Strategies for G-protein coupled receptor deorphnanization

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Abstract

G-protein coupled receptors (GPCRs) form the largest group of transmembrane proteins in mammals, and they are major regulators of multiple cellular processes. Intense research performed in the last decades revealed their implication in pathophysiology and included many of their members as important drug targets. With the advent of genomics a large number of new GPCR genes have been discovered, however many of them still encode receptors for which no ligands and biological roles have yet been discovered. They are termed orphan GPCRs (oGPCRs) and are under intense scientific investigations by the pharma industry and academia to discover ligands and thus to determine their contribution to pathophysiology. Two strategies have been used to identify ligands of GPCRs, known as the reverse and forward pharmacology. Both of them are presented and discussed in this review.

Keywords: GPCR, deorphnanization, reverse pharmacology, forward pharmacology, drug discovery

INTRODUCTION

G-protein coupled receptors are the largest superfamily of transmembrane receptors in eukaryotes. Their key structural signature is the presence of a seven transmembrane (7TM) domain which separates the extracellularly located N-terminus from the intracellular C-terminus. Another property of GPCRs is that they mediate cellular effects of a cognate ligand by coupling to intracellular heterotrimeric guanine nucleotide binding proteins, or G-proteins, composed of αβγ subunits. Functionally, G-proteins can be grouped into four subtypes based on the structural and functional similarities of their α-subunit: Gq/11, mediating intracellular calcium increase; Gs, activating adenylyl-cyclases (ACs) and increasing cytosolic cyclic adenosine monophosphate (cAMP) in cells; Gi, negative regulators of ACs, thus reducing intracellular cAMP concentration and G12/13, mediating cytoskeletal remodeling cellular motility (1, 2).

GPCRs are key mediators in the regulation of a variety of physiological processes and also participate in a multitude of diseases. They are required for the function of complex senses such as olfaction, taste, or visual perception. Moreover, their role in the regulation of blood pressure, kidney function, cancer progression and allergic reactions has been well documented (3). As a consequence of their biological importance, more than 40% of the world-wide approved drugs target GPCRs either as agonists or antagonists. Several examples of commercially approved drugs targeting GPCRs and their therapeutic effects can be found in the Table 1. Although GPCRs are the most prominent drug target, with a market value of over 65 billion dollars in annual sales (4), almost 140 genes in human genome still encode receptors for which neither biological roles nor endogenous ligands have been identified, termed orphan GPCRs (oGPCRs) (5, 6).

GPCR deorphnanization started in 1986, when the primary structure of the β-adrenergic receptor (β-AR) was characterized. Surprisingly, its amino-acid sequence not only shared high homology with the sequence of visual rhodopsin but also shared a common secondary structure, the 7TM domain (7). This seminal finding led to the concept that all plasma membrane 7TM proteins which are able to interact with G-proteins, belong to a superfamily of receptors. At that time it was already thought that a number of biologically active metabolites, such as acetylcholine, eicosanoids (especially prostaglandins), biogenic amines and hormones were inducing effects by acting as ligands of GPCRs. Newly developed molecular biology techniques, such as degenerated PCR and low-stringency hybridization, were successfully used to...
identify new receptors, with different homologies to β-AR (8). Moreover, with the expansion of genomics at the turn of the century, a large number of potential GPCR genes were identified, reaching almost 900 in human and over 1000 in mouse, including olfactory receptors (9-11). Due to the large number of GPCR genes discovered, some of them were abbreviated as “GPR” followed by a number, as in the case of GPR81 or GPR37 and so on. However, this terminology does not necessarily reflect any degree of relatedness among them.

In the last decades, huge efforts by the academia and pharma industry have been made to identify endogenous ligands and agonists of oGPCRs and by this to reveal their role in pathophysiology and to possibly include them in the druggable targets portfolio. The identification of modulators of oGPCRs can be a tedious and risky process and can reach tremendous costs at the risk of obtaining modest results, if any at all. Therapeutically, although the endogenous ligand of a potentially important oGPCR may remain unknown, the identification of a compound with agonistic properties can be considered a major success because it allows to directly testing its potential beneficial role.

