Nonclassical Ligand-Independent Regulation of Go Protein by an Orphan Class C G-Protein–Coupled Receptor

Mariana Hajj, Teresa De Vita, Claire Vol, Charlotte Renassia, Jean-Charles Bologna, Isabelle Brabet, Magali Cazade, Manuela Pastore, Jaroslav Blahos, Gilles Labesse, Jean-Philippe Pin, and Laurent Prézeau

Institute of Functional Genomics (M.H., T.D.V., C.V., C.R., J.-C.B., I.B., M.C., M.P., J.-P.P., L.P.) and CBS (G.L.), University of Montpellier, French National Center for Scientific Research (CNRS), French National Institute of Health and Medicine (INSERM), Montpellier, France; and Institute of Molecular Genetics, Academy of Sciences of the Czech Republic and Department of Pharmacology, 2nd Medical School, Charles University, Prague, Czech Republic (J.B.)

Received May 16, 2018; accepted May 28, 2019

ABSTRACT

The orphan G-protein–coupled receptor (GPCR) GPR158 is expressed in the brain, where it is involved in the osteocalcin effect on cognitive processes, and at the periphery, where it may contribute to glaucoma and cancers. GPR158 forms a complex with RGS7–β5, leading to the regulation of neighboring GPCR-induced Go protein activity. GPR158 also interacts with αo, although no canonical Go coupling has been reported. GPR158 displays three VCPWE motifs in its C-terminal domain that are putatively involved in G-protein regulation. Here, we addressed the scaffolding function of GPR158 and its VCPWE motifs on Go. We observed that GPR158 interacts with and stabilized the amount of RGS7–β5 through a 50-residue region downstream of its transmembrane domain and upstream of the VCPWE motifs. We show that two VCPWE motifs are involved in αo binding. Using a Gαo–βγ bioluminescence resonance energy transfer (BRET) sensor, we found that GPR158 decreases the BRET signal as observed upon G-protein activation; however, no constitutive activity of GPR158 could be detected through the measurement of various G-protein–mediated downstream responses. We propose that the effect of GPR158 on Go is unlikely due to a canonical activation of Go, but rather to the trapping of Gαo by the VCPWE motifs, possibly leading to its dissociation from βγ. Such action of GPR158 is expected to prolong the βγ activity, as also observed with some activators of G-protein signaling. Taken together, our data revealed a complex functional scaffolding or signaling role for GPR158 controlling Go through an original mechanism.

Introduction

G-protein-coupled receptors (GPCRs) are key players in cell-cell communication and the protein family most targeted by commercial drugs (Overington et al., 2006). GPCRs couple to heterotrimeric (αβγ subunits) G proteins that control the activity of membrane and intracellular effectors. GPCRs behave as G-protein nucleotide exchanging factors to promote the GTP-bound G-protein α subunit active state, generally considered dissociated from βγ. Furthermore, G-protein α and βγ activities can be decreased by other signalings, such as regulators of G-protein signaling (RGS) (Gerber et al., 2016) or prolonged by activators of G-protein signaling (AGS) (Blumer and Lanier, 2014). GPCRs also mediate their action through G-protein–independent pathways (such as those involving arrestins), signaling cross-talk (Prezeau et al., 2010), receptor transactivation (Milligan, 2006), or association with specific signalosomes (Bockaert et al., 2010). Thus, GPCR signaling pathways must be functionally organized to integrate so many regulatory inputs.

Cellular and physiologic functions of the orphan receptor GPR158 are largely unknown. It is expressed in the brain (Orlandi et al., 2015), where it likely regulates neuron excitability as it has been “fished out” as a partner of potassium Kv4.2 and calcium Cav2 channels (Marionneau et al., 2009; Müller et al., 2010). In the CA3 region of the hippocampus, GPR158 has recently been reported to be involved in the cognitive actions of osteocalcin (Khrimian et al., 2017) and to control the presynaptic differentiation of mossy fiber-CA3 synapses by interacting with proteoglycans of the extracellular matrix (Condomitti et al., 2018). GPR158 could also be involved in the shaping of retinal...
photoreceptor signaling (Orlandi et al., 2012). Its expression is regulated by glucocorticoids in trabecular meshwork cells, a mechanism possibly involved in the occurrence of glaucoma (Patel et al., 2013). Although its role in prostate cancer emerged recently (Patel et al., 2015), the impact of GPR158 mutations in colorectal cancers and leukemia must still be confirmed (Wood et al., 2007; Greif et al., 2011). Together, these data suggest important regulatory roles in cellular processes, highlighting the real necessity to understand GPR158’s signaling roles in healthy and disease cells.

Potential signaling functions for GPR158 are supported by the observation that GPR158 can interact with the RGS7 protein and the G-protein αo subunit (Orlandi et al., 2012, 2015). Indeed, GPR158 brings RGS7 to the plasma membrane, where RGS7 can allosterically accelerate the GTPase activity of Goα and turn it off when Goα is activated by neighboring GPCRs. Three short VCPWE motifs are present in the last third of the C-terminal domain of GPR158. Although they resemble the ICPWE motif of the retinal phosphodiesterase (PDE) γ subunit known to mediate the interaction with R7 family RGS proteins and active G-protein α subunits (Slep et al., 2001), they were not involved in RGS7 interaction with GPR158 (Orlandi et al., 2015). Thus, whereas RGS7 would interact in the first third of the C-terminal domain of GPR158, Goα may bind on two sites, one in the same region as RGS7 and another in the second half of the C-terminal domain. Surprisingly, the role of the VCPWE motifs remains unclear. Furthermore, GPR158 possesses residues signatures reported to be important for G-protein coupling of other GPCRs (Bjarnadottir et al., 2005), but the ability of GPR158 to couple to G proteins in a canonical way has not been addressed either. Thus, it seems that GPR158 plays important roles in the absence of ligand, notably through its association with RGS7. The question remains of what the scaffolding role of GPR158 toward RGS7 and Goα could be in the absence of ligand.

