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Defining the Molecular Basis for the First Potent and Selective Orthosteric Agonists of the FFA2 Free Fatty Acid Receptor*

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Background: Understanding the function of FFA2 has been slowed by a lack of selective orthosteric ligands.

Results: Residues within FFA2 that dictate the recognition and function of potent and selective orthosteric agonists are described.

Conclusion: Key aspects of ligand interaction with the orthosteric binding pocket of FFA2 are defined.

Significance: This work will be invaluable in future drug development at the FFA2 receptor.

FFA2 is a G protein-coupled receptor that responds to short chain fatty acids and has generated interest as a therapeutic target for metabolic and inflammatory conditions. However, definition of its functions has been slowed by a dearth of selective ligands that can distinguish it from the closely related FFA3. At present, the only selective ligands described for FFA2 suffer from poor potency, altered signaling due to allosteric modes of action, or a lack of function at non-human orthologs of the receptor. To address the need for novel selective ligands, we synthesized two compounds potentially having FFA2 activity and examined the molecular basis of their function. These compounds were confirmed to be potent and selective orthosteric FFA2 agonists. A combination of ligand structure-activity relationship, pharmacological analysis, homology modeling, species ortholog comparisons, and mutagenesis studies were then employed to define the molecular basis of selectivity and function of these ligands. From this, we identified key residues within both extracellular loop 2 and the transmembrane domain regions of FFA2 critical for ligand function. One of these ligands was active with reasonable potency at rodent orthologs of FFA2 and demonstrated the role of FFA2 in inhibition of lipolysis and glucagon-like peptide-1 secretion in murine-derived 3T3-L1 and STC-1 cell lines, respectively. Together, these findings describe the first potent and selective FFA2 orthosteric agonists and demonstrate key aspects of ligand interaction within the binding site of FFA2 that will be invaluable in future ligand development at this receptor.

G protein-coupled receptors (GPCRs)² activated by free fatty acids have generated significant interest in recent years for their potential use as therapeutic targets in the treatment of various metabolic and inflammatory conditions. These GPCRs include a family of four receptors: two that respond to long chain fatty acids (FFA1 (previously GPR40) and FFA4 (previously GPR120)) and two that respond to short chain fatty acids (SCFAs) (FFA2 (previously GPR43) and FFA3 (previously GPR41)). Although the long chain fatty acid receptors have received the most interest, FFA1 due to its ability to potentiate glucose-stimulated insulin release (1) and FFA4 for its ability to stimulate glucagon-like peptide-1 secretion (2) and to mediate the insulin-sensitizing and anti-inflammatory properties of n-3 fatty acids (3), the SCFA receptors, and particularly FFA2, have also generated significant interest. However, the development and validation of FFA2 as a therapeutic target has been hindered greatly by a lack of suitably selective ligands for this receptor over the closely related FFA3 (4).

Although both FFA2 and FFA3 respond to the same group of SCFA ligands, they display clear differences in the rank order of potency for these ligands. Of particular interest, human FFA2 (hFFA2) responds with equal potency to acetate (C2) and propionate (C3), whereas hFFA3 displays greater potency for C3 than for C2 (5–7). Although this observation has led to some use of C2 as a selective agonist for FFA2 (8, 9), both the potency and selectivity for C2 at hFFA2 remain low (10), making it

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2 The abbreviations used are: GPCR, G protein-coupled receptor; SCFA, short chain fatty acid; pERK, ERK phosphorylation; hFFA2 and -3, human FFA2 and -3, respectively; mFFA2 and -3, mouse FFA2 and -3, respectively; rFFA2, rat FFA2; 4-CMTB, 4-chloro-α-(1-methylethyl)-N-2-thiazolylbenzeneacetamide; CATPB, (S)-(−)-3-(2-(3-chlorophenyl)acetamido)-4-(4-(1fluoromethyl)phenyl)butanoic acid; GLP-1, glucagon-like peptide-1; TUG-891, 3-(4-fluoro-4-methyl-1,1-biphenyl-2-yl)methoxyphenylpropionic acid; GPS; guanosine 5′-3-O-(thio)triphasphate; eYFP, enhanced yellow fluorescent protein; ECL2, extracellular loop 2; BRET, bioluminescence resonance energy transfer; HBSS, Hank’s balanced salt solution.
impractical to use this ligand effectively to differentiate clearly between FFA2 and FFA3 function. More importantly, we have recently demonstrated that although the rank order difference is maintained, the selectivity for C2 is not preserved in the rodent orthologs of FFA2 and FFA3, where C2 was found to be equipotent at mouse FFA2 (mFFA2) and mFFA3 (11). Clearly, this demonstrates a need for synthetic ligands that can differentiate between the function of these receptors in both human and rodent systems.

Early attempts to identify synthetic ligands for FFA2 revealed a series of phenylaceticamides, exemplified by 4-chloro-α-(1-methylaryl)-N-2-thiazolylbenzeneacetamide (4-CMTB), as the first reasonably potent and selective agonists of this receptor (12). However, detailed examination of the pharmacology of these compounds has demonstrated that they are in fact ago-allosteric modulators of FFA2, binding to a site distinct from the endogenous SCFAs (12, 13). Although 4-CMTB is active at rodent orthologs of FFA2 (11) and has been used to define some aspects of FFA2 function (12, 14), it is also apparent that as an allosteric modulator, signaling responses to 4-CMTB at FFA2 are not identical to those of the endogenous SCFAs (13). Considering this, there is clearly a need to develop novel FFA2 selective ligands with properties more similar to, and which ideally bind to the same site as, the endogenous SCFAs. To date, the only study describing selective orthosteric agonists for FFA2 examined the structure–activity relationship of small carboxylic acids to identify a number of compounds with modest selectivity for both FFA2 and FFA3 (10). However, due to their relatively small size and resulting low binding energy, the potency of these small carboxylic acid agonists remains low.

Selective antagonists for the SCFA receptors would be equally useful in helping to define the specific functions of FFA2 compared with FFA3, and, indeed, at least one FFA2 antagonist/inverse agonist, (S)-3-(2-(3-chlorophenyl)acetonitrile)-4-(4-(3-trifluoromethyl)phenyl) butanoic acid (CATPB), has been described (15). However, the practical utility of this compound is reduced substantially by the fact that it appears to be highly selective for the human ortholog, with no significant affinity at mFFA2 (11). Given the lack of potent and selective orthosteric agonists or antagonists with activity in non-human systems, there is clearly a substantial need for novel FFA2 ligands if the function of this receptor is to be fully explored and understood.

