Small-Molecule Positive Allosteric Modulators of the β2-Adrenoceptor Isolated from DNA-Encoded Libraries

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ABSTRACT

Conventional drug discovery efforts at the β2-adrenoceptor (β2AR) have led to the development of ligands that bind almost exclusively to the receptor’s hormone-binding orthosteric site. However, targeting the largely unexplored and evolutionarily unique allosteric sites has potential for developing more specific drugs with fewer side effects than orthostERIC ligands. Using our recently developed approach for screening G protein–coupled receptors (GPCRs) with DNA-encoded small-molecule libraries, we have discovered and characterized the first β2AR small-molecule positive allosteric modulators (PAMs)—compound (Cmpd)-6 [(R)-N-(4-amino-1-(4-(tert-butyl)phenyl)-4-oxobutan-2-yl)-5-(N-isopropyl-N-methylsulfamoyl)-2-((4-methoxyphenyl)thio)benzamide] and its analogs. We used purified human β2ARs, occupied by a high-affinity agonist, for the affinity-based screening of over 500 million distinct library compounds, which yielded Cmpd-6. It exhibits a low micro-molar affinity for the agonist-occupied β2AR and displays positive cooperativity with orthostERIC agonists, thereby enhancing their binding to the receptor and ability to stabilize its active state. Cmpd-6 is cooperative with G protein and β2-arrestin1 (a.k.a. arrestin2) to stabilize high-affinity, agonist-bound active states of the β2AR and potentiates downstream cAMP production and receptor recruitment of β2-arrestin2 (a.k.a. arrestin3). Cmpd-6 is specific for the β2AR compared with the closely related β1AR. Structure–activity studies of select Cmpd-6 analogs defined the chemical groups that are critical for its biologic activity. We thus introduce the first small-molecule PAMs for the β2AR, which may serve as a lead molecule for the development of novel therapeutics. The approach described in this work establishes a broadly applicable proof-of-concept strategy for affinity-based discovery of small-molecule allosteric compounds targeting unique conformational states of GPCRs.

Introduction

The modulation of the β2-adrenoceptor (β2AR) activity plays an integral role in the treatment of a wide range of diseases. As such, GPCRs have become the target for over one third of current pharmaceuticals, the vast majority of which bind to the orthosteric site of the receptors. This region is defined as the site to which the endogenous ligand(s) for the receptor binds, such as adrenaline for the adrenoceptors or histamine for the histamine receptors (Lefkowitz, 2007; Whalen et al., 2011; Wacker et al., 2017). Most clinically used antagonists are orthosteric binders and exert their effects by competitive inhibition. Recently, however, an increasing number of negative and positive allosteric modulators [negative allosteric modulators (NAMs) and positive allosteric modulators (PAMs), respectively] for GPCRs has been described (Gentry et al., 2015), although to date only two have reached the clinic (Dorr et al., 2005; Lindberg et al., 2005). Rather than directly stimulating or inhibiting biologic effects...
on their own, these allosteric compounds exert their effects by modulating receptor responsiveness to endogenous agonists. Such allosteric ligands offer a number of potential advantages as drugs, including greater specificity among closely related receptor subtypes, and maximum or ceiling effects that can reduce adverse actions, among others (Wootten et al., 2013; Christopoulos, 2014). Such allosteric modulators can also serve as valuable reagents in the research laboratory, where, by means of their cooperative interactions with orthosteric ligands, they can help to freeze or lock specific receptor conformations so that they can be studied by biophysical techniques (Christopoulos, 2014; Wacker et al., 2017).

Selection of allosteric modulators for GPCRs using the usual cell-based functional assays such as those for cAMP generation or β-arrestin2 (a.k.a. arrestin3) recruitment (Rajagopal et al., 2010) has a number of disadvantages. These include that they can be quite laborious and difficult to interpret because one is looking for modulation of a response rather than the on or off responses that such assays are better suited to measure. Such assays are also subject to a variety of artifacts and have relatively limited compounds throughput of \( \sim 10^{5} \) to 10\(^{6} \). In contrast, interaction or affinity-based methods, in which large libraries of self-encoding potential binders are screened against a target protein molecule, circumvent these shortcomings. A particularly powerful approach is the use of DNA-encoded small-molecule libraries (DELs) potentially containing billions of compounds. Each molecule in such a library is covalently linked to a small stretch of nucleotides, which serves as a barcode that is used to identify target binders by next-generation sequencing (NGS) (Franzini and Randolph, 2016; Goodnow et al., 2017). Such approaches work well when applied to soluble protein targets but have been much more difficult to adapt to membrane proteins such as GPCRs. However, using this approach, we recently described isolation of the first NAM for the \( \beta_{2} \)-adrenoceptor (allosteric β-blocker) (Ahn et al., 2017) and identified its intracellular binding site on the receptor by X-ray crystallography (Liu et al., 2017). This molecule, compound (Cmpd)-15, was isolated by panning DELs against the inactive receptor, in which the orthosteric site was unoccupied.

An advantage of affinity-based screening methods is that one can bias the selections toward isolation of molecules with a particular functional profile by including one or another orthosteric ligand or even allosteric transducer protein molecules, e.g., G protein or β-arrestin, in complex with the receptor. In this work, we report our successful isolation of the first PAMs of the \( \beta_{2} \)-adrenoceptor (\( \beta_{2} \)-AR) by panning DELs against the purified receptor occupied by the high-affinity agonist BI-167107 (Rasmussen et al., 2011). We present a detailed pharmacological characterization of these molecules with the receptor and illustrate their potential utility as laboratory tools for interrogating biophysical properties of the receptors, as well as molecules for a new type of therapeutic agent.

**Materials and Methods**

**Materials.** Cmpd-6 and its analogs were synthesized using the methods described below. With the exception of BI-167107, which was synthesized as described previously (Wang et al., 2013), all of the orthosteric \( \beta_{2} \)-AR ligands used were purchased from Sigma-Aldrich (St. Louis, MO) and sourced at a 95% or greater purity. Nuevolution provided the DNA-encoded libraries used for screening. The \([^{1} \text{H}]R, R’\)-4-methoxyfenoterol used in the radioligand-binding studies was provided by Irving Wainer (Laboratory of Clinical Investigation, National Institute on Aging Intramural Research Program, Bethesda, MD). The \( \beta_{2} \)-AR-Gso and \( \beta_{2} \)-vasopressin 2 receptor (\( V_{2} \)-R)-β-arrestin1 fusion clones containing an N-terminal hemagglutinin signal sequence followed by a FLAG epitope tag for the receptor were generated in pcDNA3.1 by standard polymerase chain reaction (PCR) amplification and cloning procedure. β-arrestin1, a non-arrestin, was amplified by standard PCR methods and cloned in-frame with the C terminus of \( V_{2} \)-R essentially as before (Strauch et al., 2014). For the Gs fusion construct, the coding sequence for the short splice variant of human Gso subunit was used from a plasmid obtained from cDNA Resource Center (Bloomington, PA). Both Gso and \( \beta_{2} \)-AR sequences were PCR amplified separately, and the amplified fragments were assembled into a tetracycline-inducible pcDNA3.1 plasmid by using HiFi DNA assembly (NEB, Ipswich, MA) to finally generate the Gso fusion at the C terminus of the receptor. Both fusion constructs were sequence verified, and aliquots of maxi-prepared DNA were used for transfections. Previous purification methods were used to obtain rat \( \beta_{2} \)-arrestin1 (a nonvisual arrestin, a.k.a. arrestin2) and heterotrimeric Gso protein (Shuikla et al., 2013; Staus et al., 2016).

