G Protein–Coupled Receptor Oligomerization Revisited: Functional and Pharmacological Perspectives

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Abstract—Most evidence indicates that, as for family C G protein–coupled receptors (GPCRs), family A GPCRs form homo- and heteromers. Homodimers seem to be a predominant species, with potential dynamic formation of higher-order oligomers, particularly tetramers. Although monomeric GPCRs can activate G proteins, the pentameric structure constituted by one GPCR homodimer and one heterotrimeric

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G protein may provide a main functional unit, and oligomeric entities can be viewed as multiples of dimers. It still needs to be resolved if GPCR heteromers are preferentially heterodimers or if they are mostly constituted by heteromers of homodimers. Allosteric mechanisms determine a multiplicity of possible unique pharmacological properties of GPCR homomers and heteromers. Some general mechanisms seem to apply, particularly at the level of ligand-binding properties. In the frame of the dimer-cooperativity model, the two-state dimer model provides the most practical method to analyze ligand–GPCR interactions when considering receptor homomers. In addition to ligand-binding properties, unique properties for each GPCR oligomer emerge in relation to different intrinsic efficacy of ligands for different signaling pathways (functional selectivity). This gives a rationale for the use of GPCR oligomers, and particularly heteromers, as novel targets for drug development. Herein, we review the functional and pharmacological properties of GPCR oligomers and provide some guidelines for the application of discrete direct screening and high-throughput screening approaches to the discovery of receptor-heteromer selective compounds.

I. Morphologic Aspects of G Protein–Coupled Receptor Oligomerization

A. The Search for the Predominant Oligomeric G Protein–Coupled Receptor Species

Although G protein–coupled receptors (GPCRs) were initially thought to be, and function exclusively as, monomeric entities, evidence accumulated over the past two decades indicates that they can form homomers and heteromers in intact cells (Bouvier, 2001; Milligan and Bouvier, 2005; Pin et al., 2007; Ferré et al., 2009). It is now well accepted that family C GPCRs (e.g., metabotropic glutamate, calcium-sensing receptors, GABA_B, and sweet and umami taste receptors) form constitutive homo- or heteromers (Kniazeff et al., 2011). Such observations raised a long debated question about whether family A (rhodopsin-like) GPCR dimers were also constitutive and required for G protein activation. A clear demonstration that this is not the case came from studies in which monomeric entities were trapped into nanodiscs. In such experiments, it was demonstrated that \( \beta_2 \)-adrenergic, rhodopsin, and \( \mu \)-opioid receptors function effectively as monomers (Bayburt et al., 2007; Whorton et al., 2007, 2008; Kuszak et al., 2009). Also, monomeric rhodopsin in solution activated its G protein transducin at the diffusion limit (Ernst et al., 2007). Further support for the functionality of monomeric GPCR units came when isolated monomeric seven transmembrane (7TM) domains of family C GPCRs were shown to be capable of fully activating G protein when directly activated by synthetic small molecules (El Moustaine et al., 2012). Finally, a 1:1 stoichiometry has also been found to be sufficient for rhodopsin–\( \beta \)-arrestin binding (Tsukamoto et al., 2010; Bayburt et al., 2011). But those observations did not exclude that class A GPCR oligomers can spontaneously form in living cells, as it is well demonstrated with family C GPCRs, and raised the question of their functional significance. In fact, the three GPCRs shown to be functional as strict monomers were shown also to exist as dimers or higher-order oligomers in living cells (see below). One of the research groups that demonstrated the functionality of GPCR monomers in nanodiscs found evidence for the existence of stable \( \beta_2 \) receptor oligomers, mostly tetramers, after reconstitution into phospholipid vesicles (Fung et al., 2009). Also, rhodopsin–\( \beta \)-arrestin binding stoichiometry in isolated rod outer segment membranes was found to depend on the percentage of activated receptors, increasing from 1:1 to 2:1 (Sommer et al., 2012).

An explosion of data supporting the existence of homo- and heteromers of GPCRs in intact cells (stored in the GPCR Oligomerization Knowledge Base; http://www.gpcr-okb.org; Khelashvili et al., 2010) came with the widespread use of biophysical techniques, such as resonance energy transfer (bioluminescence and...
fluorescence resonance energy transfer (BRET and
FRET, respectively), fluorescence complementation, or combination of these techniques (Milligan and Bouvier,
2005; Gandía et al., 2008; Pin et al., 2007, 2009; Bacart et al., 2008; Carriba et al., 2008; Guo et al., 2008;
Cabello et al., 2009; Ferré et al., 2009; Urizar et al.,
2011). These techniques, however, have largely fallen
short in answering questions about the size of the oligomer complexes and their possible dynamic nature.
Initial evidence for GPCR oligomerization in a physiologically relevant system came from atomic force
microscopy experiments in native disk membranes from mice, which showed rhodopsin to be arranged in
paracrystalline arrays of dimers (Fotiadis et al., 2003).
These studies raised severe criticism (Chabre and le
Maire, 2005), and more recent studies using single-
molecule techniques have begun to address the details
of the spatial and temporal organization of GPCR
complexes in living cells by directly observing the state
and behavior of individual proteins in the cell. Single-
molecule total internal reflectance fluorescence micros-
copy (TIR-FM) was first used to track the position of
individual molecules of muscarinic acetylcholine M_1
and N-formyl peptide receptors using fluorescently
labeled ligands (Hern et al., 2010; Kasai et al., 2011).
Those ligands had such slow off-rates that they may be
considered essentially irreversibly bound. Both studies
suggested a transient (second-scale) formation and
dissociation of dimers, with 30–40% proportion of
dimers at any given time. These conclusions require
that the fluorescent ligands bind with similar affinity
to both protomers in the dimer (see below for
discussion of ligand cooperativity within receptor
dimers/oligomers; see Section II.B), and it is possible
that the fluorescent ligand might stabilize the receptor
in a nonrepresentative conformational state. Despite
these concerns, similar studies have reached equiva-
 lent conclusions on the state of the muscarinic
acetylcholine M_2 receptor in cardiac muscle (Nena-
sheva et al., 2013). More recently, TIR-FM was used
together with SNAP-tag technology to directly label
cell-surface GPCRs with organic fluorophores to dy-
namically monitor individual \( \beta_1 \)- and \( \beta_2 \)-adrenoceptors
as well as GABA_\text{B} receptors on the surface of living,
transiently transfected cells (Calebiro et al., 2013).
This study showed that all three receptors form dimers
and higher-order oligomers (also with estimations of
dynamic second-scale receptor–receptor interactions)
in a proportion dependent on the subtype of receptor
and on the receptor density. At low densities, mono-
meric species were predominant for the \( \beta_1 \)-adrenoceptor,
whereas \( \beta_2 \)-adrenoceptors displayed a higher
proportion of dimers. A step-wise appearance of first
dimers and successively trimers, tetramers, and
upwards to higher-order complexes appeared with progressive increase in receptor density. At densities
comparable to receptor expression in native tissue,
dimers and higher-order oligomers were the predom-
inant species for both receptors, and agonists did not
modify the oligomerization status (Calebiro et al.,
2013). As expected (Maurel et al., 2008; Comps-Agrar
et al., 2011), GABA_\text{B} receptors showed mostly dimeric
and tetrameric species at lower densities, but higher-
order species also appeared with increased expression
levels (Calebiro et al., 2013).

Fluorescence correlation spectroscopy (FCS) is an-
other indirect but useful technique to determine the
oligomer status of protein clusters. It measures the
molecular brightness of a fluorescence-tagged protein,
which provides an estimate of the number of fluorescent
molecules by recording fluctuations in fluorescence
intensity arising from individual fluorescent molecules
(Chen et al., 2003). FCS has been used to study \( \mu \)-opioid
receptor homo- and heteromerization with \( \delta \)-opioid
receptors, and number and brightness measurements
suggested that \( \mu \)-opioid receptors exist primarily as
dimers that oligomerize with \( \delta \)-opioid receptors into
tetramers (Golebiowska et al., 2011). The recent FCS
with a particle counting histogram approach by Herrick-
Davis et al. (2013) also provides support for homodimers
being the predominant, and perhaps only, species for
several GPCRs, including \( \alpha_{1B} \)-adrenoceptor, \( \beta_2 \)-adreno-
ceptor, serotonin 5-HT\text{_{2A}} and 5-HT\text{_{2c}}, muscarinic ace-
tylcholine M_1 and M_2, and dopamine D_1 receptors. As
with the results obtained with TIR-FM, the homodimer
configuration was unaltered by agonist treatment
(Calebiro et al., 2013; Herrick-Davis et al., 2013). Unlike
the fast dynamic behavior of GPCR monomers/dimers
shown in TIR-FM experiments (Hern et al., 2010; Kasai
et al., 2011; Calebiro et al., 2013), however, the dimers
were stable over a 10-fold range of receptor expression
levels (Herrick-Davis et al., 2013). This is also in line
with other findings that suggest much more stable
interactions, such as higher-order dopamine D_3 receptor
oligomers, over a high range of receptor expression
(Guo et al., 2008) and stable \( \beta_2 \)-adrenoceptor tetramers in
phospholipid vesicles (Fung et al., 2009). Stability,
at least over a substantial time period, of GPCR dimers/
oligomers is also inferred from experiments that in-
dicate these complexes are generated at an early stage
of biosynthesis. In fact, it has been suggested that early-
stage dimerization may be required for effective folding
and maturation of the receptor (Salahpour et al., 2004;
Bulenger et al., 2005; Milligan, 2010, 2013). The same
reasoning has been used based on evidence for cointern-
alization, with the capacity of a selective ligand of one of
the receptors in a GPCR heteromer to cointernalize the
two different receptors constituting the heteromer
(Hillion et al., 2002; Milligan, 2010, 2013; Ward et al.,
2011; Tadagaki et al., 2012). Negative results should
also be acknowledged, such as the recent study by
Gavalas et al. (2013), which measured the recruitment
of subsets of histidine-tagged GPCR protomers into
artificial microdomains (containing immobilized metal
affinity chromatography or streptavidin beads) on the surface of living cells and determines the simultaneous corecruitment of untagged protomers. The study only showed evidence for corecruitment of metabotropic glutamate receptor protomers, but not β2-adrenergic or μ-opioid protomers, into such artificial microdomains. Apart from differences between different receptors or when considering receptor homomers or heteromers, it might be that a number of the approaches used to study oligomerization are unable to resolve fluctuations in receptor interactions occurring on a second or subsecond scale. On the other hand, mammalian transfected cells might be lacking elements that increase the stability of the interfaces that determine the receptor–receptor interactions, which might also be different depending on the cellular compartment studied.

