. is not applicable. n.d. is not determined.

(CB₂R cAMP EC₅₀ = 49 nM for **10b** and IC₅₀ = 88 nM for **11b**, respectively).

Pharmacological characterization of fluorescent probes 12-18

Compared to their unlabeled congeners, the binding affinity of the fluorescent probes **12–18** indicated fluorescent dye dependency which was not unexpected as the structural nature of the fluorophore alters the membrane interactions of the constructs. Overall, we observed a certain drop in affinity for all dye-bearing probes, consistent with previous results.^{20,21,34} This decrease is believed to mainly originate from additional configurational and steric penalties due to conjugation.³⁵ However, most of the probes retained high affinity and selectivity for CB₂R (ESI, Table S-2†).

In functional studies (Table 2), probes **13** and **15** showed full agonism with potencies (EC_{50}) of approximately 525 nM, while **12** and **14** showed partial agonism with higher potencies (EC_{50}) of approximately 80 nM. As anticipated, the functional mode of action of probes **16–18** fully retained their inverse agonism with IC₅₀ values in the range of 114–262 nM (Table 2 and Fig. 3).

To assess the specificity of the probes and scout for putative off-targets, TAMRA-probe pairs **12** and **16** were screened against a customized panel of 50 representative receptors and



Fig. 3 Inhibition of cAMP accumulation on hCB₂R was determined with a (HTRF) cAMP assay. The maximum effect (E_{max} in %) was normalized to reference full agonist APD371.

enzymes.³⁷ Both probes were devoid of any relevant off-target interactions thus confirming their suitability for specific CB_2R detection studies (ESI, Table S-1†). Furthermore, kinetic binding characterization of agonist **12** and inverse agonist **16** using HEK293-hCB₂R cell membranes revealed fast association and appropriate residence time (ESI, Fig. S-8†).

Table 2hCB2R potency and predicted log $D_{7.4}$ of fluorescent probes 12–18										
Cmpd	EC_{50} or IC_{50}^{a} (nM)	E_{\max}^{b} (%)	Function	Parent cmpds	Dye	$c \log D_{7.4}^{c}$				
12	82	80	Partial agonist	1/10b	TAMRA	3.7				
13	528	127	Full agonist	1/10b	SiR	3.9				
14	77	79	Partial agonist	1/10b	Alexa488	-0.5				
15	523	129	Full agonist	1/10b	Alexa467	-0.9				
16	114	-29	Inverse agonist	2/11b	TAMRA	3.6				
17	129	-29	Inverse agonist	2/11b	SiR	2.2				
18	262	-26	Inverse agonist	2/11b	Alexa488	-0.9				

^{*a*} The potency (EC₅₀ or IC₅₀) of fluorescent probes **12–18** was measured using cells stably expressing hCB₂R in homogeneous time-resolved fluorescence (HTRF) cAMP assay. The data are the means of four or seven independent experiments performed in technical replicates. ^{*b*} Maximum effect (E_{max} in %) was normalized to reference full agonist APD371. ^{*c*} For computational calculation of $c \log D_{7.4}$ see ref. 36.

For high probe quality, a lower lipophilicity is crucial, as it significantly reduces non-specific binding.³⁸ Therefore, it is noteworthy that all predicted $c \log D_{7.4}$ values of our drug-derived probes showed an overall significantly lower lipophilicity compared to phytocannabinoids²¹ and are in a favorable drug-like range (Table 2). Moreover, except for highly ionized sulfonated fluorescent dyes such as Alexa488 and Alexa647 (14, 15 and 18), attachment of TAMRA (12 and 16) and SiR (13 and 17) did not lead to a significant change in the lipophilicity of the final probes compared to their unlabeled counterparts (10b $c \log D_{7.4} = 3.8$ and 11b $c \log D_{7.4} = 3.2$).

Cellular imaging by flow cytometry and confocal microscopy in live cells

The probe's specificity and suitability for imaging were further validated by flow cytometry. All the fluorescent probes were incubated at various concentrations ranging from 0.014 to 10 μ M – with live CHO cells overexpressing hCB₁R or hCB₂R and wild-type (wt) CHO cells as the control. Despite some differences observed in the mean fluorescence intensity (MFI) of probes bearing the same dyes, most of the tested probes indicated decent selectivity and specificity for hCB₂R in flow cytometry, suggesting their suitability for imaging applications.

