Molecular dynamics-guided discovery of an allosteric modulator for GPR40/FFAR1

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Edited by Barry Honig, Howard Hughes Medical Institute, Columbia University, New York, NY, and approved February 14, 2019 (received for review June 28, 2018)

The long-chain fatty acid receptor FFAR1/GPR40 binds agonists in both an interhelical site between the extracellular segments of transmembrane helix (TM)-III and TM-IV and a lipid-exposed groove between the intracellular segments of these helices. Molecular dynamics simulations of FFAR1 with agonist removed demonstrated a major rearrangement of the polar and charged anchor point residues for the carboxylic acid moiety of the agonist in the interhelical site, which was associated with closure of a neighboring, solvent-exposed pocket between the extracellular poles of TM-I, TM-II, and TM-VII. A synthetic compound designed to bind in this pocket, and thereby prevent its closure, was identified through structure-based virtual screening and shown to function both as an agonist and as an allosteric modulator of receptor activation. This discovery of an allosteric agonist for a previously unexploited, dynamic pocket in FFAR1 demonstrates both the power of including molecular dynamics in the drug discovery process and that this specific, clinically proven, but difficult, antidiabetes target can be addressed by chemotypes different from existing ligands.

Significance
Structures of G protein-coupled receptors (GPCRs) in complex with ligands mainly provide frozen pictures with little information about the actual molecular mechanism of action of the ligand in the normally highly dynamic receptor. Through computer-based molecular dynamics simulations of a receptor for long-chain fatty acids, free fatty acid receptor 1 (FFAR1), we discover that an unoccupied, solvent-exposed pocket closes on removal of the lipid-like agonist; that is, during a major conformational change of the receptor. Importantly, a compound designed to prevent closure of this previously unrecognized, dynamic pocket was identified through structure-based virtual screening and shown to function as an allosteric agonist for the receptor. The study demonstrates that molecular dynamics simulations can be used in drug discovery to identify different modes of stabilizing specific receptor states.


The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1811066116/-/DCSupplemental.

Published online March 14, 2019.

www.pnas.org/cgi/doi/10.1073/pnas.1811066116
well as mutational analyses, we describe an ago-allosteric modulator and highlight the extracellular site as an avenue for the structure-based design of FFAR1 ligands.

Results

MD Simulations Reveal a Major Rearrangement of Polar Anchor-Point Residues. Initially, we performed 600 ns of unbiased MD simulations and compared representative frames that were obtained by geometric clustering. Although FFAR1 is crystallized in complex with agonists, the overall structure of the intracellular part of the helical bundle of FFAR1 is rather similar to the structure of various published inactive receptor conformations, conceivably because of thermostabilizing mutations, which allow for high-affinity agonist binding, but also prevent the receptor from signaling in solution (9). In analogy to previous MD simulation studies of GPCRs in their inactive form (14, 15), we did not observe any substantial change of the overall helical configuration at the intracellular side, which remained similar to the inactive-like state of, for example, the 2-adrenergic receptor (16) (Fig. S2 A). However, during the course of the MD simulation of FFAR1 in the presence of TAK-875, the intracellular segment of TM-VI moved slightly (~4 Å) toward TM-V, which is likely related to the insertion of ICL3 in the FFAR1 model, which in the X-ray crystal structure is replaced with a fusion protein (PDB 4PHU).

At the extracellular solvent-exposed face of the FFAR1 receptor, we observed a major rearrangement of residues of the polar network during the MD simulations in the absence of TAK-875, which was not observed in the presence of the stabilizing carboxylic acid group of the agonist (Fig. 1 C and D). This disruption lead to a substantial conformational change of not only the side chains of the polar network but also a shifting movement of the extracellular part of TM-I, TM-II, and TM-VII, ultimately resulting in total closure (<10 Å) of the solvent-exposed extracellular site after ~340 ns of simulation time (Fig. 1 B and D). The movement of TM-VII was facilitated by the structural flexibility induced by the proline of the conserved, central NPxxY motif. In contrast to these changes observed in the extracellular site, the hydrophobic tunnel-like part of the interhelical TAK-875 site remained surprisingly stable after the ligand had been removed (Fig. 1 B and D). We compared the structural heterogeneity of all the MD trajectory structures by geometric clustering of residues forming both the interhelical and the extracellular site. Only 35 clusters were found for the trajectory in the presence of TAK-875, and 64% of all trajectory conformations belonged to a single large cluster based on a 1 Å rmsd cutoff. In comparison with this, more than a 100 different clusters were identified for the trajectory in the absence of TAK-875. This further illustrates the structural dynamics that occur in the absence of TAK-875 (SI Appendix, Fig. S3).