**Cell Culture and Transfection.** HEK-293 and HEK-293T cells were cultured at 37°C and at 5% CO\(_{2}\) in a humidified condition. Cells were cultured in standard minimum Eagle’s growth media supplemented with 10% fetal bovine serum and penicillin/streptomycin. HEK-293 cell lines stably expressing the GloSensor (Promega, Madison, WI) cAMP reporter (Nobles et al., 2011) and HEK-293T cell stably expressing the Tange assay (Barrettin 1, DNA was maintained, as described above. The HEK-293 cell line stably expressing the GloSensor cAMP reporter together with the \( \beta_{2} \)-AR was created by transfecting a hygromycin B–resistant plasmid expressing the GloSensor reporter into cells stably overexpressing the \( \beta_{2} \)-AR (Sheno et al., 2006), followed by selection with 150 \( \mu \)g/ml hygromycin B (Invitrogen, Carlsbad, CA). The clone with the greater fold over basal ratio and highest sensitivity was selected and maintained with 150 \( \mu \)g/ml G418 (Sigma-Aldrich) and 100 \( \mu \)g/ml hygromycin B. Transient transfections were performed using FuGENE 6 (Promega), according to the manufacturer’s instructions, and all assays were done 48 hours post-transfection. The \( \beta_{2} \)-AR-Gs and \( \beta_{2} \)-V\(_{2}\)-R-β-arrestin 1 fusion proteins were transfected into Exp293F cells (Invitrogen) using ExpiFectamine (Invitrogen), as described by the manufacturer.

**Expression, Purification, and High-Density Lipoprotein Reconstitution of the \( \beta_{2} \)-AR.** As previously described (Kobilka, 1995), BeatBac Baculovirus Expression System was used to express the full-length human \( \beta_{2} \)-AR containing an amino-terminal FLAG epitope tag, carboxyl-terminal His-tag, and a N187E glycosylation mutation in Sf9 insect cells. In brief, cells were infected at a density of \( 3 \times 10^{4} \) cells/ml and harvested 67 hours thereafter. The cells were then solubilized in a buffer containing 1% n-dodecyl-β-D-maltoside (Anactrace, Maumee, OH), 20 mM HEPES, pH 7.4, 150 mM NaCl, and protease inhibitors. Functional \( \beta_{2} \)-AR was purified, as previously described (Kobilka, 1995; Kahsai et al., 2016), using FLAG-M1 antibody and alprenolol affinity chromatography, followed by size exclusion chromatography using a Superdex 200 (16/600 prep grade) column. The monomeric receptor peak from the size exclusion chromatography was pooled and concentrated to 1–2 mg/ml. Purified functional \( \beta_{2} \)-AR was then reconstituted into high-density lipoprotein (HDL) particles using previously published methods (Whorton et al., 2007; Staus et al., 2016). In brief, FLAG-\( \beta_{2} \)-AR was incubated with a 50-fold molar excess of biotinylated membrane scaffold protein 1 Apo A1 and 8 mM POPC/POPG (3:2 molar ratio; 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine and 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-[1-rac-glycerol]) lipids (Avanti Polar Lipids, Alabaster, AL) for 1 hour at 4°C. Detergent was removed using BioBeads SM-2 (Bio-Rad, Hercules, CA) by incubating overnight at 4°C. Then, receptor-containing HDL particles were isolated using FLAG-M1 affinity chromatography and size exclusion chromatography.

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DNA-Encoded Small-Molecule Library. The DNA-encoded small-molecule libraries used for screening were created using a tagged-split-and-pool chemistry approach (Chemetics) at Nuevolution, as previously described (Kontijevskis, 2017).

Affinity Selection. Figure 1A schematically illustrates the library selection process. More specifically, 30 μg biotinylated β2AR-HDL particles were immobilized on 25 μl NeutrAvidin beads (Thermo Fisher Scientific, Waltham, MA). The β2ARs bound to NeutrAvidin beads were then incubated with library molecules in 50 μl binding buffer (20 mM HEPES, pH 7.4, 100 mM NaCl) supplemented with 20 μM BI-167107 (BI) and 1 mg/ml sheared salmon sperm DNA (ssDNA; Ambion, Waltham, MA) for 45 minutes at room temperature, while intensely shaken. Prior to this incubation, 1 μl library molecules were allocated for later quantitative PCR (qPCR). Following this incubation, the beads were transferred to a micro-column connected to a vacuum apparatus and subsequently washed three times with 100 μl ice-cold binding buffer containing 10 μM BI. During each of the washing steps, excess liquid was removed via vacuum suction. To elute off the bound compounds, the beads were incubated twice with 50 μl water containing 1.5% Fos-choline (Avanti Polar Lipids) at 37°C for 15 minutes and subsequently at 72.5°C for 15 minutes. Following this incubation, the solution was separated from the beads by centrifugation at 1000 × g for 1 minute. After addition of 1 μl 10 mg/ml ssDNA, the combined supernatant was applied to a nucleotide removal kit (Qiagen, Hilden, Germany) to remove denatured protein and lipid molecules. The mixture of the bound compounds was then eluted with 50 μl water from the nucleotide removal column, and 1 μl purified material was allocated for later qPCR quantification. The remaining purified sample was then either used for the next round of selection with fresh β2AR-HDLs, or applied to NGS.

Quantitative PCR. Library DNA was quantified using qPCR at the end of each round of affinity selection. Briefly, the DNA samples were either directly amplified, or amplified after being diluted in solution containing 0.1% Tween 20, 20 mg/ml ssDNA using Jump-Start Taq ReadyMix (Sigma-Aldrich), according to manufacturer's guidelines. The samples from the libraries before selection were evaluated using at least four different concentrations on a log scale. This provides a standard to determine the DNA copy number of the samples from each iterative round of selection. The reaction was done with the primer set, including the universal forward primer (5′-CAAGTCACCAAGAATTCATG-3′) and a unique reverse primer for each library, and FAM/TAMRA probe 5′-CAGACGACCTAGGATCACC-3′ using a StepOnePlus (Applied Biosystems, Waltham, MA).