Two main pharmacological strategies have been used in the last years for deorphanization, known as the reverse and the forward pharmacology. Both strategies will be presented in this article discussing their applicability in the orphan receptor research field.

**REVERSE PHARMACOLOGY**

The concept of reverse pharmacology, also known as target-based drug discovery, relies on the hypothesis that the modulatory effect of a protein (i.e., receptor, ion channel, enzyme and so on) will have important biological consequences which can be of therapeutic importance, thus becoming a pharmacological target (12). In other words, reverse pharmacology aims to identify compounds which can alter a chosen target’s activity. In the case of GPCRs, the identification of the endogenous ligand and/or modulatory compounds, as its primary goal, serves two convergent objectives: firstly, the understanding of the cellular pathway the receptor is integrated into and its role in the regulation of a particular physiological process and secondly, it gives information on its possible therapeutic value. The most straightforward way to identify ligands at oGPCRs is to ectopically express them in a cellular system and to measure dependency of ligand-induced modulation of a signaling pathway on the presence of the receptor in quest (Fig. 1). In this strategy, the selected receptor is used as bait to identify ligands. Prior to ligand screening, several preliminary studies are usually performed or taken into account: 1) receptor loss-of-function phenotyping, 2) receptor bioinformatics and 3) tissue expression analyses.

1) Loss-of-function studies by genetic deletion or mutational analysis can indicate whether an oGPCR plays an important role in the regulation of a physiological process. Several examples support the idea of genetic studies as valuable preliminary steps in the identification of potentially relevant oGPCRs: for example, mutations in the GPR101 gene, which is strongly expressed in the hypothalamus (13), have been associated with gigantism and acromegaly (14). In the case of GPR149, which shows expression in granulosa cells and oocytes, the gene-deficient mice showed increased fertility and enhanced ovulation (15). Disruption of another orphan receptor, GPR45, caused hypothalamic proopiomelanocortin (POMC) expression and obesity in mice (16) whereas mice lacking GPR21 receptor, which shows expression in the adipose tissue and liver, were resistant to diet-induced obesity and had increased glucose tolerance and insulin sensitivity (17). A compelling database of GPCR genes and known associated phenotypes can be found at IUPHAR database: http://www.guidetopharmacology.org.

2) Bioinformatics analysis can be used to compare sequences of oGPCRs with those of the liganded receptors to predict cognate ligands based on sequence homology. For example, the orphan receptor GPR105 (known as P2Y14) which showed significant homology with other nucleotide-liganded purinergic receptors was screened against a library containing nucleotide di- and triphosphate conjugates to identify UDP-glucose as ligand (18). Another example was the identification of sphingosine-1-phosphate (S1P) as a potent ligand of EDG-1 (S1P1) receptor (15) a finding which led to the subsequent discovery of multiple receptors (S1P1 to S1P5) receptors for S1P, based on sequence homology analyses (19-21). Furthermore, due to the success of resolving the β1- and β2-AR receptors crystal structures (22), followed by the X-ray structures of 20 receptors in the following years, structure-based drug design (SBDD) can be an effective tool to design potential agonists or antagonists (23, 24). Several examples where SBDD in conjuction with computer modelling has been successfully employed to find new ligands at known or orphan GPCRs are the identification of alloseric modulators and agonists of the proton receptors GPR68 and GPR65 (25). Another remarkable example with potential clinical relevance is the discovery through SBDD of a novel µ-opioid receptor agonist which selectively drives Gi/o-mediated signaling while minimizing β-arrestin-dependent signaling pathways (26). Furthermore, 3D homology modelling of GPCRs using as templates crystal structure of numerous GPCRs represents an important tool for ligand prediction and identification and it has been already succesfully used in the past, for example, in the identification of a selective agonist of M1 acetylcholine receptor (27).

3) Tissue expression analysis of an oGPCR can provide valuable information about its potential biological role.
The expression data are usually combined with genetic studies to acquire more information of the receptor's biological importance. Compelling expression studies of GPCRs in mouse (including oGPCRs) have been made (28, 29). Other resources exist, such as the Allen Brain Atlas (http://www.brain-map.org/).

