
G-protein coupled receptors (GPCRs) function as both gatekeepers and molecular messengers of the cell. They relay signals that span the cell membrane mediating nearly every significant physiological process and currently represent the target of about 30% of all drugs. The signals they transmit can arise from a remarkable variety of stimuli which includes, but is not limited to, photons, neurotransmitters and hormones. GPR55, a rhodopsin-like (Class A) GPCR, has received a great deal of attention due to its emerging involvement in a multitude of physiological processes and its putative identity as a third type of cannabinoid receptor. Characterizations of GPR55 knock-out mice reveal a role for the receptor in controlling inflammatory pain, neuropathic pain, and bone resorption. Myriad other studies indicate that GPR55 activation may play a part in oncogenesis and metathesis.

GPR55 can be found in numerous tissue types throughout the body and is also highly expressed throughout the cerebellum and surrounding central nervous system lending credence to the idea that this receptor may play a more crucial physiological role than originally thought. GPR55 has an extensive physiological profile and has been shown to respond uniquely to a great number of diverse compounds. Specifically, it has been shown to recognize many cannabinoid compounds, including CB1 and CB2 endogenous ligands, phytocannabinoids and synthetic cannabinoids. Similar to the ligands of the CB1 and CB2 receptors, the endogenous ligand of GPR55,
lysophosphatidylinositol (LPI), is a lipid-derived molecule. LPI activates ERK1/2 and increases [Ca\(^{2+}\)] and, to date, there has been no evidence that LPI interacts with the other cannabinoid receptors.

Despite innumerable prospective clinical uses hinted at by the aforementioned research no low nanomolar potency ligands of GPR55 have been identified. Nor has there been a radio-ligand developed to characterize the binding site of this receptor. Lack of such tools is a great impediment to any forward progress towards developing the GPR55 receptor as a therapeutic target for drug design.

The following research details the creation of both a GPR55 active- and a GPR55 inactive- state homology model. Towards this goal, Chapter I details the background of the discovery, pharmacological relevance and ligand scope of GPR55. Its purpose is to establish a framework for the research that follows and highlight the medical importance of this elusive receptor.

Chapter II describes the synthetic preparation of antagonists of GPR55 for use in preliminary SAR studies. The original high throughput screen that lead to the identification of novel GPR55 scaffold chemotypes from the screening of over 300,000 compounds gave rise to the piperidinyloxadiazolone compound CID23612552 and the synthetic diversification of what was then dubbed Scaffold 1.

A detailed description of the methods used in the construction of the updated R and R* state of GPR55 models is handled in Chapter III. A combination of Conformational Memories\(^4,5\) (using the CHARMM forcefield), Ligand Conformational Analysis (performed using Spartan (Wavefunction, Inc., Irvine, CA)) and
Macromodel/Maestro/Glide (from the Schrödinger suite) was used to build and refine both GPR55 model states.

Chapter IV then covers model validation and refinement. Using the phenylpiperazine (ML184 CID2440433) and mutations performed in the lab of Dr. Mary Abood (Temple University) it was shown that the current iteration of the GPR55 R* model was indeed a valid representation of the activated state of this receptor. This chapter also provides information that gives rise to the “Future Directions” chapter, Chapter V.

This final chapter is a look forward to the research that still remains to be done to ensure that these models will function as the accurate tools that they have the potential to be. We used the GPR55 R bundle to suggest antagonist structures that will maximize ligand/receptor interactions and hopefully give rise to nanomolar potency molecules. These ligands will need to be synthesized and tested. We also identified key residues in the active bundle (GPR55 R*) that could be mutated to enhance or verify ligand binding. Mutations that destroy receptor function, while interesting, would not have the same utility as the aforementioned kinds of mutations.

References:


CONSTRUCTION AND VALIDATION OF GPR55 ACTIVE AND INACTIVE STATE

IN SILICO MODELS THROUGH THE USE OF BIOLOGICAL ASSAYS,
MUTATION DATA, AND STRUCTURE

ACTIVITY RELATIONSHIPS

by

Mary A. Lingerfelt

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Approved by

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Committee Chair
For my father, J. G. Lingerfelt, whose support was resolute.
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<tr>
<td>A2-AR</td>
<td>A2-adenosine receptor</td>
</tr>
<tr>
<td>βarr2-GFP</td>
<td>β-arrestin2-green fluorescent protein</td>
</tr>
<tr>
<td>β2-AR</td>
<td>β2-adrenergic receptor</td>
</tr>
<tr>
<td>CID2745687</td>
<td>methyl-5-[(tert-butylcarbamothioylhydrazinyl-idene)methyl]-1-(2,4-difluorophenyl)pyrazole-4-carboxylate</td>
</tr>
<tr>
<td>CM</td>
<td>Conformational Memories</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s modified Eagle’s Medium</td>
</tr>
<tr>
<td>DOR</td>
<td>delta opioid receptor</td>
</tr>
<tr>
<td>ERK</td>
<td>extracellular signal related kinase</td>
</tr>
<tr>
<td>FBS</td>
<td>fetal bovine serum</td>
</tr>
<tr>
<td>GPCR</td>
<td>G-protein-coupled receptor</td>
</tr>
<tr>
<td>HBSS</td>
<td>Hanks Balanced Salt Solution</td>
</tr>
<tr>
<td>LPA</td>
<td>lysophosphatidic acid</td>
</tr>
<tr>
<td>Rho</td>
<td>rhodopsin</td>
</tr>
<tr>
<td>TMH</td>
<td>transmembrane helix</td>
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CHAPTER I
THE GPR55 STORY: DISCOVERY, PATHOPHYSIOLOGICAL RELEVANCE AND LIGAND SCOPE

Introduction

G-protein coupled receptor 55 (GPR55), first identified and cloned in 1999, is a receptor in the Class A (rhodopsin-like) G protein-coupled receptor (GPCR) family. This receptor is expressed in myriad tissue types throughout the human body and is particularly populous in human striatum\(^1\) (Genbank accession # NM-005683; Figure 1). In an attempt to discover ligands that could modulate GPR55 activity, screening assays were carried out on cannabinoid libraries by AstraZeneca\(^2\) and GlaxoSmithKline\(^3\). It was discovered that GPR55 responded not only to endogenous or plant-derived CB\(_1\)R/ CB\(_2\)R ligands but various synthetically derived cannabinoid compounds as well. These findings sparked a debate that has raged for over a decade as to the veracity of classifying GPR55 as a novel, third type of cannabinoid receptor belonging in the same subfamily as cannabinoid receptors CBR\(_1\) and CBR\(_2\).\(^4\)\(^6\)

The CB\(_1\) and CB\(_2\) receptors themselves are compelling targets for their potential as modifiers of drug abuse and addiction and it is thought that, if GPR55 is indeed related to these receptors it could have a similar role. Characterizations of GPR55\((-/-)\) (knock-out) mice\(^7\)\(^8\) have shown a definite role for the GPR55 receptor in inflammatory pain, neuropathic pain, and bone development with more recent studies indicating that the receptor may play an even more pervasive role in both energy metabolism and cancer.
motility.\textsuperscript{9,10} These results suggest that the ability to manipulate the active and inactive state of GPR55 has tremendous therapeutic potential.

Figure 1. Human Striatum.

Discussing the pharmacology of GPR55 in a precise manner is a perplexing task at best: Certain CB1 and CB2 antagonists have been reported to act as agonists at GPR55 while other CB1/CB2 agonists have been reported to act as antagonists.\textsuperscript{11} Adding to the already puzzling responses of the receptor was the identification of LPI (lysophosphatidylinositol) as the endogenous ligand for GPR55.\textsuperscript{12} This lysophospholipid has been described as having an EC50 of 200nM to 49nM\textsuperscript{13} by Henstridge et al. in 2009, but later data collected on LPI’s effect on GPR55 has appeared to differ from lab to lab and from assay to assay.

One strategy for navigating the seemingly contradictory pharmacology of GPR55 is to step back and try to understand this receptor from the “ground” up. To do this one must first understand the basic morphology of GPCRs and GPR55 and its ligands specifically.
GPR55 Structure

GPR55 is one of many integral membrane proteins made up of seven membrane-spanning domains known transmembrane helices. This receptor is related to receptors of rhodopsin-like GPCRs (Class A) and shares many of their key sequence motifs (see Figure 2).

The human GPR55 (hGPR55) sequence is relatively unique and only shares low sequence identity with the cannabinoid receptors CB1R and CB2R (13.5% and 14.4%, respectively) described previously. The GPCR proteins displaying the highest percentages of homology with GPR55 are the δ-Opioid (28%), GPR35 (27%), P2Y (29%), GPR23 (30%), CCR4 (23%), LPA4 (30%) and LPA5 (30%). A helix net representation of the GPR55 sequence is displayed in Figure 3.
Figure 3. GPR55 Sequence in Helix Net Form.

The most highly conserved residues across Class A GPCRs are as follows: TMH1 - N1.50, TMH2 - D2.50, TMH3 - E/D3.49, R3.50, Y3.51 (DRY motif), TMH4 - W4.50, TMH5 - P5.50, TMH6 - C6.47, W6.48, X (this residue varies), P6.50 (CWXP motif), and TMH7 - N7.49, P-7.50, X, X, Y7.53 (NPXXY motif).* This conservation of residues and the convenient Ballesteros-Weinstein\(^1\) numbering system make sequence patterns easily identifiable on multiple sequence alignments and allow for easy comparison among residues in the transmembrane regions of different receptors. While GPR55 possesses some of the highly conserved .50 residues (N1.50, D2.50, W4.50 and P5.50) found in many GPCRs, there are a few notable differences. GPR55 has a conservative substitution (DRF) for the TMH3 E/DRY motif, a conservative substitution (SFLP) for the TMH6 CWXP motif, and a non-conservative substitution (DVFCY) for the THM7 NPXXY motif. This sequence divergence is thought to lead to an altered level of hydration and local transmembrane flexibility and, consequently, gives rise to a distinct conformation in this region.\(^1\) Outside of the TMH region, GPR55 has an extracellular-
(EC-1) loop that is shorter than most (three residues versus the six found in β2-AR and Rhodopsin) and an EC-3 loop that is noticeably longer (fourteen residues long versus the five in β2-AR and the six residues in rhodopsin, CB1 and CB2). Located on the EC2 loop there is a cysteine (C168) that is thought to form a disulfide bond with C3.25.11,17 This EC2 disulfide bond, while present in many Class A GPCRs (CXCR4, P2Y12, DOR, etc.) is absent in both CB1 and CB2.

To date there has been no x-ray crystallographic structure resolved for GPR55. Numerous homology models based on available GPCR crystal structures have been reported however, and will be discussed later in this chapter.17,19

* [The Ballesteros–Weinstein numbering scheme is based on the presence of highly conserved residues in each of seven transmembrane (TM) helices. It consists of two numbers where the first denotes the helix, 1–7, and the second the residue position relative to the most conserved residue, defined as number 50. For example, 5.42 denotes a residue located in TMH5, eight residues before the most conserved residue, Pro5.50]

**Pharmacology**

CBR1 and CBR2 are themselves compelling medical targets for their potential as modifiers of drug abuse and addiction and it is thought that, if GPR55 is indeed related to these receptors, it could have a similar role. As mentioned in the introduction, the pharmacological reports of GPR55 activity are far from clear. The jury remains out for a large section of the research community as to whether GPR55 represents a new subtype of cannabinoid receptor or just a novel cannabinoid receptor modulator. Getting an answer to this question and finding an infallible way to operate the active and inactive
states of GPR55 will grant tremendous therapeutic power to medicinal chemists in the near future.

To discuss the pharmacology of GPR55 in a precise manner is a perplexing task at best: While cannabinoid receptors couple to G-proteins of the Gi/o subfamily, as widely demonstrated in the literature, GPR55 has been seen to associate with a number of different G-proteins: G\(\alpha_{13}\), G\(\alpha_{11}\), G\(\alpha_{q}/G\alpha_{q}\), or G\(\alpha_{12/13}\); and its lack of coupling specificity seems to depend on the ligand used and the cell line in which the receptor is expressed. A range of signaling pathways can be initiated based on GPR55’s interaction with individual G proteins: RhoA, MAPK cascades, actin filament formation and intracellular calcium release via the activity of phospholipase C (PLC). The resulting cascades ultimately end in the activation of extracellular signal-regulated kinase (ERK), RhoA activated kinase (ROCK) or phosphatidylinositol 3-kinase (PI3K). In addition, two transcription factors, NFAT or NFkB (that translocate to the nucleus and modulate the expression of different genes), can also be induced following GPR55 activation.

The evaluation of novel potential GPR55 ligands has been vigorously explored through numerous assays with differing functional endpoints: GTP\(\gamma\)S binding, analysis of intracellular calcium levels, phosphorylation of ERK1/2, and the activation of the small GTPase proteins Rac1, RhoA and Cdc42. Maintaining consistency between pharmacological assays has proven to be very problematic however, due to promiscuous nature of GPR55’s response to ligands. It is a fundamental experimental problem when GPR55 ligands modulate not only the eponymous receptor but also CBRs, transient receptor potential vanilloid channels (TRPVs), or peroxisome proliferator-
activated receptors (PPARs).\textsuperscript{29,31,32} Moreover, clear monitoring of the signaling of this receptor may be dependent on intrinsic properties of GPR55 specifically. Unknowns, such as additional active conformation states,\textsuperscript{28} oligomerization,\textsuperscript{29,31,32} allostery,\textsuperscript{33,34} or biased agonism could play a pivotal role in getting the receptor to signal robustly and consistently.\textsuperscript{8,25,30,31} These considerations, and more, have been the topic of many recent reviews written on GPR55, demonstrating the growing interest in resolving the pharmacological conundrums of this receptor.\textsuperscript{11,38,39}

**Biological Relevance of GPR55**

GPR55 is ubiquitously expressed throughout the body; however, its primary physiological function still remains to be fully understood. The receptor has been shown to co-localize with both CB\textsubscript{1}R and CB\textsubscript{2}R.\textsuperscript{29,32} This indicates that GPR55 is extensively expressed throughout the CNS (hippocampus, putamen, caudate, thalamic nuclei) as well as being found in osteoclasts, kidneys and peripheral tissues such as the immune system (spleen, tonsil)\textsuperscript{7,23,39,40,41} GPR55 is also expressed in vascular endothelial cells and this has led various research groups to study the involvement of GPR55 in the regulation of vascular functions.\textsuperscript{6,42-44}

There is increasing evidence that GPR55 plays an integral role in the modulation of inflammatory and neuropathic pain. Activation of GPR55 enhances neuronal excitability by increasing intracellular Ca\textsuperscript{2+} and suppressing the K\textsuperscript{+} current acting on M-type channels as is seen in pro-nociception (the biological amplification of pain signals).\textsuperscript{24} Moreover, Staton et al.\textsuperscript{8} showed that mice lacking GPR55 did not display hyperalgesia
upon inflammatory or neuropathic stimuli. Results from experiments in microglial cells and with neutrophils further support the role of GPR55 in the inflammatory process.

A recent emerging body of evidence that points to the contribution of GPR55 to metabolic regulation and energy homeostasis. Correlations between body weight and levels of GPR55 expression in visceral adipose tissue has been reported and it is clear that GPR55, along with CB1R, controls food intake, gut motility and insulin secretion.

GPR55 also appears to have myriad, secondary roles in other complicated disease states. There is evidence that the receptor plays a crucial part in the physio-pathology of cancer. Increased levels of LPI, the putative GPR55 endogenous ligand, have been found in plasma and ascites in patients with this pathology. Moreover, GPR55 expression is significantly increased in tumor tissues as compared with their healthy counterparts and its expression may be able to be used as a potential biomarker in oncology cases associated with poor prognosis. As mentioned previously, GPR55 signals through Rho GTPases, which control cytoskeleton organization, cell polarity and cell migration, and are closely related to tumor progression. Consequently, activation of GPR55 triggers a number of signaling cascades that stimulate cancer cell proliferation, migration, and invasion. These findings support the idea that GPR55 would make an excellent oncological therapy target in the ongoing fight against cancer.

Perhaps related to GPR55’s ability to induce fibril formation and cancer cell migration is its effect on bone physiology. Whyte et al. reported that manipulation of GPR55 signaling seems to regulate osteoclast polarization and bone resorption activity. Hence, GPR55 may represent a novel target for treatments of arthritis and bone loss
associated with osteoporosis. It is worth noting that GPR55 is not involved in the regulation of CNS development, gross motor movement or learned behavior. It does seem however, to play a role in motor coordination. The physiological, therapeutic potential of GPR55 modulation is schematically summarized in Figure 4.

**WHY GPR55?**

There is an extraordinary array of evidence that implicates GPR55 in the modulation of diverse physio-pathological conditions, but issues still remain with

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Adapted from Henstridge et al.
experimental consistency. It is clear that a drive to secure nanomolar, selective ligands of GPR55 is of utmost future importance.

**GPR55 Ligands**

Due to a lack of information of GPR55 three-dimensional structure and the complexity of interpreting GPR55 functionality assays, the design of potent and selective GPR55 ligands remains a major challenge for medicinal chemists. The following section is a categorization of different GPR55 modulators that have been reported so far in the literature.

**Endogenous Ligands**

Despite the contradictory data reported for many GPR55 ligands, the bioactive lipid lysophosphatidylinositol has demonstrated GPR55 agonism in all studies and functional assays reported so far.\textsuperscript{25,41} Compilation of this data led to the proposal that LPI is an endogenous non-CB1R/ non-CB2R ligand of GPR55. Structurally, LPI contains a glycerol core esterified with a single fatty acid in either the sn-1 or sn-2 position and an accompanying inositol substituted phosphate group.\textsuperscript{12,64} In a recent study performed by Oka et al.,\textsuperscript{26} for identification of the molecular species of LPI in the rat brain, it was found that the predominant fatty acyl moiety was stearic acid (50.5%), followed by arachidonic acid (22.1%). 2-Arachidonoyl-containing LPI species (2-AGPI) displayed the highest potency and efficacy of the LPI species published to date (Figure 5). Due to this, 2-AGPI has been proposed as the natural LPI ligand for GPR55.\textsuperscript{26,65}
The ability of LPI to activate GPR55 has been confirmed in various cellular systems and pharmacological outcomes. For instance, increase of LPI levels has been detected not only in diverse tumors but also in cancer cell proliferation and tumor progression processes. LPI induces phosphorylation of ERK and elicits a rapid Ca²⁺ transient (a brief increase in calcium ion) in GPR55-expressing cells. Further reports have supported these findings in HEK293 cells expressing GPR55 and in large-diameter DRG neurons and endothelial cells. Henstridge et al. showed that LPI stimulates Ca²⁺ release that is dependent on Gα₁₃ and RhoA activation and in β-arrestin PathHunter™ assays, LPI is also revealed to be a potent agonist.
**Cannabinoid-related Ligands of GPR55**

**Endocannabinoids and derivatives.** Several endogenous cannabinoid ligands have been identified as GPR55 modulators (Figure 6). Anandamide, the predominant endocannabinoid, has displayed inconsistent results in GPR55 assays. This lipid neurotransmitter stimulated $[^{35}\text{S}]\text{GTP} \gamma \text{S}$ binding in the nanomolar range and caused calcium mobilization in the micromolar range, yet did not affect phosphorylation of ERK, β-arrestin signaling or receptor internalization. Two-arachidonoylglycerol, another significant endocannabinoid, showed agonist efficacy in a $[^{35}\text{S}]\text{GTP} \gamma \text{S}$ binding assay but was ineffectual in β-arrestin recruitment and GPR55 internalization. These discrepancies may be due to the aforementioned incongruities between functional assays or cell systems and, in accordance with IUPHAR (International Union of Basic and Clinical Pharmacology), do not demonstrate irrefutably that these eicosanoids are in fact GPR55 agonists.