If different interfaces are involved in receptor oligomerization, we could expect dimers to be predominant species, determined by the most stable interaction, followed by tetramers (dimers of dimers), determined by a weaker interaction. Recently calculated estimates of the relative stability of different, putative dimeric interfaces of different GPCR subtypes using extended biased molecular dynamics simulations in explicit lipid-water environments argue in favor of a variable strength of association depending on the specific residue composition or shape of the interface, despite an overall transiency in receptor–receptor interactions (Johnston et al., 2012). Notably, simulations of β1- and β2-adrenoceptors suggested a model of oligomerization in which more stable homodimers involving TM1 diffuse through the membrane and transiently interact with other protomers/dimers via other TM helices (e.g., TM4). In agreement with these predictions are the results obtained in a recent study with the muscarinic acetylcholine M3 receptor, using quantitative FRET spectrometry techniques with controlled expression of the energy donor-tagged species (Patowary et al., 2013). Mathematical analysis of the FRET efficiencies obtained from spectral unmixing was compatible with the M3 receptor existing as stable dimeric complexes, a large fraction of which interacted dynamically to form tetramers that were specifically within a rhombic organization rather than a square or linear configuration (Patowary et al., 2013).

B. The Search for Oligomer Interfaces

A very intriguing set of inferences in the field of GPCR oligomerization has been based on recently obtained high-resolution crystallographic structures, including those of the chemokine CXCR4, the μ-opioid and κ-opioid receptors, the β1-adrenoceptor, and the smoothened receptor (Wu et al., 2010, 2012; Manglik et al., 2012; Huang et al., 2013; Wang et al., 2013). Each of these crystallized as parallel dimers and/or tetramers. Although some of the observed interfaces may reflect artifacts of crystal packing and the conditions employed for crystallization, a tentative first suggestion from these studies is the existence of different dimer interfaces for different GPCR homodimers. Nevertheless, some TM domains have been observed more often than others. TM5 and TM6 residues constituted the main interfaces for chemokine CXCR4 and μ-opioid receptor crystallized dimers, although there were marked differences between them, with only a few contacts between specific residues for chemokine CXCR4 and extensive contacts throughout the length of these TM helices in μ-opioid receptor dimers (Wu et al., 2010; Manglik et al., 2012). Involvement of TM6 was also suggested early on for β2-adrenoceptor dimers (Hebert et al., 1996) and for the leukotriene receptor BLT1 (Baneres and Parello, 2003) by the use of interfering synthetic peptides with the same sequence as TM6. Also, by using cysteine cross-linking techniques, TM5 was suggested to be involved in homodimerization of dopamine D2, muscarinic M3, and serotonin 5-HT2C receptors (Guo et al., 2005; Mancia et al., 2008; Hu et al., 2012).

Apart from the TM5-TM6 interface, crystallized chemokine CXCR4 dimers also showed contacts at the intracellular ends of TM3 and TM4 (Wu et al., 2010), and μ-opioid dimers also showed a second, less prominent symmetric interface, involving TM1, TM2, and, also, helix 8 (H8; the helix adjacent to TM7 running along the internal membrane surface) (Manglik et al., 2012). A TM1–TM2–H8 interface (with slightly different contact residues) was also found in crystals of κ-opioid receptor dimers (Wu et al., 2012), rhodopsin (Ruprecht et al., 2004; Salom et al., 2006), opsin (ligand-free rhodopsin; Park et al., 2008), and β1-adrenoceptor (Huang et al., 2013). An additional interface involving TM4 and TM5 was also obtained for the β1-adrenoceptor and smoothened receptor (Wang et al., 2013). Notably, the two crystallographic interfaces of the β1-adrenoceptor (TM1–TM2–H8 and TM4–TM5) were suggested to be physiologically relevant with cysteine-cross-linking experiments (Huang et al., 2013). A model of the potential rhodopsin dimer was initially built by docking the rhodopsin crystal structure into the unit cell constraints determined by atomic force microscopy. The model included a primary interface between protomers involving TM4 and TM5, which implied asymmetric binding of one heterotrimeric G protein to each protomer (Liang et al., 2003; Fotiadis et al., 2004). The model also suggested a secondary interface holding rows of dimers together, which was later shown to involve TM1, TM2, and H8 (Salom et al., 2006). The possible involvement of a TM1–TM2–H8 interface in rhodopsin quaternary organization was recently supported by cross-linking experiments of endogenous cysteines of rhodopsin in disk membranes (Knepp et al., 2012).

It is important to remember that crystal structures are not only the result of specific crystallization conditions, but they correspond to static, ligand-specific conformational states of receptors stripped of their
natural lipid environment. For GPCRs, the majority of crystal structures that are currently available refer to antagonist-bound (inactive) structures. The inferred dimeric interfaces may, therefore, depend on those specific conformational states. As suggested by Huang et al. (2013), the comparison of the differences in the interfaces observed from the crystallized structures of the antagonist-bound μ-opioid and chemokine CXCR4 receptors and the ligand-free β_1-adrenoceptor might suggest that the TM5 interface can partner in the interaction with TM4 or TM6, depending on the conformation of the receptor. As mentioned by Manglik et al. (2012), the TM5-TM6 interface inferred by the crystal structure of μ-opioid dimers could preclude either protomer from properly coupling to G protein, because the agonist-induced receptor–G protein interaction depends on rearrangements of TM5 and TM6 within the seven-helical domain bundle. Therefore, it is important to assess if, by stabilizing different receptor conformations, different ligands also promote different dimeric interfaces. By cross-linking substituted cysteines, Guo et al. (2005) suggested TM4 to be a main interface in the dopamine D_2 receptor oligomer. Cross-linking of a different set of cysteines in TM4 was slowed by inverse agonists and accelerated by agonists. In fact, cross-linking of the latter set of cysteines locked the receptor in an active state, strongly suggesting that a conformational change at the TM4 dimer interface is part of dopamine D_2 receptor activation (Guo et al., 2005). Mancia et al. (2008), also using the cysteine cross-linking approach, found two interfaces in serotonin 5-HT_2C receptor dimers involving TM4-TM5 and TM1, while TM4–TM5 being selectively sensitive to receptor activation. In summary, although a pattern of similar interfaces of GPCR homomers seems to be emerging, different interfaces can be found in different oligomers and even in different conformations of the same oligomer. Although agonists did not modify the dynamics of β_2-adrenoceptor oligomerization in TIR-FM experiments (Calebiro et al., 2013) or the stability of β_2-adrenoceptor oligomers in FCS–photon-counting histogram experiments (Herrick-Davis et al., 2013) or in FRET experiments in receptors into reconstituted phospholipid vesicles, an inverse agonist did modify the stability of β_2-adrenoceptor oligomers in this latter preparation (Fung et al., 2009). Thus, selected ligands (which can stabilize specific conformations of the receptor) may still modify GPCR oligomeric interfaces and, therefore, the dynamics of receptor oligomerization.

C. Determinants of G Protein–Coupled Receptor Heteromerization

When attempting to understand the predominant oligomeric species within receptor heteromers, an additional issue is whether the complexes exist as heterodimers or as hetero-oligomers generated from individual homodimers. The lack of detection of trimers in situations in which interconversion between dimers and tetramers of the muscarinic M_3 receptor was observed (Patowary et al., 2013) is consistent with a dimer + dimer model and with the organization of class C GABA_B dimers and tetramers (Comps-Agrar et al., 2012). Some indirect biochemical data also support this model for class A receptors. These include cooperative binding of certain ligands to adenosine A_2A and A_1 receptor heteromers (see below and Orru et al., 2011a). Furthermore, oligomerization of more than two different GPCRs has been suggested from studies with sequential BRET-FRET, BRET plus bimolecular fluorescence complementation (Carriba et al., 2008; Cabello et al., 2009; Navarro et al., 2010). Tentatively, these apparent heterotrimers could represent heteromultimers of homomers, as suggested for adenosine A_2A–dopamine D_2–cannabinoid CB_1 receptor heteromers (Navarro et al., 2010).

Apart from interactions between TM domains, several studies have provided evidence for disulfide bridges between extracellular domains of class C GPCR homomers (Kniazeff et al., 2011) and for a key role of electrostatic interactions between intracellular receptor domains in receptor heteromerization (Woods and Ferré, 2005). These electrostatic interactions depend on the very asymmetric and disordered structure of intracellular domains and have been suggested to be involved in several receptor heteromers (Ciruela et al., 2004; Woods and Ferré, 2005; Navarro et al., 2010; O'Dowd 2012, 2013). Several general features of the regions involved in these interactions have been outlined: one region, in one of the receptors, contains a series of adjacent arginine residues, and the other region, in the other receptor, contains acidic residues, with several adjacent residues or at least one phosphorylated serine. Once established, the polyarginine–phosphate electrostatic interaction possesses a strong stability. Thus, these bonds can withstand fragmentation by mass spectrometric collision-induced dissociation at energies similar to those that fragment covalent bonds, and they demonstrate an extremely low dissociation constant by surface plasmon resonance (Woods and Ferré, 2005). If electrostatic interactions between intracellular domains are more predominant in receptor heteromers (not reported yet in receptor homomers) and are stronger than those provided by TM interfaces, we could speculate that some receptor heteromers could be more stable than receptor homomers. An example of a functionally relevant electrostatic interaction is the one involved in adenosine A_2A–dopamine D_2 receptor heteromerization, with the arginine-rich domain localized in the N-terminal portion of the long third intracellular loop of the dopamine D_2 receptor and the acidic domain in the distal part of the long C terminus of the adenosine A_2A receptor (Ciruela et al., 2004; Navarro et al., 2010). Mutation- or peptide-mediated disruption of the A_2A–D_2 receptor electrostatic interaction
produces a profound destabilization of the quaternary structure of the heteromer (Navarro et al., 2010) with disappearance of significantly relevant A2A receptor interactions in brain tissue, such as the adenosine A2A receptor–mediated inhibition of dopamine D2 receptor–induced depression of striatal neuronal firing (Azdad et al., 2009).

In summary, most evidence indicates that, as for family C GPCRs, which can be found as strict homodimers, heterodimers (Doumazane et al., 2011), as well as stable heteromers (Comps-Agar et al., 2012), family A GPCRs form homo- and heteromers in heterologous systems. Homodimers seem to be a predominant species with potential dynamic formation of higher-order oligomers, particularly tetramers. It still needs to be resolved if class A GPCR heteromers are preferentially heterodimers or if they are mostly constituted by heteromers of homodimers. In the next sections, we will review the evidence that GPCR homo- and heteromers constitute functional and pharmacologic units.