Metadynamics Simulations of the Rearrangement of the Polar Network. The MD simulations described here provide an unbiased view of the dynamics of FFAR1. We supplemented these by metadynamics simulations that allow us to map out the conformational free-energy landscape (17, 18). Thus, we estimated the free-energy landscape for three selected collective variables (CVs), chosen to represent the rearrangement of the polar network as distances between key sidechains. Together, these CVs describe the major conformational changes and exchange of interaction partners we observed in the MD simulation in the absence of TAK-875 (Fig. 1 F): CV1 represents the separation of the two positively charged guanidine groups of R183(Cζ) and R258(Cζ) (from 3.5 Å in the X-ray crystal structure to 10.0 Å in the presence of TAK-875, as shown by representative structures from a 600-ns unbiased MD simulation of FFAR1 in complex with TAK-875 and AP8 docked into the interhelical site (PDB 4PHU). (B) Pocket volume of the interhelical (yellow) and extracellular (blue) site over MD simulation time as 5 ns running averages (solid line) and for every 0.2 ns (faint line). (C and D) Comparison between representative structures, obtained by 1 Å rmsd geometric clustering from a 600 ns unbiased MD simulation in the presence (snapshot at 171.40 ns) and absence (snapshot at 493.24 ns) of TAK-875. (E and F) Rearrangement of the polar network stabilized by TAK-875, as shown by representative structures from a 600-ns unbiased MD simulation of FFAR1 in the presence and absence of TAK-875. Key amino acid side chains involved in the polar network are shown as purple sticks. (G–I) Free-energy profiles of selected atomic distances (CV1-3) obtained by a well-tempered metadynamics approach, using the same MD setup as for the unbiased MD simulation in absence of TAK-875. The free-energy profile was recorded every 10 ns, as indicated by the color scale. The population of conformational states are indicated as percentages. The corresponding atomic distances are displayed as black dotted lines in F.
the representative MD snapshot), CV2 describes the rotation of Y240(OH) toward K62(NG) (from 8.67 Å to 4.5 Å), and CV3 indicates the formation of a salt bridge between R258(C6) and E65(C6) (from 9.9 Å to 3.9 Å). During the metadynamics simulation, a time-dependent bias potential was introduced to discourage previously visited conformations. Thermodynamic reweighting provides access to the unbiased free energy as a function of the simulation time to examine whether the observed conformation represents energetic “trapping” resulting from insufficient sampling or corresponds to a true global free-energy minimum (17). In general, the global energy minima for the three atomic distances that were chosen as CVs were in good agreement with the atomic distances observed in the representative structures for the unbiased simulation absence of TAK-875 (CV1: 8.1 vs. 10.0 Å; CV2: 3.9 vs. 4.5 Å; CV3: 4.6 vs. 3.9 Å; Fig. 1 G–J). For CV2, a second, less occupied local minimum is present at 9.5 Å, probably indicating a second stable rotamer of Y240. However, when plotting the 2D free-energy plots for each of the combinations of the three collective variables (SI Appendix, Fig. S3), it is apparent that on rotation of Y240 toward K62, shown by a shift to a lower value of CV2 at 3.9 Å, the free-energy landscape for CV1 and 3 converges into single minima at 8.1 Å and 4.6 Å, respectively. This suggests that the rotation of Y240 stabilizes the formation of the salt bridge between R258 and E65 and keeps R183 and R258 separated. Overall, these results point to a distribution of conformational states in which the rearranged pocket volume to rise from 9.9 Å to 3.9 Å). During the metadynamics simulation (Fig. 1F) leading to the closure of the extracellular site is in fact the dominant conformation in absence of TAK-875.

