

Identification of the GPR55 Agonist Binding Site Using a Novel Set of High-Potency GPR55 Selective Ligands

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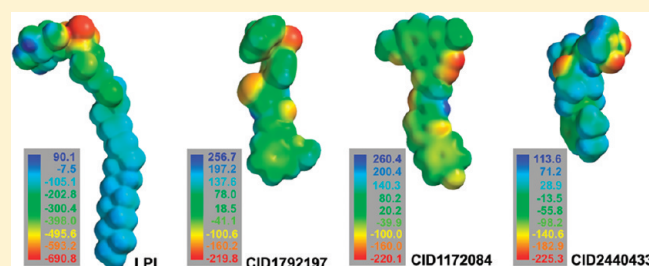
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S Supporting Information

ABSTRACT: Marijuana is the most widely abused illegal drug, and its spectrum of effects suggests that several receptors are responsible for the activity. Two cannabinoid receptor subtypes, CB1 and CB2, have been identified, but the complex pharmacological properties of exogenous cannabinoids and endocannabinoids are not fully explained by their signaling. The orphan receptor GPR55 binds a subset of CB1 and CB2 ligands and has been proposed as a cannabinoid receptor. This designation, however, is controversial as a result of recent studies in which lysophosphatidylinositol (LPI) was identified as a GPR55 agonist. Defining a biological role for GPR55 requires GPR55 selective ligands that have been unavailable. From a β -arrestin, high-throughput, high-content screen of 300000 compounds run in collaboration with the Molecular Libraries Probe Production Centers Network initiative (PubChem AID1965), we identified potent GPR55 selective agonists. By modeling of the GPR55 activated state, we compared the GPR55 binding conformations of three of the novel agonists obtained from the screen, CID1792197, CID1172084, and CID2440433 (PubChem Compound IDs), with that of LPI. Our modeling indicates the molecular shapes and electrostatic potential distributions of these agonists mimic those of LPI; the GPR55 binding site accommodates ligands that have inverted-L or T shapes with long, thin profiles that can fit vertically deep in the receptor binding pocket while their broad head regions occupy a horizontal binding pocket near the GPR55 extracellular loops. Our results will allow the optimization and design of second-generation GPR55 ligands and provide a means for distinguishing GPR55 selective ligands from those interacting with cannabinoid receptors.



GPR55 is a rhodopsin-like (class A) G protein-coupled receptor (GPCR), highly expressed in human striatum,¹ that has the potential to become an important therapeutic target [GenBank accession number NM005683 (see Figure 1)]. Characterizations of GPR55 (–/–) knockout mice^{2,3} reveal a role for GPR55 in inflammatory pain, neuropathic pain, and bone development, while other studies indicate that GPR55 activation is pro-carcinogenic.^{4–6} The GPCR proteins most homologous to GPR55 are GPR35 (27%), P2Y (29%), GPR23 (30%), and CCR4 (23%),¹ but GPR55 also exhibits a low level of amino acid identity to cannabinoid CB1 (13.5%) and CB2 (14.4%) receptors. This homology to cannabinoid receptors and the potential utility of marijuana derivatives in therapy prompted a search for GPR55 ligands among known cannabinoid receptor compounds.

Thus, some synthetic and natural cannabinoids were among the first chemicals recognized to bind GPR55.⁷

Subsequent studies of GPR55 pharmacology suggested that lysophosphatidylinositol (LPI, **1**) compounds are endogenous GPR55 agonists,⁸ with 2-AGPI, a 2-arachidonoyl-containing LPI species, possessing significantly higher potency and the maximal efficacy of the LPI species observed to date.⁹ 2-AGPI may represent the true natural ligand of GPR55. In this work, we reference efficacy relative to the commercially available LPI (**1**). Using a β -arrestin green fluorescent protein (beta arrestin-GFP)

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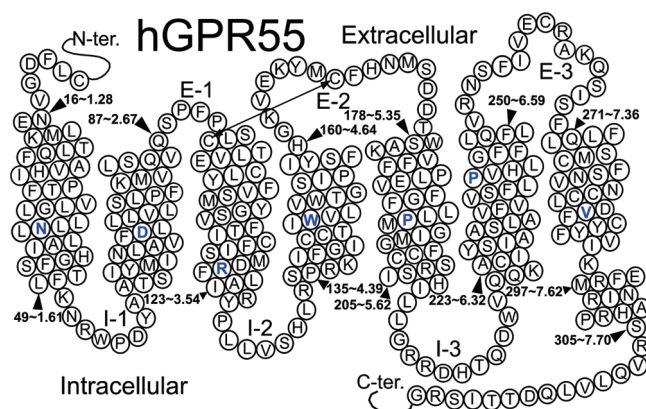


Figure 1. Helix net representation of the human GPR55 sequence.

biosensor to assess a cohort of CB1 and CB2 ligands for GPR55 activity, Kapur et al.¹⁰ confirmed LPI as a GPR55 agonist, while observing that the cannabinoid antagonists AM251 and SR141716A were also GPR55 agonists. These GPR55 ligands possess comparable efficacy in inducing β -arrestin trafficking and, moreover, activate the G protein-dependent signaling of PKC β II. Conversely, the potent synthetic cannabinoid agonist CP55940 acts as a GPR55 antagonist/partial agonist, inhibiting GPR55 internalization, the formation of β -arrestin GPR55 complexes, and the phosphorylation of ERK1/2.¹⁰

More potent and selective GPR55 ligands have been unavailable predominantly because many of the known GPR55 ligands were identified from cannabinoid receptor/lipid-biased compound libraries.^{11,12} The discovery and molecular characterization of better GPR55 chemotypes are important next steps in the design of higher-quality GPR55 ligands that can serve either as research tools or as a basis for designing novel drugs. During a collaborative project between our individual laboratories and the Sanford-Burnham screening center of the Molecular Libraries Probe Production Centers Network (MLPCN), we identified a series of GPR55 agonists that belong to novel, unreported GPR55 agonist chemotypes. These were discovered by a high-content, high-throughput β -arrestin screen (see <http://mli.nih.gov/mli/mlp-probes/>). In this work, we focus on three GPR55 agonists identified in this screen: CID1792197 (2), CID1172084 (ML185, 3), and CID2440433 (ML184, 4) (see Figure 2 for structures).

Only a few class A GPCRs have been crystallized to date. These include rhodopsin (Rho),^{13–15} the β_2 -adrenergic receptor (β_2 -AR),^{16–18} the β_1 -adrenergic receptor (β_1 -AR),¹⁹ the adenosine A2A receptor,²⁰ and, most recently, the CXCR4²¹ and dopamine D3²² receptors. These crystal structures reveal a common topology that includes (1) an extracellular N-terminus, (2) seven transmembrane α -helices (TMHs) arranged to form a closed bundle, (3) loops connecting TMHs that extend intra- and extracellularly, and (4) an intracellular C-terminus that begins with a short helical segment (helix 8) oriented parallel to the membrane surface. Although no crystal structure is available for GPR55, it is possible to build a homology model of GPR55 based on available crystal structures and refine this model on the basis of sequence-dictated differences between GPR55 and the template crystal structure. This refined model can then be used to explore the GPR55 ligand binding pocket and key amino acid interactions for GPR55 ligands. We report here the development of a refined homology model of the

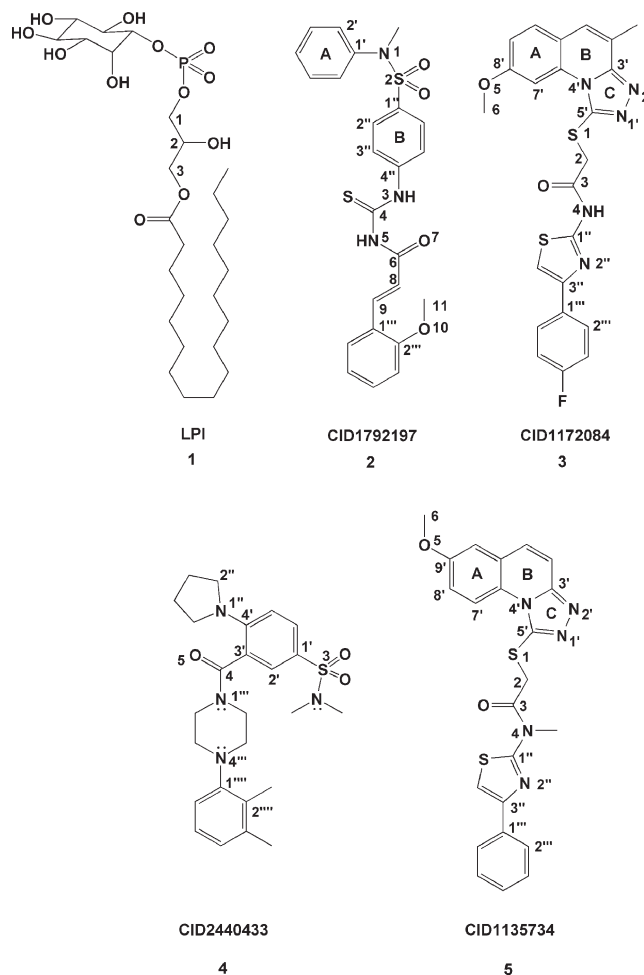


Figure 2. Structures of GPR55 agonists (1–4) and the control compound (5).

GPR55 activated state (GPR55 R*) and its subsequent use in exploring the interaction between GPR55 and the novel CID agonists (2–4) identified in our screen. Our modeling data indicate that the similarity between the CID compounds and LPI allows them all to be recognized by a single GPR55 binding pocket. The chemical diversity provided by these three lead compounds, CID (2–4), combined with the identification of key receptor interaction sites should provide a basis for the design of more potent and efficacious second-generation GPR55 ligands that retain GPR55 selectivity.

