Exploring the signaling space of a GPCR using bivalent ligands with a rigid oligoproline backbone

Nina Romantini\(^a\), Shahidul Alam\(^b,\) Stefanie Dobitz\(^b\), Martin Spillmann\(^b\), Martina De Foresta\(^c\), Roger Schibli\(^b\), Gebhard F. X. Schertler\(^b,\) Helma Wennemers\(^b,\) Xavier Deupi\(^e,\) Martin Behe\(^d\), and Philipp Berger\(^e,\)\(^f\)

\(^a\)Center for Radiopharmaceutical Sciences, Paul Scherrer Institute, CH-5232 Villigen, Switzerland;\(^b\)Laboratory of Nanoscale Biology, Paul Scherrer Institute, CH-5232 Villigen, Switzerland;\(^c\)Laboratory of Organic Chemistry, ETH Zurich, CH-8093 Zurich, Switzerland;\(^d\)Department of Biology and Chemistry, Paul Scherrer Institute, CH-5232 Villigen, Switzerland;\(^e\)Laboratory of Biomolecular Research, Paul Scherrer Institute, CH-5232 Villigen, Switzerland; and\(^f\)Condensed Matter Theory Group, Paul Scherrer Institute, CH-5232 Villigen, Switzerland

Edited by Robert J. Lefkowitz, HHMI, Durham, NC, and approved October 27, 2021 (received for review May 11, 2021)

G protein–coupled receptors (GPCRs) are one of the most important drug–target classes in pharmaceutical industry. Their diversity in signaling, which can be modulated with drugs, permits the design of more effective and better-tolerated therapeutics. In this work, we have used rigid oligoproline backbones to generate bivalent ligands for the gastrin-releasing peptide receptor (GRPR) with a fixed distance between their recognition motifs. This allows the stabilization of GPCR dimers irrespective of their physiological occurrence and relevance, thus expanding the space for medicinal chemistry. Specifically, we observed that compounds presenting agonists or antagonists at 20- and 30-Å distance induce GRPR dimerization. Furthermore, we found that 1) compounds with two agonists or antagonists at 20- and 30-Å distance induce GRPR agonistic behavior. We show the stabilisation of GPCR dimers through their transmembrane helices (9, 10). Crystal structures also point toward homodimerization based on transmembrane interactions. For instance, the μ-opioid receptor can dimerize through two different interfaces: a larger interface formed by transmembrane (TM) helices TM5 and TM6 and a smaller interface formed by TM1, TM2, and the intracellular helix 8 (11). The functional relevance of homo- and heterodimerization has been shown for many other GPCRs (12, 13). Homo- and heteromers can exhibit different pharmacology than monomers and, therefore, represent an obvious therapeutic target. Bivalent ligands for opioid receptors were already introduced in the 1980s before natural association of GPCRs was shown (14, 15). In the meantime, many bivalent ligands, especially for opioid and serotonin receptors, were synthesized. Compared to typical beta blockers. Besides the clinically approved carvedilol, several potential biased drugs are in clinical or preclinical trials (6, 7).

While most GPCRs are able to function as monomers, there is clear evidence that dimerization (or even higher-order oligomerization) represents an additional layer of regulation and a fundamental aspect of receptor function (6). Nevertheless, GPCR oligomerization is still a controversially discussed field, probably because several modes of oligomerization exist. For example, the class C GPCRs GABA\(_{B1}\) and GABA\(_{B2}\) are linked together by a cytoplasmic coiled coil to form a functional GABA\(_{B}\) receptor dimer, whereas 5-HT2A and mGluR2 interact through their transmembrane helices (9, 10). Crystal structures also point toward homodimerization based on transmembrane interactions. For instance, the μ-opioid receptor can dimerize through two different interfaces: a larger interface formed by transmembrane (TM) helices TM5 and TM6 and a smaller interface formed by TM1, TM2, and the intracellular helix 8 (11). The functional relevance of homo- and heterodimerization has been, in the meantime, shown for many other GPCRs (12, 13). Homo- and heteromers can exhibit different pharmacology than monomers and, therefore, represent an obvious therapeutic target. Bivalent ligands for opioid receptors were already introduced in the 1980s before natural association of GPCRs was shown (14, 15). In the meantime, many bivalent ligands, especially for opioid and serotonin receptors, were synthesized.