Other endocannabinoids such as palmitoylethanolamide, noladin ether, virodhamine and oleyethanolamide resulted in $[^{35}\text{S}]\text{GTP} \gamma \text{S}$ binding in transiently transfected $h\text{GPR55-HEK293}$ cells, with EC$_{50}$ values of 4, 10, 12 and 440 nM respectively. Results similar to these were previously reported by AstraZeneca in the same $[^{35}\text{S}]\text{GTP} \gamma \text{S}$ assay. According to those GTP$\gamma$S studies, anandamide activated GPR55 and CBRs with similar potencies while palmitoylethanolamide, virodhamine and 2-AG displayed selective action through GPR55.
Figure 6. Endocannabinoids as GPR55 Ligands.

**Phytocannabinoids and related molecules.** Bioactive constituents from the plant *Cannabis sativa* and synthetic analogues (Figure 7) have also shown discrepant data regarding their GPR55 pharmacology. Delta$^9$-THC exhibits activation of GPR55 in $[^{35}S]$GTPγS binding, RhoA assays and intracellular Ca$^{2+}$ mobilization in transiently transfected *h*GPR55-HEK293 cells$^{1,2,24}$ but was unable to stimulate ERK1/2 phosphorylation or β-arrestin recruitment.$^{12,63}$ It remains to be seen whether these results are a consequence of experimental variability, differences in functional readouts or GPR55 intrinsic properties. HU210, a synthetic derivative of Δ$^9$-THC, also displayed activity as GPR55 agonist in diverse assays while being inactive in others.$^{7,2,63}$ Abnormal-cannabidiol (Abn-CBD) and the structurally related O-1602 have been reported as GPR55 selective agonists in $[^{35}S]$GTPγS assays with EC$_{50}$ values in the micromolar and nanomolar ranges respectively.$^{7,6,2}$ Nevertheless, as has been observed with other GPR55 ligands, there have been research groups who reported a lack of activity when using different cellular systems or functional endpoints.$^{26,63}$
The potent CB₁R/CB₂R agonist CP55,940 has also shown conflicting data when used to stimulate GPR55, acting as an agonist in [³⁵S]GTPγS assays⁷,² and antagonist²³,¹²,⁵ in β-arrestin, ERK phosphorylation and Ca²⁺ mobilization tests.

![Figure 7. Structure of GPR55 Active Phytocannabinoids and Synthetic Derivatives Related to Δ⁹-THC.](image)

Of the other phytocannabinoids, such as cannabinol and JWH133, initial studies confirmed their inactivity towards GPR55.⁶ However, in a recent, more sensitive ERK assay, developed by Sharon Anavi-Goffer et al., it was determined that JWH133 could reduce basal pERK acting as GPR55 inverse agonist. From the same GPR55 assay, the phytocannabinoids Δ⁹-tetrahydrocannabivarin, cannabidivarin and cannabigerovarin acted as potent inhibitors of LPI and could constitute a new set of novel GPR55 ligands.
CBD, a phytocannabinoid seen recently in headlines for its omnipotent “curative” powers and whose mechanism of action is not fully understood, has also shown promising GPR55 pharmacological activity. This non-psychoactive component of marijuana acts as GPR55 antagonist preventing $[^{35}\text{S}]\text{GTP}\gamma\text{S}$ binding and Rho activation$^{7,57,66}$ and was inactive in Ca$^{2+}$ mobilization assays$^{24}$ and β-arrestin recruitment$.^{63}$ Conversely, O-1918, a synthetic derivative of CBD that does not bind to CB$\text{Rs}^{67}$ acted as a GPR55 antagonist in intracellular Ca$^{2+}$ functional analysis using endothelial cells that express GPR55.$^{42}$

**Synthetic cannabinoid ligands.**

* Arylpyrazoles. The arylpyrazole scaffold has been extensively explored in the cannabinoid world, becoming very relevant in the design of CB$\text{R}$ or CB$2\text{R}$ inverse agonists or antagonists (Figure 8). Rimonabant (SR141716A), the familiar CB$\text{R}$ arylpyrazole antagonist, has been found to behave as a GPR55 agonist in some assays$^{2,25,63,68}$ but as a GPR55 antagonist in others,$^{23,24,69}$ exerting no effect in one additional study.$^{12}$ In comparison, SR144528, a potent CB$2\text{R}$ antagonist, is inactive in all the GPR55 assays reported so far. It did not induce a rise in calcium in transiently transfected $h\text{GPR55-HEK293}$ cells nor in β-arrestin binding assays.$^{24,63}$ Also studied extensively were AM251 and AM281, CB$\text{R}$ antagonists structurally related to SR141716A. AM251 behaved as a GPR55 agonist in different biochemical assays,$^{3,5,7,13,25,63,68}$ while AM281 did not seem to interact with the receptor$^{7}$ or displayed very weak agonist effects.$^{27,45}$
Figure 8. Arylpyrazoles SR141716A, AM251, AM281, and SR144528.

The CB₁R antagonists SR141716A, AM281 and AM251 can act as agonists while also acting as inhibitors of LPI inducing activation of ERK phosphorylation within the same cellular model. This may explain the seemingly conflicting data surrounding the GPR55 pharmacology of several CB₁R and/or CB₂R ligands. It suggests that certain arylpyrazole ligands may act in a bitopic fashion at GPR55.69

In 2010 Daly et al.⁴³ reported that the fluorescent ligand T1117 activated GPR55 by promoting a characteristic oscillatory Ca²⁺ response in HEK293 cells stably expressing recombinant GPR55. This molecule, structurally analogous to the CB₁R ligand AM251 but linked to a fluorescent tetramethylrhodamine group, showed weak or no affinity towards CB₁R.

Aminoalkylindoles. The CB₁R/CB₂R agonist WIN55,212-2 has been used extensively to probe the endocannabinoid system (Figure 9). This significant amino-alkylindole does not display any activity at GPR55. Consistent data from various biological assays confirm that WIN55,212-2 does not bind GPR55 either as an agonist or an antagonist.⁷,²,²⁴ JWH015, a WIN55,212-2 analogue was, however, able to activate GPR55 in [³⁵S]GTPᵽS assay with EC₅₀ value in the nanomolar range and was effective at
micromolar concentrations in Ca\(^{2+}\) mobilization assays.\(^{24}\) Moreover, data from an ERK1/2 phosphorylation assay suggests that the CB\(_2\)R agonist GW405833 is a bitopic ligand of GPR55, acting as a partial agonist of GPR55 alone or as an allosteric modulator enhancing LPI signaling.\(^{33}\)

Figure 9. Synthetic Cannabinoid Ligands WIN55,212-2, JWH015, and GW405833.

**Coumarins.** The coumarin scaffold has shown great potential and versatility in the development of potent and highly selective CBRs ligands.\(^{70}\) Reexamining this scaffold, using β-arrestin recruitment assays, uncovered 3-substituted coumarins as novel GPR55 antagonists.\(^{34}\) Interaction with CBRs require a lipophilic substituent in position 7 (PSB-SB-487, Figure 10) while having a methyl at position 8 is favorable for GPR55 antagonism (PSB-SB-489, Figure 10). Deng et al.\(^{71}\) have developed computational QSAR models so that they might design coumarin derivatives with enhanced potency.
Magnolol derivatives. Magnolol (Figure 11) is one of the main bioactive compounds in the bark of *Magnolia officinalis*.\(^{72}\) This biphenyllic compound and other related lignans are able to modulate CB\(_1\)R and CB\(_2\)R.\(^{73}\) Magnolol acts as a partial CB\(_1\)R/CB\(_2\)R agonist whereas its major metabolite, tetrahydromagnolol, is a potent peripheral CB\(_2\)R agonist that acts as a weak antagonist of GPR55.\(^{74}\) In a notable study, Müller et al.\(^{75}\) developed structure-activity relationships of new magnolol analogues by varying the alkyl chains and the phenolic groups. They demonstrated that methylation of one of the hydroxyl groups (5´-hexyl-2´-methoxy-5-propylbiphenyl-2-ol) maximizes antagonistic potency when binding to GPR55. This structure represents an untapped potential lead compound in the development of new GPR55 antagonists.

Figure 10. 3-(2-Hydroxybenzyl)-5-isopropyl-8-methyl-2\(H\)-chromen-2-one (PSB-SB-489) and 7-(1,1-dimethyoctyl)-5-hydroxy-3-(2-hydroxybenzyl)-2\(H\)-chromen-2-one (PSB-SB-487).

Figure 11. Magnolol and Analogues.
Non-cannabinoid Related Ligands of GPR55

The growing evidence that illuminates GPR55’s involvement in innumerable biological pathways has generated great interest in this receptor as a therapeutic target.

The logical next step in maximizing this receptor’s potential was the focused drive towards the identification of new selective GPR55 ligands. This drive gave rise to a collaborative project spanning three different laboratories and the Sandford-Burnham screening center of the Molecular Libraries Probe Production Centers Network (MLPCN)\(^{76,77}\) and allowed for the identification of six different GPR55 chemical scaffolds.\(^{76,77}\) This study began with high-throughput screening of a library of compounds using β-arrestin assays in U2OS cells permanently expressing HA-GPR55E and βarr2-GFP.\(^5\) As seen in Figure 12, Phenylpiperazine (CID2440433), triazoloquinoline (CID1172084) and morpholinesulfonlyphenylamide (CID15945391), represent the potent GPR55 agonists discovered in this study\(^{77}\) while piperidinyloxadiazolone (CID23612552), thienopyrimidine (CID1434953) and quinoline aryl sulfonamide CID1261822 are representative GPR55 antagonists.\(^{76}\)

Parallel studies,\(^{68}\) undertaken by GlaxoSmithKline in 2011, validated the previously GSK developed benzoylpiperazine as a GPR55 agonists in *Saccharomyces cerevisiae* yeast and in HEK293 cells. GSK494581A, the most potent ligand, and GSK575594A, the most selective one, had been patented prior to this discovery as glycine transporter subtype 1 inhibitors (being 60-fold selective for GPR55) (Figure 13). Of note is the result that these ligands activate human but not rodent GPR55, hinting at important differences in the binding pocket of the two orthologs.
Figure 12. GPR55 Agonist Scaffolds (CID2440433, CID1172084, and CID15945391) and GPR55 Antagonist Scaffolds (CID23612552, CID1434953, and CID1261822) from HTS. CID16020046 Represents Another GPR55 Antagonist Recently Described by Kargl et al.\textsuperscript{72}

Figure 13. Benzoylpiperazines Previously Reported as Glycine Transporter Inhibitors.
In 2013, Kargl et al. identified CID16020046 (Figure 12) as a selective GPR55 antagonist.\textsuperscript{78} This compound originated from the same MLSCN (Molecular Libraries Screening Centers Network) screen mentioned above; however, it was not selected by those researchers for further development at that time. This molecule antagonizes agonist-mediated GPR55 activation in yeast cells and inhibits GPR55-mediated intracellular Ca\textsuperscript{2+} release. CID16020046 reduces LPI signaling in primary human lung microvascular endothelial cells (HMVEC-L) and in human platelets suggesting yet another novel therapeutic application for GPR55.

An analogue of \((R,R')\)-fenoterol (a short-acting β2-adrenergic agonist), with a 573-fold greater selectivity for β2-AR than β1-AR, has been shown to reduce GPR55 agonist efficiency.\textsuperscript{79} The compound, \((R,R')\)-4′-methoxy-1-naphthylfenoterol (MNF, Figure 14), exerts selective inhibition of GPR55 signaling as demonstrated in various assays with readouts for ERK phosphorilation and cell motility. \((R,R')\)-MNF has been shown to reduce the pro-oncogenic activity of GPR55 thus revealing its therapeutic potential as an antitumor agent.

![Figure 14. Naphthylfenoterol (MNF).](image-url)
An analysis of the vast range of GPR55 scaffolds reported to this point may help to elucidate the structural motifs involved in modulating the activity of this receptor. It is hoped that this information will contribute to the work being done in the rational design of new compounds able to selectively and potently bind this promising target.

The construction of a comprehensive and current model of both the active and inactive state of the GPR55 receptor was therefore necessitated by the aforementioned results. Development of a binding site template, to guide the design of novel ligands with which to control the activation of this receptor, has the potential to be a vast and relatively unexploited therapeutic source. To date, no low nanomolar potency ligands have been identified or synthesized, nor has there been a radioligand developed to characterize binding at this receptor. The goal of the following research to be presented was the building, refinement and testing of two GPR55 receptor prototypes (an active state model and an inactive state model) to be used as a tool for exploring the ligand binding site. This predictive structure is essential for screening of current synthetic analogs and for the improved design of higher potency ligand profiles. Combining the extrapolative power of modern computational methods with the structure activity feedback from collaborating biochemists allows for a complete picture of the functioning model of GPR55 to be recognized.
References


CHAPTER II

DESIGN, SYNTHESIS, AND ANALYSIS OF ANTAGONISTS OF GPR55: PIPERIDINE-SUBSTITUTED 1,3,4-OXADIAZOL-2-ONES

Introduction

The work described herein began as a collaborative project between the Reggio and Abood labs and the Sanford Burnham Screening Center of the Molecular Libraries Probe Production Centers Network (MLPCN). A series of GPR55 antagonists belonging to novel GPR55 antagonist chemotypes and having IC$_{50}$s in the 0.34 to 2.72 µM range were identified (8). These compounds were then screened for agonist activity with GPR55 along with agonist/antagonist activity at both the CB1 and CB2 receptors. Several of the GPR55 antagonists identified were completely selective. Structure-activity relationship (SAR) studies of chemotype one were began in the synthetic lab of Dr. Mitch Croatt. The results of initial ChemNavigator compound database substructure searches for Scaffolds 1-3 were used to guide the development of the research strategy to be presented in this chapter. Given GPR55’s complicated pharmacological profile it is unsurprising that search results varied widely between scaffolds. Scaffold 1 had only 5 hits in the Aldrich database with a Tanimoto similarity of 70% or greater. Scaffold 2 (which will be expounded upon in Chapter IV) had 91 unique hits from the Aldrich database, of which only 39 could truly “fit” in the original GPR55 model. In contrast, Scaffold 3 had 161 unique hits from the Aldrich database, with all regions of space
sampled quite well. The readily available commercial analogs of scaffolds 2 and 3 lead to Scaffold 1 being chosen for initial ligand diversification.

The synthetic route was only three steps from simple starting materials: amide bond formation between the carboxylic acid and piperidone,\textsuperscript{33} reductive amination of a hydrazide and the ketone,\textsuperscript{34} and a carbonylative cyclization\textsuperscript{35} (Scheme 1; Figure 15).

![Scheme 1. Synthetic Scheme for Scaffold 1.](image)

Figure 15. Scheme 1: Synthetic Scheme for Scaffold 1.

Figure 16 shows a subset of the available R-groups and Aryl-groups that were initially examined. It is also important to note that all of the commercially available analogues of parent compound 1 had the same phenyl group (Ar = Ph). The synthetic route proposed allowed for this position to be modified in hopes to observe a direct effect on the IC\textsubscript{50} values. In the original model the carbonyl of the azoxazolone interacts with K2.60 so it is hypothesized that if we place electron-rich aryl-groups next to this heterocycle, we could strengthen the interactions between the ligand and the receptor.
The overarching goal of the research presented in this chapter was the creation of a manageable library of synthetic antagonists with which to begin revision of the GPR55 model. It was postulated that results from the antagonist synthesis combined with mutation data from the Abood lab would aid in the revision of the initial GPR55 model that had been created in the Reggio lab. The revised model would then be used to guide synthesis of new ligands and allow for a chem-informatics search for additional chemotypes.
Design, Synthesis, and Analysis of Antagonists of Gpr55: Piperidine-Substituted 1,3,4-Oxadiazol-2-Ones

As submitted to Organic Medicinal Chemistry Letters*

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1) Figures/Tables were re-rendered/resized to match dissertation global style.
2) Page layout has been altered from published form to meet dissertation layout requirements (i.e. the use of 1 column vs. 2, etc.)

Abstract

A series of 1,3,4-oxadiazol-2-ones was synthesized and tested for activity as antagonists at GPR55 in cellular beta-arrestin redistribution assays. The synthesis was designed to be modular in nature so that a sufficient number of analogues could be rapidly accessed to explore initial structure-activity relationships. The design of analogues was guided by the docking of potential compounds into a model of the inactive form of GPR55. The results of the assays were used to learn more about the binding pocket of GPR55. With this oxadiazolone scaffold, it was determined that modification of the aryl group adjacent to the oxadiazolone ring was often detrimental and that the distal
cyclopropane was beneficial for activity. These results will guide further exploration of this receptor.

**Introduction**

GPR55, a recently deorphanized, rhodopsin-like (class A) G protein-coupled receptor (GPCR), is a receptor for l-α-lysophosphatidylinositol (LPI, Figure 17) which serves as the endogenous agonist (GenBank entry NM 005683).\(^1\) Initial studies noted that a variety of CB1 and CB2 ligands bind to GPR55\(^2,3\) and more recent studies have focused on physiological roles for GPR55 in inflammatory pain,\(^2\) neuropathic pain,\(^2\) bone development,\(^3\) and the potential for activation of GPR55 being pro-carcinogenic.\(^4,5,6,7,8\) Despite the important potential biological functions of GPR55, the research is limited by the lack of both potent and selective agonists and antagonists.\(^9,10\)

![Figure 17. LPI and Lead Antagonists of GPR55.\(^{12}\)](image)
Based on a high-throughput, high-content screen of approximately 300,000 compounds from the Molecular Libraries Probe Production Centers Network initiative, a few molecular scaffolds were identified that had relatively good selectivity and potency as antagonists at GPR55. These structures were then docked into the inactive state model of GPR55 to visualize the key features of the antagonists. Of the compounds that exhibited selective and moderate activity as antagonists at GPR55, three different structural families were identified as illustrated by ML191, ML192, and ML193 (Figure 17). The docking of the structures in Figure 17 into the inactive state model of GPR55 indicated a few important interactions as we previously reported. Briefly, the primary interaction was hydrogen bonding between the lysine at position 2.60(80) and the oxadiazolone carbonyl in ML191, the amide carbonyl in ML192, or an oxygen of the sulfonamide in ML193. The hypothesized interactions with K2.60(80) positioned the bottom aryl rings of all three structures, as represented in Figure 18, to maintain the toggle switch interaction between M3.36(105) and F6.48(239). The remaining interactions of the ligands presented in Figure 18 and GPR55 are primarily aromatic stacking with various residues. Specifically for ML191, the toluene ring attached to the cyclopropane stacks with F169 and the phenyl group attached to the oxadiazolone stacks with F6.55(246) and F3.33(102) (Figure 18). In addition to these interactions, moderate beneficial van der Waals interactions were identified between the oxadiazolone and both M7.39(274) and Y3.32(101). Since the interactions between ML191 and GPR55 centered on the three aromatic rings of ML191, compounds were desired that modified the electronics and steric of these areas. Hence, the ML191 synthetic studies reported herein
were undertaken to explore the SAR of this oxadiazolone class of compounds. ML191 was also chosen as the lead antagonist since there are very few structurally related compounds that could be purchased and screened compared to the available compounds for ML192 and ML193.