The conformational dynamics and apparent connectivity between the different FFAR1 binding sites prompted us to investigate whether the closure of the extracellular site is reversible. The addition of TAK-875 after 600 ns of unbiased MD simulation in fact causes a reopening of this site, causing the pocket volume to rise from <10 Å² to ∼100–250 Å² (SI Appendix, Fig. S1). Likewise, the associated relocation of TM-I, TM-II, and TM-VII to their initial position (SI Appendix, Fig. S1B) leads to a Co-rmsd decrease from 2.8 Å (representative structure from the MD simulation in absence of TAK-875) to 1.8 Å for the MD simulation in which TAK-875 was added after 600 ns, compared with the X-ray crystal structure of FFAR1 (PDB 4PHU). As a consequence, the observed structural connectivity appears to be based on a concerted dynamic allosteric mechanism in which the presence of an agonist in the interhelical site is associated with formation of a relatively large well-defined pocket; that is, the extracellular site on the opposite side of the polar network (Fig. 1E). The observed structural dynamics of this part of the receptor leads to the hypothesis that prevention of the closure of the extracellular site through binding of a ligand specifically designed for this site would stabilize the active conformation by preventing the receptor from moving into the presumably inactive conformation, and consequently would function as an agonist.

**Structure-Based Design of an Ago-Allosteric Modulator.** Accordingly, we probed the extracellular site by conducting a structure-based virtual screening of >12 million commercially available small synthetic compounds (Fig. 2). These compounds were each docked in two steps into an ensemble of four representative structures from the unbiased MD simulation in the presence of TAK-875, and filtered on the basis of general chemical criteria and docking scores (19). A library of the 99 best-scored compounds was acquired and tested for their ability to activate FFAR1-mediated signal transduction in transfected HEK293 cells. Compound 1 (ZINC9634346/Z184779911), a derivative of 1,3,5-triazine-2-amine (Fig. 3C), was the most efficacious, with similar potency and efficacy as the endogenous ligand, oleic acid (Fig. 3A). Radiolabeled competition binding experiments showed that Compound 1, in contrast to unlabeled TAK-875, was unable to displace [3H]TAK-875 from the receptor, indicating that this agonist can bind simultaneously with an agonist binding to the interhelical site, in agreement with its expected binding in the extracellular site (Fig. 3B). We then tested the ability of Compound 1 to modulate activation of FFAR1 by oleic acid and the two different types of synthetic agonists, TAK-875 and AM-5262 (Fig. 3D–F). Compound 1 acted not only as an agonist but also as an allosteric modulator by increasing the potency of oleic acid ∼52-fold and increasing its efficacy (E_max) with 35% at 100 μM of Compound 1 (Fig. 3D). However, although increasing the potency of TAK-875 and AM-5262 by fivefold and twofold, respectively, Compound 1 surprisingly had a negative effect on the efficacy of these two agonists; that is, reducing their E_max by 22% and 35%, respectively (Fig. 3 E and F). Thus, Compound 1 acts as a positive allosteric modulator on the endogenous lipid agonist, with respect to both potency and efficacy, but increases the potency of the two types of synthetic agonist types only slightly, and even decreases their maximal efficacy. In this context, it is probably important to note that the maximal efficacy of Compound 1 itself is less than that of TAK-875 and AM-5262 (Fig. 3A). Therefore, it seems possible that Compound 1, by stabilizing the extracellular site, is shifting the dynamic equilibrium to an “active-like” conformation that can be described as an intermediate between the conformations induced by oleic acid, TAK-875, and AM-5262.

During follow-up experiments, we acquired 96 commercially available analogs of Compound 1, which were subsequently experimentally screened in an inositol phosphate (IP) accumulation assay. We initially identified two compounds, the γ-lactam ZINC7064000/Z107326816 and the ester-linked 3-methyl-7-fluorobenzofuran ZINC45346728/Z154646582, with similar pharmacological activity as Compound 1 (SI Appendix, Fig. S7). In general, the available analogs were chemically rather diverse, especially around the coumarin...
motif and the attached ether linker, which was only present in three of 96 compounds. We performed a more systematic exploration of the structure–activity relationship, based on custom-made synthesis (SI Appendix, Fig. S8). We found several active compounds with lowered potency and efficacy, among which the meta- substituted Compounds 4 and 5 showed the highest EC_{50} values of ~1 μM. It was also possible to replace the coumarin motif in Compound 1 with an ester-linked 3-methyl-7-fluorobenzofuran (Compound 6) moiety to achieve a modest potency gain at the cost of some loss of efficacy.

To rule out colloid aggregation-based activation, we tested the ability of Compound 1 to inhibit AmpC β-lactamase (SI Appendix, Fig. S9). At 31.6 μM concentration, the compound showed 0.4% enzyme inhibition, comparable to TAK-875 and AM-5262. Further, Compound 1 did not show any unspecific activity on mock-transfected cells (SI Appendix, Fig. S10).