Probes carrying sulfonated dyes such as Alexa488 **14** and Alexa647 **18** showed high selectivity even at very high concentrations, which is in line with our previous data (ESI, Fig. S-1†).²⁰ To exclude unspecific binding, we further examined the effect of preincubation of cells with high-affinity competitor ligands such as agonist **JWH133** (ref. 39) and inverse agonist **RO6851228** (ref. 40) on probe **15** binding (ESI, Fig. S-2†). Both ligands competed with **15** in a dose-dependent manner confirming a high target specificity of **15** for CB₂R.

Based on the high specificity of the novel probes we continued our investigation by performing time-lapse confocal microscopy experiments to visualize hCB₂R in live CHO cells. Fig. 4 displays the frames of CHO cells overexpressing hCB₂R and hCB1R along with parental CHO cells 10 min after administration of probes 12, 14-16 and 18. All probes selectively stained CB₂R on the cell membrane, but also intracellular staining was observed to some extent, especially for probe 12. We monitored the staining of 12 in a time-lapse setting and could follow the permeation kinetics through the outer cellular membrane over time, clearly evidencing internalization (ESI, Fig. S-4[†]). For selected probes, blocking experiments with competitive non-labeled ligands RO6851228 (ref. 40) (CB2Rinverse agonist) and/or RO6871304 (ref. 41) (CB₂R-agonist) were carried out as well (ESI, Fig. S-3[†]). Both unlabelled ligands completely blocked the staining of all tested fluorescent probes in hCB₂R expressing CHO cells, confirming the high binding specificity and being fully consistent with the results obtained for WT-CHO and hCB1R expressing CHO cells.

To further validate the results obtained for CHO cells we continued with imaging human-derived live HEK293 cells expressing hCB_2R . We followed the probe 12 distribution in the same manner over 10 min (ESI, Fig. S-5†). Probe 12 exhibited a similar distribution pattern in the human-derived HEK293



Fig. 4 Time-lapse confocal microscopy in live cells. The wild type (wt) and overexpressing hCB₂R- and hCB₁R-CHO cells were co-stained with probes 12, 14–16 and 18 and Hoechst 33342 (blue, nucleus counterstain). The images were recorded 10 min after probe incubation. Images are representative of two independent experiments. Scale bars, 10 μ m.

cells as in the CHO system, predominantly staining cellular membranes but also the intracellular compartments.

We then set out to visualize endogenous CB_2R expression aiming to ascertain different patterns of activation and localization of CB_2R pools induced by agonist and inverse agonist binding by using non-transfected primary cultures of human macrophages derived from healthy donors. Upon staining of human macrophages with probes **12** and **16** a markedly different distribution pattern was observed (ESI, Fig. S-6†). While agonist probe **12** was widely distributed within the cell, inverse agonist probe **16** was mainly localized on cellular membranes and partially in perinuclear dots, suggesting a distinct localization of inactive and activated CB_2R pools.

Super-resolution confocal imaging of hCB₂R-CHO cells

To examine in more depth the localization and activation states of the CB_2R pools, we resorted to super-resolution confocal imaging of hCB_2R overexpressing CHO cells. First, we assessed the intensity and subcellular distribution of fluorescence



Fig. 5 Super-resolution confocal imaging of overexpressing hCB₂R CHO cells. The cells were stained for 15 min with (A) **12** (0.8 μ M, red) or (B) **16** (0.8 μ M, red) and Hoechst 33342 (blue, nucleus counter stain). Cells were optically sectioned using confocal laser-scanning microscopy equipped with an Airyscan detector. For quantifying the relative levels of labeling using the two TAMRA probes, identical imaging settings (*i.e.*, objective, light path, laser power, gain, offset, frame size, zoom and scan speed) were maintained throughout the acquisition process. Images are representative of three independent experiments. Scale bars, 10 μ m.