Figure 18. (A) Docking and Key Interactions between ML191 and GPR55. ML191 (Green) Has a Key H-bond Interaction with K2.60 (Pink). ML191 Also Has π-stacking or Other van der Waals Interactions with F169, F3.33, F6.55, M7.39, and Y3.32 (All Mustard). The Interactions with M7.39 and F6.55 Appear to Hinder the Rotation of M3.36 and F6.48 (Both Purple) Which Are Considered the Toggle Switch for GPR55. (B) Electrostatic Potential Map of ML191. This Figure is Adapted from Previously Published Work.12
Our synthetic approach to GPR55 antagonists was designed so that many different structures could be accessed to rapidly explore initial SAR, along with validating or modifying our current model (Figure 18). The synthesis begins with the coupling of a carboxylic acid to 4-piperidone by first forming the acid chloride (Scheme 1a; Figure 19). The different acids chosen, based on the initial hit, modify the electronics and steric at this section of the molecule. Relative to ML191, compound 2a reduces the steric impact, 2b increases the electron-density in the aromatic ring, and compounds 2c and 2d decrease the electron-density. Compounds 2e and 2f were selected to examine the influence of steric bulk at the position of the cyclopropane ring. The largest change in overall structure relates to the 1-naphthoic acid derivative (2f). Although the naphthalene ring is structurally different, this analogue can position the distal aromatic ring in a similar position as the phenyl rings of the other analogues since the bond angle for the Cα will be similar to that of the cyclopropane analogues, however, this structure is much flatter.

![Scheme 1a: Synthesis of Acylated Piperidones.](image)

Figure 19. Scheme 1a: Synthesis of Acylated Piperidones.
With a handful of acylated piperidones prepared, the final two steps first involved a reductive coupling of aryl hydrazides (3t–z) with the previously synthesized piperidones (2a–f) to yield hydrazides 4 (Scheme 2; Figure 20). These compounds were then cyclocarbonylated using triphosgene to yield oxadiazolones 5. The reductive coupling reactions proceeded smoothly but the products of that step were often unstable to silica gel chromatography. Therefore, the unpurified products were treated with triphosgene without further purification. This modification of the synthesis typically improved the yields of the final compounds (see Supplementary Data in Appendix A for individual yields).

![Scheme 2: Synthesis of GPR55 Antagonists.](image)

Similar to the cyclopropane starting materials (1a–f), the hydrazides (3t–z) were selected to probe the electronic and steric opportunities in the binding site. Based on the current model (Figure 18), the aromatic ring adjacent to the oxadiazolone is involved in
an interaction with M3.36(105) and F6.48(239). Additionally, the oxadiazolone contributes as the key interaction between the basic carbonyl oxygen with the ammonium of K2.60(80). Thus, electron rich aromatic rings adjacent to the oxadiazolone should make the carbonyl oxygen more basic and strengthen this interaction.

A targeted exploration of the SAR of all six acids (1a–f) with hydrazide 3t and all seven hydrazides (3t–z) with acid 1a (Figure 19 and Table 1) was performed instead of synthesizing and exploring the biological activity of all 42 permutations of the six acids and seven hydrazides. Acid 1a and hydrazide 3t were chosen as the constants since these were the most simplified pieces consisting of an unsubstituted phenyl ring. Unfortunately, there were solubility issues with some of the compounds (e.g., 5bt and 5bv), so additional combinations were required to elucidate the effect of the different areas of the scaffold.

Table 1
GPR55 Antagonist Activity of Compounds

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<tr>
<th>Entry, Compound</th>
<th>R</th>
<th>Ar</th>
<th>IC$_{50}^a$ (95% CI)</th>
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</thead>
<tbody>
<tr>
<td>1, 5at</td>
<td>![image]</td>
<td>![image]</td>
<td>12 µM (3.9 - 36)</td>
</tr>
<tr>
<td>2, 5au</td>
<td>![image]</td>
<td>![image]</td>
<td>0.42 µM (0.075 - 2.4)</td>
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Table 1

Cont.

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<tr>
<th>Entry, Compound</th>
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<tr>
<td>10, 5et</td>
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<td><img src="366x351" alt="Image" /></td>
<td>0.77 µM (0.39 - 1.5)</td>
</tr>
</tbody>
</table>

*Note.* IC$_{50}$ values and 95% confidence intervals (CI) were determined by running the sample in triplicate versus a 6 µM concentration of LPI. $^b$Compound was not completely soluble at the concentrations of the assay.
Compounds were initially screened via an image-based cell assay to identify antagonist activity. The rationale for using the β-arrestin recruitment assay was to provide a fair comparison of IC$_{50}$ values since our initial report employed this assay.$^{11,12}$ Briefly, U2OS cells overexpressing GPR55 and βarr2-GFP were exposed to LPI (6 μM; EC$_{80}$) resulting in the recruitment of β-arrestin. Antagonist activity was evaluated by ligand-mediated inhibition of LPI-induced receptor activation. This strategy quickly identified the compounds that had IC$_{50}$ values higher than 15 μM which were excluded from further analysis (Figure 21).

Concentration response curves were generated for compounds that were active at concentrations below 15 μM employing both the image-based β-arrestin recruitment assay and the DiscoveRx PathHunter® chemiluminescent β-arrestin complementation assay. In the DiscoveRx PathHunter® system, CHO-K1 cells stably expressing GPR55

![Compounds with Poor Activity](image)

Figure 21. Analogues with Poor Activity (>15 μM).
(fused with a β-galactosidase enzyme fragment), and β-arrestin (fused to an N-terminal deletion mutant of β-galactosidase) were used to quantitate the inhibition of LPI-induced β-arrestin activity (Figure 22, Table 1). Hence, antagonist activity was evaluated through the use of two differential means of β-arrestin quantitation, in two different cellular backgrounds (see Supplemental information, Biological Assay). IC₅₀ values were similar in both methodologies.

Figure 22. Representative Images of Antagonist Screening. (A) 0.1 μM 5dt + LPI; (B) 1.0 μM 5dt + LPI; (C) 10 μM 5dt + LPI; (D) 3 μM LPI; (E) 10 μM 5dt; (F) DMSO.

Screening of the compounds allowed for a number of interesting SAR observations. First of all, pyridyl analogue 5au demonstrated that a pyridine ring would be beneficial for both c log P as well as increasing the potency (entry 2 versus entry 1). Electron-poor aryl groups next to the oxadiazolone were detrimental (Figure 21), but electron-rich groups were also not beneficial (entry 3). As discussed earlier, it was
anticipated that the more electron-rich aryl groups next to the oxadiazolone would be beneficial electronically, but the results obtained could be validated by the electron-donating groups being larger and creating some detrimental steric interactions.

It was found that electron-rich cyclopropylaryl groups had relatively good activities (entry 7), but also typically had solubility problems (entries 5 and 6). Fortunately, analogue 5bw was soluble, but the moderate activity illustrates that the \( p \) -methoxy group is detrimental since the most closely related analogue (ML191, Figure 17) was more active and the electron-withdrawing \( p \)-chloro analogue (5dt) was even more active. Dichlorophenyl analogue 5ct had good activity, but was not as potent as compared to the monochloroaryl compound (entries 8 and 9) which could be justified based on the larger steric bulk of the second chlorine atom. Structure 5et with the dimethylcyclopropyl group was similarly active (entry 10) as compared to the parent compound, ML191 (Figure 17) which is interesting since this compound adds more steric bulk to the cyclopropyl aryl section of the molecule, albeit in a slightly different location than analogue 5ct. It should be noted that analogue 5et is the only structure analyzed that is chiral. The synthesis of 5et was racemic and the model indicates that there are no major anticipated differences in activities between the two enantiomers.

In conclusion, this letter presents initial SAR for piperidine-substituted oxadiazolone antagonists at GPR55, a recently deorphanized G protein-coupled receptor that lacks a potent and selective ligand of nanomolar potency. These data help to better define areas for improvement of this family of GPR55 antagonists since both halves of the molecule were independently modified. The activities spanned about two orders of
magnitude and will be used as a guide for future efforts which will be published in due time.

Acknowledgments

This research was supported by National Institutes of Health Grants R21NS077347, R01DA023204, T32DA007237, and P30DA013429. The authors thank Dr. Franklin J. Moy (UNCG) for assisting with analysis of NMR data and Dr. Brandie M. Ehrmann (UNCG) for acquisition of the high-resolution mass spectrometry data at the Triad Mass Spectrometry Laboratory at the University of North Carolina at Greensboro.
References


CHAPTER III

BUILDING OF THE ACTIVE AND INACTIVE STATE MODELS OF GPR55

Introduction

The previous GPR55 homology model (Evangelia et al.)\textsuperscript{1}, was created using the crystal structure of the β2-adrenergic receptor (Protein Data Bank entry 2RH1)\textsuperscript{2} as a template. That model was later altered to reflect several structural elements that were observed in the recently crystallized structure of the CXCR4 receptor in its inactive state.\textsuperscript{3} The impetus for the current revision of the receptor originates from the pursuit of two goals: (a) A desire to explain new mutation results, suggesting that K2.60 and E3.29 are crucial residues for binding in GPR55 and (b) The need to have an up-to-date model that reflects the high degree of sequence homology GPR55 shares with a logically chosen crystal structure template and that incorporates the structural information gleaned from both the agonist and antagonist ligands that have been synthesized thus far.

In the work discussed here, the Ballesteros-Weinstein numbering system for Class A GPCRs will be used.\textsuperscript{4} In this numbering system, the position of each residue is labeled by two numbers. The first (1 through 7) corresponds to the transmembrane helix (TMH) in which the residue is located. The number following the helix indicates the residue’s position relative to the most highly conserved residue (among Class A GPCRs) in that helix. The conserved residue in each helix being given the number .50 and residues before or after it are numbered accordingly.
Sequence alignments (Figure 23) have been compiled which compare GPR55 with CB1, CB2, the h-δopioid receptor and other Class A GPCRs. These alignments allow for a more facile comparison of amino acid sequence similarities and deviations between receptors. The same highly conserved residues that have been used in the past to align the sequences of CB1⁵⁻⁹ and CB2¹⁰,¹¹ to that of bovine rhodopsin were used as alignment guides for GPR55. These residues are the most highly conserved residues across Class A GPCRs: TMH1 - N1.50, TMH2 - D2.50, TMH3 - E/D3.49, R3.50, Y3.51 (DRY motif), TMH4 - W4.50, TMH5 - P5.50, TMH6 - C6.47, W6.48, X (this residue varies), P6.50 (CWXP motif), and TMH7 - N7.49, P-7.50, X, X, Y7.53. Sequence patterns are easily identifiable on multiple sequence alignments and allow for easy comparison among residues in the transmembrane regions of different receptors. While GPR55 possesses some of the highly conserved .50 residues (N1.50, D2.50, W4.50 and P5.50) found in many GPCRs, there are a few notable differences. GPR55 has a conservative substitution (DRF) for the TMH3 E/DRY motif, a conservative substitution (SFLP) for the TMH6 CWXP motif, and a non-conservative substitution (DVFCY) for the THM7 NPXXY motif. Outside of the TMH region, GPR55 has an extracellular-1 (EC-1) loop that is shorter than most (three residues versus the six found in β2-AR and Rhodopsin) and an EC-3 loop that is noticeably longer (fourteen residues long versus the five in β2-AR and the six residues in rhodopsin, CB1 and CB2). Based upon sequence alignments from the Reggio lab, the template chosen for the new GPR55 models was the δ-Opioid Receptor.¹² See Figure 24.
Figure 23. GPCR Multi-Sequence Alignment.

Figure 24. δ-Opioid Receptor.
Constructing the GPR55 R and GPR55 R* TMHs

Updating the receptor models mandated an assessment of sequence-dictated conformational divergences of the GPR55 transmembrane helices (TMH) from the chosen 1.8 angstrom δ-opioid receptor crystallographic template. The Monte Carlo/simulated annealing technique CM13 was used to study the conformations of four GPR55 TMHs (TMHs 1, 5, 6, 7) with important sequence divergences from the DOR template. This technique allows for thorough exploration of conformations of TMHs containing helix deforming residues such as Proline, Glycine, Serine or Threonine.

Figure 25. CM Output for Active-State GPR55 TMH6 Clustered, Showing F6.48.
Conformational Memories explores the low free energy conformations possible for a helix of interest using Monte Carlo simulated annealing. The CM method, developed by Guarnieri and Wilson\textsuperscript{13} and expanded by Guarnieri and Weinstein,\textsuperscript{14} was used originally to thoroughly explore the dihedral conformational space of a molecule, independent of the dihedral conformation of the initial molecular structure. The method has been expanded to allow the variation of bond angles in addition to dihedral angles.\textsuperscript{15} CM combines Monte Carlo exploration of the dihedral angle space with simulated annealing (MC/SA) to determine the range of values in which each dihedral angle is capable of existing within a broad temperature range. In the CM calculations used to construct the GPR55 TMHs under consideration, the backbone dihedrals of each helix were set to the standard $\phi$ (-63°) and $\psi$ (-41.6°) for transmembrane helices. The established protocol is then to allow all torsion angles to vary ±10°, and to allow a larger variation of ±50° in regions containing flexible series of residues.\textsuperscript{16}

The CM calculation is performed in two phases: an exploratory phase and a biased annealing phase. In the exploratory phase, a random walk is used to first identify the region of conformational space most probable for each torsion angle and bond angle. The initial temperature for each run was 3000 K with 50,000 Monte Carlo steps applied to each torsion or bond angle variation with step-wise cooling, over 18 steps, to a final temperature of 310 K. Each step consists of varying two dihedral angles and one bond angle chosen at random from the entire set of variable angles. The torsion angles and bond angles are randomly picked at each temperature and each move is accepted or rejected using the Metropolis Criterion (Figure 26).\textsuperscript{17}
Accepted conformations in the Exploratory Phase are used to create “memories” of torsion angles and bond angles that were accepted (Figure 27). This information provides a map of the accessible conformational space of each TMH as a function of temperature. In the biased annealing phase, the only torsion angle and bond angle moves attempted are those that would keep the angle in the “populated conformational space” mapped at 310 K in the exploratory phase. The biased annealing phase for the calculations reported here began at 749.4 K with cooling to 310 K in 7 steps.
One hundred eight structures were output at 310 K. The output from each TMH study was then superimposed on the corresponding template helix in the δ-Opioid receptor template from the intracellular (IC) end of the receptor up to the .50 residue. This particular superposition was to allow for examination of how CM represented the variation that could occur at the extracellular (EC) end of each TMH. The only exception to this superposition was TMH6 which had to be superimposed from the IC portion of the helix downward so as to represent the GPR55 R* “kicked out” region, occurring upon receptor activation, of the bottom of the helix. A helix was selected for inclusion in the revised GPR55 model that fit in the bundle with no van der Waals overlaps with residues on other TMHs.

**Extracellular and Intracellular Loops**

Extracellular and intracellular loops connecting all 7 TMHs were then added using MODELLER v8.2 26. MODELLER is a computer program used in producing homology models of protein tertiary structures. It employs a technique originating in nuclear magnetic resonance known as satisfaction of spatial restraints, by which a set of geometrical criteria are used to create a probability density function for the location of each atom in the protein.

Energy minimizations were performed using Macromodel and the OPLS2005 all-atom force field (version 9.8, Schrödinger, LLC, New York, NY). A distance-dependent dielectric, 8.0 Å extended non-bonded cutoff, 20.0 Å electrostatic cutoff, and 4.0 Å hydrogen bond cutoff were used. The minimization was performed in two stages. In the first stage, a harmonic constraint was placed on all the TMH backbone torsions (ϕ, ψ, and ω), with this constraint gradually reduced to zero in 500-step increments (using a total of
5000 steps to reach zero). In addition, a 500 kcal/mol harmonic constraint was placed on the backbone torsions of the loops. The minimization consisted of a conjugate gradient minimization using a distance-dependent dielectric, performed in 1000-step increments until the bundle reached the 0.05 kJ/mol gradient. In the second stage of the calculation, the TMH portion of the bundle was 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.

**Conformational Assessment of the Ligands**

A complete systematic conformational analysis of ligands (both agonists and antagonists) to be docked in the GPR55 models was performed, in parallel to the CM and prior to the docking of each ligand in its respective receptor, using *ab initio* Hartree-Fock calculations at the 6–31G* level as encoded in the Spartan molecular modeling program (Wavefunction, Inc., Irvine, CA). Specifically, HF 6–31G* 6-fold conformer searches were performed for all rotatable bonds within a molecule (Figure 28). 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. As an example of the output, the global minimum energy conformer of ML184 and the actual docked conformer are shown below superimposed on all heavy atoms (Figure 29).

Figure 28. GPR55 Agonist ML184 with Two of Six Rotatable Bonds Indicated.

Figure 29. GRPR55 Agonist ML184 Docked (Cyan) and Global Minimum (Copper) Conformations.
Docking of Ligands

The lowest energy conformation of all ligands was used, unless otherwise noted, for all receptor docking studies. The Schrödinger module, Induced Fit, (Schrödinger Inc. Portland, OR) was used to explore possible binding conformations and receptor site interactions in the GPR55 R and R* models. The binding site of the receptor was defined by the area enclosed in a box with dimensions 26Å x 26Å x26Å (the default value).

Because the endogenous ligand, LPI, is negatively charged, it is likely that a positively charged amino is a primary interaction site for GPR55 ligands. The only positively charged amino acid in the binding pocket of the original model was K2.60 (80). Recent K2.60A and E3.29A mutation studies suggest that both of these charged residues are crucial for ligand activation at GPR55 (M. Lingerfelt et al., manuscript submitted to Biochemistry). For this reason, both K2.60 (80) and E3.29A (98) were set as the center of the box and a hydrogen bond with K2.60 (80) was defined as a constraint. The two constraints (enclosing box and hydrogen bond) were used for both steps that Induced Fit Docking (IFD) performs.

The initial receptor minimization that IFD implements in preparation for docking ligands was omitted because the receptor model had already been energy minimized. For the first Glide calculation, receptor and ligand Van der Waals radii were set to the default value of 0.50 and the number of maximum poses to be produced was also set to 40. During the Prime stage of the calculation, amino acids within 5.0Å of the ligand were refined to better accommodate the ligand. Since the GPR55 receptor is a transmembrane protein, an implicit membrane was used during the Prime refinement step. During the
second step, the actual docking of the ligand performed by the module GLIDE, the top 30 poses produced by the first calculation were re-docked in the receptor. Docked poses that were within 40.0 kcal/mol of the absolute relative lowest one were kept.