Once the receptor is chosen based on its potential biological relevance, deorphanization process requires the application of two technologies: a) generation of a suitable compound/metabolite library to be screened and b) measuring receptor activation following ligands stimulation.

a) Generation of a suitable and complex ligand library can sometimes be challenging due to the limited availability and high costs of some compounds of interest. However, in the last years a number of companies provide ready-to-use or even custom-designed libraries. If the library is to be custom-designed, the selection criteria of the substances and/or metabolites are usually based on public information on described effects of metabolites.
One advantage of the library-based approach is that the purified compounds can be tested at relatively high concentrations. By screening ligand libraries, a large number of receptors were deorphanized such as receptors for short-chain fatty acids (30) or the receptor for polyunsaturated fatty acids (PUFAs) expressed in the pancreatic islets, known as FFAR1 (GPR40), which enhances glucose-induced insulin secretion after PUFAs stimulation (31, 32). A complementary approach to library-based strategy is to test fractions from extracts of tissues/organs where the receptor is highly expressed followed by the isolation and purification of the endogenous ligand(s). Examples are the identification of novel neuropeptides from tissue fractions and their receptors, especially those involved in the regulation of food intake and behavior such as orexin and orexin receptors (33) and the melanin-concentrating hormone (MCH) and its receptor, MCH1 (34, 35).

Orexin and MCH systems are also orexigenic, as genetic inactivation of one of the receptors or peptide ligands led to lean phenotypes in mouse. However, the use of tissue/organ extract as a source of endogenous ligands can be complicated by the fact that some ligands act in a paracrine manner, being synthesized in a different organ than their site of action. An example is the anorexigenic peptide ghrelin which is produced in the stomach and functions as an endogenous ligand of the growth hormone secretagogue (GHS) receptor expressed primarily in the pituitary gland and the hypothalamus (36, 37).

(b) Determination of receptor activation after stimulation with ligands or tissue fractions relies on canonical intracellular signaling pathways modulated by heterologously expressed GPCRs and G-proteins (31). However, one of the first challenges the investigator has to face is to ensure that the overexpressed receptor to be studied traffics to the plasma membrane. It is well known that some GPCRs, when ectopically expressed in one of the standard reporter cell lines such as COS-1, HEK293 or CHO fail to reach plasma membrane because of a lack of an interaction partner such as receptor activity-modifying proteins (RAMPs) (39, 40).

Methods to determine the activity of a GPCR rely on the G-protein-promoted activation of different intracellular signaling cascades after receptor activation and typically include: (i) GTP\(\gamma\)S binding assay on membranes of cells expressing an oGPCR; (ii) determination of intracellular calcium transients as a result of receptor activation; and (iii) modulation of intracellular cAMP levels after activation of Gs- and Gi-coupled receptors.

(i) GTP\(\gamma\)S binding assay measures the guanine nucleotide exchange of G\(\gamma\) subunit (GDP to GTP\(\gamma\) exchange) as it is the earliest event of the activated receptor. Thus, it is not subject to amplification by other cellular processes providing “true” determinations of the efficacies and potencies of a ligand. GTP\(\gamma\)S binding measures the accumulation of a non-hydrolysable analog of GTP, \[^{35}\text{S}\text{-GTP}\(\gamma\) on the membrane of cells expressing a receptor. The main advantage of this assay is that it allows discriminating between agonists, antagonists and inverse agonists (an agonist which decreases the basal activity of a constitutively active receptor). GTP\(\gamma\)S binding assay depends also on the relative abundance