G protein–coupled receptors | cell signaling | receptor dimerization

G protein–coupled receptors (GPCRs) are one of the most important drug–target classes in pharmaceutical industry. Their diversity in signaling, which can be modulated with drugs, permits the design of more effective and better-tolerated therapeutics. In this work, we have used rigid oligoproline backbones to generate bivalent ligands for the gastrin-releasing peptide receptor (GRPR) with a fixed distance between their recognition motifs. This allows the stabilization of GPCR dimers irrespective of their physiological occurrence and relevance, thus expanding the space for medicinal chemistry. Specifically, we observed that compounds presenting agonists or antagonists at 20- and 30-Å distance induce GRPR dimerization. Furthermore, we found that 1) compounds with two agonists or antagonists at 20- and 30-Å distance induce GRPR agonistic behavior. We show the stabilisation of GPCR dimers through their transmembrane helices (9, 10). Crystal structures also point toward homodimerization based on transmembrane interactions. For instance, the μ-opioid receptor can dimerize through two different interfaces: a larger interface formed by transmembrane (TM) helices TM5 and TM6 and a smaller interface formed by TM1, TM2, and the intracellular helix 8 (11). The functional relevance of homo- and heterodimerization has been shown for many other GPCRs (12, 13). Homo- and heteromers can exhibit different pharmacology than monomers and, therefore, represent an obvious therapeutic target. Bivalent ligands for opioid receptors were already introduced in the 1980s before natural association of GPCRs was shown (14, 15). In the meantime, many bivalent ligands, especially for opioid and serotonin receptors, were synthesized.
Critical factors in the development of these compounds are the length, rigidity, and water solubility of the linker. Most of the time, flexible linkers with a length of 18 to 25 atoms, corresponding to ~20- to 30-A˚ distance, have been used (reviewed, for example, in refs. 16 and 17).

Oligoprolines are conformationally well-defined molecular scaffolds that can be functionalized at defined sites to obtain bivalent ligands. Oligoproline derivatives are well soluble in water under physiological conditions and therefore ideally suited scaffolds for applications in aqueous media. In aqueous environments, oligoprolines adopt a highly symmetric polyproline II helix conformation in which every third residue is stacked on top of each other at a distance of ~10 Å (18). The functionalization pattern of this molecular scaffold can be easily fine-tuned by modular chemical synthesis reminiscent to an “LEGO” approach. In addition to their application for targeting GPCRs, oligoprolines have also shown their value in the development of inhibitors of protein–protein interactions, cell-penetrating peptides, hierarchical supramolecular assemblies, and the controlled formation of silver nanoparticles (19).

In this work, we use functionalized oligoproline backbones to induce artificial homodimers of the gastrin-releasing peptide receptor (GRPR) (now properly known as bombesin BB2 receptor) to explore its signaling space. GRPR is a class-A GPCR that belongs to the bombesin family, and its homodimerization has not been documented so far. It is mainly expressed in different regions of the gastrointestinal (GI) tract and the brain (20, 21). GRPR does not only regulate various functions in the GI tract, such as gut hormone secretion and GI motility (22), but also many processes of the central nervous system like the regulation of memory, fear, and itching (23). Interestingly, GRPR was found to be overexpressed on different cancer types, among them prostate cancer and breast cancer, and its overexpression is accompanied by an ability to accelerate growth of cancer cells and increase their invasive potential (24, 25). Studies in tumor xenograft mouse models already showed that GRPR antagonists are able to decrease tumor growth rate (26), suggesting the interest of GRPR as a drug target. In addition, the specific delivery of radiolabeled peptides to tumor cells overexpressing the receptor enables targeted radiotherapy and nuclear imaging of these cancers.

