(4-(Bis(4-Fluorophenyl)Methyl)Piperazin-1-yl)(Cyclohexyl) Methanone Hydrochloride (LDK1229): A New Cannabinoid CB1 Receptor Inverse Agonist from the Class of Benzhydryl Piperazine Analogs

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ABSTRACT

Some inverse agonists of cannabinoid receptor type 1 (CB1) have been demonstrated to be anorectic antiobesity drug candidates. However, the first generation of CB1 inverse agonists, represented by rimonabant (SR141716A), otenabant, and tariabant, are centrally active, with a high level of psychiatric side effects. We generated a new CB1 inverse agonist, (4-(bis(4-fluorophenyl)methyl)piperazin-1-yl)(cyclohexyl)methanone hydrochloride (LDK1229), from the class of benzhydryl piperazine analogs. This compound binds to CB1 more selectively than cannabinoid receptor type 2, with a Kᵢ value of 220 nM. Comparable CB₁ binding was also observed by analogs 1-[bis(4-fluorophenyl)methyl]-4-cinnamylpiperazine dihydrochloride (LDK1203) and 1-[bis(4-fluorophenyl)methyl]-4-tosylpiperazine hydrochloride (LDK1222), which differed by the substitution on the piperazine ring where the piperazine of LDK1203 and LDK1222 are substituted by an alkyl group and a tosyl group, respectively. LDK1229 exhibits efficacy comparable with SR141716A in antagonizing the basal G protein coupling activity of CB₁, as indicated by a reduction in guanosine 5’-O-(3-thio)triphosphate binding. Consistent with inverse agonist behavior, increased cell surface localization of CB₁ upon treatment with LDK1229 was also observed. Although docking and mutational analysis showed that LDK1229 forms similar interactions with the receptor as SR141716A does, the benzhydryl piperazine scaffold is structurally distinct from the first-generation CB₁ inverse agonists. It offers new opportunities for developing novel CB₁ inverse agonists through the optimization of molecular properties, such as the polar surface area and hydrophilicity, to reduce the central activity observed with SR141716A.

Introduction

The cannabinoid receptors are members of the class A superfamily of G protein–coupled receptors (GPCRs). The cannabinoid receptor 1 (CB₁) is present in high abundance throughout the central nervous system (Howlett, 1995) but is also expressed in a number of peripheral tissues, such as the cardiovascular and reproductive systems as well as the gastrointestinal tract (Croci et al., 1998; Batkai et al., 2001; Engeli et al., 2005), and is involved in substance addiction, chronic pain, memory, and metabolic and inflammatory disorders (Howlett et al., 2004; Mackie, 2006; Pertwee, 2006). A second subtype of the cannabinoid receptors, the cannabinoid receptor 2 (CB₂), is predominantly found in immune cells and non-neuronal tissues (Gallegue et al., 1995) and is implicated in a variety of modulatory functions, including immune suppression, induction of apoptosis, and induction of cell migration (Basu and Dittel, 2011). The CB₁ receptor preferentially couples to the Gᵢₒ type of G proteins (Howlett and Fleming, 1984) and has been functionally linked to the inhibition of adenylyl cyclase (Slipetz et al., 1995) and the activation of mitogen-activated protein kinases, including extracellular signal-regulated kinase-1 and -2, p38 mitogen-activated protein kinase, and c-Jun N-terminal kinase.
the CB1 receptor possesses agonist-independent constitutive activity. Upon binding to receptor agonists, such as the endogenous arachidonyl-ethanolamide and 2-arachidonoyl glycerol, or the synthetic agonists 5-(1,1-dimethylheptyl)-2-(5-hydroxy-2-(3-hydroxypropyl) cyclohexyl)phenol (CP55,940) and (R(+)-[2,3-dihydro-5-methyl-3-(4-morpholinylmethyl)pyrrole-1,2,3-d]-1,4-benzoxazin-6-yl]-1-naphthalenyl-methanone (WIN55,212-2), the active form of the receptor predominates. Interestingly, the CB1 receptor possesses agonist-independent constitutive or basal activity that can be induced by inverse agonists (Fertwey, 2005). This ligand-independent activity led to a receptor model that accounts for multiple activation states (Gether and Kobilka, 1998; Ghanouni et al., 2001) with distinct biochemical characteristics, including extent and selectivity of G protein or β-arrestin coupling (Kenakin, 1995).

Inverse agonists of the CB1 receptor have attracted considerable attention in drug discovery because of their ability to regulate appetite and manage substance addiction (Janero, 2012; Janero and Makriyannis, 2009; Silvestri and Di Marzo, 2012). Consequently, considerable effort has been invested in discovering compounds that can regulate the constitutive activity of the CB1 receptor. However, SR141716A (rimonabant) (Rinaldi-Carmona et al., 1994), the only CB1 inverse agonist to be briefly clinically marketed in Europe for the control of obesity (Moreira and Crippa, 2009), was removed from use due to its severe neuropsychiatric side effects, including mood-depressant actions (Després et al., 2005; Traynor, 2007). The first generation of CB1 inverse agonists are commonly derived from diaryl analogs of pyrazole (e.g., SR141716A) or pyrazole bioisosteres, such as imidazole, triazole, thiazole, and pyrazoline (Lange and Kruse, 2005, 2008; Muccioli and Lambert, 2005). However, most of these are brain penetrant due to their physicochemical nature (Chorvat, 2013) and will likely generate unwanted side effects in the central nervous system. In an effort to develop new CB1 inverse agonist scaffolds, we analyzed the common pharmacophore of this class of compounds, which was proposed by Lange and Kruse (2005). This showed that the biaryl groups connecting to a central heteroaromatic ring are pivotal in forming aromatic stacking interactions with CB1 receptors (McAllister et al., 2004; Shim et al., 2012). This brought our attention to the benzhydryl piperazine scaffold, which exhibits a similar structure to the common pharmacophore of CB1 inverse agonists in lieu of the biaryl heteroaromatic ring moiety. Hence, we synthesized a group of benzhydryl piperazine analogs, and demonstrated that some of the synthesized analogs, i.e., LDK1203, LDK1222, and LDK1229, exhibited inverse agonist binding profiles for the CB1 receptor. In addition to its binding profile to the CB1 receptor, LDK1229’s inverse agonism is evident from its effect on basal as well as agonist-induced G protein coupling and its impact on the internalization and cell surface expression of CB1. Docking studies using a model of the inactive CB1 receptor showed that LDK1229 forms interactions with the receptor that were consistent with the inverse agonist SR141716A. Discovering new and improved means for inhibiting the activity of CB1 is critical for understanding the constitutive activity of CB1 and developing new therapeutic agents for treating substance addiction and disorders associated with CB1 activity.