To address this need, we identified compounds from the patent literature with reported activity at FFA2 that we hypothesized were likely to be orthosteric agonists, based primarily on the presence, as in the SCFAs, of a carboxylate moiety (16). We synthesized two compounds based on the structures described and examined their function at human and rodent orthologs of the FFA receptors. This confirmed these compounds to be potent and selective orthosteric FFA2 agonists. We then used a combination of ligand-based structure activity, detailed pharmacological and functional analysis, homology modeling, species ortholog comparisons, and mutational analysis to define key molecular determinants of how these ligands interact with FFA2 and then used the ligand with similar potency at human and mouse FFA2 to define the role of this receptor in both the regulation of lipolysis in model adipocytes and in the release of the incretin glucagon-like peptide-1 (GLP-1) from enteroendocrine cells.

EXPERIMENTAL PROCEDURES

Materials—FFA2 ligands were synthesized based on the methods described previously (16). The identity of each compound was confirmed by NMR spectroscopy. Synthesized compound, compound 1 (see Fig. 1), was 3-benzyl-4-(cyclopentyl)-2-thiazolylbenzeneacetamide (TUG-891), was synthesized as described by Shimpukade et al. (17). Tissue culture reagents were from Invitrogen. Molecular biology enzymes and reagents were from Promega (Southampton, UK). The radiochemical [35S]GTPγS was from PerkinElmer Life Sciences. All other experimental reagents were from Sigma.

Plasmids and Mutagenesis—All plasmids used encoded either human, mouse, or rat FFA1–4 receptors with enhanced yellow fluorescent protein (eYFP) fused to their C termini in the

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cultured for at least 24 h before initiating GLP-1 secretion experiments.

[^55]GTPγS Incorporation Assay—Cell membrane preparations were generated from Flp-In^TM T-REx^TM 293 cells induced to express the receptor of interest, as described previously (18).[^55]GTPγS binding experiments were performed according to a method described previously (10). Briefly, cell membrane preparations containing 5 or 10 μg of protein were added to assay buffer (50 mM Tris-HCl, pH 7.4, 10 mM MgCl₂, 100 mM NaCl, 1 mM EDTA, 1 mM GDP, and 0.1% fatty acid-free bovine serum albumin) containing the appropriate concentrations of ligand and allowed to reach equilibrium by preincubating for 15 min at 25 °C. To initiate the assay, 50 nCi of[^55]GTPγS was added to each tube, and the reaction was terminated by rapid filtration through GF/C glass filters using a 24-well Brandel cell harvester (Alpha Biotech, Glasgow, UK) after a 1-h incubation at 25 °C. Unbound[^55]GTPγS was washed from filters by three washes with ice-cold wash buffer (50 mM Tris-HCl, pH 7.4, and 10 mM MgCl₂) before the remaining bound[^55]GTPγS was measured by liquid scintillation spectrometry.

Bioluminescence Resonance Energy Transfer (BRET) β-Arrestin-2 Recruitment Assay—A plasmid encoding an eYFP-tagged form of the receptor to be assayed was co-transfected in a 4:1 ratio with a β-arrestin-2 Renilla luciferase (Rluc) plasmid. Cells were then transferred into white 96-well plates 24 h post-transfection. Then 48 h post-transfection, cells were washed, and the culture medium was replaced with Hanks’ balanced salt solution (HBSS) immediately prior to conducting the assay. To measure β-arrestin-2 recruitment, the Rluc substrate coelenterazine h was added to a final concentration of 2.5 μM, and then cells were incubated for 10 min at 37 °C, test compounds were then added, and cells were incubated for a further 5 min at 37 °C. BRET resulting from receptor-β-arrestin-2 interaction was then assessed by measuring the ratio of luminescence at 535 and 475 nm using a Phorstar FS fitted with the BRET1 optic module (BMG Labtech, Aylesbury, UK).

Intracellular Ca²⁺ Mobilization Assay—All Ca²⁺ experiments were carried out using Flp-In^TM T-REx^TM stable inducible cell lines treated with doxycycline (100 ng/ml) to induce expression of the receptor of interest. Cells were plated 50,000/well in black 96-well plates with clear bottoms and then allowed to adhere for 3–6 h. Doxycycline was then added to induce receptor expression, and cells were maintained in culture overnight. Prior to the assay, cells were labeled for 45 min with the calcium-sensitive dye Fura-2 AM and then washed and incubated for 15 min with HBSS. Fura-2 fluorescent emission at 510 nm resulting from 340- or 380-nm excitation was then monitored using a Flexstation (Molecular Devices, Sunnyvale, CA) plate reader. Base-line fluorescence was measured for 16 s, test compounds were then added, and fluorescence was measured for an additional 74 s. The base line-subtracted maximum 340/380-nm ratio obtained after the compound addition was then used to plot concentration-response data.

Extracellular Signal-regulated Kinase 1/2 (ERK) Assay—All ERK phosphorylation experiments were carried out using Flp-In^TM T-REx^TM stable inducible cell lines treated with doxycycline (100 ng/ml) to induce expression of the receptor of

pcDNA5 FRT/TO expression vector as described previously (11, 13, 17, 18). A chimeric form of hFFA2, in which extracellular loop 2 (ECL2) was replaced with ECL2 from hFFA3 was also as reported previously (13). All individual point mutations described were introduced using the QuikChange method (Stratagene).

Cell Culture, Transfection, and Stable Cell Lines—For experiments utilizing transient heterologous expression, HEK293T cells were used. Cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FBS at 37 °C and 5% CO₂. Transfections were carried out using polyethylenimine, and experiments were conducted 48 h post-transfection. In experiments where stable cell lines were employed, the Flp-In^TM T-REx^TM system (Invitrogen) was used to generate 293 cells with tetracycline- or doxycycline-inducible expression of the receptor of interest. To generate these cell lines, Flp-In^TM T-REx^TM 293 cells were co-transfected with the pOG44 vector and the receptor of interest in pcDNAs/FRT/TO. Transfection with pOG44 drives expression of Flp recombinase, which, in turn, allows for recombination between FRT sites in pcDNA5/FRT/TO and in the genome of the Flp-In^TM T-REx^TM 293 cells, thus allowing identification of stable inducible cells for the receptor of interest to be generated by appropriate antibiotic selection. All experiments carried out using these cells were conducted after a 24-h treatment with 100 ng/ml doxycycline to induce expression of the receptor of interest.