Here we identified the interaction site of RGS7 in the 714–764 region of GPR158 C-terminal domain and confirmed that the three VCPWE motifs are not involved in this interaction. In contrast, these motifs were required for Goα association. We also show that GPR158 likely induced Goα-βγ dissociation as measured by a BRET assay, reflecting possible activation by the receptor; however, we found no evidence of canonical coupling of WT and GPR158 mutants to G protein under basal condition using second-messenger functional assays. Indeed, the GPR158-induced increased levels of dissociated Goα is possibly due to trapping of GPR158 by the VCPWE motifs. By trapping Goα, GPR158 is expected to prolong the action of βγ, as observed with some group II AGS proteins. These data, which do not exclude the possible direct G-protein activation upon ligand activation of GPR158, unravel new ways for GPCRs to locally regulate G-protein pathways.

Materials and Methods

Compounds. All compounds and reagents were purchased from the most appropriate sources and companies.

Plasmids and Site-Directed Mutagenesis. To generate C-terminal Flag- or HA-tagged GPR158 constructs, HA or Flag tag sequences were introduced downstream of GPR158 coding region after a unique MluI restriction site, followed by a stop codon. To generate N-terminal tagged HA- and Flag-GPR158 constructs, MluI and XbaI restriction sites were inserted by polymerase chain reaction after the N-terminal peptide signal and after the C-terminal stop codon of the GPR158 coding sequence, respectively. The resulted MluI-XbaI fragment was then inserted into pRK-HA and pRK-Flag RAB4A (OB1) plasmids (Kniazeff et al., 2004; Rives et al., 2009) digested using the MluI and XbaI sites, which excised the RAB4A coding sequence. We generated HA-Snap-GPR158 and Flag-Clip-GPR158 by inserting the sequence encoding Snap tag and Clip tag of HA subunit of HA-GPR158 and Flag-GPR158, respectively, using a Quick-Change strategy (Stratagene, San Diego, CA). The RGS7 (s2 form) and β5 plasmids were purchased from (UMR cDNA Resource Center, MO). The HA tag was introduced at the C-terminal end of the coding sequence of RGS7 upstream of both a XhoI restriction site and a stop codon using a Quick-Change strategy. To generate RAB4A-HA, the HA tag sequence was inserted at the C-terminal end of the RAB4A coding sequence between a XhoI site and the stop codon using a Quick-Change strategy. Chimeras formed by exchanging domain between metabotropic glutamate (mGlu1a and GPR158 or mGlu2 and GPR158 were generated using polymerase chain reaction overlap strategies, and point mutations in GPR158 sequence were generated using the Quick-Change strategy. All final constructs were verified by sequencing (MWG, Ebersberg, Germany).

Cell Culture and Transfection. Human embryonic kidney 293 cell cultures (from American Type Culture Collection) were cultivated in Dulbecco’s modified Eagle’s medium (Invitrogen Life Technologies, Gaithersburg, MD), supplemented with 10% fetal calf serum (Sigma, Lisle-D’Aube, Saint-Quentin Fallavier, France), were checked each month for mycoplasma. The cells were seeded in 100-mm plates and incubated at 37°C in a CO2 incubator. Cells were transiently transfected using LipofectAMINE 2000 according to the manufacturer’s protocol (Thermo Fisher Scientific, Waltham, MA). Cells were seeded in 96-well plates (Greiner Bio-One, Frickhausen, Germany) at 50,000 or 100,000 cells per well. Alternatively, cells were transfected by electroporation as previously described (Maurel et al., 2004). Ten million cells were electroporated with indicated plasmids containing the coding sequence of the proteins of interest and completed to a total amount of 10 μg of plasmid DNA with pRK6 empty vector before being plated in 96-well plates. All media used for cell culture were purchased from Life Technologies/Thermo Fisher Scientific.

Extracellular and Intracellular Antibody Time-Resolved Förster (TRF) Resonance Energy Transfer Assay (Homogeneous TRF). Based on a luminescence resonance energy transfer technology, time-resolved Förster resonance energy transfer (TR-RET) experiments using labeled antibodies were performed in 96-well plates in homogeneous conditions (HTRF), as previously described (Maurel et al., 2008). Twenty-four hours after transfection, cells were washed with cold Tris-Krebs buffer (20 mM, Tris pH 7.4, 118 mM NaCl, 5.6 mM glucose, 1.2 mM KH2PO4, 1.2 mM MgSO4, 4.7 mM KCl, and 1.8 mM CaCl2) and incubated overnight at 4°C with antibodies, 3 nM, and d2- labeled antibody, 6 nM) (Cisbio Bioassays, Codolet, France). The cells were then incubated for 5 minutes at room temperature with KF (200 mM). The fluorescence of the europium cryptate–labeled antibody, 3 nM, and d2- labeled antibody, 6 nM) (Cisbio Bioassays, Codolet, France). The cells were then incubated for 5 minutes at room temperature with KF (200 mM). The fluorescence of the europium cryptate–labeled antibody, 3 nM, and d2- labeled antibody, 6 nM) (Cisbio Bioassays, Codolet, France).
Cell-Surface Protein-Level Quantification by Enzyme-Linked Immunosorbent Assay and SNAP Labeling. Enzyme-linked immunosorbent assay (ELISA) experiments were performed as previously described (Maurel et al., 2008). Briefly, cells were fixed with 4% paraformaldehyde and blocked with phosphate-buffered saline (PBS) containing 1% fetal calf serum and then incubated 30 minutes at 0.5 mg/liter with monoclonal anti-Flag M2 antibodies (Sigma), anti-HA antibodies (clone 3F10; Roche Applied Science, Basel, Switzerland) or anti-Myc antibodies (clone 9E10; University of Iowa, Iowa City, IA). When these primary antibodies were not conjugated themselves with horseradish peroxidase (HRP), they were further washed and incubated (30 minutes) with HRP-conjugated goat anti-rat IgG (0.5 mg/liter; Jackson ImmunoResearch Laboratories, West Grove, PA) or anti-rabbit IgG or anti-mouse IgG (0.5 mg/liter; Amersham Biosciences GE Healthcare, Chicago, IL) for 30 minutes. After washes, bound antibody was detected by chemiluminescence using SuperSignal substrate (Pierce, Rockford, IL) and a Mithras LB 940 plate reader (Berthold Technologies, Bad Wildbad, Germany). As a control for intracellular ELISA quantification, cells were permeabilized for 5 minutes with 0.05% Triton X-100 just after being fixed, SNAP-tag labeling was performed as described previously (Doumazane et al., 2011). Briefly, 24 hours after transfection, human embryonic kidney 293 cells (HEK293) cells were incubated at 37°C for 1 hour with a solution of 100 nM of Lumi4-Tb (Cisbio Bioassays). After labeling, the cells were washed three times with Krebs buffer, and drugs were added as described. TR-FRET measurements were performed on INFINITE 500 (TECAN, Mannele, Switzerland) or PHERAstar FS (BMG Labtechnologies) microplate readers, which are equipped with standard TR-FRET optical modules.