METHODS

Chemical Library Screening. Potent GPR55 selective agonists were identified using a β -arrestin (see below), high-throughput, high-content screen of 300000 compounds run in collaboration with the Molecular Libraries Probe Production Centers Network program. For more details about this library of compounds, see <http://mli.nih.gov/mli/compound-repository/mlsmr-compounds/>. Compounds were screened for agonism (PubChem AIDs 2347) at GPR55 and for both agonism and antagonism at GPR35 (PubChem AIDs 2397 and 2338), CB1 (PubChem AIDs 2341 and 2339), and CB2 (PubChem AIDs 434928 and 434922). The high-content imaging assay utilized in each case employs a cell line permanently expressing a β -arrestin

GFP biosensor and an enhanced receptor of interest (i.e., GPR55, GPR35, CB1, or CB2). A full description of each of these assay protocols (according to AID number) is accessible at the PubChem website (<http://pubchem.ncbi.nlm.nih.gov/>). Potent GPR55 agonists lacking agonism or antagonism at GPR35, CB1, or CB2 were further evaluated for pERK activation and PKC β II translocation (see below). A set of novel GPR55 agonist molecular scaffolds were selected from the screen [CID1792197 (2), CID1172084 (3), and CID2440433 (4)], including a chemically related inactive compound [CID1135734 (5)], and the binding of each compound was explored using a computer model of the GPR55 activated state (see below).

Biochemical Studies. *β -Arrestin Assessment of GPR55 Activation.* U2OS cells permanently expressing HA-GPR55E and β arr2-GFP have previously been described.¹⁰ Cells were plated onto 384-well assay plates (Greiner) at 80–85% confluency and maintained at 37 °C in 5% CO₂ overnight. Prior to drug application, serum-containing medium was removed and replaced with serum-free MEM. Agonist-stimulated redistribution of β arr2-GFP was assessed following drug treatment for 75 min at 37 °C in 5% CO₂. Cells were then fixed with 4% paraformaldehyde for 40 min at room temperature followed by two washes with PBS. The nuclear stain, DAPI (Invitrogen), was added, and plates were sealed for imaging. Image acquisition was performed on an Opera QEHS instrument (Perkin-Elmer) using the following settings: 20 \times 0.45 NA air objective, camera set to 2 \times 2 binning for an image size of 688 \times 512 pixels, two channels acquired sequentially, 488 nm excitation for β -arrestin-GFP and 365 nm excitation for DAPI (details in PubChem AID 1965).

Data Analysis. β arr2-GFP aggregates were identified using the Acapella spot detection algorithm (Perkin-Elmer). Briefly, the analysis protocol extracts from two-dimensional images those pixels that generate objects of interest that fall within a predetermined range of sizes and intensities and that are embedded among widely varying local backgrounds. The primary assay read-out was based on the number of detected fluorescent spots representing pits or vesicles per cell. Concentration–effect curves for agonist-mediated receptor activation were analyzed by nonlinear regression techniques, and data were fitted to sigmoidal dose–response curves to obtain EC₅₀ values. Activity values were normalized to the positive controls [LPI (1) as 100% at 10 μ M] and negative controls (vehicle only as 0%) for each well plate. Statistical analysis was performed using one-way analysis of variance (ANOVA) followed by a Dunnett's post-test or a two-tailed unpaired Student's *t* test, and *p* values of <0.05 were considered significant.

Western Blot Analysis of ERK Activity. GPR55E-expressing U2OS cells were grown to subconfluence in 60 mm plates and serum-starved overnight before being assayed. After drug treatment, the cells were disrupted in lysis buffer [50 mM Hepes, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 10% glycerol, 1% Triton X-100, 10 μ M MgCl₂, 20 mM *p*-nitrophenyl phosphate, 1 mM Na₃VO₄, 25 mM NaF, and a protease inhibitor mixture (1:25, pH 7.5)]. Lysates were immediately placed on ice for 10 min and then centrifuged at 16000g for 30 min at 4 °C. Supernatants, corresponding to the cytosolic fraction, were collected, and protein concentrations were determined by the Bradford assay (Bio-Rad) using bovine serum albumin as a standard. Cytosolic fractions (20 μ g) were separated on a 10% gel by sodium dodecyl sulfate–polyacrylamide gel electrophoresis followed by immunoblotting.²³ Antibodies against doubly phosphorylated ERK1/2

(1:5000) were detected using a LI-COR Odyssey IR Imager (LI-COR Biotechnologies, Lincoln, NE). A polyclonal antibody against total ERK1/2 (Cell Signalling 9102, 1:1000) was used to confirm equal protein loading. Densitometric analysis was performed using LI-COR Odyssey IR Imager. The value obtained for both ERK1 and ERK2 was normalized to total ERK1/2 levels. The data were normalized to the control and presented as percentage stimulation.

PKC β II Assay of GPR55 Activation. HEK 293 cells plated in 35 mm glass well Matek plastic dishes were transiently transfected with 175 μ L of a solution containing 1.5 μ g/mL PKC β II-GFP cDNA or the PKC plasmid and 5 μ g/mL human GPR55 cDNA in pCMV-Sport6 (Open Biosystems, Huntsville, AL) using a standard calcium phosphate protocol. Cells expressing GPR55 and PKC β II-GFP were utilized 24 h after transfection. Cells were washed with warm MEM and maintained at 37 °C in 5% CO₂ for 30–45 min after drug application. Agonist-stimulated redistribution of PKC β II-GFP was assessed after drug treatment at room temperature.

Receptor Model Development. *Modeling of GPR55 Using GPCR Crystallization Data.* A model of the activated form of GPR55 was created using the 2.4 Å crystal structure of β_2 -AR.¹⁶ First, the GPR55 sequence was aligned with the sequences of β_2 -AR, Rho, the A2 adenosine receptor, and the CB1 and CB2 receptors. The same highly conserved residues that have been used in the past to align CB1^{24–29} and CB2^{30,31} receptors with the sequence of bovine Rho were used as alignment guides for the GPR55 sequence. GPR55 possesses most of the conserved class A patterns in TMH1, -2, -4, and -5 (N1.50, D2.50, W4.50, and P5.50, respectively). There are a few notable conserved motif differences between the GPR55 sequence and the sequence of not only the cannabinoid receptors but also the crystallized GPCRs. (1) GPR55 has a conservative substitution (DRF) for the TMH3 E/DRY motif. (2) It has a conservative substitution (SFLP) for the TMH6 CWXP motif. (3) It has a nonconservative substitution (DVFCY) for the THM7 NPXXY motif. Finally, the GPR55 extracellular-1 (EC-1) loop is shorter than most [three amino acids (aa) vs 6 aa in β_2 -AR and Rho], and the GPR55 EC-3 loop is noticeably longer than most (14 aa long vs 5 aa in β_2 -AR and 6 aa in Rho, CB1, and CB2).

Conformational Memories (CM) Technique for Calculating GPCR Structure. We explored the possible sequence-dictated conformational deviations of GPR55 TMH1, -2, -5, -6, and -7 from the β_2 -AR template using the CM technique.^{32–34} The CM method uses multiple Monte Carlo/simulated annealing random walks employing the CHARMM force field. Backbone ϕ and ψ torsions in regions of interest were allowed to vary $\pm 50^\circ$, while all other backbone torsion angles were allowed to vary $\pm 10^\circ$. Side chain torsions were allowed to vary $\pm 180^\circ$. All bond angles were allowed to vary $\pm 8^\circ$ except for the C–S–C angles that were allowed to vary $\pm 15^\circ$. A minimum of 105 conformers was generated for each GPR55 helix independently in a distance-dependent dielectric at 310 K. Output helices were analyzed using ProKink.³⁵ This program, which is embedded in Simulaid Conversion,³⁶ was used to calculate the face shift, wobble angle, and bend angle of each helix.

TMH1 in GPR55 has a Pro at position 1.41 not found in other sequences. An ideal helix ($\phi = -62.9^\circ$, and $\psi = -41.6^\circ$) with the GPR55 sequence was built, and the region around the Pro (A1.37–P1.41) was varied using CM.

TMH2 in GPR55 has a Pro at position 2.58 instead of at position 2.59 as in the β_2 -AR. β_2 -AR TMH2 was mutated to the

GPR55 sequence, and residues V2.54–P2.58 were considered the variable region. S2.56 near P2.58 was set to be in g minus χ_1 .

TMH5 in GPR55 has two prolines, the conserved P5.50 that both β_2 -AR and Rho have and an additional one at position 5.41. β_2 -AR TMH5 was mutated to the GPR55 sequence. Only the region from K5.37 to P5.41 was varied because β_2 -AR TMH5 already has a proline kink at P5.50.

TMH6 in GPR55 does not have the highly conserved class A CWXP motif. Instead, it has the conservative SFXP substitution. Because TMH6 is implicated in conformational changes during receptor activation,^{37–42} it was very important to explore the conformational space of GPR55 TMH6. X-ray crystal structures of Rho and others have shown that an “ionic lock” salt bridge is formed between R3.50 near the intracellular (IC) end of TMH3 and D/E6.30 at the IC end of TMH6. This salt bridge is facilitated by the proline kink in TMH6 that bends the IC end of TMH6 toward TMH3. Biophysical studies have indicated that this salt bridge breaks upon activation and TMH6 straightens.⁴⁰ An ideal helix ($\varphi = -62.9^\circ$, and $\psi = -41.6^\circ$) with the GPR55 TMH6 sequence was built, and the region around the Pro (V6.46–P6.50) was varied using CM while the S6.47 χ_1 was held in the trans position (see the Supporting Information for further details).

