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Development of new chemical tools to study the cannabinoid receptor type 2

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Chapter 2

Structural Basis of Selective Cannabinoid CB₂ Receptor Activation

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Structural Basis of Selective Cannabinoid CB₂ Receptor Activation

Introduction

Preparations of the plant *Cannabis sativa* have been used for centuries in the treatment of various diseases, including cancer and neuropathic pain.¹ The synthetic version of its psychoactive constituent, Δ^9 -tetrahydrocannabinol (THC, Figure 2.1), is in FDA approved drugs Marinol[®] or Syndros[®] (dronabinol). The extracted version of THC is one of the active constituents of oromucosal spray Sativex[®] (nabiximols). These drugs are primarily used for the treatment of chemotherapy-induced nausea, enhancement of appetite in cachexic AIDS-patients, and to alleviate the spasticity and pain associated with multiple sclerosis.²⁻⁶ However, THC-based therapies are associated with clinically undesired psychotropic and cardiovascular adverse effects and challenging pharmacokinetic properties due to their high lipophilicity that may limit their therapeutic efficacy.⁷⁻¹⁰

THC exerts its therapeutic effects mostly via the G protein-coupled receptors (GPCRs) cannabinoid CB₁ and CB₂ receptors (CB₁R and CB₂R), which have 68% sequence identity in their seven transmembrane (TM) domains.¹¹ Both receptors are activated by the endogenous signaling lipids anandamide (AEA) and 2-arachidonoylglycerol (2-AG) (Figure 2.1), the two main endocannabinoids. The CB₁R, which is the most abundantly expressed GPCR in the central nervous system (CNS) is responsible for the psychotropic side effects of THC.¹²⁻¹⁴ It plays a role in memory, learning, neurogenesis, neuronal migration, and synaptogenesis. Furthermore, its presence in many organ tissues belies more non-neurological functions.¹⁵ The CB₂R is mainly found on the cells of the immune system and is upregulated under pathophysiological conditions.^{16,17} Its activation in general is associated with anti-inflammatory responses in tissue injury of the liver, heart, kidney, colon, and brain as determined in various preclinical models.¹⁸⁻²² Based on preclinical studies it is thought that selective CB₂R agonists may retain and exceed certain therapeutic properties of THC without inducing psychotropic side effects.²³

Various academic and industrial groups have developed selective CB₂R ligands.²⁴ HU308 (Figure 2.1) was the first selective CB₂R agonist to be reported that displayed anti-inflammatory and analgesic properties in mouse models without inducing CNS-side effects.¹⁸ However, poor physico-chemical properties (*e.g.* low solubility, high lipophilicity) of HU308, which has a calculated logarithm of octanol-water partition coefficient (cLogP) of 8.0²⁵, and its analogs prevented the successful clinical translation of this class of cannabinoid-based drugs.

A next generation of CB₂R ligands was developed with improved drug-like properties. For instance, Olorinab[®] (APD371, Figure 2.1) is the most polar CB₂R agonist reported to date with a cLogP of -0.4.²⁶ A phase 2a small-scale safety and tolerability trial in 14 patients with chronic abdominal pain associated with Crohn's disease showed mild-to-moderate adverse events and an improvement in abdominal pain scores.²⁷ Pyridinylbenzylimidazolidine-2,4-dione derivatives were previously disclosed as selective CB₂R agonists and their affinity studied, along with target binding kinetics and potency as a function of their lipophilicity, which resulted in the discovery of the orally available and peripherally restricted selective CB₂R agonist LEI-101 (Figure 2.1).²⁸⁻³⁰ It is intriguing that the CB₂R binding pocket tolerates a wide array of ligands with very different scaffolds and hydrophobicity. For example, HU308 has a 2-billion-fold higher lipophilicity than APD371. Despite the tremendous progress in the field of CB₂R drug discovery, there is still a poor molecular understanding on how these CB₂R agonists selectively activate CB₂R over CB₁R.

Recently, three-dimensional structures of the CB₁R and CB₂R have been elucidated in both the active and inactive states by crystallography or cryo-electron microscopy (cryo-EM) and the binding modes of

diverse ligands and their activation mechanism were reported.^{31–35} Remarkably, those structures revealed that CB₁R and CB₂R possess a highly similar, lipophilic orthosteric agonist binding pocket, which makes it challenging to explain the selective activation of CB₂R. To date, no structural studies with selective CB₂R agonists have been reported that could aid in understanding the molecular basis of CB₂R selectivity.

Here, LEI-102 is introduced as a novel, potent and selective CB₂R agonist with excellent physico-chemical and biological properties. LEI-102 was used in conjunction with CB₂R selective agonists APD371 and HU308, and non-selective agonist CP-55,940 to investigate the activation mechanism of CB₂R. This study combined ligand-target binding kinetics, site-directed mutagenesis, and cryo-EM methods. It was found that CB₂R has a distinct activation mechanism compared to CB₁R. Additionally, the physico-chemical properties of the ligands was found to influence their entry pathway into the receptor. Highly lipophilic ligands, such as HU308 and the endocannabinoids, may reach the binding pocket through the membrane, whereas more polar ligands, such as LEI-102, APD371 and CP-55,940, enter the receptor via an alternative route. Furthermore, it was shown that the favorable physico-chemical properties of LEI-102 and CB₂R selectivity underscore its promising *in vivo* efficacy via oral administration in a chemotherapy-induced nephropathy model without inducing CNS-mediated side effects. Together, these studies enhance the current insights of how certain physico-chemical properties of ligands translate to *in vivo* activity and changes their engagement to GPCRs.

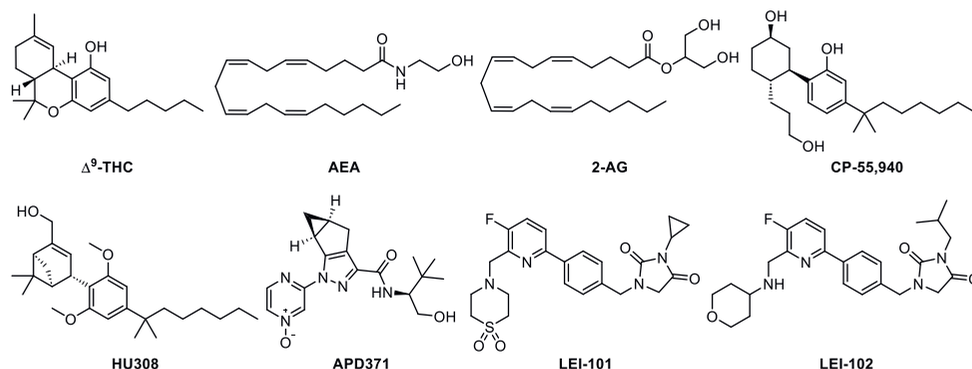
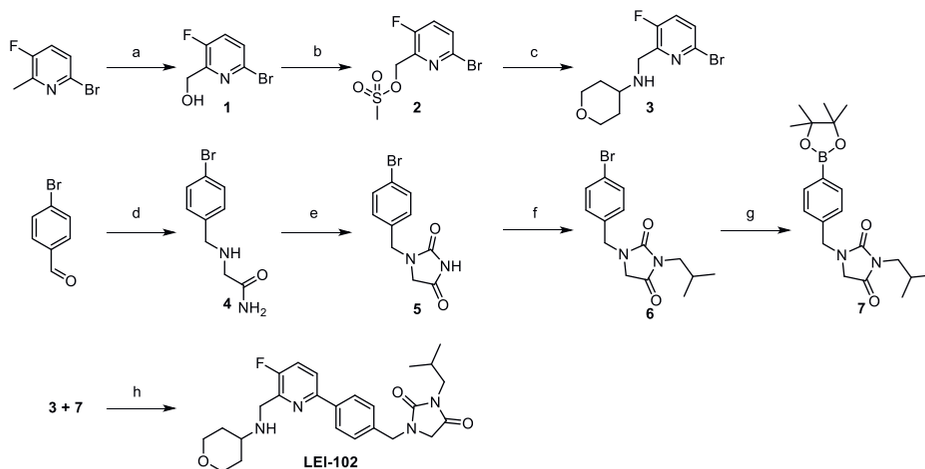


Figure 2.1 The chemical structures of the main constituent of *Cannabis sativa* Δ⁹-tetrahydrocannabinol (THC), and the two major endocannabinoids anandamide (AEA) and 2-arachidonoylglycerol (2-AG), as well as of non-selective CBR agonist CP-55,940 and CB₂R agonists HU308, APD371, LEI-101 and LEI-102.

Results

LEI-102 as a high affinity and potent CB₂R–selective agonist

To obtain a novel CB₂R agonist with beneficial physico-chemical properties, LEI-102, a pyridinylbenzylimidazolone-2,4-dione derivative, was designed and synthesized (Scheme 2.1). LEI-102 combined an isobutyl substituent on the imidazolone with an aminotetrahydropyran to replace the cyclopropyl and thiomorpholine 1,1-dioxide in LEI-101, respectively.³⁰



Scheme 2.1 Synthetic route of LEI-102. Reagents and conditions: a) Step 1: *m*-CPBA (1.8 eq), 0 °C-RT, DCM, 4 days; Step 2: TFAA (2.2 eq), 55 °C, 3 h; Step 3: K₂CO₃ (2.4 eq), THF:MeOH (20:1), 17 h, 35% (three steps); b) Et₃N (2.3 eq), MsCl (1.7 eq), THF, 0 °C-RT, 1 h, 75%; c) K₂CO₃ (2.2 eq), tetrahydro-2H-pyran-4-amine (1.3 eq), ACN, 50 °C, 3 h, 67%; d) Step 1: 2-aminoacetamide hydrochloride (1.0 eq), NaOH (1.1 eq), MeOH:H₂O (5:1), RT, 18 h; Step 2: NaBH₄ (2.1 eq), 18 h, 91% (two steps); e) CDI (2.1 eq), DMAP (2.1 eq), ACN, 60 °C, 70 h, 37%; f) K₂CO₃ (3.0 eq), 1-bromo-2-methylpropane (2.0 eq), DMF, RT, 20 h, 88%; g) KOAc (4.4 eq), bis(pinacolato)diboron (1.5 eq), Pd(dppf)Cl₂ (0.06 eq), DMF, 75 °C, 20 h; h) **3** (1.0 eq), **7** (1.5 eq), K₂CO₃ (6.0 eq), Pd(PPh₃)₄ (0.1 eq), toluene:EtOH (4:1), 75 °C, 18 h, 45% (two steps).

LEI-102 has a cLogP of 2.1 as calculated by ChemDraw 19.0 (Table 2.1). The inhibitory constant (pK_i), potency (pEC₅₀) and intrinsic activity (E_{max}) of LEI-102 were determined in [³H]-RO6957022 displacement assays on stably expressing CB₂R membranes and [³⁵S]GTPγS G protein activation assays using HEK293T membranes transiently expressing recombinant hCB₂R or hCB₁R, respectively (Table 2.2). APD371, HU308, CP-55,940 and the endocannabinoids AEA and 2-AG were also explored. LEI-102 had a high binding affinity for CB₂R (pK_i = 8.0 ± 0.1) and was more potent than the selective CB₂R agonists APD371 and HU308. LEI-102 did not bind CB₁R, thereby showing at least 1000-fold selectivity (Table 2.5). In G protein activation assays, LEI-102 activated the receptor as a partial agonist (E_{max} 76 ± 1 %) with a pEC₅₀ value of 6.9 ± 0.2 (Table 2.2).

Table 2.1 Physico-chemical properties of the investigated ligands.

Compound	CLogP	SLogP	TPSA (Å)	MW (g/mol)	Num RotatableBonds	NumHBD	NumHBA
Δ ⁹ -THC	7.24	5.74	29.5	314.5	4	1	2
2-AG	6.89	5.03	66.8	378.6	17	2	4
AEA	6.18	5.24	49.3	347.5	16	2	2
CP-55,940	5.82	5.66	60.7	376.6	10	3	3
HU308	8.00	6.63	38.7	414.6	10	1	3
APD371	-0.35	0.70	107.0	357.4	4	2	6
LEI-102	2.07	3.58	74.8	454.5	8	1	5
LEI-101 ³⁰	0.89	-	90.4	472.5	6	0	6

All values calculated using RDKit KNIME nodes version 4.5.0.v202207051536, apart from CLogP which was calculated using ChemDraw 19.0. For LEI-101 all values were calculated with Chemdraw 22.2.0.

Table 2.2 Functional Activity, Affinity and Kinetic Parameters of Synthetic Agonists and Endocannabinoids on hCB₂R.