Materials and Methods

Synthesis

The benzhydrylpiperazine analogs LDK1203, LDK1222, and LDK1229 were synthesized by alkylation, tosylation, and acylation of the 1-(4,4'-difluorobenzhydryl)piperazine (Fig. 1A). The employed 1-(4,4'-difluorobenzhydryl)piperazine (II) was prepared by mono-alkylation of piperazine with 4,4'-difluorobenzhydrol chloride (I) that was obtained by halogenation of 4,4'-difluorobenzhydrol methanol (I) with oxaly chloride, as previously reported (Weiwer et al., 2012). The obtained free bases of the benzhydrylpiperazine analogs were then converted to their corresponding hydrochloric salts by reacting with an ethereal solution of HCl. The chemical identity of the newly synthesized compounds is as follows: LDK1203 is 1-(bis[4-fluorophenyl)methyl]-4-cinnamylpiperazine dihydrochloride, LDK1222 is 1-(bis[4-fluorophenyl)methyl]-4-tosylpiperazine hydrochloride, and LDK1229 is 1-(bis[4-fluorophenyl)methyl]piperazin-1-yl)cyclohexyl)methanone hydrochloride. The structure confirmation data for the free bases of LDK1203, LDK1222, and LDK1229 are as follows. LDK1203: 1H NMR (500 MHz, chloroform-d) 7.28–7.40 (m, 8H), 7.23 (t, J = 7.5 Hz, 1H), 6.99 (t, J = 7.5 Hz, 4H), 6.53 (d, J = 15 Hz, 1H), 6.28 (d, J = 15 Hz, 1H), 4.26 (s, 1H), 3.22 (d, J = 6 Hz, 2H), 2.24–2.78 (m, 8H). Mass (MS) (electron impact [EI]): m/z = 404.21 (M·'). LDK1222: 1H NMR (500 MHz, chloroform-d) 7.64 (d, J = 7.9 Hz, 2H), 7.36 (d, J = 7.9 Hz, 2H), 7.22–7.31 (m, 4H), 6.94 (t, J = 8.4 Hz, 4H), 4.22 (s, 1H), 2.91–3.10 (m, 4H), 2.48 (s, 3H), 2.38–2.47 (m, 4H). MS (EI): m/z = 442.15 (M·').

LDK1229: 1H NMR (500 MHz, chloroform-d) 7.35 (dd, J = 8.2, 5.5 Hz, 4H), 6.99 (t, J = 8.2 Hz, 4H), 4.22 (s, 1H), 3.61 (t, J = 5 Hz, 2H), 3.42 (t, J = 5 Hz, 2H), 2.43 (t, J = 5 Hz, 2H), 2.42 (t, J = 5 Hz, 2H), 2.31–2.40 (m, 4H), 1.74–1.82 (m, 2H), 1.65–1.73 (m, 3H), 1.44–1.56 (m, 2H). MS (EI): m/z = 398.2 (M·').

Plasmid Construction

All mutants were generated by site-directed mutagenesis (Quick-Change Kit; Stratagene, La Jolla, CA) using the human CB1 cDNA cloned into pcDNA3.1 as a template, according to the manufacturer’s instructions. All mutations were confirmed by DNA sequencing.

CB1 Expression and Membrane Preparation

Human embryonic kidney 293 (HEK293) cells were maintained in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum and 3.5 mg/ml glucose at 37°C in 5% CO2. For transient expression of the receptors, HEK293 cells were seeded at 800,000 cells/100-mm dish on the day prior to transfection and transfected with 5–10 μg of the wild-type or mutant human CB1 receptor cloned into pcDNA3.1 using the calcium phosphate precipitation method (Chen and Okaya, 1987). At 24 hours post-transfection, membranes of transfected cells expressing either the wild-type or mutant receptors were prepared as described previously (Ahn et al., 2009).