Communoprecipitation and Western Blotting. At 48 hours after transfection in 100-mm plates, cells were washed with ice-cold PBS-GAB (PBS supplemented with glucose and antibiotics). After washes, cells were scraped with PBS-GAB and centrifuged for 5 minutes at 2000 rpm. The supernatant was removed, and the pellet was resuspended in lysis buffer (Hepes 1 M, NaCl 5 M, NP40 20%, glycerol, dodecyl maltoside, and protease inhibitors cocktail (Sigma)). The lysate was incubated for 2 hours at 4°C with mild shaking and then clarified by centrifugation at 15,000g for 15 minutes. Clarified lysate was incubated with monoclonal anti-HA conjugate agarose beads, (Clone HA-7; Sigma) overnight at 4°C and then centrifuged for 2 minutes at 13,000 rpm. The supernatant was removed, and the pelleted beads were washed four times with PBS 1× before elution by addition of loading buffer. The samples were loaded on NuPAGE MES buffer (Amersham Biosciences GE Healthcare, Chicago, IL) and subjected to immunoblotting.

For the primary rabbit anti-HA (Invitrogen Life Technologies) antibody was used at 0.6 mg/liter and the mouse anti-Flag antibody (F3165; Sigma) at 2 mg/liter. Secondary antibodies anti-rabbit HRP-linked IgG (0.5 mg/liter; Amersham Biosciences GE Healthcare) and anti-mouse HRP-linked IgG (Santa Cruz, Dallas, TX) were applied for 30 minutes. Immunoreactive bands were visualized by ECL detection kit (Amersham Biosciences GE Healthcare) on Kodak ML light films.

IP-One and eCAMP Assays. Experiments were performed in a 96-well plate format. The IP-One HTRF kit (Cisbio Bioassays) was used according to the recommendations of the manufacturer to measure the production of inositol phosphate second messenger (IP3) through assessment of inositol monophosphate (IP1) accumulation, a downstream metabolite of IP3. Cells were incubated in the presence of indicated receptor agonist for 30 minutes at 37°C and then incubated in the presence of a cryptate-labeled anti-IP1 or eCAMP antibodies and D2-labeled IP1 or CAMP for 1 hour at room temperature. The fluorescence of the europium cryptate and d2, 620, and 665 nm respectively, was measured (without washing) 50 micro-seconds after excitation at 337 nm using RubyStar or PHERAstar plate readers (BMG Labtechnologies).

BRET Experiments. As previously described (Ayoub et al., 2007) for saturation curves, a constant amount of a plasmid encoding one of the proteins of interest bearing the donor-Luc was coexpressed with a range of expression of the plasmid encoding the second protein of interest bearing the acceptor-YFP. After washing of the cells, the Luciferase substrate Coelenterazine h (cat. no. C-6780 Invitrogen/Thermo Fisher Scientific) was added (5 mM in 50 µl per well) to initiate the BRET process. Readings were recorded using the Mithras LB940 reader (Berthold Biotechnologies) (Rluc filter: 485 ± 20 nm and YFP filter: 530 ± 25 nm), data were collected using the MicroWin2000 software, and BRET signal expressed in milliBRET units of BRET ratio. Dose-response curves were fitted with a linear regression or sigmoid dose-response equation, using Prism (GraphPad software, San Diego). The Go, experiments were performed as previously described (Rives et al., 2009). The Go activation/dissociation BRET assays were performed as described previously (Brulé et al., 2014). The transfected HEK293 cells were washed with PBS, and readout was performed on a Mithras LB940 plate reader (Berthold Technologies) at 37°C after the addition of coelenterazine h (5 µM) and ligand. When indicated, treatment with pertussis toxin (100 ng/ml) was performed for 16 hours before stimulation of the cells. The BRET ratio was calculated based on the difference between the emission at 530/485 nm of co-transfected Rluc and YFP fusion proteins and the emission at 530/485 nm of the Rluc fusion protein alone.

Data Analysis and Statistical Tests. Data were analyzed using Prism 7.0e (GraphPad) or Excel 16.16.8 (Microsoft), and statistical tests were performed using Prism tutorial and R software.

Results

GPR158 Forms Homodimers at the Cell Surface. The orphan receptor GPR158 shares homology within its seven-transmembrane (7TM) domain with the class C GPCRs (Bjarnadottir et al., 2005), including the mGlU and GABA (GABAB) receptors. Like most class C GPCRs, GPR158 also has a large N-terminal extracellular domain linked to the 7TM domain via a cysteine-rich domain; however, neither part of the N-terminal domain is related to those of the class C GPCRs (Kniazeff et al., 2011). The N-terminal part of most class C GPCRs, called the Venus flytrap domain, contains the agonist binding site and is responsible for the constitutive dimerization of these receptors (Pin and Bettler, 2016; Koehl et al., 2019).