Compound	pEC ₅₀ ^a	E _{max} (%) ^b	pK _i (K _i , nM) ^c	k _{on} (M ⁻¹ s ⁻¹) ^d	ET (s) ^e	k _{off} (s ⁻¹) ^f	RTD (min) ^g	K _D (nM) ^h
LEI-101 ^{30,i}	6.6 ± 0.2	65 ± 8	6.51 ± 0.1 (309.0)	(3.0 ± 1.1) × 10 ⁴	33.3 ± 8.91	(2.1 ± 0.5) × 10 ⁻³	8.8 ± 1.6	76 ± 10
LEI-102	6.9 ± 0.2	76 ± 1	8.0 ± 0.1 (9.7)	(6.3 ± 1.1) × 10 ⁴	16.0 ± 2.82	(1.0 ± 0.2) × 10 ⁻³	16 ± 3.3	16.5
APD371	7.9 ± 0.1	134 ± 12	7.5 ± 0.1 (35.3)	(2.5 ± 0.4) × 10 ⁴	40.1 ± 5.90	(3.7 ± 0.5) × 10 ⁻⁴	45 ± 6.6	14.9
HU308	7.1 ± 0.2	91 ± 8	7.0 ± 0.1 (92.4)	(7.0 ± 2.3) × 10 ³	143 ± 47.9	(2.3 ± 0.4) × 10 ⁻⁴	71 ± 11	33.7
CP-55,940	8.5 ± 0.3	98 ± 4	8.9 ± 0.1 (1.4)	(1.8 ± 0.4) × 10 ⁶	0.57 ± 0.13	(5.2 ± 0.9) × 10 ⁻⁴	32 ± 5.5	0.3
AEA	6.3 ± 0.2	60 ± 5	6.3 ± 0.1 (484.5)	(6.6 ± 0.5) × 10 ³	152 ± 10.4	(2.4 ± 0.1) × 10 ⁻³	6.8 ± 0.4	371.7
2-AG	5.9 ± 0.1	93 ± 22	7.0 ± 0.1 (97.3)	(5.3 ± 1.0) × 10 ⁴	18.8 ± 3.61	(2.3 ± 0.3) × 10 ⁻³	7.4 ± 1.0	42.7

^{a, b} Potency (pEC₅₀) and efficacy (E_{max}) values were obtained from [³⁵S]GTPγS assays on HEK293T membranes transiently expressing CB₂R WT. The percentage maximum effect (E_{max} in %) was calculated compared to CP-55,940. ^{c, d, f} Binding affinities (pK_i), association (k_{on}) and dissociation (k_{off}) rate constants were determined in [³H]-RO6957022 binding assays on CHO_K1_hCB₂R_bgal membranes at 10 °C. ^g Engagement time (ET) of the compound at 1 μM agonist was determined by ET = 1/(k_{on} · 1 × 10⁻⁶) and is expressed in seconds (s), whereas k_{on} is expressed in M⁻¹s⁻¹. ^h Residence time (RTD) was determined by RTD = 1/(60 · k_{off}) and is expressed in min, whereas k_{off} is expressed in s⁻¹. ^h Kinetic K_D values, defined by K_D = k_{off}/k_{on}. Values represent the mean ± SEM of at least three independent experiments performed in duplicate. LEI-101 data was obtained from literature and values were measured on CHO_K1_hCB₂R_bgal membranes.

Distinct target binding kinetic profiles of CB₂R agonists

For the quantification of the ligand-target binding kinetic parameters of the agonists, displacement and competition association assays with [³H]-RO6957022 were performed on CHO membranes stably over-expressing hCB₂R_bgal (Table 2.1). The equilibrium K_i and kinetic K_D values were well correlated, validating the competition association assay. The determined dissociation rate constants (*k_{off}*) of all agonists was converted into Residence Time Distribution (RTD). LEI-102 had an RTD of 16 min, which was around half that of APD371 (45 min) and CP-55,940 (32 min), whereas HU308 had the longest RTD at the receptor of 71 min (Table 2.1). Endocannabinoids 2-AG and AEA had the shortest RTD, both approximately 7 min. The association rate constants (*k_{on}*) varied greatly between the different agonists, ranking from fast to slow engagement CP-55,940 > LEI-102 > 2-AG > APD371 > HU308 = AEA. The calculated engagement time (ET) to CB₂ at 1 μM of each agonist further emphasized that CP-55,940 arrived at CB₂R within one second, whereas APD371, LEI-102 and 2-AG needed between 16 and 40 s to reach the CB₂R binding site. Interestingly, HU308 and AEA took 143 and 152 s to bind CB₂R, respectively. Due to the distinct target-binding kinetic profiles found for the four synthetic CB₂R agonists the choice was made to elucidate their binding poses in CB₂R with cryo-EM.

Overall similar structural comparison of CB₂R-G_i in complex with different agonists

To obtain the stable complex sample of CB₂R-G_i bound with LEI-102, APD371, HU308, or CP-55,940, a similar procedure was used as for the complex preparation previously described for AM12033-CB₂R-G_i (PDB: 6KPF). Single particle analysis of the cryo-EM samples yielded a normal global map for CB₂R-LEI-102-G_i-scFv16, CB₂R-APD371-G_i-scFv16, CB₂R-HU308-G_i-scFv16 and CB₂R-CP-55,940-G_i-scFv16, complex, at 2.9 Å, 3.0 Å, 3.0 Å and 2.9 Å, respectively (Figure 2.2). The ligand, receptor and G protein in the isolated complex were clearly visible in the cryo-EM maps (Figure 2.2). The overall structures of the four complexes were comparable, with root mean square deviation (RMSD) of the Cα atoms of the receptors are around 0.35 Å. The ligand binding interfaces of the four CB₂R and G_i complexes were similar to each other, and to those of the previous AM12033-CB₂R-G_i or WIN55212-2-CB₂R-G_i complex structures.

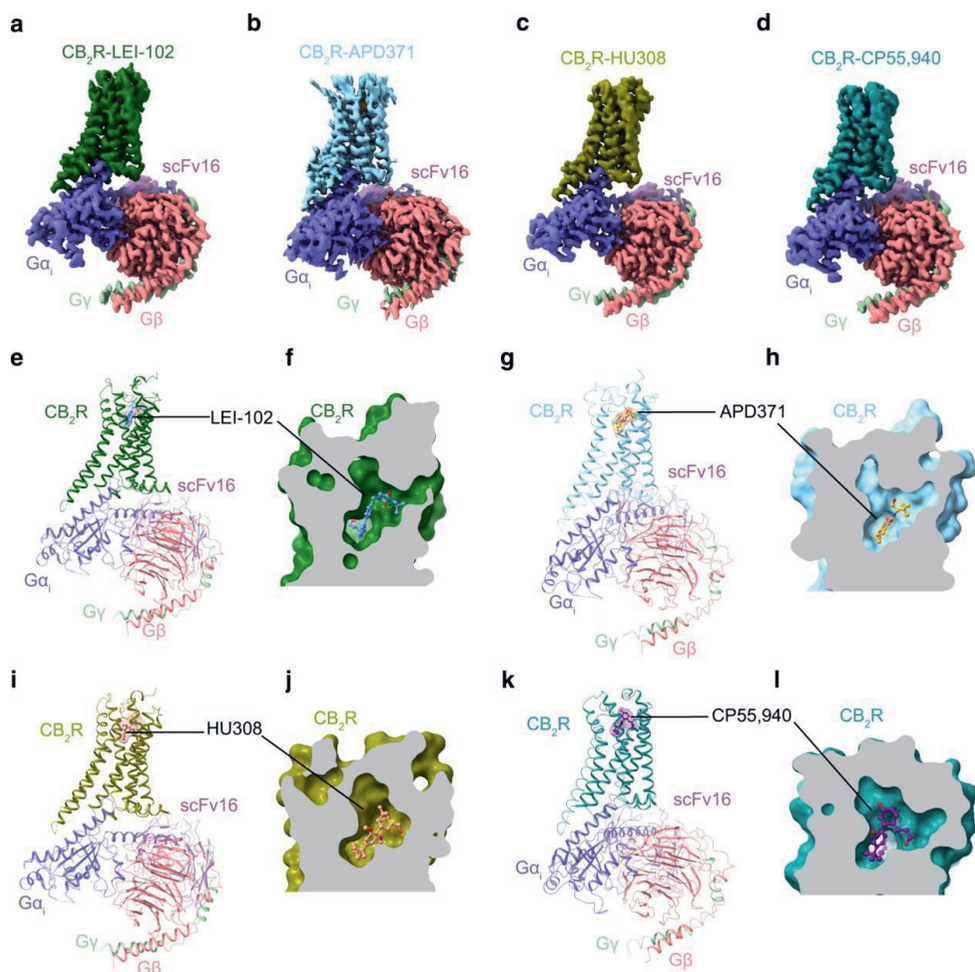


Figure 2.2 Cryo-EM structures of CB₂R-G_i complexes (A-D) Cryo-EM density maps of (A) LEI-102 (Dark green), (B) APD371 (Sky blue), (C) HU308 (Olive) and (D) CP-55,940 (Teal) bound CB₂R in complex with G_{αi} (Slate), G_β (Salmon), G_γ (Pale green), scFv16 (Violet purple). (E-L) Overall structures of CB₂R-G_i complexes and enlarged view of orthosteric pocket of (F) LEI-102, (H) APD371, (J) HU308 and (L) CP-55,940 using the same color codes as (A-D), with agonists shown as Cornflower blue (LEI-102), Orange (APD371), Dark salmon (HU308) and Purple (CP-55,940) sticks, respectively.

The binding mode of LEI-102 in CB₂R

A clear electron density in the orthosteric ligand binding pocket in the LEI-102-CB₂R-G_i complex resulted in the unambiguously defined binding pose of LEI-102. LEI-102 predominantly interacted with the residues in the binding pocket via hydrophobic interactions (Figure 2.3A). The isobutyl substituent of LEI-102 showed interactions with residues S90^{2.60} (Ballesteros-Weinstein numbering in superscript), F106^{3.25}, K109^{3.28}, and I110^{3.29} in CB₂R. The imidazolidine-2,4-dione forms π - π interaction with F94^{2.64} and showed further hydrophobic interactions with F106^{3.25} and P184^{ECL2}. The benzyl formed an aromatic interaction with F183^{ECL2}, and hydrophobic interactions with F87^{2.57} and S285^{7.39}. The phenyl ring in the core of LEI-102 formed a cation π interaction with F183^{ECL2} and T-shaped π - π interaction with F281^{7.35}. The pyridine had hydrophobic contacts with F117^{3.36} and W258^{6.48}. The aminotetrahydropyran sidechain protruded into the long channel and formed hydrophobic interactions with residues I110^{3.29},

T114^{3.33}, I186^{ECL2}, Y190^{5.39}, L191^{5.40}, W194^{5.43} and M265^{6.55}. Additionally, a hydrogen bond was formed with T114^{3.33} (Figure 2.3A).

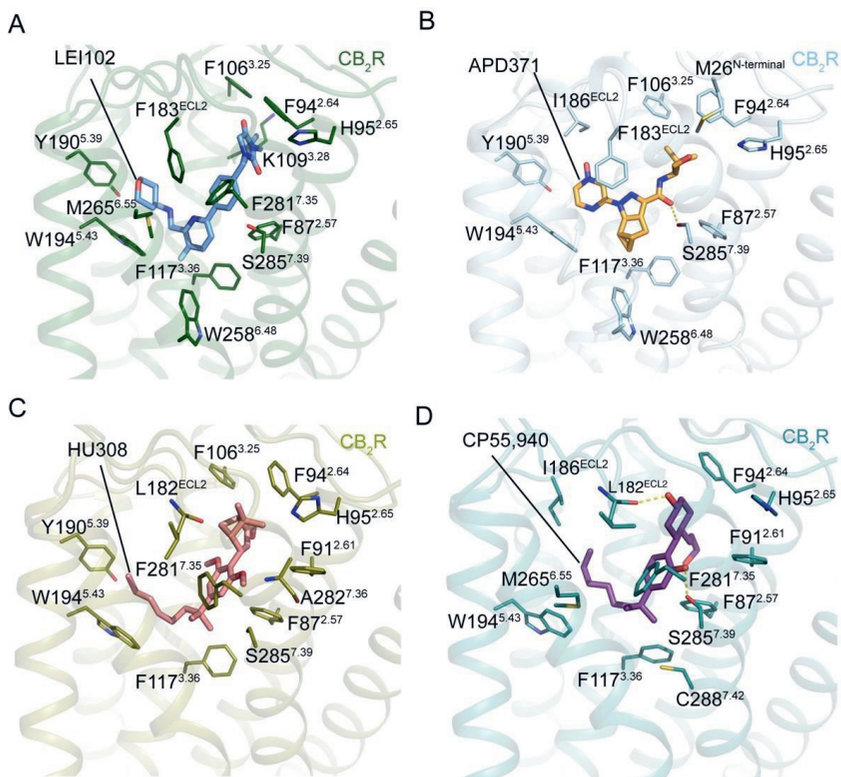


Figure 2.3 Key interactions between CB₂R structure and agonists (A-D) Key residues involved in (A) LEI-102 (cornflower blue) with CB₂R-G_i (dark green), (B) APD371 (orange) with CB₂R-G_i (sky blue), (C) HU308 (dark salmon) with CB₂R-G_i (olive) and (D) CP-55,940 (purple) with CB₂R-G_i (teal) binding in the complex structures. The amino acids involved interactions are showing sticks, hydrogen bonds are highlighted with yellow dashed lines.