NGS and Analysis. To increase the yield and then append the required sequencing adapters for emulsion PCR, the affinity-selected materials were amplified by two rounds of PCR. The first round of PCR was done with the same oligonucleotide primer set as used for qPCR. The second round of PCR was performed with oligonucleotide primers made from fusing the Ion Torrent adapter sequences to the universal forward primer, to which a sorting code was inserted to allow for sample pooling, and a unique reverse primer for each library. The unique sequence of the reverse primer provides precise sample tracking and a distinct identifier for each library. Following gel purification, the final PCR products were subjected to

![Fig. 1. Hit compounds from DEL screening with the agonist-occupied β2AR in HDL particles.](image)
single-direction ampiclon sequencing using the Ion Torrent platform (PrimBio, Exton, PA). Sequences having significant copy numbers (high signal-to-noise ratio), determined by analyzing the NGS output using a custom-built algorithm, were deconvoluted to their corresponding chemical structures from the database.

**Isothermal Titration Calorimetry.** Isothermal titration calorimetry (ITC) experiments were performed with maltose neopentyl glycol (MNG; Anatrace)–solubilized βAR on a MicroCal Auto-iTC200 system (Malvern, Malvern, UK), according to the previously reported method (Ahn et al., 2017). Dialysis of the purified βAR was carried out against a dialysis buffer (20 mM HEPES, pH 7.5, 100 mM NaCl, 0.01% MNG, and 0.001% cholesteryl hemisuccinate). Titrations were performed at 25°C, in which 40 μl 200 μM Cmpd-6 in the aforementioned dialysis buffer was loaded into the syringe, followed by an initial injection of 0.2 μl, and then subsequent 2 μl injections (0.4-second duration, 150-second spacing, and 5-second filter period) into the 200 μl sample cell containing βAR (at 30 μM final) prestimulated with isoproterenol (ISO; 2 mM final). During the experiment, the reference power was set to 7 μcal × s⁻¹ and sample cell was stirred continuously at a speed of 1000 rpm. ITC raw data were baseline corrected, peak area integrated, and fitted by using a one-site nonlinear least-squares fit model using the MicroCal Origin software program, to provide affinity constant (Kd), stoichiometry (N), and thermodynamic parameters such as enthalpy (ΔH) and entropy (ΔS).

**Radioligand Binding.** Binding experiments were done as previously detailed (Ahn et al., 2017). In brief, competition radioligand binding assays were done using the radiolabeled agonist [³H]cytanopindol (CYP; 2200 Ci/mmol; PerkinElmer, Waltham, MA) at a concentration of 60 pM. The βAR-HDL particles were used at ~0.7 ng per reaction. Reactions consisted of the βAR-HDL particles, [³H]-CYP, Cmpd-6 at varying concentrations, and a serial dilution of a competitor βAR agonist, most often ISO. All of the components were diluted in an assay buffer (20 mM HEPES, pH 7.4, 100 mM NaCl, 0.1% bovine serum albumin (BSA), and 1 mM ascorbic acid). Each reaction was allowed to reach equilibrium by incubating for 90 minutes at room temperature. Assays were then terminated by rapid filtration onto GF/B glass-fiber filters (Brandel, Gaithersburg, MD) treated with 0.3% polyethyleneimine and washed with 8 ml cold binding buffer (20 mM HEPES, pH 7.4, 100 mM NaCl) using a harvester (Brandel). [³H]-CYP bound to the βAR-HDL particles was measured using either a Packard Cobra Quantum gamma counter (Packard; GMI, Ramsey, MN) or a WIZARD2-2 Detector Gamma Counter (PerkinElmer). Data were expressed as specific binding.

For [³H]-methoxyfenoterol ([³H]-FEN) (Toll et al., 2012) binding, membrane preparations from Sf9 cells expressing either βAR or β₂AR were used. For in cell phosphorylation of β₂AR, GRK2-CAAX was coexpressed and, prior to harvest, cells were stimulated with the agonist ISO (10 μM) for 20 minutes. Membranes for βAR and phosphorylated β₂AR were essentially prepared, as described earlier (Strachan et al., 2014; Ahn et al., 2017). For [³H]-FEN binding with Gs or nanobody-80 (Nb80), βAR membranes were incubated in the G protein assay buffer (50 mM Tris-HCl, pH 7.4, 2 mM EDTA, 12.5 mM MgCl₂). For binding assays containing rat β-arrestin1, phosphorylated β₂AR membranes were incubated in the β-arrestin assay buffer (50 mM Tris-HCl, pH 7.4, 50 mM potassium acetate, 5 mM MgCl₂). Both assay buffers were supplemented with 0.05% BSA and 0.018% L-ascorbic acid. [³H]-FEN (12.6 Ci/mmol) was used at its Kᵢ (4.3 nM) in binding assays testing for PAM activity of Cmpd-6 (and its analogs) and Cmpd-43. In [³H]-FEN saturation-binding assays, testing for the cooperativity of Cmpd-6 (20 μM) with Gs (100 nM), β₂AR (1 μM), or Nb80 (1 μM), [³H]-FEN was used in the range of 0.39–50 nM to saturate the high-affinity agonist binding sites in the receptor. All binding reactions were incubated to equilibrium (90 minutes) at room temperature and then harvested onto polyethyleneimine-soaked GF/B filters, followed by four rapid washes of 2 ml with prechilled G protein wash buffer (50 mM Tris-HCl, pH 7.4, 2 mM EDTA, 12.5 mM MgCl₂) or β-arrestin wash buffer (50 mM Tris-HCl, pH 7.4, 50 mM potassium acetate). Bound [³H] was extracted overnight with 5 ml scintillation fluid and quantified using a Packard Cobra Quantum gamma counter (Packard; GMI). Nonspecific radioligand binding was assessed in reactions that contained the antagonist propranolol (20 μM).