Here, we show that bivalent oligoproline-based ligands with a distance of 20 and 30 A˚ between the recognition motifs are indeed able to induce dimerization of GRPR. The consequences for signaling of these changes in the oligomerization state were evaluated by measuring recruitment of six different adaptor proteins from three gene families (Gαq, β-arrestin-1/2, and GRK2/3/5). We show that dimerization can, indeed, influence the recruitment profile of GRPR, and we observe different recruitment efficacies and potencies of our tested drugs. In addition, we observed a strong effect on Gq recruitment of our bivalent ligand that was unable to induce dimerization, suggesting an allosteric effect.

Results

Ligands. For our studies, we used divalent ligands consisting of oligoprolines as rigid scaffolds that allow for tailoring distances of 10, 20, and 30 Å between agonistic and/or antagonistic ligands. In particular, as recognition motifs, we used truncated derivatives of bombesin, such as the targeting motif from RM1 (27) as antagonist and the targeting motif from AMBA (28) as agonist, in which we replaced the C-terminal methionine with a norleucine. Our oligoproline backbones were modified with a 2,2′,2′′,2′′′-(1,4,7,10-Tetraazacyclododecane-1,4,7,10-tetrayl)-tetraacetic acid (DOTA) chelator that can be loaded with radionuclides, since they were initially designed for radiopharmaceutical applications (Fig. 1A) (29). The recognition motifs coupled to oligoproline backbones yielded monovalent agonists/antagonists with the recognition motif in different distances to the DOTA chelator (30) and bivalent agonists/antagonist with 10-, 20-, and 30-Å distance between recognition motifs [(30, 31); see SI Appendix, Fig. 1 for characterization of previously unpublished ligands]. In addition, hybrid compounds with the agonist and the antagonist separated by 10, 20, and 30 Å were used (31). The schematic structure and nomenclature of the studied compounds are given in Fig. 1.

Bivalent Ligands with 20- and 30-Å Distance Induce Dimerization. Dimerization of GRPR was measured by coexpressing an NLuc-tagged and a Cherry-tagged GRPR (Fig. 2A). Bringing these two receptors in close proximity on the plasma membrane by dimerization leads to bioluminescence resonance energy transfer [BRET (32)]. No increase in the BRET signal was observed when cells were stimulated with AMBA or RM1, our reference compounds, indicating that these two compounds do not influence the oligomerization status of GRPR. This was also observed when a single agonist or antagonist was bound to an oligoproline backbone (Fig. 2B and C). However, stimulation with 20- and 30-Å ligands (agonists, antagonists, and hybrids) resulted in an increase of the BRET signal within minutes, indicating dimerization (Fig. 2 D–F). For hybrid and antagonistic bivalent ligands, saturation was reached after 1 to 3 min and for bivalent agonists within 10 min. As a control, we also used a GRPR mutant (GRPR-R288A-NLuc) that was previously shown to be unable to bind to ligands (33, 34). No dimerization was observed with this mutant, indicating that ligand binding is necessary for dimerization (SI Appendix, Fig. 2A).

These different kinetics might be related to the internalization of the receptors stimulated by agonists. We also observed that the BRET signal for the bivalent hybrids and the antagonist with 30-Å distance is lower. Nevertheless, these values (kinetics and signal strength) should not be compared directly, since BRET signals depend on the distance and relative arrangement between BRET donor and acceptor, which are unknown. Stimulating cells with bivalent ligands at high concentrations can lead to “high-dose inhibition,” in which a bivalent ligand recruits only one receptor molecule due to saturation of receptors (35). However, we do not see a decrease in the dimerization signal at high concentrations as observed with the unconjugated ligands (10 μM). Stimulation at high concentrations does not occur in our experimental setting (SI Appendix, Fig. 2B). The 10-Å bivalent ligands, on the other hand, did not induce a substantial change in the BRET ratio. These results indicate that the 20- and 30-Å ligands can induce receptor dimerization by “catching” the receptors on the plasma membrane and forcing them into close proximity. Conclusively, a distance of 20 and 30 Å between the ligands is suitable for accessing the binding pockets of two GRPR molecules at the same time, while 10 Å is too short to allow the concurrent interaction with two receptors.