II. Functional Aspects of G Protein–Coupled Receptor Oligomerization

A. Principles of Allosterism and Properties of Allosteric Modulators

Allosterism can be defined as the process by which the interaction of a chemical or protein at one location on a protein or macromolecular complex (the allosteric site) influences the binding or function of the same or another chemical or protein at a topographically distinct site (Christopoulos and Kenakin, 2002). This definition provides a framework to understand the biochemical properties of GPCR homomers and heteromers. As suggested by Kenakin and Miller (2010), it is useful to define allosterism in terms of three interacting species: the “modulator,” a ligand or protein that binds to the “conduit” (usually a protein; the GPCR protomer or oligomer in this review), which transmits the thermodynamic allosteric energy to the “guest,” the target of the allosteric modulation. With GPCRs, we can then consider three different kinds of allosterism depending on the location of the target of the allosteric modulation. If the target of the allosteric modulation is another ligand cobinding with the allosteric modulator, this is referred to as classic allostery. If the target of the allosteric modulation resides in the cytosol, it may be called cytosolic allosterism. Finally, if the target of the allosteric modulation interacts with the conduit of the allosteric modulation along the plane of the membrane, this is referred to as lateral allosterism, with a main example being allosterism in receptor oligomers (Kenakin and Miller, 2010).

An example of classic allosterism would be the case of a ligand that modulates allosterically the effect of an orthosteric agonist. An orthosteric agonist has two main independent properties: an affinity for the receptor and an intrinsic efficacy, which determines the power of the agonist to induce a functional response. The allosteric modulator can have different and independent effects on these properties of distinct orthosteric agonists (Kenakin and Miller, 2010; Smith and Milligan, 2010). Therefore, there may be two general effects of allosteric modulators on orthosteric ligands: on their affinity and on intrinsic efficacy. Analysis of functional responses will determine the contribution of both allosteric effects. Another property of allosteric modulators is saturability. A negative allosteric modulator that selectively modifies the affinity of an orthosteric antagonist, such as a “right shift” would continue ad infinitum. This ceiling effect of classic allosteric modulators can have important therapeutic implications by reducing overdose effects, compared with orthosteric ligands (Kenakin and Miller, 2010; Smith and Milligan, 2010).

In cytosolic allosterism (as defined by Kenakin and Miller, 2010), modulated proteins are cytosolic signaling proteins, such as G proteins, GPCR kinases (GRKs), and β-arrestins. Importantly, this type of allosterism can lead to functional selectivity, i.e., the ability of a ligand to selectively promote a specific cellular signaling event (Reiter et al., 2012). The rationale is that different ligands can stabilize different conformations of the receptor. If different cytosolic proteins that mediate different signaling interact with different residues or phosphorylated residues of the GPCR, it may be expected that some conformations can favor or impair the binding of a particular signaling protein or that they can induce a different conformation of the cytosolic protein, leading to biased agonism or biased antagonism (Reiter et al., 2012). This can also have important therapeutic implications, i.e., when a particular signaling pathway or end point is associated with a therapeutic response, whereas another is associated with unwanted or side effects. Agonist binding to GPCRs and G protein activation are rapidly followed by several coordinated events common to most GPCRs. These include recruitment of GRKs that phosphorylate the receptor at multiple intracellular sites, followed by the recruitment of β-arrestins, which trigger receptor endocytosis. But, in addition to canonical G protein–mediated signaling, GPCRs can also bind to other cytosolic adaptors, including β-arrestins, which elicit G protein–independent signaling through activation of mitogen-activated protein kinase (MAPK) and Akt. Most known endogenous and exogenous ligands can signal through both signaling mechanisms. There are already, however, several examples of biased ligands that preferentially signal through β-arrestins over G proteins. In some cases, this may be a therapeutically beneficial effect, in others an unwanted one. Carvedilol, for instance, has been reported to be a β-arrestin–biased ligand acting at β2-adrenoceptors (but
also at β1-adrenoceptors), which may contribute to its clinical value in heart failure beyond its β-blocker property (Reiter et al., 2012). The work by Lefkowitz and coworkers indicates that a mechanism behind functional selectivity can be a ligand-dependent “phosphorylation barcoding” (concept developed by Tobin et al., 2008). Thus, the β-arrestin–biased agonism of carvedilol appears to depend on its ability to impair GRK2- and promote GRK6-mediated phosphorylation (Nobles et al., 2011). Another mechanism is the differential structural conformations that lead to a preferential binding for specific G proteins. For instance, different agonists of the cannabinoid CB1 receptor differentially regulate the binding of the three homologous G1 proteins to the receptor. Whereas one ligand (desacetylevonantradol) behaved as an agonist for G11 and G12 and an inverse agonist for G13, another ligand [(R)-methanandamide] behaved in exactly the opposite manner, as an inverse agonist for G11 and G12 and an agonist for G13 (Mukhopadhyay and Howlett, 2005).

**B. Allosterism in Receptor Oligomers: Modulation of Ligand Affinity**

Evidence for the existence of GPCR homomers (and heteromers) could already be deduced some time ago from analysis of radioligand-binding experiments, with the evidence of complex binding, such as upward concave nonlinear Scatchard plots in saturation experiments and as biphasic curves in agonist–antagonist competitive-inhibition experiments. Such complex curves were usually explained by two nonoverlapping models (reviewed in Casadó et al., 2007). First is the monomer–G protein model, which considers receptors as monomers with two independent populations not in equilibrium. One population would be bound and the other would not be bound to G proteins, with the receptors coupled to G proteins having higher affinity for the agonist. Receptors with high affinity could then be converted into low affinity with the addition of GTP, because it would uncouple the G protein from the receptor (De Lean et al., 1980). To explain the complex radioligand binding curves, this first mechanism would assume a preexisting proportion of both populations of receptors and, therefore, a limited pool of G proteins. The second model, the dimer-cooperativity model, considers oligomerization, or at least GPCR dimers. In this case, allosteric communication through the two protomers allows negative cooperativity, meaning that the binding of a ligand to the first protomer decreases the affinity of the ligand for the second protomer. Cooperativity (positive or negative) is a particular type of allosteric modulation in receptor oligomers, where the protomers of a homodimer are the conduit of the allosteric modulation and the same ligand is the allosteric modulator (binding to the first protomer) and the modulated target (binding to the second protomer). This mechanism does not assume a limited pool of G proteins, which is always present and acts as an additional allosteric modulator that increases the affinity of the agonist, and provides the conformation of the dimer that allows negative cooperativity of a ligand through the protomers.

In apparent support for the first mechanism, in the study by Whorton et al. (2007) with reconstitution of monomers of β2-adrenoceptors and G proteins in high-lipoprotein nanoparticles, antagonist/agonist competitive inhibition curves showed biphasic curves, and the addition of a GTP analog converted the low affinity population into high affinity. The study also showed, however, that the proportion of receptors with high affinity state increases up to 100% by increasing the G protein pool, indicating that in situ, in membrane preparations of brain tissue, the detection of two populations of receptors would indicate a limited pool of G proteins. In this case, the monomeric model with two G protein–dependent populations of receptors could explain the upward concave Scatchard plots and the biphasic antagonist/agonist competition curves (Casadó et al., 2007). The upward concave Scatchard curves would be the result of the addition of two curves that describe the two independent populations, with high and low affinities for the agonist. The biphasic competition curves would be the result of two different affinities of the agonist, which would identify both populations, displacing the antagonist that would have the same affinity for both populations. However, some early studies indicated that antagonist binding could also be modulated by guanine nucleotides, which, according to the monomer–G protein model, would suggest that antagonist binding would also depend on G protein coupling (Burgisser et al., 1982; De Lean et al., 1982; Klotz et al., 1990). Dissociation kinetic experiments can resolve the enigma. Investigation of the dissociation kinetics of a tracer ligand in the absence and presence of a second ligand represents a sensitive method to detect cooperative interactions between two topographically distinct binding sites. Thus, ligands that compete for the same site on a monomeric receptor should not influence one another’s dissociation kinetics. In contrast, allosteric modulation between two simultaneously bound and interacting sites, either within a receptor monomer or across a receptor dimer or oligomer, should alter ligand dissociation (May et al., 2007). This analysis has been used to demonstrate homomerization of several GPCRs (Urizar et al., 2005; Albizu et al., 2006, 2010; Springael et al., 2006; May et al., 2011). Therefore, at present, there is no doubt that negative cooperativity is an allosteric property of some ligands that bind to GPCR homomers.

It could be argued that the dimer-cooperativity model is an artifact of membrane preparations, where there would not be a limited pool of G proteins for the specific receptor under study compared with the in situ situation. Negative cooperativity has not only been
observed in membrane preparations from different tissues, including brain, and from artificial systems, such as transfected cells, but by measuring dissociation kinetics with fluorescent GPCR ligands, negative cooperativity can also be demonstrated in living cells (May et al., 2011), indicating that it is not an artifact of membrane preparation and that it can be of functional and pharmacological significance. Finally, a significant study that strongly supports the dimer-cooperativity model not only in living cells, but also in native tissues, involved homogeneous time-resolved FRET with fluorescent ligands (Albizu et al., 2010). Homogeneous time-resolved FRET is based on an energy transfer between a lanthanide (europium or terbium) and a compatible fluorophore (lanthanide’s long-lasting fluorescence allows measurement of its fluorescence after a laser pulse at time points when background fluorescence has disappeared, greatly increasing signal-to-noise ratio) (Gandia et al., 2008). Having ligands for oxytocin receptors separately fused to donor and acceptor molecules demonstrated the existence of dimers, both in transfected cells and in native tissue (mammary glands; Albizu et al., 2010). Furthermore, the differential FRET obtained with labeled agonists and antagonists, with significantly less FRET obtained with agonists, indicated a stoichiometry of one agonist molecule per dimer consistent with negative cooperativity, and this was only observed with agonists and not antagonists (Albizu et al., 2010).