**Measurements of cAMP Production.** cAMP production, an indirect marker of Gs protein activation, was measured using the GloSensor (Promega), a chemiluminescence-based cAMP biosensor, as previously described (Ahn et al., 2017). In brief, HEK-293 cells stably expressing the GloSensor luciferase enzyme alone, or together with the βARs, were plated in 96-well, white clear-bottom plates at a density of ~80,000 cells/well. Cells were given at least a 24-hour incubation to recover cell surface receptor expression before the assay was started. Cells were then treated with the GloSensor reagent (Promega) and incubated at 27°C and ~100% relative humidity for ~1 hour. Cells were then treated with either a varying dose of Cmpd-6 or a vehicle control (dimethylsulfoxide (DMSO)) diluted in Hank’s balanced solution (Sigma-Aldrich), supplemented with 20 mM HEPES, pH 7.4, 0.05% BSA, and 3-isobuty-1-methylxanthine (Sigma-Aldrich) at a final concentration of 100 μM. For most of the cAMP assays, cells were then incubated further for 20 minutes, before a serial dilution of the β-agonist was added. For the assays with HEK-293 cells stably overexpressing βAR, Cmpd-6 and the β-agonist serial dilution were added to the cells simultaneously. Upon stimulation of the cells with the β-agonist, changes in luminescence were read using a NOVOstar microplate reader (BMG Labtech, Cary, NC) at various time points ranging from 5 to 35 minutes.

**Measurement of β-Arrestin Recruitment.** β-arrestin2 recruitment to the test receptor is measured using the previously described Tango assay (Barnea et al., 2008). HEK-293T cells stably expressing the β₁V₁R tethered to the tetracycline transactivator transcription factor by a tobacco etch virus protease cleavage site, the human β-arrestin2 protein fused to the tobacco etch virus protease, and the tetracycline transactivator–driven luciferase reporter were used for this assay. Cells were plated on a 96-well, white clear-bottom plate at a density of ∼50,000 cells/well and were given at least a 24-hour incubation at 37°C, 5% CO₂, and ~100% relative humidity to recover surface receptor expression. Cells were treated with either a varying dose of Cmpd-6 or a vehicle control (DMSO) diluted in Hank’s balanced solution (Sigma-Aldrich), supplemented with 20 mM HEPES, pH 7.4, and 0.05% BSA, and then incubated at 37°C, 5% CO₂, and ~100% relative humidity for ~20 minutes. After the incubation, a serial dilution of the β-agonist was added, following which the cells were incubated for 6 hours at 37°C and ~100% relative humidity. At the end of the incubation, the plate was cooled to room temperature. After adding the Bright-Glo reagent (Promega), chemiluminescence signals were read using a NOVOstar microplate reader (BMG Labtech) at 5–10 minutes.

**Bimane Assay.** The minimal cysteine βAR (Yao et al., 2009) was used in bimane fluorescence experiments. For bimane labeling at cysteine-265 of the βAR, threefold molar excess of monobromobimane (Sigma-Aldrich) was used, as previously described (Yao et al., 2009), and HDL reconstitution of the βAR-bimane was carried out, as described above. Bimane-labeled βAR-HDLs at 250 nM were incubated in black, solid-bottom 96-well microplates with the vehicle (DMSO) or 10 μM ISO, either alone or together with 1 μM Nb80 or 20 μM Cmpd-6 for 30 minutes at room temperature. All of the components were diluted in buffer comprised of 20 mM HEPES, pH 7.4, and 100 mM NaCl. A CLARIOstar plate reader (BMG Labtech) was used to collect fluorescence emission spectra using the top-read mode with excitation at 370 nm (16-nm bandpass) and emission scanning from 400 to 600 nm (10 nm bandpass) in 1-nm increments.

**Nanoβ-Enzyme-Linked Immunosorbent Assay.** As previously described, the 6X-His tagged nanobodies, Nb80 (Rasmussen et al., 2011) and Nb6B9 (Ring et al., 2013), were purified from periplasmic extracts of Escherichia coli WK6 cells. Nanobodies were affinity purified using Ni-NTA agarose beads (Qiagen). Purified nanobodies were then dialyzed overnight in 20 mM HEPES, pH 7.4, 100 mM NaCl, followed by size-exclusion chromatography. Nb80 (10 μg/ml) was passively adsorbed onto Maxisorp (NUNC, Roskilde, Denmark).
Denmark) 96-well plates (Thermo Fisher Scientific, St. Louis, MO) in nanobody buffer (20 mM HEPES, pH 7.4, 100 mM NaCl), and plates were incubated overnight at 4°C. Enzyme-linked immunosorbent assay was performed essentially as described before (Stau et al., 2014). Purified β2AR was preincubated with 0.2% DMSO or 10 μM final of either ICI-118551 or BI-167107 ligands, and Cmpd-6 (20 μM) or N6689 (1 μM) for 30 minutes in assay buffer (20 mM HEPES, pH 7.4, 100 mM NaCl) containing 0.01% MNG (Anatrace), 0.001% cholesteryl hemisuccinate (Sigma-Aldrich), and 0.5% BSA. The pre-incubated reactions were then overlaid on Nb80-adSORbed 96-well plates for 90 minutes at room temperature. Following incubation, the unbound material was washed with assay buffer, and the captured β2AR was detected using a horseradish peroxidase–conjugated anti-Flag (M2) antibody (1:5000) diluted in assay buffer. Following antibody incubation (1 hour, room temperature), plates were washed with assay buffer, and signal was developed using 100 μl Ultra-TMB (Pierce, Rockford, IL). The developed signal was quenched with 100 μL acidified assay buffer, and the absorbance was measured at 450 nm.

**Synthesis and Characterization of Cmpd-6 and Its Derivatives.** Complete details of chemical syntheses are described in Supplemental Material.

**Results**

**Screening and Identification of Primary Hits Including Cmpd-6 and -43.** Using our recently developed approach for screening DELs against GPCRs (Ahn et al., 2017), in this study we screened ~500 million unique DNA-encoded small molecules (Supplemental Table 1) to obtain PAMs at the β2AR. To increase the chance of obtaining PAMs, the orthosteric site of the receptor was occupied by a high-affinity β-agonist BI-167107, which shifted the β2AR population toward active conformations (Rasmussen et al., 2011; Manglik et al., 2015) (Fig. 1A). Furthermore, purified human β2ARs were reconstituted in detergent-free HDL particles (Fig. 1A). The HDL reconstitutions were performed using a biotinylated version of the membrane scaffolding protein ApoA1 (Whorton et al., 2007). In addition to providing the receptor with a native-like membrane environment, the biotinylated HDL particles provide an excellent immobilization scheme that avoids any physical perturbations to the receptor during the screening process. The β2ARs in biotinylated HDL particles can be efficiently captured on NeutrAvidin beads (Supplemental Fig. 1A) and have a comparable affinity for antagonist binding to that of β2ARs in membrane preparations (Supplemental Fig. 1B). Furthermore, by competitive radioligand-binding assays, we show that β2ARs in HDL particles can functionally couple to heterotrimeric Gs (Supplemental Fig. 1C). G protein coupling to the β2AR substantially increases the affinity of the competing agonist—ISO. This high-affinity coupling of Gs to the β2AR can be completely blocked by the addition of GTPγS—a nonhydrolyzable GTP analog.