<table>
<thead>
<tr>
<th>Trade Name</th>
<th>GPCR Target</th>
<th>Effect</th>
<th>Therapeutic Applications</th>
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<tbody>
<tr>
<td>Albuterol</td>
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<td>agonist</td>
<td>asthma</td>
</tr>
<tr>
<td>Alprenolol</td>
<td>adrenergic-(\beta_1,\beta_3/\beta_1A)</td>
<td>antagonist</td>
<td>angina pectoris</td>
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<td>Amthamine</td>
<td>histamine-(H_3)</td>
<td>agonist</td>
<td>gastric dysfunctions</td>
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<td>Baclofen</td>
<td>gamma-aminobutyric-GABAB</td>
<td>agonist</td>
<td>skeletal muscle relaxant</td>
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<tr>
<td>Clonidine</td>
<td>adrenergic-(\alpha_1)</td>
<td>agonist</td>
<td>high blood pressure; anxiety, migraine</td>
</tr>
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<td>histamine-(H_1)</td>
<td>antagonist</td>
<td>allergy</td>
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<td>antagonist</td>
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<td>hypertension</td>
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<td>(\mu)-opioid</td>
<td>agonist</td>
<td>pain</td>
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<td>HCAR(_2)</td>
<td>agonist</td>
<td>antiatherogenic</td>
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</tr>
<tr>
<td>Ricinoleic acid</td>
<td>prostaglandin (E_2-\text{EP}_3)</td>
<td>agonist</td>
<td>laxation; labour</td>
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Table 1. Example of approved drugs targeting GPCRs and their therapeutic applications.
of endogenously expressed G-proteins. As a results, it is more suitable for studying Gi/o-coupled receptors due to the higher cellular levels of Gi/o proteins (41). Beside the need of a radioactive tracer, a disadvantage of this method is the use of a final filtration step to separate bound from free \[^{35}S\]GTP, thus limiting its high-throughput capabilities. However, this assay has been successfully used to identify ligands at oGPCRs (42). To address some of these issues, Perkin Elmer has released a non-radioactive time-resolved fluorescence (TRF) GTP-binding assay using europium-labeled GTP (DELFIA™). The fluorescence recorded is directly proportional with the amount of Eu-GTP accumulated in membranes from cells expressing a receptor and thus reflects receptor activation. Although it still requires the washing step, it has been successfully validated on a number of receptor targets such as serotonin 5-HT\(5a\), \(\alpha\)2-adrenergic, dopamine and muscarinic receptors (43).

(ii) Intracellular calcium assay is one of the most versatile and cost-efficient screening methods used in reverse pharmacology for receptor deorphanization and characterization. It relies on the property of Gq-coupled receptors to activate an intracellular signaling pathway which includes activation of phospholipase C, isoform \(\beta\) (PLC\(_{\beta}\)) with generation of diacylglycerol (DAG) and inositol triphosphate (IP\(_3\)) which in turn triggers the release of endoplasmic-reticulum (ER)-stored calcium by binding to IP3 receptors. The basis of this assay is the heterologous expression of a target receptor in a cell line and determination of \([Ca^{2+}]_i\) by specific probes which can be chemical (fluorescent) or genetic. Chemical fluorescent calcium indicators have been used for decades to measure free intracellular \(Ca^{2+}\) concentrations in cells. They are cell permeant acetoxyethyl (AM) esters such as Fura-2/AM, Fluo-4/AM to Fluo-8/AM generation of indicators (44). Their main advantage is that they do not require being transfected. It is the method of choice for fluorometric imaging plate readers (FLIPR)-based assays used in the pharma industry where large compound libraries are screened. The main disadvantage of the fluorescent calcium indicators is the lower signal-to-noise ratio and their tendency to mislocalize to subcellular compartments. In contrast to chemical probes, calcium-sensitive genetic probes offer several advantages over fluorescent probes. They are based on the calcium sensitive photoprotein, aequorin, which is expressed in the jellyfish *Aequorea victoria*. One of the best probes consists of a fusion protein between aequorin and the green fluorescent protein (GFP) separated by a short linker (45). The main advantage over chemical fluorescent probes is the absence of a significant background and the very high signal-to-noise ratio. In this assay, a receptor of interest is transfected together with the genetic probe and 48 h later the functional aequorin is reconstituted by the addition of a cofactor, coelenterazine h. The functional assay is usually performed in a luminometer equipped with ligand dispensing capabilities in a 96, 384 or 1536 well format. This type of assay has been successfully used to find a large number of ligands and agonists of orphan and known GPCRs (46).