The binding mode of APD371 in CB₂R

APD371 mainly formed hydrophobic and aromatic interactions with residues from ECL2, TM2, TM3, TM5, TM6 and TM7 (Figure 2.3B). The carbonyl group of APD371 formed a putative hydrogen bond with S285^{7.39} and a hydrophobic interaction with F87^{2.57}. The pyrazole and pyrazine cores of APD371 formed aromatic interaction with F183^{ECL2}. Furthermore, the pyrazine core formed hydrophobic contacts with T114^{3.33}, I186^{ECL2}, L191^{5.40} and W194^{5.43}. The (*S*)-1-hydroxy-3,3-dimethylbutyl head formed hydrophobic contacts with residues M26^{N-terminus}, S90^{2.60}, F94^{2.64}, F106^{3.25}, I110^{3.29} and V113^{3.32}. The cyclopropyl group formed hydrophobic contacts with F117^{3.36}, W194^{5.43}, W258^{6.48} and V261^{6.51}.

The binding mode of HU308 in CB₂R

The interactions between HU308 and CB₂R were hydrophobic, including residues from ECL2, TM2, TM3, TM5, TM6 and TM7 (Figure 2.3C). The phenyl of 2,6-dimethoxyphenyl core formed hydrophobic interactions with F87^{2.57}, F183^{ECL2} and S285^{7.39}, the C2-methoxy formed hydrophobic contacts with A282^{7.36} and S285^{7.39}, and the C6-methoxy formed hydrophobic contacts with I110^{3.29}, V113^{3.32} and T114^{3.33}, respectively. The dimethylheptyl chain of HU308 extended into the long channel and formed hydrophobic interactions with residues from ECL2 (F183^{ECL2}), TM3 (T114^{3.33}, F117^{3.36}), TM5 (W194^{5.43}). The 1,1-dimethyl formed hydrophobic interactions with residues F87^{2.57}, F117^{3.36}, F281^{7.35} and S285^{7.39}.

The bicyclic head of HU308 formed hydrophobic interactions with M26^{N-terminus}, F106^{3.25}, I110^{3.29}, S90^{2.60}, F94^{2.64}, P184^{ECL2} and the 2-methanol formed a hydrophobic interaction with F94^{2.64}.

The binding mode of CP-55,940 in CB₂R

CP-55,940 adopted an L-shape conformation in the orthosteric binding pocket (Figure 2.3D). The cyclohexanol group formed hydrophobic interactions with F94^{2.64}, L182^{ECL2}, F183^{ECL2}, and P184^{ECL2}. The hydroxyl group established a hydrogen bond with L182^{ECL2} and the hydroxypropyl formed hydrophobic contacts with F87^{2.57}, S90^{2.60}, F91^{2.61}, I110^{3.29}, and V113^{3.32}. The phenol core formed hydrophobic interactions with F87^{2.57}, F183^{ECL2}, F281^{7.35} and S285^{7.39}, and its hydroxyl additionally formed a hydrogen bond with S285^{7.39}. The dimethyl formed hydrophobic interactions with F183^{ECL2}, F281^{7.35}, M265^{6.55}, F87^{2.57}, F117^{3.36} and C288^{7.42}. The dimethylheptyl alkyl chain of CP-55,940 extended into the long channel and formed hydrophobic interactions with residues I110^{3.29}, F183^{ECL2}, I186^{ECL2}, W194^{5.43}, T114^{3.33} and F117^{3.36}.

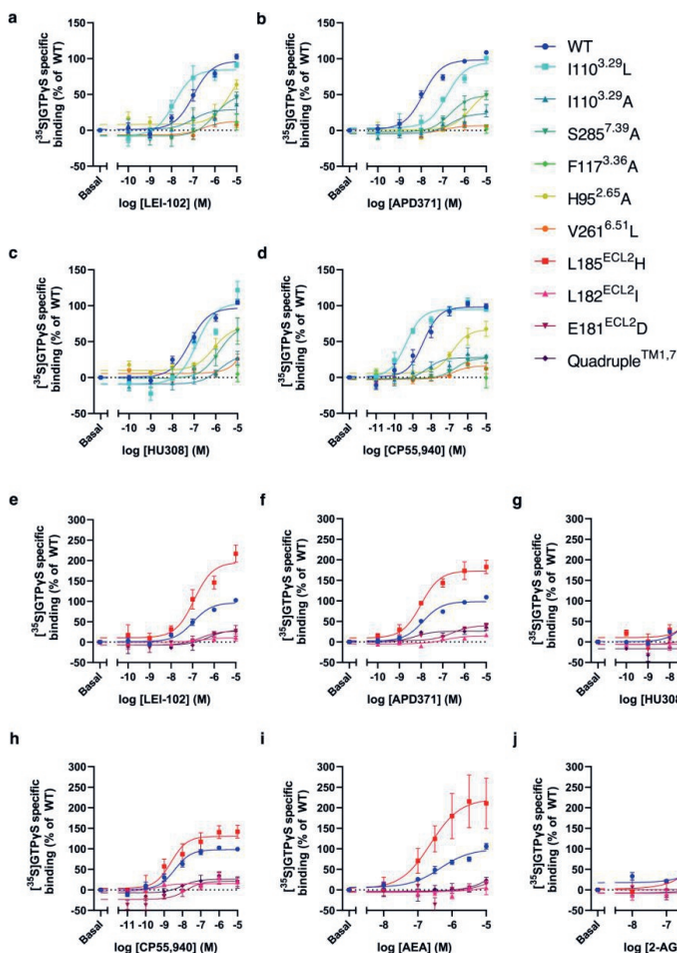


Figure 2.4 Characterization of G Protein activation of wild type (WT) and mutant CB₂R by synthetic agonists and endocannabinoids. (A-D) Dose-response curves for G protein activation of WT and mutants that are located in the CB₂R binding pocket by (A) LEI-102, (B) APD371, (C) HU308 and (D) CP-55,940. (E-J) Dose response curves for G protein activation of WT and mutants that are proposed to be involved in ligand entry of CB₂R via either the ECL2 or membrane access by (E) LEI-102, (F) APD371, (G) HU308, (H) CP-55,940, (I) AEA and (J) 2-AG. (A-J) The maximum activation level of WT CB₂R was set

to 100% while the basal levels were set to 0%. Data are presented as mean \pm SEM of at least three individual experiments performed in duplicate.

LEI-102 and APD371 require H95^{2.65} for G protein activation in CB₂R

To study the mechanism of CB₂R activation, five residues in the binding pocket were further characterized based on the complex structures (Figure 2.3). Six CB₂R mutants were created, *i.e.* four residues (S285^{7.39}, H95^{2.65}, I110^{3.29} and F117^{3.36}) were replaced by alanine, as these are conserved between CB₂R and CB₁R, and two others (I110^{3.29}, V261^{6.51}) were substituted by the CB₁R reciprocal residue leucine. All mutants were sufficiently expressed at the cell surface as determined with an ELISA (Figure 2.5, Table 2.3). To characterize the binding mechanisms of LEI-102, APD371, HU308 and CP-55,940, their responses were investigated by [³H]CP-55,940 displacement and [³⁵S]GTP γ S binding assays. Of note, in the [³H]CP-55,940 displacement assay, only the CB₂R-I110^{3.29}L mutant showed a sufficient binding window (data not shown). This prevented the affinity determination of the four agonists on other mutant receptors. Five mutant receptors, except CB₂R-F117^{3.36}A, were still active in the [³⁵S]GTP γ S functional assay for study of the receptor activation mechanism (Figure 2.4A-D, Table 2.4). All four synthetic agonists were unable to activate CB₂R-F117^{3.36}A, which indicated an important role of this residue in the activation of CB₂R.

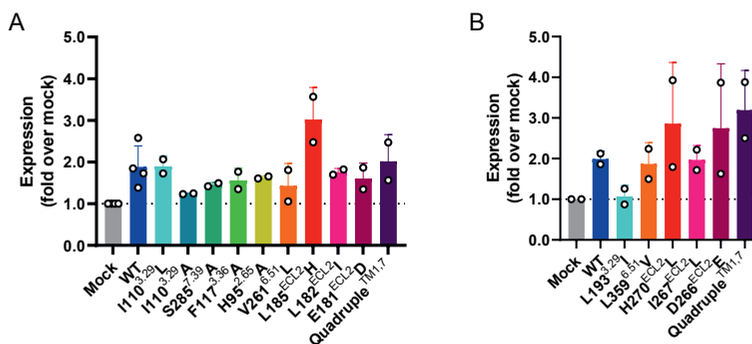


Figure 2.5 Cell Surface Receptor Expression. Receptor expression as determined by ELISA for (A) CB₂R and (B) CB₁R wild type (WT) and mutants. Data are expressed as mean \pm SD of at least two experiments performed in quintuplicate.

Table 2.3 Cell surface expression levels of cannabinoid receptor constructs in ELISA.

	CB ₂ R	Expression (fold over Mock)	CB ₁ R	Expression (fold over Mock)
Binding pocket	WT	1.9 \pm 0.5	WT	2.0 \pm 0.2
	I110 ^{3.29} L	1.9 \pm 0.2	L193 ^{3.29} I	1.1 \pm 0.3
	I110 ^{3.29} A	1.2 \pm 0.0	L193 ^{3.29} A	n.d.
	S285 ^{7.39} A	1.5 \pm 0.1	S383 ^{7.39} A	n.d.
	F117 ^{3.36} A	1.6 \pm 0.3	F200 ^{3.36} A	n.d.
	H95 ^{2.65} A	1.6 \pm 0.0	H178 ^{2.65} A	n.d.
	V261 ^{6.51} L	1.4 \pm 0.5	L359 ^{6.51} V	1.9 \pm 0.5
Ligand entry	L185 ^{ECL2} H	3.0 \pm 0.8	H270 ^{ECL2} L	2.9 \pm 1.5
	L182 ^{ECL2} I	1.8 \pm 0.1	I267 ^{ECL2} L	2.0 \pm 0.4
	E181 ^{ECL2} D	1.6 \pm 0.4	D266 ^{ECL2} E	2.7 \pm 1.6
	Quadruple ^{TM1,7}	2.0 \pm 0.6	Quadruple ^{TM1,7}	3.2 \pm 1.0

Mutations are shown in the numbering of the cannabinoid CB₂ (CB₂R) or CB₁ receptor (CB₁R) amino acid sequence as well as the Ballesteros and Weinstein GPCR numbering system. Data are presented as fold over mock (empty pcDNA3.1 vector) and are mean \pm SD of at least two individual experiments performed in quintuplicate. n.d. is not determined.

Table 2.4 The Affinity and Potency of the Agonists on the CB₂R mutants.

Construct	LEI-102			CP-55,940			HU308			APD371		
	pK _i	pEC ₅₀										
WT	7.5 ± 0.1	6.9 ± 0.2	9.2 ± 0.2	8.5 ± 0.3	7.8 ± 0.2	7.1 ± 0.2	8.0 ± 0.0	8.0 ± 0.0	7.9 ± 0.1	7.9 ± 0.1	8.0 ± 0.0	7.9 ± 0.1
I110 ^{3,29} L	7.9 ± 0.0	7.8 ± 0.1**	9.1 ± 0.1	9.4 ± 0.2	7.3 ± 0.1	6.7 ± 0.4	7.1 ± 0.0***	7.1 ± 0.0***	6.7 ± 0.3	6.7 ± 0.3	7.1 ± 0.0***	6.7 ± 0.3
I110 ^{3,29} A	n.d.	7.0 ± 0.6	n.d.	8.7 ± 0.4	n.d.	6.4 ± 0.5	n.d.	6.4 ± 0.5	6.6 ± 0.6	6.6 ± 0.6	n.d.	6.6 ± 0.6
S285 ^{7,39} A	n.d.	6.3 ± 0.3	n.d.	6.7 ± 0.1**	n.d.	5.9 ± 0.1***	n.d.	5.9 ± 0.1***	6.5 ± 0.3	6.5 ± 0.3	n.d.	6.5 ± 0.3
F117 ^{3,39} A	n.d.	n.d.										
H95 ^{2,65} A	n.d.	<5	n.d.	6.9 ± 0.2**	n.d.	6.6 ± 0.6	n.d.	6.6 ± 0.6	5.7 ± 0.2**	5.7 ± 0.2**	n.d.	5.7 ± 0.2**
V261 ^{6,51} L	n.d.	6.4 ± 0.0	n.d.	<5	n.d.	<5	n.d.	<5	6.9 ± 0.2*	6.9 ± 0.2*	n.d.	6.9 ± 0.2*
L185 ^{5,62} H	7.3 ± 0.1	6.9 ± 0.1	9.4 ± 0.1	8.9 ± 0.4	7.2 ± 0.0	6.8 ± 0.2	7.7 ± 0.1	6.8 ± 0.2	8.0 ± 0.1	8.0 ± 0.1	7.7 ± 0.1	8.0 ± 0.1
L182 ^{5,62} L	n.d.	7.4 ± 0.8	n.d.	8.8 ± 0.8	n.d.	<5	n.d.	<5	6.6 ± 0.4	6.6 ± 0.4	n.d.	6.6 ± 0.4
E181 ^{5,62} D	n.d.	6.9 ± 0.6	n.d.	7.9 ± 0.5	n.d.	7.0 ± 0.7	n.d.	7.0 ± 0.7	6.9 ± 0.6	6.9 ± 0.6	n.d.	6.9 ± 0.6
Quadruple ^{TM1,7}	6.8 ± 0.2	6.2 ± 0.2	9.2 ± 0.1	8.1 ± 0.2	7.1 ± 0.4	n.d.	6.7 ± 0.3	n.d.	8.4 ± 0.1	8.4 ± 0.1	6.7 ± 0.3	8.4 ± 0.1

Mutants that gave no dpm window could not have their potency determined (n.d. not determined). Mutations are shown in the numbering of the cannabinoid receptor 2 amino acid sequence as well as the Ballesteros and Weinstein GPCR numbering system. pEC₅₀ <5 was reported when the curve fit was not finished. Values are presented as the mean ± SEM of at least three independent experiments performed in duplicate. One-way Welch ANOVA with Dunnett's T3 posthoc test or Welch's t-test was used to analyze differences in pEC₅₀ and E_{max} values compared to WT (*p < 0.05, ** p < 0.01, *** p < 0.001).