**Ligand/Receptor Minimization**

To optimize the ligand/receptor interaction in the receptor/ligand complexes, each was minimized using Macromodel 9.1 (Schrödinger Inc.; Portland, OR). As described above in the section for bundle construction, 500 steps of Polak-Ribier conjugate gradient minimization using force field defined dielectric were performed. A harmonic constraint was placed on all the TMH backbone torsions (φ, ψ, and ω) to preserve the general shape of the helices during minimization. The sidechains of the TMH region, the loops and termini were allowed to relax. The Generalized Born/Surface Area (GB/SA) continuum solvation model for water as is implemented in Macromodel 9.1 was used for minimizing the loop regions. An 8.0Å nonbonded cutoff (updated every 10 steps), a 20.0Å electrostatic cutoff, and a 4.0Å hydrogen bond cutoff were used in each stage of the calculation. The transmembrane region and the docked ligand were then frozen while the loops were minimized. An example of the resultant ligand/receptor interaction energies for GPR55 agonist ML184 and the GPR55 R* bundle can be seen in Supplemental Table 1 in Chapter IV.
References


CHAPTER IV
IDENTIFICATION OF CRUCIAL AMINO ACID RESIDUES INVOLVED IN AGONIST SIGNALING AT THE GPR55 RECEPTOR

As Submitted to Biochemistry

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1) Figures/Tables were re-rendered/resized to match dissertation global style.
2) Page layout has been altered from published form to meet dissertation layout requirements (i.e. the use of 1 column vs. 2, etc.)

Abstract

GPR55 is a newly de-orphanized Class A GPCR that has been implicated in inflammatory pain, neuropathic pain, metabolic disorder, bone development, and cancer. Few potent GPR55 ligands have been identified to date. This is largely due to an absence of information about salient features of GPR55, such as residues important for signaling and residues implicated in the GPR55 signaling cascade. The goal of the work reported here was to identify residues that are key for the signaling of the GPR55 endogenous ligand, 1-α-lysophosphatidylinositol (LPI), as well as the signaling of the GPR55 agonist, ML184, (CID 2440433, 3-[4-(2,3-dimethylphenyl)piperazine-1-carbonyl]-N,N-dimethyl-
4-pyrrolidin-1-ylbenzenesulfonylamide). SRE and SRF luciferase assays were used as read-outs for studying LPI and ML184 signaling at the GPR55 mutants. A GPR55 R* model based on the recent DOR crystal structure was used to interpret the resultant mutation data. Two residues were found to be crucial for agonist signaling at GPR55, K2.60 and E3.29, suggesting that these residues form the primary interaction site for ML184 and LPI at GPR55. Y3.32F, H(170)F and F6.55A/L mutation results suggested that these residues are part of the orthosteric binding site for ML184, while Y3.32L, M3.36A and F6.48A mutation results suggest the importance of a Y3.32/M3.36/F6.48 cluster in the GPR55 signaling cascade. C(10)A and C(260)A mutations suggest that these residues form a second disulfide bridge in the extracellular domain of GPR55, occluding ligand extracellular entry in the TMH1-TMH7 region of GPR55. Taken together, these results provide the first set of discrete information on residues important for LPI and ML184 signaling and for GPR55 activation. This information should aid in the rational design of next generation GPR55 ligands and hopefully the creation of the first high affinity GPR55 radioligand, a tool that is sorely needed in the field.

**Introduction**

GPR55 [GenBank accession number NM005683] is a Class A G-protein coupled receptor (GPCR) that recognizes a sub-set of cannabinoid CB1 and CB2 ligands, suggesting that GPR55 may be a cannabinoid receptor. Subsequently, 1-α-lysophosphatidylinositol was reported to be the endogenous ligand of GPR55. GPR55 has been found to be implicated in inflammatory pain, neuropathic pain, metabolic disorder, bone development, and cancer, indicating the real potential of GPR55 ligands
as therapeutics. In search of more potent GPR55 ligands and in collaboration with the Sanford-Burnham Institute, we performed high throughput screens for GPR55 agonists and antagonists using the NIH library of 300,000 compounds and identified several novel GPR55 chemotypes for each.4 One of these compounds from the agonist screen is ML184, (CID 2440433, 3-[4-(2,3-dimethylphenyl) piperazine-1-carbonyl]-N,N-dimethyl-4-pyrrol-idin-1-ylbenzenesulfonamide)5 (EC50=263nM). However, despite these ligand development efforts, the best GPR55 ligands (like ML184) remain active at sub-micromolar concentrations, concentrations not low enough for the development of a GPR55 radioligand.

To address this situation, we reasoned that knowledge of GPR55 structure, particularly binding pocket residues important for ligand signaling and those involved in the receptor activation would greatly aid rational GPR55 drug design approaches. To this end, we report here a GPR55 mutation study guided and analysed using a GPR55 R* model based on the recent delta opioid receptor (DOR) crystal structure. This study identifies multiple residues important for ML184 signaling and for GPR55 activation.

Results

Biological Evaluation

**SRE responses of mutant and wild-type GPR55 receptors.** HEK293 cells transiently transfected with both pGL4.33 [luc2P/SRE/Hygro] and each of the mutants, as well as, WT were made to assess agonist-induced receptor activation of SRE. The SRE assay assesses the contribution of MAPK/ERK signaling pathway for GPR55 activity.
We found that the WT GPR55 has an EC\textsubscript{50} value of 56 nM for the agonist ML184 (Figure 30 and Table 2).

Figure 30. SRE Responses Induced by ML184 in GPR55 Wild Type and Mutant Transfected Cells. (A) WT GPR55, Y3.32F and Y3.32L. (B) WT GPR55, F6.55A and F6.55L. (C) WT GPR55, Q6.58M and C(10)A. (D) WT GPR55, Q7.36A and Q7.36N. (E) WT GPR55, H(170)F and C(260)A. (F) WT GPR55, M3.36A, F6.48A, K2.60A, E3.29A and E3.29L.
Table 2

ML184 Induced SRE Responses of Wild-type and Mutant Transfected Cells

<table>
<thead>
<tr>
<th>ML184</th>
<th>EC50 (nM)</th>
<th>CI (nM)</th>
<th>Fold/WT</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>56</td>
<td>24 to 133</td>
<td>-</td>
</tr>
<tr>
<td>Y3.32F</td>
<td>1659</td>
<td>670 to 4110</td>
<td>30</td>
</tr>
<tr>
<td>Y3.32L</td>
<td>9514</td>
<td>718 to 126100</td>
<td>170</td>
</tr>
<tr>
<td>F6.55A</td>
<td>840</td>
<td>345 to 2045</td>
<td>15</td>
</tr>
<tr>
<td>F6.55L</td>
<td>5131</td>
<td>1034 to 25460</td>
<td>92</td>
</tr>
<tr>
<td>Q6.58M</td>
<td>72</td>
<td>22 to 237</td>
<td>1</td>
</tr>
<tr>
<td>C(10)A</td>
<td>635</td>
<td>197 to 2044</td>
<td>11</td>
</tr>
<tr>
<td>H(170)F</td>
<td>341</td>
<td>158 to 734</td>
<td>6</td>
</tr>
<tr>
<td>C(260)A</td>
<td>657</td>
<td>222 to 1941</td>
<td>12</td>
</tr>
<tr>
<td>Q7.36A</td>
<td>230</td>
<td>57 to 920</td>
<td>4</td>
</tr>
<tr>
<td>Q7.36N</td>
<td>154</td>
<td>69 to 342</td>
<td>3</td>
</tr>
<tr>
<td>M3.36A</td>
<td>900</td>
<td>290 to 2790</td>
<td>16</td>
</tr>
<tr>
<td>F6.48A</td>
<td>514</td>
<td>266 to 993</td>
<td>9</td>
</tr>
<tr>
<td>K2.60A</td>
<td>NO response</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>E3.29A</td>
<td>&gt;30000</td>
<td>very wide</td>
<td>-</td>
</tr>
<tr>
<td>E3.29L</td>
<td>&gt;30000</td>
<td>Very wide</td>
<td>-</td>
</tr>
</tbody>
</table>

Note. From \( n \geq 3 \) experiments

We determined that the EC50 value did not change regardless of the level of receptor expression (See Figure S1 in Supporting Information). Mutants Y3.32F and Y3.32L showed right shifted SRE response curves with EC50’s higher than WT (Figure 30 A). The mutants F6.55A and F6.55L induced a similar SRE response to ML184 with right shifted curves. The Q6.58M curve was almost identical to that of WT indicating this mutant is not important to GPR55 receptor function. The mutant C(10)A showed an increased EC50 value (636 nM). Very similarly, the mutant C(260)A induced an
increased EC₅₀ value (657 nM), suggesting the existence of a disulfide bridge (see below). Q7.36A and Q7.36N showed similar responses to WT indicating this amino acid is not critical for agonist induced GPR55 receptor activation. The response curve of H(170)F only showed a slight difference from that of WT. Mutant M3.36A, F6.48A, E3.29A and E3.29L all showed increased EC₅₀ values. E3.29A and E3.29L showed very little responses. Among all the mutants, K2.60A did not show any response in the SRE assay indicating that K2.60A rendered the GPR55 receptor non-functional.

In addition to the agonist ML184, the endogenous ligand (LPI) was used in the SRE assay for all the mutants (Figure 31). K2.60A did not show any LPI induced responses compared with WT. E3.29A and E3.29L showed a slight response at high concentration of LPI. F6.55A, F6.55L, Q6.58M, Q7.36A and Q7.36N all showed similar curves as to WT. Y3.32F, Y3.32L, C(10)A, H(170)F, C(260)A, M3.36A and F6.48A showed right shifted curve compared with WT. The LPI induced SRE response did not reach plateau for all the constructs including WT GPR55, thus no EC₅₀ value was calculated.

**SRF responses of mutant and wild-type GPR55 receptors.** In addition to the SRE assay, the SRF assay was used to investigate another signaling pathway of GPR55. The SRF response is an indicator of G12-RhoA pathway activation which GPR55 has been reported to induce.⁶ The study of two distinct signaling pathways of GPR55 broadens the understanding of effects of these mutants on GPR55 function. Thus, the SRF assay was used to test GPR55 mutants in addition to the SRE assay (Figure 32 and Figure 33).
Figure 31. SRE Responses Induced by LPI in GPR55 Wild Type and Mutant Transfected Cells. (A) WT GPR55, Y3.32F and Y3.32L. (B) WT GPR55, F6.55A and F6.55L. (C) WT GPR55, Q6.58M and C(10)A. (D) WT GPR55, Q7.36A and Q7.36N. (E) WT GPR55, H(170)F and C(260)A. (F) WT GPR55, E3.29A and E3.29L. (G) WT GPR55 and M3.36A. (H) WT GPR55, F6.48A and K2.60A.
Figure 32. SRF Responses Induced by ML184 in GPR55 Wild Type and Mutant Transfected Cells. (A) WT GPR55, Y3.32F and Y3.32L. (B) WT GPR55, F6.55A and F6.55L. (C) WT GPR55, C(10)A and C(260)A. (D) WT GPR55, M3.36A, F6.48A, K2.60A, E3.29A and E3.29L.
Figure 33. SRF Responses Induced by LPI in GPR55 Wild Type and Mutant Transfected Cells. (A) WT GPR55, Y3.32F and Y3.32L. (B) WT GPR55, F6.55A and F6.55L. (C) WT GPR55, C(10)A and C(260)A. (D) WT GPR55, E3.29A and E3.29L. (E) WT GPR55 and M3.36A. (F) WT GPR55, F6.48A, K2.60A.
Y3.32F and Y3.32L showed right shifted curves compared with WT. Y3.32L had a higher EC50 value (864 µM; Figure 32). Similar to the SRE assay, the F6.55A and F6.55L had higher EC50 values than WT (Figure 32, Table 3). Both C(10)A and C(260)A showed less response than WT. M3.36A, F6.48A, E3.29A and E3.29L all behaved in the SRF assay similar to in the SRE assay with reduced responses. K2.60A was not able to induce any response in the SRF assay similar to its inactivity in the SRE assay.

Table 3
ML184 Induced SRF Responses of Wild-type and Mutant Transfected Cells

<table>
<thead>
<tr>
<th>ML184</th>
<th>EC50 (nM)</th>
<th>CI (nM)</th>
<th>Fold/WT</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>33</td>
<td>15 to 75</td>
<td>-</td>
</tr>
<tr>
<td>Y3.32F</td>
<td>851</td>
<td>449 to 1615</td>
<td>26</td>
</tr>
<tr>
<td>Y3.32L</td>
<td>864000</td>
<td>very wide</td>
<td>26182</td>
</tr>
<tr>
<td>F6.55A</td>
<td>550</td>
<td>338 to 893</td>
<td>17</td>
</tr>
<tr>
<td>F6.55L</td>
<td>10500</td>
<td>4663 to 23630</td>
<td>318</td>
</tr>
<tr>
<td>C(10)A</td>
<td>554</td>
<td>254 to 1206</td>
<td>17</td>
</tr>
<tr>
<td>C(260)A</td>
<td>363</td>
<td>201 to 656</td>
<td>11</td>
</tr>
<tr>
<td>M3.36A</td>
<td>2044</td>
<td>1415 to 2952</td>
<td>62</td>
</tr>
<tr>
<td>F6.48A</td>
<td>335</td>
<td>177 to 633</td>
<td>10</td>
</tr>
<tr>
<td>K2.60A</td>
<td>NO response</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E3.29A</td>
<td>&gt; 30000</td>
<td>very wide</td>
<td>-</td>
</tr>
<tr>
<td>E3.29L</td>
<td>&gt; 30000</td>
<td>very wide</td>
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*Note. n ≥ 3 experiments.*

**Molecular Modeling**

No x-ray crystal structure for GPR55 has been reported. For this reason, a GPR55 activated state (R*) model based upon the crystal structure of the delta-opioid receptor
(DOR) was used to interpret the mutation results reported here. Glide docking studies in this model focused on ML184. These studies were used to identify a docking site for ML184 in the GPR55 R* model that involves residues on TMHs2-3-5-6 and the EC-2 loop.

![Figure 34. GPR55 Agonist ML184, Numbered to Show Torsional Angles Varied for Conformational Analysis.](image)

**Primary interactions for agonist signaling.**

**ML184 docked into the GPR55 R* bundle.** Figure 35 describes the ML184 binding site identified by Glide. Figure 35 A shows all hydrogen bonding interactions for ML184 at GPR55. Here, the ML184 sulfonamide oxygen hydrogen bonds directly with K2.60 (H bond (N-O) distance, 2.8 Å; (N-H—O) angle, 174°). H(170) in the EC-2 loop also forms a hydrogen bond with an ML184 sulfonamide oxygen (H bond (N-O) distance, 2.8 Å; (N-H—O) angle, 150°). Y3.32 forms a hydrogen bond with the ML184 carboxamide oxygen (H bond (N-O) distance, 2.8 Å; (N-H—O) angle, 163°).
Figure 35. All Hydrogen and Aromatic Stacking Interactions for ML 184 at GPR55. (A) This Figure Shows All Hydrogen Bonding Interactions for ML184 at GPR55. The View is Looking from TMHs 4 and 5 towards TMH7. Hydrogen Bonding is Shown with Dotted Yellow Lines. (B) This Figure Shows All Aromatic Stacking Interactions for ML184 at GPR55. The View is Looking from TMH 4 towards TMH6. Aromatic Stacking Interactions are Shown with Light Blue Dotted Lines between Ring Centroids.
In addition to hydrogen bonding, ML184 also has aromatic stacking interactions in its binding site. Figure 35 B shows all of these aromatic stacking interactions. Y3.32 and F3.33 form aromatic stacks with the ML184 benzene ring proximal to the ML184 sulfonamide moiety (Y3.32 ring centroid to centroid distance 5.6 Å; angle 70°); (F3.33 ring centroid to centroid distance 6.5 Å; angle 47°). F6.55 forms an aromatic stacking interaction with the distal, dimethyl-phenyl ring of ML184 (ring centroid to centroid distance 5.4 Å; angle 88°), while F5.39 and F5.47 also form aromatic stacks with the central ML184 aromatic ring (F5.39 ring centroid to centroid distance 5.3 Å: angle 70°; F5.47 ring centroid to centroid distance 4.5 Å: angle 41°).

M7.39 is engaged in a Met-aromatic ring interaction with the central benzene of ML184 (Distances to central benzene ring centroid from M7.39 sidechain atoms CG 5.1 Å, SD 5.3 Å, CE 4.4 Å) (see Figure 36 A). In this orientation, the sulfur points up, with adjacent carbons pointing down towards phenyl ring carbons on ML184. ML184 also has an indirect interaction with E3.29, as this residue holds K2.60 (with which ML184 has a direct hydrogen bond) in a salt bridge that directs K2.60 towards the ML184 binding site (see Figure 36 B). The ML184 conformer docked here is 1.91 kcal/mol above the ML184 global minimum.
Figure 36. (A) M7.39 is engaged in a Met-aromatic Ring Interaction With the Central Benzene of ML184. In this orientation, the sulfur points up, with adjacent carbons pointing down towards phenyl ring carbons on ML184. (B) Although there is no direct interaction between ML184 and E3.29 in the current model, modeling suggests that E3.29 forms a salt bridge with K2.60. This salt bridge positions K2.60 for interaction with ML184. The view here is from TMH6 looking towards TMH2/3.
**Toggle switch residues.** GPR55 shares with the P2Y12\(^9\) and hPAR-1\(^{10}\) receptors, a Y/F3.32-M3.36-F6.48 cluster of residues that likely acts as the toggle switch for activation of GPR55. Figure 37 illustrates these toggle switch residues contoured at their Van der Waals radii in the inactive (R) and the activated (R\(^*\)) state.

Figure 37. Positions of the Toggle Switch Residues Y3.32, M3.36, and F6.48 (Contoured at Their van der Waals Radii) in the GPR55 R and R\(^*\) Models. Additional Aromatic Residues in the Region of the Toggle Switch Help Stabilize the Cluster of Residues. These are F3.33 and Y3.37.

**Inactive state.** In the inactive state (Figure 37 A), Y3.32 (\(\chi_1=178^\circ\)) sits extracellular to M3.36 (\(\chi_1=-170^\circ\)), maintaining Van der Waals interactions with M3.36. M3.36 (\(\chi_1=-170^\circ\)) is stacked over F6.48 (\(\chi_1=-90^\circ\)). M3.36 also has a methione/aromatic ring interaction with F6.48 (Distances to F6.48 benzene ring centroid from M3.36 sidechain atoms CG 3.6 Å, SD 4.6 Å, CE 3.8 Å). Additional aromatic residues in the
region of the toggle switch help stabilize the cluster of residues. These are F3.33 \( (\chi_1 = -74^\circ) \) and Y3.37 \( (\chi_1 = -57^\circ) \).