Fig. 6 Co-staining studies utilizing super-resolution confocal microscopy images of hCB_2R -CHO cells. (A–F) Double-staining of hCB_2R -overexpressing CHO cells exposed for 15 min with 12 (A) or 16 (D) together with LysoTracker, a lysosome marker (B and E, green). Merged images are shown in the panels (C) and (F). The superimposition of the two stainings revealed that 12 and LysoTracker did not colocalize, while 16 and LysoTracker partially co-stained in several dot structures (white arrows in the magnified box). The images are representative of three independent experiments, and in each case, five fields were examined. Scale bars, 10 μ m.

staining produced by the matched TAMRA-probe pair **12** and **16**. Image acquisition at higher magnification and resolution was performed after 15 minutes of incubation with **12** and **16** and revealed that both probes yielded a similar localization pattern, yet with distinct labeling intensity. Indeed, for both probes the CB₂R staining was largely observed intracellularly, especially in the perinuclear membranes that are suggestive of the Golgi apparatus and the endoplasmic reticulum; however, appreciable fluorescence was also present on the cell membrane (Fig. 5A and B). Notably, these results corroborate and extend the findings previously reported by den Boon *et al.*¹⁸

It is noteworthy that the analysis of MFI of **12**-labeled CB₂R showed a plasma membrane labelling intensity of CB₂R approximately six-fold higher than that of **16**-labeled CB₂R (**12**labeled CB₂R = 2090 \pm 175 MFI; **16**-labeled CB₂R = 350 \pm 4 MFI; unpaired *t*-test, *t* = 19.94, df = 4, and *p*-value < 0.0001). Similarly, when examining the intracellular compartments, a marked difference in MFI values was observed (**12**-labeled CB₂R = 2575 \pm 100 MFI; **16**-labeled CB₂R = 340 \pm 13 MFI; unpaired *t*-test, *t* = 37.57, df = 4, and *p*-value < 0.0001). This marked difference in labelling intensity still persisted even when differences in binding affinities were compensated for by increasing the concentration of **16** *versus* **12** (ESI, Fig. S-7†). Moreover, comparable quantum yields of probes **12** and **16** in more lipophilic media (ESI, Table S-3†) ruled out potential interference with the observed differences in MFI.

Given that probes 12 and 16 are conjugated with the identical fluorophore and exhibit comparable physicochemical features, including their binding affinity for CB₂R, respectively (ESI, Table S-2[†]), the more pronounced signal detected with probe 12 relative to probe 16 indicates that under steady-state conditions in living cells, CB₂R is present predominantly in its active conformation. It is also plausible that the inactive form of the receptor is less accessible for interaction with probe 16 resulting in weaker labeling when compared to probe 12.42,43 We further examined the different distribution patterns of the two probes by co-staining with AlexaFluor488 LysoTracker. Fig. 6 indicates that probe 12 had partial overlap but no clear colocalization with the LysoTracker staining. In contrast, costaining with probe 16 exhibited a clear co-localization pattern. These results are in line with the observations made on human macrophages (vide supra), indicating a distinct intracellular distribution of CB₂R pools, which is influenced by the activation state and may have further implications on cellular signalling pathways. This is consistent with previous extensive studies showing that localization of G protein-coupled receptors (GPCRs) is not restricted to the plasma membrane, but rather is in dynamic exchange with localization in various intracellular compartments, including endosomes, the Golgi apparatus, the endoplasmic reticulum, the nucleus and the mitochondria.44 This dynamic localization is often influenced by the receptor's activation state, with both active and inactive



Fig. 7 Endocytosis blocking studies using super-resolution confocal microscopy in hCB₂R-CHO cells. Cells were stained for 10 min with **12** (A) or **16** (C). In order to minimize endocytosis, hCB₂R-CHO cells were pretreated for 30 min with 400 μ M sucrose and 5 μ g mL⁻¹ filipin and then incubated for 10 min either with **12** (B) or **16** (D). Association curve of **12** on the plasma membrane and internal membranes of hCB₂R-CHO cells without sucrose and filipin (E) and in the presence of both (F). Images are representative of three independent experiments. Scale bars, 10 μ m. (E) and (F) demonstrate internal and external fluorescence over time.

forms potentially residing within intracellular organelles. CB₂R, similar to CB₁R, undergoes constitutive endocytosis, resulting in significant intracellular accumulation.^{18,45-47} This distribution pattern has been confirmed in leukocytes using anti-CB₂R antibodies.⁴⁸ Functionally, CB₂R can be either active or inactive within these subcellular compartments.^{17,47} Our findings reveal that inactive CB₂R is presumably partially sequestered in lysosomes and in an as-yet uncharacterized vesicular compartment. We propose that this sequestration may serve as a regulatory mechanism, preventing CB₂R activation and subsequent signalling until the receptor is recycled to the plasma membrane or degraded in the lysosome.