Table 2.5 The Affinity and Potency of the Agonists on the CB₁R mutants.

Construct	LEI-102			CP-55,940			HU308			APD371		
	Displacement (%)	Potency (fold over basal)	pK _i	Potency (fold over basal)	Displacement (%)	Potency (fold over basal)						
WT	6 ± 7	1.0 ± 0.1	8.2 ± 0.0	1.2 ± 0.1	17 ± 7	0.8 ± 0.0	-19 ± 17	0.8 ± 0.0	1.0 ± 0.0	1.0 ± 0.0	1.0 ± 0.0	1.0 ± 0.0
L359 ^{6,51} V	41 ± 10	1.1 ± 0.1	8.3 ± 0.1	1.4 ± 0.2	25 ± 6	0.9 ± 0.1	18 ± 9	0.9 ± 0.1	1.1 ± 0.1	1.1 ± 0.1	1.1 ± 0.1	1.1 ± 0.1
H270 ^{6,52} L	46 ± 3	1.2 ± 0.0	8.3 ± 0.1	1.6 ± 0.1	47 ± 5	1.2 ± 0.1*	8 ± 3	1.2 ± 0.1*	1.1 ± 0.0	1.1 ± 0.0	1.1 ± 0.0	1.1 ± 0.0
I267 ^{6,52} L	-7 ± 19	1.0 ± 0.0	8.4 ± 0.1	1.0 ± 0.0	25 ± 3	0.8 ± 0.0	-24 ± 17	0.8 ± 0.0	0.9 ± 0.0	0.9 ± 0.0	0.9 ± 0.0	0.9 ± 0.0
D266 ^{6,52} E	34 ± 5	1.1 ± 0.1	8.4 ± 0.1	1.4 ± 0.1	38 ± 10	0.8 ± 0.1	12 ± 7	0.8 ± 0.1	0.9 ± 0.1	0.9 ± 0.1	0.9 ± 0.1	0.9 ± 0.1
Quadruple ^{TM1,7}	-13 ± 22	1.2 ± 0.1	8.0 ± 0.2	1.8 ± 0.1*	26 ± 10	0.9 ± 0.0	0 ± 25	0.9 ± 0.0	1.1 ± 0.0	1.1 ± 0.0	1.1 ± 0.0	1.1 ± 0.0

Affinity is expressed as pK_i for CP-55,940, but as displacement (receptor occupancy by agonist at 10 μM) for the CB₂R-selective agonists. The Potency is displayed as fold over basal (with basal the endogenous activity of the mutant). Mutations are shown in the numbering of the cannabinoid receptor 2 amino acid sequence as well as the Ballesteros and Weinstein GPCR numbering system. pEC₅₀ <5 was reported when the curve fit was not finished. Values are presented as the mean ± SEM of at least three independent experiments performed in duplicate. One-way Welch ANOVA with Dunnett's T3 posthoc test or Welch's t-test was used to analyze differences in pEC₅₀ and E_{max} values compared to WT (*p < 0.05, ** p < 0.01, *** p < 0.001).

The potency of LEI-102 was significantly increased at the CB₂R-I110^{3,29}L mutant to a pEC₅₀ value of 7.8 ± 0.1 in the G protein activation assay, while the binding affinity remained similar to wild type (WT) receptor (Figure 2.4A, Table 2.4). Three mutations CB₂R-I110^{3,29}A, CB₂R-S285^{7,39}A and CB₂R-V261^{6,51}L had no significant effect on the potency of LEI-102 in the functional assay. In contrast, the potency on mutant receptor CB₂R-H95^{2,65}A was significantly reduced for LEI-102. No gain in binding affinity for the swap mutant in CB₁R-L359^{6,51}V was found with LEI-102 (Table 2.5).

APD371 acted as a full CB₂R agonist with a pEC₅₀ value of 7.9 ± 0.1 and a higher maximal activation compared to that of CP-55,940 in the functional assay (Table 2.2). Mutant receptor CB₂R-I110^{3,29}L did not affect the G protein response of APD371 (Figure 2.4B, Table 2.4), while the binding affinity was significantly reduced to a pK_i of 7.1 ± 0.0 (Table 2.4). APD371 potency was not affected by mutant receptors CB₂R-I110^{3,29}A or CB₂R-S285^{7,39}A. The responses of APD371 for CB₂R-H95^{2,65}A and CB₂R-V261^{6,51}L were significantly impacted with 158-fold and 10-fold drop in potency, respectively (Table 2.4).

It is apparent that CB₂R-H95^{2,65} has a crucial role in G protein activation of CB₂R by LEI-102 and APD371. Additionally, LEI-102 activation was increased for the CB₂R-I110^{3,29}L mutant, while APD371 activation relied on CB₂R-V261^{6,51}.

An important role for S285^{7,39} and V261^{6,51} in CB₂R activation by HU308 and CP-55,940

The potency and affinity of HU308 on CB₂R were not affected by the CB₂R-I110^{3,29}L swap mutant (Figure 2.4C, Table 2.4). In addition, activation of mutant receptors CB₂R-I110^{3,29}A and CB₂R-H95^{2,65}A by HU308 was not affected with pEC₅₀ values of 6.4 ± 0.5 and 6.6 ± 0.6, respectively. The maximum activation level of mutant receptor CB₂R-S285^{7,39}A was unaffected compared to WT receptor, but a significant 15-fold loss in potency was observed. Lastly, CB₂R-V261^{6,51}L had a significant loss of potency, *i.e.* more than 120-fold lower (Figure 2.4C, Table 2.4).

Similar to HU308, the potency of CP-55,940 on CB₂R was not affected by the CB₂R-I110^{3,29} mutations compared to WT in the G protein activation assay, nor was its binding affinity for CB₂R-I110^{3,29}L (Figure 2.4D, Table 2.4). In response to CP-55,940, mutant receptors CB₂R-S285^{7,39}A and CB₂R-V261^{6,51}L were significantly affected with decreased pEC₅₀ values of 6.7 ± 0.1 and <5, respectively. Moreover, the potency of CP-55,940 was significantly affected on the CB₂R-H95^{2,65}A with a 40-fold decrease compared to WT receptor (Figure 2.4D, Table 2.4). No gain in potency or affinity was observed for the swap mutant CB₁R-L359^{6,51}V for either HU308 or CP-55,940 (Table 2.5).

Taken together, this showed that CB₂R-S285^{7,39} and CB₂R-V261^{6,51} were crucial for HU308 and CP-55,940 to activate the G protein at CB₂R, where CP-55,940 additionally required an interaction with CB₂R-H95^{2,65}.

Table 2.6 The Affinity and Potency of the Endocannabinoids on both CB₂R and CB₁R.

CB ₂ R	Construct	AEA		2-AG	
		pK _i	pEC ₅₀	pK _i	pEC ₅₀
	WT	6.2 ± 0.1	6.3 ± 0.2	5.8 ± 0.1	5.9 ± 0.1
	L185 ^{ECL2} H	6.3 ± 0.1	6.6 ± 0.0	6.4 ± 0.2	5.9 ± 0.2
	L182 ^{ECL2} I	n.d.	<5	n.d.	<5
	E181 ^{ECL2} D	n.d.	<5	n.d.	<5
	Quadruple ^{TM1,7}	5.6 ± 0.3	<5	<5	<5

CB ₁ R	Construct	pK _i	Potency (fold over basal)	pK _i	Potency (fold over basal)
	WT	6.2 ± 0.1	1.2 ± 0.1	5.4 ± 0.2	1.3 ± 0.1

H270^{ECL2L}	6.1 ± 0.0	1.4 ± 0.1	5.0 ± 0.4	1.5 ± 0.1
I267^{ECL2L}	6.2 ± 0.1	1.0 ± 0.1	5.9 ± 0.2	1.1 ± 0.0
D266^{ECL2E}	6.4 ± 0.1	1.3 ± 0.1	5.7 ± 0.1	1.5 ± 0.2
Quadruple^{TM1,7}	6.3 ± 0.0	1.6 ± 0.1	5.2 ± 0.0	1.8 ± 0.1*

pEC₅₀ <5 was reported when the curve fit was not finished. Values are presented as the mean ± SEM of at least three independent experiments performed in duplicate. One-way Welch ANOVA with Dunnett's T3 posthoc test or Welch's t-test was used to analyze differences in pEC₅₀ and E_{max} values compared to WT (*p < 0.05, ** p < 0.01). n.d. not determined.

HU308 and endocannabinoids gain access via membrane entry

The ligand-target binding kinetics (Table 2.2) revealed that the highly lipophilic HU308 and anandamide had a very slow on-rate compared to the other ligands. It has been postulated that ligands of lipid receptors may gain access to the binding pocket via a membrane channel. Both potential ligand entry pathways, *i.e.* either via ECL2 or via a membrane channel between TM1 and TM7, were examined. To this end, four additional mutant receptors were created. Three residues in the ECL2 of CB₂R, which were different from CB₁R, were mutated towards the reciprocal CB₁R residues, *i.e.* CB₂R-L185^{ECL2H}, CB₂R-L182^{ECL2I} and CB₂R-E181^{ECL2D}. In the fourth mutant receptor, four residues in TM1 and TM7 that align the potential membrane channel in CB₂R were mutated to the reciprocal CB₁R residues and combined as a quadruple mutant, *i.e.* CB₂R-K279^{7.33T}, CB₂R-K33^{1.32Q}, CB₂R-V36^{1.35I} and CB₂R-C40^{1.39S} (termed "CB₂R-Quadruple^{TM1,7}"). Next, all four synthetic agonists and the two endocannabinoids were tested on these four CB₂R mutant receptors in [³H]CP-55,940 and [³⁵S]GTPγS assays. Only CB₂R-L185^{ECL2H} and CB₂R-Quadruple^{TM1,7} were evaluated in the [³H]CP-55,940 displacement assays due to insufficient binding window for the other two mutant receptors (data not shown). The binding affinities of the agonists were not affected for mutant receptors CB₂R-L185^{ECL2H} and CB₂R-Quadruple^{TM1,7} (Table 2.6. Interestingly, the potencies of LEI-102, APD371 and CP-55,940 in the functional assay were not significantly affected for any of the mutant receptors, whereas HU308 and the endocannabinoids were less potent on CB₂R-L182^{ECL2I} (Figure 2.4E-J, Tables 2.4 and 2.6). Additionally, the endocannabinoids showed a decreased potency on CB₂R-L181^{ECL2D}, but not on CB₂R-L185^{ECL2H}. Of note, HU308 and both endocannabinoids completely lost their ability to activate CB₂R in the CB₂R-Quadruple^{TM1,7} mutant, suggesting that this may be an important access point to the receptor binding pocket for these agonists (Figure 2.4G, I-J).

Discussion

Currently, several crystal and cryo-EM CBR structures have been resolved in which non-selective agonists adopt a nearly identical binding position in the orthosteric pocket, regardless of the receptor.^{34–38} The results of the site-directed mutagenesis, ligand-target binding kinetics and cryo-EM studies were used to generate a better understanding of the binding and activation mechanism of CB₂R-selective agonists.

F117^{3.36} has an important role in CB₂R activation that does not mimic F200^{3.36} in CB₁R

The data revealed a crucial role for CB₂R-F117^{3.36} as replacement by alanine resulted in a complete loss of G protein activation by all tested agonists (Figure 2.4A-D, Table 2.4). It has been shown that the CB₁R counterpart F200^{3.36} plays an important regulatory role in activation as part of the "twin toggle switch" with CB₁R-W356^{6.48,39}. In contrast, CB₂R-W258^{6.48} has been described to be solely responsible for activation as a toggle switch without the help of CB₂R-F117^{3.36} in structural studies, since the conformation of CB₂R-F117^{3.36} in agonist-bound structures is comparable to the conformation in the antagonist-bound CB₂R structure as well as the CB₁R agonist-bound structures.^{33,36} The mutational data supports this hypothesis, as the constitutive activity pattern observed by McAllister *et al.* for the reciprocal CB₁R-F200^{3.36A}, is not observed here. This excludes CB₂R-F117^{3.36} from having a suppressive

function.³⁹ Together, this data provides strong support for a different, but important, role for F117^{3,36} in CB₂R activation.