**Agonist activated state.** ML184 binding causes Y3.32 to change conformations \( (\chi_1 = 178^\circ \text{ to } \chi_1 = -157^\circ) \), because Y3.32 is a direct ligand binding site residue. Figure 37 B illustrates that the movement of Y3.32 frees M3.36 to undergo a conformational change \( (\chi_1 = -170^\circ \text{ trans to } \chi_1 = -75^\circ \text{ g}^+) \), which then allows F6.48 to undergo its \( (\chi_1 = -90^\circ \text{ g}^+ \text{ to } \chi_1 = -177^\circ \text{ trans}) \) conformational change. The F6.48 conformational change causes flexing in the SFXP hinge region of GPR55, straightening TMH6 and breaking the ionic lock (R3.50/Q6.30) producing an opening at the intracellular end of GPR55 for G protein coupling. F3.33 \( (\chi_1 = -81^\circ) \) and Y3.37 \( (\chi_1 = -55^\circ) \) remain members of the extended cluster. Figure 38 illustrates the position of ML184 to the extended toggle switch residues in the GPR55 activated state.

**Disulfide bridge residues.** As described in the Methods section, the high degree of sequence homology between GPR55 and the CXCR4 receptor, particularly in the EC regions, dictated several modifications to our initial set of GPR55 models. These were (1) the introduction of EC helical extensions on TMH5-7 of GPR55; (2) the introduction of a β sheet motif into the EC-2 loop in GPR55; and (3) the introduction of a disulfide bridge between Cys(10) in the N-terminus and Cys(260) in the EC-3 loop near the top of TMH7. In the current paper, we tested the importance of a Cys(10)-Cys(260) disulfide bridge via single point mutations. Figure 39 illustrates the extracellular end of the receptor and the disulfide bridge between N-terminal C(10) and C(260) at the EC end of TMH7.
Figure 38. Position of ML184 Relative to the Extended Toggle Switch Residues in the GPR55 Activated State. ML184 (Contoured at its Van der Waals Radii) is Positioned above Y3.32 and Interacts Directly with it. The Movement of Y3.32 for This ML184 Interaction, Permits M3.36 ($\chi_1$ trans to $g^+$) and, in Turn, F6.48 ($\chi_1$ $g^+$ to trans) to Change to Their R* Conformations.

Figure 39. The Extracellular End of the Receptor and the Disulfide Bridge Between N-Terminal C(10) and C(260) at the EC End of TMH7.
**Binding pocket position.** Figure 40 illustrates the positions of Q6.58 and Q7.36 relative to the ML184 binding site. These residues are positioned relatively high on the EC end of TMHs 6 and 7 and above the ML184 binding pocket in our current GPR55 R* model. In our first GPR55 model, Q6.58 was near the nitrogen in the pendant five membered ring of ML184 and able to form a hydrogen bonding interaction with this nitrogen. As discussed in the next section, mutation results suggested that Q6.58 and Q7.36 do not interact with ML184. This necessitated a re-orientation of ML184 in the binding pocket in the current model.

Figure 40. Positions of Q6.58 and Q7.36 Relative to the ML184 Binding Site. Both Residues are Clearly Above the ML184 Binding Site.
Discussion

MI184 Docked into the GPR55R* Bundle

When one begins the characterization of the ligand binding pocket in a newly de-orphanized receptor, like GPR55, mutation studies are crucial to understanding the binding pocket. Our original GPR55 models (R and R*) were based on a beta-2-adrenergic receptor template. In this model, we docked ML184 so that the ligand piperazine and di-methyl phenyl ring segment was vertical in the binding pocket, while the rest of the ligand occupied a horizontal space near the extracellular loops. Mutation results reported here clearly necessitate that a reorientation of ML184 occur in the binding pocket. In the refined model reported here, we also took advantage of crystal structures that were unavailable at the time of the original model creation and found that the delta-opioid receptor (DOR) crystal structure was a more appropriate template for the next generation model. The DOR, for example has a PRO at position 4.59, as does GPR55. The beta-2-adrenergic receptor, on the other hand, has a PRO shifted by one residue to 4.60. In addition, the DOR has the same toggle switch partner as GPR55, M3.36.

Ligand Interactions within the Binding Crevice

As predicted by previous studies conducted on the GPR55 receptor in the Reggio lab, the current model has a crucial hydrogen bonding interaction between ML184 and K2.60 on TMH 2 (see Figure 35A or 35B) that results in the strongest ligand Interaction Energy (-9.4 kcal/mol) of any residue in the GPR55 R* bundle (see Table S-1 in Appendix B). A strong hydrogen bond between the electronegative sulfonamide and the
positive lysine is consistent with the experimental results that show a complete loss of signaling upon the K2.60 mutation to an alanine (see Tables 2 and 3).

Mutation of E3.29 to either an alanine or a leucine resulted in a profound reduction of receptor signaling as well. Though there is no direct interaction between ML184 and E3.29 in the current model, modelling suggests that E3.29 forms a salt bridge with K2.60 (see Figure 36 B). This salt bridge positions K2.60 for interaction with ML184. The loss of this residue’s directing capability for K2.60 via the E3.29/A/L mutations effectively make K2.60 less available in the binding pocket, leading to a considerable loss of function.

At the ML184 binding site, the EC-2 loop residue, H(170), serves as a hydrogen bond donor to the ML184 sulfonamide oxygen (see Figure 35 A). The Interaction Energy for ML184 with H(170) is -3.5 kcal/mol (see Table S-1 in Appendix B). Simultaneously, the sulfonamide oxygen also receives a hydrogen bond from K2.60. Because hydrogen bonding is stronger when the donating residue is charged, the H170 hydrogen bond with ML184 is weaker (-3.5 kcal/mol) than that with K2.60 (-9.1 kcal/mol) (see Table S-1 in Appendix B). The pKa of histidine is 6.0. At physiological pH, about 10% of histidine residues are protonated. We assume here that H(170) is uncharged. An H(170)F mutation removes the hydrogen bonding ability of residue 170, but not its ability to form aromatic interactions. Mutation of H(170) to a phenylalanine resulted in a 6-fold reduction in SRE responses (see Table 2). The magnitude of the effect upon mutation is consistent with the loss of a hydrogen bond. In addition, the fact that H(170) mutation affected ML184’s signaling suggests indirectly that there is also a C(168)/C3.25 disulfide bond in GPR55.
This bond requires that the EC-2 loop extend over to TMH3 to link with C3.25 and therefore is responsible for the H(170) location in the GPR55 R* model. In addition, inspection of GPCR crystal structures that contain the analogous disulfide bridges reveal that the second residue after the disulfide bridge typically points down into the binding crevice. H(170) is the second residue after the C(168)/C3.25 disulfide bridge.

Y3.32 serves two functions in GPR55. It is a binding site residue (see Figure 35 A and B) and also part of the extended toggle switch for GPR55 activation (see Figure 37). A stepwise loss of function is seen when Y3.32 is first mutated to a phenylalanine (30-fold loss, see Table 2) and then a leucine (170-fold loss, see Table 2). At the ML184 binding site, Y3.32 donates a hydrogen bond to the ML184 carboxamide oxygen (Figure 35 A) and forms an aromatic stacking interaction with the ML184 central benzene ring (Figure 35 B) proximal to the ML184 sulfonamide moiety. The first mutation, Y3.32F, removes the residue’s potential to donate a hydrogen bond. The magnitude of the effect (30-fold loss) is consistent with Y3.32 serving as the lone hydrogen bond donor to ML184 in this region. The second mutation, Y3.32L, removes both hydrogen bonding and aromatic stacking interactions from the ML184 binding site at this position. The 170-fold reduction in EC₅₀ (see Table 2) is consistent with the loss of two important interactions for ML184. Y3.32F mutation results are also consistent with Y3.32’s participation in the extended toggle switch region in GPR55, as loss of aromaticity at 3.32 in the Y3.32L mutant should have a profound effect on signaling (170-fold reduction in EC₅₀; see Table 2).
The current model of GPR55 places F6.55 facing inward towards the binding crevice. This residue forms an aromatic stacking interaction with the distal, dimethyl-phenyl ring of ML184 (Figure 35 B; Interaction Energy = -5.1 kcal/mol; see Table S-1 in Appendix B). The F6.55A mutation resulted in a 15-fold loss in EC₅₀ for ML184, consistent with the loss of this aromatic stacking interaction. The F6.55L resulted in an even larger loss in EC₅₀ (92-fold). This mutation not only removes an aromatic stacking interaction, but also causes crowding in the ML-84 binding pocket. This crowding causes ML184 to shift position and the net result would be reduced binding site interactions. The 92-fold loss in EC₅₀ is consistent with such an alteration.

The x-ray crystal structure of the beta-1-adrenergic receptor complexed with Gs protein shows that TMH6 has straightened by flexing its proline kink at P6.50.¹⁵ This conformational change also impacts binding pocket residues. W6.48 has been shown to change its conformational state within the binding pocket upon receptor activation (χ₁ g⁺ → trans). In the inactive state, W6.48 is typically held in its χ₁=g⁺ conformation by another binding pocket residue. Together, this pair of residues is known as the “toggle switch.”¹⁶ In the GPR55 activated state model reported here, Y3.32, M3.36 and F6.48 form an extended toggle switch (see Figure 37), with additional interactions from F3.33 and Y3.37. The mutation effects of Y3.32 are discussed above in the context of their effects on the ML184 binding pocket interactions and upon signaling. For both the SRE and SRF readouts used here, mutation of M3.36 to alanine had a significant impact on signaling (16-fold, 62-fold). Mutation of the toggle switch residues M3.36 and F6.48 resulted in increased EC₅₀ values, but these mutants were able to ultimately reach SRE
induction comparable to wild-type. As ML184 is a GPR55 agonist, it is expected that its position in the bundle would not cause it to interfere with the receptor’s ability to activate. This is verified by the experimental data and is what is observed in the current ligand receptor complex.

Class A GPCRs typically have a disulfide bridge between a residue in the EC-2 loop and Cys3.25 near the top of TMH3. The sequence of GPR55 suggests that it likely has this disulfide bridge as well. Mutation studies of these Cys residues typically result in loss of function as the EC-2 – C3.25 disulfide bridge is important for binding pocket structure. The CXCR4\(^1\)\(^1\)\(^1\) and CCR5\(^1\)\(^\text{19}\) crystal structures reveal the existence of a second disulfide bridge that links the N terminus of the receptor to the EC end of TMH7, forming what has been called the fourth EC loop.\(^1\)\(^\text{19}\) This loop has been proposed to shape the entrance of the ligand-binding pocket and to add rigidity to the overall surface of the receptor.\(^1\)\(^\text{19}\) The sequences of ~30% of Class A GPCRs contain such Cys residues, including the lysophospholipid (LPA), bradykinin (B1-2), endothelin (ETA-B), melanocortin (MC1-5), serotonin (5-HT), purinergic (P2Y), and orphan receptors, such as GPR55. The structure of one of the latest resolved rhodopsin-like receptors, P2Y12, revealed the presence of such an EC-4 loop;\(^9\) however, the conservation of these residues does not necessarily imply the formation of a EC-4 loop (see crystal structures of dopamine D3\(^2\)\(^\text{0}\) and serotonin, 5HT1B\(^2\)\(^\text{1}\) receptors). Because the GPR55 sequence suggests the presence of an EC-4 loop, this second disulfide bridge was incorporated into our GPR55 model and was tested via C(10)A and C(260)A mutations here. Results reported here show that each mutation impacts ML184 activation of GPR55, although
these mutations are not devastating (11-fold for C(10)A; 12-fold for C(260)A). We have taken these results as evidence that this disulfide bridge is present in GPR55 and have retained this bridge in our model (see Figure 39). It should be noted that the EC-4 loop may have larger effects on GPR55 agonists other than ML184, as it has been proposed that this fourth loop shapes the entrance of the ligand-binding pocket.19

Q6.58 and Q7.36 are located at the extreme EC ends of TMH6 and TMH7 (see Figure 40). In our first GPR55 model, Q6.58 was near the nitrogen in the pendant five membered ring of ML184 and able to form a hydrogen bonding interaction with this nitrogen.12 A Q6.58M mutation should have resulted in the loss of this hydrogen bond and a significant reduction in ML184 EC50. However, at the Q6.58M/A mutations, ML184 retained WT signaling. This result is a key result from a modelling perspective because it clearly suggested that a different binding mode for ML184 should be sought.

Also in our previous GPR55 R model, Q7.36 interacted simultaneously with K2.60 and with antagonist, ML192.13 The re-orientation of ML184 dictated by the Q6.58M/A mutations dictated that Q7.36 should have no ligand interactions. Consistent with this result, experimental data reported here shows that at both the Q7.36A and Q7.36N mutant, ML184 retains WT signaling.

Quite recently, additional synthetic ligands, as well as another endogenous agonist for GPR55 have been described.22 The synthetic ligands employed our previous model as a guide for ligand design. The mutation results reported here will be important for the design of more potent and efficacious agonists for GPR55.
Conclusions

Results reported here identify key GPR55 residues that are important for agonist signaling, as well as residues implicated in the agonist activated signaling cascade. Two residues crucial for ML184 signaling at GPR55 are K2.60 and E3.29. Three additional residues, Y3.32, H(170) and F6.55, are important for ML184 signaling. Further, results suggest that a cluster of residues, F3.32/M3.36/F6.48 serves as the toggle switch for activation of GPR55. GPR55 also likely possesses, a second disulfide bridge that links the N terminus of the receptor to the EC end of TMH7. This loop has been called the fourth EC loop.19 All of these results provide, for the first time, structural information that should aid in the rational design of next generation GPR55 ligands. It is hoped that this will lead to a high affinity GPR55 radioligand, a tool that is sorely needed in the field.

Experimental Section

Material and Reagents

Soy LPI (purchased from Sigma-Aldrich), and ML184 (MolPort) were dissolved in dimethyl sulfoxide to a concentration of 10 mM. The SRE reporter, pGL4.33[luc2P/SRE/Hygro] and the SRF-RE reporter, pGL4.34[luc2P/SRF-RE/Hygro] were from Promega.

Mutagenesis and Cell Culture

directed mutagenesis kit (Stratagene). DNA sequencing subsequently confirmed the presence of the desired mutation only.

**Serum Response Element (SRE) and Serum Response Factor (SRF) Assay**

HEK293 cells were transiently transfected with GPR55 and pGL4.33 [luc2P/SRE/Hygro] or pGL4.34 [luc2P/SRF-RE/Hygro] vector reporter plasmids using Lipofectamine 2000 as described by the manufacturer (Invitrogen). Transfected HEK293 cells were seeded (60,000 cells per well) in 96-well plates. Five hours later, medium was changed to 1% FBS/DMEM. Cells were incubated overnight. The next day cells were treated with ligands for 5 h in serum-free DMEM medium at 37°C. After treatment, cells were lysed by 1X lysis buffer for 10 min at room temperature. Plates were read to record bioluminescent light immediately after the injection of 40µl Luciferin (≥250µM) per well. Luminescence was measured in an Envision 2104 multilabel Reader (PerkinElmer). Luminescence values are given as relative light units. Concentration-effect curves for agonist-mediated receptor activation were analyzed by nonlinear regression techniques using GraphPad Prism 5.0 software (GraphPad) and data were fitted to sigmoidal dose-response curves to obtain EC50 values.

**Modeling**

**Amino acid numbering system.** The amino acid numbering system used here is the Ballesteros-Weinstein numbering system in which the most highly conserved residue across Class A GPCRs in each TMH is assigned a number .50. This number is preceded by the TMH number and can be followed by the absolute sequence number in parentheses. For example, the most highly conserved residue in TMH4 is W4.50. For
GPR55, this residue is W4.50(146). The residue preceding this residue is I4.49(145) and the residue following it is V4.51(147). Loop residues in this system are identified by their absolute sequence numbers only.

**Modeling of hGPR55 active-state bundle (GPR55 R*) using GPCR x-ray crystallography data.** The model of the activated form of GPR55 (GPR55 R*) described in the current work was created using the 1.8 Å crystal structure of the human delta opioid receptor (hDOR, PDB id: 4N6H) as a template. Transmembrane regions of GPR55 vs. hDOR in which the placement of prolines differed (GPR55 transmembrane helices (TMHs) 1, 5, 6 and 7) were explored using the Conformational Memories (CM) method described below.

**Conformational Memories (CM) Method for Calculating TMH Conformation**

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 (i to i-4 of a proline) were allowed to vary +/- 50°, while all other backbone torsion angles were allowed to vary +/- 10°. Side chain torsions were allowed to vary +/- 180°. All bond angles were allowed to vary +/- 8° except for C-S-C angles that were allowed to vary +/- 15°. A minimum set of 108 conformers was generated for each GPR55 helix, independently, in a distance dependent dielectric at 310 K.

**TMH1** in GPR55 has a Pro at position 1.41 not found in DOR. An ideal helix (φ = -62.9°, and ψ = -41.6°) with the GPR55 sequence was built, and the region containing the proline and 4 residues prior to the Pro (i to i-4, A1.37 - P1.41) was varied using CM.
**TMH5** in GPR55 has two prolines, the highly conserved P5.50 and an additional one at position 5.41. TMH5 of the DOR crystal structure was mutated to the GPR55 sequence and the backbone dihedrals from P5.41 to K5.37 were varied to explore the possible conformations caused by this second, proline.

**TMH6** in GPR55 has a conservative SFXP substitution in place of the highly conserved Class A CWXP motif. Biophysical studies have indicated that there is a salt bridge, or ionic lock (between R3.50 near the intracellular (IC) end of TMH3 and D/E6.30 at the IC end of TMH6), common to all Class A GPCRs that is broken upon activation. The breaking of this ionic lock allows TMH6 to straighten, moving its intracellular (IC) end away from the TMH bundle. For this reason, it was crucial to explore the conformational space of GPR55 TMH6 which has a glutamine (Q) at position 6.30 in place of a typical aspartic (D) or glutamic (E) acid. The GPR55 TMH6 sequence was built, using the ideal helix values ($\phi = -62.9^\circ$, and $\psi = -41.6^\circ$) and the i to i-4 region around the Pro (P6.50-V6.46) was varied using CM.

**TMH7** in GPR55 lacks the highly conserved Class A GPCR NPXXY motif having instead a DVXXY sequence. Traditionally the NPXXY motif influences the conformation of TMH7 and places Y7.53 in the correct position to interact with F7.60 on Helix 8 (Hx8, a short intracellular extension of TMH7 that lies usually parallel to the cell membrane). For receptors that possess an NPXXY motif, we have typically designated the backbone region of P7.50- X7.46 region as variable. For GPR55 TMH7, an ideal helix ($\phi = -62.9^\circ$, and $\psi = -41.6^\circ$) was built and the i to i-4 region in which there is normally a Pro (V7.50 - C7.46) was varied using CM.
Construction of the GPR55 Active-state Bundle

Transmembrane helices 2, 3 and 4 of the DOR were mutated to the corresponding GPR55 residues, while helices chosen from the CM output that would fit in the bundle were substituted in the bundle. This included a straightened TMH6 conformer for which the TMH3-TMH6 ionic lock was broken, reflecting the R* state. Incorporated into this new bundle were updates from previous homology models of GPR55\textsuperscript{13-14} created in this lab: The high degree of sequence homology between GPR55 and the CXCR4 receptor,\textsuperscript{11} particularly in the EC regions, dictated several modifications to the model. These were (1) the introduction of
EC helical extensions on TMH5-7 of GPR55; (2) the introduction of a β sheet motif into the EC-2 loop in GPR55; and, (3) the introduction of a disulfide bridge between Cys(10) in the N-terminus and Cys(260) in the EC-3 loop near the top of TMH7.