Using the BI-167107–occupied β2AR in HDL particles, we iteratively screened four different DELs (Kontijevskis, 2017), each of which comprised over 100 million unique compounds (Supplemental Table 1), to isolate molecules that specifically bound to the active state of the receptor. The total number of molecules in each library was ~0.5–1 × 10^14. Three rounds of iterative selection (Fig. 1A) were performed with each library until the total number of target-bound molecules was decreased to ~1 × 10^6, which was monitored by qPCR. Following amplification of preserved DNA barcodes by PCR, the samples were subjected to NGS to identify compounds that outlasted the entire selection procedure. Sequences having significant copy numbers (i.e., high signal-to-noise ratio) were deconvoluted to their corresponding chemical structures from the database. Through this analysis, we determined 50 compounds as primary candidates that possibly bind to the β2AR (Supplemental Table 1) and named them Cmpd-1 to Cmpd-50. These 50 candidate compounds were synthesized on a small scale without their DNA barcodes to evaluate their activity as PAMs in secondary screens.

PAMs are expected to potentiate the binding of orthosteric agonists to GPCRs and even plausibly the coupling of transducer proteins, e.g., G protein and β-arrestin, to receptors (Wootten et al., 2013; Christopoulos, 2014). Accordingly, these 50 potential hits were tested for their ability to increase the binding of the radiolabeled agonist, ^3^H-FEN (Toll et al., 2012), to the β2AR in membrane preparations, both in the absence and presence of transducers (Supplemental Fig. 1D). Through this secondary screen, we identified seven structurally related compounds, as shown in Fig. 1B, including Cmpd-6, which showed the strongest PAM activity among the compounds (Supplemental Fig. 1E). These compounds not only increased ^3^H-FEN binding to the β2AR alone, but also, to varied extents, potentiated the transducer-induced high-affinity ^3^H-FEN binding at the receptor. Interestingly, between these compounds, only subtle structural differences were observed, which were confined to one variable region designated as R1 (blue colored in Fig. 1B). Two of the seven compounds (Cmpd-6 and -43) were chosen for further characterization of their PAM activity and were synthesized on a large scale.

To assess direct molecular interaction between Cmpd-6 and the agonist-bound, active β2AR, ITC was employed. The values summarizing binding affinity (Kd), stoichiometry (N), and thermodynamic parameters are shown in Fig. 1C. ITC video indicates that the process of interaction between Cmpd-6 and active β2AR is exothermic, therefore enthalpically favored with a stoichiometry of ~1 and Kd of 5.2 ± 0.5 μM.

**PAM Activity of Cmpd-6 and -43 in β2AR-Mediated Downstream Signaling.** To evaluate the PAM activity of Cmpd-6 and -43 in β2AR-mediated downstream functions, we monitored their effects on both agonist-induced Gs protein–cAMP production (Binkowski et al., 2011; Rajagopal et al., 2011) and β-arrestin2 recruitment to the receptor (Rajagopal et al., 2011; Bassoni et al., 2012) using cellular assays. An issue encountered in these functional assays is the differential levels of the signal produced by virtue of the high amplification process downstream of Gs protein activation, compared with the stoichiometric recruitment of β-arrestin2 to the receptor (Rajagopal et al., 2011). To circumvent this problem and achieve similar levels of the signal in the two cellular assays, we used the endogenously expressed β2AR in cAMP production assays, whereas β-arrestin2 recruitment was measured using cells stably overexpressing the β2V2R. This chimeric receptor has the V2R tail recombinantly appended at the C terminus of the β2AR, retaining the pharmacological traits of the native β2AR, but displaying a more stable interaction with β-arrestin, which is an advantage for β-arrestin recruitment assays (Tohgo et al., 2003). Both Cmpd-6 and -43 increased the ability of the agonist ISO to activate G protein–mediated cAMP production through the β2AR in a dose-dependent way (Fig. 2, A and B). We observed that Cmpd-6 (Fig. 2A) and Cmpd-43 (Fig. 2B) increased the maximal response induced by ISO, as well as potentiating the EC50 value of ISO, which was apparent in its left-shifted dose–response curve. In this assay,
Cmpd-6 shows stronger activity than Cmpd-43, which is consistent with the preliminary data showing the extent of dose-dependent increases in ³H-FEN binding to the β₂AR induced by these compounds, shown in Supplemental Fig. 1E. We also obtained a comparable pattern of agonist-induced β-arrestin2 recruitment to the β₂V₂R with Cmpd-6 (Fig. 2C) and Cmpd-43 (Fig. 2D), respectively.

Increases in the ISO-induced maximal response by Cmpd-6 and -43 in both assays suggest that ISO may act as a partial agonist, which does not reach the maximum response possible in these systems (Langmead, 2011), allowing Cmpd-6 and -43 to further increase the maximal agonist-induced response. To verify this, we monitored cAMP production by overexpressed β₂AR (a system that has much higher amplification; Supplemental Fig. 2) in the presence or absence of Cmpd-6. We observed that Cmpd-6 led to dose-dependent leftward shifts of the ISO dose-response EC₅₀ values, with increases in the basal activity, but did not increase the ISO-stimulated maximal response. This shows that even a full agonist such as ISO can act as a partial agonist depending on the assay system used, which would not have been suspected without the cooperation displayed by these new PAMs. Overall, these results strongly demonstrate that Cmpd-6 and -43 have PAM activity for β₂AR-mediated downstream functions, and that Cmpd-6 has stronger PAM activity than Cmpd-43.

**Cmpd-6 and -43 Potentiate the Binding Affinity of Agonists for the β₂AR.** A hallmark of PAMs is that they allosterically stabilize the agonist-bound active conformation of the receptor (Langmead, 2011), as do transducer proteins, G protein, and β-arrestin, as illustrated in the GPCR ternary complex model (De Lean et al., 1980). Because PAM-mediated stabilization of active GPCR conformation leads to potentiation of agonist-binding affinity for the receptor, we next tested whether Cmpd-6 and -43 increase the binding of an agonist to the β₂AR. For this, we monitored the competition binding of the orthosteric agonist ISO against the radiolabeled antagonist ¹²⁵I-cyanopindolol (CYP) to the β₂AR-HDL in the presence or absence of Cmpd-6 (Fig. 3A) and Cmpd-43 (Fig. 3B). As expected, both compounds potentiated the binding of ISO to the β₂AR in a dose-dependent way, as evidenced by the robust left shifts in ISO competition curves. Consistent with the results obtained in the cellular assays (Fig. 2), at their highest concentration tested (Fig. 3, A and B), Cmpd-6 potentiated the IC₅₀ value of ISO close to 50-fold, which was substantially more than the ~30-fold change elicited by Cmpd-43. Additionally, we obtained comparable shifts in the ISO dose–response curve induced by Cmpd-6 and -43 in radioligand competition binding done with membranes prepared from β₂AR-overexpressing cells (Supplemental Fig. 3, A and B).