Results from radioligand binding experiments, therefore, can indicate the existence of GPCR oligomerization. Furthermore, apart from negative cooperativity, the monomeric–G protein model fails to explain other complex radioligand-binding data, such as downward concave Scatchard plots, as reported with the mixed vasopressin-oxytocin receptor antagonist OTA (d(CH2)5[Tyr(Me)2-Thr4-Orn8-Tyr(NH2)9]-oxytocin) (Albizu et al., 2006, 2010). A dimer-cooperativity model could explain these findings by the existence of positive cooperativity (Albizu et al., 2006, 2010). Other results of radioligand-binding experiments that cannot be explained by the monomer–G protein model are biphasic antagonist/antagonist competition curves—for instance, those recently described for the adenosine A2A receptor antagonists ZM-241385 [4-(2-[7-amino-2-(2-furyl)[1,2,4]triazolo[2,3-a][1,3,5]triazin-5-ylamino]ethyl)phenol] and SCH-442416 [2-(2-furanyl)-7-[3-(4-methoxyphenyl)propyl]-7H-pyrazolo[4,3-e][1,2,4]triazolo[1,5-c]pyrimidin-5-amine] as radioligand and competing compounds, respectively (Orru et al., 2011a). These complex competitive-inhibition curves could be explained by means of the dimer-cooperativity model. In this case, we have two possibilities: first, allosteric interactions with the same ligand, i.e., negative cooperativity, where the binding of one ligand, SCH-442416, to the first protomer modifies the affinity of the binding of the same ligand to the second protomer; or second, allosteric interactions between different ligands in the receptor homomer, where the binding of one ligand, i.e., ZM-241385, to the first protomer changes the affinity of the other ligand, SCH-442416, to the second protomer. This implies different types of allosterism, and their differentiation could have important therapeutic implications (see below and Fig. 1). Negative cooperativity has also been reported for some serotonin 5-HT2A receptor antagonists, like the atypical antipsychotics clozapine and risperidone, but not the typical antipsychotic haloperidol (Brea et al., 2009). Importantly, there was a correlation between the binding behavior of the different antipsychotics and their properties as antagonists of a serotonin 5-HT2A receptor-mediated signaling. Thus, negative cooperativity could explain a biphasic inhibition of serotonin-stimulated arachidonic acid release (Brea et al., 2009).

What would be the functional significance of negative (or positive) cooperativity of a receptor homomer? In a framework of symmetric signaling properties of both protomers in a receptor homodimer, we would expect two different levels of ligand-mediated signaling that would depend on the concentration of the ligand. Negative cooperativity could provide a mechanism that protects the biologic system against acute elevations of the endogenous ligand (Agnati et al., 2005). Positive cooperativity, on the other hand, could provide an amplificatory mechanism, although to our knowledge, no clear examples of positive cooperativity of endogenous ligands have been reported (but see below about symmetry).

Regarding allosterism in receptor oligomers, and according to Kenakin and Miller (2010), two different scenarios have to be considered. First, the receptor dimer becomes the new conduit of the allosteric modulation, the allosteric modulator binds to one of the protomers and the modulated target binds to the other protomer. When the allosteric modulator and the target of the allosteric modulation are the same ligand and the protomers are identical (GPCR homomers), the result of this allosterism is positive or negative cooperativity. When the allosteric modulator and the target of the allosteric modulation are different and the protomers identical, the result is allosteric interactions between different ligands in the receptor homomer. The same can apply, with different ligands and different protomers, in receptor heteromers. As an example, adenosine A2A receptor ligands modulate the affinity of dopamine D2 receptor ligands in the A2A–D2 receptor heteromer. This is a well known receptor heteromer localized in one of the two main neuronal populations in the striatum and suggested to be a target for the treatment of Parkinson’s disease (Ferré et al., 2008; Azdad et al., 2009). In the case of the δ-μ-opioid receptor heteromer, it was shown that binding and signaling by morphine or μ receptor agonists...
were potentiated by δ-opioid receptor antagonists, and reciprocally, binding and signaling by δ-opioid receptor agonists were potentiated by μ receptor selective antagonists (Gomes et al., 2004, 2011). Studies carried out with the δ-opioid–cannabinoid CB1 receptor heteromer have also revealed allosteric modulations of cannabinoid CB1 receptor ligands on δ-opioid receptor ligand binding properties (Bushlin et al., 2012; Rozenfeld et al., 2012). Both in recombinant systems expressing both receptors and endogenous tissues, binding and consequently signaling by δ opioid receptor could be potentiated by a low, nonsignaling dose of cannabinoid CB1 receptor agonist or a selective antagonist (Bushlin et al., 2012). These unique properties, taken with the fact that the δ-opioid–CB1 receptor heteromers are upregulated during neuropathic pain (Bushlin et al., 2012), make this receptor heteromer an attractive target for the treatment of this disease.

In the second scenario of allostery in receptor oligomers (Kenakin and Miller, 2010), one of the protomers becomes the allosteric modulator, which is sometimes referred in the literature as ligand-independent allosteric modulation in the receptor heteromer. As we will see later, dopamine D2 receptor selectively modifies the binding of SCH-442416 to the adenosine A2A receptor in the A2A–D2 receptor heteromer (Orru et al., 2011a). The melatonin MT1-GPR50 receptor heteromer constitutes a particular example of this allosteric modulation, where the presence of the orphan GPR50 receptor has a negative allosteric effect on melatonin binding to the MT1 receptor (Levoye et al., 2006a).

### C. Allostery in Receptor Oligomers: The Two-State Dimer Model

In view of the existence of clear experimental evidence supporting the dimer-cooperativity model, there is a need to develop new models of analysis of radioligand-binding experiments that consider GPCRs as oligomers or, at least, GPCR dimers. From all the models that consider receptors as monomers, the most commonly used is the two-independent-site model, which can explain the upward concave Scatchard plots and the biphasic competition curves. This is assuming the existence of two independent interconvertible populations of receptors with two different affinities for the agonists. This model, however, has serious drawbacks. When trying to resolve complex radioligand-binding data, when using the two-independent-site model, the values of the equilibrium dissociation constants and number of receptors obtained vary significantly depending on the concentration of the radioligand (Casadó et al., 2009a), indicating a lack of robustness.

For the analysis of radioligand-binding experiments, several models that consider receptors as dimers/oligomers have been developed (Durroux, 2005; Casadó et al., 2007; Rovira et al., 2009; Giraldo, 2013). These models, however, involve quite complex initial mechanistic equations, with a high number of constants, which make them quite unpractical for the analysis of radioligand binding experiments. Nevertheless, the further
development of empirical equations from one of these models, the two-state dimer model, makes it particularly insightful and easy to use (Franco et al., 2005, 2006; Casadó et al., 2007, 2009a,b). This is not entirely obvious when looking at the initial equations that include all the microscopic equilibrium isomerization and dissociation constants: seven for the binding of one ligand for saturation experiments, and 11 for two ligands for competition experiments (described in detail in Casadó et al., 2007). Hence empirical equations were derived that disclose a lower number of constants, macroscopic equilibrium dissociation constants, and this allows for a practical analysis of radioligand-binding data when considering the receptors as dimers (analyzed in Casadó et al., 2007).

Equation for saturation experiments:

\[ A_{\text{bound}} = \frac{(K_{D2}A + 2A^2)}{K_{D1}K_{D2} + K_{D2}A + A^2} \]

\( A \) is the concentration of radioligand and \( R_{T} \) is the total amount of receptor dimers (the traditional \( B_{\text{max}} \) from the two-independent-site model would be twice this value). \( K_{D1} \) and \( K_{D2} \) are the macroscopic equilibrium dissociation constants, which define the dissociation equilibria involved in the binding of a ligand to the receptor dimer as a whole (not to be confused with the intrinsic equilibrium dissociation constants) (Fig. 1). \( K_{D1} \) represents the equilibrium between free ligand, empty dimer, and semi-occupied dimer (the first ligand occupying a dimer). \( K_{D2} \) represents the equilibrium between free ligand and the occupied and semi-occupied dimer (the second ligand occupying a dimer in which one binding site is already occupied by the first ligand). Assuming symmetry, the rate at which the ligand binds to the dimer is proportional to the number of unoccupied receptors in the dimer, therefore twice for the empty compared with the semi-occupied dimer. The rate at which the ligand dissociates from the dimer is proportional to the number of occupied receptors in the dimer, therefore twice in the occupied compared with the semi-occupied dimer. This implies, and it is easy to demonstrate, that when the intrinsic affinity of both receptors in the dimer is the same, i.e., when there is no cooperativity, \( K_{D2} = 4K_{D1} \). The two-state dimer model also introduces a dimer cooperativity index, \( D_{C} \), which is defined as log \((4K_{D1}/K_{D2})\). \( D_{C} = 0 \) implies no cooperativity, whereas positive and negatives values imply positive and negative cooperativity, respectively (Fig. 1).

Equation for competitive-inhibition experiments:

\[ A_{\text{bound}} = \frac{(K_{DA2}A + 2A^2)}{K_{DB1}K_{DA2} + K_{DA2}A + A^2} \]

\[ + \frac{K_{DA2}B/K_{DAB}}{K_{D1}K_{DA2} + K_{DA2}A + A^2} \]

\[ + \frac{K_{DA2}B/K_{DAB}}{K_{D1}K_{DA2}B/K_{DB1}} \]

\[ + \frac{K_{DA1}K_{DA2}B^2}{(K_{DB1}K_{DB2})} \]

Although the equation might look complex, there are just a few constants to be determined. \( A \) is again the concentration of radioligand (fixed in competition experiments). \( B \) is the concentration of the competing compound. \( R_{T} \) is again the total amount of receptor dimers. \( K_{DA1} \) and \( K_{DA2} \) are the macroscopic equilibrium dissociation constants for the binding of \( A \), the radioligand, to the first and second receptor in the dimer, equivalent to \( K_{D1} \) and \( K_{D2} \) described above for saturation experiments. These constant values would have been calculated from previous saturation experiments with the radioligand and, therefore, they are already known. \( K_{DB1} \) and \( K_{DB2} \) correspond to the macroscopic equilibrium dissociation constants for the binding of \( B \), the competing compound, to the first and second receptor in the dimer (Fig. 1). And there is one more constant, \( K_{DAB} \), which can be considered as a hybrid equilibrium dissociation constant, a value of the association and dissociation of \( B \) on a dimer semi-occupied by \( A \) (Fig. 1). \( K_{DAB} \) is instrumental to explain a biphasic curve in competitive-inhibition experiments by means of a non-cooperative binding. It measures an allosteric interaction between the two different ligands in the dimer, by which the binding of the radioligand to the first protomer decreases or increases the affinity of the competitor to the second protomer. It can easily be demonstrated that when the binding of the radioligand to one protomer in the dimer does not modify the binding of the competing ligand to the other empty protomer, \( K_{DAB} = 2K_{DB1} \) (Figs. 1 and 2). Reciprocally, a \( K_{DBA} \) value, which is the equilibrium dissociation constant of \( A \) binding to a receptor dimer semi-occupied by \( B \), can be deduced (Figs. 1 and 2). The relation between these parameters is:

\[ K_{DBA} = K_{DAB}K_{DA1}/K_{DB1} \]

The two-state dimer model also introduces two dimer radioligand/competitor modulation indexes, \( D_{AB} \) and \( D_{BA} \), which are defined as log \((2K_{DB1}/K_{DAB})\) and log \((2K_{DA1}/K_{DBA})\), respectively. \( D_{AB} \) or \( D_{BA} = 0 \) implies no modulation, whereas positive and negatives values imply positive and negative modulation, respectively (Figs. 1 and 2). Furthermore, we can also define a dimer cooperativity index for the competing ligand, \( D_{CB} \), as log \((4K_{DB1}/K_{DB2})\). \( D_{CB} = 0 \) implies no cooperativity, whereas positive and negatives values imply positive and negative cooperativity, respectively (Figs. 1 and 2). Finally, the two-state dimer model allows the calculation of the concentration of the competitor providing half saturation = \((K_{DB1}K_{DB2})^{1/2}\), independent of the biphasic or monophasic nature of the competition curve or of the radioligand concentration (Casadó et al., 2009a).