(iii) cAMP assay is suitable for measuring activated Gs- and Gi-coupled receptors which are known to either increase (Gs) or decrease (Gi/o) intracellular cAMP concentration after activation by a ligand. To determine the activation of Gs-coupled receptors, cells overexpressing a receptor are stimulated with the candidate compound and the intracellular accumulation of cAMP is determined 15 minutes later. To determine the activation of Gi-coupled receptors, cells need to be pre-stimulated with the ligand of interest for 5 minutes followed by stimulation with a Gi-independent activator of AC, forskolin and determination of cAMP levels 15 minutes later. Although forskolin works well in most of the cases, a more physiological approach would be to use a GPCR-mediated cAMP increase induced by prostacyclin, PGE2, or isoproterenol which activate ACs through Gs-coupled receptors endogenously or heterologously expressed. There are numerous methods to determine intracellular cAMP concentrations but the most reliable are ELISA (commercially available) and newly introduced genetic probes, such as GloSensor™, 22F CAMP from Promega. The genetic probes offer great advantage due to the low costs of the assay and also because they allow real-time visualization of cAMP accumulation in living cells.

The selection of the appropriate functional assay during the preparation of a screening campaign is a critical step. Needless to say, in the case of oGPCRs, no information is in general available as to what type of G-protein the receptor is coupled to. An alternative approach, with very good results, is to use the property of the promiscuous Gx15 and Gx16 subunits to enforce Gi- and Gs-coupled receptors to signal via PLC\(_{\beta}\)-IP\(_3\)-Ca\(^{2+}\) pathway (47, 48). In this type of assay, reporter cells are transfected with the receptor, the promiscuous \(\alpha\) subunits G15 or 16 and the calcium sensitive probe. It is a very versatile method which has been successfully used to identify ligands of Gi- and Gs- and G12/13-coupled receptors (49). Another way to elucidate signaling properties of an oGPCR with unknown G-protein coupling is to overexpress it in a chosen cell line and measure ligand-independent activation (constitutive activity) of specific signaling pathways. This phenomenon has been described to be a result of the receptor overexpression which leads to constitutive activity and it has been successfully used to identify intracellular signaling characteristics for receptors that induce, for example, cAMP increase after their overexpression (50, 51).

\(\beta\)-arrestins are intracellular scaffold proteins that interact with the activated receptor to promote receptor desensitization (52). Due to their high interaction...
specificity, β-arrestins can be used as probes to measure receptor activation following ligand stimulation, in a G-protein independent manner. An elegant way of measuring GPCR activation by using β-arrestins translocation has been recently developed and successfully used to discover novel ligands of oGPCRs (53, 54).

Although reversed pharmacology was successfully employed to discover endogenous ligands and agonists of many GPCRs, there are cases where its limits are evidenced due to a different pharmacology and/or biological role of certain receptors. Some of the orphan receptors may simply have ligand-independent functions. Several examples include GABA<sub>B1</sub>, GPR50 and the constitutively active receptors.

The metabotropic gamma-aminobutyric acid B (GABAB) receptor which is expressed in the brain, is an obligatory heterodimer composed of two GABA<sub>B1</sub> and GABA<sub>B2</sub> subunits. Interestingly, GABA<sub>B1</sub> subunit provides GABA-binding site whereas GABA<sub>B2</sub> facilitates trafficking of GABA<sub>B1</sub> to the cells surface and is required for G-protein coupling but is unable to bind GABA (55).

GPR50 is an orphan receptor predominantly expressed in the brain and it shares structural homology with MT1 and MT2 melatonin receptor subfamily which are activated by the circadian hormone melatonin. Although GPR50 forms <i>in vitro</i> heterodimers with both MT1 and MT2, only interaction with MT1 appears to have functional effects as GPR50 impairs MT1-induced activation of adenyl-cyclase and subsequent cAMP-dependent pathway. Interestingly, GPR50/MT1 heterodimer completely inhibits high-affinity melatonin binding and receptor-induced G-protein activation (56).

Constitutively activated receptors are receptors which are able to induce the activation of specific G-protein-mediated signaling pathways in the absence of a ligand (57). Thus they are thought to be still orphan receptors and have ligand-independent functions. Constitutive signaling of some GPCRs seems to be induced by mutations which are also associated to various pathological states (58-60) and therefore this pathway is under intense pharmacological surveillance. For example, activating mutations in the thyroid stimulating hormone (TSH) receptor, which plays critical roles in thyroid gland growth and functions, cause toxic adenoma as well as familial and sporadic nonautoimune hyperthyroidism (61).