Additionally, results shown in Fig. 3C further confirm the PAM activity of Cmpd-6 and -43 for increasing the binding of an orthosteric agonist to the β₂AR. Cmpd-6 and -43 dose dependently increased the binding of the radiolabeled agonist ³H-FEN to the β₂AR expressed in cell membranes, consistent with what we observed in our preliminary experiments with these compounds (Supplemental Fig. 1E). Again, Cmpd-6 is more efficacious than Cmpd-43 in increasing ³H-FEN binding to the β₂AR. Furthermore, the low micro-molar affinity (EC₅₀) value of Cmpd-6 obtained in this assay (Fig. 3C) is comparable to its Kᵣ value measured for its direct interaction with the β₂AR by ITC analyses (Fig. 1C). We also observed another feature of allosteric molecules, the ceiling effect, with these compounds in both binding experiments (Fig. 3). The increases in the binding of both agonists, ISO (Fig. 3, A and B) and FEN (Fig. 3C), were saturated with increasing concentrations of these allosteric compounds.

**Cmpd-6 Stabilizes the Agonist-Induced Active Conformation of the β₂AR.** We further demonstrated that Cmpd-6 stabilizes active conformational ensembles of the β₂AR.
β2AR through a biophysical assay. Agonist-induced activation of the β2AR causes the outward movement of transmembrane helix 6 (TM6), which can be detected by labeling of cysteine-265 at the intracellular base of TM6 with monobromobimane, an environmentally sensitive fluorescent label. Following receptor activation, the outward movement of TM6 leads to a decrease in fluorescence intensity with a concomitant increase in the maximum wavelength for emission (Rasmussen et al., 2011). Cmpd-6 alone induced decreases in overall fluorescence intensity, but not increases in the maximum wavelength for emission from the bimane-labeled β2AR (Fig. 4). In contrast, ISO decreased fluorescence to a similar extent, but also increased the maximum wavelength. This suggests that the conformational ensemble of the β2AR when bound to Cmpd-6 alone is similar to, but distinct from, that induced by orthosteric agonists. Interestingly, Cmpd-6 further potentiated ISO-induced decreases in the fluorescence intensity and increases in the maximum wavelength from the bimane-labeled β2AR (Fig. 4). Importantly, Cmpd-6-mediated potentiation of ISO effects was similar in magnitude to that observed with an allosteric nanobody (Nb80) that mimics the G protein–mediated potentiation of ISO affinity compared with the uncoupled receptor (Fig. 5, B and D). Additionally, we assessed the PAM activity of Cmpd-6 by measuring binding of the radiolabeled orthosteric agonist 3H-FEN aimed at saturating high-affinity sites on the β2AR (Fig. 5, E and F). Compared with no transducer controls, addition of Cmpd-6 or the exogenous transducers, heterotrimeric Gs (at β2AR membranes; Fig. 5E) and β-arrestin1 (at phosphorylated β2V3R membranes; Fig. 5F), robustly increased the high-affinity 3H-FEN binding to the receptor. Interestingly, addition of Cmpd-6 together with Gs or β-arrestin1 further enhanced the maximal high-affinity 3H-FEN binding. Although there was noticeable cooperativity between Cmpd-6 and Gs, this potentiation in 3H-FEN binding was also prominent in the presence of the G protein mimic Nb80 (Rasmussen et al., 2011) (Supplemental Fig. 4A). Together with our findings from cellular assays, these binding studies clearly demonstrate a positive cooperativity between Cmpd-6 and transducers to modulate high-affinity state agonist binding to the β2AR.

Of note, the data in Supplemental Fig. 4A also suggest that Cmpd-6 does not occlude transducer coupling to the β2AR and most likely binds to a potentially unique allosteric site in the receptor. Accordingly, to test whether Cmpd-6 physically competes for binding to the intracellular transducer-binding pocket, we performed an enzyme-linked immunosorbent assay to capture the β2AR with the G protein mimic Nb80 that recognizes agonist-bound active state of the receptor (Supplemental Fig. 4B). In the presence of the high-affinity agonist BI-167107, and compared with DMSO or the antagonist ICI-118551, there was a marked increase in receptor capture by Nb80. This receptor capture was significantly inhibited in the presence of saturating amounts of a competing nanobody Nb6B9 (Ring et al., 2013), which is an affinity-matured version of Nb80 and thus competes for a common binding epitope on the β2AR. Interestingly, and in contrast to Nb6B9, under these experimental conditions the addition of a saturating concentration of Cmpd-6 did not alter the capture of β2AR by Nb80. These data suggest that presence of Cmpd-6 does not interfere with transducer coupling to the β2AR, which further establishes positive cooperativity between transducers and the compound.
The PAM Activity of Cmpd-6 Is Specific for the β2AR. The specificity of Cmpd-6 for the β2AR was first evaluated through in vitro competition radioligand (125I-CYP) binding at the β1AR, the most closely related subtype of the adrenoceptor. In this assay, Cmpd-6 induces a minimal left shift of the ISO competition curve for binding to the β1AR (Supplemental Fig. 5A) unlike the robust ISO curve shift by Cmpd-6 observed with the β2AR (Fig. 3A). This shows that Cmpd-6 specifically induces the high-affinity binding of the orthosteric agonist ISO to the β2AR, but not to the β1AR. Furthermore, we observed only marginal changes promoted by Cmpd-6 in the ISO dose–response pattern of β1AR-mediated cAMP production (Supplemental Fig. 5B), which is markedly different from that of the β2AR-mediated response (Supplemental Fig. 2). We also detected minimal allosteric effects of Cmpd-6 on cAMP production mediated by other receptors. These are the transiently overexpressed V2R (Supplemental Fig. 5C), as well as the prostaglandin E2 (Supplemental Fig. 5D) and the vasoactive intestinal peptide (Supplemental Fig. 5E) receptors, both endogenously expressed in the assay cells. These findings clearly demonstrate that the PAM activity of Cmpd-6 is specific for the β2AR relative to the β1AR and other receptors tested in this work.