For experiments using differentiated mouse adipocytes, subconfluent 3T3-L1 fibroblasts were maintained in DMEM supplemented with 10% newborn calf serum and incubated at 37 °C and 10% CO₂. At 2 days postconfluence, growth-arrested cells were differentiated to 3T3-L1 adipocytes by the addition of DMEM containing 10% FBS, 0.5 mM methyl isobutylxanthine, 0.25 μM dexamethasone, 1 μg/ml bovine insulin, and 5 μM troglitazone. After a 72-h incubation (day 3), cells were supplemented with DMEM containing 10% FBS, 1 μg/ml bovine insulin, and 5 μM troglitazone. At day 6, the cells were changed into DMEM containing 10% FBS for 2 days, prior to use 9–14 days postdifferentiation.

The human-derived liposarcoma cell line SW872 was maintained in DMEM supplemented with 10% FBS and maintained at 37 °C and 10% CO₂. To differentiate these cells into adipocytes, cells were plated in 12-well plates and cultured until confluent. Culture medium was then replaced with medium supplemented with 0.5 mM methyl isobutylxanthine, 1 μM dexamethasone, 10 μg/ml bovine insulin, and 500 μM oleic acid. After a 72-h incubation, culture medium was replaced with medium containing 10 μg/ml bovine insulin and 500 μM oleic acid. Culture medium was then replaced every 48 h with fresh medium supplemented only with 500 μM oleic acid until cells were used for experiments between days 10 and 14. To confirm differentiation, cells were fixed with 4% paraformaldehyde and stained with oil red O to demonstrate lipid accumulation.

STC-1 enteroendocrine cells were maintained in DMEM supplemented with 10% FBS and maintained at 37 °C and 10% CO₂. For experiments, cells were plated in 24-well plates and
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We synthesized two compounds based on a 4-oxobutanoic acid backbone (compounds 1 and 2; Fig. 1A) that are related to structures described in the patent literature as agonists of FFA2 (16) and examined their potency and selectivity to activate FFA2 across various functional assays. First, a [35S]GTPγS incorporation assay was employed to assess Gq/11-mediated signaling (Fig. 1B). In this assay, compounds 1 and 2 both effectively stimulated incorporation of [35S]GTPγS into membranes of Flp-InTM T-RExTM 293 cells to express hFFA2-eYFP with pEC50 values of 7.14 ± 0.08 and 6.98 ± 0.12, respectively. These compounds were significantly (p < 0.001) more potent than the endogenous ligand, C3 (pEC50 = 4.27 ± 0.10), but had efficacy similar to that of C3 (114 ± 4 and 94 ± 4% of the C3 response, respectively). Because the closely related receptor FFA3 is also a Gq-coupled GPCR, the potential of 1 and 2 to stimulate [35S]GTPγS incorporation into equivalent membranes from Flp-InTM T-RExTM 293 cells induced to express hFFA3-eYFP was used to determine their selectivity for hFFA2 over hFFA3 (Fig. 1C). In these experiments, although C3 promoted binding of [35S]GTPγS with the anticipated potency (pEC50 = 3.62 ± 0.07), neither compound 1 nor 2 produced a measurable response at up to 10 μM, indicating that both compounds are highly selective for hFFA2 over hFFA3.

To examine the ability of 1 and 2 to stimulate Gq/11-mediated pathways via activation of hFFA2, changes in intracellular Ca2+ were monitored in intact Flp-InTM T-RExTM 293 cells induced to express hFFA2-eYFP (Fig. 1D). As with the [35S]GTPγS assay, both 1 and 2 did so and with significantly higher potency (pEC50 = 6.68 ± 0.06 and 6.39 ± 0.06, respectively) than C3 (pEC50 = 3.50 ± 0.10). However, in this case, both 1 and 2 were found to have significantly greater efficacy than C3 (p < 0.01), with Emax values of 132 ± 4 and 130 ± 4% of the C3 response, respectively. To assess further the selectivity of 1 and 2 for FFA2, Ca2+ assays were also employed in Flp-InTM T-RExTM 293...
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293 cells induced to express equivalent C-terminally eYFP-tagged forms of either of the long chain free fatty acid-sensitive GPCRs, hFFA1 and hFFA4, because both of these couple predominantly to G_{q/11}, G proteins. Compared with the previously described FFA1-selective ligand TUG-424 (20), which elevated Ca^{2+} with pEC_{50} = 7.19 ± 0.06 in hFFA1-eYFP-expressing cells (Fig. 1E), 2 produced no measurable response at concentrations up to 10 μM, whereas 1 did generate a Ca^{2+} response but only at the highest concentrations tested (3–10 μM). Because FFA1 agonists have at times been shown to possess PPAR\gamma activity, we also tested whether 1 displayed agonism or antagonism of PPAR\gamma (testing performed by Cerep, Poitiers, France). In these experiments, 1 (10 μM) was found to have no measurable activity either as an agonist or antagonist (data not shown). When experiments were conducted on cells induced to express hFFA4-eYFP (Fig. 1F), the previously described FFA4-selective agonist TUG-891 (17) effectively increased intracellular Ca^{2+} (pEC_{50} = 6.68 ± 0.08), whereas neither 1 nor 2 had any effect at concentrations up to 10 μM, indicating that 1 and 2 are extremely selective for FFA2 over both FFA1 and FFA4.

FFA2 activation has also been linked to increased phosphorylation of ERK (13), a pathway that may be initiated via activation of G_{i} or G_{q/11}, as well as by G protein-independent signaling pathways. Therefore, we explored ERK phosphorylation stimulated by C3, 1, and 2 in cells induced to express hFFA2-eYFP (Fig. 1G). In these experiments, C3 stimulated increased ERK phosphorylation (pEC_{50} = 3.56 ± 0.11), as did both 1 (pEC_{50} = 6.94 ± 0.16) and 2 (pEC_{50} = 6.48 ± 0.14). Both 1 and 2 were significantly more potent than C3 (p < 0.001), but, as in the [^{35}S]GTP\gammaS assay, neither 1 or 2 displayed efficacy significantly different from that of C3. The final FFA2 signaling pathway examined was the recruitment of β-arrestin-2. Previous work has shown that activation of hFFA2 does stimulate recruitment of β-arrestin-2 (21), an event that may be linked with GPCR internalization and/or G protein-independent signaling. In order to assess this, a BRET-based assay was employed in which hFFA2-eYFP and β-arrestin-2 Renilla luciferase were co-expressed transiently in HEK293T cells (Fig. 1H). In this assay, C3 stimulated β-arrestin-2 recruitment to human FFA2-eYFP (pEC_{50} = 3.22 ± 0.10), and so did both 1 (pEC_{50} = 5.72 ± 0.10).