Potential polar network of H95^{2,65} and S285^{7,39} in CB₂R

In CB₁R, water-mediated interactions between CB₁R-H178^{2,65}, CB₁R-S383^{7,39} and bound ligands have previously been shown with *in silico* modelling.^{40,41} The importance of CB₁R-S383^{7,39} for classical synthetic cannabinoids such as AM11542, AM841 and CP-55,940 was further emphasized in CB₁R-S383^{7,39}A mutants.³⁶ This is in line with the observation that removal or methylation of the phenolic OH on classical cannabinoids, such as in L-759656, JWH-133 and HU308, always affords selectivity over CB₁R.^{18,42} Non-classical agonists, such as WIN55,212-2, do not form a hydrogen bond with CB₁R-S383^{7,39} and consequently are not affected by an alanine mutation.⁴³ This translates to the observation that CP-55,940 and HU308 are more affected by the CB₂R-S285^{7,39}A mutation than LEI-102 and APD371 (Figure 2.4A-D, Table 2.4). The decrease in activation is at least 30-fold smaller for CB₂R than CB₁R.³⁶ The elucidated cryo-EM structures of the four agonists chosen in this study did not show direct interactions with CB₂R-H95^{2,65}, though its role in stabilizing the surrounding residues cannot be ruled out. The large effect seen on G protein activation of CB₂R-H95^{2,65}A by LEI-102, APD371 and CP-55,940 (Figure 2.4A-D, Table 2.4) must therefore stem from an indirect interaction, supporting the polar network hypothesis between CB₂R-H95^{2,65} and CB₂R-S285^{7,39} in CB₂R.

V261^{6,51} as a potential selectivity hotspot for CB₂R over CB₁R

Residues at position 6.51 have previously been described to be involved in the binding sites of μ , δ , and κ opioid receptors, the dopamine D2 receptor and adenosine receptors, and could play a role in ligand binding selectivity between different subtypes.⁴⁴⁻⁴⁶ The results show a reduction of G protein activation by APD371, HU308, and CP-55,940 with introduction of the bulkier CB₁R leucine in CB₂R-V261^{6,51}L. LEI-102 could still be accommodated in the binding pocket and was not affected (Figure 2.4A-D, Table 2.4). Additionally, in the swap mutant CB₁R-L359^{5,61}V partial recovery of [³H]CP-55,940 replacement was found for CB₂R-selective agonists LEI-102, HU308 and APD371, albeit not significant (Table 2.5). This supports the role of this residue in selectivity of agonists in CB₂R.

The lowly-conserved ECL2 is a large effector of GPCR selectivity

The ECL2 has frequently been implicated to be important for GPCR activation and some GPCRs even use their ECL2 as a ligand to auto-activation.⁴⁷ There are distinct differences between the conformations of ECL2 in CB₁R and CB₂R. In antagonist-bound CB₁R crystal structures, the ECL2 dips into the binding pocket, interacting with the ligand and inducing the inactive conformation.^{31,32} The inactive state of CB₂R, however, does not expand like CB₁R and instead the ECL2 acts more as a lid on the binding pocket in active and inactive CB₂R, akin to active CB₁R.³³ A key distinction seen in the CB₁R crystal structures with AM6538 and Taranabant, is the ionic lock formed by CB₁R-E100^{N-terminus} (CB₂R-L17) and CB₁R-H270^{ECL2} (CB₂R-L185).^{31,32} The results showed improved binding of [³H]CP-55,940 for LEI-102 and HU308 with the CB₁R-H270^{ECL2}L mutation, while the non-selective agonists showed no change (Table 2.5). Through the loss of this ionic lock, selectivity over CB₁R is partially lost, showing that expulsion of ECL2 upon ligand entry may play an important role in selectivity.

An alternative entry pathway for lipophilic agonists via a membrane channel

In recent years, computational studies have suggested that lipophilic ligands for various GPCRs, such as the opsin receptor, sphingosine-1-phosphate receptor 1 (S1P₁) and cannabinoid receptors, might gain access to the binding pocket through lateral diffusion via a membrane channel between TM1 and TM7.^{32,40,48-51} The membrane entry pathway was evaluated with a CB₂R quadruple mutant (K33^{1,32}Q, V36^{1,35}I, C40^{1,39}S and K279^{7,33}T), which showed a significant loss in potency and a corresponding decrease in affinity, albeit not significant, for HU308 and the endocannabinoids (Figure 2.4E-J, Table 2.4

and 2.6). These compounds are more lipophilic than LEI-102 and APD371, making them more suitable to traverse the membrane to enter between TM1 and TM7. Notably, HU308 and anandamide also showed a substantially longer ET in the kinetic assays compared to the other agonists (Table 2.2). This might suggest a possible relationship between slow association and membrane channel entry at the CB₂R. Likewise, for a peptide GPCR a trend in reduced association rate was found with increasing lipophilicity.⁵² Alternatively, this is in contrast with the mechanism at the α 2-adrenoceptor at which lipophilic compounds had a faster association rate.⁵³ This shows the diversity in drug-target binding kinetics as receptor-specific properties and thus the importance of investigating these mechanisms for individual receptors.⁵⁴

The discovery of a membrane access channel for endocannabinoids on the CB₂R is also intriguing from a physiological perspective. Endocannabinoids are produced on demand and act as autocrine or paracrine effectors in the immune system regulating the migration of CB₂R-expressing immune cells.¹⁷ The results suggest that endocannabinoids first have to travel through the plasma membrane via lateral diffusion to reach the receptor. This may suggest that the trafficking and cellular uptake of endocannabinoids could be regulated through extracellular or intracellular vesicles that merge with the plasma membrane. Regardless of the exact mechanism of endocannabinoid trafficking, the results provide experimental evidence of a membrane channel located between TM1 and TM7 in CB₂R that is used by the endocannabinoids to enter the receptor.

Conclusion

In silico and mutational studies on LEI-102 and five other CBR agonists have shown that physicochemical properties determine both pharmacokinetic properties of the ligand and their manner of engagement with the target. The ECL2 and CB₂R-V261^{5,61} are hotspots for inducing selectivity over CB₁R, and CB₂R-F117^{3,36} is vital for receptor activation. Additionally lipophilic ligands including the endocannabinoids may enter the binding pocket through a membrane entrance between TM1 and TM7. Altogether, these discovered molecular mechanisms for selective receptor engagement and activation can have implication for drug design and lipid signaling at GPCRs in general.

Experimental Section

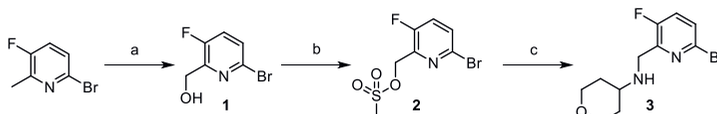
Chemistry

General Remarks

All reagents and solvents were purchased from commercial sources and were of analytical grade (Sigma-Aldrich, BroadPharm®). Reagents and solvents were not further purified before use. All moisture sensitive reactions were performed under inert atmosphere. Solvents were dried using 4 Å molecular sieves prior to use when anhydrous conditions were required. Water used in reactions was always demineralized. Analytical Thin-layer Chromatography (TLC) was routinely performed to monitor the progression of a reaction and was conducted on Merck Silica gel 60 F254 plates. Reaction compounds on the TLC plates were visualized by UV irradiation (λ 254) and/or spraying with potassium permanganate solution (K₂CO₃ (40 g), KMnO₄ (6 g), and H₂O (600 mL)), ninhydrin solution (ninhydrin (1.5 g), n-butanol (100 mL) and acetic acid (3.0 mL)) or molybdenum solution ((NH₄)₆Mo₇O₂₄·4H₂O (25 g/L) and (NH₄)₄Ce(SO₄)₄·2H₂O (10 g/L) in sulfuric acid (10%)) followed by heating as appropriate. Purification by flash column chromatography was performed using Screening Devices B.V. silica gel 60 (40-63 μ m, pore diameter of 60 Å). Solutions were concentrated using a Heidolph laborata W8 4000 efficient rotary evaporator with a Laboport vacuum pump. Analytical purity was determined with Liquid Chromatography-Mass Spectrometry (LC-MS) using a Finnigan LCQ Advantage MAX apparatus with electrospray ionization (ESI), equipped with a Phenomenex Gemini 3 μ m NX-C18 110Å column (50x4.6mm), measuring absorbance at 254 nm using a Waters 2998 PDA UV detector and the m/z ratio by using an Acquity Single Quad (Q1) detector. Injection was with the Finnigan Surveyor Autosampler

Plus and pumped through the column with the Finnigan Surveyor LC pump plus to be analyzed with the Finnigan Surveyor PDA plus detector. Samples were analyzed using eluent gradient 10% → 90% ACN in MilliQ water (+ 0.1% TFA (v/v)). For purification by mass guided preparative High-Performance Liquid Chromatography (Prep-HPLC) was performed on a Waters AutoPurification HPLC/MS apparatus with a Gemini prep column 5 μm 18C 110 Å (150x21.2mm), Waters 2767 Sample manager, Waters 2545 Binary gradient module, Waters SFO System fluidics organizer, Waters 515 HPLC pump M, Waters 515 HPLC pump L attached to a Waters SQ detector Acquity Ultra performance LC. A five column volume purification protocol was applied with the eluents A: 0.2% aq. TFA, B: ACN, flow 25 mL/min, with a minimum start gradients of 0% to maximum end gradient of 100% of B. ¹H, ¹³C, ¹H-COSY and HSQC Nuclear Magnetic Resonance (NMR) spectra were recorded on a Bruker AV 300 (300/75 MHz), AV 400 (400/100 MHz) or AV 500 (500/125 MHz) spectrometer at ambient temperature using CDCl₃ as solvent. Chemical shifts (δ) are referenced in parts per million (ppm) with tetramethylsilane (TMS) or CDCl₃ resonance as the internal standard peak (CDCl₃/TMS, δ 0.00 for ¹H (TMS), δ 77.16 for ¹³C (CDCl₃)). Multiplicity is reported as bs = broad singlet, s = singlet, d = doublet, bd = broad doublet, dd = doublet of doublet, t = triplet, q = quartet, p = quintet, m = multiplet. Coupling-constants (*J*) are reported in Hertz (Hz).

Synthesis of LEI-102



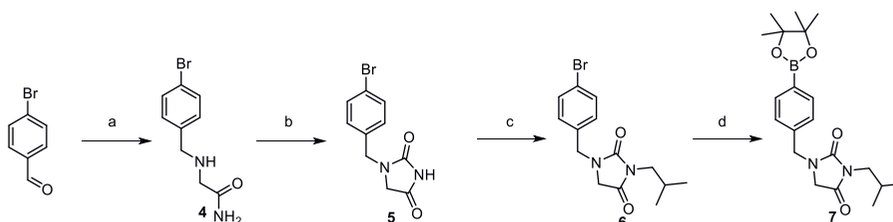
Scheme 2.2 Reagents and conditions: a) Step 1: *m*-CPBA (1.8 eq), 0 °C-RT, DCM, 4 days; Step 2: TFAA (2.2 eq), 55 °C, 3 h; Step 3: K₂CO₃ (2.4 eq), THF:MeOH (20:1), 17 h, 35% (three steps); b) Et₃N (2.3 eq), MsCl (1.7 eq), THF, 0 °C-RT, 1 h, 75%; c) K₂CO₃ (2.2 eq), tetrahydro-2H-pyridin-4-amine (1.3 eq), ACN, 50 °C, 3 h, 67%.

(6-Bromo-3-fluoropyridin-2-yl)methanol (1): To a stirred and cooled (0 °C) mixture under inert atmosphere of 6-bromo-3-fluoro-2-methylpyridine (10.7 g, 56.3 mmol, 1 eq) in DCM (370 mL) was added portion-wise *m*-CPBA (23.6 g, 70-75%, 100 mmol, 1.8 eq). The reaction mixture was stirred at room temperature (RT) for 4 days. Sat. NaHCO₃ (aq) and sat. Na₂S₂O₃ (aq) was added (1:1, v/v) and the layers were separated. The aqueous layer was extracted thrice with DCM. The combined organic layer was dried (MgSO₄), filtered, and concentrated under reduced pressure. To the residue was added TFAA (17 mL, 122 mmol, 2.2 eq) at 0 °C. After 15 minutes the temperature was increased to 55 °C for 3 h. The mixture was concentrated under reduced pressure, redissolved in DCM and sat. Na₂CO₃ (aq) was added. The layers were separated and the organic layer was washed with sat. NaHCO₃ (aq). The solvent was evaporated and the crude was dissolved in THF:MeOH (20:1, v/v) and K₂CO₃ (18.2 g, 132 mmol, 2.3 eq) was added. After 17 h H₂O was added and the layers were separated. The aqueous layer was extracted thrice with EtOAc. The combined organic layer was dried (MgSO₄), filtered, and the solvent evaporated under reduced pressure. The crude product was purified with flash column chromatography (10-20% EtOAc in pentane) to yield a white solid (5.79 g, 19.7 mmol, 35%). ¹H-NMR (500 MHz, CDCl₃) δ 7.42 (ddt, *J* = 8.5, 3.5, 0.7 Hz, 1H), 7.29 (t, *J* = 8.5 Hz, 1H), 4.80 (d, *J* = 3.3 Hz, 2H). ¹³C-NMR (126 MHz, CDCl₃) δ 156.10 (d, *J* = 256.2 Hz), 148.74 (d, *J* = 19.1 Hz), 135.01 (d, *J* = 2.9 Hz), 128.17 (d, *J* = 4.2 Hz), 126.09 (d, *J* = 19.8 Hz), 59.07.