Recruitment of Adaptor Proteins to Activated GRPR. Since GRPR homodimerization has not been reported, we were interested to find if our dimer-inducing ligands can also influence the balance between the receptor signaling pathways (i.e., induce biased signaling). Signal transduction bias can arise at the level of the ligand, the receptor, or the cell (6). As we are, in this work, exclusively interested in the effect of the ligands, we recorded time-resolved recruitment of adaptor proteins (G proteins, arrestins, and G protein-coupled receptor kinases [GRKs]) as a measure of their ability to alter receptor signaling. These assays are either based on enzyme complementation [split NanoLuc (36)] or BRET (37) (SI Appendix, Fig. 3 B–D). With this strategy, we reduce the possible influence of the cell type that we are interested to find if our dimer-inducing ligands can also influence the balance between the receptor signaling pathways (i.e., induce biased signaling).
Data were acquired at different ligand concentrations between 0.001 and 5 μM. Time-resolved data for the reference compound AMBA is shown in Fig. 3; data for all other ligands is shown in SI Appendix, Fig. 4. For measuring Gq recruitment, we used a recently published BRET assay based on engineered Gα proteins (37). These probes provide an excellent signal-to-noise ratio but have the disadvantage that they are not released after binding. GRPR recruited the mGαq probe efficiently, reaching saturation within 10 min after stimulation with 1-μM agonist (Fig. 3A and SI Appendix, Fig. 4A).

GRPR interacts with both nonvisual arrestins (38). In the present study, β-arrestin-1 and β-arrestin-2 recruitment was measured using a split Nanoluc assay (39) A sharp increase in luminescence was observed after stimulation with high-agonist concentrations, reaching maximal signal within 3 min after stimulation (Fig. 3B and C and SI Appendix, Fig. 4B and C). The following decay of the signal was first steep and exponential but flattened after ~10 min to a slow decay. This biphasic reaction might be explained by the continuous synthesis/recycling/degradation of the receptor as described previously (36, 40).

Fig. 1. Overview of used compounds. (A) AG-1-BBN-1 as an example for a monovalent compound. All monovalent compounds contain three blocks of three prolines. Agonists/antagonists were coupled to block 1, 2, or 3. (B) Bivalent ligands consists of two to four oligoproline blocks. The first ligand is always attached to block 1. The second ligand is linked to block 2, 3, or 4, resulting in bivalent ligands with a distance of 10, 20, and 30 Å between recognition motifs. Nomenclature and distances are given on the right side. (C) Chemical structures of the agonist (AMBA) and the antagonist (RM1).

Fig. 2. Bivalent ligands with 20- and 30-Å distance induce receptor dimerization. (A) Dimerization was measured using HEK293 cells coexpressing an NLuc- and Cherry-tagged GRPR. (B and C) Monovalent agonists and antagonists were not able to induce receptor dimerization. (D–F) Bivalent agonists, antagonists, and hybrid compounds with 20- and 30-Å distance between ligands induced dimerization, whereas bivalent compounds with 10-Å distance failed to induce dimerization. The curves represent time-resolved measurements of one representative experiment out of three (mean ± SEM of triplicate). The orange dotted line indicates the time point of the ligand addition. Control (Ctrl): no ligand addition.
At lower-ligand concentrations, luminescence increased more slowly and reached saturation after \( \approx 10 \) min.

GRKs are also known interaction partners of GPCRs. We used again split NanoLuc-based probes to measure GRK2, GRK3, and GRK5 recruitment to activated GRPR. GRK2/3 and GRK5 represent two branches of the GRK family that target different sequences leading to different phosphorylation patterns at the C-terminal tail of receptors (41). As for \( \beta \)-arrestins, a transient interaction with all three tested GRKS was observed (Fig. 3 D–F and SI Appendix, Fig. 4 D–F). Also, as in \( \beta \)-arrestins, maximal activation at high-ligand concentrations was reached within 3 min after stimulation, while at lower concentrations, the peak of the signal was shifted to later time points (\( \approx 10 \) min). Additionally, at lower concentrations, the recruitment of the adaptor proteins was no longer transient but remained at saturation during the whole time span of the assays.