TMH7 in GPR55 lacks the highly conserved NPXXY motif of class A GPCRs and instead has a DVXXY sequence. The NPXXY motif not only influences the conformation of TMH7 but also places Y7.53 in the correct position to interact with F7.60 on the intracellular extension of TMH7, helix 8 (Hx8). For receptors that possess an NPXXY motif, we typically designate the 7.46–7.50 region as variable. For GPR55 TMH7, an ideal helix ($\varphi = -62.9^\circ$, and $\psi = -41.6^\circ$) was built and the region around the Pro (C7.46–V7.50) was varied using CM.

Construction of the TMH Region of GPR55. The outputs of the CM runs were used to build the TMH region of the receptor. TMH1, -2, -5, and -7 were aligned with the intracellular part of the corresponding helices of the β_2 -AR; TMH6 was aligned with the intracellular end of β_2 -AR TMH6 for the inactive form of the receptor and with the extracellular end of β_2 -AR TMH6 for the activated form of the receptor. The GPR55 inactive and activated bundles were then energy minimized to relieve any side chain clashes and to pack the TMHs. The OPLS2005 all atom force field in MacroModel 9.1 (Schrodinger Inc., Portland, OR) was used employing a distance-dependent dielectric with extended cutoffs (8.0 Å for nonbonded, 20.0 Å for electrostatic, and 4.0 Å for hydrogen bonding). A harmonic constraint was placed on all the TMH backbone torsions (φ , ψ , and ω) to preserve the general shape of the helices and let the side chains adjust. The constraint on the TMH backbone torsions was gradually reduced to zero during the course of the minimization.

Modeling of the Intra- and Extracellular Loops and Receptor Termini. Modeler 8.2^{43,44} was used to add the N- and C-termini (Met1–Asn16 and Ser305–Gly319 respectively) and the intra- and extracellular loops (IC1, Leu49–Trp53; IC2, Leu127–Arg136; IC3, Gly210–Gln220; EC1, Ser88–Phe90; EC2, Phe159–Ser178; and EC3, Asp253–Ile266). The final loops were energy minimized to a gradient of 0.01 kcal/mol in a high dielectric using the Generalized Born/Surface Area (GB/SA) continuum solvation model for water, while the TMHs were frozen. The same force field, cutoffs, and conjugate gradient method mentioned above were used.

Molecular Dynamics (MD) Simulation of the GPR55 Activated State (R*) in a Palmitoylphosphatidylcholine

Bilayer. The MD simulation cell for the GPR55 R* model embedded in a bilayer containing 151 palmitoylphosphatidylcholine (POPC) lipid molecules was built as described by Hurst et al.⁴⁵ The membrane was generated *de novo* using a library of lipid conformations from previous neat lipid simulations, as described by Grossfield et al.⁴⁶ The system was neutralized with 10 Cl[−] ions, and NaCl was added to achieve an ionic strength of 0.15 M. Three internal waters were included, two waters flanking Asp2.50 as seen in other crystal structures of GPCRs^{13–20} and one water close to Asp7.49 (usually Asn in other GPCRs) that is surrounded by hydrophobic residues and has no partner. The system was then solvated with 13454 bulk waters and was run in the NPAT ensemble (using an experimental area per lipid value for POPC of 68.3 Å²)⁴⁷ for a total of 70 ns. The NAMD2 simulation package⁴⁸ was used to run the dynamics employing the CHARMM27 force field for proteins and lipids.^{49–51} A plot of the root-mean-square deviation (rmsd) of the TMH bundle versus time was prepared to identify the range of time points for which the TMH bundle geometry has stabilized in the lipid bilayer (plateau region in plot).

Ligand Conformational Analysis of LPI. LPI is a very flexible molecule and can assume a very large number of conformations. CM^{32–34} was used to explore the conformational space of LPI because CM has been shown to achieve complete sampling of the conformational space of very flexible ligands, it overcomes energy barriers efficiently, and it converges in a reasonable number of steps.³² The CM output was then analyzed and separated into groups (clusters) of similar conformations using X-Cluster as implemented in MacroModel.⁵² An increasing rmsd from the first structure of the CM output is used as a criterion for partitioning the output into clusters.

Conformational Analysis of CID Compounds. Complete conformational analyses of the CID compounds were performed using *ab initio* Hartree–Fock calculations at the 6-31G* level as encoded in Jaguar (version 9.0, Schrodinger, LLC, New York, NY). HF 6-31G* 6-fold conformer searches were performed for the rotatable bonds [CID1792197 (2), N1–S2, S2–C1'', C4''–N3, C6–C8, C9–C1''', and C2'''–O10 bonds; CID1172084 (3), C8'–O5, C5'–S1, S1–C2, C2–C3, N4–C1'', and C3''–C1''' bonds; CID2440433 (4), N2–S3, S3–C1', C4'–N1'', C3'–C4, C4–N1'', and N4'''–C1'''' bonds; CID1135734 (5), C9'–O5, C5'–S1, S1–C2, C2–C3, N4–C1'', and C3''–C1'''' bonds]. In each conformer search, local energy minima were identified by rotation of a subject torsion angle through 360° in 60° increments (6-fold search), followed by HF 6-31G* energy minimization of each rotamer generated. To calculate the difference in energy between the global minimum energy conformer of each compound 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.

Electrostatic Potential Map Calculation. The electrostatic potential density surface was calculated using Spartan '08 (Wave function, Inc., Irvine, CA). The electrostatic potential energy was calculated using the Hartree–Fock method at the 6-31G* level of theory and was mapped on the 0.002 isodensity surface of each molecule. The surface was color-coded according to the potential, with electron rich regions colored red and electron poor regions colored blue.

Docking of Ligands. A low-free energy conformer of LPI generated by CM and the lowest-energy conformation of each

Table 1. High-Throughput β -Arrestin Screen Results

PubChem CID	compound	agonist EC ₅₀ (μ M)	agonist assay EC ₅₀ (μ M)				antagonist assay IC ₅₀ (μ M)		
		GPR55	GPR35	CB1	CB2		GPR35	CB1	CB2
1792197	2	0.11/0.12 (avg 0.11)	>32	>32	>32		>32	>32	>32
1172084	3	0.15/0.18 (avg 0.16)	>32	>32	>32		>32	>32	>32
2440433	4	0.25/0.27 (avg 0.26)	>32	>32	>32		>32	21.8	15.1
1135734	5	>32	>32	>32	>32		>32	>32	>32

CID compound were used as inputs for receptor docking studies. Initially, each compound was manually docked in the binding site of the GPR55 R* model that had been equilibrated in POPC (see above). The automatic docking program, Glide (Schrodinger Inc.), was then used to explore other possible binding conformations and receptor site interactions. Glide was used to generate a grid based on the centroid of select residues in the binding site (from the manual dock). Extra precision (XP) and flexible docking with ring sampling were selected for the docking setup. K2.60(80), the only positively charged residue in the transmembrane region, was defined as a hydrogen bond donor and was used as the only constraint for the automatic docking of all the compounds.

Ligand–Receptor Complex Minimization. The receptor–ligand complexes were minimized using the OPLS2005 all atom force field in Macromodel 9.1 (Schrodinger Inc.). An 8.0 Å nonbonded cutoff (updated every 10 steps), a 20.0 Å electrostatic cutoff, and a 4.0 Å hydrogen bond cutoff were used in each stage of the calculation. The first stage consisted of 20000 steps of conjugate gradient minimization in 1000-step increments using a distance-dependent dielectric. No harmonic constraints were placed on the ligand or the receptor side chains to optimize the ligand–receptor interaction. A harmonic constraint was placed on all the TMH backbone torsions (φ , ψ , and ω) to preserve the general shape of the helices, with this gradually reduced to zero in the course of the first stage of the minimization. The loops and N- and C-terminal backbone torsions were also constrained with a 250 kcal/mol force during the first stage. In the second stage of the minimization, the Generalized Born/Surface Area (GB/SA) continuum solvation model for water was used as implemented in Macromodel. This stage of the calculation consisted of 20000 steps of Polak–Ribier conjugate gradient minimization in 1000-step increments or until the bundle reached the 0.05 kJ/mol gradient, in which the ligand and TMH bundle were frozen, but the loops and termini were allowed to relax.

RESULTS

Identification and Biochemical Characterization of GPR55 Agonists. The three CID agonists (2–4) (Figure 2) were identified on the basis of the ability of a β -arrestin-GFP reporter to distinguish the active and inactive conformations of GPCRs in general and GPR55 in particular.⁵³ These ligands, (E)-3-(2-methoxyphenyl)-N-({4[methyl(phenyl)sulfamoyl]phenyl} carbamothioyl)prop-2-enamide (CID1792197, 2), N-[4-(4-fluorophenyl)-1,3-thiazol-2-yl]-2-[(8-methoxy-4-methyl[1,2,4]-triazolo[4,3-a]quinolin-1-yl)sulfanyl]acetamide (CID1172084, 3), and 3-[4-(2,3-dimethylphenyl)piperazine-1-carbonyl]-N,N-dimethyl-4-pyrrolidin-1-ylbenzenesulfonamide (CID2440433, 4), were further chosen for computer modeling of the GPR55 binding pocket because of their potencies, EC₅₀ values of 110–250 nM, and >100-fold selectivity for GPR55 over GPR35, CB1,

and CB2 (see Table 1) in the β -arrestin translocation assay in U2OS cells permanently expressing HA-GPR55E and β arr2-GFP.¹⁰ Figure 3A illustrates concentration response curves for agonist activity in the β -arrestin trafficking assay. CID1792197 (2), CID1172084 (ML185, 3), and CID244043 (ML184, 4) induced trafficking with EC₅₀ values of 0.11 \pm 0.01, 0.16 \pm 0.02, and 0.26 \pm 0.01 μ M, respectively [mean \pm standard error of the mean (SEM); n = 2]. Activity values were normalized to LPI (1) as 100% at 10 μ M. No trafficking was observed with CID1135735 (5). The dose–response curves in Figure 3A reach 100% activation as compared to LPI and then decrease to a slightly lower plateau. This is likely due to a combination of the biology of the pit/vesicle formation with the image analysis assay read-out. The primary assay read-out is based on the well average of the “Number of Spots Detected per Cell” parameter. As higher concentrations of the compounds are added, the pits either cannot be resolved properly and multiple pits may be detected as one larger pit or the pits merge into larger vesicles, resulting in an apparent decrease in activity at higher concentrations. See Figure S-1 of the Supporting Information for an illustration of this point (zooming in to >250% is necessary to see the smaller dimer spots well). In addition, it is possible that compound solubility problems at higher concentrations may also be contributing to this decrease to a lower plateau.