Radioligand Binding Assay

In the homologous and heterologous competition binding experiments, approximately 7.5 μg of wild-type CB1, CB2, or mutant CB1 membrane was incubated at 30°C for 60 minutes with a fixed tracer ([3H]CP55,940, 141.2 Ci/mmol; [3H]SR141716A, 56 Ci/mmol; or [3H]WIN55,212-2, 52.2 Ci/mmol, PerkinElmer Life Sciences, Boston, MA) concentration typically at its Kd, which was determined from saturation-binding isotherms (see Results for details). Binding assays were performed with at least nine concentrations of unlabeled competitor ligand (ranging between 100 pM and 100 μM), as described previously (Ahn et al., 2012). Nonspecific binding was determined in the absence of a competitor. Results from the competition binding experiments were analyzed using a one-site saturation binding model to determine the Kd and Bmax of the CB1 receptor. The equilibrium dissociation constant (Kd) was determined as the concentration of radioligand that produced a 50% displacement of the radioligand from the receptor.
Confocal Microscopy and Image Quantification

HEK293 cells were transfected with the CB1 receptor carboxyl terminally fused to green fluorescent protein (GFP) using Lipofectamine (Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions. The CB1-GFP expressing cells were seeded onto 35-mm glass-bottomed dishes (MatTek, Ashland, MA) precoated with poly(d-lysine). Cells were treated with different compounds for various lengths of time and then washed twice with phosphate-buffered saline, followed by fixation with 4% paraformaldehyde for 10 minutes at room temperature. Images were acquired using a Leica confocal laser scanning microscope (Leica, Buffalo Grove, IL), and detection of GFP was carried out following excitation at 488 nm. Quantification of the fluorescence intensity was achieved by using the Quantitative Imaging of Membrane Proteins software (http://go.warwick.ac.uk/bretschneider/quimp) (Dormann et al., 2002; Bosgraaf et al., 2009), a set of plug-ins for the open source program ImageJ (http://rsb.info.nih.gov/ij/). The Boa plug-in was used to detect the cell surface and checked against the cell edge in the transmitted image of each cell. The Ana plug-in was then used to read the cell contours produced by the Boa plug-in and compute the ratio of fluorescence intensity on the cell surface to the average intensity of the cell interior fluorescence. The results are representative of at least four independent transfections and six different images for each condition. Untransfected cells exhibited no apparent fluorescence under the experimental conditions that were used. The parameters for all of the acquired images and their consequent analysis were kept constant throughout.

Ligand and GTP-S Binding Data Analysis

All ligand-binding assays and GTP-S-binding assays were carried out in duplicate. Data are presented as the mean ± S.E. value or the mean with the corresponding 95% confidence limits from at least three independent experiments. K_i values were calculated using the Cheng-Prusoff equation (Cheng and Prusoff, 1973) based on K_d values obtained from saturation-binding analyses. The binding constants for the wild-type and mutant receptors were compared using analysis of variance followed by Bonferroni’s post hoc test for significance. P values of < 0.05 were considered to be statistically significant.

Computational Methods

Conformational Analysis of LDK1229. To generate a library of low-energy conformers of LDK1229, the Spartan Conformation Distribution protocol was used (Wavefunction, Inc., Irvine, CA). In this protocol, the algorithm systematically searches through all rotatable bonds and ring conformations (e.g., alternate chair conformations for flexible rings). The energy of each conformation generated was calculated using the Merck Molecular Force Field (MMFF94S). This calculation yielded 68 unique conformations of LDK1229. The geometry and energy of these 68 conformations was refined by performing ab initio HF-6-31G* energy minimizations on each conformer. To calculate the difference in energy between the global minimum energy conformer and its final docked conformation, rotatable bonds in the global minimum energy conformer were driven to their corresponding value in the final docked conformation and the single-point energy of the resultant structure was calculated at the HF-6-31G* level.

Template Rationale. Our CB1 inactive state model was initially constructed by using the 2.8-Å X-ray crystal structure of bovine rhodopsin as a template (Palczewski et al., 2000). We chose rhodopsin for several reasons: 1) Rhodopsin has an intact ionic lock (R3.50F14/E/D6.30N89), which is the hallmark of the class A GPCR inactive state. 2) The cannabinoid receptors and rhodopsin have very hydrophobic-binding pockets. Crystal structures reveal that the N-terminus of rhodopsin/opsin is closed over the binding pocket, by rapid filtration through Whatman GF/C filters. The radioactivity trapped in the filters was determined by liquid scintillation counting.

Guanosine 5’-O-(3-Thio)Triphosphate Binding Assay

Guanosine 5’-O-(3-thio)triphosphate (GTP*S) binding assays were performed as described previously (Ahn et al., 2012). Briefly, 7.5 µg of membranes were incubated for 60 minutes at 30°C in a total volume of 200 µl GTP*S binding assay buffer (50 mM Tris-HCl, pH 7.4, 3 mM MgCl₂, 0.2 mM EGTA, and 100 mM NaCl), with unlabeled test compounds as indicated: 0.1 nM guanosine 5’-O-(3-thio)triphosphate (C(35)S-GTP*S) (1250 Ci/mmol; PerkinElmer Life Sciences, Boston, MA), 3 µM GDP (Sigma-Aldrich, St. Louis, MO), and 0.1% (w/v) bovine serum albumin. Three micromolar GDP was used to increase the window of basal activity. Nonspecific binding was determined with 10 µM unlabeled GTP*S (Sigma-Aldrich). The reaction was terminated