(6-Bromo-3-fluoropyridin-2-yl)methyl methanesulfonate (2): To a cooled (0 °C) mixture of **1** (1.6 g, 7.8 mmol, 1 eq) and Et₃N (2.5 mL, 17.9 mmol, 2.3 eq) in dry THF (40 mL) was added dropwise MsCl (1.0 mL, 12.9 mmol, 1.7 eq). After stirring at RT for 1 h the solution was concentrated under reduced pressure. DCM and H₂O were added and the layers were separated. The aqueous layer was extracted thrice with DCM. The combined organic layer was washed with brine, dried (MgSO₄), filtered, and the

solvent evaporated under reduced pressure to yield a yellow solid (1.65 g, 5.8 mmol, 75%). ¹H-NMR (500 MHz, CDCl₃) δ 7.52 (dd, *J* = 8.6, 3.5 Hz, 1H), 7.37 (t, *J* = 8.5 Hz, 1H), 5.33 (d, *J* = 2.1 Hz, 2H), 3.13 (s, 3H). ¹³C-NMR (126 MHz, CDCl₃) δ 157.82 (d, *J* = 261.3 Hz), 142.15 (d, *J* = 16.0 Hz), 130.74 (d, *J* = 4.4 Hz), 127.06 (d, *J* = 20.4 Hz), 65.50 (d, *J* = 1.6 Hz), 38.39.

***N*-(6-Bromo-3-fluoropyridin-2-yl)methyltetrahydro-2*H*-pyran-4-amine (3):** A stirred suspension of **2** (1.49 g, 5.3 mmol, 1 eq), K₂CO₃ (1.6 g, 11.6 mmol, 2.2 eq) and tetrahydro-2*H*-pyran-4-amine (0.66 mL, 6.7 mmol, 1.3 eq) in acetonitrile (25 mL) was heated (50 °C) for 6 h, then stirred an additional 3 days at RT. After dilution with DCM and H₂O the layers were separated. The aqueous layer was extracted thrice with DCM. The combined organic layer was dried (MgSO₄), filtered, and the solution evaporated under reduced pressure. The crude product was purified with flash column chromatography (20-100% EtOAc in pentane) to yield a yellow oil (1.01 g, 3.5 mmol, 67%). ¹H-NMR (300 MHz, CDCl₃) δ 7.40 (dd, *J* = 8.6, 3.6 Hz, 1H), 7.35 – 7.26 (m, 1H), 4.08 – 3.95 (m, 4H), 3.42 (td, *J* = 11.6, 2.2 Hz, 2H), 2.74 (tt, *J* = 10.5, 4.1 Hz, 1H), 1.89 (ddd, *J* = 12.7, 4.5, 2.3 Hz, 2H), 1.52 (dtd, *J* = 13.1, 11.0, 4.5 Hz, 2H). ¹³C-NMR (75 MHz, CDCl₃) δ 157.12 (d, *J* = 255.9 Hz), 149.21 (d, *J* = 17.0 Hz), 127.83 (d, *J* = 4.2 Hz), 125.97 (d, *J* = 21.2 Hz), 66.76, 53.64, 44.90, 33.59.



Scheme 2.3 Reagents and Conditions: a) Step 1: 2-aminoacetamide hydrochloride (1.0 eq), NaOH (1.1 eq), MeOH:H₂O (5:1), RT, 18 h; Step 2: NaBH₄ (2.1 eq), 18 h, 91% (two steps); b) CDI (2.1 eq), DMAP (2.1 eq), ACN, 60 °C, 70 h, 37%; c) K₂CO₃ (3.0 eq), 1-bromo-2-methylpropane (2.0 eq), DMF, RT, 20 h, 88%; d) KOAc (4.4 eq), bis(pinacolato)diboron (1.5 eq), Pd(dppf)Cl₂ (0.06 eq), DMF, 75 °C, 20 h.

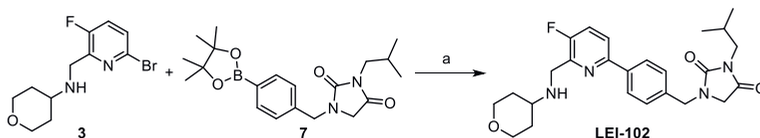
2-((4-Bromobenzyl)amino)acetamide (4): To a stirred mixture of 4-bromobenzaldehyde (9.2 g (49.7 mmol, 1.1 eq) and 2-aminoacetamide hydrochloride (5.06 g, 45.8 mmol, 1.0 eq) in MeOH:H₂O (170 mL, 5:1, v/v) was added NaOH (2.06 g, 51.5 mmol, 1.1 eq). After stirring at RT overnight, NaBH₄ (3.6 g, 95.2 mmol, 2.1 eq) was added and the solution was stirred overnight at RT. The solution was acidified to pH 3 with 2 M HCl, then neutralized with sat. NaHCO₃ (aq). Methanol was evaporated under reduced pressure and the resulting slurry was filtered to yield a white solid (11.0 g, 45.2 mmol, 91%). ¹H-NMR (300 MHz, MeOD) δ 7.69 – 7.59 (m, 2H), 7.47 – 7.38 (m, 2H), 4.22 (s, 2H), 3.81 (s, 2H).

1-(4-Bromobenzyl)imidazolidine-2,4-dione (5): To stirred suspension of **4** (10.0 g, 40.1 mmol, 1.0 eq) in acetonitrile (300 mL) were added CDI (13.86 g, 85.5 mmol, 2.1 eq) and DMAP (10.2 g, 83.5 mmol, 2.1 eq). The mixture was heated (60 °C) under inert atmosphere for 70 h. 1 M HCl (aq, 250 mL) was added and the aqueous layer extracted thrice with EtOAc. The combined organic layer was washed with H₂O and brine, dried (MgSO₄), filtered, and the solvent evaporated under reduced pressure. The crude product was purified with flash column chromatography with dry loading over Celite (5-10% acetone in DCM) to yield a yellow solid (3.95 g, 14.7 mmol, 37%). ¹H-NMR (300 MHz, CDCl₃) δ 7.83 (bs, 1H), 7.56 – 7.45 (m, 2H), 7.20 – 7.10 (m, 2H), 4.49 (s, 2H), 3.79 (s, 2H). ¹³C-NMR (75 MHz, CDCl₃) δ 132.41, 129.95, 77.58, 77.16, 76.74, 50.36, 46.01.

1-(4-Bromobenzyl)-3-isobutylimidazolidine-2,4-dione (6): To a stirred solution of **5** (2.00 g, 7.4 mmol, 1.0 eq) in anhydrous DMF (18 mL) were subsequently added K₂CO₃ (3.08 g, 22.3 mmol, 3.0 eq) and 1-bromo-2-methylpropane (1.62 mL, 14.9 mmol, 2.0 eq). After stirring at RT for 20 h, the mixture was

filtered and the filtrate diluted with diethyl ether and washed thrice with H₂O (3 x 50 mL). The combined organic layer was washed with brine, dried (MgSO₄), filtered, and concentrated under reduced pressure. The crude product was purified with flash column chromatography (10-40% EtOAc in pentane) to yield a white solid (2.12 g, 6.52 mmol, 88%). ¹H-NMR (300 MHz, CDCl₃) δ 7.47 (d, *J* = 8.3 Hz, 2H), 7.14 (d, *J* = 8.3 Hz, 2H), 4.52 (s, 2H), 3.74 (s, 2H), 3.33 (d, *J* = 7.4 Hz, 2H), 2.15 – 2.04 (m, 1H), 0.91 (d, *J* = 6.8 Hz, 6H). ¹³C-NMR (75 MHz, CDCl₃) δ 169.57, 156.78, 134.41, 131.79, 129.48, 121.77, 60.01, 48.61, 45.98, 45.71, 28.57, 19.70. LCMS (LCQ Fleet, 10-90): *t_r* = 7.00 min, *m/z*: 325.17 [M+H]⁺, 327.08 [M+H]⁺ (Br).

3-Isobutyl-1-(4-(4,4,5,5-tetramethyl-1,3,2-dioxaborolan-2-yl)benzyl)imidazolidine-2,4-dione (7): A mixture of **6** (0.50 g, 1.54 mmol, 1 eq), KOAc (0.66 g, 6.76 mmol, 4.4 eq) and bis(pinacolato)diboron (0.59 g, 2.31 mmol, 1.5 eq) in DMF (10 mL) was sonicated for 15 min under argon flow. Subsequently, Pd(dppf)Cl₂ (0.07 g, 0.09 mmol, 0.06 eq) was added and the mixture was heated (75 °C) for 20 h. The mixture was cooled to RT, diluted with EtOAc (100 mL) and H₂O (10 mL) and the layers were separated. The aqueous layer was extracted thrice with EtOAc (3 x 20 mL). The combined organic layer was extracted with sat. NaHCO₃ (aq), H₂O and brine, dried (MgSO₄), filtered, and concentrated under reduced pressure. The crude product was co-evaporated with CHCl₃ and used in the next step without further purification.



Scheme 2.4 Reagents and Conditions: a) **3** (1.0 eq), **7** (1.5 eq), K₂CO₃ (6.0 eq), Pd(PPh₃)₄ (0.1 eq), toluene:EtOH (4:1), 75 °C, 18 h, 45% (two steps).

1-(4-(5-Fluoro-6-(((tetrahydro-2H-pyran-4-yl)amino)methyl)pyridin-2-yl)benzyl)-3-isobutylimidazolidine-2,4-dione (LEI-102): To a degassed mixture of **3** (0.29 g, 1.0 mmol, 1.0 eq), **7** (0.56 g, ~1.5 mmol, crude) and K₂CO₃ (1.29 g, 6.0 mmol, 6.0 eq) in toluene:ethanol (10 mL, 4:1, v/v) was added under argon atmosphere Pd(PPh₃)₄ (0.18 g, 0.10 mmol, 0.1 eq). The resulting mixture was heated (75° for 18 h, then cooled to RT and filtered. The filtrate was diluted with EtOAc and washed with H₂O and brine, dried (MgSO₄), filtered, and concentrated under reduced pressure. The crude product was purified with flash column chromatography (0-20% MeOH in EtOAc) to yield a white solid (0.24 g, 0.53 mmol, 53%). Further purification with preparative HPLC resulted in a yield of 0.204 g (0.45 mmol, 45%). ¹H-NMR (400 MHz, CD₃CN) δ 8.05 (d, *J* = 8.3 Hz, 2H), 7.86 (dd, *J* = 8.7, 3.6 Hz, 1H), 7.61 (t, *J* = 9.0 Hz, 1H), 7.34 (d, *J* = 8.1 Hz, 2H), 4.53 (s, 2H), 4.45 (s, 2H), 3.93 (dd, *J* = 11.4, 4.4 Hz, 2H), 3.77 (s, 1H), 3.47 (tt, *J* = 11.8, 3.8 Hz, 2H), 3.30 (td, *J* = 11.9, 1.9 Hz, 2H), 3.24 (d, *J* = 7.3 Hz, 2H), 2.05 (bd, *J* = 13.3 Hz, 2H), 1.99 (dt, *J* = 13.2, 6.6 Hz, 1H), 1.83 (qd, *J* = 12.1, 4.5 Hz, 2H), 0.88 (d, *J* = 6.7 Hz, 6H). ¹³C-NMR (100 MHz, CD₃CN) δ 171.54, 157.25 (d, *J* = 226.6 Hz), 156.12, 153.03 (d, *J* = 4.5 Hz), 140.51 (d, *J* = 16.1 Hz), 138.83, 137.70, 129.13, 128.22, 125.56 (d, *J* = 18.8 Hz), 122.81 (d, *J* = 4.3 Hz), 118.38, 66.55, 55.55, 50.30, 46.81 (d, *J* = 7.9 Hz), 42.95, 30.02, 28.32, 20.32. LCMS (LCQ Advantage, 10 90%): *t_r* = 5.32 min, *m/z*: 455.27 [M+H]⁺, 908.93 [2M+H]⁺. HRMS (ESI⁺) *m/z*: calcd. for C₂₅H₃₂FN₄O₃ [M+H], 455.245; found, 455.245.