Figure 3B presents representative images showing β -arrestin translocation in GPR55 (seen as a punctate pattern in cell interiors) produced by application of LPI (10 μ M) versus CID1172084 (3) (1 μ M) compared to the lack of translocation seen for the DMSO control. It is clear here that both LPI and CID1172084 (3) induce translocation in HA-GPR55E-expressing U2OS cells compared to the control (HA-GPR55E-expressing U2OS cells treated with DMSO) [see Table S-1 of the Supporting Information for images for all compounds (1–5)].

Treatment of HEK or U2OS cells expressing GPR55 with the CID (2–4) compounds or LPI (1) also activates at least two other intracellular signaling pathways common to agonists of many GPCRs, pERK activation and translocation of PKC β II to the plasma membrane. Figure 3C illustrates that CID compounds elicit phosphorylation of ERK1/2. U2OS cells expressing GPR55E were incubated with 1 μ M CID2440433, 1 μ M CID1792197, 1 μ M CID1172084, and 10 μ M LPI. ERK1 and -2 were monitored and normalized to LPI (10 μ M as 100%). Notably, treatment with either of the CID compounds at 1 μ M exhibited phosphorylation of ERK1/2 that was at least as strong as the previously reported activation upon LPI (10 μ M) treatment. A representative gel is shown, and the mean \pm SEM from three different experiments is shown. Figure 3D presents a table summarizing PKC β II translocation results for application of CID compounds (2–4) to GPR55-expressing cells.

GPR55 Receptor Model Development. A homology model of the GPR55 inactive state was developed using the β_2 -AR crystal

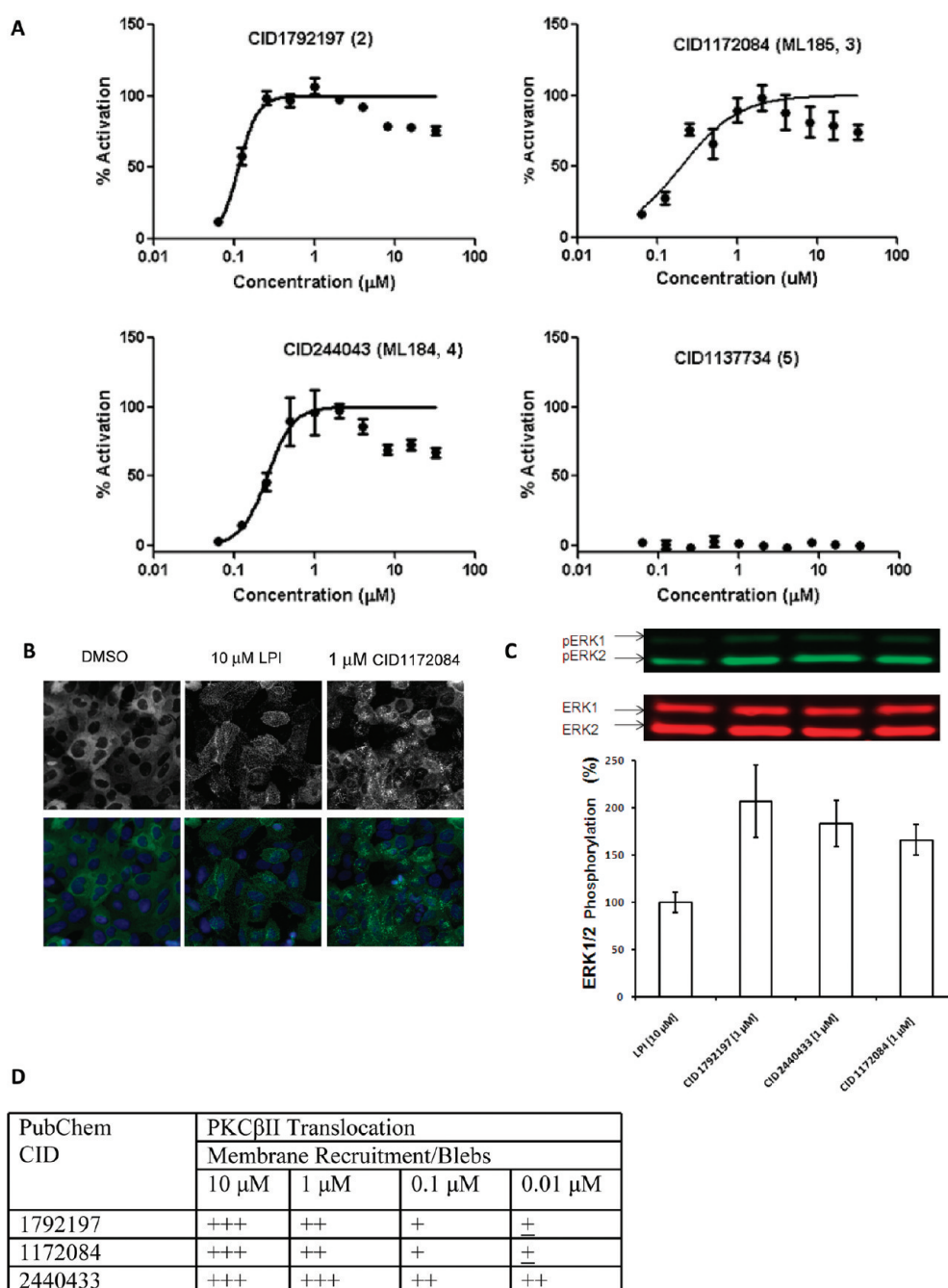


Figure 3. Agonist activity produced by CID compounds. (A) Concentration–response curves for agonist activity in the β -arrestin trafficking assay. CID1792197 (2), CID1172084 (ML185, 3), and CID244043 (ML184, 4) induced trafficking with EC_{50} values of 0.11 ± 0.01 , 0.16 ± 0.02 , and $0.26 \pm 0.01 \mu\text{M}$, respectively (mean \pm SEM; $n = 2$). Activity values were normalized to LPI (1) as 100% at $10 \mu\text{M}$. No trafficking was observed with CID1135735 (5). (B) Representative images showing β -arrestin translocation of LPI vs CID1172084 [see Table S-1 of the Supporting Information for images for all compounds (1–5)]. (C) CID compounds elicit ERK1/2 phosphorylation. U2OS cells expressing GPR55E were incubated with $1 \mu\text{M}$ CID2440433, $1 \mu\text{M}$ CID1792197, $1 \mu\text{M}$ CID1172084, and $10 \mu\text{M}$ LPI. ERK1/2 were monitored and normalized to LPI ($10 \mu\text{M}$ as 100%). Notably, treatment with either of the CID compounds at $1 \mu\text{M}$ revealed ERK1/2 phosphorylation that was at least as strong as the previously reported activation upon LPI ($10 \mu\text{M}$) treatment. A representative gel is shown, and the mean \pm SEM from three different experiments is shown. (D) PKC β II translocation produced by CID compounds: (+) $\sim 1\%$ GFP membrane recruitment or blebs formation, (++) $\sim 10\%$ GFP membrane recruitment or blebs formation, and (+++) $\sim 30\%$ GFP membrane recruitment or blebs formation.

structure [Protein Data Bank (PDB) entry 2RH1¹⁶] as the template. This initial model was refined on the basis of sequence-dictated differences between GPR55 and the β_2 -AR crystal structure. Figure 4 illustrates a comparison of the β_2 -AR crystal structure (Figure 4A) and the GPR55 inactive state model

(Figure 4B). Here the view is from the extracellular side of each receptor. The key differences between these two bundles in the transmembrane helices are described below.

TMH1 in the β_2 -AR pulls away from the TMH bundle on the EC side of the receptor, and hydrophobic residues on TMH2 and

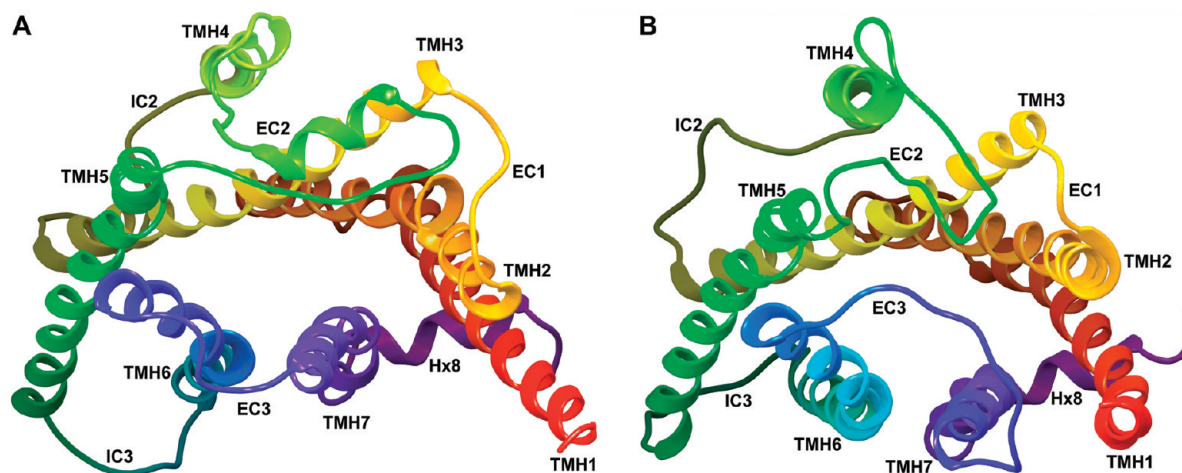


Figure 4. Comparison of (A) the β_2 -AR crystal structure and (B) the GPR55 inactive state model. The view is from the extracellular side of each receptor.