Fig. 1. Compound structures. (A) Synthesis of benzhydryl piperazine analogs LDK1203, LDK1222, and LDK1229. (a) Oxalyl chloride, dichloromethane (DCM), catalytic N,N-dimethylformamide (DMF). (b) Piperazine, CH₂CN. (c) 3-bromo-1-phenyl-1-propene, K₂CO₃, DMF. (d) TsCl, N,N-diisopropylethylamine (DIPEA), DCM. (e) Cyclohexanecarbonyl chloride, DIPEA, DCM. (f) 1.0 M HCl in ether. (B) Representative members of the first generation of CB1 inverse agonists are shown for comparison.
preventing access from the extracellular milieu (Palczewski et al., 2000; Park et al., 2008; Scheerer et al., 2008). It is very likely that CB₁, with its 112-residue N-terminus, is also closed off to the extracellular milieu. Instead, rhodopsin/opsin have been reported to have lipid portals that are used for entry and exit via the lipid bilayer for 11-cis-retinal/11-trans-retinal as they are shuttled into and out of the receptor (Hildebrand et al., 2000). There is evidence from simulations (Hurst et al., 2010) and from experimental covalent labeling studies (Picone et al., 2005; Pei et al., 2008) that the cannabinoid receptors also possess a portal between transmembrane helix (TMH) 6 and TMH7 through which ligands enter. 3) In addition, the cannabinoid receptors and rhodopsin share an unusual GWNC amino acid sequence motif at the extracellular end of TMH4. Here, a Trp forms an aromatic stacking interaction with Y5.39275. This interaction influences the extracellular positions of TMH3-4-5.

**CB₁ Inactive State Model.** A sequence alignment between the sequence of the human CB₁ receptor (Gerard et al., 1991) and the sequence of bovine rhodopsin was constructed using highly conserved residues as an alignment guide. In addition, the hydrophobicity profile of the sequence was also considered when constructing the sequence alignment. Residues in the bovine rhodopsin structure were then mutated to those of human CB₁. The Monte Carlo–simulated annealing technique Conformational Memories was used to sample the conformational space of TMH6 (Barnett-Norris et al., 2002). This is because TMH6 is known to undergo a functionally necessary conformational change for G protein–mediated signaling. For the inactive state model, the chosen TMH6 conformer was one that enabled the formation of a salt bridge between the highly conserved TMH6 residue D6.30³³⁸ and the highly conserved TMH3 residue R3.50²¹⁴. This salt bridge (also termed the ionic lock) has been shown to be important in maintaining the inactive state in the ß₂-adrenergic (Ballesteros et al., 2001) and serotonin receptor type 2a (5HT-2a) receptors (Visiers et al., 2002). Extracellular and intracellular loops were added to the model using Modeller (Maruc et al., 2013). SR141716A was docked (within the TMH3-4-5-6 region) in this CB₁ inactive state model, and the energy of the complex was minimized, as previously described (Hurst et al., 2006).

**Glide Docking of LDK1229 at the CB₁ Receptor (Inactive State Conformation).** The docking program Glide (version 5.7, Schrödinger, New York, NY), was used to explore possible receptor binding modes of LDK1229. First, the SR141716A-CB₁ binding site was chosen as a starting point for Glide docking studies because of the similar pharmacology between SR141716A and LDK1229. Second, LDK1229 displaces SR141617A in competitive binding experiments, suggesting some commonality between their binding sites. Thus, Glide was used to generate a grid centered on the center of mass of our previously reported binding site for SR141716A at the CB₁ receptor (Hurst et al., 2002; McAllister et al., 2003, 2004). The grid dimensions were 26 x 26 x 26 Å. This grid size allowed Glide to thoroughly explore the receptor for possible binding site(s). In addition, the results of previously reported mutagenesis and synthetic studies suggest that SR141716A forms an important interaction with K3.281⁹² that is necessary for its ability to act as an inverse agonist of G protein–mediated signaling (Hurst et al., 2002, 2006). Therefore, Glide was required to dock LDK1229 in such a way so that it formed a hydrogen bond with K3.281⁹². The only other constraint used was the requirement that LDK1229 must be docked within the exploration grid. Standard precision was selected for the docking setup. A total of 68 conformations of LDK1229 was flexibly docked using Glide. The best Glide dock was chosen for subsequent calculations. The chosen Glide dock was minimized using the minimization protocol described below.

**Receptor Model Energy Minimization Protocol.** The energy of the LDK1229-CB₁ complex, including loop regions, was minimized using the Optimized Potentials for Liquid Simulations 2005 force field in Macromodel 9.9 (Schrödinger). An 8.0-Å extended nonbonded cutoff (updated every 10 steps), a 20.0-Å electrostatic cutoff, and a 4.0-Å hydrogen bond cutoff were used in the calculation. The minimization was performed in two stages. In the first stage, the TMH backbone was frozen. This constraint was used to preserve the secondary structure while allowing the ligand and TMH side chains to relax. In addition, loop residues were frozen until they could be minimized using an appropriate dielectric in the next stage of the minimization. No constraints were placed on the ligands during this stage. The minimization consisted of a conjugate gradient minimization using a distance-dependent dielectric of 2.0, performed in 1000-step increments until the bundle reached the 0.05 kJ/mol gradient. In the second stage of the calculation, the helices and ligand were frozen, but the loops were allowed to relax. The generalized Born/surface area continuum solvation model for water as implemented in Macromodel was used. This stage of the calculation consisted of a Polak-Ribier conjugate gradient minimization in 1000-step increments until the bundle reached the 0.05 kJ/mol gradient.

**Results**

**Chemistry.** The free base forms of LDK1203, LDK1222, and LDK1229 were synthesized according to the scheme illustrated in Fig. 1A, and then purified by CombiFlash chromatography (Teledyne Isco, Lincoln, NE) followed by conversion to hydrochloric salts to increase shelf life and aqueous solubility. Confirmation data for their structure by mass spectroscopy and ¹H NMR are presented in Materials and Methods. The representative first-generation inverse agonists rimonabant, otenabant, taranabant, and ibipinabant are shown for comparison (Fig. 1B).