We observed that, like most class C GPCRs, GPR158 also exists as homodimers at the surface of transfected HEK293 cells. Indeed, Western blots showed two major bands possibly corresponding to GPR158 monomers and dimers, regardless of which antibodies were used to reveal the protein (Fig. 1A). Moreover, N-terminally HA epitope-tagged GPR158 (HA-GPR158) coimmunoprecipitated N-terminally Flag epitope-tagged GPR158 (Flag-GPR158) coexpressed in HEK293 cells (Fig. 1B). In contrast, neither Flag-mGlU2 nor the Flag-GB2 subunit of the GABAB receptor coimmunoprecipitated with HA-GPR158 receptor (Fig. 1B), whereas HA-GB2 association to its Flag-GB1 subunit partner could easily be detected (Fig. 1B). Cellular dimeric TR-FRET signals (Maurel et al., 2008) were detected using anti-HA and anti-Flag antibodies labeled with luminescence resonance energy transfer–compatible dyes in cells coexpressing HA-GPR158 and Flag-GPR158, but not in cells coexpressing Flag-GPR158 and HA-GB2 (Fig. 1C and D). Finally, Flag-GPR158 homodimer formation was disrupted by an increasing amount of HA-GPR158, but not by HA-GB2 (Fig. 1E). Taken together, these data indicate that GPR158, like the other class C GPCRs, forms homodimers at the cell surface. Given that stoichiometry is a mandatory feature of functional class C GPCRs (El Moustaine et al., 2012), we
analyzed any possible functional role of GPR158. We first further explored the role of its intracellular domain already reported to interact with G-protein–mediating signaling proteins, including Go and RGS7.

**RGS7 Protein Level Is Stabilized by a Specific Region of GPR158 C-Terminal Domain.** In transiently transfected HEK293 cells, GPR158, but not the related GB2 GABAB subunit, interacted with RGS7 and not with RGS4, as shown by coimmunoprecipitation (Fig. 2, A and B). In these experiments the G-protein subunit β5 was coexpressed with RGS7 as it forms a stable complex with RGS7 and protects it from proteolytic degradation (Anderson et al., 2009) (Supplemental Fig. 1A). The RGS7:β5 interaction with GPR158 was confirmed by a TR-FRET approach (Supplemental Fig. 1B). Indeed, a large TR-FRET signal was measured in cells expressing C-terminally Flag-tagged GPR158 (GPR158-Flag) and RGS7-HA, incubated with TR-FRET compatible anti-HA or anti-Flag antibodies, after cell permeabilization. Interestingly, no signal was observed between GPR158 and RGS4-HA. The differential signal measured between RGS7-HA and RGS4-HA expressing cells was not due to different levels of these two proteins nor to a differential amount of GPR158 (Fig. 2A; Supplemental Fig. 1B).
Using membrane-targeting microscopy and in vitro coimmunoprecipitation assays, Orlandi et al. (2015) reported that RGS7 interacted with the C-terminal domain of GPR158. In agreement with this observation, we found that the C-terminal truncated forms of GPR158, which ended at residues Ser692 (GPR158-ΔC1) or Ser699 (GPR158-ΔC2) (Fig. 2C), did not coimmunoprecipitate RGS7 (Fig. 2, D and E). Additionally, no significant intracellular TR-FRET signal with RGS7 could be measured with these cropped GPR158 mutants (Fig. 2F), despite protein levels being similar to those of wild-type GPR158 (Fig. 2A).  

Fig. 2. GPR158 C-terminal domain is required for RGS7 interaction. (A) C-terminal HA-tagged RGS7 (RGS7-HA), but not the C-terminal tagged RGS4 (RGS4-HA) was coimmunoprecipitated by Flag-GPR158 when coexpressed in HEK293 cells. (B) Flag-GPR158, but not Flag-GB2, was coimmunoprecipitated with RGS7-HA in transfected HEK293 cells. (C) Schematic representation of the WT and truncated ΔC1 and ΔC2 (with the last residues mentioned) versions of GPR158 receptor. (D–F) RGS7 binds to the C-terminal domain of GPR158, as shown with coimmunoprecipitation (D and E) and TR-FRET (F) approaches. HEK293 cells were transfected with WT or C-terminal domain truncated versions ΔC1 and ΔC2 of GPR158 together with RGS7-HA. In (D), the coimmunoprecipitation was performed using the C-terminal Flag-tagged versions of the WT (GPR158-Flag) and truncated (GPR158-ΔC1-Flag, GPR158-ΔC2-Flag) GPR158 receptor, coexpressed in HEK293 cells with RGS7-HA. In (E), the coimmunoprecipitation was performed using HA-GPR158 or HA-GPR158-ΔC1, coexpressed in HEK293 cells with the C-terminal Flag-tagged RGS7 (RGS7-Flag). (F) For TR-FRET experiments, HEK293 cells expressing truncated or WT GPR158-Flag and RGS7-HA were permeabilized with tritonX-100 (0.1%) and incubated with antibodies against HA and Flag epitopes bearing the donor and acceptor fluorophores. Each experiment shown is representative of four independent experiments, and data in (F) are the mean ± S.E.M. of triplicates.
(Fig. 2, D–F). Unexpectedly, the three conserved VCPWE motifs of GPR158 (Fig. 3A; Supplemental Fig. 2) were not involved in RGS7 binding (Fig. 3, B and C), despite their similarity to the PDE γ-subunit motif known to participate in complex formation with the RGS7-related RGS9 protein (Slep et al., 2001). Indeed, mutation of the three motifs individually (GPR158-Mut1, -Mut2, and -Mut3) or in combination (GPR158-Mut4) (Fig. 3A) did not suppress GPR158:RGS7 interaction as measured by either coimmunoprecipitation (Fig. 3B) or TR-FRET (Fig. 3C) approaches.