Table 2. ProKink Analysis Results for the Chosen TMH Conformers

helix	hinge residue	bend angle (deg)	wobble angle (deg)	face shift (deg)
TMH1	P1.41	21.1	163.7	43.2
TMH2	P2.58	28.4	−100.1	67.3
TMH5	P5.41	14.6	−160.6	20.9
TMH6 R state	P6.50	32.2	−97.8	54.0
TMH6 R* state	P6.50	25.7	−98.3	68.4
TMH7	V7.50	31.9	173.0	−0.9

TMH7 form a wall that excludes the EC portion of TMH1 from the binding site crevice. CM results show that the lipid-facing residue P1.41 in GPR55 bends the EC end of TMH1 instead toward the binding crevice, filling the space between TMH1/2 and TMH1/7. See Table 2 for helix geometry details.

Because TMH1 in GPR55 bends toward the binding pocket on the EC end, TMH2 must take a different conformation as well to avoid steric overlap with the EC end of TMH1. CM results show that GPR55 has this built into its sequence because GPR55 has a proline at position 2.58 instead of at position 2.59 as in the β_2 -AR, which bends the extracellular portion of TMH2 away from TMH1 and toward TMH3. See Table 2 for helix geometry details. This TMH2 helix exposes residues S2.64(84), L2.63(83), and K2.60(80) to the binding site crevice. It is important to note that had TMH2 retained the β_2 -AR TMH2 conformation, the EC-1 loop would not have been able to join the EC ends of TMH2 and TMH3 because the GPR55 EC-1 loop is only three amino acids long (SPF). In contrast, the EC-1 loop of the β_2 -AR is six amino acids in length (KMWTFG).

Because β_2 -AR TMH5 has the conserved P5.50 as does GPR55 TMH5, the backbone torsions for that Pro kink were preserved and only the region around P5.41 in GPR55 TMH5 was explored via CM. P5.41 bends the extracellular end of TMH5 inward toward the binding crevice and toward the extracellular end of TMH6. See Table 2 for helix geometry details. TMH5 was chosen from the CM output to avoid a clash with the EC end of TMH6.

The CM output gives helices that bend inward toward the binding crevice and also helices that bend toward TMH7. From

the helices generated by CM, we picked two helices; the more bent one was chosen for the inactive GPR55 bundle. This conformer allowed a hydrogen bond to form between R3.50 and Q6.30 at the IC end of the TMH bundle. A second TMH6 conformer was chosen that was straighter that we incorporated in the activated GPR55 model. See Table 2 for helix geometry details. This conformer was chosen so that the R3.50–Q6.30 hydrogen bond was broken. In the β_2 -AR, aromatic residues at positions 6.44 and 6.52 flank the toggle switch residue, W6.48. These residues, along with C6.47, have been proposed to be part of a larger rotamer toggle switch within the binding site crevice of the β_2 -AR.⁵⁴ GPR55 also has aromatic residues flanking F6.48, F6.44, and H6.52, while the residue at position 6.47 is a serine. The CM results showed that there is a correlation between the conformation of F6.48(239) and H6.52(243) (see Table S-2 of the Supporting Information).

GPR55 TMH7 lacks the highly conserved class A GPCR NPXXY motif, having instead a DVFCY motif. The lack of a proline residue in TMH7 resulted in CM output that did not have the usual proline-induced bend and deformation seen in crystal structures of other class A GPCRs. See Table 2 for helix geometry details.

GPR55 Model Activated Form (R*). The hallmark of GPCR activation is the breaking of the ionic lock between R3.50 and E/D6.30 that allows TMH6 to straighten. The result is an intracellular opening of the receptor, exposing residues that can interact with the C-terminus of G α . The active form of the GPR55 receptor model differs from the inactive form mainly in changes at the intracellular side of the receptor. The intracellular end of TMH6 has straightened and moved away from the TMH bundle, breaking the hydrogen bond between R3.50 and Q6.30. The distance between the C α atoms of R3.50 and Q6.30 is 13.3 Å in the GPR55 R* model, indicating a loss of interaction between the IC ends of TMH3 and -6. This distance is comparable to the opening formed in the very recent rhodopsin meta II (activated state) crystal structure (R3.50–E6.30 C α distance of 14.7 Å)⁵⁵ in which the C-terminal fragment of transducin's G α is inserted. The size of this same opening in another recent activated GPCR crystal structure, the nanobody-stabilized β_2 -AR crystal structure, is larger (R3.50 and E6.30 C α distance of 17.2 Å),⁵⁶ because of the physical insertion of the nanobody higher than the C-terminal fragment of transducin's G α that causes the IC opening to grow.

Table 3. Dihedral Angle Definitions of the Minimum Energy Conformers of the CID Compounds

dihedral angle	CID1792197	CID1172084	CID2440433	CID1135734
C4''–N3–C4–N5	–176.6°			
N3–C4–N5–C6	4.2°			
N5–C6–C8–C9	180.0°			
C8–C9–C1'''–C2'''	161.4°			
C3''–C4''–N3–C4	–66.5°			
N1–S2–C1''–C2''	86.2°			
C1'–N1–S2–C1''	70.2°			
C2'–C1'–N1–S2	81.2°			
C5'–S1–C2–C3		–179.8°		
S1–C2–C3–N4		–178.4°		
N4'–C5'–S1–C2		–178.8°		
C3–N4–C1''–N2''		–179.6°		
C7'–C8'–O5–C6		179.9°		
N2''–C3''–C1'''–C2'''		–18.8°		
C2'–C1'–S3–N2			–90.0°	
C1'–S3–N2–LP			–179.7°	
C3'–C4'–N1''–C2''			173.7°	
C2'–C3'–C4–O5			114.1°	
O5–C4–N1'''–LP			87.4°	
LP–N4'''–C1'''–C2'''			28.0°	
C5'–S1–C2–C3				–179.3°
S1–C2–C3–N4				178.2°
N4'–C5'–S1–C2				179.4°
C3–N4–C1''–N2''				178.1°
C8'–C9'–O5–C6				–0.1°
N2''–C3''–C1'''–C2'''				–19.2°

Molecular Dynamics, GPR55 R in a POPC Simulation.* To equilibrate our GPR55 R* model in a realistic environment, a 70 ns MD simulation was performed in a POPC bilayer environment. The most noticeable change that took place during the MD simulation involved the shape of TMH7 and the location of Hx8. Because of the lack of an NPXXY motif (GPR55 has a DVXXY motif instead), CM produced TMH7 conformations that lacked the pronounced bend and discontinuity seen in the β_2 -AR template structure. Energy minimization of the GPR55 bundle produced a distortion in TMH7 as described above. Further changes in TMH7 occurred during the MD simulation of GPR55 R*. Both the wobble angle and face shift angles for TMH7 changed (bend angle of 31.2°, wobble angle of –127.3°, and face shift of 17.9°), and its intracellular end along with Hx8 moved away from the bundle, increasing the size of the IC opening of the receptor. This change is not unexpected because the R3.50–Q6.30 interaction is no longer present in R* and the IC ends of the TMH domain must open during activation to receive the C-terminus of G protein G α . A plot of rmsd versus time (data not shown here) revealed that the rmsd of the GPR55 R* model stabilized by 35 ns. A frame taken at the 36 ns point of this simulation was selected for initial ligand docking.

Ligand Conformational Analysis. *Binding Conformation of LPI from CM Calculations.* For LPI (1), the CM distance-dependent dielectric calculation produced conformers with bent headgroups containing two hydrogen bonds between the inositol C5 hydroxyl (equatorial) and a phosphate oxygen and also between the glycerol hydroxyl and a phosphate oxygen. An LPI

conformer with this bent headgroup orientation and an all-trans fatty acid chain (lowest-energy conformation for the saturated fatty acid chain) was used for initial docking studies in the GPR55 R* model.

Global Minimum Energy Conformations of Compounds 2–5. The key dihedral angles defining the minimum energy conformations of compounds 2–5 are listed in Table 3. See the Supporting Information for a more detailed description of the minimum energy conformations.

GPR55 Agonist Molecular Electrostatic Potential Maps. Figure 5 illustrates the molecular electrostatic potential maps (ranges in kilojoules per mole given next to each ligand) of the docked conformations of LPI (1) and CID compounds (2–4) at GPR55 R*. These ligands have inverted-L or T shapes with the most electronegative atom (or group of atoms) in each compound located in the “elbow” of the L or at one end of the T cross bar, with the rest of the molecule assuming a nearly straight conformation. The phosphate oxygens of LPI, the adjacent nitrogens of the triazole ring of CID1172084 (3), and the sulfonamide oxygens of CID1792197 (2) and CID2440433 (4) are the most electron rich regions of the ligands and were used as the interaction point with K2.60(80). LPI's phosphate group has the most negative electrostatic potential region, which is expected for a negatively charged group.