Considering the evidence of receptor oligomerization and the validity of the dimer-cooperativity model, the two-state dimer model provides a better approach than the monomer–G protein models for fitting data that eventually give more accurate and physiologic relevant parameters. In terms of accuracy, the two-state dimer model is significantly more robust than the
two-independent-site model. Thus, with the two-state dimer model the same parameters ($R_T$, equilibrium dissociation constants) are obtained irrespective of the concentration of radioligand (Casadó et al., 2009a). In terms of physiologic relevance, the two-state dimer model does not only give equilibrium dissociation constants for high- and low-affinity binding to receptor dimers, but provides indexes of ligand cooperativity and of allosteric modulation between different ligands simultaneously binding to the dimer. For instance, the analysis with the two-state dimer model of the complex competition data obtained with the ligand SCH-442416 displacing $[^3H]$ZM-241385 indicated the existence of a strong negative cooperativity of the binding of SCH-442416 to the adenosine $A_2A$ receptor, without evidence for allosteric interactions between both ligands (Orru et al., 2011a) (Fig. 2, B and C). This behavior of SCH-442416 was not observed with other adenosine $A_2A$ receptor ligands and was only observed in cells transfected with dopamine D$_2$ receptors, which therefore act as allosteric modulators of adenosine $A_2A$ receptor dimers (Orru et al., 2011a). Thus, the observation of dopamine D$_2$ receptor-mediated negative binding cooperativity of SCH-442416 suggests that the minimal $A_2A$–D$_2$ receptor heteromer unit is made of heteromers of homodimers (at least heteromers with $A_2A$ receptor dimers). This has important pharmacological implications that will be discussed below. The robust and significant fit of the equations provided by the two-state dimer model agrees with the above-reviewed evidences supporting dimers as predominant GPCR species and functional units. It is also of importance to underscore that the two-state dimer model should provide a good approximation, even in the context of a significant proportion of high-order oligomers, because we would probably be dealing with dimers or other multiples of dimers (see above).

Radioligand competition curves are usually analyzed for drug screening to determine the affinity constants

![Figure 2](image_url)

**Fig. 2.** Application of the two-state dimer model. Two different competing ligands, the adenosine $A_2A$ receptor agonist CGS 21680 (A) and the $A_2A$ receptor antagonist SCH-442416 (B and C), are used to displace the $A_2A$ receptor antagonist $[^3H]$ZM-241385 from membrane preparations of sheep striatum (A) or mammalian cells stably transfected with adenosine $A_2A$ and $A_1$ receptors (B) or $A_2A$ and dopamine D$_2$ receptors (C). (A) CGS 21680 displaces the binding of $[^3H]$ZM-241385 in a biphasic manner; although not obvious by just looking at the graph, the analysis with a monomeric model gives a statistically significant better fit for two than for one binding site; analysis with the two-state dimer model indicates that the agonist does not show negative cooperativity ($D_{CB} = 0$); in fact, previous studies with saturation experiments with $[^3H]$CGS 21680 usually show noncurvilinear Scatchard plots (Jarvis et al., 1989; Borea et al., 1995); the analysis nevertheless indicates that the biphasic displacement can be explained by an allosteric modulation between ligands, by which the binding of $[^3H]$ZM-241385 facilitates the binding of CGS 21680 to the $A_2A$ receptor dimer ($D_{AB} = 0.5$). (B) Typical antagonist/antagonist competition, with SCH-442416 displacing in a monophasic manner $[^3H]$ZM-241385 ($D_{CB} = 0$; $D_{AB} = 0$). On the other hand, in C, SCH-442416 displaces $[^3H]$ZM-241385 in an obvious biphasic manner; the analysis with the two-state dimer model indicates that in this case, with coexpression of D$_2$ receptors, SCH-442416 binding to $A_2A$ receptors displays a strong negative cooperativity ($D_{CB} = -2.30$; $D_{AB} = 0$). Results are modified from Casadó et al. (2009a) and Orru et al. (2011), where details of the methods, including radioligand concentrations, can be found. CGS 21680, 4-(2-[6-amino-9-(N-ethyl-β-D-ribofuranuronamidosyl)-8H-purin-2-yl]amino)ethyl]benzenepropanoic acid.
for putative agonists and antagonist. If the target receptor is assumed to be primarily in its monomeric form, monomeric models should be used. However, if homomers have already been described for the target receptor, the two-state dimer model should be the most appropriate choice. In the absence of previous assumptions and in the absence of obvious biphasic competition curves, monomeric models could be used to provide initial estimates of drug affinities, because they imply a lesser number of constants to handle. When not obvious as biphasic, monomeric models can also be used to establish if the competition curves fit significantly better for one or two binding sites. In the latter case we should proceed with the analysis with the two-state model (see Fig. 2A). Nevertheless, when looking for the most appropriate model, the best approach would be implementing both a monomer-based model and a two-state dimer model and comparing results (preferably using a statistical test; see Casadó et al., 2009a). It is important to realize that choosing either model should not imply discarding the possibility of the existence of mixtures of monomers and dimers/oligomers.

D. Allosterism in Receptor Oligomers: Allosteric Modulation of Intrinsic Efficacy and Functional Asymmetry

The two-state dimer model assumes that both protomers have the same initial ligand binding properties and, therefore, the same probability to bind the first ligand molecule and the same ability of the first ligand–protomer complex to allosterically modulate the binding properties of the second protomer. That is, there should be an initial symmetry of both protomers in a receptor dimer at the binding level. Are there symmetric signaling properties of both protomers in a receptor dimer? This requires examining functional responses and to address questions about the minimal functional GPCR-G protein unit. Furthermore, above we only considered allosteric modulation of the affinity of ligands, whereas allosteric modulation of the intrinsic efficacy of the ligands should also be considered. When dealing with dimers, the most accepted model, at present, is that two protomers can only accommodate one heterotrimeric G protein complex (reviewed in Maurice et al., 2011). Extensive biochemical evidence indicates the involvement of at least two distinct G protein regions in interactions with the receptor. The primary docking site is in the C terminus of the Go subunit, which penetrates into the crevice created in the intracellular surface of the receptor by the movement of cytoplasmic regions of TM5 and TM6 upon binding of the agonist (Oldham and Hamm, 2008; Rasmussen et al., 2011). Another putative site is in the C terminus of the Gy subunit. These regions are 55 Å apart in the Gaβγ heterotrimer, a distance greater than the width of the monomeric GPCR (approximately 45 Å), indicating that for both contacts to take place simultaneously, one heterotrimeric G protein must contact two receptor protomers (Oldham and Hamm, 2008). Rasmussen et al. (2011) recently resolved the crystal structure of the active state ternary complex composed of agonist-occupied monomeric β2-adrenoceptor and nucleotide-free Gα heterotrimer. One of the most interesting findings was the lack of direct interactions between the β2-adrenoceptor and Gβγ. Given that the heterotrimer is required for efficient coupling to a GPCR, these results are consistent with the existence of β2-adrenoceptor dimers in cell membranes with one protomer interacting with Ga and the second protomer interacting with the Gβγ subunit (Rasmussen et al., 2011). Although a sequential interaction of simultaneously incompatible contacts could still be possible (Herrmann et al., 2004), this sequential fit mechanism would still be compatible with the existence of GPCR homodimers and with the pentameric structure consisting of one GPCR homodimer and one heterotrimeric G protein as a minimal functional unit (Baneres and Parello, 2003; Han et al., 2009; Pellissier et al., 2011). Even if we consider tetramers as the predominant oligomeric species, the most common minimal receptor–G protein stoichiometry would still probably be 2:1, and oligomeric entities can be viewed as multiples of dimers, as, for instance, suggested by the crystallographic structure of μ-opioid receptor and β1-adrenoceptor (Manglik et al., 2012; Huang et al., 2013).

The apparent asymmetric pentameric structure of GPCRs (homodimer plus heterotrimeric G protein) might seem incompatible with the two-state dimer model and the dimer-cooperativity model, because they assume there is an initial symmetry between both protomers regarding ligand binding properties (the same probability to bind the first ligand molecule) and that the G protein is always interacting with the receptor (Casadó et al., 2007). In this respect, two opposing models have been presented to explain the encounter between G proteins and the activated receptor. In the “collision coupling” model, the interactions occur as a result of the free lateral diffusion, and G proteins only interact with the agonist-bound receptor. In the “precoupling” model, G proteins are already interacting with receptors before agonist binding and the ligand modifies this interaction by creating the conformational change in the receptor (crevice) that allows the α subunit to “tightly” bind the receptor and induce G protein activation (reviewed in Oldham and Hamm, 2008). About the symmetry, the two-state dimer model only assumes an initial symmetry (at least in terms of ligand-binding probability) and fits very well with the precoupling model of receptor–G protein interaction. The two-state dimer model accepts that the ligand binding to the first protomer determines an asymmetric function of the pentameric functional unit. According to the model, this ligand-induced asymmetry
determines the allosteric modulations at the binding level, such as the reduced affinity of the ligand for the second protomer (negative cooperativity).

What is the functional response that a ligand can produce when binding to one or both protomers in a GPCR homodimer? Is the binding of an agonist to one of the protomers enough to elicit a full functional response? Does the asymmetric constraint of G protein coupling to the two protomers determine an asymmetric functional response? Most experimental data agree with the model that proposes that ligand occupancy to the first protomer is enough to produce a significant G protein activation and functional response. Occupancy of the second protomer will then usually potentiate (Kniazeff et al., 2004; Pellissier et al., 2011), but could also reduce (Han et al., 2009) or not alter significantly (Hlavackova et al., 2005), the functional response (irrespective of the allosteric modulations at the binding level).

Therefore, when the ligand binds to the second protomer in a homodimer, it will often act as an allosteric modulator of the intrinsic efficacy of the ligand when binding to the first protomer.