![Diagram](image)

**Fig. 3.** RGS7 interaction requires a short region of the proximal C-terminal domain but not the VCPWE motifs. (A) Schematic representation of the WT and mutated forms of GPR158 used for coimmunoprecipitation (B) and TR-FRET (C) experiments. (B) Coimmunoprecipitation was performed from cells expressing WT, ΔC1, or mutated Mut1-4 GPR158-HA receptor, together with RGS7-Flag and β5. (C) For TR-FRET experiments, HEK293 cells expressing WT, ΔC1, Mut1, Mut2, Mut3, or Mut4 GPR158-HA, together with either RGS7-Flag and β5 or Flag-β-arrestin 1, were permeabilized with tritonX-100 (0.1%). The cells were then incubated with antibodies against HA and Flag epitopes bearing the donor and acceptor fluorophores. Arrestin was used as a negative control for interaction with GPR158. (D) Last residues of the truncated versions (ΔC1–ΔC11) of GPR158. Coimmunoprecipitation (E) and TR-FRET experiments (F) were performed from cells coexpressing WT or truncated (ΔC1–ΔC11) GPR158-Flag, together with RGS7-HA and β5 (E and F) or HA-β-arrestin (F). Each experiment is representative of three independent experiments, and the data in (C and F) are the mean ± S.E.M. of triplicates.
Using a series of GPR158-Flag C-terminal deletion mutants (GPR158-ΔC1 to -ΔC11, Fig. 3D), we identified the region encompassing residues 714 to 764 as the RGS7 binding site using both coimmunoprecipitation and TR-FRET approaches (Fig. 3, E and F). The 714–764 region overlaps with the region reported by Orlandi et al. (2015) to contribute to RGS7 binding. We found that this GPR158-RGS7 interaction enhanced RGS7 abundance in both the absence or presence of its β5 partner (Fig. 4; Supplemental Fig. 3). Using a constant amount of transfected RGS7-coding plasmid, RGS7 protein level was increased in HEK293 cells coexpressing increasing levels of GPR158 (Fig. 4), but not in cells expressing either mGlu2 (Fig. 4A) or GPR158-ΔC1 (Fig. 4B). Of note, the expression of mGlu2 actually led to decreased level of RGS7 protein (Fig. 4A). Whether this is due to mGlu2 itself or to its constant activity due to ambient glutamate in cell culture media remains to be clarified. The enhancing effect exerted by GPR158 was less pronounced in the absence of β5 (Supplemental Fig. 3). Such a stabilizing effect involves the RGS7 binding domain of GPR158 as revealed using a series of mGlu2 chimeric constructs containing the WT or mutated C-terminal domain of GPR158 (Supplemental Fig. 4, A and B).

These data identified a specific region in the C-terminal domain of GPR158, not including the VCPWE motifs, interacting with and stabilizing RGS-β5. We then wondered whether these VCPWE motifs were involved in the interaction with Gαo, a RGS7-regulated G-protein subunit that has been reported to interact with the C-terminal domain of GPR158 (Orlandi et al., 2012, 2015).

**VCPWE Motifs Contribute to Gαo Binding.** We first confirmed the interaction between GPR158 and Gαo using both TR-FRET (Fig. 5A) and BRET approaches (Fig. 5B). Indeed, the GPR158 and Gαo interaction was supported by the generation of a saturating BRET signal curve (Fig. 5B) in HEK293 cells expressing various expression ratios of GPR158-Venus and Gαo-RLuc. In contrast, a nonspecific, linear low BRET signal curve was obtained in control cells expressing GPR158-Venus and Homer3-RLuc (Fig. 5B), an mGlu receptor interacting intracellular protein (Rives et al., 2009). Interestingly, a saturating BRET curve was also observed in cells expressing GPR158-Venus and the Gαo-related Gα11-RLuc protein, whereas no BRET signal could be detected with Gαq-RLuc (Fig. 5C), suggesting that GPR158 can interact with the Gαi/o protein family members, but not with Gαq. We then showed that the C-terminal domain of GPR158 was required for this interaction since the TR-FRET signal was greatly reduced in cells expressing Gαo-Flag and the truncated GPR158-ΔC1-HA or GPR158-ΔC2-HA (Fig. 5A). The VCPWE motifs located in the cytoplasmic domain of GPR158 were identified as key elements for the interaction with Gαo, as the mutation of all three motifs (GPR158-Mut4) largely decreased the TR-FRET signal with Gαo (Fig. 5A). Whereas Mut1 and Mut3 show a lower TR-FRET signal than WT GPR158 with Gαo, the signal with Mut2 was not affected (Fig. 5A lower panel). These results revealed that motifs 1 and 3 were involved in the association of Gαo with GPR158; however, the Mut1 protein level was significantly reduced compared with that of Mut3 (Fig. 5A, middle panel), suggesting that Mut1 is involved to a lesser extent than is Mut3 in αo interaction. When using a BRET approach (Fig. 5D), and to a lesser extent using a TR-FRET approach (Fig. 5A), Gαo interaction with GPR158-Mut4 could still be measured; however, the signal was lower than that obtained with the WT GPR158, consistent with the existence of a second Gαo interacting site in GPR158 (Orlandi et al., 2015). The difference between the observed TR-FRET and BRET signals is likely related to photophysical properties of the techniques, as the dyes used for each technique display different Ro (5 and 10 nm, respectively). This difference would be enhanced by the larger distance contributed by the size of antibodies used in the TR-FRET approach. Because GPR158 behaves as a scaffolding protein for G-protein signaling proteins and can interact with G-protein subunits, even in absence of ligand, we then asked whether GPR158 displayed GPCR canonical basal coupling to G proteins, as described for many GPCRs (Oh et al., 2006; Thathiah et al., 2009).