The resulting homology model was then optimized using the following protocol\textsuperscript{13}: The energy of the GPR55 R* bundle was minimized using the OPLS 2005 force field in Macro-model 9.9 (Schrödinger Inc., Portland, OR). An 8.0-Å extended non-bonded 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 minimization was performed in three stages. Each stage consisted of a Polak- Ribier conjugate gradient minimization in 1000-step increments until the bundle reached a 0.05 kJ/mol gradient. In the first stage of the calculation, the TMH region of the receptor was held stationary and the loops were allowed to relax using the generalized born/surface area continuum solvation model for water (Macro-model). In the second stage, the loops were frozen and the side chains of
the TMHs were allowed to adjust. A distance dependent dielectric was used for this minimization. In the third stage, the N and C termini were minimized using the protocol described, for the loops, above. In this stage, only the termini were minimized.

**Conformational Analysis of ML184**

A complete conformational analysis of the GPR55 agonist ML184 (CID2440433) was performed using *ab initio* Hartree-Fock (HF) calculations at the 6–31G* level as encoded in Jaguar (version 9.0, Schrodinger, LLC, New York, NY). Hartree-Fock 6–31G* six-fold conformer searches were performed for the rotatable bonds, N1-S2, S2-C3, C4’-N5’, C4-C5, C5-N6, N7-C8 (see numbering system in Fig 29), of ML184 as follows: 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 energy difference between the global minimum energy conformer of ML184 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. This difference was calculated to be 1.91 kcal/mol.

**Docking of ML184**

A low free-energy conformer of ML184 was used as input for receptor docking. ML184 was initially docked manually in the binding site of the GPR55 R* model, after which the automatic docking program, Glide (Schrodinger Inc.), was used to explore other possible binding conformations and receptor site interactions. Extra precision (XP)
and flexible docking with ring sampling were selected for the docking setup. Lysine 2.60(80), a previously identified ligand interaction site,\textsuperscript{12} was defined as a hydrogen bond donor and was used as a constraint for the automatic docking of the ligand. The energy of the ligand/GPR55 R* complex was minimized using the OPLS 2005 force field in Macromodel 9.9 (Schrödinger Inc., Portland, OR). An 8.0-Å extended non-bonded 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 as described previously. The dock with the best Glide score (-9.1) was selected as the final ML184/GPR55 R* model.

**Assessment of Pair-wise Interaction Energies**

After defining the atoms of ML184 as one group (Group 1) and the atoms corresponding to a residue that lines the binding site in the final ligand/GPR55 R* complex as another group (Group 2), Macromodel (version 8.6, Schrödinger, LLC, New York, NY) was used to output the pair-wise interaction energy (coulombic and Van der Waals) for a given pair of atoms. The pairs corresponding to Group 1 (ligand) and Group 2 (residue of interest) were then summed to yield the interaction energy between the ligand and that residue. The ML184/GPR55 R* complex was found to have interaction energies totaling to -55.9 kcal/mol. Taking the conformational energy cost for ML184 (1.91 kcal/mol) into account, the final total energy for the ML184/GPR55 R* complex was found to be -54.0 kcal/mol. A breakdown of interaction energies is provided in Table S-1.
Acknowledgments

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References


CHAPTER V

FUTURE CONSIDERATIONS

Now that there are models of the GPR55 active and inactive states it is imperative that the models continue to be refined. It is also necessary that they be used as the tools that they were designed to be to suggest new directions for ligand development.

Chemo-informatics Approach

A shift in focus to other scaffolds will likely be needed owing to the fact that the analogs synthesized thus far have not reached low nanomolar efficacies. One approach to scaffold diversification is receptor-based core hopping. New scaffolds for both GPR55 agonists and antagonists can be explored using the Schrodinger suite module CoreHopper. This task option in the Schrodinger small-molecule discovery suite allows for rapid screening of novel cores to help overcome undesirable properties of a ligand by creating new lead compounds with improved core properties while preserving key R-group interactions. Core hopping can also potentially be used for generation of novel derivatives to a known ligand (such as LPI). Chemotypes produced by the CoreHopper program can be used to explore the 102 million-compound library of commercially available compounds in the IReasearch library in ChemNavigator. Once a subset of structures has been identified, these structures can then be converted to three-dimensional form, energy minimized and then subjected to high throughput docking in the updated models using the Induced Fit Docking module discussed in Chapter III.
Validating Chemo-informatics Virtual Search Results

It would be necessary for select compounds from the virtual search to be purchased and evaluated in Dr. Abood’s lab. Antagonist compounds (initially at 10 μM) could be screened for their ability to inhibit the β-arrestin trafficking caused by the GPR55 agonist, LPI, using high content imaging by established methodology (described in Chapter II. Compounds found to inhibit LPI’s β-arrestin trafficking can then be 1) counter-screened for their ability to inhibit receptor internalization and 2) their ability to antagonize LPI induced ERK1/2 signaling in a 96-well plate format using a LI-COR Odyssey imager for quantification of responses. Agonist compounds (initially at 10 μM) will be screened for their ability to stimulate β-arrestin trafficking, using high content imaging as described in Chapter II. Compounds found to produce β-arrestin trafficking will be counter-screened for their ability to produce receptor internalization as described above.

This would be the point in this endeavor where heretofore unassessed analogs could be designed using each new scaffold identified. If a large number of compounds is generated and found to be active, factors such as cost and future synthetic practicality will be used to filter the final range of compounds to be assessed.

Unutilized Synthesis of Analogues for Scaffold 2

The structure of Scaffold 2 (from the original Burnham Screen mentioned in Chapters I–IV previously) can be divided into two primary areas, the furyl amide and a tricyclic heteroaromatic group, either of which could be modified (Figure 41). The proposed synthetic plans are shown in Scheme 3 (Figure 42). The amide section of 18 can
be connected using the corresponding acid chloride,\(^1\) and there are a multitude of variants possible at this position. The fused, heteroaromatic section can be put together from readily available ketones, ethyl cyanoacetate, and elemental sulfur (S\(_8\)) using the Gewald synthesis.\(^2\) The resulting 2-aminothiophene can be reacted with a nitrile under acidic conditions and then dehydrated using phosphoryl chloride to provide electrophilic coupling partners.\(^3\) These two sections can then be brought together to provide the desired analogues (21).\(^4\) This synthesis can be modified as needed to give rise to the different compounds, but the basic route can be followed for most of the structures proposed in Figure 43.

Figure 41. Scaffold 2.

![Scaffold 2](image)
Figure 42. Scheme 3: Synthetic Scheme for Scaffold 2.

Figure 43. Analogs Proposed for Scaffold 2 Development.
The analogs that should be targeted for synthesis initially are ones that would modify the furan group due to the fact that there are very few commercially available structures that extend out in this area of the binding pocket. This area of the binding pocket is of known importance due to mutation data highlighting ligand interactions with K2.60 and Y3.32. Compounds 25 and 26 were designed to have an intramolecular hydrogen-bond to rigidify the structure in the binding pocket and to allow other moieties on the ligand to explore available space. Alcohol 27 explores the potential for additional hydrogen bonds (hydrogen bond donation to S7.32 and hydrogen bond acceptance with Q6.58).

It is now known from the mutation data detailed in Chapter IV that Q6.58 does not interact with GPR55 agonists in the binding pocket as it sits high in the EC side of the receptor. It remains to be seen if that mutation would have an effect on antagonists of GPR55. Analogs in Figure 43 were proposed based on the fit they had with the original receptor model and so will need to be re-evaluated to confirm their validity in the current model.
References


APPENDIX A
SUPPLEMENTAL INFORMATION FOR CHAPTER II

Supporting Information for:

Design, Synthesis, and Analysis of Antagonists of GPR55: Piperidine-Substituted 1,3,4-Oxadiazol-2-Ones

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Graphical Abstract

General Information

Solvents and reagents were obtained from commercial sources and used without further purification; anhydrous solvents were dried following standard procedure, reaction progress was monitored by TLC (Silica gel 60 F254) glass plates visualized with UV light and permanganate stain. The anhydrous reactions were performed in oven dried glassware under nitrogen atmosphere. Chromatographic purification was performed using silica gel (60 Å, 32-63µm). NMR spectra were recorded in CDCl3 using a Bruker AVANCE DRX 300 spectrometer (300 MHz for ¹H), JEOL ECA spectrometer (500 MHz for ¹H and 125 MHz for ¹³C). Chemical shifts reported in ppm using tetramethylsilane as reference for the ¹H NMR and the residual solvent peak for ¹³C (77 ppm). The abbreviations used to describe peak splitting patterns are: s = singlet, d = doublet, t = triplet, sept = septet, dd = doublet of doublets, m = multiplet. Coupling constants, J, are reported in hertz (Hz). IR was obtained with Perkin Elmer FTIR Spectrometer One and Spectrometer 65 with ATR sampling accessories. Frequencies are
in cm$^{-1}$. High Resolution Mass Spectra were acquired on a ThermoFisher Scientific LTQ Orbitrap XL MS system.

**Synthesis of Hydrazides 3a-f**

![Chemical structures of hydrazides 3t to 3w](attachment:chemical_structures.png)

The hydrazides 3v-x were synthesized following the reported procedure in the literature and hydrazides 3t, 3u, 3y, and 3z were purchased from Acros. Compound 3v matched the data previously reported and 3w and 3x, are fully characterized below.

**Characterization of Hydrazides 3w and 3x**

![Chemical structure of 4-Trifluoromethyl-benzoic acid hydrazide (3w)](attachment:chemical_structure_3w.png)

4-Trifluoromethyl-benzoic acid hydrazide (3w):

White solid (2.54 g, 12.4 mmol, 75%).
\[^1\text{H} \text{NMR}\] (CDCl\(_3\), 400 MHz): \(\delta = 10.01\) (s, 1H), 8.01 (d, \(J = 8.2\) Hz, 2H), 7.81 (d, \(J = 8.2\) Hz, 2H), 4.59 (s, 2H).

\[^{13}\text{C} \text{NMR}\] (CDCl\(_3\), 100 MHz): \(\delta = 164.6, 137.1, 131.2\) (q, \(J = 31.5\) Hz, 1C), 127.9 (2C), 125.3 (q, \(J = 3.8\) Hz, 2C), 123.9 (q, \(J = 270.0\) Hz, 1C).

\[\text{IR: } 3335, 1621, 1577, 1536, 1504, 1318, 1136, 1112, 1065, 928, 865, 771, 690, 607\text{ cm}^{-1}.\]

\[\text{HRMS (ESI): } \text{C}_8\text{H}_8\text{F}_3\text{N}_2\text{O}^+ [\text{M}+\text{H}]^+ \text{ calculated: } 205.05832; \text{ found: } 205.05817.\]

\[\text{3-Trifluoromethyl-benzoic acid hydrazide (3x):}\]

White solid (85 mg, 0.42 mmol, 60%).

\[^1\text{H} \text{NMR}\] (CDCl\(_3\), 400 MHz): \(\delta = 10.1\) (s, 1H), 8.16 (s, 1H), 8.12 (d, \(J = 7.8\) Hz, 1H), 7.85 (d, \(J = 7.8\) Hz, 1H), 7.69 (t, \(J = 7.8\) Hz, 1H), 4.95 (s, 2H).

\[^{13}\text{C} \text{NMR}\] (CDCl\(_3\), 100 MHz): \(\delta = 164.5, 134.3, 131.1, 127.8, 129.3\) (q, \(J = 31.5\) Hz, 1C), 127.8 (q, \(J = 3.8\) Hz, 1C), 124.1 (q, \(J = 270.8\) Hz, 1C), 123.7 (q, \(J = 3.8\) Hz, 1C).

\[\text{IR: } 3305, 1639, 1615, 1518, 1482, 1437, 1321, 1308, 1283, 1161, 1119, 1090, 1070, 818, 691, 642, 594\text{ cm}^{-1}.\]

\[\text{HRMS (ESI): } \text{C}_8\text{H}_8\text{F}_3\text{N}_2\text{O}^+ [\text{M}+\text{H}]^+ \text{ calculated: } 205.05832; \text{ found: } 205.05809.\]

**General Method for Synthesis of Piperidones 2a-f**

Thionyl chloride (4 eq) was added dropwise to an oven-dried round bottomed flask containing commercially available acid 1a-f (0.212 mmol – 29.0 mmol) and
dichloromethane (0.3 M). The reaction was stirred at 50 °C for one hour. The resulting solution was concentrated under vacuum and re-dissolved in dichloromethane (20 mL, 0.11 M) and added dropwise to a round bottom flask at 0 °C containing 4-piperidone (1.26 eq), triethylamine (7.5 eq), and THF (0.06 M). After 15 minutes the solution was refluxed at 50 °C for 24 hours. The reaction was carefully quenched with saturated aqueous ammonium chloride and extracted with chloroform (3 x 20 mL). The resultant organic layers were combined and dried over Na₂SO₄, decanted and concentrated under vacuum. Product purification was achieved by silica gel flash chromatography (40% ethyl acetate/60% hexane).

**Characterization of Piperidones 2a-f**

![Structure of Piperidone 2a](image)

1-(1-Phenyl-cyclopropanecarbonyl)-piperidin-4-one (2a): White solid (798 mg, 3.3 mmol, 36%).

**^1H NMR** (CDCl₃, 400 MHz): δ = 7.35 - 7.26 (m, 2H), 7.25 - 7.21 (m, 3 H), 3.95 - 3.67 (br, 4H), 2.50 - 2.25 (br, 2H), 2.05 - 1.91 (br, 2H), 1.46 (dd, J = 6.9, 4.6 Hz, 2H), 1.24 (dd, J = 6.9, 4.6 Hz, 2H).

**^13C NMR** (CDCl₃, 100 MHz): δ = 207.4, 171.5, 140.3, 129.2 (2C), 127.0, 125.7 (2C), 44.5, 41.9, 40.8 (2C), 29.8, 14.8 (2C).
IR: 3008, 2873, 2359, 2341, 1714, 1633, 1427, 1316, 1270, 1232, 1188, 1079, 981, 760, 738, 698, 600, 572 cm\(^{-1}\).


1-[1-(4-Methoxy-phenyl)-cyclopropanecarbonyl]-piperidin-4-one (2b):
Pale yellow crystalline solid (2.51 g, 13.0 mmol, 52%).

\(^1\)H NMR (CDCl\(_3\), 500 MHz): \(\delta = 7.16\) (dt, \(J = 5.2, 2.9\) Hz, 2H), \(6.84\) (dt, \(J = 5.7, 2.3\) Hz, 2H), \(3.80 - 3.89\) (m, 2H), \(3.70 - 3.80\) (br m, 2H), \(3.78\) (s, 3H) \(2.39\) (br m, 2H), \(1.97\) (br m, 2H), \(1.39\) (dd, \(J = 6.9, 4.6\) Hz, 2H), \(1.16\) (dd, \(J = 6.9, 4.6\) Hz, 2H).

\(^{13}\)C NMR (CDCl\(_3\), 125 MHz): \(\delta = 207.5, 171.8, 158.6, 132.2, 127.2\) (2C), 114.5 (2C), 55.5, 41.8 (2C), 40.7 (2C), 29.1, 14.3 (2C).

IR: 2957, 1716, 1633, 1509, 1468, 1435, 1425, 1312, 1277, 1239, 1180, 1031, 979, 831, 811, 652, 560 cm\(^{-1}\).

HRMS (ESI): \(\text{C}_{16}\text{H}_{20}\text{NO}_3\) [M+H]\(^+\) calculated: 274.14377; found: 274.14304.
1-[1-(2,4-Dichloro-phenyl)-cyclopropanecarbonyl]-piperidin-4-one (2c):

White solid (540 mg, 1.73 mmol, 68%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta = 7.40$ (s, 1H), 7.20 - 7.27 (m, 2H), 3.68 (br m, 4H), 2.15 (br m, 4H), 1.67 (dd, $J = 6.9, 4.6$ Hz, 2H), 1.14 (dd, $J = 6.9, 4.6$ Hz, 2H).

$^{13}$C NMR (CDCl$_3$, 125 MHz): $\delta = 207.1, 171.0, 137.8, 137.5, 133.8, 130.3, 130.0, 127.7, 43.8$ (2C), 40.8 (2C), 28.9, 14.6 (2C).

HRMS (ESI): C$_{15}$H$_{15}$Cl$_2$NO$_2$ [M+H]$^+$ calculated: 312.05581; found: 312.05524.

![Chemical structure of 1-[1-(2,4-Dichloro-phenyl)-cyclopropanecarbonyl]-piperidin-4-one (2c)](image)

1-[1-(4-Chloro-phenyl)-cyclopropanecarbonyl]-piperidin-4-one (2d):

Pale yellow solid (5.06 g, 25.4 mmol, 21%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta = 7.29$ (d, $J = 8.6$ Hz, 2H), 7.18 (d, $J = 8.6$ Hz, 2H), 3.90-3.65 (br m, 4H), 2.41 (br m, 2H), 2.08 (br m, 2H), 1.47 (dd, $J = 6.9, 4.6$ Hz, 2H), 1.21 (dd, $J = 6.9, 4.6$ Hz, 2H).

$^{13}$C NMR (CDCl$_3$, 125 MHz): $\delta = 207.0, 171.1, 138.9, 132.8, 129.4$ (2C), 127.2 (2C), 44.5, 41.9, 40.8 (2C), 29.4, 15.0 (2C).

IR: 3009, 1726, 1685, 1491, 1388, 1371, 1338, 1247, 1178, 1096, 1045, 1014, 951, 931, 828, 762, 753, 716, 678, 544 cm$^{-1}$.

HRMS (ESI): C$_{15}$H$_{17}$ClNO$_2$ [M+H]$^+$ calculated: 278.09423; found: 278.09387.
1-(2,2-Dimethyl-1-p-tolyl-cyclopropanecarbonyl)-piperidin-4-one (2e):

White solid (160 mg, 0.560 mmol, 38%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta = 7.32$ (d, $J = 9.2$ Hz, 2H), 7.10 (d, $J = 9.2$ Hz, 2H), 4.12-3.98 (br m, 2H), 3.82-3.77 (br m, 1H), 3.55-3.49 (br m, 1H), 2.35-2.27 (br, 5H), 2.15-2.09 (br m, 1H), 1.88-1.79 (br m, 1H), 1.31 (d, $J = 5.2$ Hz, 2H), 1.29 (s, 3H), 1.10 (d, $J = 5.2$ Hz, 2H), 0.86 (s, 3H).