Biology

General Remarks

Monoclonal M2 mouse anti-FLAG primary antibody (#F3165) was purchased from Sigma-Aldrich (Zwijndrecht, the Netherlands), while secondary goat anti-mouse HRP-conjugated antibody (#115-035-003) was bought from Jackson ImmunoResearch Laboratories (West Grove, PA, USA). Bicinchoninic acid

(BCA) ad BCA protein assay reagent was obtained from Pierce Chemical Company (Rockford, IL, USA). [³H]-RO6957022 (specific activity 82.83 Ci mmol⁻¹) was custom synthesized at F. Hoffman-La Roche Ltd (Basel, Switzerland). [³⁵S]GTPγS (specific activity 1250 Ci mmol⁻¹ #NEG030H250UC), [³H]-CP-55,940 (specific activity 108.5 Ci mmol⁻¹ #NET1051250UC) and GF/C filter plates (#6055690) were purchased from PerkinElmer (Waltham, MA, USA). CP-55,940 (#C1112), AM630 (#SML0327) and DL-dithiothreitol (DTT, #646563) were obtained from Sigma-Aldrich, HU308 (#H800010) was from LKT Laboratories (St. Paul, MN, USA), APD371 was provided by F. Hoffmann-La Roche Ltd, anandamide (AEA, #1339), 2-Arachidonylglycerol (2-AG, #1298) and phenylmethylsulfonyl fluoride (PMSF, #4486) were purchased from Tocris Bioscience (Bristol, UK) and GDP (#J61646) was from Thermo Fisher Scientific (Waltham, MA, USA). All buffers and solutions were prepared using Millipore water (deionized using a MilliQ A10 Biocel with a 0.22 μm filter) and analytical grade reagents and solvents. Buffers are prepared at room temperature (RT) and stored at 4 °C, unless stated otherwise.

Quantification and statistical analysis

All experimental data were analyzed using GraphPad Prism 9.0 (GraphPad Software Inc., San Diego, CA). All values obtained are means ± standard error of the mean (SEM) of at least three independent experiments performed in duplicate, unless stated otherwise.

From [³H]-RO6957022 competition association assays, the k_{on} and k_{off} were determined by non-linear regression analysis, using the “kinetics of competitive binding” model as described by Motulsky and Mahan⁵⁵:

$$\begin{aligned}
 K_a &= k_1 \cdot [L] \cdot 10^{-9} + k_2 \\
 K_b &= k_3 \cdot [I] \cdot 10^{-9} + k_4 \\
 S &= \sqrt{(K_a - K_b)^2 + 4 \cdot k_1 \cdot k_3 \cdot [L] \cdot [I] \cdot 10^{-18}} \\
 K_f &= 0.5 \cdot (K_a + K_b + S) \\
 K_s &= 0.5 \cdot (K_a + K_b - S) \\
 Q &= \frac{Bmax \cdot k_1 \cdot [L] \cdot 10^{-9}}{K_f - K_s} \\
 [Y] &= Q \cdot \left(\frac{k_4 \cdot (K_f - K_s)}{K_f \cdot K_s} + \frac{k_4 - K_f}{K_f} \cdot e^{(-K_f \cdot X)} - \frac{k_4 - K_s}{K_s} \cdot e^{(-K_s \cdot X)} \right)
 \end{aligned}$$

Where [L] is the radioligand concentration per experiment (~1.5 nM), I is the IC₅₀ concentration of agonist (nM), X is the time (s), and Y is the specific binding of the radioligand (dpm). K_a and K_b are the observed association rate constants (k_{obs}) of the radioligand and the agonist of interest, respectively. k_1 and k_3 are the association rate constants (k_{on} in M⁻¹s⁻¹) of [³H]-RO6957022 (determined per experiment) and the agonist of interest, respectively. Similarly, k_2 and k_4 are the dissociation rate constants (k_{off} in s⁻¹) of [³H]-RO6957022 (experimentally determined at 4.3×10^{-4} s⁻¹, data not shown) and the agonist of interest, respectively. The engagement time (ET in seconds) of the agonists of interest was determined at 1 μM of agonist using the equation $ET = 1/(k_{on} \cdot 1 \times 10^{-6})$. The residence time distribution (RTD in min) was calculated using the equation $RTD = 1/(60 \cdot k_{off})$.⁵⁶ The association and dissociation rate constants were used to calculate the kinetic K_D using: $K_D = k_{off}/k_{on}$.

[³⁵S]GTPγS agonist responses on CB₂R constructs were baseline-corrected for the individual mutant's basal activity. The responses were normalized to the basal activity of the construct (0%) and top of the CP-55,940 (for WT responses only) or WT curve (for mutants, 100%). The potency (pEC₅₀) and efficacy (E_{max}) values were obtained by non-linear regression to a sigmoidal concentration-effect curve with a

Hill slope of 1 by using the “log(agonist) vs response (three parameters)” model. [³⁵S]GTPγS data from CB₁R constructs were expressed as fold over the mutant’s basal activity to also quantify the effects of CB₂R selective agonists.

Displacement assays were baseline-corrected with NSB and normalized to this value (0%) and TB (100%). The equilibrium dissociation constants (K_D) of [³H]-CP-55,940 on different mutants were calculated from homologous displacements by non-linear regression analysis, using the “one-site homologous” model. The half-maximal inhibitory concentrations (pIC₅₀) of the agonists in [³H]-CP-55,940 and [³H]-RO6957022 assays were obtained by non-linear regression analysis of the homologous and heterologous displacement curves and further converted into inhibitory constant pK_i using the Cheng-Prusoff equation.⁵⁷ In which the experimentally determined K_D for each construct was used for [³H]-CP-55,940 assays or 0.78 nM for [³H]-RO6957022 assays (data not shown).

Differences in pEC₅₀, E_{max}, pK_D and pK_i values for each mutant compared to WT were analyzed using a one-way Welch’s ANOVA with Dunnett’s T3 multiple comparisons test or an unpaired Student’s t-test with Welch’s correction. Significant differences are displayed as * p < 0.05; ** p < 0.01, *** p < 0.001 and **** p < 0.0001.

Cell Culture

Cells were cultured and prepared similar to previously described.⁴⁴ *Spodoptera frugiperda* (Sf9) cells were used for CB₂R-Gi co-expression for cryo-EM studies. Sf9 cells were grown in ESF 921 medium (Expression systems) at 27 °C and 125 rpm. For transfections, human embryonic kidney 293 T (HEK293T; female, ATCC #CRL-3216) cells were grown as monolayers in culture medium i.e. Dulbecco’s Modified Eagle’s Medium (Sigma-Aldrich #6546), supplemented with 10% fetal calf serum (Sigma-Aldrich #F7524), 2 mM L-glutamine (Sigma-Aldrich #G8541), 100 IU/mL penicillin and 100 µg/mL streptomycin (Duchefa Biochemie #P0142 and #S0148) under a humidified atmosphere at 37 °C with 5% CO₂. Subculture was done twice a week at 80 – 90% confluence on 10 cm ø plates by trypsinization. CHO cells stably expressing hCB₂ (CHOK1_hCB₂R_bgal; PathHunter EA Parental Cell line, female, DiscoverX #93-0706C2) were cultured in Ham’s F12 Nutrient Mixture (Sigma-Aldrich #4888) supplemented with 10% fetal calf serum, 2 mM L-glutamine, 100 IU/mL penicillin, 100 µg/mL streptomycin, 300 µg/mL hygromycin (Bio-Connect #ANT-HG-5) and 800 µg/mL G418 (Bio-Connect #SC-29065B) in a humidified atmosphere at 37 °C with 5% CO₂. Cells were subcultured twice a week when reaching 80 - 90% confluence on 10 or 15 cm ø plates by trypsinization.

Constructs and Expression of CB₂R and G_i heterotrimer

A N-BRIL fused WT human CB₂R construct was cloned into a modified pFastBac1 vector with an HA signal sequence at the N-terminus followed by a 10×His-tag and a FLAG-tag. Human G_{α1} and G_{β1γ2} subunits were cloned into pFastbac1 and pFastDual vector individually. The CB₂R and G_i heterotrimer were co-expressed in Sf9 insect cells using the Bac-to-Bac Baculovirus Expression System (Invitrogen). Sf9 cells were infected at a cell density of 2-2.5×10⁶ cells/ml with three separate virus preparations for CB₂R, G_{α1} and G_{β1γ2} at a ratio of 1:2:2. The infected cells were cultured at 27 °C for 48 h before collection by centrifugation and the cell pellets were stored at -80 °C for future use.

Constructs, Expression and Purification of scFv16

Methods of complex expression and purification in the current study have been described previously.³⁴ The CB₂R and G_i heterotrimer were co-expressed in Sf9 insect cells using the Bac-to-Bac Baculovirus Expression System (Invitrogen). Cells were infected with three separate virus preparations for CB₂R, G_{α1} and G_{β1γ2} at a ratio of 1:2:2 at a cell density of 2.5×10⁶ cells/mL. After 48 h, the cell culture was collected by centrifugation and the cell pellets were stored at -80 °C until use. The cell pellets were thawed and lysed in the hypotonic buffer of 10 mM HEPES (pH 7.5), 10 mM MgCl₂, 20 mM KCl with EDTA-free

complete protease inhibitor cocktail tablets (Roche, #5056489001). The CB₂R-G_i complex was formed in membranes by addition of 25 μM agonist (LEI-102, APD371, HU308 and CP-55,940, respectively) and 2 units of apyrase (NEB, #M0398S) in the presence 500 μg scFv16. The lysate was incubated for overnight at 4 °C and discard the supernatant by centrifugation at 40,000 rpm for 30 min. Subsequently, the solubilization buffer containing 50 mM HEPES (pH 7.5), 100 mM NaCl, 0.75% (w/v) lauryl maltose neopentyl glycol (LMNG, Anatrace, #4216588), 0.15% (w/v) cholesterol hemisuccinate (CHS, Sigma-Aldrich, #C6512) supplemented with 25 μM agonist and 2 units of apyrase (NEB) were added to solubilize complexes for 2 h at 4 °C. Insoluble material was removed by centrifugation at 40,000 rpm for 30 min and the supernatant was immobilized by batch binding to TALON IMAC resin (Clontech, #635507) including 20 mM imidazole over 6 h at 4 °C. Then, the resin was packed and washed with 15 column volumes (CVs) of washing buffer I containing 25 mM HEPES (pH 7.5), 100 mM NaCl, 10% (v/v) glycerol, 0.1% (w/v) LMNG, 0.02% (w/v) CHS, 30 mM imidazole and 20 μM agonist, and 15 CVs of washing buffer II containing 25 mM HEPES (pH 7.5), 100 mM NaCl, 10% (v/v) glycerol, 0.03% (w/v) LMNG, 0.006% (w/v) CHS, 50 mM imidazole and 20 μM agonist. After that, the protein was eluted using 3 CVs of elution buffer containing 25 mM HEPES (pH 7.5), 100 mM NaCl, 10% (v/v) glycerol, 0.01% (w/v) LMNG, 0.002% (w/v) CHS, 250 mM imidazole and 25 μM agonist. Finally, the complex was concentrated using the centrifugal filter with 100 KD molecular weight cutoff and loaded onto a Superdex200 10/300 GL column (GE Healthcare) with buffer containing 20 mM HEPES (pH 7.5), 100 mM NaCl, 0.00075% (w/v) LMNG, 0.00025% GDN (Anatrace, #GDN101), 0.0001% (w/v) CHS, 100 μM TCEP. The fractions consisting of purified CB₂-G_i complex were collected and concentrated to 0.8-1.0 mg/ml for electron microscopy experiments.

CB₂R-G_i-scFv16 Complex Formation and Purification

The cell pellets corresponding to 1 L CB₂R-G_i co-expression culture were thawed and lysed in the hypotonic buffer of 10 mM HEPES, pH7.5, 10 mM MgCl₂, 20 mM KCl with EDTA-free complete protease inhibitor cocktail tablets (Roche). The CB₂R-G_i complex was formed in membranes by addition of 20 μM agonist (LEI-102, APD371, HU308 or CP-55,940) and 2 units of apyrase (NEB) in the presence of 500 μg scFv16. The lysate was incubated overnight at 4 °C and the supernatant discarded after centrifugation at 40,000 rpm for 30 min. The complex was separated from the membranes in buffer containing 50 mM HEPES (pH 7.5), 100 mM NaCl, 0.75% (w/v) lauryl maltose neopentyl glycol (LMNG, Anatrace), 0.15% (w/v) cholesterol hemisuccinate (CHS, Sigma-Aldrich), 20 μM agonist and 2 units of apyrase (NEB) at 4 °C for 2 h. The supernatant was isolated by ultracentrifugation, and then incubated with TALON IMAC resin (Clontech) and 20 mM imidazole over 6 h at 4 °C. The resin was washed with 15 column volumes of washing buffer I (25 mM HEPES (pH 7.5), 100 mM NaCl, 10% (v/v) glycerol, 0.1% (w/v) LMNG, 0.02% (w/v) CHS, 30 mM imidazole and 20 μM agonist) and 15 column volumes of washing buffer II (25 mM HEPES (pH 7.5), 100 mM NaCl, 10% (v/v) glycerol, 0.03% (w/v) LMNG, 0.006% (w/v) CHS, 50 mM imidazole and 20 μM agonist). The protein was eluted using 3 column volumes of elution buffer (25 mM HEPES (pH 7.5), 100 mM NaCl, 10% (v/v) glycerol, 0.01% (w/v) LMNG, 0.002% (w/v) CHS, 250 mM imidazole and 25 μM agonist). The purified CB₂R-G_i-scFv16 complex was concentrated, then injected onto a Superdex200 10/300 GL column (GE Healthcare) equilibrated in buffer (20 mM HEPES (pH 7.5), 100 mM NaCl, 0.00075% (w/v) LMNG, 0.00025% GDN, 0.0001% (w/v) CHS, 100 μM TCEP). The complex peak fractions were collected and concentrated individually to 0.8-1.0 mg/mL for electron microscopy experiments.