We should also consider G protein–independent signaling, such as β-arrestin–dependent signaling. Similar to G proteins, GPCR oligomerization raises the question of whether GRK and β-arrestin binding to GPCRs occur in an asymmetric manner. The possible asymmetric nature of the receptor–β-arrestin interaction is still a matter of debate. As with G protein binding and activation, studies with artificially reconstituted monomeric GPCRs show that receptor dimers are not required for their functional interaction with GRK and β-arrestins (Tsukamoto et al., 2010; Bayburt et al., 2011). These results, however, do not exclude the possibility that GRK or β-arrestins bind to GPCR dimers (Modzelewski et al., 2006; Sommer et al., 2012). GPCR oligomerization potentially provides a larger platform to accommodate the different GPCR interacting proteins necessary for receptor function (reviewed in Maurice et al., 2011). This is demonstrated in the complex of melatonin MT₁ receptor dimers, Gᵢ protein and RGS20 (a protein that regulates the speed of G protein signal transduction), which both bind directly to helix 8 of the receptor (Maurice et al., 2010). In this complex, protomer 1 binds the ligand and the G protein and protomer 2 binds RGS20. In this case, RGS20, by slowing down the decay time of G protein inactivation, thus participates in prolonged signal transduction (Maurice et al., 2010).

Several examples of allosteric modulation of intrinsic efficacy have been reported for GPCR heteromers. As for ligand affinity, there are cases of allosteric modulation of the intrinsic efficacy of a ligand of one of the protomers by the other protomer or by a ligand that binds to the other protomer. As an example of the latter, in the α₂A-adrenoceptor–μ-opioid receptor heteromer, morphine binding to the μ-opioid receptor inhibits α₂A-adrenoceptor signaling (Jordan et al., 2003). This has been related to a morphine-induced conformational change of the α₂A-adrenoceptor, as detected by dynamic intramolecular FRET techniques (Vilardaga et al., 2008). These studies indicated that activation of the μ-opioid receptor component in the α₂A-adrenoceptor–μ-opioid receptor heteromer modulates α₂A-adrenoceptor receptor signaling by a direct conformational change that propagates from the μ to the α₂A-adrenoceptor within 0.4 seconds, slightly faster than the rate of G protein activation, implicating a direct allosteric modulation through the receptor heteromer (Vilardaga et al., 2008).

As an example of an allosteric modulation of intrinsic efficacy of a ligand of one of the protomers by the other protomer in the GPCR heteromer, in the metabotropic glutamate mGlu₂-serotonin 5-HT₂A receptor heteromer, glutamate produces a stronger mGlu₂-mediated signaling and serotonin produces a weaker serotonin 5-HT₂A-mediated signaling compared with when each receptor is expressed alone (Fribourg et al., 2011). In the same receptor heteromer, the serotonin 5-HT₂A receptor antagonist clozapine also induced a positive allosteric modulation of the intrinsic efficacy of glutamate (Fribourg et al., 2011). These results could have implications for the understanding of the clinical efficacy of this compound in schizophrenia and establish the mGlu₂–5-HT₂A receptor heteromer as a promising target for the treatment of this disease.

Considering, therefore, receptor heteromers in the frame of allosteric modulation of GPCRs, there are examples of allosteric modulation by specific ligands of some receptors of the affinity and intrinsic efficacy of ligands for other receptors (see above). There are also significant examples of functional selectivity, where one of the protomers of the heteromer acts as an allosteric modulator that “forces” the other receptor protomer to signal predominantly through a distinct signaling pathway. An example is that of the δ-μ-opioid receptor heteromer (Rozenfeld and Devi, 2010). Under normal physiologic conditions, the μ-opioid receptor is found mostly in the homomeric state, and morphine stimulation induces G protein–mediated signaling, which is involved in its analgesic effects, and low β-arrestin–mediated signaling, which promotes unwanted effects, such as tolerance to the analgesic effects (Raehal et al., 2011). In contrast, upon chronic treatment with morphine, the abundance of δ-μ-opioid receptor heteromers increases (Gupta et al., 2010), and morphine stimulation induces β-arrestin–mediated signaling (Rozenfeld and Devi, 2007), which is thought to contribute to the tolerance to the analgesic effect of morphine upon its chronic administration (Raehal et al., 2011). Notably, occupancy of the δ-opioid receptor protomer with a selective δ-opioid receptor antagonist (Rozenfeld and Devi, 2007) or with a bivalent ligand in the δ-μ-opioid
receptor heteromer (see below and Daniels et al., 2005) allows for the restoration of G protein–biased agonism of morphine and, therefore, its analgesic effects (Rozenfeld and Devi, 2010).

Other dramatic examples of functional selectivity in receptor heteromers are changes in G protein coupling, such as with the angiotensin AT1–cannabinoid CB1 receptor heteromer (Rozenfeld et al., 2011). Although, in the normal liver angiotensin AT1 receptor signals via Gq, under conditions of alcohol-induced liver fibrosis cannabinoid, CB1 receptor is upregulated and heteromerizes with angiotensin AT1 receptor and this, in turn, leads to G1-mediated signaling by angiotensin AT1 receptor that can be blocked by cannabinoid CB1 receptor antagonist, suggesting that AT1–CB1 receptor heteromer represents a disease-specific and potentially tissue-specific therapeutic target. Another example of change in G protein coupling is the dopamine D1–histamine H3 receptor heteromer, where dopamine D1 receptor agonists activates G1 instead of Gα proteins (inhibition of cAMP formation) (Ferrada et al., 2009). Another important allosteric property of the D1–H3 heteromer is the ability of H3 receptor agonists to inhibit dopamine D1 receptor–mediated G protein and β-arrestin signaling, which allows histamine to provide a brake on dopamine D1 receptor–mediated effects, including cell death (Moreno et al., 2014). This heteromer also binds the two-transmembrane sigma σ1 receptors, which are well known targets for cocaine. Binding of cocaine to the D1–H3–σ1 complex disrupts the allosteric properties of the D1–H3 heteromer allows dopamine D1 receptor to couple to Gα and to eliminate the histamine H3 receptor–mediated signaling brake, which promotes cell death (Moreno et al., 2014). Therefore D1–H3 heteromers may provide a new therapeutic target for the D1 receptor–mediated neurotoxic effects of cocaine.

Another recent example of functional selectivity in receptor heteromers is the dopamine D2–ghrelin GHS1a receptor heteromer (Kern et al., 2012). These receptors are colocalized in the brain, in the hypothalamus and brain stem (Guan et al., 1997). Interestingly, ghrelin receptors seem to be orphan receptors in certain localizations, because the peptide ghrelin, which is produced in the stomach, can only reach the arcuate nucleus but not other hypothalamic nuclei and other brain regions. In the hypothalamus, heteromerization with ghrelin GHS1a receptor modifies dopamine D2 receptor signaling, resulting in Gβγ-dependent mobilization of Ca2+ (Kern et al., 2012). The anorexigenic effects of dopamine D2 receptor agonists seem to be mediated by the hypothalamic D2–GHS1a receptor heteromer, which therefore might become an important therapeutic target for eating disorders.

In summary, the allosteric analysis of GPCR homomers and heteromers provides overwhelming additional evidence for the results of in vitro experiments and experiments in heterologous systems, supporting the existence of functionally and pharmacologically relevant GPCR oligomers. The evidence points to the pentameric structure constituted by one GPCR homodimer and one heterotrimeric G protein as a main minimal functional unit and oligomeric entities can be viewed as multiples of dimers. Allosteric mechanisms determine a multiplicity of possible unique pharmacological properties of receptor homomers and heteromers. Some general mechanisms seem to apply, particularly at the level of ligand-binding properties. When considering receptor homomers, the two-state dimer model provides the most practical method to analyze ligand–GPCR interactions. If possible, similar practical models would need to be developed to study ligand–GPCR interactions for receptor heteromers and for dynamically changing oligomers. In addition to ligand-binding properties, unique properties for each GPCR oligomer seem to emerge in relation to different intrinsic efficacy of different ligands (intrinsic efficacy) for different signaling pathways (functional selectivity). This provides a rationale for the use of GPCR oligomers, and particularly heteromers, as novel targets for drug development.

III. G Protein–Coupled Receptor Oligomers as Novel Targets for Drug Development

A. Localization of Receptor Oligomers in Native Tissues

Before considering the possible role of GPCR oligomers in vivo, and particularly receptor heteromers, as targets for drug development, we must first readdress questions about the demonstration of receptor homo- and heteromers in native tissues, because most of the studies discussed above have been performed in heterologous expression systems, particularly in transfected mammalian cells. Next, we need to address the questions about localization in situ and significance of receptor homomers and heteromers in vivo (Levoye et al., 2006b; Pin et al., 2007; Ferré et al., 2007, 2009). Efforts toward addressing these important questions have been hampered by the lack of appropriate tools and techniques that would allow their clear demonstration in native tissues. Biochemical techniques such as coimmunoprecipitation cannot provide unambiguous evidence for direct protein–protein interactions, and biophysical techniques, such as BRET and FRET, are difficult to implement in tissues. In contrast to heterologous systems, where wild-type or mutated receptors fused with tags or fluorescent proteins can be easily expressed, the expression level of receptors in native tissues is difficult to modify and their sequences cannot be modified unless using knock-in strategies. Nevertheless, quantitative FRET in situ has been used to explore the presence of dopamine D1–D2 receptor heteromers in brain tissue (Hasbi et al., 2011).
An alternative approach to label and detect native GPCR oligomers has been the use of antibodies or fluorescent ligands. However, they need to be highly specific and to provide a reliable signal, and the levels of GPCR oligomers should be abundant or highly regulated. Recent efforts to identify tools that fit some of these criteria have begun to yield promising results. Using labeled ligands that could demonstrate proximity of endogenous receptors, the study by Albizu et al. (2010) provided quite a clear demonstration of the presence of oxytocin receptor homomers in the mammary gland. They showed differential FRET with labeled agonists and antagonists in tissue as they had seen with transfected cells, implying the existence of oxytocin receptor homomers. This is therefore an indirect but clear demonstration of the existence of receptor homomers in native tissues.

Efforts to generate heteromer-selective antibodies using a subtractive immunization strategy have begun to yield useful monoclonal antibodies with heteromer selectivity, because they recognize the epitope only in wild-type animals but not in animals lacking either protomers of the heteromer (Gomes et al., 2013a,c). These heteromer-selective antibodies have helped not only studies examining heteromer levels and regulation in endogenous tissue but also studies investigating the extent of heteromer-specific signaling (Gomes et al., 2013a,c). In the case of δ–μ-opioid receptor heteromers, the antibodies were helpful in demonstrating morphine-induced increases in heteromer abundance, as well as characterizing heteromer-mediated signaling (Gupta et al., 2010). The AT₁–CB₁ receptor heteromer selective antibody was useful in demonstrating upregulation of the heteromer in hepaticstellate cells after alcohol-induced fibrosis (Rozenhofeld et al., 2011). The δ-opioid–CB₁ receptor heteromer selective antibody was useful in demonstrating increases in δ-opioid–CB₁ receptor heteromers in the brain after peripherally elicited neuropathic pain (Bushlin et al., 2012). Finally, the δ–κ-opioid receptor heteromer selective antibody was critical in demonstrating a role for this heteromer in sensory pain transmission, because the antibody was able to augment the effect of δ receptor agonist-mediated antinociception (Berg et al., 2012). Together these studies demonstrate that GPCR heteromer antibodies, if found to be highly selective, are critical and important tools in studies of GPCR heteromers in native tissue.