**Effects on Gq Recruitment.** The data of the recruitment assays were then used to generate dose–response curves for all ligands (Fig. 4). From these dose–response curves, potencies (half maximal effective concentration, \( EC_{50} \)) and efficacies (\( E_{max} \)) of all drugs were obtained (Fig. 5). We observed a significantly increased efficacy toward miniGq recruitment for both dimer-inducing bivalent agonists (AG-2-BBN-2 and AG-2-BBN-3; Fig. 5A). The bivalent compound that is unable to induce receptor dimerization (AG-2-BBN-1) exhibits impaired miniGq potency and efficacy; this compound induced G-protein recruitment with an efficacy of only \( 57 \pm 6\% \) and was therefore significantly less efficient than monovalent ligands and the bivalent compounds AG-2-BBN-2 and AG-2-BBN-3. Furthermore, AG-2-BBN-1 also exhibited a reduced potency for G-protein recruitment as compared with AMBA, AG-2-BBN-2, and AG-2-BBN-3.

Interestingly, all three monovalent agonists with oligoproline backbones show a clearly enhanced potency when compared to AMBA, whereas efficacies are similar to the values of AMBA. This suggests that the oligoproline backbone with its linker or the DOTA moiety may be in contact with the receptor and have an effect on Gq recruitment or lead to higher affinity.

**Effects on \( \beta \)-arrestin/GRK Potency and Efficacy.** We observed that the efficacies of all bivalent agonists toward \( \beta \)-arrestin-1 and \( \beta \)-arrestin-2 are similar to the reference agonist AMBA. This is contrast to the observed increased efficacy of the dimer-inducing compounds (AG-2-BBN-2 and AG-2-BBN-3) for miniGq recruitment. Monovalent agonists show slightly reduced efficacies toward \( \beta \)-arrestin-1 and \( \beta \)-arrestin-2. In contrast, we observed a clearly reduced potency for \( \beta \)-arrestin-2 recruitment after stimulation with the bivalent agonist with 20-A distance (Fig. 5C). The monovalent agonists AG-1-BBN-2 and AG-1-BBN-3 showed both increased potencies for \( \beta \)-arrestin-2 recruitment. For GRK recruitment, we observed an increased potency with the dimer-inducing compound with 30-A distance (AG-2-BBN-3) for GRK2/3 but not for GRK5 (Fig. 5 D–F). In addition, increased efficacies for GRK5 recruitment were measured for all monovalent compounds (Fig. 5F) that correlate with the increase in Gq potency (Fig. 5A).

**Implications for Signaling Balance.** The balance between different signaling pathways plays an important role in drug action (6). For better visualization of the observed effects toward effector recruitment, we use a two-dimensional bias plot to display efficacy and potency (Fig. 6; see SI Appendix, Fig. 5 for a spider web representation). In our study, we use AMBA as reference agonist and consider it as unbiased. Our analysis reveals several interesting shifts in the signaling balance.

First, we observe that dimerization of GRPR leads to G-protein recruitment bias. Bivalent ligands show a clearly enhanced Gq efficacy with minor effects on \( \beta \)-arrestin and GRK2/3. In Fig. 6A, AG-2-BBN-2 and AG-2-BBN-3 are clearly shifted to the right of the graph, indicating G-protein bias (i.e., these compounds preferentially lead to G-protein recruitment over \( \beta \)-arrestin recruitment). Second, the nondimer–inducing bivalent ligand with 10-A distance (AG-2-BBN-1) has lower potency and lower efficacy for Gq recruitment. This leads to a signaling shift toward \( \beta \)-arrestin and GRKs (Fig. 6B). The effector recruitment properties of AG-2-BBN-1 also clearly differ from those of the monovalent compounds. Finally, we also observe different recruitment between \( \beta \)-arrestin-1 and

