G protein–coupled receptors (GPCRs) activate cellular responses ranging from odorants to neurotransmitters. Binding an agonist leads to activation of a heterotrimeric G protein (GP) that stimulates external signaling. Unfortunately, the mechanism remains unknown. We show for 15 class A GPCRs, including opioids, adrenergics, adenosines, chemokines, muscarinics, cannabinoids, serotonin, and dopamines, that interaction of an inactive GP, including Gs, Gi, Go, G11, and Gq, to the inactive GPCR, containing the intracellular ionic lock between transmembrane (TM) helices 3 and 6, evokes thermally to form a precoupled GPCR-GP complex with an opened TM3-TM6 and the GP-α5 helix partially inserted into the GPCR but not activated. We show that binding of agonist to this precoupled GPCR-GP complex causes the Gα protein to open into its active form, with the guanosine diphosphate exposed for signaling. This GP–first paradigm provides a strategy for developing selective agonists for GPCRs since it is the pharmacophore for the precoupled GPCR-GP complex that should be used to design drugs.

G protein activation | molecular metadynamics | biased agonists | opioids | adrenergic

G protein–coupled receptors (GPCRs) constitute the largest group of membrane receptors in eukaryotes. They are responsible for activating cellular responses to numerous bioactive molecules, including odorants, pheromones, hormones, and neurotransmitters (1). Binding these external signaling molecules activates a GPCR through two pathways: one involving heterotrimeric G proteins (GPs) and the other involving arrestins, often with quite different consequences. Since ~34% of all modern medicinal drugs (2, 3) act on one or more of the 800 human GPCRs (4), there is great interest in understanding how binding of a ligand elicits this signaling. The aim of this paper is to elucidate this mechanism.

Generally, it has been assumed that binding agonists to the inactive conformation of a GPCR shifts the equilibrium toward the activated conformation (5, 6), allowing the liganded GPCR to recruit the inactive GP with bound guanosine diphosphate (GDP) and then activate the GP to mediate specific cellular signaling (7). We refer to this paradigm as the ligand-first mechanism of GP activation. There is support for this mechanism from experiments that excluded GP to allow agonist to bind first, leading to subsequent full activation after adding GP. On the other hand, many experimental observations (8–13) and computational studies (10, 11, 14–16) have revealed that agonists alone often do not stabilize the active conformation of the GPCR, hindering the subsequent recruitment of GP. Thus, complex formation between class A GPCRs and their cognate GPs greatly relies on random collisions between the pair (17, 18). Given that different types of GPs exist in a cell, there is tight competition between different subtypes of GPs to possibly couple with a given liganded GPCR, which makes a complex formation between liganded GPCR and GP relatively slow. However, the cellular response through the activation of GPs was shown to be rather rapid (7, 19, 20). In addition, the intrinsic basal activity of GPCRs, which leads to constitutive activation of GPs, is another phenomenon that the ligand-first mechanism of GP activation cannot describe. Many of the GPCRs that display this intrinsic basal activity in the absence of a ligand (the apo state) are required for normal physiological functions (21). For example, serotonin receptors (5-HT_2A and 5-HT_2C) exhibit a high level of constitutive activity (22). Suppressing the constitutive activity of 5-HT_2A and 5-HT_2C triggers mechanisms underlaying depression and anxiety (22).

In this paper, we propose an alternate paradigm (Fig. 1) in which prior to ligand binding, the inactive GP interacts with the inactive GPCR to open the intracellular region by breaking the transmembrane (TM) 3-TM6 tight coupling (an interaction from R^350, part of the DRY motif, to a conserved residue at the cytosolic end of TM6) to form a stable precoupled complex. This precoupled complex remains at this resting state until an agonist binds to the GPCR-GP complex to open the tightly

**Significance**

We report the detailed atomistic mechanism for how molecules such as morphine, dopamine, or epinephrine binding outside of a cell to a G protein–coupled receptor (GPCR) in the cell membrane cause a G protein (GP) bound at the inside of the cell to break apart and signal the cell to influence appetite, anxiety, memory, cognition, learning, and sleep. Most surprising is that the GP binds first to the GPCR to form a precoupled complex that remains at rest until the drug binds to induce the signaling. Most important, it is the precoupled GPCR-GP structure that provides the basis for the design of therapeutics to maximize activity and selectivity.