$^{13}$C NMR (CDCl$_3$, 125 MHz): $\delta = 207.6, 170.9, 136.8, 134.7, 129.4 (2C), 129.1 (2C), 44.7, 41.6, 41.2, 40.8, 38.5, 25.0 (2C), 24.5, 22.7, 21.2.$

IR: 2981, 2929, 2357, 2341, 1714, 1634, 1449, 1418, 1307, 1263, 1231, 1219, 1135, 1081, 807, 760, 739, 657, 573 cm$^{-1}$.

HRMS (ESI): C$_{18}$H$_{23}$NO$_2$ [M+Na]$^+$ calculated: 308.1621; found:308.1639.

1-(1-Naphthalen-2-yl-cyclopropanecarbonyl)-piperidin-4-one (2f):

Clear oil oil (5.00 g, 29.0 mmol, 20%).
**1H NMR** (CDCl₃, 500 MHz): δ = 7.94 - 7.89 (m, 2H), 7.88 – 7.85 (m, 1H), 7.57 - 7.77 (m, 4H), 4.21 (t, J = 6.3 Hz, 2H), 3.51 (t, J = 6.0 Hz, 2H), 2.69 (t, J = 6.3 Hz, 2H), 2.32 (q, J = 6.3 Hz, 2H).

**13C NMR** (CDCl₃, 100 MHz): δ = 206.5, 169.6, 133.4, 133.3, 129.4, 129.3, 128.5, 127.2, 126.5, 125.1, 124.3, 123.6, 45.7, 41.5, 40.9, 40.8.

**IR:** 3053, 2964, 2360, 1713, 1631, 1507, 1470, 1435, 1366, 1314, 1279, 1244, 1199, 1150, 973, 800, 780, 627 cm⁻¹.


### General Method for Synthesis of Oxadiazolones 5

Ketones 2a-f (0.083 mmol – 5.40 mmol) was combined with the corresponding hydrazine (3t-z; 1.09 eq) and methanol (0.05 M) in an oven-dried round bottomed flask and stirred at 30 °C for 1 hour. The temperature was raised to 50 °C for an additional 2 hours. The reaction was cooled to 0 °C and sodium borohydride (1.05 eq) was added. The reaction was allowed to warm to room temperature, stirred for 2 hours, quenched with brine, and extracted with dichloromethane (3 x 20 mL). The resultant organic layer was dried over Na₂SO₄, decanted, and concentrated under vacuum. The crude hydrazide (4) was then solubilized in THF (0.011 M) at 0 °C and triethylamine (2.6 eq) and triphosgene (0.67 eq) were sequentially added. The reaction was stirred at room temperature for 24 hours, quenched with brine, and extracted with ethyl acetate (3 x 20 mL). The resultant organic layer was dried over Na₂SO₄, decanted and concentrated under vacuum. Product purification was achieved by silica gel flash chromatography (gradient elution starting
with 25:75 ethyl acetate:hexane and ending with a 50:50 ethyl acetate:hexane) to yield azoxazolones 5.

**Characterization of Oxadiazolones 5**

![Chemical structure of 5-Phenyl-3-[1-(1-phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-3H-[1,3,4]oxadiazol-2-one (5at):](image)

5-Phenyl-3-[1-(1-phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-3H-[1,3,4]oxadiazol-2-one (5at):

White solid (100 mg, 0.257 mmol, 74%).

**$^1H$ NMR** (CDCl$_3$, 500 MHz): $\delta = 7.83$-$7.80$ (m, 2H), 7.54 - 7.42 (m, 3H), 7.36 - 7.31 (m, 2H), 7.25-7.18 (m, 3H), 4.76-4.75 (br m, 1H), 4.28-4.10 (br m, 2H), 2.97-2.75 (br m, 2H), 2.01-1.90 (br m, 2H), 1.75-1.66 (br m, 1H), 1.65-1.52 (br, 1H), 1.50-1.41 (br m, 2H), 1.25-1.18 (br m, 2H).

**$^{13}C$ NMR** (CDCl$_3$, 100 MHz): $\delta = 171.1, 153.5, 153.0, 140.6, 131.8, 129.1$ (2C), 129.0 (2C), 126.6, 125.8 (2C), 125. 5 (2C), 123.9, 53.2, 44.6, 41.3, 29.8 (2C), 29.6, 15.3 (2C).

**IR:** 2934, 2359, 2341, 1772, 1635, 1449, 1431, 1353, 1154, 993, 739, 690, 580 cm$^{-1}$.

**HRMS** (ESI): C$_{46}$H$_{47}$N$_6$O$_6$ [2M+H]$^+$ calculated: 779.35516; found: 779.3574.
3-[1-(1-phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-5-pyridin-3-yl-3H-[1,3,4]oxadiazol-2-one (5au):

Light yellow-orange solid (105 mg, 0.269 mmol, 13%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta = 9.06$ (br s, 1H), 8.74 (br s, 1H), 8.06 (dt, $J = 2.0, 8.0$ Hz, 1H), 7.42 (dd, $J = 5.1, 8.0$ Hz, 1H), 7.33 - 7.31 (m, 2H), 7.23-7.18 (m, 3H), 4.77-4.72 (br m, 1H), 4.28-4.10 (br m, 2H), 2.95-2.72 (br m, 2H), 2.01-1.88 (br m, 2H), 1.75-1.66 (br, 1H), 1.65-1.60 (br, 1H), 1.49-1.40 (br m, 2H), 1.25-1.18 (br m, 2H).

$^{13}$C NMR (CDCl$_3$, 100 MHz): $\delta = 171.1$, 152.6, 152.4, 151.5, 147.1, 140.6, 133.0, 129.0 (2C), 126.7, 125.5 (2C), 123.9, 120.5, 53.5, 44.6, 41.3, 29.8 (2C), 29.6, 15.3 (2C).

IR: 3179, 2955, 2863, 1784, 1625, 1610, 1456, 1432, 1412, 1328, 1160, 988, 740, 702, 580 cm$^{-1}$.

5-(4-Methoxy-phenyl)-3-[1-(1-phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-3H-[1,3,4]oxadiazol-2-one (5av):

White solid (223 mg, 0.53 mmol 43%).

$^1$H NMR (CDCl$_3$, 400 MHz): $\delta = 7.76$ (d, $J = 9.0$, 2H), 7.37-7.29 (m, 2H), 7.25 -7.17 (m, 3H), 6.96 (d, $J = 9.0$ Hz, 2H), 4.74 (br s, 1H), 4.32-4.17 (br s, 1H), 4.12 (sept, $J = 5.0$ Hz, 1H), 3.87 (s, 3H), 2.98-2.70 (br m, 2H), 2.05-1.84 (br m, 2H), 1.80-1.65 (br m, 1H), 1.64-1.50 (br, 1H), 1.51-1.35 (br, 2H), 1.30-1.14 (br, 2H).

$^{13}$C NMR (CDCl$_3$, 100 MHz): $\delta = 171.2$, 162.4, 153.6, 153.2, 140.7, 129.0 (2C), 127.6 (2C), 126.6, 125.5 (2H), 116.4, 114.6 (2C), 55.7, 53.1, 44.6, 41.3, 29.9 (2C), 29.6, 15.3 (2C).

IR: 2936, 1765, 1632, 1508, 1440, 1352, 1249, 1176, 1161, 1029, 1017, 836, 744, 693, 609, 574 cm$^{-1}$.


3-[1-(1-phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-5-(4-trifluoromethyl-phenyl)-3H-[1,3,4]oxadiazol-2-one (5aw):

White solid (85.1 mg, mmol 87%).

$^1$H NMR (CDCl$_3$, 400 MHz): $\delta = 7.94$ (d, $J = 8.1$ Hz, 2H), 7.75 (d, $J = 8.1$ Hz, 2H), 7.37-7.32 (m, 2H), 7.25-7.17 (m, 3H), 4.77 (br, 1H), 4.30-4.13 (br m, 2H), 2.97-2.73 (br m,
2H), 2.05-1.83 (br m, 2H), 1.80-1.65 (br m, 1H), 1.64-1.50 (br, 1H), 1.50-1.35 (br, 2H), 1.32-1.18 (br, 2H).

$^{13}$C NMR (CDCl$_3$, 100 MHz): $\delta = 171.2, 152.7, 152.3, 140.6, 133.3 (q, J = 32.4 Hz, 1C), 129.0 (2C), 127.2, 126.7, 126.2 (q, J = 3.8 Hz, 2C), 126.1 (2C), 125.5 (2C), 123.7 (q, J = 271.0 Hz, 1C), 53.5, 44.6, 41.3, 29.9 (2C), 29.6, 15.2 (2C).

IR: 2846, 2863, 2354, 2341, 1776, 1633, 1437, 1321, 1170, 1150, 1119, 1107, 1063, 1011, 992, 856, 847, 736, 699, 570, 515 cm$^{-1}$.

HRMS (ESI): C$_{24}$H$_{22}$F$_3$N$_3$NaO$_3$ [M+Na]$^+$ calculated: 480.15054 found: 480.1529

![Chemical Structure](image)

3-[1-(1-phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-5-(3-trifluoromethyl-phenyl)-3$H$-[1,3,4] oxadiazol-2-one (5ax):

White solid (117 mg, 0.253 mmol, 62%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta = 8.09$ (s, 1H), 8.00 (d, $J = 7.8$ Hz, 1H), 7.75 (d, $J = 7.8$ Hz, 1H), 7.63 (t, $J = 7.8$ Hz, 1H), 7.34 (t, $J = 7.8$ Hz, 2H), 7.25 -7.20 (m, 3H), 4.77 (br, 1H), 4.30-4.13 (br m, 2H), 3.00 - 2.80 (br m, 2H), 2.05-1.83 (br m, 2H), 1.80-1.65 (br m, 1H), 1.64-1.50 (br, 1H), 1.50-1.35 (br, 2H), 1.32-1.18 (br, 2H).

$^{13}$C NMR (CDCl$_3$, 100 MHz): $\delta = 171.8, 152.7, 152.3, 140.6, 132.0 (q, J = 33.4 Hz, 1C), 129.9, 129.0 (2C), 128.8, 128.2 (q, J = 3.8 Hz, 1C), 126.7, 125.6 (2C), 124.8, 123.6 (q, J = 271.0 Hz, 1C), 122.7 (q, $J = 3.8$ Hz, 1C), 53.5, 44.6, 41.3, 29.9 (2C), 29.6, 15.2 (2C).
IR: 2934, 2360, 2340, 1778, 1736, 1276, 1169, 1155, 1128, 1073, 1032, 994, 743, 693, 575 cm\(^{-1}\).

HRMS (ESI): C\(_{24}H_{23}F_3N_3O_3\)\([\text{M}+\text{H}]^+\) calculated: 458.16860, found: 458.16885.

**5-Furan-2-yl-3-[1-(1-phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-3\text{H}-[1,3,4]oxadiazol-2-one (5ay):**

White solid (79 mg, 0.208 mmol, 51%).

**\(\text{\textsuperscript{1}H NMR (CDCl}_3, 500 MHz): \(\delta = 7.59 (d, J = 1.8 Hz, 1H), 7.34-7.31 (m, 2H), 7.23-7.17 (m, 3H), 6.97 (d, J = 3.7 Hz, 1H), 6.56 (dd, J = 3.7, 1.8 Hz, 1H), 4.72 (br, 1H), 4.29-4.10 (br m, 2H), 2.95-2.70 (br m, 2H), 2.05-1.90 (br m, 2H), 1.80-1.57 (br m, 2H), 1.45 (br, 2H), 1.21 (br, 2H).**

**\(\text{\textsuperscript{13}C NMR (CDCl}_3, 125 MHz): \(\delta = 171.1, 152.1, 149.9, 145.8, 140.6, 139.0, 129.0 (2C), 126.6, 125.3 (2C), 113.9, 112.2, 53.5, 44.6, 42.2, 29.8 (2C), 29.5, 15.3 (2C).**

IR: 2928, 2861, 2359, 2341, 1778, 1634, 1434, 1326, 1151, 1038, 995, 949, 904, 741, 699, 572 cm\(^{-1}\).

HRMS (ESI): C\(_{42}H_{43}N_6O_8\) [2\text{M}+\text{H}]^+ calculated: 759.31369 found: 759.3155.
3-[(1-Phenyl-cyclopropanecarbonyl)-piperidin-4-yl]-5-thiophene-2-yl-3H-[1,3,4]oxadiazol-2-one (5az):

White solid (80 mg, 0.202 mmol, 49%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta$ = 7.56 (dd, $J = 1.3$, 3.8 Hz, 1H), 7.50 (dd, $J = 1.3$, 5.0 Hz, 1H), 7.35-7.31 (m, 2H), 7.24-7.18 (m, 3H), 7.12 (dd, $J = 3.8$, 5.0 Hz, 1H), 4.80-4.65 (br, 1H), 4.25-4.15 (br, 1H), 4.15-4.05 (m, 1H), 2.95-2.70 (br m, 2H), 2.00-1.85 (br, 2H), 1.75-1.62 (br, 1H), 1.62-1.50 (br, 1H), 1.45 (br, 2H), 1.22 (br, 2H).

$^{13}$C NMR (CDCl$_3$, 100 MHz): $\delta$ 171.0, 152.2, 150.0, 140.3, 129.7, 129.2, 128.8 (2C), 128.0, 126.5, 125.3, 125.2 (2C), 53.1, 44.4, 41.1, 29.6 (2C), 29.4, 15.1 (2C).

IR: 3085, 2928, 2860, 2358, 2341, 1768, 1633, 1613, 1430, 1324, 1272, 1153, 1045, 1012, 987, 857, 738, 699, 578 cm$^{-1}$.

HRMS (ESI): C$_{21}$H$_{22}$N$_3$O$_3$S[M+H]$^+$ calculated: 396.13764; found: 396.1381.
3-\{1-\{1-(4-Methoxy-phenyl)-cyclopropanecarbonyl\}-piperidin-4-yl\}-5-phenyl-3H-[1,3,4]oxadiazol-2-one (5bt):

White solid (190 mg, 0.453 mmol 62%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta = 7.81$ (br d, $J = 6.7$ Hz, 2H), 7.51-7.45 (m, 3H), 7.15 (br d, $J = 9.0$ Hz, 2H), 6.86 (br d, $J = 9.0$ Hz, 2H), 4.80-4.65 (br, 1H), 4.35-4.20 (br m, 1H), 4.14 (tt, $J = 4.6$, 10.9 Hz, 1H), 3.78 (s, 3H), 3.00-2.70 (br m, 2H), 2.08-1.51 (br m, 4H), 1.47-1.35 (br m, 2H), 1.20-1.12 (br m, 2H).

$^{13}$C NMR (CDCl$_3$, 100 MHz): $\delta$ 171.4, 158.3, 153.5, 153.1, 132.6, 131.8, 129.1, 126.8, 125.8 (4C), 123.9, 114.4, 114.3, 55.5, 53.3, 53.1, 44.5, 41.3, 29.9, 29.1, 14.7 (2C).

IR: 2906, 2869, 1763, 1629, 1513, 1449, 1430, 1322, 1245, 1181, 1154, 1038, 1021, 997, 856, 823, 731, 657, 592, 572 cm$^{-1}$.


5-(4-Methoxy-phenyl)-3-\{1-\{1-(4-methoxy-phenyl)-cyclopropanecarbonyl\}-piperidin-4-yl\}-3H-[1,3,4]oxadiazol-2-one (5bv):

White solid (91.4 mg, 0.209 mmol 81%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta = 7.75$ (br d, $J = 9.2$ Hz, 2H), 7.14 (br d, $J = 9.2$ Hz, 2H), 6.96 (br d, $J = 9.2$ Hz, 2H), 6.87 (br d, $J = 9.2$ Hz, 2H), 4.80-4.65 (br, 1H), 4.35-4.18 (br, 1H), 4.12 (tt, $J = 10.9$, 4.6 Hz, 1H), 3.87 (s, 3H), 3.79 (s, 3H), 3.00-2.70 (br m, 2H), 2.02-
1.80 (br, 2H), 1.80-1.64 (br, 1H), 1.64-1.50 (br, 1H), 1.48-1.36 (br m, 2H), 1.20-1.12 (br m, 2H).

$^{13}$C NMR (CDCl$_3$, 125 MHz): $\delta =$ 171.4, 162.4, 158.3, 153.6, 153.2, 132.6, 127.6 (2C), 126.9 (2C), 116.4, 114.6 (2C), 114.3 (2C), 55.7, 55.5, 53.1, 44.6, 41.3, 29.9 (2C), 29.1, 14.7 (2C).

IR: 2948, 2357, 2341, 1762, 1632, 1509, 1436, 1246, 1176, 1156, 1025, 996, 832, 740, 733, 607, 571, 525 cm$^{-1}$.

HRMS (ESI): C$_{25}$H$_{28}$N$_3$O$_5$ [M+H]$^+$ calculated: 450.20235; found: 450.2014.

3-{1-[1-(4-Methoxy-phenyl)-cyclopropanecarbonyl]-piperidin-4-yl}-5-(4-trifluoromethyl-phenyl)-3H-[1,3,4]oxadiazol-2-one (5bw):

White solid (104 mg, 0.213 mmol, 88%).

$^1$H NMR (CDCl$_3$, 500 MHz): $\delta =$ 7.94 (d, $J =$ 8.1 Hz, 2H), 7.74 (d, $J =$ 8.1 Hz, 2H), 7.15 (br d, $J =$ 8.6 Hz, 2H), 6.87 (br d, $J =$ 8.6 Hz, 2H), 4.80-4.66 (br, 1H), 4.35-4.20 (br, 1H), 4.16 (tt, $J =$ 10.9, 4.0 Hz, 1H), 3.78 (s, 3H), 3.00-2.70 (br m, 2H), 2.05-1.85 (br, 2H), 1.80-1.64 (br, 1H), 1.64-1.48 (br, 1H), 1.42-1.34 (br m, 2H), 1.21-1.10 (br m, 2H).

$^{13}$C NMR (CDCl$_3$, 125 MHz): $\delta =$ 171.5, 158.4, 152.7, 152.3, 133.3 (q, $J =$ 32.5 Hz, 1C), 132.6, 127.2, 127.0 (2C), 126.2 (q, $J =$ 3.8 Hz, 2C), 126.1 (2C), 123.7 (q, $J =$ 272.5 Hz, 1C), 114.4 (2C), 55.5, 53.5, 44.5, 41.3, 29.9 (2C), 29.1, 14.60 (2C).
IR: 2954, 2924, 2359, 2341, 1784, 1622, 1513, 1415, 1321, 1250, 1157, 1127, 1064, 1029, 994, 854, 819, 748, 735, 590, 560 cm\(^{-1}\).

HRMS (ESI): C\(_{25}\)H\(_{25}\)F\(_3\)N\(_3\)O\(_4\) \([\text{M+H}]^+\) calculated: 488.17917; found: 488.1795.

3-\{1-[1-(2,4-Dichloro-phenyl)-cyclopropanecarbonyl]-piperidin-4-yl\}-5-phenyl-3\(H\)-[1,3,4]oxadiazol-2-one (5ct):

White solid (35.9 mg, 0.078 mmol 48%).