**GPR158 Did Not Display Detectable Constitutive Coupling to Gq, Gs, or Gi/o Proteins.** To assess the putative basal G-protein activation by GPR158, we used a BRET assay. This assay monitors the association state of the Gαo-βγ protein complex composed of Gαi/o-RLuc and βγ-Venus. The BRET signal decreases when a Go-coupled GPCR is activated, as illustrated with the μ-opioid receptor (MOR) (Fig. 6A) and the GABAB receptor (Fig. 6B). In contrast, the BRET signal was not affected by the
vasopressin receptor V2, known not to couple to Go (Fig. 6A).
In addition, a basal coupling to Go of both MOR (Fig. 6A) and
GABAB (Fig. 6B) in the absence of ligand could be detected,
but not for V2 (Fig. 6A). Moreover, the basal Go coupling of
MOR increased as a function of MOR cell-surface protein
level measured using nonpermeable fluorescent labeling of
Snap tag domains inserted at the N-terminal end of the
receptor (Fig. 6C). Increasing GPR158 expression also led to
a proportional decrease of BRET signal (Fig. 6C), suggesting
a ligand-independent basal coupling of GPR158 to Go.
This GPR158 basal coupling to G protein was not, how-
ever, associated with any change in a second-messenger
production-based readout (Fig. 6D), in contrast to what was
observed with either MOR or GABAB. The chimeric GqTop
protein containing portions of Gi/o protein was used to allow
Gi/o-coupled receptors to activate PLC, leading to the pro-
duction of IP1, as illustrated with MOR or GABAB (Fig. 6D;
Supplemental Fig. 5). In contrast to the large basal increase in
IP1 production measured in GABAB receptor–expressing cells
or, to a lesser extent, in MOR-expressing cells, no effect was
observed with GPR158, despite a similar protein level of both
receptors (Fig. 6D). These data bring no evidence for a
constitutive canonical GqTop protein activation by GPR158,
suggesting no activation of constitutive Gi/o or Gq proteins

Fig. 5. Goa association with GPR158 involves the VCPWE motifs. (A) TR-FRET–based analysis of GPR158 and Goa (Goa isoform) association in
HEK293 was measured in cells transfected with Goa-Flag and either the WT, ΔC1, ΔC2, or Mut1-4 GPR158-HA (lower panel). The amount of Goa-Flag
and GPR158-HA versions were quantified by ELISA (upper and middle panels, respectively) against the Flag and HA epitopes and expressed as % of
either Goa-Flag or WT GPR158-HA protein levels detected in the Goa-Flag or GPR158-HA control conditions (black bars). The amount of the Goa-Flag
protein is not significantly different (P = nonsignificant (ns)) in the various tested conditions (upper panel). Similarly, no significant difference was
observed between the amount of the various GPR158 protein versions (middle panel), except between Mut1 and Mut3 versions (* on the graph,
P = 0.0185). Statistical analysis of the HTRF signal is indicated directly on the lower-panel graph. For each of the three panels, data from six experiments are
pooled on the same graph, and values are mean ± S.E.M. Data statistics were analyzed using a one-way analysis of variance (ANOVA) test (no difference
in the variance was checked with a Brown-Forsythe method), and a multiple comparison correction was performed by Dunnett method; adjusted F values
are reported (*P < 0.05; **P < 0.01. (B) Interaction of GPR158 with Goa (Goa isoform) was assessed by BRET assay in HEK293 cells transfected with
Goa-RLuc or Homer3-RLuc and increasing amounts of GPR158-Venus. Only the association GPR158-Venus and Goa-RLuc generated a saturating curve
suggesting a specific association. Protein levels of GPR158 and Goa or Homer3 were monitored by determination of the specific Venus fluorescence and RLuc luminescence signals, the ratios of which were used for plotting the x-axis. (C) Selective association of GPR158 with various Go subunits. The BRET
signal was monitored in cells expressing Goa-RLuc, Goa1-RLuc, or Goaq-RLuc and increasing amounts of GPR158-Venus. Protein levels of GPR158 and
Goa were monitored by determination of the specific Venus fluorescence and RLuc luminescence signals, the ratios of which were used for plotting the
x-axis. (D) BRET saturation curves were established from cells transfected with increasing amounts of plasmids coding for WT GPR158-Venus or
GPR158-Mut4-Venus and constant amounts of Goa-RLuc. The data from three independent experiments were pooled for BRET experiments in (B–D), and
values are means ± S.E.M. of triplicate determinations.

This GPR158 basal coupling to G protein was not, how-
ever, associated with any change in a second-messenger
production-based readout (Fig. 6D), in contrast to what was
observed with either MOR or GABAβ. The chimeric GqTop
protein–containing portions of Gi/o protein was used to allow
Gi/o-coupled receptors to activate PLC, leading to the pro-
duction of IP1, as illustrated with MOR or GABAβ (Fig. 6D;
Supplemental Fig. 5). In contrast to the large basal increase in
IP1 production measured in GABAβ receptor–expressing cells
or, to a lesser extent, in MOR-expressing cells, no effect was
observed with GPR158, despite a similar protein level of both
receptors (Fig. 6D). These data bring no evidence for a
constitutive canonical GqTop protein activation by GPR158,
suggesting no activation of constitutive Gi/o or Gq proteins
either. Moreover, no detectable GPR158 constitutive coupling to Gs or Gi, as assessed by the adenylyl cyclase-driven production of the second-messenger cAMP (Supplemental Fig. 5), could be detected.

Further analysis of GPR158 7TM amino-acid sequence identified residues K502 and R505 in TM3 (Supplemental Fig. 6), highly conserved in class C receptors from fishes to humans and the mutations of which (such as mutations equivalent to K502E or R505A of GPR158 in GABAB receptor) do affect the ligand-induced and constitutive activity of GABAB, mGlu1, or calcium-sensing GPCRs (Francesconi and Duvoisin, 1998; Ango et al., 2001; Duthey et al., 2002; Pin et al., 2004; Binet et al., 2007; Rondard et al., 2011). Surprisingly, the effect of GPR158 on Go activation/dissociation was not significantly altered when the receptor bore mutations K502E or R505A (Fig. 7, A and B). The preceding data showed that GPR158 affects the heterotrimERIC Gαo-βγ association state and that this effect is unlikely due to a ligand-independent canonical coupling to Go protein. As additional evidence in support of this conclusion, the effect of GPR158 on Go is slightly diminished after treatment with the Gi/o inhibitor pertussis toxin (PTX), but the PTX effect is similar to that observed with mock-transfected cells, suggesting that the effect of GPR158 on Go is PTX-independent. In contrast, both the basal and agonist-induced Go activation observed with MOR is, as expected, largely inhibited by PTX (Fig. 7C).