To verify the binding sites and mode of action of the three different types of synthetic agonists, we introduced a series of, respectively, four, two, and seven mutations in the suspected extrahelical, intrahelical, and extracellular binding sites and tested the ability of the agonists to activate FFAR1, as listed in SI Appendix, Table S1 and shown for key selected mutations in Fig. 4. A very clear result was obtained by steric hindrance mutations A83F, in the hydrophobic tunnel of the interhelical site, which, as expected, impaired the potency of TAK-875 considerably (i.e., 1,740-fold; Fig. 4A), without affecting the function of AM-5262 or Compound 1 (Fig. 4C and D). Conversely, mutations in the lipid-exposed extrahelical site impaired the potency of AM-5262 as anticipated, without affecting TAK-875, as shown for Y44F (Fig. 4A and D). However, two of the mutations in the extrahelical site (Y114F and S123A) unexpectedly decreased the potency and efficacy of Compound 1 (Fig. 4C and SI Appendix, Table S1), although this less lipophilic compound is expected to act through binding to the extracellular site distant from the mutations in the lipid-exposed extrahelical site. Although the possibility of Compound 1 binding to the extrahelical site cannot be excluded, the other two mutations in this site (Y44F and G95F) actually improved potency of Compound 1 while dramatically decreasing potency of AM-5262. Specifically, the bulky phenylalanine, which is introduced for G95, is located in a central position in the extrahelical site and would be expected to occupy a large section of this pocket. This steric hindrance mutation has a major effect on the pharmacologic activity of AM-5262 (67-fold potency decrease), as well as the positive allosteric modulator AP8 (10), but remarkably, it has an unexpected positive effect on the activity of Compound 1, roughly doubling both its potency and efficacy. This observation might be related to an effect on a potential allosteric network that controls the dynamic structural connectivity between these two sites. A similar finding has been described in a mutational mapping study involving the M1 and M4 muscarinic receptors, in which mutation of a large number of residues involved in an allosteric network, but not in direct ligand contact, led to reduction of the activity of the positive allosteric modulator LY2033298 (21, 22).

This notion of an allosteric connectivity between the three FFAR1 binding sites was further supported by the mutational analysis of the extracellular site (Fig. 4 and SI Appendix, Table S1), in which all seven mutations had a profound effect on the activity of Compound 1, ranging from 34.6% reduction of EC_{max} (K259A) to a complete loss of activity (K259W, A66W, and Y12W). However, we also observed that several of these mutations impaired the potency of, in particular, TAK-875 (400–500-fold decrease), and in one case (K259W) also decreased the potency of AM-5262 (20-fold). A single mutation (K259A) could be identified that selectively affected the efficacy of Compound 1 without affecting the two other compounds (blue curves in Fig. 4A, C, and D). Because of their size and lipophilicity, it is not conceivable that TAK-875 or AM-5262 bind to the solvent-exposed extracellular site, also with respect to the crystallographically confirmed binding mode of TAK-875 in the tunnel-like interhelical site. It is more likely that the close proximity between these two sites (L262 is, e.g., a close neighbor to R258, a residue in direct contact with TAK-875) is the underlying cause for the observed effects, and that inserting a mutation into the extracellular site affects the dynamic coordination of the neighboring polar network required for anchoring the carboxylic acid moiety of the TAK-875 ligand.

![Fig. 3. Pharmacologic profile of Compound 1.](Image 301x643 to 384x705)

(A) Dose-response curves of Compound 1 in comparison with known synthetic and endogenous FFAR1 agonists. (B) Radiolabeled ligand binding of [H]TAK-875 in combination with Compound 1. (C) Chemical structure of Compound 1 (D–F) Dose-response curves of Compound 1 alone and in combination (10 and 100 μM fixed concentration) with oleic acid, TAK-875, and AM-5262. Values for log EC_{50} efficacy, and potency fold-changes are indicated for each compound in the table (Right).
Discussion