\(^1\)H NMR (CDCl\(_3\), 300 MHz): \(\delta = 7.82\) (dd, \(J = 7.8, 1.5\) Hz, 2H), 7.53-7.45 (m, 4H), 7.26-7.23 (m, 2H), 4.49-4.18 (br m, 2H), 4.11 (tt, \(J = 10.9, 4.6\) Hz, 1H), 2.81 (td, \(J = 11.5, 2.3\) Hz, 2H), 1.87-1.76 (br m, 2H), 1.74 – 1.59 (m, 4H), 1.18 - 1.12 (br m, 2H).

\(^{13}\)C NMR (CDCl\(_3\), 125 MHz): \(\delta = 170.7, 153.5, 153.1, 138.0, 137.5, 133.5, 131.8, 130.4, 130.1, 129.2 (2C), 127.5, 125.8 (2C), 123.9, 53.0, 43.7 (2C), 29.8 (2C), 28.9, 14.4 (2C).

IR: 2924, 2852, 2358, 2341, 1815, 1770, 1717, 1615, 1449, 1350, 1275, 1066, 957, 920, 772, 729, 688, 572 cm\(^{-1}\).

HRMS (ESI): C\(_{23}\)H\(_{22}\)Cl\(_2\)N\(_3\)O\(_3\) \([\text{M+H}]^+\) calculated: 458.10382; found: 458.10321.
3-{1-[4-Chloro-phenyl]-cyclopropanecarbonyl]-piperidin-4-yl]-5-phenyl-3H-[1,3,4]oxadiazol-2-one (5dt):

White solid (0.404 mg, 0.95 mmol, 19%).

\(^1\)H NMR (CDCl\(_3\), 500 MHz): \(\delta = 7.84-7.80\) (m 2H), 7.54-7.45 (m, 3H), 7.31 (br d, \(J = 8.6\) Hz, 2H), 7.14 (br d, \(J = 8.6\) Hz, 2H), 4.85-4.70 (br, 1H), 4.20-4.11 (br, 1H), 4.15 (tt, \(J = 10.9, 4.6\) Hz, 1H), 3.00-2.70 (br m, 2H), 2.04-1.85 (br, 2H), 1.85-1.67 (br, 1H), 1.67-1.54 (br, 1H), 1.54-1.40 (br m 2H), 1.25-1.15 (br m, 2H).

\(^1\)C NMR (CDCl\(_3\), 100 MHz): \(\delta = 170.6, 153.5, 153.0, 139.3, 132.4, 132.0, 129.1\) (3C), 127.0, 125.8 (4C), 123.9, 53.1, 44.6, 41.4, 29.9 (2C), 29.2, 15.4 (2C).

IR: 2939, 2864, 2359, 2341, 1765, 1636, 1613, 1492, 1430, 1401, 1356, 1324, 1154, 1097, 1036, 1021, 1011, 992, 813, 738, 691, 583 cm\(^{-1}\).

HRMS (ESI): C\(_{23}\)H\(_{23}\)ClN\(_3\)O\(_3\) [M+H]\(^+\) calculated: 424.14225; found: 424.1425.

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3-{1-(2,2-Dimethyl-1-p-tolyl-cyclopropanecarbonyl]-piperidin-4-yl]-5-phenyl-3H-[1,3,4]oxadiazol-2-one (5et):

White solid (72 mg, 0.172 mmol, 49%).

\(^1\)H NMR (CDCl\(_3\), 500 MHz): \(\delta = 7.83\) (d, \(J = 7.0\) Hz, 1H), 7.76 (d, \(J = 7.0\) Hz, 1H), 7.53-7.43 (br m, 3H), 7.34 (d, \(J = 8.1\) Hz, 2H), 7.16-7.11 (br m, 2H), 4.68-4.43 (br m, 2H), 4.17-
4.08 (br m, 1H), 3.21-2.90 (br m, 1H), 2.70-2.63 (br m, 1H), 2.32 (s, 3H), 1.95-1.70 (br m, 4H), 1.32-1.22 (br, 4H), 1.12-1.06 (m, 1H), 0.87 (s, 3H).

**13C NMR** (CDCl₃, 125 MHz, multiple peaks observed due to rotational isomers): δ 170.7 & 170.6 (1C, C12), 153.5 & 153.2 (1C, C8), 153.0 (1C, C7), 136.5 & 136.4 (1C, C17), 135.1 (1C, C20), 131.7 & 131.6 (1C, C3), 129.4 (2C, C19, C21), 129.2 (2C, C2, C4), 129.1 (2C, C1, C5), 125.7 & 125.6 (2C, C18, C22), 123.9 (1C, C6), 53.3 & 53.1 (1C, C9), 44.7 & 44.4 (1C, C11), 40.9 & 40.8 (1C, C11’), 38.6 (1C, C13), 30.3 & 30.2 & 30.1 & 29.5 (2C, C10, C10’), 25.2 (2C, C16, C14), 24.5 & 24.1 (1C, C15), 22.8 & 22.7 (1C, C16’), 21.2 (1C, C23).

**IR:** 2931, 2867, 1776, 1618, 1447, 1431, 1354, 1325, 1038, 1019, 992, 736, 686, 569, 518 cm⁻¹.


![Chemical Structure](image)

3-[1-(1-Naphthalen-1-carbonyl)-piperidin-4-yl]-5-phenyl-3H-[1,3,4]oxadiazol-2-one

(5ft):
White solid (0.677 mg, 1.70 mmol, 60%).
$^{1}H$ NMR (CDCl$_3$, 500 MHz): $\delta = 8.09 - 7.78$ (m, 5H), 7.64 – 7.40 (m, 7H), 5.06 (d, $J = 13.0$ Hz, 1H), 4.33 - 4.23 (m, 1H), 3.62 – 3.53 (m, 1H), 3.17 – 3.02 (m, 2H), 2.29 – 2.11 (m, 2H), 2.01 – 1.74 (m, 2H).

$^{13}C$ NMR (CDCl$_3$, 100 MHz, multiple peaks observed due to rotational isomers): $\delta = 169.5, 153.5, 152.9, 134.0, 133.5, 131.7, 129.7, 129.5, 129.3, 129.0$ (2C), 128.6, 127.2, 126.6, 125.7 (2C), 125.2, 124.8, 123.7, 53.1, 46.0, 40.5, 31.0, 30.1.

IR: 2933, 2359, 2341, 1768, 1629, 1448, 1436, 1352, 1323, 981, 796, 777, 732, 687, 675, 632, 584 cm$^{-1}$.

HRMS (ESI): C$_{24}$H$_{21}$N$_{3}$NaO$_{3}$ [M+Na]$^+$ calculated: 422.14751; found: 422.1471.

Spectra Data
Chemical Shift (ppm)

Normalized Intensity
Chemical Shift (ppm)

Normalized Intensity

15.2457
29.5857
29.8145
41.2846
44.6027
53.1647
76.9058
77.2300
77.5446
123.9113
125.4464
125.7610
126.6287
128.9837
129.1267
131.7583
140.6064
153.0204
153.4685
171.1456
ML-126-cryst HSQC
**Biological Assay**

Antagonist activity was evaluated based on the inhibition of LPI-induced β-arrestin activity through the use of two differential means of β-arrestin quantitation, and in two different cellular backgrounds. Compounds were initially screened in an image based cell assay to identify their antagonist activity on their ability to inhibit LPI-induced β-arrestin recruitment. In this assay, U2OS cells over-expressing GPR55 (modified at the C-terminus, GPR55E) and GFP tagged β-arr2, are grown on glass coverslips and pre-incubated (30 minutes) with candidate antagonists prior to LPI exposure (6μM; 40 minutes). Images of cells were taken using a fluorescent microscope and analyzed with Image J (http://rsbweb.nih.gov/ij/) using a custom written plug-in. The modified GPR55E receptor, concentration of agonist (EC₈₀), and details of the β-arrestin recruitment assay have been previously described.¹¹,¹² Compounds with an IC₅₀ below 15μM were also evaluated employing DiscoveRx PathHunter® Complementation technology. CHO-K1 cells stably expressing GPR55 (fused to a β-galactosidase enzyme fragment), and β-arrestin (fused to an N-terminal deletion β-galactosidase mutant), were grown in selection media and were passaged up to 10 times according to manufacturer protocols. In the presence of agonist (LPI, 90 minutes), and PathHunter® detection reagents containing β-galactosidase substrate, a chemiluminescent signal is generated due to the forced complementation of the GPR55 fused β-galactosidase fragment and the β-arrestin fused N-terminal deletion mutant fragment of β-galactosidase. A concentration response curve was generated, and antagonist activity was evaluated based on the inhibition of the LPI-
induced chemiluminescence signal. Prior to exposure with LPI (37 °C, 90 minutes) cells were preincubated with putative antagonists for 30 minutes (37 °C).

The chemiluminescence signal was measured using a Perkin Elmer Envision plate reader for 1 second. Experiments were run in triplicate and repeated at a minimum of 3 times. Ligand readings, expressed as relative luminescence units, were subtracted from corresponding vehicle readings and analyzed in GraphPad Prism version 5.0 (GraphPad, San Diego, CA). IC₅₀ values of antagonists were similar in both β-arrestin methodologies, and in both cellular backgrounds.

**Modeling Methods**

**Conformational Analysis of ML191** ((CID23612552), 5-phenyl-3-(1-(1-(p-tolyl)cyclopropane-carbonyl)piperidin-4-yl)-1,3,4-oxa-diazol-2(3H)-one). A complete conformational analysis of ML191 was performed using *ab initio* Hartree-Fock calculations at the 6-31G* level as encoded in Spartan '08 (Wavefunction, Inc., 18401 Von Karman Ave., Suite 370, Irvine, CA 92612). 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 energy difference between the global minimum energy conformer of ML191 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 for ML191 was calculated using Spartan '08 (Wavefunction, Inc., 18401 Von Karman Ave., Suite 370, Irvine, CA 92612). The electrostatic potential energy was calculated using the *ab initio* Hatree-Fock method at 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.

**Receptor Model Development** - The construction of our initial GPR55 receptor homology model was described previously in Kotsikorou *et al.* This homology model used the crystal structure of β2-AR as the template. This initial model has been modified to reflect several structural elements found in the x-ray crystal structure of CXCR4 in its inactive state, a receptor with which the GPR55 receptor has high homology. 1) TMH2 and TMH4 of GPR55 were modeled using the corresponding helices in the CXCR4 structure. GPR55 and CXCR4 have prolines in the positions 2.58 and 4.59 that would lead to very different conformations of TMHs 2 and 4 compared to the β2-AR template which has prolines at 2.59 and 4.60. In addition, preceding the prolines at positions 2.56 and 4.57, CXCR4 has threonines in a g- conformation that influence the overall bend of each helix. Analogously, GPR55 has serines at 2.56(76) and 4.57(153) that can assume a g- conformation. Therefore, the use of the CXCR4 TMH2 and TMH4 as templates is well-justified. 2) The high sequence homology between GPR55 and CXCR4 in the extracellular ends of TMHs 6 and 7, dictated the introduction of extracellular (EC) helical extensions to the EC3 loop of GPR55 to match that of
CXCR4. 3) The EC2 loop in CXCR4 contains a beta sheet. Due to high sequence similarity in this region, as well, the beta sheet motif was built into the EC-2 loop in GPR55. 4) In addition to the disulfide bridge between the EC-2 loop and C3.25, CXCR4 has another disulfide bridge between a Cys in the N-terminus Cys(28) and Cys(274) in the EC-3 loop. The same residues (Cys(10) and Cys(260)) are found in the GPR55 sequence, so this second disulfide bridge was added to the GPR55 model. Recent C(10)A and C(260)A mutation studies suggest that the C(10)/C(260) disulfide bridge exists in GPR55 (M. Lingerfelt et al., 2016, manuscript submitted). The GPR55 inactive state model is also characterized by an intracellular hydrogen bond between R3.50(119) and Q6.30(221) that closes off the intracellular domain of the receptor, preventing G-protein interaction. The resultant homology model was energy minimized to relieve any steric overlaps.

**Docking of Ligands** - The lowest energy conformation of ML191 was used as input for receptor docking studies. The Schrödinger workflow, Induced Fit, (Schrödinger Inc. Portland, OR) was used to explore possible binding conformations and receptor site interactions in the GPR55 R model. The Induced Fit Docking workflow was used to dock subject ligands in the binding site of the GPR55 R receptor model. The binding site of the receptor was defined by the area enclosed in a box with dimensions 26Å x 26Å x26Å (the default value). Because the endogenous ligand, LPI, is negatively charged, it is very likely that a positively charged amino is the primary interaction site for GPR55 ligands. The only positively charged amino acid in the binding pocket is K2.60(80). Recent K2.60A mutation studies suggest that this residue is essential for ligand activation at
GPR55 (M. Lingerfelt et al., manuscript in preparation). For this reason, K2.60(80) was set as the center of the box and a hydrogen bond with K2.60(80) was defined as a constraint. The two constraints (enclosing box and hydrogen bond) were used for both docking steps that Induced Fit Docking performs. The initial receptor minimization that Induced Fit Docking performs in preparation for docking the ligand was omitted since the receptor model was already energy minimized. For the first Glide calculation, the receptor and ligand Van der Waals radii were set to the default value of 0.50 and the number of maximum poses to be produced was also set to 40. During the Prime stage of the calculation, amino acids within 5.0Å of the ligand were refined to better accommodate the ligand. Since the GPR55 receptor is a transmembrane protein, an implicit membrane was used during the Prime refinement step. Then, during the second Glide step, the top 30 poses produced by the first calculation were redocked to the receptor. Docked poses that were within 40.0 kcal/mol from the lowest one were kept.

**Ligand/Receptor Minimization** - To optimize the ligand/receptor interaction in the receptor/ligand complexes, each was minimized using Macromodel 9.1 (Schrödinger Inc.; Portland, OR). Since the receptor model was minimized before the docking of the ligands, a brief minimization was adequate to resolve any steric clashes after docking. To this end, 500 steps of Polak-Ribier conjugate gradient minimization using force field defined dielectric were performed. A harmonic constraint was placed on all the TMH backbone torsions (φ, ψ, and ω) to preserve the general shape of the helices during minimization. The sidechains of the TMH region, the loops and termini were allowed to relax. The Generalized Born/Surface Area (GB/SA) continuum solvation model for water
as is implemented in Macromodel 9.1 was used for minimizing the loop regions. An 8.0Å nonbonded cutoff (updated every 10 steps), a 20.0Å electrostatic cutoff, and a 4.0Å hydrogen bond cutoff were used in each stage of the calculation. The transmembrane region and the docked ligand were frozen while the loops were minimized.

**Results**

**ML191 Global Minimum Energy Conformer.** Overall, the GPR55 probe molecules have three regions. The first two regions include a broad head connected to a central portion of the ligand that is vertical. Together, this gives ML191 the shape of the number “7”. The third section at the end of the vertical segment is a pendant aromatic or heterocyclic ring that is nearly perpendicular to the vertical segment. In the global minimum energy conformer of ML191, the para-methylphenyl ring is at an angle with the carbonyl group that joins the cyclopropyl and the piperidine rings (C1-C2-C3-C4 = 53.5° and C2-C3-C4-N6 = 72.5°). The piperidine, which is in a chair conformation, and the attached 1,3,4-oxadiazo-2-one ring form the vertical segment of the “7”. The 1,3,4-oxadiazo-2-one ring is perpendicular to the plane of the piperidine ring (H-C9-N10-C11 = -179.9°) and the pendant phenyl ring is in plane with the 1,3,4-oxadiazo-2-one ring (O12-C13-C14-C15 = -0.4°). The global minimum energy conformer of ML191 is illustrated in Figure 15B (bottom).

**ML191 Molecular Electrostatic Potential Map** - Figure 15B (top) illustrates the molecular electrostatic potential map (ranges in kJ/mol given next to ligand) of the docked conformation of ML191 at GPR55 R. Below this map, the ML191 conformer used to calculate the map is shown in tube display (Figure 15B (bottom). ML191
possesses a broad head region connected to a central portion of the ligand that has a thin profile. The most electronegative region of ML191 is located close to the end of this central section and this portion is followed by a pendant ring that juts out nearly perpendicular to the central portion of the molecule. The carbonyl oxygen of the 1,3,4-oxadiazoo-2-one ring of ML191 is the most electronegative region of the ligand and forms the part of ML191 that interacts with K2.60(80) (see docking results below).

**Compound Docking in GPR55 Inactive State (R) Model** – Docking studies identified the putative binding site for ML191 to be the TMH 2-3-5-6-7 region of 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. Figure 15A illustrates the final ML191/GPR55 R complex obtained using Induced Fit. The ML191 1,3,4-oxadiazoo-2-one carbonyl oxygen forms a hydrogen bond with K2.60(80). The hydrogen bond (N-O) distance and (N-H—O) and angle are 2.81Å and 154º respectively. ML191 forms a number of aromatic stacking interactions. The methylphenyl ring next to the cyclopropyl group forms a stack with the EC2 loop residue F169 (ring centroid to centroid distance is 5.08Å and the angle between ring planes is 55°). The pendant phenyl group adjacent to the 1,3,4-oxadiazoo-2-one ring stacks with F6.55 (246). The ring centroid to centroid distance is 5.41Å and the angle between ring planes is 50°. The total pairwise interaction energy for ML191 with GPR55 is -33.16 kcal/mol. The major interaction energy contributions for this compound are the hydrogen bonding interaction with K2.60(80) and Van der Waals interactions with M7.39 (274). Aromatic stacking interactions with F169 and Van der Waals interactions with Y3.32
(101) also contribute to the interaction energy. The remainder of the aromatic stacking interactions identified contribute less to the overall interaction energy. The pendant phenyl group prevents a change in the $\chi_1$ of the toggle switch residue, M3.36 (104) and therefore keeps GPR55 in the inactive state.

**Literature Cited**


APPENDIX B
SUPPLEMENTAL INFORMATION FOR CHAPTER IV

Supporting Information For

Identification of Crucial Amino Acid Residues Involved in Agonist Signaling at the GPR55 Receptor*

* This material is available free of charge via the Internet at http://pubs.acs.org.

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Author Contributions

MEA and PHR designed the study and wrote the paper and contributed equally to this work. MAL and PZ also contributed to writing the manuscript and contributed equally to this work. MAL and DPH constructed the molecular model in consultation with PHR. PZ designed and constructed vectors for expression of mutant proteins and Mary A. Lingerfelt, Pingwei Zhao, Haleli P. Sharir, Dow P. Hurst, Patricia H. Reggio and Mary E. Abood analyzed the mutant phenotypes in vitro. HPS designed and constructed vectors for expression of mutant proteins. All authors reviewed the results and approved the final version of the manuscript.
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Figure S1.
Table S1

ML184/GPR55 R* Complex Interaction Energy

<table>
<thead>
<tr>
<th>GPR55 B&amp;W</th>
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<th>Vdw (kcal)</th>
<th>Total Interaction Energy (kcal)</th>
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-55.9 Interaction E (kcal/mol)
1.91 Ligand Conformational Cost (kcal/mol)
-54.0 Total Interaction Energy (kcal/mol)