GPR158 VCPWE Motifs Constitutively Increased Dissociated Go Levels in Cells Independently to Classic G-Protein Coupling. We found that the VCPWE motifs are essential for the GPR158 effect on Go. First, deletion of the C-terminal domain of GPR158 (GPR158-ΔC1) completely abolished the GPR158-induced change in the Go BRET signal (Fig. 8A). Second, the mutation of all three motifs (GPR158-Mut4) suppressed the effect of GPR158 on the Go BRET sensor (Fig. 8B); however, the mutation of only one of these motifs was not sufficient to suppress the GPR158 effect (Fig. 8, C and D) or the combined mutation of motifs 1 and 2 or 2 and 3 (Fig. 8, E and F). Indeed, only the combined mutation of both VCPWE motifs 1 and 3 led to a suppression of the GPR158 effect on the Go BRET sensor (Fig. 8E).
Discussion

Signaling and functions of GPR158 remain poorly characterized, although it has been proposed to be involved in the effect of osteocalcin in the brain (Khrimian et al., 2017). The scaffolding ability of GPR158 to interact with RGS7 allows it to regulate Go signaling induced by neighboring receptors when expressed in the same cells (Orlandi et al., 2012). Besides, GPR158 binds Go, but its ability to couple to Go is still unclear. Furthermore, although GPR158 possesses three conserved VCPWE motifs, their function is still not elucidated.

Here we addressed the role of scaffolding and signaling of GPR158. We show that RGS7 interacts in the proximal part of the C-terminal intracellular domain, whereas Go interacts downstream of this site with two of the three VCPWE motifs. Despite its ability to interact with Go, we did not find any evidence for a canonical basal activation of this G protein by GPR158. Instead, we propose that GPR158, under basal conditions, can regulate Go signaling by trapping the Go subunit, leaving βγ to act on its effectors. Although our data were all obtained in a recombinant system, our analysis using various expression levels of the partners and the use of different tags, inserted at different location and various approaches, provides a good indication that what is reported here is likely also occurring in native systems. With that said, we cannot exclude that the described process can be further controlled by other partners not expressed in HEK293 cells. A possible canonical G-protein activation upon agonist binding to GPR158 cannot be excluded, as we did not examine this possibility in the present study.

We delineated the RGS7 binding site in the 714–764 region of GPR158 C-terminal domain, proximal to the T7M domain (Fig. 3). This small region overlaps with the CD1 region defined by Orlandi et al. (2015) as containing a binding site for RGS7 and displaying homology with R7BP protein. Indeed, GPR158 and R7BP compete for interacting with RGS7. RGS7 is composed of the RGS, GGL, and DEP domains. The latter is proposed to interact with GPR158 (Orlandi et al., 2012), as well as other proteins, but no clear DEP-binding consensus sequence has been identified. As observed with R7BP, GPR158 also stabilizes the RGS7 protein, leading to an increase in RGS7 protein level (Fig. 4). Consistent with this observation in HEK293 cells, a decrease of RGS7 protein level has been reported in GPR158 KO mice, with diminution of the pool of RGS7 in the membrane fraction and relocation into the cytoplasm, as revealed by electron microscopy in native and transfected models (Orlandi et al., 2012, 2015). Accordingly, GPR158 appears to regulate the pool of RGS7 and tunes its localization to the plasma membrane. RGS7 protein level is also known to depend on the coexpression of β5 (Supplemental Fig. 1) (Anderson et al., 2009), which binds to RGS7 GGL domain to form a putative G-protein βγ complex. As such, RGS7 binds GPR158 and β5 via two independent domains, DEP and GGL, respectively, leaving its RGS domain free to bind active Go and deactivate it.

We also demonstrated that two of the VCPWE motifs (i.e., motifs 1 and 3) of GPR158 were important for Go binding, even in the absence of RGS7 (Fig. 5). Motif 2 in primates, including human GPR158, does not contain the conserved
proline residue, suggesting it has lost its ability to bind Go during evolution (Supplemental Fig. 2) (Slep et al., 2001). Mutation of the three motifs did not completely suppress Go binding, as revealed with the BRET approach, suggesting that there might be another site. Interestingly, Orlandi et al. (2015) reported two Go binding sites in GPR158 C-terminal domain, one close to the RGS7 binding site and another one in the distal part of the C-terminal domain. Thus, according to our data, VCPWE motifs may correspond to this second distal site, whereas the other corresponds to the proximal site. The remaining Go interaction observed after mutating the VCPWE motifs could also simply be indirect owing to a
proximity to GPR158 resulting from Go association with RGS7 or other proteins or GPCRs interacting with GPR158. We further showed that the VCPWE motifs are not only important for Go binding but also impact Go-βγ association. The observed decreased of BRET between Go-RLuc and βγ-Venus in the presence of GPR158 (Fig. 6) could reflect the dissociation or the conformational change usually observed upon G-protein activation (Gales et al., 2006). Because we obtained no evidence for a canonical activation of G proteins by GPR158 in absence of ligand, however, we favored alternative hypotheses. The VCPWE motifs may bind Go-RLuc, reducing its association with βγ-Venus, or Go-βγ complexes activated by other endogenous GPCRs may have led to the release of Go, which can be trapped by VCPWE motifs, preventing their reassociation with βγ. Both situations likely occur, as the first explanation is supported by the PTX-insensitive component of the GPR158 effect on Go BRET sensor, whereas the second is supported by known action of VCPWE-related motifs. Indeed: 1) the ICPWE motif of the γ subunit of the retinal PDE binds the active α subunit (Slep et al., 2001), and 2) the distal site in the GPR158 C-terminal domain has been proposed to preferentially bind an active form of Go (Orlandi et al., 2015). Taken together, these findings suggest that VCPWE motifs likely trap isolated Go and inhibit them from activating their effectors while leaving βγ free to activate its own effectors. GPR158 would then induce a ligand-independent signaling bias of βγ versus α subunits, an effect reminiscent of the function of some group II AGS proteins (Blumer and Lanier, 2014). Of note, two VCPWE motifs (1 and 3) are required for this effect (Fig. 8), suggesting that either one Go binds to both motifs or, alternatively, that each motif binds one Go independently of each other, possibly differently regulated by RGS7. Further experiments will be necessary to clarify this point. Because GPR158 associates with Cav2 calcium channel in the rat brain (Müller et al., 2010) and Kv4.2 potassium channel in the mouse brain (Marionneau et al., 2009), which are both regulated by βγ and RGS proteins, such a Go-trapping mechanism by the VCPWE motifs may change the kinetics of such regulatory effects of GPR158. In the retina, where both RGS7 and the GPR158-related GPR179 containing 21 VCPWE motifs are expressed (Audo et al., 2012; Orlandi et al., 2012), such a mechanism could control the spatiotemporal regulation of signaling of photoreceptors and ON bipolar cells. This process may be reminiscent of the control of the PDE response by rhodopsin and transducin that involves RGS9/7 and the ICPWE motif of PDE γ subunit (Slep et al., 2001).