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coupled Gα-GDP complex while further opening the intracellular region of the GPCR to form the fully activated agonist-GPCR-GP complex with the GDP available for exchange. We refer to this as the GP-first mechanism of GP activation.

Previous observations have shown that a number of GPCRs form a precoupled complex with their cognate GPs (23–28), but the detailed molecular mechanism of subsequent activation by an agonist is not understood. In this study, to pursue the GP-first mechanism of GP activation, we examined coupling of 15 class A GPCRs to a total of six different GPs and show that in all 15 cases, the inactive GP couples with the inactive GPCR to form a precoupled-GPCR-GP complex in which the intracellular TM3-TM6 tight link is broken and the GP-α5 helix is partially inserted into the GPCR. We refer to this as the precoupled GPCR-GP complex. The combinations examined are as follows:

a) 5-HT2A-serotonin receptor–Gq protein
b) A2A adenosine receptor–Gs protein
c) β2-adrenergic receptor–Gs protein
d) μ-opioid receptor–Gi1 protein
e) κ-opioid receptor–Gi1 protein
f) δ-opioid receptor–Gi1 protein
g) CCR5-chemokine receptor–Gi1 protein
h) CB1-cannabinoid receptor–Gi1 protein
i) A1-adenosine receptor–Gi2 protein
j) 5-HT1B-serotonin receptor–Go protein
k) D2-dopamine receptor–Go protein
l) M1-muscarinic receptor–Gi1 protein
m) M3-muscarinic receptor–Gq protein
n) α2A-adrenergic receptor–Gq protein
o) 5-HT2C-serotonin receptor–Gq protein

The predicted precoupled complexes are discussed in detail below.

Moreover, for 5-HT2A-serotonin receptor–Gq protein, A2A adenosine receptor–Gs protein, and μ-opioid receptor–Gi1 protein, we predicted how binding a full agonist to the precoupled tightly coupled Gα-GDP complex causes the Gα subunit to open and becomes activated with the GDP available for exchange.

In this paper, we investigate the GP-first mechanism of GPCR activation using long-scale molecular dynamics simulations (an aggregate of ∼20 μs) and metadynamics (metaD) simulations to follow the sequence of structural and energetic steps involved in activation of class A GPCRs and their cognate GP. The activation process goes through several metastable states in which GPCR and GP undergo several structural changes on the sequence toward activation. Some metastable states may be separated by high energy barriers that could take microseconds or longer. Thus, we used metaD simulations (29) incorporating collective variables to describe the slow degrees of freedom but biased to encourage each GPCR and its cognate GP to explore large regions of conformational phase space to track the activation in much reduced time.

Results

We expect that prior to ligand binding, inactive GPCRs and their cognate inactive GPs have sufficient time to interact and form a precoupled complex. Thus, we considered the interactions between class A inactive apo-GPCRs (which generally have tight cytosolic TM3-TM6 coupling, such as an ionic lock) with their cognate inactive GP-bound GDP complex. The cryogenic electron microscopy (cryo-EM) structure of apo-5-HT1A-Gi protein (30) indicates that GPs couple to unliganded GPCR in a similar fashion and geometry as they do with liganded GPCRs in the fully active state. Thus, we placed the inactive GPs close enough (31) to the inactive apo-GPCRs (SI Appendix, Methods), adopting a similar geometry and orientation revealed by cryo-EM and X-ray crystallography for the fully activated agonist-GPCR-GP complex, such that GPCRs and GPs could start interacting (Fig. 2A and SI Appendix, Figs. S2–S4). The inactive state of the GP does not have the Gα5 helix fully extended, so there is no steric clash between GP and