FORWARD PHARMACOLOGY

Forward pharmacology (classical pharmacology or phenotypic drug discovery) is based on the phenotypic screening of libraries consisting of natural or synthetic compounds to identify those which have a desirable effect. Historically, forward pharmacology precedes reverse pharmacology in the sense that the biological effects of numerous substances were discovered long before even the receptor concept was proposed. For example, as early as in the 19<sup>th</sup> century, pilocarpine and atropine were thought to “compound” with a physiological substance present in the target organ to trigger observed effects (such as salivary secretion in dogs). With the advance of metabolomics, the number of metabolites identified in body fluids steadily increased to over 5000. Many of them have unknown biological roles and may even serve as ligands of known or orphan GPCRs. A comprehensive list of metabolites and their description can be found at http://www.hmdb.ca/. Another useful approach to generate a portfolio of putative ligands of a GPCR relies on computer-based virtual ligand screening (VLS) using GPCR’s crystal structure as template, when available (62). Structure-based VLS has been successful in identifying new ligands of GPCRs (63).

The central task of the forward pharmacology in GPCR deorphanization is the identification of the mechanism of action of a ligand which already exist solid evidences of its biological role. As opposite to reverse pharmacology, in this case a ligand is used to find the cognate receptor. In its simplest way, a collection of receptors is individually expressed in a reporter cell line and ligand-induced activation of each receptor is determined and compared to the effect on cells that do not express the receptor. The selection of the receptors is usually based on the information regarding the expression of receptors in the tissues where the ligand is active. A more complex strategy in the identification of a receptor for a particular ligand starts with finding a cellular system where the metabolite/compound is able to activate a GPCR-mediated pathway (Fig. 1). It is then followed by pharmacological analyses of the cellular effect of the ligand at different concentrations (concentration-dependency), its sensitivity to several inhibitors of G-proteins and, where possible, determination of stereospecific effects. All these experiments conclude with a pharmacological profile of the candidate ligand and it is known as ligand “validation”. The next step is the discovery of the putative receptor. A number of considerations have to be taken into account at this stage. First of all, from all of the above experiments there is no indication that only one receptor mediates the observed cellular effects. Secondly, it cannot be excluded a more complex signaling pathway involving a GPCR which is necessary but not critical in the mechanism of action of the metabolite/compound. Lastly, the ligand can induce the observed effects after being metabolized to another active species which in turn can activate a GPCR (secondary effect). In order to identify the cognate receptors several converging strategies may be employed.

Expression analysis of all known and predicted GPCRs in the cell type where the ligand is active can provide very useful information. Based on the G-protein coupling studies in the ‘validation’ step, the list of potential receptors can be significantly narrowed down. For example, if the
ligand induces cellular cAMP accumulation, the candidate receptors to be further tested are the ones which are known to be Gs-coupled. However, as in the case of the unknown intracellular coupling of the oGPCRs, individual testing of each oGPCR is the safest approach. On the other hand, there is a very high chance that the ligand activates a signaling pathway only in one cell type as it is inactive in others. Expression analysis can be effectively used in this situation to compare receptors expressed in the responsive cells with those in the cells the ligand was inactive. Differential expression analysis can be a powerful tool, especially if more than one cell type is analyzed for responsiveness to a ligand and can be expanded for a portfolio of metabotolites/compounds. Such a strategy has been used to identify the receptor which mediates cAMP accumulation in a human megakaryocytic leukemia cell line (MEG-01) and vasorelaxation of arterial segments from mouse induced by the arachidonic acid metabolite, 19-hydroxyeicosatetraenoic acid, 19(S)-HETE (64).