---

**Fig. 3.** Time-resolved measurement of adaptor protein recruitment to GRPR. Cells were stimulated with different concentrations of the reference compound AMBA at the indicated time point (dashed line). (A) BRET-based assay for miniGq recruitment. Nanoluciferase complementation assay for \( \beta \)-arrestin-1 (B), \( \beta \)-arrestin-2 (C), GRK2 (D), GRK3 (E), and GRK5 (F). Measurements for all other compounds are shown in SI Appendix, Fig. 4.
β-arrestin-2 for the two dimer-forming ligands. AG-2-BBN-2 (20 Å) has a lower potency toward β-arrestin-2. This leads to preference of AG-2-BBN-2 signaling toward β-arrestin-1 (yellow area in Fig. 6C). In contrast to that, the potencies of AG-2-BBN-3 (30 Å) toward β-arrestin-1 and β-arrestin-2 are similar to AMBA (orange area in Fig. 6C). AG-2-BBN-1, the bivalent agonist that does not induce receptor dimerization, did not exhibit a recruitment bias for any of the β-arrestins and recruited them with a similar potency to AMBA. Finally, we clustered the ligands according to their recruitment and dimerization potential. Data matrices were normalized, and k-means clustering was subsequently used to classify ligands. We observed that AG-2-BBN-1 forms a subtree with AMBA and that all monovalent ligands cluster together, as well as the bivalent agonists (SI Appendix, Fig. 6).

Effects of Hybrid Compounds on Effector Recruitment. We then also tested our hybrid compounds that are composed of an agonist and an antagonist. We observed for all compounds a clearly reduced efficacy for most assays (SI Appendix, Fig. 7). The mechanistic origin of this effect needs further investigation. Possible explanations are impaired internalization of the receptor or structural effects. Unfortunately, these low efficacies led to a reduced signal-to-noise ratio, which prevented us from getting concise potency data. Nevertheless, bivalent compounds with an agonist and antagonist could be promising when a reduced efficacy is desired.

Discussion

In this work, we aimed to induce forced dimers of GRPR to overcome the limitations of its physiological signaling profile and thereby expand the possibilities for medicinal chemistry toward this receptor. The oligomerization state of GRPR under resting conditions is not known, but we could show that binding of monovalent ligands does not change the oligomerization state. This does not exclude the idea that dimers or oligomers of GRPR may exist under resting conditions, as it was shown for several class A GPCRs, but they are probably not relevant for the activation process (42, 43).

Previous studies using bivalent ligands aimed to induce naturally occurring dimers and, therefore, mainly used flexible linkers to allow for a natural assembly of two receptors. In contrast, the oligoproline scaffold used in this work forms a stiff backbone that allows establishing defined distances between the attached ligands. Thus, binding of such more rigid bivalent ligands presumably reduces the mobility of attached GPCR dimers. We were able to induce dimers with oligoproline-based bivalent ligands at distance of 20 and 30 Å but not when the two recognition motifs were only 10 Å apart. This is in line with previous studies with flexible linkers that showed dimerization when ligands were spaced more than 15 Å (16, 17, 30).

We then investigated how these dimers may influence cellular signaling of GRPR by measuring recruitment of signaling partners. Surprisingly, we did not only observe effects of dimerization but also allosteric effects. In general, the effects of our ligands on Gq recruitment were larger than on β-arrestin/GRK, indicating that they preferably induce and stabilize the open receptor state that binds the G protein. On the other hand, β-arrestins are able to bind intermediate states of the opening process and thus are less sensitive to the allosteric effect of our ligands (44).