![Fig. 1. GP-first mechanism of GP activation. Prior to ligand binding, the inactive GP interacts with the inactive GPCR to open the intracellular region by breaking the TM3-TM6 tight link to form a stable precoupled complex. This precoupled complex remains at this resting state until an agonist binds to the GPCR-GP complex to open the intracellular region of GPCR and the tightly coupled Gα-GDP complex to form the fully activated agonist-GPCR-GP complex with the GDP available for exchange or release.](https://doi.org/10.1073/pnas.2110085119)
GPCRs. Finally, we compare this optimized geometry between GP and GPCRs after activation directly to the structures resolved by experiments.

**Gq Protein Precoupling to the Inactive 5-HT₂A Receptor.** We used the serotonin 5-HT₂A receptor as the prototype for our proposed GP-first activation pathway, and then we examined a similar molecular mechanism of activation for 14 other class A GPCRs, including the A₂A adenosine receptor and β₂ adrenergic receptor, the archetypes for class A GPCRs.

The inactive conformation of 5-HT₂A receptor has a salt bridge R173³⁵⁰-E318⁽⁶⁾₃⁰ (the superscript is Ballesteros-Weinstein numbering for GPCRs (32), taken from Pády-Szekerés et al. (33)), the ionic lock, that inhibits activation. Breaking and disruption of the ionic lock is believed to be an important step toward the activation of class A GPCRs (34-36). We hypothesize that during the formation of precoupled state between 5-HT₂A and Gq protein, the α₅ helix of Gq protein partially penetrates to the intracellular region of 5-HT₂A and makes a salt bridge with R173³⁵⁰ and E318⁽⁶⁾₃⁰, opening the ionic lock. The rearrangements in ionic interactions between the cytoplasmic end of 5-HT₂A receptor and the α₅ helix of Gq protein facilitates opening the ionic lock. To examine this hypothesis, we performed two independent metaD simulations for an aggregated ~1 μs, in which we evaluated the energetics of forming two salt bridges: V358 (terminal CO₂⁻)-R173³⁵⁰(CZ) and K353(NZ)-E318⁽⁶⁾₃⁰ (CD) and their consequences on the ionic lock between R173³⁵⁰(CZ)-E318⁽⁶⁾₃⁰ (CD).

We find that during formation of the precoupled complex (Fig. 2A), as the GP approaches the GPCR, the terminal CO₂⁻ at the end of the Gq₅ helix forms a salt bridge with R173³⁵⁰ (Fig. 2D), initiating breaking of the ionic lock. However, our ~400-ns metaD simulation (Fig. 2B and C) reveals that this salt bridge V358-R173³⁵⁰ is endothermic by 4.2 ± 0.2 kcal/mol (Fig. 2B). Thus, after opening the ionic lock between TM3-TM6, the V358 terminal carboxylate disengages from R173³⁵⁰ and penetrates deeper into the core of the receptor (Fig. 2E). Interestingly, K353 on the Gq₅ helix forms a persistent salt bridge with E318⁽⁶⁾₃⁰ (Fig. 2D and B) with a binding affinity of -1.0 ± 0.1 kcal/mol (Fig. 2C) that contributes substantially to breaking the R173³⁵⁰-E318⁽⁶⁾₃⁰ ionic lock while stabilizing the position of Gq₅ inside the intracellular region of 5-HT₂A.