The proximity ligation assay (PLA) is an antibody-based method that also provides an approximation to the demonstration of receptor heteromers in tissues. It has been used for instance for the demonstration of close proximity of adenosine A₂A and dopamine D₂ receptors in brain slices (Trifilieff et al., 2011). In PLA, primary antibodies labeling the two putative protomers of the receptor heteromer are labeled with two different species-specific secondary antibodies conjugated to complementary oligonucleotides. When the antibodies are in close proximity, the complementary DNA strands can be ligated, amplified, and visualized with a fluorescent probe. The maximal distance between the secondary antibodies in this assay is ~16 nm, only slightly larger than that for resonance energy transfer between fluorophores (~10 nm) (Soderberg et al., 2006). However, PLA does not demonstrate GPCR heteromerization, but allows a validation in vivo of the molecular proximity of two GPCRs, giving the frame for heteromerization to take place in vivo.

Another indirect but easier and straightforward demonstration depends on the identification of a biochemical property ("biochemical fingerprint") of the receptor oligomer (Ferré et al., 2009). To ascertain a "biochemical fingerprint" of the receptor heteromer, the putative biochemical property should be disrupted with molecular or chemical tools that significantly destabilize the quaternary structure of the heteromer (such as introducing mutations of key determinant residues at the oligomerization interfaces or using competing peptides that harbor the same sequence as that of the receptor heteromer interface). The recently reported dopamine D₂–D₄ receptor heteromer serves as a good example. A human polymorphism of the dopamine D₄ receptor, D₄,7 (with seven repeats of a proline-reach 16-amino acid sequence in the third intracellular loop) differs from the most common variant, D₄,4 (with four repeats) in that it does not heteromerize with dopamine D₂ receptor and it does not show a pharmacological D₂–D₄ receptor interaction (Gonzalez et al., 2012). This interaction consists of a dopamine D₂ receptor agonist-mediated potentiation of MAPK activation induced by a dopamine D₄ receptor agonist (Gonzalez et al., 2012). Thus, a synergistic MAPK activation could be used as a biochemical fingerprint of the heteromer, which could only be observed in mice expressing wild-type dopamine D₄ receptor (which also heteromerized with mouse wild-type dopamine D₂ receptor), but not in knock-in transgenic mice expressing the dopamine D₄,7 human variant (Gonzalez et al., 2012). The use of transgenic animals is therefore a useful indirect method to study receptor heteromer function. An additional example is the recent study about the role of MT₁–MT₂ receptor heteromers on the effect of melatonin on rod photoreceptor light sensitivity, which was abolished in melatonin receptor MT₁,Δ/Δ and MT₂,Δ/Δ mice (Baba et al., 2013). The involvement of MT₁–MT₂ receptor heteromers in this effect was further underscored by the lack of effect of melatonin in transgenic mice overexpressing a nonfunctional melatonin MT₂ receptor mutant that competes with the formation of functional MT₁–MT₂ heteromers in photoreceptor cells (Baba et al., 2013). These experiments are of therapeutic interest for the improvement of photoreceptor functioning in several clinical conditions and applications in the pathogenesis of age-related macular degeneration and glaucoma.
Therefore, use of several complementary approaches is needed to address the identification of GPCR oligomers in native tissues. It is essential that these approaches are critically evaluated and validated using artificial systems, such as mammalian transfected cells, especially when looking for specific biochemical properties of a GPCR oligomer. It is not enough just reproducing some findings in native tissues previously found in the artificial systems. A unique biochemical finding in mammalian transfected cells that express receptor heteromers (which can be demonstrated with properly applied biophysical techniques) could well be related to interactions between the two receptors that do not depend on oligomerization. As mentioned above, we need to obtain and implement tools that specifically disrupt GPCR oligomerization or the quaternary structure of the GPCR oligomer. When possible, those tools should also be applied in situ. For instance, by selectively interfering with the oligomer interfaces, particular mutations or the in vitro application or in vivo delivery of peptides altering the quaternary structure of a GPCR oligomer (as first demonstrated with biophysical techniques in artificial systems) should selectively alter the detection of the putative biochemical property of the GPCR oligomer.

**B. Do Receptor Heteromers Constitute Possible Targets for Drug Development?**

GPCRs are targeted by many therapeutic drugs aiming at restoring or ameliorating a variety of health disorders and remain a primary focus of biomedical research and programs of pharmacological drug discovery. However, despite the proven success of GPCRs as drug targets, clinically useful ligands do not exist for the majority of GPCRs. A main contributing factor is the lack of selectivity of orthosteric ligands across members of a GPCR subfamily (or that certain GPCRs do not have roles in major disease phenotypes), with the obvious selectivity problem, which leads to unwanted side effects. An alternative approach already found successful for ligand-gated ion channels, such as benzodiazepines, positive allosteric modulators of GABA<sub>A</sub> receptors, is the development of GPCR allosteric modulators (ligands that modulate the affinity or intrinsic efficacy of orthosteric agonists of the same GPCR unit) (Conn et al., 2009). Allosteric modulators can potentially overcome the selectivity problem. The orthosteric binding sites across members of a GPCR subfamily for a particular endogenous ligand, such as noradrenaline or dopamine, are highly conserved, but GPCRs offer numerous possible selective allosteric binding sites across different receptor subtypes. Furthermore, allosteric modulators possess pharmacologically appealing properties, such as saturability. In recent years, significant progress has been made in the discovery of various GPCR allosteric modulators, although not without difficulties, such as generating a structure–activity relationship in an allosteric compound series, or likelihood of lack of cross-species activity for validation/efficacy studies (Conn et al., 2009).

Receptor heteromers, with their allosteric properties, give rise to a new kind of pharmacological target. The ability of one of the protomers to act as an allosteric modulator of the second protomer in the receptor heteromer gives the possibility of finding selective ligands for the protomer acting as conduit of the allosteric modulation. A proof of concept was provided by the dramatically decreased binding affinity (strong negative cooperativity) of the adenosine A<sub>2A</sub> receptor antagonist SCH-442416 in cells cotransfected with dopamine D<sub>2</sub> receptors compared with cells only expressing adenosine A<sub>2A</sub> receptors or also expressing adenosine A<sub>1</sub> (Orru et al., 2011a) (Fig. 2, B and C). Several other antagonists, such as MSX-3, also showed a less dramatic yet significantly decreased affinity for adenosine A<sub>2A</sub> receptors when cotransfected with dopamine D<sub>2</sub> receptors (Orru et al., 2011a). These results are important from a practical pharmacological point of view, because different striatal subpopulations of adenosine A<sub>2A</sub> receptors modulate different striatal functions (Orru et al., 2011a,b), and these subpopulations are constituted by different adenosine A<sub>2A</sub> receptor heteromers. A<sub>2A</sub>–D<sub>2</sub> receptor heteromers are selectively localized postsynaptically in GABAergic striatopallidal neurons, where they exert a key role in the modulation of the activity of these neurons (see above and Azdad et al., 2009). On the other hand, A<sub>1</sub>–A<sub>2A</sub> receptor heteromers are selectively localized presynaptically in the striatal glutamatergic terminals that make synaptic contact with GABAergic strionigral neurons (Quiroz et al., 2009). Through these heteromers, postsynaptic blockade of adenosine A<sub>2A</sub> receptors produces locomotor activation, and presynaptic blockade of adenosine A<sub>2A</sub> receptors produces inhibition of corticostriatal neurotransmission (Ciruela et al., 2006; Quiroz et al., 2009; Orru et al., 2011a). The postsynaptic effect has implications for Parkinson’s disease and, based on the existence of A<sub>2A</sub>–D<sub>2</sub> receptor interactions, adenosine A<sub>2A</sub> receptor antagonists are being used for the treatment of Parkinson’s disease (see below and Armentero et al., 2011). On the other hand, presynaptic effects can have implications for drug addiction, but a concomitant postsynaptic adenosine A<sub>2A</sub> receptor blockade will determine opposing effects. In this regard, a low dose of the selective adenosine A<sub>2A</sub> receptor antagonist MSX-3 (shown to be preferentially presynaptic in rodents; Orru et al., 2011a) decreased tetrahydrocannabinol and anandamide self-administration in squirrel monkeys, whereas a slightly higher dose had potentiating effects (Justinová et al., 2011). The appearance of potentiating rather than suppressing effects on cannabinoid reinforcement at the slightly higher dose of MSX-3 would likely preclude the use of such
a compound as a pharmacologic treatment of cannabis abuse. As expected, recent results showed a significant antagonistic effect of SCH-442416 over a much wider range of doses than with MSX-3 (Justinová et al., submitted manuscript), strongly supporting the use of this or other similar compounds for the treatment of drug abuse.

When the receptor heteromer is the conduit for the allosteric modulation, the allosteric modulator acts very much the same way as a classic allosteric modulator. The difference is that the binding of the allosteric modulator takes place in the orthosteric site of one of the receptors in the heteromer and the modulated ligand binds to the orthosteric site of the other receptor. For instance, adenosine A2A receptor antagonists increase the affinity of the dopamine D2 receptor for dopamine, which as mentioned earlier is being used as a strategy for the treatment of Parkinson's disease. In fact, evidence has already been provided for an increase in the therapeutic index of L-dopa with adenosine A2A receptor antagonists (Armentero et al., 2011). Furthermore, adenosine A2A receptor antagonists can potentially be useful as monotherapy in early stages of Parkinson's disease where some symptoms seem to be related to the loss of a tonic dopamine D2 receptor-mediated inhibition of adenosine A2A receptor signaling (Ferré et al., 2008; Armentero et al., 2011). Schizophrenia is another pathologic condition where the A2A–D2 receptor heteromer could be used as a therapeutic target (Ferré, 1997). Thus, different from dopamine D2 receptor antagonists, adenosine A2A receptor agonists behave as atypical antipsychotics (antipsychotics with low extrapyramidal side effects) in animal models (Rimondini et al., 1997). This profile could be related to the allosteric and therefore saturable effect on dopamine D2 receptor-mediated signaling. So far, the cardiovascular effects of adenosine A2A receptor agonists preclude their use as antipsychotics, but ligands with selectivity for the A2A–D2 receptor heteromer would be devoid of these peripheral side effects because of the lack of peripheral A2A–D2 receptor interactions (Schindler et al., 2004). In the A2A–D2 receptor heteromer as target for Parkinson's disease and schizophrenia, we deal with an allosteric modulation of the affinity of dopamine D2 receptor ligands, although changes in intrinsic efficacy might also take place. As mentioned above, another candidate target for antipsychotic treatment is the mGlu2–5HT2A receptor heteromer, where the atypical antipsychotic clozapine, in addition to blocking serotonin 5-HT2A, induces an allosteric modulation of the intrinsic efficacy of glutamate (Fribourg et al., 2011). Functional selectivity is also being addressed in the search for antipsychotics with low liability for side effects.