FIGURE 1. Compounds 1 and 2 are potent and selective agonists of FFA2. Chemical structures of compounds 1 and 2 are shown in A. Concentration-response curves for C3, 1, and 2 across various function assays are shown for hFFA2 (B, D, G, and H), hFFA3 (C), hFFA1 (E), and hFFA4 (F). Assays chosen represent [^{35}S]GTP\gammaS to assess G_{q/11} coupling (B and C), Ca^{2+} mobilization to assess G_{q/11} signaling (D–F), ERK phosphorylation (G), and β-arrestin-2 recruitment (H). Error bars, S.E.
Compounds 1 and 2 were significantly more potent than C3 (p < 0.001), and, although not statistically significant (p = 0.05), there was a trend toward increased efficacy for both 1 and 2 compared with C3 in this assay.

Compounds 1 and 2 Bind to the Orthosteric Pocket of FFA2—

The only series of synthetic small molecule agonists previously published and with reasonable potency at FFA2 are allosteric in nature and, as such, bind to a different site than the SCFAs. These compounds, exemplified by 4-CMTB, display reduced signaling in some pathways and lack the carboxylate moiety that is critical in the SCFAs for interaction with the orthosteric binding site (13). Because 1 and 2 both contain a carboxylate functionality and display broadly similar signaling responses to C3 across all assays tested, we hypothesized that these compounds are probably orthosteric agonists of FFA2. To assess this, we first examined whether the carboxylate was critical to ligand function by generating both methyl and tert-butyl ester analogs of 2.

Compounds 1 and 2 are orthosteric agonists of FFA2. Concentration-response curves at hFFA2 for C3, 2, a methyl ester of 2 (ME-Cmp 2), and a tert-butyl ester of 2 (TBE-Cmp 2) are shown in [35S]GTPγS (A) and β-arrestin-2 recruitment (B) assays. The ability of the hFFA2 antagonist CATPB as well as methyl and tert-butyl esters of 2 to inhibit an EC50 concentration of compound 2 in the β-arrestin-2 recruitment assay is shown in C. Concentration-response curves were generated using wild type and H140A, R180A, H242A, and R255A mutants of hFFA2 in the β-arrestin-2 recruitment assay, and the results are shown for C3 (D), 1 (E), and 2 (F). Antagonism experiments where concentration-response curves were generated to agonist in the presence of increasing fixed concentrations of CATPB are shown using C3 (G), 1 (H), and 2 (I) as agonists. Error bars, S.E.

and 2 (pEC50 = 5.35 ± 0.09). As in the other assays, both 1 and 2 were significantly more potent than C3 (p < 0.001), and, although not statistically significant (p > 0.05), there was a trend toward increased efficacy for both 1 and 2 compared with C3 in this assay.

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residues is important for function of these ligands. As additional support for an orthosteric mode of binding of 1 and 2, we compared the ability of CATPB to limit the function of C3, 1, and 2. In ERK phosphorylation assays, increasing concentrations of CATPB resulted in higher concentrations of each agonist being required to cause half-maximal promotion of phosphorylation (Fig. 2, G, H, and I). Furthermore, in each case, the effect of CATPB was fully overcome by sufficiently high concentrations of the agonist. Global curve fitting of the data yielded similar pA2 values for CATPB of 7.23 ± 0.09, 7.58 ± 0.15, and 7.24 ± 0.14 for experiments conducted with C3, 1, and 2, respectively. These data are consistent with competitive interactions between each agonist and CATPB and, therefore, with C3, 1, and 2 sharing an overlapping binding site.

Compounds 1 and 2 Are Not Modulated Allosterically by 4-CMTB—There were, however, differences between the behavior of compounds 1 and 2 and C3. The compound 4-CMTB is recognized as an ago-allosteric modulator of FFA2 (12, 13). As a result, as well as direct activation of signaling, increasing submaximal, fixed concentrations of 4-CMTB increase the potency of C3 (Fig. 3A), and vice versa (Fig. 3B). By contrast, increasing fixed concentrations of 4-CMTB did not alter the potency of 2 (Fig. 3C), nor did fixed concentrations of 2 alter the potency of 4-CMTB (Fig. 3D). This observation could be consistent with either of two possibilities: 1) with so-called “probe dependence” (22) of 4-CMTB allosterism, such that it modulates the binding of C3 but not 2, or 2) with the binding sites of 4-CMTB and 2 overlapping each other. Although C3 and 2 appear to bind to the same site, whereas 4-CMTB and C3 do not, given the very small size of C3 relative to 2, it is still at least conceivable that 2 could overlap the binding sites of both C3 and 4-CMTB. To address this possibility, we first examined whether the hFFA2 antagonist CATPB was competitive with 4-CMTB by measuring the 4-CMTB concen-

FIGURE 3. The FFA2 allosteric ligand 4-CMTB does not modulate function of compounds 1 and 2. Concentration-response curves for C3 in the β-arrestin-2 recruitment assay in the presence of increasing fixed concentrations of 4-CMTB are shown in A. The reciprocal experiment showing the concentration response to 4-CMTB with increasing fixed concentrations of C3 is shown in B. Similar experiments are shown using a [35S]GTP-γ-S assay with 2 and 4-CMTB (C and D). An antagonism experiment measuring the concentration response to 4-CMTB in a pERK assay in the presence of increasing fixed concentrations of CATPB is shown in E. In F, a competition experiment is shown using a fixed concentration (10 μM) of the full agonist 1 with increasing concentrations of the partial agonist 4-CMTB in the β-arrestin-2 BRET assay. Error bars, S.E.
tration response in the presence of increasing concentrations of CATPB (Fig. 3E). Interestingly, in these experiments, CATPB did inhibit the 4-CMTB response; however, unlike with C3, 1, and 2, where the inhibition resulted in reduced potency with no effect on efficacy, the inhibition of 4-CMTB by CATPB had the opposite effect, reducing efficacy with no effect on potency. Because we had previously shown that CATPB is competitive with C3, 1, and 2 and therefore does not bind irreversibly, this pattern of CATPB inhibition of the ERK response to 4-CMTB indicates that these two molecules must bind to separate sites and that CATPB is a negative allosteric modulator of 4-CMTB efficacy, further strengthening the case that 2 and 4-CMTB bind to distinct sites. Finally, to examine this more directly, we took advantage of the fact that 4-CMTB is only a partial agonist in the BRET assay. Therefore, if competitive, 4-CMTB would be expected to antagonize the response to the full agonist, and indeed there was a small increase in response, with increasing concentrations of 4-CMTB. Taken together, these results suggest strongly that both 1 and 2 are binding to a distinct site to 4-CMTB and that, therefore, the lack of allosterism between these two compounds represents a probe dependence in the ability of 4-CMTB to modulate function of FFA2 orthosteric agonists.