To determine whether disruption of the ionic lock R173³⁵⁰-E318⁽⁶⁾₃⁰ in the precoupled 5-HT₂A-Gq protein complex is statistically significant, we carried out an independent free energy metaD simulation for ~600 ns (Fig. 2E). In this study, we separately evaluated the energetics of opening the ionic lock, finding that precoupling of Gq protein to 5-HT₂A breaks open the ionic lock to ~7 Å while reducing the energy by -1.8 ± 0.1 kcal/mol, a thermodynamically favorable process. Indeed, this disruption with breaking of TM3-TM6 coupling is well known to be a critical step in activation of class A GPCRs (34-36). We find that a persistent charge-charge or salt bridge interaction K353-E318⁽⁶⁾₃⁰ emerges in the precoupled complex concomitant with the Gq₅ terminal carboxylate penetrating into the core of the GPCR after the ionic lock is broken (Fig. 3O), a result similar to our first simulation.

There remains a possibility that the rigid-body orientation of Gq protein could be different in the precoupled state from that in the fully active complex. To eliminate the possibility that the specific rigid-body orientation of Gq protein is solely responsible for opening the TM3-TM6 coupling, we carried out an independent third metaD free energy calculation for ~1.5 μs in which only the Gq₅-α₅ peptide (the last 26 residues: 33⁽³⁾-V358) is placed in close proximity (K353 and V358 10 Å away from R173³⁵⁰ and E318⁽⁶⁾₃⁰, respectively) to the inactive 5-HT₂A (SI Appendix, Fig. S1). The increased degrees of freedom for the Gq₅-α₅ peptide enabled the metaD to explore numerous positions and orientations that would emerge from various orientations of the whole Gq protein in complex with...
the 5-HT2A. We find that prior to ligand binding, a salt bridge contact from the terminal carboxylate, V358, to R1733.50 ([SI Appendix](https://doi.org/10.1073/pnas.2110085119 pnas.org), Fig. S1A–C), contributes to opening the 5-HT2A ionic lock to 7 Å, consistent with the precoupled state of the whole Gq protein (Figs. 2E and 3O). These calculations confirm that formation of the precoupled 5-HT2A-Gq protein complex is not an artifact resulting from a specific rigid-body orientation of the Gq protein.

**Generalization of the GP-First Mechanism for Activation of Class A GPCRs.** We expect that inactive GPs generally have sufficient time to couple to inactive GPCRs prior to drug binding and that formation of a precoupled state applies generally to all class A GPCRs and their cognate GP. In our proposed GP-first paradigm:

1) The apo-GPCR initially exhibits a tight cytoplasmic region due to the interaction of R3.50 (part of the DRY motif) with TM6. This TM3-TM6 coupling for the inactive conformation of class A GPCRs can be either an ionic lock (Fig. 3A, B, and G–O), a hydrogen bond (Fig. 3C–E), or a hydrophobic interaction (Fig. 3F). Thus, the TM3-TM6 coupling constitutes major slow degrees of freedom along the activation path that need to be disturbed to accommodate the Gα5 helix for the emergence of the precoupled state and later activation (34–36).

2) During the formation of the precoupled state between class A GPCRs and their cognate GPs, the α5 helix of GPs partially penetrates the intracellular region of GPCRs and makes a salt bridge with R3.50 (part of the DRY motif), which breaks the tight TM3-TM6 coupling, opening up the cytoplasmic region of GPCRs.

To further validate our GP-first paradigm, we examined whether Gs protein alone can open up the TM3-TM6 coupling of the apo-β2 adrenergic receptor from its inactive conformation. We tested the β2 adrenergic receptor because it is one of the best-characterized class A GPCRs. Although the X-ray crystallographic study of the inactive β2 adrenergic receptor found a hydrophobic coupling between R1313.50-L2726.34, a previous computational study (37) revealed that the inactive conformation of the β2 adrenergic receptor with its native intracellular loop 3 forms an ionic lock between R1313.50-E2686.30. To optimize the inactive state of the β2 adrenergic receptor, we first inserted the native intracellular loop 3 into the crystallographic inactive state and then performed a ∼2.2-μs metaD simulation to find that R1313.50 makes an ionic lock with E2686.30. Subsequently, to examine if partial insertion of the Gα5 helix into the core of the β2 adrenergic receptor perturbs the ionic lock, we performed a ∼800-ns metaD simulation in which we evaluated the energetics of salt bridges involving the Gα5 helix: R389 to E2686.30 and E392 to R1313.50 ([SI Appendix](https://doi.org/10.1073/pnas.2110085119 pnas.org), Fig. S2). Our free energy calculations show that E392 forms a salt