A particular value of receptor heteromers as targets for drug development is that they can be involved in the pathophysiology of the disorder being targeted. In this regard, the receptor heteromer would be disease specific (Ferré et al., 2010; Gomes et al., 2013a). The dopamine D1–D3 receptor heteromer seems to be involved in the very common and troublesome secondary effect of L-dopa treatment in Parkinson's disease, L-dopa–induced dyskinesia (reviewed in Ferré et al., 2010). The dopamine D1 receptor in the D1–D3 receptor heteromer has significantly more affinity for dopamine or other agonists than when not forming heteromers with dopamine D3 receptors (Fiorentini et al., 2008; Marcellino et al., 2008), which results in a significant increase in the potency of dopamine D1 receptor agonists to signal through the G protein and potentiate cAMP accumulation (Fiorentini et al., 2008). This is a very significant finding, because it is well known that, under normal conditions, striatal dopamine D1 receptors have a much lower affinity for dopamine than dopamine D2 or D3 receptors (Missale et al., 1998). There is a substantial amount of evidence that indicates that dopamine D1 and D3 receptors are colocalized in GABAergic striatonigral neurons (which constitutes the direct striatal efferent pathway), where they play a key role in the development of L-dopa–induced dyskinesia in Parkinson's disease. The working hypothesis is that dopamine denervation followed by chronic L-dopa treatment leads to an increased dopamine D1 receptor signaling promoted by an upregulation of dopamine D3 receptors and by an increase in the expression of D1–D3 receptor heteromers, which produces an imbalance in basal ganglia processing due to an increased activation of the direct pathway (reviewed in Ferré et al., 2010). Significantly, treatment with dopamine D3 receptor partial agonists or antagonists attenuates L-dopa–induced dyskinesia in 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP)–treated monkeys (Bezard et al., 2003; Visanji et al., 2009). Finally, an increase in dopamine D3 receptor density has been observed in human cocaine fatalities (Staley and Mash, 1996), suggesting that a concomitant increase in D1–D3 receptor heteromers may also occur. The D1–D3 receptor heteromer therefore emerges as a target for the treatment of L-dopa–induced dyskinesia and cocaine addiction.

As mentioned earlier, the availability of the heteromer-selective antibodies has helped explore disease-specific modulation of their abundance and hence define them as novel therapeutic targets (Gomes et al., 2013c). A potential role for δ-μ-opioid receptor heteromers in morphine tolerance was implied by studies that demonstrated an increase in the abundance of δ-μ-opioid receptor heteromers in the brain and in dorsal root ganglion cells in response to a paradigm of morphine treatment that leads to development of tolerance (Gupta et al., 2010). AT1–CB1 receptor heteromers are upregulated in alcohol-induced liver fibrosis, and this leads to a shift in signaling from a Gi to Gq-coupled pathway implicating...
the CB₁R–AT₁R heteromer as a potential target for the treatment of liver fibrosis (Rozenfeld et al., 2011). Finally, the δ-opioid–CB₁ receptor heteromer was found to be upregulated during neuropathic pain and the heteromer exhibited unique allosteric modulation, making this an attractive target for the development of therapeutic for the treatment of pain (Bushlin et al., 2012).

C. Approaches for the Identification of Receptor-Heteromer Selective Compounds

Despite their pharmacologic potential, there are very few studies reporting the identification of receptor-heteromer selective compounds. A receptor-heteromer selective compound would not only include ligands with significantly different affinity or intrinsic efficacy for one or both receptors in a receptor heteromer (compared with the same biochemical properties of the ligands for the same receptors when not forming heteromers), it would also include compounds that would exert significant allosteric modulations in the receptor heteromer. Two general approaches were recently used for the discovery of receptor-heteromer selective compounds: discrete direct screening (DDS) and high-throughput screening (HTS). DDS is used when a series of ligands targeting either of the receptors in the heteromer is available. In this case, relatively smaller number of ligands would allow the use of labor-intensive techniques to demonstrate differences in affinity (for agonists or antagonists) or intrinsic efficacy (agonists). The most important step for this strategy to be successful is the availability of appropriate cell lines. The study by Orru et al. (2011a) provides a successful example of DDS of adenosine A₂A receptor antagonists in different comparable cell lines expressing only adenosine A₂A receptors or A₁–A₂A or A₂A–D₂ receptor heteromers. As mentioned above, radioligand-binding experiments and analysis of the data with the two-state dimer model demonstrated a very specific behavior, a negative cooperativity of the adenosine A₂A receptor antagonist SCH-442416 in cells expressing A₂A–D₂ receptor heteromers (Fig. 2, B and C). This biochemical result predicted a preferential striatal presynaptic profile of SCH-442416, which totally correlated with in vivo experiments in rodents and monkeys. Notably, preclinical data suggest that this compound could be useful in the treatment of drug addiction (see Section III.B). SCH-442416 can now be used as a lead compound for further chemical optimization leading to novel compounds exhibiting even lower affinity for the A₂A–D₂ receptor heteromer (striatal postsynaptic adenosine A₂A receptors), whereas keeping a significant affinity for A₁–A₂A receptor heteromers (striatal presynaptic adenosine A₂A receptors). The same DDS approach could be used to find adenosine A₂A receptor antagonists useful for Parkinson’s disease that would have preferential affinity for the A₂A–D₂ versus the A₁–A₂A receptor heteromer or that would have stronger allosteric modulatory effects of the affinity or intrinsic efficacy of dopamine D₂ receptor agonists. Other receptor-heteromer selective compounds obtained by the DDS approach include the putative D₁–D₂ receptor heteromer compound SKF-83959 [6-chloro-7, 8-dihydroxy-3-methyl-1-(3-methylphenyl)-2,3,4,5-tetrahydro-1H-3-benzazepine] (Rashid et al., 2007; but see Chun et al., 2013) and clozapine, which not only acts as a potent serotonin 5-HT₂A receptor antagonist but also increases the intrinsic efficacy of glutamate in the mGlu₅–5-HT₂A receptor heteromer (Fribourg et al., 2011).

HTS using bivalent ligands constitutes another approach for targeting receptor-heteromers (recently reviewed in Hiller et al., 2013). A bivalent ligand consists of two pharmacophoric entities linked by an appropriate spacer. In principle, it should be possible to target GPCR heteromers by selection of adequate, potent, and receptor-selective pharmacophores. The proof of principle came from the work of Daniels et al. (2005), who reported the effects of bivalent ligands incorporating the pharmacophores of the μ-opioid receptor agonist oxymorphine and the δ-opioid receptor antagonist naltrexone tethered by spacers of different lengths to selectively label δ-μ-opioid receptor heteromers. Only ligands with appropriate spacer length produced no tolerance to their analgesic effects (Daniels et al., 2005). Absence of tolerance was therefore most likely due to the appropriate tethering of the μ-opioid receptor agonist and δ-opioid receptor antagonist and again supported the involvement of δ-μ-opioid receptor heteromer in the tolerance to the analgesic effects of opioids. The same research group recently published results that suggest that some bivalent ligands can very selectively interact with the putative μ-opioid–glutamate metabotropic mGlu₅ receptor heteromer, with potent antinociceptive effects (Akgun et al., 2013). Although clinical application of these big molecules might not be feasible for pharmacokinetic reasons, the same as any receptor-heteromer selective compound, bivalent ligands can provide important tools for the identification and study of the functional significance of receptor heteromers. HTS is the most current initial strategy in the process of drug development. Availability of a simple, quick, and robust assay is critical for enabling HTS (reviewed in Franco et al., 2010). A frequently used assay for initial screens is the second-messenger cell-based assay using promiscuous G protein that is able to elicit a calcium response. Another common strategy is based on β-arrestin translocation that occurs as a result of receptor activation-dependent recruitment of β-arrestins. This latter approach was recently used for the identification of δ-μ-opioid receptor heteromer selective compounds, because a previous study suggested that this heteromer preferentially recruits β-arrestin (Rozenfeld and Devi, 2007). Screening of a small-molecule library was performed with the β-arrestin recruitment assay in cell lines expressing...
δ-opioid receptor heteromers; the hits were subject to a secondary screening using cells expressing only δ-opioid or μ-opioid receptor heteromers (Gomes et al., 2013b). The compound named CYM51010 [ethyl 1-[4-(acetyl-amino)benzyl]-4-(2-phenylethyl)piperidine-4-carboxylate] was identified as a ligand that preferentially activates δ-opioid receptor heteromer (Gomes et al., 2013b). The compound showed strong intrinsic efficacy not only for β-arrestin but also for G protein–mediated signaling [guanosine 5′-O-(thio)triphosphate binding]. The δ-opioid receptor-heteromer selective antibody was able to block not only the CYM51010-mediated signaling in heterologous cells, but also intrathecal analgesia in a rodent model of thermal pain (Gomes et al., 2013b). The significant intrinsic efficacy of the compound for δ-opioid receptor–induced β-arrestin and G protein–mediated signaling (compared with morphine) makes CYM51010 an attractive candidate for the development of a new analgesic with lower tolerance than morphine. In fact, using an in vivo assay of analgesia (tail-flick assay), CYM51010 was found to exert similar antinociceptive effects than morphine but with less tolerance (Gomes et al., 2013b). Compounds such as these that selectively target the heteromers are likely to serve as valuable tools to explore the physiologic role of heteromers in biologic systems.

In summary, GPCR heteromers offer very promising new targets for drug development. First, they tend to exhibit unique localization (for instance a type-specific neuron in the brain). Second, their distinct allosteric properties provide a framework for saturability, ligand selectivity, and functional selectivity. These are all properties that are now eagerly pursued in possibly new marketable compounds. Very few studies have addressed the search for receptor-heteromer selective compounds, which is quite surprising given the continuous discovery of functionally significant receptor heteromers (for recent review, see Gomes et al., 2013a). Compounds or reagents (monovalent antibodies or membrane permeant peptides) specifically targeting the heteromer interface are likely not only to serve as much needed tools to explore the physiologic significance of GPCR heteromers, but also serve as “leads” for development of highly selective drugs for the treatment of a variety of disorders. We provide some guidelines for the application of DDS and HTS approaches to the discovery of receptor-heteromer selective compounds. This should definitively serve as a key element in GPCR drug discovery programs.

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