**LDK1229 and Its Analogs Exhibit Inverse Agonist Binding Profiles.** To elucidate the nature of these new compounds, we performed ligand-binding studies using membrane preparations from HEK293 cells transfected with CB₁. All three compounds competed with CP55,940 and SR141716A, an agonist and inverse agonist of CB₁, respectively. Using [³H]CP55,940 (Kᵣ = 2 nM) as a tracer, LDK1203, LDK1222, and LDK1229 bound wild-type CB₁ with Kᵣ values of 260, 331, and 220 nM, respectively. Similarly, while using [³H]SR141716A as a tracer (Kᵣ = 5 nM), the compounds also bound with comparable affinities (Kᵣ = 297, 366, and 246 nM, respectively; Tables 1–3). The structures of LDK1203 and LDK1222 differed from that of LDK1229 by the substitution groups on the piperazine ring where the piperazine ring of LDK1203 differ from that of LDK1229 by the tosyl group. The free base forms of LDK1203, LDK1222, and LDK1229 were synthesized according to the scheme illustrated in Fig. 1A, and then purified by CombiFlash chromatography (Teledyne Isco, Lincoln, NE) followed by conversion to hydrochloric salts to increase shelf life and aqueous solubility. Confirmation data for their structure by mass spectroscopy and ¹H NMR are presented in Materials and Methods. The representative first-generation inverse agonists rimonabant, otenabant, taranabant, and ibipinabant are shown for comparison (Fig. 1B).

Since LDK1229 exhibited the strongest binding affinity among the compounds, it was further investigated for its capability to bind CB₁ receptors in the active and inactive states to confirm its nature as a CB₁ inverse agonist. The CB₁ receptor in the active and inactive states can be readily mimicked by our previously engineered mutant CB₁ receptor models (T210I and T210A) (D’Antona et al., 2006). CB₁ with the T210I mutation was shown to adopt a fully active form in comparison with the CB₁ wild type. In contrast, CB₁ with the T210A mutation adopts an inactive state. A GPCR in its active state displays enhanced affinity toward agonists but decreased affinity toward inverse agonists, whereas the inactive state of the receptor exhibits the opposite binding properties such that inverse agonists show higher binding affinity than agonists (Cotecchia et al., 1990; McWhinnie et al., 2000; Wade et al., 2001; D’Antona et al., 2006). Hence, our mutant CB₁ receptor models provide ideal tools to evaluate the inverse agonism of LDK1229.
As expected for an inverse agonist, using [3H]SR141716A as the tracer, the binding affinity of LDK1229 to the inactive T210A receptor was enhanced (Kᵢ = 68 nM) relative to its affinity to the wild-type CB₁ receptor (Kᵢ = 246 nM; Table 2). A comparable trend is observed with the inverse agonist SR141716A, which binds T210A with the highest affinity (Kᵢ = 1.47 nM) and a progressively weaker affinity for the wild-type CB₁ (Kᵢ = 5.23 nM) and then the T210I receptor (Kᵢ = 14.7 nM). The ratio of binding affinity for the wild-type receptor relative to the mutant receptor for LDK1229 and SR141716A also reflects the preference of these two compounds for the inactive T210A receptor (Table 2).

Using [3H]CP55,940 as the tracer and examining binding of the agonist CP55,940, the pattern is reversed (Table 3). CP55,940 binds T210A most weakly and then progressively exhibits enhanced affinity for the wild-type CB₁ and the T210I receptors. This is consistent with its agonist properties. In contrast, LDK1229 binds CB₁ wild type more tightly than the T210I receptor, which is consistent with inverse agonist binding profiles (D’Antona et al., 2006). For the constitutively active T210I receptor, a large decrease in the binding affinity of LDK1229 compared with the binding affinity of the agonist CP55,940 (5831 nM versus 0.835 nM) is observed. The ratio of binding affinity for the wild-type receptor relative to the mutant receptor for LDK1229 and CP55,940 is also given for comparison (Table 3). At the extremes, when using [3H]CP55,940 as a tracer, we cannot detect any specific binding with T210A, nor can we detect any specific binding with T210I up to 32 μM of LDK1229 when using [3H]SR141716A as a tracer. Taken together, these data suggest that LDK1229 increased binding affinity to T210A over wild type, and its decreased binding affinity to T210I results from the compounds’ inverse agonist mode of action.

**LDK1229 Inhibits Basal and Agonist-Induced G Protein Coupling in a Concentration-Dependent Manner.**

Given the inverse agonist properties observed in the binding profile of LDK1229 to the wild-type and mutant CB₁ receptors, [35S]GTPγS binding assays were performed in its presence. This assay monitors the level of G protein activation by determining the extent of binding of the nonhydrolyzable GTP analog to Gα subunits. We investigated the effects of LDK1229 on the basal G protein coupling activity levels of the wild-type CB₁ receptor (Fig. 2A). Interestingly, using 1 μM of LDK1229 in the absence of CP55,940 substantially reduced the basal level of [35S]GTPγS binding from 110 to 70 fmol/mg.

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>LDK1203 and LDK1222 binding to the wild-type CB₁ receptor</th>
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<tr>
<td><strong>Kᵢ</strong></td>
<td>nM</td>
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<tr>
<td><strong>Versus [3H]CP55,940</strong></td>
<td></td>
</tr>
<tr>
<td>CP55,940</td>
<td>2.56 (1.45–4.51)</td>
</tr>
<tr>
<td>LDK1203</td>
<td>260 (80.7–837)</td>
</tr>
<tr>
<td>LDK1222</td>
<td>331 (177–619)</td>
</tr>
<tr>
<td><strong>Versus [3H]SR141716A</strong></td>
<td></td>
</tr>
<tr>
<td>SR141716A</td>
<td>5.23 (3.29–8.32)</td>
</tr>
<tr>
<td>LDK1203</td>
<td>297 (192–458)</td>
</tr>
<tr>
<td>LDK1222</td>
<td>366 (144–928)</td>
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</table>

*Data are the median and corresponding 95% confidence limits of three independent experiments performed in duplicate. Kᵢ values were determined from competition binding assays using [3H]CP55,940 or [3H]SR141716A as tracers at their respective Kᵢ values, as described in Results.