**ECL2 Is Critical to the Binding and Selectivity of Compounds 1 and 2**—We next considered whether the probe-dependent effect of 4-CMTB might be instructive in defining the mode or location of binding of 1 and 2. A recent study demonstrated that replacement of ECL2 of hFFA2 with the equivalent region from hFFA3 has no negative effect on the potency of C3 or 4-CMTB but essentially eliminates allosteric communication between the binding sites of these ligands (13). We considered, therefore, if the lack of allosteric interactions between 4-CMTB and 2 reflects that these ligands interact with ECL2 in such a way as to prevent allosteric communication. We tested this by comparing the potency of C3, 1, and 2 in a β-arrestin-2 recruitment assay at wild type hFFA2 and the variant in which ECL2 is replaced by this sequence from hFFA3 (13). As observed previously, this modification resulted in a modest increase in the potency of C3, pEC50 of 3.84 ± 0.013 at the ECL2 swap mutant compared with 3.23 ± 0.07 at wild type (Fig. 4A). By contrast, for both 1 (Fig. 4B) and 2 (Fig. 4C), marked reductions in potency were observed. In addition to showing broadly that ECL2 is important for allosteric communication between C3 and 4-CMTB, Smith et al. (13) also identified a single point mutation in ECL2 of hFFA2, L173A, which also eliminated allosteric communication. This alteration did not affect the potency of C3 in the β-arrestin-2 interaction assay (Fig. 4D) but resulted, however, in a significant (p < 0.01) decrease in potency for both 1 (pEC50 of 5.29 ± 0.11 at L173A compared with 5.99 ± 0.07 at wild type (Fig. 4E)) and 2 (pEC50 values of 4.93 ± 0.11 and 5.54 ± 0.08 at L173A and wild type, respectively (Fig. 4F)).

Because the hFFA2/hFFA3 ECL2 swap mutant substantially lost potency to both 1 and 2, we next explored if ECL2 represents a key region in defining the selectivity of these compounds for hFFA2. To do so, we employed sequence alignment to identify residues in ECL2 conserved between FFA1 and FFA3, but not FFA2. This search yielded one obvious residue, where a
glutamic acid is present in hFFA1 (position 145) and hFFA3 (position 152) but is a glutamine in hFFA2 (position 148). Previous work has shown that alteration of Glu-145 to alanine in FFA1 breaks an extracellular ionic lock and increases ligand-independent constitutive activity (23). We therefore generated both Q148E and Q148A mutants of hFFA2 to test whether a similar ionic lock might influence the FFA2 selectivity of 1 and 2. These mutants were first assessed in the /H9252-arrestin-2 assay with C3 (Fig. 5A), 1 (Fig. 5B), and 2 (Fig. 5C). Although the potency of C3 was reduced some 10-fold at the Q148E mutant (pEC$_{50}$ = 3.27 ± 0.10 at wild type and 2.34 ± 0.11 at Q148E), both 1 and 2 were virtually inactive at this mutant. In contrast, C3 did not show reduced potency at the Q148A mutant (pEC$_{50}$ = 3.35 ± 0.08), and although both had reduced efficacy, neither 1 nor 2 displayed significant loss in potency at the Q148A mutant. Because the potency of agonists in the β-arrestin-2 assay generally reflects receptor occupancy and hence is a surrogate measure of affinity, this suggests that the presence of a glutamate but not alanine at this position significantly reduces the binding of both 1 and 2 to hFFA2. To explore this further, we also employed the ERK assay because this often provides higher agonist potency values due to the effects of receptor reserve (24). Here, the potency of C3 (Fig. 5D) was little affected at Q148E compared with wild type hFFA2 (pEC$_{50}$ = 3.89 ± 0.11 for wild type and 3.76 ± 0.07 at Q148E), whereas for both 1 (Fig. 5E) and 2 (Fig. 5F), there was again a nearly complete loss of agonist function. These results confirm that the lack of glutamate at this position in FFA2 is critical to the binding of 1 and 2 and therefore potentially could be involved in the selectivity of these compounds. To test whether this position contributes to the selectivity, we generated the reciprocal E145Q and E154Q mutants of hFFA1 and hFFA3, respectively. Interestingly, although the E145Q mutant did not alter the potency of the FFA1 agonist TUG-424 in a β-arrestin-2 recruitment assay (Fig. 5G), it did result in a clear gain of function for 1 when compared against wild type hFFA1.

**FIGURE 5.** The absence of a glutamate residue in ECL2 of FFA2 that is present in both FFA1 and FFA3 defines the FFA2 selectivity of compound 1 and 2. Concentration responses are shown using the β-arrestin-2 recruitment assay for C3 (A), 1 (B), and 2 (C) at wild type, Q148E, and Q148A variants of hFFA2. Similar ERK experiments are shown with wild type and Q148E hFFA2 (D–F). β-Arrestin-2 recruitment to wild type and the E145Q mutant of hFFA1 is shown for TUG-424 (G), 1 (H), and 2 (I). Error bars, S.E.
but did not significantly alter the response for 2 (Fig. 5I). Similar experiments were conducted on the equivalent E154Q mutant of hFFA3 in [35S]GTPγS experiments; however, this mutant was found to completely lack function to all ligands tested (data not shown).

Molecular Modeling Suggests Further Interactions within the Transmembrane Domains of hFFA2 for Compounds 1 and 2—Predictions and modeling of extracellular loop regions of GPCRs remain highly speculative and challenging, particularly for receptors well removed phylogenetically from those for which atomic level structures are available. By contrast, the overall architecture of the transmembrane helix bundle is generally well preserved among GPCRs. As such, confidence in homology models of these regions is increasing, and well validated models are becoming widely used in suggesting likely ligand-receptor interactions within these regions. FFA2 homology models have provided important insights into modes of orthosteric ligand binding and differences in this between species orthologs (13, 18, 21). We therefore employed homology modeling of hFFA2 and ligand docking methods to suggest additional residues that might contribute to the enhanced potency of 1 and 2 compared with C3 (Fig. 6, A and B). Based on such models, five transmembrane residues and one additional ECL2 residue were highlighted for subsequent analysis. Each of these was mutated to alanine and assessed with C3, 1, and 2 in a [35S]GTPγS incorporation assay (Table 1). From these studies, it was apparent that several mutants (i.e. Y90A3.29, Y165AECL2, and Y238A6.51) displayed significantly lower potency to each ligand, indicating their likely contribution to the orthosteric binding pocket by forming hydrophobic and aromatic interactions. In addition, Tyr-165 might also contribute in steering the ligand into the binding cavity. The I145A4.61 mutant appeared to not be expressed and produced no response to any ligand. Most interestingly in the current context, although the V179A5.38 mutant did not alter potency for C3 (Fig. 6C), it did result in a significant reduction in potency for both 1 (p < 0.01; Fig. 6D) and 2 (p < 0.05; Fig. 6E). Our docking results predict that the cyclopropyl ring of 1 and 2 forms hydrophobic contacts with V179A5.38. Overall, the ligands 1 and 2 have more contacts with the residues of helices 3, 5, and 6 in the extracellular part of the helical bundle that remarkably impact the potency of the ligands when the residues are mutated to alanine.