Another efficient strategy to identify the cognate receptor for a ligand is to determine its effect in cells transfected with siRNA targeting individual known and predicted GPCR mRNAs (functional siRNA screening). If the ligand has a significantly lower effect in cells transfected with siRNA against mRNA of a particular GPCR, then this receptor becomes a strong candidate as mediator of the observed cellular effects. Confirmation of the candidate receptor should be performed by expressing candidate receptors in a reporter cell line and determination of receptor activation by the ligand. Although a powerful method, functional siRNA screening requires intense preparative steps. For instance, it is useful to choose the siRNA library/assay conditions that efficiently allow the “re-discovery” of a known receptor by using its endogenous ligand/agonist. As an example, U46619, an agonist of thromboxane A2 (TXA2R) receptor (65), can be tested on responsive cells to determine whether after screening on cells transfected with the siRNA library only siRNAs against TXA2R can strongly reduce the effect of U46619. Accordingly, several other known ligands, activating distinct intracellular signaling pathways should be analyzed before entering a large deorphanization campaign with an unpaired ligand. As proof-of-principle example, by applying this technology it was shown that ricinoleic acid, which is a major component of castor oil used for millennia for its laxative and labor-inducing effects, induces pharmacological effects such as intestinal and uterus contraction by acting as a ligand of prostaglandin E2 receptor, subtype EP3 (66).

CONCLUSIONS

In a practical sense, reverse and forward pharmacology are not just separated strategies. For example, a receptor can be deorphanized by forward pharmacology and then the reverse pharmacology can be used to further discover agonists and antagonists. An example can be the identification of a family of receptors involved in nutrient sensing by binding hydroxy-carboxylic acids resulted from β-oxidation of fatty acids (67).

Although the deorphanization of oGPCRs by reverse and forward pharmacology has fundamentally contributed to our understanding of pathophysiology, it is worth noting that the peak of ligand-receptor pairing has been reached in 2003. After that, the deorphanization rate has steadily declined. A reason for the lower rate of deorphanization in the last years may be the increasing costs of ligand screening and the likelihood of the ligand-identification failure. Although pharma companies face the expiration of patents for blockbuster drugs and thus are forced to find new drug targets, they also face increase safety rules and regulations resulting in a high number of leads that fail to reach the higher phases in drug approval. Even if an important receptor is deorphanized and modulators of its activity are of therapeutic importance, it would take many years and very high costs to reach final phases of drug approval. Thus, the interest of the major companies appears to be shifted back towards known targets where development of new compounds with a better pharmacology may be more beneficial. Still, this leaves academia an important role to try understanding the biological function of the remaining oGPCRs.

Another explanation for the decrease rate of deorphanization is that many oGPCRs are phylogenetically distant from known GPCRs, suggesting that they might have completely new ligands and biological roles. From an academic point, this is of the utmost interest. Furthermore, it is also possible that classic deorphanization assays, which rely on the G-protein-dependent signaling pathways, may fail to uncover oGPCRs that can signal in a G-protein-independent manner (68).

The deorphanization process is not always flawless. Initial identification claiming successful oGPCR-ligand pairing requires confirmation by independent groups thus leading to possible controversies sometimes persisting for several years. Several examples can be used to learn more about the difficulties encountered during receptor deorphanization. Such examples include GPR68 (OGR1) and GPR4 which have been initially described as a lipid receptors but could not be confirmed by others, data being thus retracted (69). Another interesting example is GPR30 where controversial features include its subcellular localization and its capacity to bind estrogen thus supporting the notion of non-genomic effects of steroids mediated by GPCRs. Despite intense research in the last years, no clear consensus has been reached so far (70-72).

GPCR heterodimerization has been described as another structural feature with signaling and pharmacology consequences (73, 74). It would be interesting to choose
oGPCRs that show overlapping expression patterns and identify possible oGPCR heterodimers which could be further exploited in screening campaigns. On the other hand, the recent development of metabolomics resulted in the identification of a large number of new metabolites which should also be included in the compound libraries to be tested for potential agonistic/antagonistic effects on known or orphan GPCRs.

The success in receptor-ligand pairing in the last years required the convergence of at least three scientific areas: genomics, metabolomics and cellular biology, through the development of versatile high-throughput screening methods. More challenges as well as interesting questions and answers lie ahead as still a large number of oGPCR await pharmacological characterization.

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G-protein coupled receptor deorphanization