*Binding values are from Ahn et al. (2009) for comparison.

We also evaluated its impact on agonist CP55,940-induced [35S]GTPγS binding in the presence of various concentrations of LDK1229. We observed a progressive decrease in the specific GTPγS binding, with increasing concentration of LDK1229 up to 32 μM for the CB₁ wild-type receptor (Fig. 2B).

Like LDK1229, LDK1203 produced an antagonizing effect on the basal and agonist-induced levels of [35S]GTPγS and is included for comparison (Fig. 2). The antagonizing effect is consistent with the properties of an inverse agonist that promotes the inactive form of the receptor and suggests that this compound inhibits G protein coupling.

**LDK1229 Promotes Cell Surface Expression of the CB₁ Receptor.** Upon prolonged exposure to an agonist, GPCRs become desensitized and subsequently internalize in the cell. Inverse agonists, however, act in the opposite manner by promoting GPCR localization to the cell surface (Rinaldi-Carmona et al., 1998b; Marion et al., 2004). To assess the effect of LDK1229 on CB₁ cellular localization, we determined the cellular response of CB₁ upon LDK1229 treatment using confocal microscopy of cells expressing GFP-tagged CB₁ receptors. Previous observations (Leterrier et al., 2004; Martini et al., 2007; Rozenfeld and Devi, 2008) indicate that CB₁ is partially constitutive and that much of the wild-type CB₁ receptor (≈85%) is localized on intracellular vesicles. Using the CB₁-GFP chimera expressed in HEK293 cells and treated with vehicle alone (0.03% dimethylsulfoxide), we found that about 15% of CB₁ is present at the plasma membrane but that the pattern of receptor fluorescence within the cells was intracellularly punctate, suggesting that a substantial proportion of receptors is internalized, which is consistent with its basal activity (Fig. 3). Treatment with 0.1 μM of the agonist CP55,940 resulted in a further shift toward internalization and a punctate appearance of the CB₁-GFP receptor within the cells (data not shown). In contrast, treatment with 1 μM SR141716A or 10 μM LDK1229 resulted in approximately 20% cell surface localization evident within 3 hours and an increase to about 65% of cell surface localization after 5 hours (Fig. 3B). This is strikingly different from the cell surface localization of 13% for basal cells at 3 hours and an increase to about 65% of cell surface localization after 5 hours (Fig. 3B). Taken together, these data further support the inverse agonist activity of LDK1229.

**LDK1229 Docked in the CB₁ Receptor (Inactive State) Model.** Since LDK1229 exhibited the strongest binding affinity among the compounds, Glide docking studies were performed and suggest that LDK1229 binds in the TMH3-4-5-6 region of CB₁. This is the same region that forms
the binding site for SR141716A at CB1. Figure 4, A and C illustrates SR141716A (shown in cyan) docked in the CB1 inactive state model. The hydrogen bonding interaction is consistent with our prior CB1 mutant cycle studies, which indicated that the amide oxygen of SR141716A interacts directly with K3.28192 (McAllister et al., 2004). This interaction is critical for the inverse agonist properties of SR141716A (Hurst et al., 2002, 2006).

At its binding site, SR141716A also forms several aromatic-stacking interactions. First, the SR141716A dichlorophenyl ring forms aromatic T-stacks with F3.36200 and W5.43279 (shown in orange; see Fig. 4A). In addition, the SR141716A chlorophenyl ring forms offset parallel aromatic stacks with Y5.39275 and W5.43279 (shown in orange). In addition, SR141716A forms hydrophobic interactions with W6.48356 (shown in lavender). These aromatic interactions are consistent with our prior mutagenesis studies (McAllister et al., 2003, 2004), which indicated that SR141716A interacts within an aromatic microdomain in CB1, which is comprised of F3.36200, Y5.39275, W5.43279, and W6.48356. Finally, SR141716A forms hydrophobic interactions with L3.29193, V3.32196, and L6.51359 (shown in lime). The geometry of this hydrogen bonding interaction is instrumental to the maintenance of the CB1-inactive state (McAllister et al., 2004). The binding of SR141716A stabilizes the F3.36200/W6.48356 aromatic stacking interaction through the formation of an extensive network of aromatic stacks within the ligand-receptor complex (Shim et al., 2012).

Figure 4, B and D illustrates the final docked conformation of LDK1229 (shown in lavender) in the CB1 inactive state model (Supplemental Material). Like SR141716A, the LDK1229 amide oxygen forms a hydrogen bond with K3.28192 (shown in yellow). The geometry of this hydrogen bond is not optimal, however, because the hydrogen bond distance is longer than that found for SR141716A (2.8 Å versus 2.6 Å) and because the hydrogen bond angle deviates more from linearity than that formed with SR141716A (157° versus 171°). This suggests that the interaction between LDK1229 and K3.28192 is weaker than the interaction between SR141716A and K3.28192. The importance of this difference between SR141716A and LDK1229 is discussed in more detail below.