Compounds 1 and 2 Display Differential Function at Rodent Orthologs of FFA2—Recent work has demonstrated that FFA2 displays significant species ortholog variation with respect to both the endogenous SCFA ligands and to the synthetic antagonist CATPB (11, 21). Considering this, it was important to
establish whether similar species ortholog variation might affect the usefulness of 1 and 2 in non-human systems. For this, we compared the activity of C3, 1, and 2 at each of the human, mouse, and rat orthologs of FFA2 across multiple assay end points (Table 2). Initially, we examined the responses of C3 (Fig. 7A), 1 (Fig. 7B), and 2 (Fig. 7C) at each ortholog in the G_{i/o}-dependent [35S]GTPγS assay. Consistent with previous reports (11), C3 displayed significantly (p < 0.01) lower potency at mFFA2 and rFFA2 compared with hFFA2. Interestingly, although 1 also displayed a reduction in potency (p < 0.05) of approximately the same magnitude as C3 at mFFA2 (3.9-fold) and rFFA2 (5.4-fold) compared with hFFA2, 2 displayed a much larger reduction in potency at both mFFA2 (20-fold) and rFFA2 (22-fold) compared with hFFA2. Similar results were obtained when these compounds were assessed at the species orthologs in a Ca^{2+} assay to measure G_{i/o}-coupled signaling (Fig. 7, D–F), with a much greater reduction in potency observed for 2 than for C3 or 1. In addition, because 1 was found to have some activity at hFFA1, we also tested these compounds at mFFA1 in a Ca^{2+} assay, finding no activity for 2 and only weak activity for 1, requiring at least 10 μM before any Ca^{2+} response was observed (data not shown).

In ERK phosphorylation assays, similar potencies were observed for C3 at hFFA2, mFFA2, and rFFA2 (Fig. 7G), and a small reduction in potency at the rodent orthologs was observed with 1 (Fig. 7H), whereas, once more, a substantial reduction in potency was seen for 2 (Fig. 7I). We have recently demonstrated that utilizing the variation in pharmacology between species orthologs of FFA2 can be a valuable approach in defining key aspects of receptor function (11, 21). Therefore, we next examined whether we could take advantage of the species differences in the function of C3, 1, and 2 to help further define important and differential aspects of their function. First we considered a possible explanation for the broad decrease in potency we observed across all ligands. Recently, we have demonstrated that the absence in hFFA2, or presence in mFFA2, of an ionic lock between a glutamate residue at position 159 in ECL2 and arginine residues within the orthosteric binding pocket accounts for both reduced constitutive activity and lower SCFA potency at the mouse ortholog (11). Therefore, we hypothesized that this same residue might account for the broad decrease in potency we observed across all ligands. To assess this, we compared the ability of C3 (Fig. 8A), 1 (Fig. 8B), and 2 (Fig. 8C) to produce a Ca^{2+} response via either wild type hFFA2 or a G159E hFFA2 mutant. In these experiments, despite similar levels of expression, C3 displayed significantly reduced potency at the G159E mutant compared with wild type (pEC_{50} = 4.26 ± 0.06 at wild type and 3.69 ± 0.06 at G159E). Similarly, 1 displayed reduced potency (7.06 ± 0.11 at wild type compared with 6.32 ± 0.07 at G159E), as did 2 (6.66 ± 0.06 at wild type and 6.28 ± 0.07 at G159E). Interestingly, the magnitude of potency decreases observed at the G159E mutant were similar for C3 (3.7-fold) and 1 (5.5-fold) to the potency decreases observed for these compounds at the two rodent orthologs compared with human FFA2 in the Ca^{2+} assay (~4.5-fold for C3 and ~12-fold for 1). In contrast, the potency decrease for 2 was only 2.4-fold at the G159E hFFA2 mutant, substantially less than the decrease observed for rodent compared with human orthologs in the Ca^{2+} assay using this ligand (416-fold for mFFA2 and 154-fold for rFFA2). These results indicate that the presence or absence of this “ionic lock” accounts for the broad decrease in potency seen for all agonist ligands at the rodent orthologs but does not account for the substantially greater loss in potency observed with 2.

To attempt to understand this selective reduction in potency for 2, we re-examined the homology models with 1 or 2 docked (Fig. 6, A and B) with the aim of identifying key residues predicted to be involved in ligand interaction in the human ortholog that were not conserved in the mouse and rat orthologs. This yielded only one residue in proximity to the defined orthosteric pocket, serine 86^{3,29} in hFFA2, which is glycine in both rFFA2 and mFFA2. We hence generated an S86G mutant of hFFA2 and examined its function across several assays. Initially, we tested this mutant in the β-arrestin-2 recruitment assay, where, although this alteration had no significant effect on the potency of either C3 (Fig. 9A) or 1 (Fig.
9B), it did result in a significant reduction of potency of 2 (Fig. 9C). Similar results were observed when the S86G hFFA2 mutant was tested in both Ca\textsuperscript{2+} (Fig. 9, D–F) and ERK (Fig. 9, G–I) assays. Although the decreases in potency for 2 at S86G compared with wild type hFFA2 (7-fold in the β-arrestin-2 assay, 10-fold in the Ca\textsuperscript{2+} assay, and 8-fold in the pERK assay) were not as large as the decrease observed for 2 at the rodent ortholog compared with human FFA2, this site clearly contributes to the specific reduction in potency for 2 in the rodent orthologs. Indeed, it is likely that a combination of the effects of the presence of Gly-86 and Glu-159 residues in rat and mouse FFA2 accounts for the bulk of potency loss observed for 2 at the rodent orthologs.