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**Fig. 3.** Precoupled complexes of class A GPCR-GPs. Detailed atomic interactions between Gα5 helix and class A GPCRs in the precoupled state resulted from extensive metaD simulations for (A) β2-adrenergic receptor–Gs, (B) A2A-adenosine receptor–Gs, (C) μ-opioid receptor–Gi1, (D) κ-opioid receptor–Gi1, (E) δ-opioid receptor–Gi1, (F) CCR5-chemokine receptor–Gi1, (G) CB1-cannabinoid receptor–Gi1, (H) A1-adenosine receptor–Gi2, (I) 5-HT1B-serotonin receptor–Gα, (J) D2-dopamine receptor–Go, (K) M1-muscarinic receptor–Gi1, (L) mouse M3-muscarinic receptor–Gq, (M) α2A-adrenergic receptor–Gq, (N) 5-HT2C-serotonin receptor–Gq, and (O) 5-HT2A-serotonin receptor–Gq. The details of the calculations are represented in SI Appendix, Fig. S4 and Table S3. (C–E) Adapted from figure 6A of ref. 14.
bridge with R131\(^{3.50}\) (Fig. 3A) with a high binding affinity of \(-10.2 \pm 0.2\) kcal/mol. Simultaneously, E268\(^{6.30}\) (the partner in the ionic lock) establishes a high-affinity (\(-2.8 \pm 0.3\) kcal/mol) salt bridge with R389. As a result, the TM3-TM6 ionic lock opens fully to \(\sim 12\) Å (Fig. 3A), leading to the precoupled state. The significant roles of R389 and E392 in the precoupled complex agree with a recent mutagenesis (38) study indicating that R389 and E392 are essential for efficient formation of a complex between Gs protein and \(\beta_2\) adrenergic receptor. In fact, mutation of E392 to an Ala residue perturbed the initiation of GDP release (38). Additionally, a series of E392\(^{205}\) mutants to Ala, Arg, Gln, Val, Leu, and Ser exhibited impaired cyclic adenosine monophosphate accumulation (39), confirming that E392 serves a crucial role in the activation of Gs protein.

To examine the GP-first mechanism of activation for other class A GPCRs, we studied precoupling of 13 additional GPCRs to Gs (Fig. 3B), Gi/o (Fig. 3C–F), or Gq/11 (Fig. 3K–O). We also took all sorts of TM3-TM6 couplings—an ionic lock (Fig. 3B and G–O), a hydrogen bond (Fig. 3C–E), or a hydrophobic interaction (Fig. 3F)—into consideration. All sorts of TM3-TM6 couplings must be opened to accommodate the Gq5 helix for emergence of the precoupled state. Thus, we performed \(\sim 7,2\)-\(\mu\)s metaD simulations, with the main idea that the salt bridge interactions from the Gq5 helix to counterparts of the TM3-TM6 couplings, particularly R\(^{3.50}\) (part of the DRY motif), open up the cytoplasmic regions. To eliminate the probable impacts of the chosen collective variable from our results, we used various combinations of collective variables (SI Appendix, Figs. S3 and S4 and Table S3). For these calculations, we used either of two well-validated force fields [ChARMm36m (40) and AMBER14 (41); SI Appendix, Tables S2–S4] to eliminate the possibility that formation of the precoupled GPCR-GP complex results solely from the choice of a specific force field. Indeed, our free energy calculations find for 13 cases that the salt bridge interactions between the Gq5 helix and the intracellular region of the GPCR (particularly to the conserved R\(^{3.50}\)) break open the tight TM3-TM6 coupling to accommodate the Gq5 helix partially inside the core of receptors, leading to emergence of the precoupled GPCR-GP complex. Fig. 3 depicts the molecular interactions in the GPCR-GP precoupled complexes for all 15 cases. The GPCRs studied include the A\(_{2A}\) adenosine receptor (42, 43), D2 dopamine receptor (25, 44), \(\alpha\)-adrenergic receptor (23, 25), M3 muscarinic receptor (26), and AI adenosine receptor (25), which were previously shown to make a precoupled complex with their cognate GP.