To determine the degree of endocytosis-mediated receptor internalization influenced by the receptor functional state, we performed blocking experiments. Although the experimental setting did not allow for an entire block of endocytosis, a significant reduction was achieved by treating the cells with 0.4 M sucrose and 5 μ g mL⁻¹ filipin,²¹ which can indirectly abrogate multiple pathways of endocytosis.49 This treatment reduced the intracellular staining of 12 by 70% without affecting that of 16 (see Fig. 7). We recorded the fluorescence intensity of probe 12 by time-lapse imaging and observed signal saturation in both native and blocker-pre-treated cells, both for cytoplasm and internal membranes. In cells pretreated with the endocytotic blockers sucrose and filipin the plasma membrane signal reached the same intensity plateau as in untreated cells, but was negligible for the internal membranes during the entire course of the experiment. The observation that suppression of endocytosis affects the internalization of 12 but does not reduce that of 16 aligns with the behavior of agonists and inverse agonists. Unlike agonists, inverse agonists stabilize the inactive form of a GPCR, which does not promote endocytosis from the membrane.46,50,51 However, this finding also clearly demonstrates that both probes 12 and 16 are able to reach a significant intracellular concentration by passive permeation in almost the same concentration range. Further investigation into the molecular basis of the differences in localization and activity states might provide valuable insights into the interaction dynamics between CB₂R and its ligands, potentially informing the design of more effective probes or therapeutic agents targeting this receptor.

Conclusions

In this study, we have developed the first general CB_2R selective matched agonist or inverse agonist platform using validated drug-derived CB_2R ligands 1 and 2, respectively, as starting points. We designed fluorescent probes sharing highly similar drug-derived chemotypes but addressing either the active or inactive state of CB_2R . Due to our reverse-design approach using highly drug-like precursors, all labeled probes exhibited favorable physicochemical properties. The key for the receptor recognition element and fluorophore attachment was a common centrepiece that allowed straightforward linker attachment by amide coupling. The probes retained good binding affinity towards CB_2R and high selectivity against CB_1R upon conjugation of fluorescent dyes. When investigating the

functional responses in a cellular cAMP accumulation assay, the labeled probes were able to evoke a similar functional response as their unlabeled congeners. In particular, the labeled inverse agonist enabled us to address the resting state of CB₂R. This is of great benefit and importance for various control experiments in live cells which are now possible. With matching TAMRA probes in hand we performed time-lapse confocal microscopy imaging studies in various biological models, including hCB₂R-CHO cells, hCB₂R-HEK293 cells and human primary macrophages. In all settings, it was clearly demonstrated that hCB₂R has different localization and internalization patterns, depending on the functional state. Most notably, our probes allowed the assessment of localization and distribution of the active and inactive conformations of CB2R through superresolution confocal microscopy analysis. These studies indicate that within living cells, a considerable number of CB2Rs are located intracellularly and are in an active state. The differential content of the active and inactive states of CB₂R suggests a complex regulatory mechanism governing its activity and interactions with intracellular signalling pathways. These observations, enabled by our matched fluorescent agonist and inverse agonist probe pair, suggest significant implications for understanding the receptor's role in physiological and pathological processes. Furthermore, our approach demonstrates the potential for super-resolution imaging of our CB₂R probes in studying membrane receptors representing a powerful tool for future research in cellular biology and pharmacology.

Data availability

Primary data for probe synthesis, characterization, photophysical measurements, and fluorescence imaging are provided in the ESI.†

Author contributions

M. N. and U. G. conceived the research and acquired funding for this project. M. W.-K., A. O., L. M., Y. M., T. G., B. B., W. G., D. V., L. H. H., S. O., M. Ma., U. G., and M. N. designed the research approach. M. W.-K., A. O., L. M., L. S., J. B., X. L., S. R., Y. M., M. S., T. G., C. v. d. H., B. B., A. H., Y. K., W. G., D. S., J. P. v. K., J. Br., T. H., D. V., and S. O. performed experiments, analyzed the raw data, and worked on probe validation. M. W.-K., A. O., L. M., Y. M., S. O., M. Ma., U. G., and M. N. analyzed all data generated and wrote the manuscript. J. B., Y. M., W. G., J. Br., T. H., D. V., and L. H. Provided useful comments and feedback for the manuscript. All authors have given approval to the final version of the manuscript.

Conflicts of interest

The authors declare no competing financial interest.

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