Compounds 1 and 2 Demonstrate FFA2-mediated Inhibition of Lipolysis in Human Adipocytes, whereas Only Compound 1 Effectively Inhibits Lipolysis in Mouse Adipocytes—Having defined the potency and activity of C3, 1, and 2 in cells trans-
ected to express each of the human, mouse and rat orthologs of FFA2 in a heterologous manner, this provided the opportunity to define the specific role of FFA2 in functional responses of cells endogenously expressing the receptor. To do so, we examined the effects of these ligands on stimulated lipolysis in both a human-derived liposarcoma cell line, SW872, and in mouse 3T3-L1 cells, after differentiating each cell line into adipocyte-like phenotypes. Initial RT-PCR studies on mRNA isolated from differentiated SW872 and 3T3-L1 cells confirmed expression of FFA2 but not FFA3 in each cell type (Fig. 10, A and B).

To examine lipolysis in the differentiated SW872 line, cells were treated with 10 µM forskolin to stimulate lipolysis, which was then measured by assessing the concentration of glycerol released into the cell culture supernatant (Fig. 10, C–E). Unexpectedly, treatment with C3 (10 mM) enhanced lipolysis in these cells markedly. However, this effect was not blocked by the hFFA2 antagonist CATPB. In contrast, both 1 (10 µM) and 2 (10 µM) significantly reduced (p < 0.05) glycerol release to 79 ± 9 and 79 ± 15% of the forskolin alone control, respectively. Importantly, for both 1 and 2, the effect of the ligand was completely reversed by the hFFA2 antagonist CATPB.

To assess lipolysis in the 3T3-L1 adipocytes, we first treated cells with high concentrations of C3 (10 mM), 1 (10 µM), or 2 (10 µM) and measured inhibition of isoprenaline-stimulated glycerol release (Fig. 10F). Both C3 (p < 0.01) and 1 (p < 0.05) significantly inhibited glycerol release, whereas 2 did not. Although failing to reach statistical significance, C3 tended to produce a greater inhibition of glycerol release, 54 ± 10%, compared with only 38 ± 5.0% for 1. Finally, in concentration-response experiments, C3 inhibited glycerol release with a pIC50 = 3.24 ± 0.36 (Fig. 10G), whereas 1 was some 480-fold more potent (pIC50 = 5.92 ± 0.43) (Fig. 10H), and 2 was again without significant effect (Fig. 10I). Because the only FFA2 antagonist currently available, CATPB, is highly human-selective, it was not possible to perform studies to block these effects with an FFA2 antagonist. However, because we had demonstrated that 1 has low potency agonism at FFA1, we did confirm that the effect of compound 1 was not blocked by the FFA1 antagonist GW1100 (data not shown). These results demonstrate that FFA2 inhibits lipolysis in both human- and rodent-derived adipocyte models and that, despite their chemical similarity, compound 1 but not compound 2 is a novel and valuable tool to specifically assess the function of FFA2 in murine cells and tissues.

Compound 1 Stimulates GLP-1 Release from Murine STC-1 Enteroendocrine Cells—Having shown that 1 can be used to assess FFA2 function in mouse cells endogenously expressing

FIGURE 9. The reduced function of compound 2 at rodent FFA2 orthologs results from the presence of glycine at position 86, compared with serine at this position in hFFA2. Concentration-response curves are shown at wild type and S86G hFFA2 in β-arrestin-2 (A–C), Ca2+ mobilization (D–F), and pERK (G–I) assays using C3, 1, or 2. Error bars, S.E.
the receptor, we finally set out to use this compound to selectively examine FFA2 versus FFA3 function in cells endogenously co-expressing both receptors. For this, we examined the release of GLP-1 from the mouse-derived enteroendocrine cell line STC-1, which we confirmed by RT-PCR to express both FFA2 and FFA3 (Fig. 11A). Treatment of these cells with C3 (1 or 10 mM) or 1 (10 μM) significantly increased GLP-1 secretion \( (p < 0.001) \). Interestingly, there was no difference in the maximum GLP-1 response produced by C3 or 1, indicating that C3 is probably stimulating GLP-1 release primarily through FFA2 and not FFA3 in this cell line.

**DISCUSSION**

In cases where closely related GPCRs respond to the same endogenous ligand(s), developing selective synthetic agonists or antagonists capable of differentiating between the receptors is critical to defining their individual biological roles. This is the case for FFA2 and FFA3 because each responds to the same endogenous SCFA ligands. Differences in the rank order of potency have been described for the SCFAs at FFA2 compared with FFA3 (4). However, the potency and selectivity of these compounds remains low. As a result, very high concentrations of SCFAs must be used, and, because the SCFAs are known to have additional off-target effects, including, for example, inhibition of histone deacetylases (25), this has made attributing the biological effects of SCFAs directly to FFA2 or FFA3 challenging. Further complicating the issue is the recent observation that C2, a SCFA with a degree of selectivity for hFFA2 over hFFA3, does not show this selectivity at the rodent orthologs of these receptors (11), making it effectively impossible to clearly define FFA2 versus FFA3 functions using SCFAs in rodent models without knock-out approaches. Although such knock-out studies have been informative in this respect (8, 9, 14, 26, 27), there is also evidence that knock-out of FFA3 results in reduced FFA2 expression (8), clearly complicating the interpretation of data based upon such animals. Moreover, in studies on the possible contribution of FFA2 to gut inflammatory disease, two distinct lines of FFA2 knock-out mice generated entirely contradictory results, in that a model of colitis, one line dis-
played reduced and the other exaggerated inflammation (26, 27). To address such issues, there is a clear need for synthetic ligands that can effectively differentiate between FFA2 and FFA3 in both human and non-human experimental systems. Although several synthetic ligands have been reported with selectivity for either FFA2 or FFA3 (10–13, 28), each of these suffers from either poor potency, low selectivity, altered function due to an allosteric mode of action, or lack of function at non-human orthologs of the receptors. Therefore, our description of compounds 1 and 2 as the first potent and selective orthosteric agonists of hFFA2 and of 1 as the first orthosteric compound displaying reasonable activity at rodent orthologs of FFA2 suggests that these compounds will be extremely useful tools in future studies designed to define the specific biological role of this receptor.

To demonstrate the usefulness of compounds 1 and 2 in this respect, we demonstrated that both compounds inhibit lipolysis in human-derived SW872 cells, whereas only compound 1 does in murine 3T3-L1 adipocytes. Previous studies have indicated that SCFAs inhibit lipolysis in adipocytes and that this occurs via a G\textsubscript{i/o}-dependent mechanism (9, 12). However, at least in SW872 cells, this was not the case because C3 actually markedly enhanced lipolysis, although via a mechanism that did not involve FFA2 because this effect was not blocked by the hFFA2 antagonist CATPB. This observation indicates that C3 has additional cellular targets in SW872 cells and that any effect on lipolysis it may have through FFA2 is effectively masked by these much larger “off-target” effects. Clearly, this highlights a major challenge to using only SCFA ligands to assess FFA2 function that can only be addressed through the development of more potent and selective ligands, such as compound 1.