**Agonist Activation of GP.** To determine the role of agonist in the GP-first activation paradigm, we inserted a full agonist, 25CN-NBOH, into the precoupled complex of 5-HT\(_{2A}\)-Gq protein such that a salt bridge from the conserved D155\(^{3.32}\) to the protonated N atom of the agonist locks the ligand into the orthosteric binding pocket of 5-HT\(_{2A}\) (SI Appendix; Fig. S5). Given that the cellular signaling through the GP activation arises from the exchange of a GDP for a guanosine triphosphate (GTP), we assessed the energetics of opening the Gq5 subunit from the cleft between \(\alpha\)-helical (AH) (the center of mass of C\(_{\alpha}\) for the residues 154 to 161 and 175 to 182) and Ras-like (the center of mass of C\(_{\alpha}\) for the residues 51 to 62) domains, which defines the nucleotides’ (GDP and GTP) binding pocket. Opening the tight Gq5 makes the GDP release or exchange facile (45, 46). Our free energy calculations show that the Gq5 subunit subsequently undergoes a remarkable opening, increasing the separation between AH and Ras-like domains from \(\sim 16\) Å (tight conformation) to \(\sim 23\) Å (open conformation) while opening the GDP binding site (Fig. 4A and B), a remarkable structural rearrangement induced by ligand binding. This process is energetically favorable (\(-7.5 \pm 0.6\) kcal/mol; Fig. 4A) and leaves the GDP exposed to water, making it susceptible to dissociation or GTP exchange. This opening of Gq5 expedites GDP release, a critical event in activation of GP and GP signaling (45, 46). Our metaD simulation finds that although opening the Gq5 protein provides an exit path for GDP dissociation, the GDP remains bound to the Ras-like domain in our simulations, consistent with previous experimental (47, 48) and computational (49, 50) studies showing that GDP still remained only bound to the Ras-like domain even when the Gq subunit opens up.

To examine whether the opening of GP from the GDP binding site prevails only in the presence of agonist, we also estimated the energetics for opening of the Gq5 subunit for

- a partial agonist, lysergic acid diethylamide (LSD);
- an inverse agonist, methiothepin; and
- the case with no ligand present in the orthosteric pocket of 5-HT\(_{2A}\) (SI Appendix; Fig. S5).

Our free energy calculations reveal that the presence of the partial agonist also induces the Gq5 subunit to open up from 16 to 23 Å (Fig. 4C and D) but in a less favorable process (\(\Delta G = -3.8 \pm 0.6\) kcal/mol) compared to the full agonist binding (\(\Delta G = -7.5 \pm 0.6\) kcal/mol). To compare the relative energy barrier of activation between these types of ligands in the same activation pathway, we reweighted (51) the metaD free energies to estimate the error. We find that the free energy barrier associated with activation for a full agonist (\(\Delta G = +3.3 \pm 0.6\) kcal/mol) is far more favorable than for a partial agonist (\(\Delta G = +6.8 \pm 0.5\) kcal/mol) (Fig. 4B and D).