PAM Activity of Cmpd-6 When the β2AR Is Stimulated with a Range of Different Agonists. Some allosteric modulators show differential activity depending on the orthosteric agonist stimulating the receptor, a phenomenon known as probe dependence (Wootten et al., 2013;
We examined whether Cmpd-6 displays such differential activity when the orthosteric site of the $\beta_2$AR is occupied with a range of agonists, namely epinephrine (EPI) and fenoterol (FEN), which are very strong partial agonists compared with ISO, and clenbuterol (CLEN), which is a weak partial agonist (Rajagopal et al., 2011). We first evaluated the extent of the dose–response curve (IC$_{50}$ value) shift induced by Cmpd-6 in radioligand ($^{35}$S-L-CYP) competition binding to the $\beta_2$AR (Supplemental Fig. 6, A–D). This allowed us to test the allosteric activity of Cmpd-6 solely for binding of an agonist in the absence of transducer coupling to the receptor. We observed that the extent of the curve shift in the presence of Cmpd-6 in this assay essentially followed the efficacy of the tested agonists to induce downstream signaling.

We next compared the PAM activity of Cmpd-6 for downstream signaling of the $\beta_2$AR when stimulated with these four agonists using cell-based functional assays, monitoring cAMP accumulation (Supplemental Fig. 6, E–H) and $\beta$-arrestin2 recruitment to the receptor (Supplemental Fig. 6, I–L). In general, full and strong partial agonists show greater affinity (EC$_{50}$ value) shifts by Cmpd-6 compared with that observed with the weak partial agonist CLEN. However, CLEN displayed a substantially greater Cmpd-6–mediated increase in maximal response than did the full agonists. Interestingly, no direct relationship between the extent of the EC$_{50}$ shift by Cmpd-6 and the efficacy of ISO, EPI, and FEN was seen. In functional assays, Cmpd-6 induced a noticeably greater shift with EPI (Supplemental Fig. 6, F and J) than with ISO (Supplemental Fig. 6, E and I) and FEN (Supplemental Fig. 6, G and K), whereas the fold increases by Cmpd-6 in the maximal response induced by these agonists were comparable. Thus, we did not observe any unique probe dependence of Cmpd-6 with this small panel of agonists.

**Structure–Activity Relationships of Cmpd-6 Analogs.**

To determine structure–activity relationships (SAR) around Cmpd-6, a rational design approach and synthesis of a series of Cmpd-6 analogs (Table 1) were carried out. We evaluated the allosteric effect of these derivatives on orthosteric agonist $^3$H-FEN binding to the $\beta_2$AR in the absence and presence of transducers, either trimeric G protein or $\beta$-arrestin1. We also tested them for their allosteric activity in ISO-stimulated $\beta_2$AR downstream signaling, that is G protein–mediated cAMP production and $\beta$-arrestin2 recruitment to the activated receptor. For convenience of presenting the SAR analyses, Cmpd-6 and its analogs have been broken down into a common core scaffold, 5-hydroxysulfonflyl-2-mercaptobenzaldehyde (black), and three diverse substituents (R): N-methylpropan-2-amine (R1, blue), 4-methoxy-benzene (R2, red), and (R)-3-amino-4-(4-tert-butyl)phenyl) butanamide (R3, purple). The R1 region is common in this subset of analogs although varying in the initial analogs from the screening, as shown in Fig. 1B. The main focus of this initial analog series was to probe the importance of modifications at the R2 and R3 positions of Cmpd-6; these modifications are shown in Table 1. When the whole R3 portion of Cmpd-6 was replaced with a much smaller piperidine moiety as in analog A3, the PAM activity of Cmpd-6 was abolished, indicating that the bulkiness imparted by the tert-butyl at this region of the molecule is critical for the PAM activity of the compound. Further exploration of the relationship between activity and structure at R3, achieved by replacing the 4-tert-butyl on the benzene ring with 4-OH, and the amide tail with carboxyl as in analog A6, as well as with 4-H as in analog A7, revealed the importance of the large lipophilic ring system for the activity of Cmpd-6. This was more evident with the highly polar analog A6, which was considerably less active than the original compound, whereas the relatively small, but lipophilic, analog A7 showed moderate activity. SAR around R2 also showed significant decreases in the PAM activity of Cmpd-6, when its electron-donating group 4-OCH$_3$ was replaced with an electron-withdrawing 4-OCF$_3$ group at its para position to yield analog A4. Interestingly, when the para-OCH$_3$ group in Cmpd-6 was replaced with a meta-OCH$_3$ substitution, as in A5, there was a moderate decrease in the PAM activity of Cmpd-6. The results from these two analogs suggest that there may be a polar interaction, like H-bonding type, located in the area surrounding this 4-position of R2, which interacts with the putative binding site on the $\beta_2$AR.

**Discussion**

In the present study, DEL screening with the agonist-occupied $\beta_2$AR in HDL particles has yielded the first small-molecule PAMs for the $\beta_2$AR. We had isolated a small-molecule $\beta_2$AR NAM, Cmpd-15, from a previous screening (Ahn et al., 2017), but until now no PAM small molecule for this receptor had been described. Among these isolated PAM molecules, the most efficacious one, Cmpd-6, was characterized in detail through multiple assays. Cmpd-6 has a low micro-molar binding affinity for the agonist-occupied $\beta_2$AR and displays potent PAM activity for this receptor. Cmpd-6 positively cooperates with $\beta_2$AR agonists to enhance downstream signaling responses such as cAMP production and $\beta$-arrestin recruitment to the receptor. It not only potentiates the affinity (EC$_{50}$ values) of agonists for these responses, but also increases the agonist-induced maximal level of the responses. This is in agreement with pharmacological studies wherein Cmpd-6 shows positive cooperativity with transducers in mediating the agonist-bound high-affinity state of the $\beta_2$AR. Furthermore, Cmpd-6 stabilizes the agonist-induced active conformation of the $\beta_2$AR, leading to potentiation of the agonist-binding affinity for the receptor, which is a hallmark of PAMs (Langmead, 2011; Wootten et al., 2013; Christopoulos, 2014). Although Cmpd-6 does not show unique probe dependence or obvious biased activity toward either G protein or $\beta$-arrestin signaling, it clearly displays a ceiling effect for its activity, another pharmacological characteristic of PAMs (Wootten et al., 2013; Christopoulos, 2014). As expected for a PAM, Cmpd-6 also shows strong specificity for the $\beta_2$AR relative to the $\beta_1$AR and other receptors tested in this study. It only minimally potentiates the binding affinity of agonists to the $\beta_2$AR, as well as $\beta$ agonist–induced downstream functional activity.