In the 3T3-L1 cells, it is interesting to note that the potency of both C3 and 1 to inhibit lipolysis (3.24 ± 0.36 and 5.92 ± 0.43, respectively) was very similar to the values we measured at mFFA2 for these compounds in the [\textsuperscript{35}S]GTP\gammaS assay (3.57 ± 0.10 and 6.27 ± 0.13, respectively), a measure of G\textsubscript{i/o}-dependent signaling. We also noted that, although not quite reaching statistical significance, there was a trend toward C3 producing a greater maximum inhibition of lipolysis than 1. This could reflect either that 1 acts as a partial agonist relative to C3 in this signaling response or that C3 also has additional non-FFA2 targets in murine adipocytes that also modulate lipolysis. However, given that we observe little sign of partial agonism for 1 in other assays employing mFFA2, off-target effects of C3 perhaps represent the more likely explanation. Although we were unable to detect mFFA3 mRNA in the differentiated 3T3-L1 model adipocytes, certain studies have reported FFA3 expression in adipocytes (5), whereas another noted altered adipocyte function in FFA3 knock-out mice (8). Indeed, although the expression of FFA3 in adipocytes remains controversial (4), it is at least conceivable that, despite a lack of detection of mRNA, low levels of mFFA3 protein may be expressed in these cells and could account for the greater efficacy of C3 relative to 1.

Another critical feature of compound 1 in addition to retaining activity at the rodent orthologs of FFA2 is that it is extremely selective for FFA2 over FFA3. This should allow for the delineation of FFA2-mediated responses in cells that co-express both receptors as we were able to do in demonstrating that FFA2, rather than FFA3, mediates GLP-1 release from STC-1 cells. The similar efficacy of C3 and 1 in this assay suggests that despite expressing FFA3, GLP-1 secretion by these cells in response to C3 is predominantly through FFA2, a conclusion consistent with previous work (29).

It is somewhat surprising that such substantial differences in activity were observed between 1 and 2 at the rodent orthologs of FFA2, given their structural similarity. Indeed, the primary structural difference between the two compounds (in addition to different positions for their chlorine substituents) is the presence of a phenyl substituent in 1, which is cyclopentyl in 2. Interestingly, closer examination of the docking of these compounds to our homology model of hFFA2 shows that this phenyl/cyclopentyl substituent is predicted to be in close proximity to Ser-86, the position we have found that when mutated to glycine (the amino acid present in rodent forms of FFA2) results in profoundly different functional consequences between species orthologs of FFA2, any future ligands in this series will also need to be assessed empirically across species before being considered as useful tool compounds.
Our observation that ECL2 contributes to both the binding and selectivity of compounds 1 and 2 is also of note. Interestingly, the structure of the extracellular loops, and in particular ECL2, varies greatly among the GPCRs for which crystal structures are available (30, 31). However, despite (or perhaps because of) this, there are several examples of GPCRs where ligand selectivity is defined at least in part by ECL2 (30, 31). It is, therefore, perhaps not surprising that mutation of a single residue in ECL2 of FFA2, Gln-148, to glutamate, as it is in FFA1 and FFA3, could result in loss of potency to these ligands, whereas the reciprocal mutation, at least in FFA1, would result in a gain in function. The specific mechanism underlying how this residue may dictate FFA2 versus FFA1 selectivity may also be of interest. Previous work has shown that an ionic lock between this position in FFA1 and one of two positively charged arginine residues within the fatty acid binding pocket of this receptor regulates constitutive activity of FFA1 (23). We have also recently shown that similar ionic locks involving additional positions within ECL2 regulate constitutive activity in human and rodent orthologs of FFA2 and FFA3 (11). Therefore, it is tempting to hypothesize that mutation of Gln-148 to glutamate in hFFA2 allows for the formation of an ionic lock at this position similar to that present in FFA1 and that this effectively closes off the binding pocket to 1 and 2. By contrast, in FFA1, mutation of Glu-145 to glutamate would alleviate this ionic lock, thus allowing for increased binding by these ligands. To more directly test this, we also generated the Q144A mutant of hFFA2 and found that, unlike the Q148E mutant, the ala nine mutant did not show reduced potency to 1 and 2. This observation indicates that it is the gain of the negatively charged glutamate, and not the loss of glutamine that results in reduced potency to 1 and 2, a finding that appears to support the ionic lock hypothesis.

It was interesting that 1 did possess some activity at wild type FFA1, as well as increased activity at the E145Q mutant. This is perhaps surprising, given that there is no overlap in the endogenous ligands activating these receptors, nor have any previously described synthetic ligands for FFA1 been found to have activity at FFA2 (20, 32–37). The fact that 1 does show some activity at FFA1 indicates that there is at least some similarity in the tertiary structure of the binding pockets of these receptors and perhaps indicates that counterscreening FFA1 agonists against FFA2 and FFA3 as well as for FFA4 and PPARγ activity should be more routinely carried out in the future. The observation that 1 was more active at both wild type and E145Q FFA1 than was 2 is also strikingly similar to the pattern of activity these compounds have at rodent FFA2. Because we have demonstrated that a glycine instead of serine at position 86 accounts for this effect in rodent FFA2, we note that FFA1 possesses alanine at this 3.29 position, a residue that is also nonpolar and smaller compared with the serine present in hFFA2. This may suggest that the preference for 1 over 2 in FFA1 is perhaps dictated by factors similar to those that dictate this same preference in rodent FFA2.

We have described the first potent and selective orthosteric agonists of FFA2 and defined the molecular basis for their interaction with the receptor. Our demonstration that at least one of these ligands can be used to delineate specific FFA2 function in murine cell systems highlights the importance these compounds may have, particularly in possible future, preclinical drug development and proof-of-principle studies at this receptor. FFA2 represents an interesting therapeutic target for the treatment of various metabolic and inflammatory conditions (28), for which development has been slowed primarily by a lack of reasonably potent and selective ligands (4). Therefore, the description of compound 1 as tool to study FFA2 function as well as our detailed examination of the molecular basis for its potency and selectivity, which should aid in future ligand development at these receptors, will be invaluable to exploring the potential of FFA2 as a therapeutic target.

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Selective FFA2 Orthosteric Agonists


Defining the Molecular Basis for the First Potent and Selective Orthosteric Agonists of the FFA2 Free Fatty Acid Receptor
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