Interestingly, we observed opposite behavior on Gq efficacy of bivalent ligands depending on their ability to induce receptor dimers. Dimer-inducing compounds (at 20 to 30 Å) led to an increased efficacy, whereas the agonist with 10-Å distance had a reduced efficacy instead. This efficacy is independent of the receptor affinity of the ligand itself and depends mostly on the affinity of the adaptor protein to the receptor (45). Other works have shown that targeting GPCR homodimers with bivalent ligands often exhibit a decreased efficacy for G-protein
activation, becoming partial agonists or even antagonists (46, 47). In this regard, AG-2-BBN-2 and AG-2-BBN-3 differ from these common observations, suggesting that GRPR forced dimers may arrange in a different manner than natural homodimers, perhaps allowing positive cooperativity between two Gq adaptors. In contrast, the low efficacy of AG-2-BBN-1 is probably linked to decreased affinity of ligand-bound GRPR to Gq that is likely to result from a less favorable receptor active conformation. Furthermore, this ligand also showed reduced potency for Gq recruitment. In comparison to the efficacy, the potency depends on the affinity of the receptor for both the adaptor protein and the ligand. The 10-A˚ distance between two agonists in the polyproline scaffold is too short to recruit two receptors. Therefore, we speculate that one ligand occupies the receptor binding pocket, whereas the second ligand has an allosteric effect on the same receptor, for example, by interfering with the opening of the receptor during the activation process, thereby specifically preventing Gq recruitment but without affecting β-arrestin binding. Consequently, oligoprolines could be useful for delivering orthosteric and allosteric ligands simultaneously.

Interestingly, we observed a reduced potency of AG-2-BBN-2 toward β-arrestin-2, whereas its potency toward β-arrestin-1 is similar to AMBA. This difference could be caused by changes in potencies toward GRK2/3, resulting in different phosphorylation patterns that alter the competition of arrestins for the receptor. Even though these changes are currently relatively small, these compounds could be the basis to develop drugs that are able to switch the signaling balance of arrestin in vivo. These effects were obtained with two identical ligands, but it is possible to use two different pharmacophores to amplify these effects. For example, we observed clearly reduced efficacies when we combined an agonist with antagonist (hybrid compounds). Even though the mechanistic origin of this effect is currently not clear, it clearly shows that the future combination of different ligands can further expand the possibilities of oligoprolines.

In summary, we have shown that rigid oligoproline backbones can be used to induce artificial GPCR dimers with altered signaling properties. In addition, we have shown that the stiff linkage of a second ligand at a short distance can also alter signaling properties, possibly through allosteric effects (Fig. 6D). Together, this work identifies oligoproline backbones as an interesting tool for the development of biased drugs for GPCRs that can be useful in the pharmaceutical industry and in scientific research (e.g., to develop ligands that can stabilize GPCR dimers for structural studies). Obvious candidates for such an approach are opioid and dopamine receptors for which biased ligands are in clinical trials (6).

### Materials and Methods

**Synthesis.** The recognition motifs were synthesized by regular solid-phase peptide synthesis and inserted into the oligoproline backbone by Cu(I)-catalyzed Huisgen’s 1,3-dipolar cyclo-addition reactions (“click reaction”), as described before (31). They were purified by preparative reverse-phase high-performance liquid chromatography (HPLC) and characterized by mass spectrometry. The purity of all compounds was above 90% (30).

---

**Fig. 5.** Potency and efficacy. Potency and efficacy of tested ligands for recruitment of miniGq (A), β-arrestin-1 (B), β-arrestin-2 (C), GRK2 (D), GRK3 (E), and GRK5 (F). The asterisks indicate statistically significant differences to the reference compound AMBA. See SI Appendix, Fig. 8 for pairwise comparisons. *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001, ****P ≤ 0.0001.
Cell Culture and Creation of Stable Cell Lines. Human embryonic kidney (HEK293) cells (Merck, 85120602) or stable clones thereof were cultured in Dulbecco’s Modified Eagle Medium (DMEM) with high glucose (No. 1-26F03-I, BioConcept) supplemented with 10% fetal bovine serum and 100 U/mL penicillin/0.1 μg/mL streptomycin (Sigma-Aldrich). Cells were propagated in a humidified atmosphere at 37 °C and 5% CO₂.