The main discovery of the present study is that an extracellularly facing, solvent-exposed empty pocket, present in all three agonist-bound X-ray crystal structures of FFAR1, closes during MD simulations on removal of the prototype agonist TAK-875 from its interhelical site, and that a ligand designed to occupy the extracellular pocket functions as an agonist and allosteric modulator of the receptor. Compound 1 has been designed to bind to the extracellular site, and by presumably preventing the closure of this pocket, it might stabilize an active, signaling conformation of the receptor. In this case, Compound 1 was identified on the basis of its favorable docking score (−28.04) in a representative receptor conformation originating from the largest cluster from the 600-ns MD trajectory in the presence of TAK-875. Docking of Compound 1 into the unrefined X-ray crystal structure (PDB 4PHU), in contrast, resulted only in a score of −18.37, well above the chosen cutoff (−25.00) that was applied in the virtual screening protocol. The MD simulations were thus necessary for the identification of Compound 1, which might underline the importance of accounting for receptor dynamics in structure-based drug discovery efforts and the potential of using MD simulations to study the dynamics of unexploited pockets, which deliberately can be targeted.

Although rational drug design efforts for GPCRs have traditionally focused on pockets between the extracellular segment of the transmembrane helices and loops (23), a number of recent X-ray crystal structures has revealed that ligands in fact can act as agonists or antagonists through binding to multiple alternative sites in the receptors. This includes allosteric sites located more superficially between the extracellular loops (24); extrahelical sites in the transmembrane region (25), similar to the binding site of AP8 in FFAR1 (10); or even sites at the intracellular face of the receptor, involving interactions with the intracellular poles of the TM-III and helix 8 (26, 27). This is in accordance with hypotheses dating more than 2 decades ago, in which it was proposed that GPCRs would have multiple different agonist binding sites, based on the notion that an agonist did not have to interact with a specific active site, but only needed to stabilize an active conformation of the dynamic receptor structure (27). FFAR1 is a prime example of this, as it appears to possess at least three very distinct binding sites for agonists: the interhelical site for the hydrophobic para-substituted O-phenyl-alkyl acids, such as TAK-875 and MK-8666; an extrahelical lipid-exposed site for meta-substituted O-phenyl-alkyl acids, such as AM-5262 and AP8; and a solvent-exposed extracellular site with more polar properties (Fig. 4). It is, however, still unclear to which sites the endogenous long-chain fatty acid (LCFA) agonists, including the newly identified 20-HETE (28, 29), bind, as they readily can be docked into both the interhelical and the extrahelical site (SI Appendix, Fig. S4) (7). From a signal transduction point of view, endogenous LCFA function as Gq-only agonists (i.e., similar to TAK-875 and MK-8666), which would indicate that the interhelical binding site is the orthosteric site. Further, it is possible, and even likely, that two LCFA molecules may in fact bind to FFAR1 at the same time, and access both the inter- and extrahelical sites via the lipid bilayer. This might also be the case for some of the synthetic agonists, such as AM-5262, as they dock very well into both sites (7).

In contrast, the extracellular solvent-exposed site is more solvent-exposed and of a rather polar character, as well as a different shape, in relation to both of the two other agonist binding sites, and it is unlikely that ligands binding to either the extrahelical or the interhelical site are able to act on the extrahelical site as well. Compared with other GPCRs with crystallographically resolved allosteric binding sites, such as the muscarinic M2 receptor with its modulator binding site in the extrahelical vestibule (24), FFAR1’s extracellular site is located in a partially overlapping location, but slightly smaller in size (30) and of more polar character. The M2 allosteric binding site is largely preformed in the presence of an agonist, similar to the stabilizing effect that we observed for the extracellular allosteric pocket in FFAR1 during the MD simulations in the presence of TAK-875.

The MD simulations of FFAR1 suggested that the presence of the agonist TAK-875 in the interhelical site between the extracellular segments of TM-III and TM-IV induced a helical conformation of ICL2, connecting the intracellular poles of these two helices. Secondary structure analysis of ICL2, which initially started in a random coil conformation, revealed that it was more likely to adopt an α-helical conformation in the presence (32%) of all ICL2 residues over all frames than in the absence (6%) of TAK-875 (SI Appendix, Fig. S1 A and B). This long-range allosteric effect appeared to be related to the relatively tight binding of the octadecanoyl tail of a 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine molecule from the artificial membrane in the extrahelical groove between the intracellular segments of TM-III and TM-IV over the entire simulation time in a similar binding mode as the crystallographic pose of AP8 in FFAR1 (PDB 5TZY). A helical conformation of ICL2 has been observed in many X-ray crystal structures of agonist-bound GPCRs, and appears to be associated with active-state receptor conformations and interactions with the G proteins (16). Therefore, the extrahelical site of FFAR1 might play a role in stabilizing the important helical formation of ICL2. It is likely that this extrahelical site could function as an allosteric binding site in many other GPCRs, as a similar binding groove is found in many structures of GPCRs and is often occupied with a lipid or detergent molecule (11, 31, 32).