Docking of LPI and CID Compounds in the GPR55 R Model.* K2.60(80), the only positively charged TMH residue in the putative binding site, was used as the primary interaction site for all the ligands docked in GPR55 R* here. The putative binding site for LPI (1) and CID compounds 2–4 was identified to be

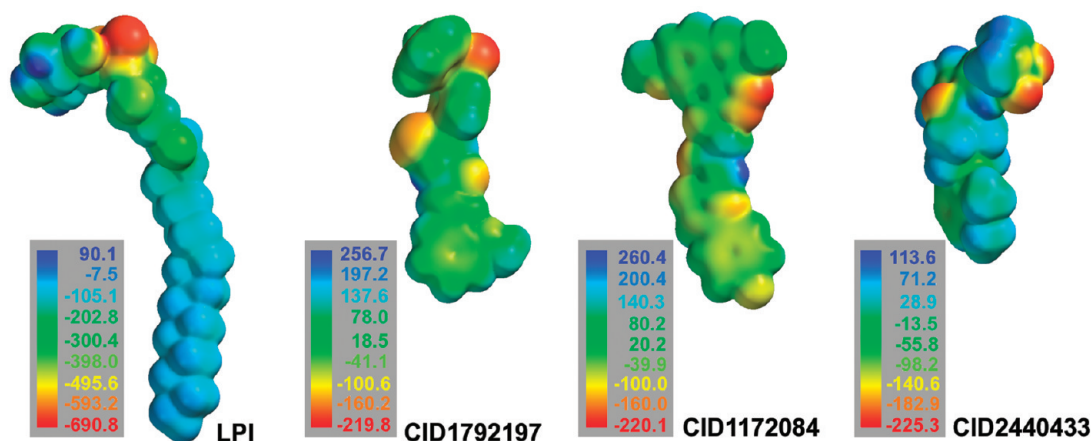


Figure 5. Molecular electrostatic potential maps of the docked conformations of compounds 1–4. The range of the electrostatic potential (in kilojoules per mole) is given next to the map of each compound.

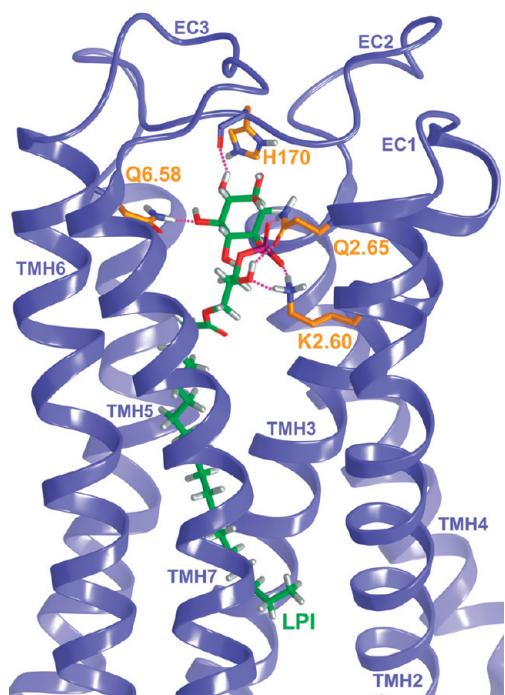


Figure 6. LPI–GPR55 R* complex. The view is from the lipid bilayer with TMH1 removed to simplify the view. TMH2 and -7 are closest to the viewer here. K2.60 has been used as the primary interaction site for the most electronegative region of LPI.

the TMH2-3-5-6-7 region of GPR55 R*, with only the CID compounds reaching TMH5.

Figure 6 illustrates the final LPI (1)–GPR55 R* complex obtained using Glide. The binding site for LPI (1) is located among TMH2, -3, -6, and -7. The primary interaction site for LPI (1) involves a salt bridge between the exposed, charged phosphate oxygen and K2.60(80). The hydrogen bond (N–O) distance and hydrogen bond (N–H–O) angle are 2.45 Å and 168°, respectively. K2.60(80) also forms a hydrogen bond to the glycerol hydroxyl of LPI. The hydrogen bond (N–O) distance and hydrogen bond (N–H–O) angle are 2.68 Å and 157°, respectively. Additionally, Q2.65(85) can form a hydrogen bond to the other exposed oxygen of the LPI (1)

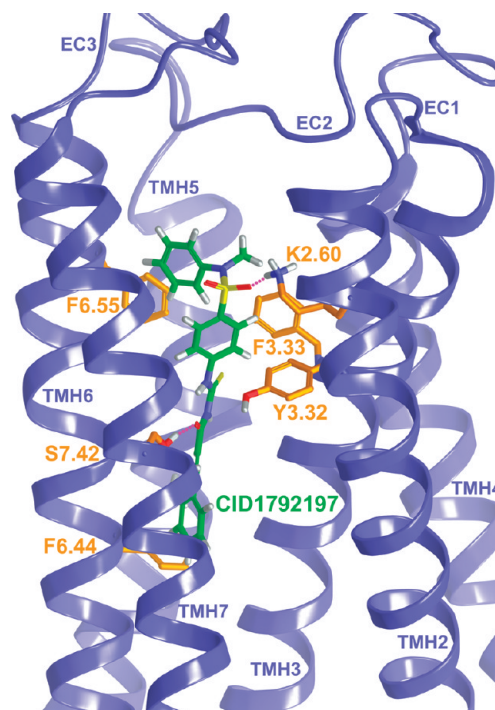


Figure 7. CID1792197 (2)–GPR55 R* complex. The view is from the lipid bilayer with TMH1 removed to simplify the view. TMH2 and -7 are closest to the viewer here. K2.60 has been used as the primary interaction site for the most electronegative region of CID1792197 (2).

phosphate and to the glycerol hydroxyl that is involved with K2.60(80). The hydrogen bond (N–O) distances and hydrogen bond (N–H–O and O–H–O) angles are 2.58 Å, 2.67 Å, 175°, and 141°, respectively. The inositol hydroxyls participate in hydrogen bonding with the backbone carbonyl of the extracellular EC2 loop residue H170 [hydrogen bond (O–O) distance and hydrogen bond (O–H–O) angle of 3.07 Å and 129°, respectively] and also Q6.58(249) at the top of TMH6 [hydrogen bond (N–O) distance and hydrogen bond (N–H–O) angle of 2.82 Å and 159°, respectively]. The total pairwise interaction energy for LPI (1) at this docking site in GPR55 is –89.31 kcal/mol, making this the ligand with the best interaction energy within

the set studied here (see Table S-3 of the Supporting Information). The charge–charge interaction of the LPI (1) phosphate group with K2.60(80) is a major contributor to the interaction energy; however, hydrogen bonding interactions with H170 and Q6.58(249) as well as van der Waals interactions with F3.33(102), M3.36(105), F169, L7.35(270), and M7.39(274) (residues not shown in Figure 6) also contributed significantly to the overall interaction energy.

Figure 7 illustrates the final CID1792197 (2)–GPR55 R* complex obtained using Glide. The CID1792197 (2) sulfonamide oxygen forms a hydrogen bond with K2.60(80), and S7.42(277) forms a hydrogen bond with the carbonyl oxygen adjacent to the thiourea group. The hydrogen bond (N–O and O–O) distances and hydrogen bond (N–H–O and O–H–O) angles are 2.66 Å, 2.60 Å, 166°, and 164°, respectively. CID1792197 (2), just like the other two CID compounds, forms a number of aromatic stacks. The pendant phenyl ring next to the sulfonamide group forms an offset parallel stack with F6.55(246) (ring centroid–ring centroid distance of 5.66 Å), and the phenyl group between the sulfonurea and sulfonamide stacks with Y3.32(101) and F3.33(102). The ring centroid–ring centroid distances are 5.43 and 4.87 Å, respectively, and the angles between ring planes are 74° and 30°, respectively. Finally, the methoxy phenyl group stacks with F6.44(235) with a ring centroid–ring centroid distance of 5.22 Å and an angle between the ring planes of 87°. The total pairwise interaction energy for CID1792197 (2) with GPR55 is –57.84 kcal/mol (see Table S-4 of the Supporting Information). The major interaction energy contributions for this compound are the hydrogen bonding interactions with K2.60(80), the aromatic stacking interactions with F3.33(102), and the van der Waals interactions with M3.36(105), F5.47(190), and L7.35(270) (residues not shown in Figure 7). Aromatic stacking interactions with F6.44(235) and van der Waals interactions with S6.47(238) and V6.51(242) (not pictured in Figure 7) also contribute a small amount to the interaction energy. The remainder of the hydrogen bonding and aromatic stacking interactions identified contribute much less to the overall interaction energy.