On the other hand, in the absence of an agonist the free energy to activate the apo-5-HT\(_{2A}\)-Gq protein precoupled complex is endothermic by \(\Delta G = +4.6 \pm 0.6\) kcal/mol (Fig. 4E and F), consistent with a previous study (44) showing a low level of constitutive activity for 5-HT\(_{2A}\) even with a broken ionic lock. Indeed, we find that insertion of an inverse agonist selectively disfavors the opening of Gq protein (Fig. 4G and H) even further, increasing the free energy by \(\Delta G = +9.7 \pm 0.6\) kcal/mol (Fig. 4J) and producing a response opposite to the full agonist, which is consistent with the physiological role of an inverse agonist. Comparing the energy barrier of activation (Fig. 4F and H) for the apo-5-HT\(_{2A}\) (\(\Delta G = +9.8 \pm 0.6\) kcal/mol) and inverse agonist (methiothepin-5-HT\(_{2A}\)) \(\Delta G = +12.1 \pm 0.6\) kcal/mol) suggests that the inverse agonist suppresses constitutive activation of 5-HT\(_{2A}\) and Gq protein.

Opening the Gq5 subunit in the presence of a full agonist has a dramatic effect on the position of the Gq5 helix and consequently on the position of TM6. We find that in the fully activated 25CN-NBOH-5-HT\(_{2A}\)-Gq complex, the Gq5 helix undergoes a pronounced \(\sim 8\)-Å upward movement along its axis into the receptor core (Fig. 5A and B), allowing the Gq5 helix to rotate 50° around its axis relative to the Gq5 helix of the inactive conformation (Fig. 5A) and leading to extensive interactions with the cytoplasmic region of 5-HT\(_{2A}\) receptor. This translation along and rotation of the Gq5 helix around its axis are known hallmarks of GP activation (49, 52, 53), playing a key role in nucleotide release. Indeed, this outward movement of the Gq5 helix is associated with \(\sim 9\)-Å outward displacement of the cytosolic end of TM6 (Fig. 5C) to match closely the fully active state of 5-HT\(_{2A}\) resolved in the cryo-EM structure (54) with root mean-square displacement (rmsd) of 2.4 Å (Fig. 5D).

To independently determine if the presence of full agonists can activate the precoupled GPCR-GP complex for other GPCRs, we inserted
The full agonist 5'-N-ethylcarboxamide adenosine (NECA) into the A2A adenosine receptor–Gs protein complex (Fig. 6A), and the full agonist morphine into the μ-opioid receptor–Gi protein complex (Fig. 6D and SI Appendix, Fig. S3).

Our free energy calculations show that the full agonists

- open the distance between the G α s subunits for the A2A adenosine receptor by ~10 Å in the cleft between AH and RAS-like domains (Fig. 6A)
- open the distance between the G α i subunits for the μ-opioid receptor by ~8 Å in the cleft between AH and RAS-like domains (Fig. 6D).

These structural rearrangements make GDP exchange and subsequent cellular signaling facile, confirming that agonists convert the precoupled to the fully active state in a thermodynamically favorable pathway.

An important implication of the GP-first mechanism of GP activation is that it is the structure the precoupled GPCR-GP complex that should be used to provide the pharmacophore for structure-based drug design of agonists to maximize activity and selectivity. Indeed, most important may be the structure at the transition state for opening the G α , which probably dominates activity. Drugs must aim at quickly activating the precoupled complex of the target GPCR-GP complex while blocking activity of the precoupled complex for all other GPCR-GP complexes to reduce undesirable side effects so common in GPCR-targeted therapies (the target-antitarget strategy).

Discussion

We expect that prior to ligand binding, inactive GPCRs and their cognate inactive GP have sufficient time to associate, producing a precoupled complex. This coupling disrupts the intrinsic tight coupling of the cytoplasmic ends of TM3-TM6 that keeps the class A GPCRs inactive. Although this disruption of TM3-TM6 coupling is essential to activation, it need not necessarily result in a remarkable outward displacement of TM6 from TM3, the well-known structural rearrangement associated with activation of class A GPCRs (53). This precoupled GPCR-GP complex remains at rest until the agonist arrives to drive the precoupled complex to its final activated state, during which TM6 experiences the large outward movement necessary to fully accommodate the G α  helix. Concomitantly, ligand binding opens the G α protein to expose the GDP and allows G α 5 to move outward while rotating about its axis to interact extensively with the intracellular GPCRs region.