Like SR141716A, LDK1229 also docks in the CB1 TMH3-4-5-6 aromatic microdomain, but the geometry of its interactions differ somewhat from SR141716A. The fluorophenyl ring of LDK1229 forms aromatic T-stacking interactions with W6.48356 and W5.43279 (shown in orange; see Fig. 3B). In addition, the other fluorophenyl ring of LDK1229 forms offset parallel aromatic stacking interactions with Y5.39275 and W5.43279 (shown in orange). However, LDK1229 does not form an aromatic stacking interaction with F3.36200. This is due to a difference in how the two compounds are oriented within the receptor. Finally, LDK1229 forms several hydrophobic interactions with residues F3.36200 (shown in orange), L3.29193, V3.32196, and L6.51359 (shown in lime).
Mutational Analysis of Computationally Predicted Residues in the LDK1229 Binding Pocket. To test the LDK1229 binding-site hypothesis generated by the receptor model, K3.28192A, W5.43279A, W6.48356A, and C7.42386M mutant CB1 receptors were generated. Our results showed that the K3.28192A mutation resulted in total ablation of [3H]CP55,940 binding at wild-type CB1 and that this residue is critical for CP55,940 binding consistent with previous data (Song and Bonner, 1996). Therefore, competition binding assays on the K3.28192A mutant were performed using [3H]WIN55,212-2 as a tracer since [3H]WIN55,212-2 binding to the K3.28192A mutant was not significantly different than its binding to the wild-type receptor ($K_d = 8.6$ nM versus 4.5 nM). Competition binding analysis revealed that the binding of LDK1229 ($K_i = 1316$ nM) was affected 6-fold by this mutation compared with wild type ($K_i = 324$ nM; Table 4), suggesting that this residue may be important for a hydrogen bonding interaction with LDK1229. These results alone cannot tell us the specific region on LDK1229 with which K3.28192 interacts, but modeling studies reported here suggest that K3.28192 hydrogen bonds with the carboxyl oxygen of LDK1229. The K3.28192A mutation results reported here are consistent with previous CB1 K3.28192A mutation studies, which showed that K3.28192A is an important interaction site (Hurst et al., 2002) for the carboxamide oxygen of SR141716A (Hurst et al., 2006).

Saturation (equilibrium) binding analysis for [3H]CP55,940 at the other three mutant receptors, W5.43279A, W6.48356A, and C7.42386M, showed that the affinity parameters obtained for the mutant receptors were not significantly different from the $K_d$ at the wild-type CB1 receptor ($K_d$ for mutants = 6, 4, and 3 nM, respectively), suggesting that these receptors were properly folded. Therefore, competition binding assays on these mutants were performed using [3H]CP55,940 as a tracer. The W5.43279A mutation had the most profound effect on LDK1229 binding compared with any of the other mutations and showed only 39% displacement of [3H]CP55,940 when using 32 μM of LDK1229. This result is consistent with the modeling studies that suggest that W5.43279A is central in the formation of the aromatic T-stacking interactions with the two fluorophenyl rings in LDK1229. The W6.48356A mutant affected LDK1229 binding 9-fold ($K_i = 1987$ nM) and the C7.42386M mutant 5-fold ($K_i = 1010$ nM; Table 4). The result for the W6.48356A mutation further supports the modeling results, which suggest that this residue participates in direct aromatic stacking interactions.

Fig. 3. Effect of LDK1229 on internalization of the CB1 receptor. (A) HEK293 cells expressing the CB1-GFP receptor were incubated with vehicle alone (0.03% dimethylsulfoxide [DMSO]), 0.5 μM SR141716A (SR), or 10 μM LDK1229. SR141716A is shown for comparison. (B) Quantification of CB1 receptors on the cell surface. The Quantitative Imaging of Membrane Proteins software with plug-ins was used as described in Materials and Methods. Images are representative of at least four independent transfections. The scale bar is 5 μm.

Fig. 4. Model of LDK1229 or SR141716A docked in the inactive CB1 receptor model. (A and C) illustrate SR141716A (shown in cyan) docked in the CB1 inactive state model. (B and D) illustrate the final docked conformation of LDK1229 (shown in lavender) in the CB1 inactive state model. Residues that form hydrogen bonds are shown in yellow, aromatic microdomain residues are shown in orange, and residues important for hydrophobic interactions are shown in lime. A PDB homology model illustrating the final docked conformation of LDK1229 in the CB1 inactive state model can be found in the Supplemental Data.
with one of the fluorophenyl rings of LDK1229, consistent with prior studies (McAllister et al., 2003). As shown in Table 4, we found here that enlarging the residue at position 7.42 via the C7.42386M mutation results in a 5-fold loss in affinity for LDK1229, which is consistent with previous studies by Farrrens and colleagues (Fay et al., 2005). Taken together, the results of all mutation studies are consistent with the modeling results reported here, specifically that LDK1229 occupies the same general binding region as SR141716A.

**LDK1229 Has Diminished Activity for CB2**. Based on both binding and functional data, LDK1229 showed some selectivity for the CB1 receptor over the CB2 receptor (Fig. 5). This is evident by LDK1229 reducing the binding affinity 3-fold (Kᵢ = 633 nM; Fig. 5A) to the CB2 receptor compared with the CB1 receptor (Kᵢ = 220 nM; Table 3) using [³H]CP55,940 as a tracer. This selectivity is also evident by the small magnitude of the reduction in basal and agonist-induced G protein coupling using CB2. Using 1 μM of LDK1229 in the absence of CP55,940 only reduced the basal level of [³H]GTPγS binding from 95 to 88 fmol/mg (Fig. 5B) compared with the 2-fold decrease observed with CB1 (Fig. 2A). We also observed a small decrease in the CP55,940-induced [³H]GTPγS binding with an increasing concentration of LDK1229 (Fig. 4B), with up to 32 μM of LDK1229 (to 110 fmol/mg from the original 120 fmol/mg). The CB2 selective inverse agonist N-[(LS)-endo-1,3,3-trimethylcycloheptan-2-yl]-5-(4-chloro-3-methylphenyl)-1-(4-methylbenzyl)-pyrazole-3-carboxamide (SR144528) (Rinaldi-Carmona et al., 1998a) antagonized the basal level of [³H]GTPγS binding and is shown for comparison. We also found that LDK1229 can dock in the CB2 inactive state model but in a higher energy conformation (data not shown). This is likely the origin of the 3-fold loss of affinity at CB2.