Figure 8 illustrates the final CID1172084 (3)–GPR55 R* complex obtained using Glide. The CID1172084 (3) 1,2,4-triazolo[4,3-*a*]quinoline ring nitrogens (positions 1' and 2') form hydrogen bonds with K2.60(80). The hydrogen bond (N–N1' and N–N2') distances and hydrogen bond (N–H–N1' and N–H–N2') angles are 2.92 Å, 2.82 Å, 155°, and 159°, respectively. Y3.32(101) forms a hydrogen bond to the amide nitrogen (N4), and S7.38(273) forms a hydrogen bond to the carboxamide oxygen. The hydrogen bond (O–N and O–O) distances and hydrogen bond (O–H–N and O–H–O) angles are 2.77 Å, 2.57 Å, 158°, and 175°, respectively. Additionally, three phenylalanines form aromatic stacking interactions with the ligand. F3.33(102) and F7.41(276) form a T-stack with the 1,2,4-triazole and the thiazole rings of the compound, respectively (ring centroid–ring centroid distances of 4.96 and 5.94 Å and angles between ring planes of 76° and 25°, respectively), and F5.39(182) forms an offset parallel stack with the first ring of the 1,2,4-triazolo[4,3-*a*]quinoline (distance of 6.21 Å). The total interaction energy for CID1172084 (3) with GPR55 was found to be –65.19 kcal/mol (see Table S-5 of the Supporting Information). While the hydrogen bonding interactions with K2.60(80) and Y3.32(101) and the aromatic stacking interactions with F3.33(102) and F7.41(276) contribute to the interaction energy, the van der Waals interactions with hydrophobic

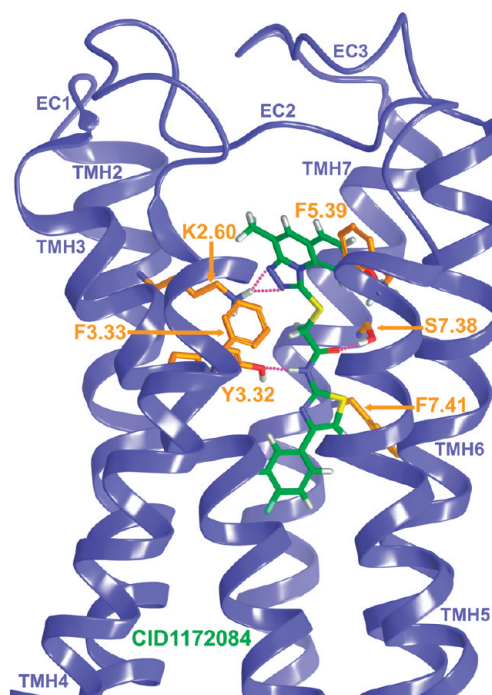


Figure 8. CID1172084 (3)–GPR55 R* complex. The view is from the lipid bilayer with TMH1 removed to simplify the view. TMH4 and -5 are closest to the viewer here. K2.60 has been used as the primary interaction site for the most electronegative region of CID1172084 (3).

residues M3.36(105), L7.35(270), M7.39(274), and S7.42(277) (residues not shown in Figure 8) make a major contribution to the total interaction energy of this complex.

Figure 9 illustrates the final CID2440433 (4)–GPR55 R* complex obtained using Glide. The CID2440433 (4) sulfonamide oxygen forms a hydrogen bond with K2.60(80), and Q6.58(249) forms a hydrogen bond with the pyrrolidine nitrogen. The hydrogen bond (N–O and N–N) distances and hydrogen bond (N–H–O and N–H–N) angles are 2.70 Å, 2.96 Å, 147°, and 172°, respectively. CID2440433 (4) also forms π – π stacking interactions with the receptor. The benzene ring adjacent to the sulfonamide forms an aromatic T-stack with F169 of the EC-2 loop (ring centroid–ring centroid distance of 6.24 Å and angle between ring planes of 58°), and the dimethyl benzyl ring adjacent to the piperazine forms multiple aromatic stacks with F5.47(190) (ring centroid–ring centroid distance of 5.46 Å and angle between ring planes of 23°). The total interaction energy for CID2440433 (4) at this docking site in GPR55 is –52.88 kcal/mol (see Table S-6 of the Supporting Information). The hydrogen bonding interactions with K2.60(80) and Q6.58(249) make major contributions to the interaction energy, as well as the aromatic stacking interactions with H169. However, the van der Waals interactions with hydrophobic residues F3.33(102), V6.51(242), and L7.35(270) (residues not shown in Figure 9) make a substantial contribution to the overall interaction energy.

Assessment of GPR55 R* Binding with the Negative Control CID1135734 (5). CID1135734 (5) is structurally related to CID1172084 (3), differing only in the absence of a fluorine, the presence of an *N*-methyl group in the amide group, and the position of the methoxy group on ring A (see Figure 2). Despite the fact that the differences are small, CID1135734 (5) is inactive

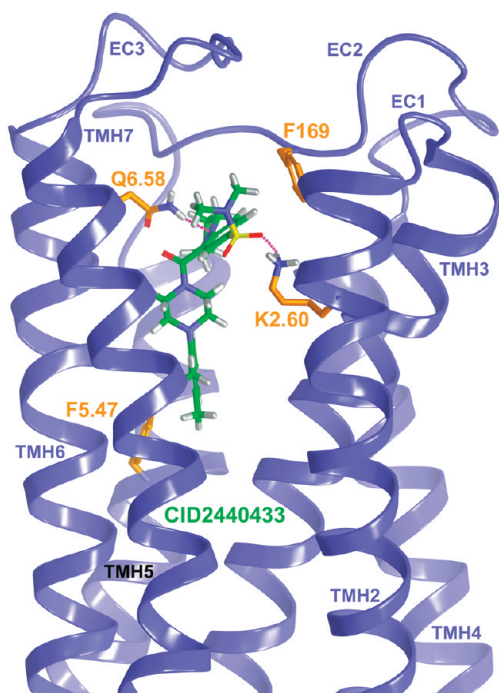


Figure 9. CID2440433 (4)–GPR55 R* complex. The view is from the lipid bilayer with TMH1 removed to simplify the view. TMH2 and -7 are closest to the viewer here. K2.60 has been used as the primary interaction site for the most electronegative region of CID2440433 (4).

at GPR55 whereas CID1172084 (3) is an agonist. CID1135734 (5) was used here as a negative control to test the GPR55 R* model. Figure 10 illustrates the structure of CID1135734 (5) superimposed on the CID1172084 (3)–GPR55 R* complex. Of the structural differences between these closely related compounds, only one appears to interfere with CID1135734 (5) binding. The methoxy attached on the 1,2,4-triazolo[4,3- α]quinoline ring does not have an impact on the interaction of ring N1 and N2 atoms with K2.60. This methoxy group is located in the spacious extracellular loop region where it does not have any steric clashes with the receptor. Additionally, the replacement of the 4-fluoro phenyl ring with the smaller, phenyl ring does not affect interactions with GPR55 R*. However, the CID1135734 (5) *N*-methyl group has major steric overlap with Y3.32(102), a residue that forms a hydrogen bond with the CID1172084 (3) amide moiety. The magenta circle in Figure 10 indicates the region of major steric overlap with Y3.32 of the receptor. This overlap likely explains why CID1135734 (5) does not activate GPR55 (see Table 1).

DISCUSSION

High-Throughput Screening. High-throughput screening remains a powerful tool for the identification of new chemotypes acting at GPCRs. This is especially true in the GPR55 field where many of the known GPR55 ligands were identified from cannabinoid receptor/lipid-biased compound libraries, leading to a low degree of chemical diversity and a lack of receptor selectivity. Here we have identified three novel chemotypic GPR55 ligands (2–4) that lack agonism or antagonism at GPR35, CB1, or CB2. Despite their unique structures, we show that all of these ligands adopt a similar binding mode at GPR55 that is related to the

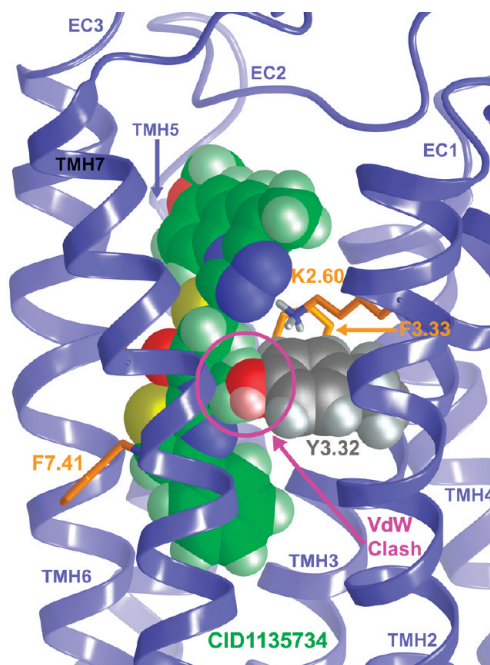


Figure 10. Structure of CID1135734 (5) superimposed on the CID1172084 (3)–GPR55 R* complex. The view is from the lipid bilayer with TMH1. The magenta circle indicates major steric overlap with Y3.32 of the receptor. This overlap likely explains why CID1135734 (5) does not activate GPR55 (see Table 1).

binding mode of the endogenous GPR55 ligand, LPI (1). Our modeling indicates that the molecular shapes and electrostatic potential distributions of these CID agonists mimic that of LPI (1).

Receptor TMH Bundle Shape. It is clear in Figure 4 that while the GPR55 model preserves the general topology of a class A GPCR, the TMH bundle is not identical in shape to the template structure, the β_2 -AR. The differences are dictated here by sequence divergences between GPR55 and the β_2 -AR that our model development protocol (see Methods) is designed to elucidate. There are numerous examples in the literature demonstrating how divergences in a single transmembrane helix, such as TMH1, affect the overall structure of the TMH bundle. For example, Rho TMH1^{13–15} contains a proline at position 1.48, while β_2 -AR TMH1^{16–18} lacks a proline. The proline in TMH1 of Rho bends the EC end of TMH1 back toward the TMH bundle. In contrast, β_2 -AR TMH1 is straight and tilts away from the TMH bundle on the EC side, and this extreme tilt away from the binding pocket is accommodated by the presence of a “hydrophobic wall” of residues on β_2 -AR TMH2 and -7 that shields the binding pocket.

GPR55 is similar to Rho in that it also contains a proline in TMH1, P1.41, and that TMH1 leans into the TMH bundle as a consequence. Likewise, sequence differences in other GPR55 helices and loop segments relative to the β_2 -AR template also permit the formation of TMH bundles by accommodating the difference between GPR55 and β_2 -AR TMH1. Whereas the β_2 -AR^{16–18} contains a TMH2 proline at position 2.59 that causes the TMH2 EC end to lean toward TMH1, GPR55 TMH2 displays a difference in sequence from β_2 -AR^{16–18} that allows it to lean away from TMH1. For GPR55, this is a proline at position 2.58 placed such that TMH2 leans toward TMH3. This is a

critical sequence change because of the very short length of the EC-1 loop in GPR55. The new CXCR4 crystal structure²¹ provides evidence of the consequence of the shift in the position of proline in TMH2 to P2.58. Here it is clear that this change causes TMH2 to lean toward TMH3. The conformation of TMH2 is also critical for the GPR55 ligand binding pocket. The only positively charged TMH residue in GPR55 is K2.60. This residue faces into the binding pocket in our model, as does residue 2.60 in the CXCR4 crystal structure.²¹ It is important to note that residue 2.60 in the β_2 -AR structure¹⁶ is in the TMH2–TMH3 interface and not available from the interior of the binding pocket.