Through a set of Cmpd-6 derivatives, we were able to discern the SAR patterns of the positive allosteric modulation of the $\beta_2$AR agonist activities and where the potential pharmacophore regions of the compound might be. As is apparent from our SAR studies, the N-isopropyl-N-methyl group is the preferred substituent at the R1 position attached to the common core chemical scaffold, 5-hydroxysulfonflyl-2-mercaptobenzaldehyde. At this region, bulkier groups such as N-cyclopentyl and N-phenyl result in increasingly poor PAM activity. At the R2 position, an electron-donating methoxy group, in the para position on the phenyl ring, is favored. This suggests that there may be a polar interaction...
TABLE 1
Structure–activity relationships of Cmpd-6 analogs
Different chemical scaffolds in the R2 and R3 regions between Cmpd-6 and its analogs are illustrated. Changes in the Vmax value by Cmpd-6 or each analog at 32 μM are expressed as percentages of the maximal level of the ISO-induced activity in the vehicle (DMSO) control in each assay. Changes in the EC50 value are expressed as fold shifts compared with the control value obtained in the vehicle (DMSO)-treated curve in each assay. Every value represents mean ± S.D. obtained from four independent experiments done in duplicate. Statistical analyses were performed using one-way analysis of variance with Dunnett’s multiple comparison post-tests compared with the control Cmpd-6–treated value in each assay. Adjusted *P < 0.05; **P < 0.01; ***P < 0.001.

| Cmpds | R2       | R3       | Re alone (%) | + Ge (%) | Re alone (%) | + βarr-1 (%) | E-Max (%) | EC50 shift (fold) | E-Max (%) | EC50 shift (fold) |
|-------|----------|----------|--------------|----------|--------------|--------------|-----------|-------------------|-----------|-------------------|
| DMSO  | 100      | 100      | 100          | 100      | 100          | 100          | 100       | 1.0               | 100.0     | 1.0               |
| Cmpd-6|          |          | 845.9 ± 129.0 | 159.3 ± 29.7 | 960.6 ± 145.5 | 284.9 ± 42.5 | 136.7 ± 18.1 | 5.2 ± 0.89*** | 168.7 ± 10.7*** | 5.1 ± 0.76*** |
| A3    |          |          | 163.5 ± 52.6*** | 93.9 ± 13.2*** | 111.6 ± 19.9*** | 100.4 ± 12.5*** | 79.0 ± 25.7*** | 0.9 ± 0.34*** | 58.6 ± 2.9*** | 0.9 ± 0.22*** |
| A4    |          |          | 207.7 ± 63.0*** | 81.6 ± 16.2*** | 194.9 ± 42.7*** | 138.5 ± 32.0*** | 115.5 ± 5.7* | 1.2 ± 0.32*** | 91.0 ± 6.9*** | 1.1 ± 0.16*** |
| A5    |          |          | 370.0 ± 106.5*** | 102.8 ± 7.0*** | 242.9 ± 49.7*** | 153.8 ± 22.2*** | 102.4 ± 16.4*** | 1.9 ± 0.55*** | 238.3 ± 8.2*** | 3.2 ± 0.64*** |
| A6    |          |          | 144.8 ± 20.3*** | 82.5 ± 9.2*** | 123.1 ± 23.5*** | 104.2 ± 22.3*** | 108.7 ± 2.2** | 1.0 ± 0.34*** | 103.7 ± 8.8*** | 0.8 ± 0.39*** |
| A7    |          |          | 249.3 ± 85.4*** | 103.8 ± 25.1*** | 207.0 ± 38.5*** | 129.7 ± 20.8*** | 76.6 ± 8.1*** | 1.8 ± 0.64*** | 207.4 ± 7.6*** | 2.7 ± 0.47*** |

Rc, receptor.

3H-Fenoterol High-Affinity Binding

- β2AR membrane
- Phosphorylated β2AR membrane
- G protein cAMP accumulation
- β-Arrestin recruitment

Cell-Based Assays

Visual representation of chemical structures is shown.
involving the methoxy functional group at the R2 position that interacts with the amino acid residues of the β2AR site. In the case of the R3 position, our analysis indicates that bulky and hydrophobic groups, such as tert-butyl benzene chemical scaffolds, are favored. This finding therefore suggests that this region of the molecule may occupy a hydrophobic pocket, deep within a putative β2AR allosteric binding site to establish contacts with core hydrophobic residues. Solution of an X-ray crystallographic structure of the β2AR in complex with Cmpd-6 will provide further insights into the binding modes of the compound.

Because allostERIC ligands are able to freeze or lock receptors into specific conformations by virtue of their cooperative interactions with orthosteric ligands, they can facilitate the study of receptors by biophysical techniques. Recently, atomic-level structural features of several GPCRs occupied by their allosteric ligands have been elucidated through X-ray crystallographic analyses (Kruse et al., 2013; Zhang et al., 2015; Jazayeri et al., 2016; Oswald et al., 2016; Zheng et al., 2016; Liu et al., 2017; Wacker et al., 2017). Such biophysical studies have revealed mechanisms by which allosteric ligands modulate the binding and action of orthosteric ligands. To date, most of the solved GPCR crystal structures together with allosteric ligands have been achieved with allosteric antagonists (or NAMs). An X-ray crystallographic study of the M2 muscarinic acetylcholine receptor (mACHr) occupied by its high-affinity orthosteric agonist and PAM, in a complex together with a transducer receptor (mAChR) occupied by its high-affinity orthosteric agonist, is currently used as a therapeutic drug for hyperplasia (2AR), hold great potential for the development of better therapeutic treatments for diseases like asthma, for which the clinical use of current 2AR agonists is limited by adverse side effects (National Asthma Education and Prevention Program, 2007).

The present study yielding PAMs, together with our previous work isolating a NAM (Ahn et al., 2017), strongly demonstrates that our current DEL screening approach with purified GPCRs can be used to accomplish target conformation-specific selection through in vitro manipulation of the receptors. Accordingly, we successfully isolated a NAM using the unoccupied β2AR in an inactive conformation, but obtained PAMs using the high-affinity agonist-occupied receptor in active conformations. In the future, complexes of the receptors with transducers G protein and β-arrestin, which are also allosteric molecules, could be used to isolate allosteric molecules that might have even more unique biased functional profiles.

In conclusion, in this study we introduce the discovery of the first small-molecule PAMs for the β2AR through an in vitro affinity-based iterative selection of highly diverse DELs against the agonist-occupied receptor in HDL particles. Characterization of the strongest PAM among these molecules reveals its positive cooperativity with orthosteric agonists in a wide range of receptor functions and its high selectivity for the β2AR. A number of pharmacological features of this compound suggest potential advantages of such a PAM over orthosteric agonists as a therapeutic drug. Finally, our current findings, together with our previous isolation of the first β2AR NAM (Ahn et al., 2017), establish a proof-of-concept strategy to isolate allosteric molecules with tailored functional profiles.

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