**Discussion**

In an effort to develop new modulators of the CB1 receptor, we synthesized a group of benzhydryl piperazine analogs, including the compounds LDK1203, LDK1222, and LDK1229, and describe their inverse agonist properties in this study. In addition to their inverse agonist binding profiles to the CB1 receptor and their preference to bind the inactive T210A CB1 receptor over the constitutively active wild-type CB1 or fully active T210I receptor, the inverse agonism exhibited by LDK1229 was also evident from its antagonistic effect on basal and agonist-induced G protein coupling and its ability to increase the CB1 localization to the cell surface. LDK1229 exhibited a lower affinity for the CB2 receptor, with a 3-fold relative selectivity for the CB1 receptor. Because the CB1 receptor is constitutively active both in vitro and in vivo (Landsman et al., 1997; Meschler et al., 2000), discovering new and improved means for inhibiting the activity of the receptor is therapeutically useful and relevant for modulating activity of the CB1 receptor system in the brain.

Our results show that the benzhydryl piperazine analogs represented by LDK1229 behave as inverse agonists of the CB1 receptor. Structurally, the benzhydryl piperazine analogs are distinct from the first generation of CB1 inverse agonists, which generally possess nitrogen-containing five- or six-member aromatic rings as their central connecting units (Lange and Kruse, 2005; Vemuri et al., 2008; Chorvat, 2013). In contrast, benzhydryl piperazine analogs have a central core of piperazine, which is nonaromatic and possesses basic amino group(s). A pharmacophore model of the first generation of CB1 inverse agonists was proposed (Lange and Kruse, 2005), of which the biaryls (e.g., SR141716A) form favorable aromatic stacking interactions with two subpockets surrounded by residues V5.39277-W4.64250-F5.42278 and residues W5.43278-F3.36200-W6.48256 of CB1. The central core (e.g., pyrazole in SR141716A) then connects to a lipophilic moiety through a hydrogen bond acceptor (e.g., the carbonyl of SR141716A). The hydrogen bond acceptor stabilizes the D6.58266-K3.28192 salt bridge in the inactive state of the CB1 receptor. The lipophilic moiety (e.g., the methylene groups of piperidine ring in SR141716A) fits in a pocket formed by V3.32276-F2.57270-L7.43387 and M7.44384 of the CB1 receptor (Lange and Kruse, 2005). In spite of the structural difference between our benzhydryl piperazine analogs and the first generation of CB1 inverse agonists, docking SR141716A and LDK1229 into the CB1 inactive state model suggest that both compounds bind in similar receptor regions and form similar interactions with the receptor. This is also consistent with the results of competitive displacement assays that suggest LDK1229 displaces SR141716A, implying an overlapping binding site. However, the results of the docking studies suggest one major difference between SR141716A and LDK1229. The geometry of the SR141716A hydrogen bond with K3.28192 is much better than that of LDK1229. Conformational analysis of LDK1229 suggests that it does not have the conformational freedom to adopt a conformation that would allow it to improve its hydrogen bond geometry. In addition, the extra ring hydrogen (cyclohexyl ring versus piperidine ring) introduced in LDK1229 forces the ligand to position itself differently. Together, these effects lead to a less favorable hydrogen bond with K3.28192. This is consistent with the results of the binding experiments that show that SR141716A has a higher affinity for CB1 than does LDK1229. Notably, LDK1229 exhibits an efficacy comparable to SR141716A in antagonizing basal GTPγS binding to the CB1 receptor, although it shows a weaker binding affinity to the receptor (Fig. 2; Tables 2 and 3). One advantage of the benzhydryl piperazine analogs is that the piperazine ring provides two amino groups that can be readily used in further derivatization. This opens up rich opportunities for structural advantages over that of SR141716A.
hydrophilicity. In comparison with the conventional scaffold as well as introducing acidic functional groups or increasing H-bonding capacity, molecular weight, and polar surface area, several strategies to reduce a ligand and other related metabolic syndromes. In general, there are potential therapeutic agents for the treatment of obesity antagonists (Janero, 2012) or peripherally restricted CB1 receptors in human ileum. Evidence has begun to emerge on the involvement of the CB1 inverse agonists and is attractive for the design and optimization to achieve the desired peripheral restriction of CB1 inverse agonists.

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Wrote or contributed to the writing of the manuscript: Mahmoud, Lu, Kendall, Shore, Hurst, Reggio.

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Fig. 5. Binding parameters of LDK1229 and LDK1203 to CB2. (A) Binding of LDK1229 and LDK1203 to CB2 wild-type receptors and (B) their effect on the stimulation of [35S]GTPγS binding. Statistical significance of the differences (*compared with basal and * compared with 0.1 μM CP55,940 alone) was assessed using one-way analysis of variance and Bonferroni’s post hoc test. ***P < 0.001; **P < 0.001.

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