The large majority of class A GPCRs contain an NPXXY motif with Pro at position 7.50. However, GPR55, GPR35, and GPR18 are rare exceptions and instead have a DXXXY motif that lacks proline. This small change in sequence, nonetheless, has an appreciable consequence on hydration and local transmembrane flexibility. For GPR55, although the TMH7 conformation determined from CM is relatively straight, bundle energy minimization induces a bend in TMH7 (see Figure 4) due to N7.43 interacting with the backbone of TMH7. The proximity of two methionine residues from TMH7 (M7.39) and TMH2 (M2.61) causes the EC end of TMH7 to bend away from the bundle as TMH2 and TMH7 pack during the minimization. When the GPR55 receptor model was then placed in the fully hydrated lipid bilayer environment, D7.49 (a residue that is normally N7.49 in other class A GPCRs), along with D2.50, recruits water and this water further perturbs the backbone structure on TMH7 by forming hydrogen bonds to backbone carbonyls in the S7.42–D7.49 region of TMH7. The net result is an increase in flexibility allowing TMH7 to bend at S7.42 and compensate for the lack of a proline at position 7.50.

Receptor Binding Pocket Shape. In most class A GPCR crystal structures, the cocrystallized ligand is oriented horizontally in the binding pocket. This is true for the initial template crystal structure for the GPR55 homology model, the β_2 -AR structure that has the partial inverse agonist carazolol oriented horizontally.¹⁶ In contrast, ZM241385 is oriented nearly vertically in the adenosine A2A crystal structure.²⁰ Because the structure of the β_2 -AR was used as the template to build the model of activated GPR55, the model initially contained a “horizontal” space for binding that did not collapse during minimization of the TMH bundle. However, as the GPR55 R* model equilibrated in the hydrated lipid bilayer, the shape of the binding pocket changed. In addition to the retained horizontal opening, the GPR55 R* model developed a rather straight and deep vertical channel reaching as far as N7.46, the residue that forms the floor of the binding pocket. In the MD simulations, the GPR55 R* binding pocket is highly hydrated, with a string of waters that extends to the IC side, which would be expected for an R* state of the receptor.^{45,57}

The GPR55 R* model agonist binding pocket is lined with hydrophilic residues almost to the intracellular side of the receptor [S3.39(108), S3.47(116), S6.36(227), and S6.40(238)], in contrast to the highly hydrophobic binding pockets of the cannabinoid CB1 and CB2 receptors. The horizontal part of the GPR55 binding pocket underneath the EC loops is lined by K2.60(80), S2.64(84), Q2.65(85), E3.29(98), Q6.58(249), S7.32(267), and Q7.36(271), and the deep vertical channel is lined by Y3.32(101), S6.47(238), S7.42(277), N7.43(278), and D7.49(284).

The resultant shape of the GPR55 R* binding site accommodates ligands that are inverted-L or T shapes with long, thin

profiles that can fit vertically deep in the receptor binding pocket, while their broad head regions occupy the horizontal binding pocket opening near the EC loops. The vertical pore is narrow enough that it cannot accommodate the *N*-methyl group of CID1135735 (5). For GPR55 agonist ligands (1–4), the highest negative electrostatic potential region is exposed either at the elbow of the L or at one end of the T cross bar (in Figure 5, the red regions are the most negative regions). It is this region that interacts with K2.60 in each of the docks pictured in Figures 6–9.

For LPI (1), the highly negative electrostatic potential is formed by the phosphate group (see Figures 2 and 6). In the LPI (1) docked conformation pictured in Figure 6, the phosphatidylinositol (bulky headgroup) is bent at the phosphate and is perpendicular to the 18-carbon saturated fatty acid tail. The elbow of CID1792197 (2) is formed by a sulfonamide group. The two oxygens of the sulfonyl are exposed, and the pendant, *N*-methyl-*N*-benzylamino group from the sulfur is positioned almost perpendicular to the long part of the molecule, placing the sulfur in the elbow (see Figures 2 and 7). The exposed sulfonyl oxygens are the most electronegative regions of CID1792197 (2) that mimic the exposed phosphate oxygens of LPI. CID1172084 (3) does not have a sulfonamide in the elbow but rather a 1,2,4-triazolo[4,3-*a*]quinoline group that is attached at the 5 position to the long segment of the molecule placing the fused ring system perpendicular to the long segment of the molecule and exposing the adjacent nitrogens of the 1,2,4-triazole. The adjacent nitrogens of the 1,2,4-triazole of CID1172084 (3) form the electronegative region that mimics the exposed phosphate oxygens of LPI (see Figures 2 and 8). CID2440433 (4) is T-shaped and has an *N,N*-dimethylsulfonamide group at the end of the cross bar of the T (see Figures 2 and 9), which is the most electronegative region in CID2440433 (4).

Similarity to Other Reported GPR55 Ligands. The cannabinoid CB1 inverse agonist, *N*-(piperidin-1-yl)-1-(2,4-dichlorophenyl)-5-(4-iodophenyl)-4-methyl-1*H*-pyrazole-3-carboxamide (AM251), has been reported to act as an agonist at GPR55.¹⁰ Glide docking studies of this compound show that this compound also can adopt a T shape in the GPR55 binding pocket. The 2,4-dichlorophenyl and pyrazole rings form the cross bar of this T shape, with the 4-iodophenyl ring penetrating into the vertical section of the binding pocket (see Figure S-2 of the Supporting Information). The energy of interaction (EOI) of AM251 with GPR55 is also comparable to the EOI of CID2440433 [CID2440433 (4)–GPR55 EOI = –52.88 kcal/mol; AM251–GPR55 EOI = –53.64 kcal/mol]. Further details concerning the modeled AM251–GPR55 R* complex are provided as Supporting Information.

The structure of CID2440433 (4) is similar to that of the benzoylpiperazine GPR55 agonist, 1-[2-fluoro-4-[1-(methoxy)-ethyl]phenyl]-4-[[4-fluoro-4-(methylsulfonyl)-2-biphenyl]carbonyl]piperazine (GSK494581A), that was reported by Brown and co-workers while our paper was under review.¹² Our GPR55 R* docking study shows that GSK494581A adopts a T shape at GPR55 R*, binding in the same receptor region and in an orientation similar to that of CID2440433 (4) (see Figure 9 and Figure S-3 of the Supporting Information), with a comparable EOI [CID2440433 (4)–GPR55 EOI = –52.88 kcal/mol; GSK494581A–GPR55 EOI = –52.74 kcal/mol]. Further details concerning the modeled GSK494581A–GPR55 R* complex are provided as Supporting Information.

Future Directions. The chemical diversity provided by the three lead compounds, 2–4, combined with the identification

of key receptor interaction sites should provide a basis for the design of more potent and efficacious second-generation GPR55 ligands that retain GPR55 selectivity. Such studies are underway in our laboratories. In addition, mutation studies currently underway include mutations to test the effects of key amino acids in the ligand pocket, including S7.42, S7.38/Y3.32, and Q6.58, which should affect the binding of CID1792197, CID1172084, and CID2440433/LPI, respectively. Preliminary results for K2.60 (K2.60A) suggest that this residue is essential for ligand activation at GPR55 (unpublished observations).

CONCLUSIONS

As a result of a high-throughput β -arrestin screen, we have identified three potent novel chemotypic GPR55 ligands (2–4) that lack agonism or antagonism at GPR35, CB1, or CB2. Their shapes mirror the bioactive conformation of the endogenous GPR55 ligand LPI (1), and docking experiments indicate that CID compounds (2–4), as well as LPI (1), bind GPR55 in the region of TMH2-3-5-6-7. The primary interaction for each of these agonists with GPR55 occurs between K2.60 and an exposed highly electronegative region on each ligand. The chemical diversity of the three agonist lead compounds not only provides a basis for designing improved second-generation GPR55 ligands but also provides the tools for separating the biological activity of GPR55 from that of cannabinoids.

ASSOCIATED CONTENT

S Supporting Information. Example β -arrestin images for compounds 1–5 and the DMSO control (Table S-1), an example of the functional read-out results that corresponds to a dose–response curve in Figure 3A (Figure S-1), a description of the conformations of the selected TMHs that were incorporated in the GPR55 model, a table that shows the correlation of χ_1 F6.48 (239) with F6.44 (235) and H6.52 (243) (Table S-2), a description of the minimum energy geometries of the CID compounds, a set of tables that include all the pairwise interactions of compounds 1–4 at the GPR55 R* binding site (Tables S-3–S-6), and a description with an image of the GPR55 R* binding site and resultant energies of interaction for two other known GPR55 agonists, AM251 (Figure S-2 and Table S-7) and GSK494581A (Figure S-3 and Table S-8). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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ABBREVIATIONS

GPCR, G protein-coupled receptor; CB1 and CB2, cannabinoid type 1 and type 2 receptors, respectively; LPI, lysophosphatidylinositol; 2-AGPI, 2-arachidonoyl-*sn*-glycero-3-phosphoinositol; MLPCN, Molecular Libraries Probe Production Centers Network; TMH, transmembrane helix; EC, extracellular; IC, intracellular; β_2 -AR, β_2 -adrenergic receptor; GFP, Green Fluorescent Protein; CM, Conformational Memories.

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