Many GPCRs display constitutive activity in absence of ligand, leading to constitutive canonical G-protein coupling, and mutation-driven constitutive activity of some GPCRs leads to various diseases (Tao, 2008). GPR158 possesses class C GPCR features required for G-protein coupling (Bjarnadóttir et al., 2005), like lysine and arginine residues in TM3 previously shown to be important for G-protein coupling in GABAb (Galvez et al., 2001; Binet et al., 2007) and mGlul (Doré et al., 2014; Koehl et al., 2019); however, we did not detect any constitutive G-protein activation when measuring Goi, Gq, orGs activity in HEK293 cells expressing WT or mutated GPR158. One can envisage that GPR158 displays no constitutive canonical G-protein coupling or that GPR158 couples to other pathways that have not been addressed in this work, like those resulting from G12/13 protein or the G-independent arrestin pathway, although no direct coupling to arrestin has been clearly demonstrated for any class C GPCRs so far (Pin and Bettler, 2016). Another possibility is that GPR158 needs a GPCR partner, like the GABAg receptor, requiring the association of two different proteins GB1 and GB2 (Pin et al., 2004; Rondard et al., 2011). More work is needed to clarify this important issue. As mentioned, such data do not exclude a direct G-protein activation by GPR158 upon agonist binding.

Dimer formation is required for activation of the multidomain class C mGlu, calcium sensing, taste T1Rs, and GABAb receptors. Indeed, the intersubunit movement of the extracellular domain (ECD) resulting from ligand binding changes the interaction mode of the 7TM domains, leading to the activation of one of them (Xue et al., 2015; Pin and Bettler, 2016; Koehl et al., 2019). We have shown here that GPR158 also forms homodimers, making possible a similar activation process with ligands interacting in the GPR158 ECD; however, not only does this ECD not share similarity with that of other class C GPCRs, but also with any other protein of known structure, making it impossible to predict the mode of action of such a domain. It is clearly important to elucidate whether and how GPR158 can directly activate G protein upon activation with a ligand.

Not all 7TM proteins couple to G proteins, like the adiponectin receptor (Vasiliauskaitė-Brooks et al., 2017), the GB1 subunit of GABAg, or both T1R1 and T1R2, which need to be associated with T1R3 to form the umami and sweet taste receptors, respectively (Kniazeff et al., 2011). Some orphan GPCRs are also considered regulatory associated proteins that control the activity of functional GPCRs, as shown elegantly for the orphan receptor GPR50 that inhibits the melatonin receptor MT1 (Levoye et al., 2006) or controls TGFβ signaling ( Wojciech et al., 2018). A recent article proposed that GPR158 mediates the action of the hormone osteocalcin (Khrimian et al., 2017), which has also been reported to activate GPRC6A, another class C GPCR (Pi et al., 2005). In cells deleted of GPR158, osteocalcin did not trigger an increase in brain-derived neurotrophic factor expression (Khrimian et al., 2017), whereas the production of second-messenger IP3 was decreased, suggesting that osteocalcin action on GPR158 modulates IP3 production. Although we did not observe any Goq binding or constitutive activity toward IP3 production in transfected HEK293 cells, this does not exclude a ligand-induced activation of the Gq pathway by osteocalcin. A signaling-partner protein may be missing in HEK293 cells to allow GPR158 to couple to Gq, as illustrated by the class C mGlu7 glutamate receptor, which needs to interact with Pick1 to couple to the PLC-IP3 pathway in neurons (Perroy et al., 2000). Moreover, mGlul, reported to be Go coupled in the recombinant system, is endogenously coupled to the Gq pathway in parallel fiber-Purkinje cell synapses (Abitbol et al., 2012).

According to our data and previous studies, GPR158 behaves as a scaffolding platform that tunes the Go pathway in an original way. In the absence of ligand, GPR158 displays various roles. First, it associates, stabilizes, and brings RGS7 to the plasma membrane, where RGS7 deactivates Go proteins activated by surrounding receptors (Orlandi et al., 2012). Second, the GPR158 VCPWE motifs-mediated trapping action on Go could impact the Go signaling in the surrounding microenvironment and favor βγ-mediated signaling. Third,
GPR158 can be associated with both RGS7 and Gα signaling proteins in synaptic plasticity and learning. Mol Pharmacol 89:273–286.


Address correspondence to: Laurent Prézeau, Univ. Montpellier, IGF, 141, rue de la Cardonille, 34094 Montpellier, cedex 05, France. E-mail: Laurent.prezeau@igf.cnrs.fr