On the other hand, NMR spectroscopy (8, 11, 12), double electron-electron resonance spectroscopy (12), crystallography (10), and computational (10, 11, 16) studies indicate that full agonists (high-efficacy ligands) alone do not stabilize the active conformation of GPCRs. Rather, they significantly increase the basal activity of GPCRs (imposing outward TM6 displacement), which was shown to be just sufficient for the recruitment and later activation of the GP (9). Indeed, our analysis shows that agonists alone cannot stabilize the active state conformation since they cannot break the tight TM3-TM6 coupling (14) in the inactive apo-GPCR (SI Appendix, Fig. S8).
This is in stark contrast to our GP-first paradigm in which the strong affinity between GP and GPCRs leads to opening the tight TM3-TM6 coupling. Thus, prior to ligand binding, GP can interact directly with GPCRs to stabilize a precoupled complex (23–28). Indeed, it was shown experimentally for the M3 muscarinic receptor that the precoupled complex between inactive GP and inactive GPCR eventually leads to rapid GP activation once the agonist binds the receptor (26), consistent with our results. Moreover, the basal activity of many GPCRs in the absence of an agonist leads to constitutive activation of the GP (21), showing that GP activation can proceed without agonists, further evidence supporting our results that GP precoupling to GPCRs is a viable activation pathway. Interestingly, the cryo-EM structure of apo-5-HT1A-Gi protein (30) indicates that GP precoupling to the GPCR results in a complex similar to the one resolved in the presence of a full agonist.

Thus, agonists must bind sufficiently strongly to the precoupled GPCR-GP complex to force opening of the Gα subdomain. Indeed, for 5-HT2A, we found that the full agonist leads to a smaller barrier for Gα opening, compared to a partial agonist or inverse agonist, to induce release of GDP while progressing toward the activated structure.

Unfortunately, the only knowledge about this ligand bound precoupled structure is from our simulations. The structure for

Fig. 5. Fully active state of 5-HT2A-Gq protein. Comparison of Gqα in our optimized fully active 25CN-NBOH-5-HT2A-Gq protein with (A) inactive Gqα protein-bound GDP and (B) fully active nucleotide free mini-Gqα subunit (54) resolved by cryo-EM. Comparison of the cytoplasmic region of 5-HT2A in our optimized fully active 25CN-NBOH-5-HT2A-Gq protein with (C) the inactive conformation (54) (Protein Data Bank [PDB]: 6WH4) resolved by X-ray crystallography and (D) the active conformation (54) (PDB: 6WHA) resolved by cryo-EM.
the agonist bound to the precoupled complex has not yet been observed experimentally.

These insights on the mechanism of activation provide strategies for designing agonists. Thus, the key to agonist design is for the ligand to bind strongly to the target precoupled GPCR-GP complex in such a way as to reduce the barrier for opening the $\Gamma^\alpha$-bound GDP to release or exchange with GTP. Concomitantly, we need to examine the precoupled GPCR-GP complexes for all off-target GPCRs to ensure that the agonist does not activate them. Thus, armed with the precoupled GPCR-GP complexes for all relevant GPCRs, we should be able to design drugs optimally for the target signaling while suppressing all off-target signaling (the target-antitarget strategy for drug design).

Our calculations predict at least 10 to 20 key interactions for each GPCR-GP combination (Fig. 3) to motivate mutation experiments to validate (or not) our predictions.

**Data Availability.** All study data are included in the article and/or SI Appendix. Model structures have been deposited in GitHub (https://github.com/amafi-gpcr/G-protein-first-mechanism-of-activation-for-class-A-GPCRs-PNAS-2022), (31).

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