Cryo-EM grid Preparation and data collection

For cryo-EM grids preparation of the CB₂R-G_i complexes, 3 μL of the concentrated protein was loaded to a glow-discharged holey carbon grid (CryoMatrix Amorphous alloy film R1.2/1.3, 300 mesh), and subsequently were plunge-frozen in liquid ethane using a Vitrobot Mark IV (Thermo Fisher Scientific).

The chamber of Vitrobot was set to 100% humidity at 4 °C. The sample was blotted for 2.5 s with blot force 2. Cryo-EM images were collected on a Titan Krios microscope operated at 300 kV equipped with a Gatan Quantum energy filter, with a slit width of 20 eV, a Gatan K2 summit direct electron camera (Gatan). Images were taken at a dose rate of 8e-/Å²/s with a defocus range of -0.8 to -2.0 μm using SerialEM software⁵⁸ in EFTEM nanoprobe mode, with 50 μm C2 aperture, at a calibrated magnification of 130,000 corresponding to a magnified pixel size of 1.04 Å. The total exposure time was 8.1 s and 45 frames were recorded per micrograph.

Image Processing and 3D Reconstruction

The cryo-EM data processing was performed with CryoSPARC.⁵⁹ For CB₂R-G_{αi}-scFv16-APD371/LEI-102/HU308/CP-55,940 dataset, a total of 7443, 5282, 7530 and 6473 movies were collected, respectively. For all datasets, patch motion correction was used for beam-induced motion correction. Contrast transfer function (CTF) parameters for each micrograph were determined by patch CTF estimation. Using Blob Picker in CryoSPARC to auto pick particles in the first 500 micrographs of CB₂R-G_{αi}-scFv16-APD371 complex dataset and then 258347 particles were extracted to conduct 2D classification. 9277 particles in good 2D patterns were selected as templates to pick better particles. 5,239,870, 3,398,611, 4,653,294 and 3,595,875 particles extracted, respectively, in a 256 Å box were divided into three hundred two-dimensional (2D) class averages with a maximum alignment resolution of 6 Å. Then, 1,152,146, 762,471, 355,832 and 440,292 particles were selected from good 2D classification after two round 2D classification, individually. Following 2D classification, these particles were subjected for ab initio reconstruction into four classes. After heterogeneous refinement, homogeneous refinement, non-uniform refinement and local refinement of the best-looking dataset in CryoSPARC, the final map has an indicated global resolution of 3.08 Å, 2.98 Å, 2.97 Å and 2.84 Å at a Fourier shell correlation (FSC) of 0.143, respectively. Local resolution was determined using the Bsoft package with half maps as input maps.⁶⁰

Model Building and Refinement

The CB₂R-AM12033 cryo-EM structure and G_i protein in CB₂R were used as the starting model for model refinement. The model was docked into the CB₂R-agonist-G_i-scFv16 EM density map using Chimera⁶¹, followed by iterative manual adjustment and rebuilding in COOT⁶² and phenix.real_space_refine in Phenix.⁶³ The model statistics were validated using MolProbity.⁶⁴ Structural figures were prepared in Chimera and PyMOL (<http://www.pymol.org>). The extent of any model overfitting during refinement was measured by refining the final model against one of the half-maps and by comparing the resulting map versus model FSC curves with the two half-maps and full model.

Generation of Mutants

The WT CB₁R and CB₂R genes were subcloned into vector pcDNA3.1 with an N-terminal HA signal peptide and FLAG-tag. Mutations were introduced by QuikChange PCR (as described by supplier).

Transfection

24 h prior to transfection, HEK293T cells were seeded on 10 cm ø plates to reach approximately 50% confluence at the start of transfection. The cells were transfected with 10 μg plasmid DNA of WT hCB₂R or hCB₁R receptor, or mutant receptor using the calcium phosphate precipitation method.⁶⁵ In short, a DNA-calcium mix was made containing 270 mM CaCl₂ and 10 μg plasmid DNA to which Hank's Balanced Salt Solution (HBSS; 280 mM NaCl, 10 mM KCl, 1.5 mM Na₂HPO₄ and 50 mM HEPES) was added in a 1:1 (v/v) ratio and mixed by aeration to create consistent calcium phosphate precipitates. For transfection, 1 mL DNA-calcium mix was added per 10 cm ø plate, followed by a 24/48 h incubation under a humidified atmosphere at 37 °C with 5% CO₂.

Enzyme-Linked Immunosorbent Assay (ELISA)

Receptor expression after transfection was measured with ELISA. A control was made by transfecting HEK293T cells transfected with an empty pcDNA3.1 vector. HEK293T cells were transfected 24 h prior to seeding. A clear, culture-treated 96-wells plate was treated with 100 μ L/well 0.1 mg/mL poly-D-lysine for 30 min. after which the wells were washed twice with 200 μ L/well PBS and the plate air-dried for 1 h. Cells were detached by incubation for 15 min. at 37 °C with 1.5 mL warm PBS/EDTA. 3 mL medium (DMEM, D6546 Sigma Aldrich) was added, cells resuspended and centrifuged at 200 x g for 5 min. The supernatant was aspirated and cells resuspended in medium to 1,000,000 cells/mL. 100 μ L is added to each well so that a total of 100,000 cells have been seeded per well.

24 h after seeding the plate was aspirated and washed with 200 μ L PBS/well. The cells were fixed with 100 μ L/well 4% formaldehyde for 10 min. Then washed twice with 100 μ L/well Tris-buffered saline (TBS). The plate was incubated with 100 μ L/well blocking buffer (2% (w/v) BSA in TBST) for 30 minutes at 400 rpm. The blocking buffer was aspirated and the plate was incubated with 100 μ L/well mouse anti-FLAG primary antibody (1 μ g/mL, Sigma F3165) for 2 h at 400 rpm. The antibody was aspirated and after washing thrice with 200 μ L/well TBST prior to addition of goat anti-mouse HRP-conjugated secondary antibody (1:10,000 dilution. Jackson ImmunoResearch Laboratories 115-035-003) and 1 h incubation at 400 rpm. The plate was aspirated and washed thrice with 200 μ L/well TBS. The plate was incubated with 100 μ L/well TMB in the dark for 10 min. until the reaction was quenched with 100 μ L 1 M H₃PO₄. Absorbance was measured at 450 nm with the Wallac EnVision™ (Perking Elmer).

Membrane Preparation

For membrane preparation, HEK293T cells were harvested 48 h after transfection. Cells were detached by scraping into 3 mL of phosphate-buffered saline (PBS) and subsequently centrifuged at 2000 x g for 5 min. Pellets were resuspended in ice-cold Tris buffer (50 mM Tris-HCl, pH 7.4) and homogenized with an Ultra Turrax homogenizer (IKA-Werke GmbH & Co. KG, Staufen, Germany). Cytosolic and membrane fractions were separated using a high-speed centrifugation step of 31,000 rpm in a Beckman Optima LE-80K ultracentrifuge with Ti70 Rotor for 20 min at 4 °C. After a second cycle of homogenization and centrifugation, the final pellets were resuspended in 50 mM Tris-HCl (pH 7.4), 5 mM MgCl₂ and stored in 100 μ L aliquots at -80 °C until use. CHOK1_hCB₂R_bgal cells were harvested when reaching 90% confluence in 15 cm ϕ plates after one week subculture at a 1:6 ratio. Membrane preparation followed a similar procedure as described above. Final membrane pellets were resuspended in 50 mM Tris-HCl (pH 7.4) and stored in 100 μ L aliquots at -80 °C until use. Membrane protein concentrations were determined using a BCA protein determination assay as described by the manufacturer (Smith et al., 1985).

[³H]-RO6957022 Competition Association Assays

Prior to kinetic assessment of agonist binding, the affinity (IC₅₀) of the agonists at the hCB₂R was determined in [³H]-RO6957022 displacement assays. CHOK1_hCB₂R_bgal was thawed, homogenized, and subsequently diluted to 1 μ g protein per well. When studying endocannabinoids, membranes were preincubated with 50 μ M PMSF for 30 minutes. Membranes were incubated with ~1.5 nM [³H]-RO6957022 and six increasing concentrations of competing agonists in a total volume of 100 μ L assay buffer (50 mM Tris-HCl (pH 7.4), 0.1% (w/v) BSA). Incubations were done for 2 h at 10 °C to reach equilibrium. Subsequently, in competition association assays, agonists were incubated at their IC₅₀ concentration in the presence of ~1.5 nM [³H]-RO6957022 in a total volume of 100 μ L assay buffer at 10 °C. Competition was initiated by addition of membrane homogenates at different time points for 2 h. Non-specific binding was determined with 10 μ M AM630. Organic solvent (DMSO or acetonitrile) concentrations were <1% in all samples. Incubations were terminated by rapid vacuum filtration with ice-cold 50 mM Tris-HCl (pH 7.4), 0.1% (w/v) BSA buffer through Whatman GF/C filters using a

Filtermate 96-well harvester (PerkinElmer). Filters were dried for at least 30 min at 55 °C and subsequently 25 µL MicroScint scintillation cocktail was added per well. Filter-bound radioactivity was measured by scintillation spectrometry using a Microbeta2 2450 counter (PerkinElmer).

[³⁵S]GTPγS Binding Assays

G protein activation was measured with agonists LEI-102, APD371, HU308, CP-55,940 AEA and 2-AG. Transient HEK293T membrane homogenates (10 µg/well) were diluted in assay buffer (50 mM Tris-HCl (pH 7.4), 5 mM MgCl₂, 150 mM NaCl, 1 mM EDTA, 0.05% BSA (w/v) and 1 mM DTT) and pretreated with 10 µg saponin and 1 µM GDP. For endocannabinoid samples, the membranes were additionally pretreated for 30 minutes with 50 µM phenylmethylsulfonyl fluoride (PMSF) before agonist addition. The membranes were incubated with 10 µM CP-55,940 (E_{max}) or six increasing concentrations of agonist (ranging from 0.01 nM to 1 µM) for 30 minutes at rt. Basal receptor activity was determined in the presence of vehicle only (0.2% DMSO/acetonitrile). [³⁵S]GTPγS (0.3 nM) was added for 100 µL final volume. The mixture was co-incubated for 90 minutes at 25 °C while shaking at 400 rpm. The plate was filtered with wash buffer (50 mM Tris-HCl (pH 7.4), 0.1% (w/v) BSA) through Whatman GF/C filters using a Filtermate 96-well harvester (PerkinElmer). Filters were dried for at least 30 min at 55 °C and subsequently 25 µL MicroScint scintillation cocktail was added per well. Filter-bound radioactivity was measured by scintillation spectrometry using a Microbeta2 2450 counter (PerkinElmer).

[³H]-CP-55,940 Homologous and Heterologous Displacement Assays

To determine agonist affinity (K_i) on WT and mutant receptors. The amount of transient HEK293T membrane was 0.75 µg to 10 µg protein per well, chosen to reach a specific [³H]-CP-55,940 binding window of 1200-1500 disintegrations per minute (dpm). Only the CB₂R-quadruple mutant was used with 20 µg/well for a window of ~500 dpm. Membranes were thawed and subsequently homogenized using an Ultra Turrax homogenizer. For AEA and 2-AG the membranes were preincubated for 30 minutes with 50 µM PMSF. Homologous displacement assays were performed with 1.5 nM final concentration [³H]-CP-55,940 against six increasing concentrations (0.1 nM to 10 µM) of cold CP-55,940. Heterologous displacement assays were done with LEI-102, APD371, HU308, AEA and 2-AG using 1.5 nM final concentration [³H]-CP-55,940 with one concentration (10 µM) or six increasing concentrations (ranging from 0.1 nM to 10 µM) in assay buffer. To a 96-well roundbottom plate was added 25 µL assay buffer, 25 µL agonist (or buffer for negative control), 25 µL 6 nM [³H]-CP-55,940, and 25 µL membrane, in order. Incubation was for 2 h at 25 °C. Separation of bound from free radioligand was performed by rapid filtration through GF/C filters (Whatman, Clifton, NJ) using a Filtermate Harvester (Brandel Inc., Gaithersburg, MD). Filters were dried for at least 30 min at 55 °C and subsequently 25 µL MicroScint scintillation cocktail was added per well. Filter-bound radioactivity was measured by scintillation spectrometry using a Microbeta2 2450 counter (PerkinElmer).

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