Cell lines stably expressing GRPR-11S and either 114-β-arrestin-1 or 114-β-arrestin-2 were created from HEK293 cells by sequential transfection of pSi-AK1-GRPR-11S and pSi-AA-114-β-arrestin-1/-2 with Lipofectamine 3000 (Thermo Fisher Scientific) according to the manufacturer’s recommendations. Stable cell lines were obtained by culturing cells in 5 mg/mL Geneticin (G418 sulfate, US Biological Life Sciences) or/and Zeocin (100 μg/mL, Invitrogen).

Dimerization Assays. The dimerization assay was based on a NanoBRET system using pSi-AG10-GRPR-NLuc (donor) and pSi-AG10-GRPR-mCherry (acceptor). HEK293 cells were seeded in 6-well tissue culture plates at a density of 1.5 × 10⁶ cells per well 12 h prior to transfection. A total of 2 μg DNA containing 90% carrier DNA and 5% of each construct was transfected using Lipofectamine 3000. Following transfection, the assay was performed according to the same protocol as the recruitment assays with a ligand concentration of 1 μM. The measured emission wavelengths were 610 nm (mCherry) and 450 nm (NanoLuc). Then, the emission of these nonstimulated cells was measured in parallel and over the same time span as the emission of ligand-stimulated cells. The BRET ratio of nonstimulated cells (background) was subtracted from the BRET ratio of ligand-stimulated cells.

Recruitment Assays. β-arrestin recruitment was measured with cell lines expressing GRPR-11S and 114-β-arrestin-1/-2. GRK2/3/5 recruitment was measured by cotransfecting pSi-AG10-GRPR-11S and pSi-AA-GRK2-114, pSi-AA-GRK3-114, or pSi-AA-GRK5-114. These assays are based on split Nanoluc enzyme complementation (36). MiniGq recruitment was measured by a NanoBRET assay by coexpressing pSi-AG10-GRPR-NLuc as the BRET donor and venus-mGq as the BRET acceptor (37). For all recruitment assays, 8–10³ HEK293 cells were seeded in 10-cm culture dishes 12 h prior to transfection and then transiently transfected. Transfection was performed with Lipofectamine 3000 using 5 μg total DNA. For GRK recruitment, the ratio of constructs was 1:1, while for miniGq recruitment, a donor:acceptor ratio of 3:1 was chosen. One day after transfection, the cells were reseeded in white 96-well microplates with clear bottoms (PerkinElmer) at a density of 80,000 cells per well and cultured for additional 24 h. For the assay, the culture medium was removed by inverting the microplate on a paper towel and replaced by 80 μL assay buffer (20 mM Heps in DMEM [high glucose, no phenol red, BioConcept]) containing furimazine according to manufacturer’s recommendations (Promega, N2012). A removable white bottom was added to the microplate, and the baseline signal was measured in a PHERAsar FSX microplate reader (BMG Labtech) for 10 min at 37 °C. For the NanoBRET assays, emission was measured at 515 nm (Venus) and 410 nm (NanoLuc), while for the NanoLuc assay, the whole emission was measured from 100 to 1,000 nm. The ligand solutions for receptor stimulation were prepared in a separate 96-well microplate in assay buffer containing furimazine. A total of 20 μL ligand solution was added to the cells directly after baseline measurement using a 96-channel bench-top pipette (Integra, Viaflo6), and the measurement was continued for at least 30 min. Each assay was done in triplicate and repeated three or more times.

Data Availability. Excel files with recruitment data have been deposited in the Open Science Framework (https://osf.io/mh4bn/) (50). All other study data are included in the article and/or SI Appendix.
ACKNOWLEDGMENTS. This work was supported by a Sinergia grant from the Swiss National Science Foundation (CRSII2_160805) to M.B., X.D., H.W., and P.B., and a National Centre of Competence in Research (NCCR) Molecular Systems Engineering (S1NFP0-182899) grant to G.F.X.S. We thank Dr. Nevin Lambert for providing plasmids, Dr. Gregor Ciccletti for critical reading of the manuscript, and Daniela Buttker for help with data analysis and figure preparation.