Compound 1 represents an exciting starting point for a different type of FFAR1 ago-allosteric modulators, with respect to both chemotype and allosteric mechanism, which could translate into alternative therapeutic strategies for the treatment of type 2 diabetes (29, 33). The modulator properties of Compound 1...
might further enable a more fine-tuned effect on physiological signaling while maintaining the spatial and temporal attributes of endogenous free fatty acid ligands (34).

Materials and Methods

MD Simulations. All MD simulations were performed in GROMACS 4.6.5 (35), using the CHARMM36 force field (36) and the TIP3P water model. The X-ray crystal structure of FFAR1 (PDB 4PHU) was embedded into a 50-ns pre-equilibrated palmitoyl-2-oleoylsn-glycero-3-phosphatidylcholine membrane, using g_membed (37). After solvation in a cuboid box, Na+ and Cl− ions were added to neutralize the overall electric charge and to model an ion concentration of 0.15 M. A steepest-descent energy minimization was conducted and the system containing TAK-875 was equilibrated in the NPT ensemble for 40 ns, using decreasing position restraints. Representative structures for the extracellular and interhelical sites were selected by performing a geometric clustering based on the 48 residues that form the extracellular site and the interhelical binding site for TAK-875 and MK-8666, using a Co-rmsd cutoff value of 1 Å to determine cluster membership (SI Appendix, Fig. S3). Well-tempered metadynamics simulations were performed in GROMACS 5.1.2 PLUMED 2.2 (18) using the MD setup described above. We obtained the free-energy profiles of three selected atomic distances: CV1 = E65(Co)-R258(Ci), CV2 = Y240(OH)-K62(Ni), and CV3 = E65(Co)-R258(Cj) (atom naming according to PDB standard). Additional details are included in the SI Appendix.

Molecular Docking and Virtual Ligand Screening. We docked the ∼10,000 member set of the ZINC database (38), using SurflexDock (version 2.5) (39), into FFAR1’s extracellular site. The 100,000 best-scored complexes (prefiltered for binding scores >2, polar score >2) were selected for a second docking protocol, using ICM 4D docking and an ensemble of four representative receptor structures (version 3.8-4a, Molsoft L.L.C.). A set of chemical filtering criteria was applied (MW ≤ 500, logP ≤ 5, nrotb ≤ 12, VLS score <−25, deux like score <0.2, internal ligand conformation energy ≤10) to yield 315 compounds from which a small library of 99 compounds was cherry-picked. Additional details are included in the SI Appendix.

Experimental Functional and Binding Studies. FFAR1 was cloned into the expression vector pCMV-Tag 2B encoding a N-terminal FLAG epitope (Stratagene), using the QuickChange method. COS7 cells were transfected using calcium phosphate precipitation method. IP accumulation assays were conducted as described by Hauge et al. (7). Competition binding assays were performed by adding increasing doses of TAK-875 and Compound 1 to the binding buffer (Hepes wash buffer + 100 μg/mL bacitracin), and immediately after, 50 μL tracer solution containing 1h-TAK-875 (5,000 cpm/well) was added. Plates were incubated for 3 h and washed twice, and γ-radiation was counted on a Packard Top Count NXT counter. Additional details are included in the SI Appendix.

Acknowledgments. We thank Susanne Hummelgaard for performing the experimental compound screening. We would further like to acknowledge an anonymous referee for proposing additional MD simulations to further substantiate our conclusions about the allosteric mechanism between the different FFAR1 binding sites. The Novo Nordisk Foundation Center for Basic Metabolic Research (NNF10CC1016515) and the Novo Nordisk Foundation Center for Protein Research (NNF14CC0013655) are supported by an unconditional grant from the Novo Nordisk Foundation to the University of Copenhagen. M.L. received a faculty scholarship from the Faculty of Health and Medical Sciences, University of Copenhagen, supported by Challenge Grant NNF14OC0013655 from the Novo Nordisk Foundation (T.W.S.). K.L-L. and J.M. were supported by a Sapere Aude Starting grant from the Independent Research Fund Denmark (DFF).


