Novel Interactions Involving the Mas Receptor Show Potential of the Renin–Angiotensin system in the Regulation of Microglia Activation: Altered Expression in Parkinsonism and Dyskinesia

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Abstract

The renin–angiotensin system (RAS) not only plays an important role in controlling blood pressure but also participates in almost every process to maintain homeostasis in mammals. Interest has recently increased because SARS viruses use one RAS component (ACE2) as a target-cell receptor. The occurrence of RAS in the basal ganglia suggests that the system may be targeted to improve the therapy of neurodegenerative diseases. RAS-related data led to the hypothesis that RAS receptors may interact with each other. The aim of this paper was to find heteromers formed by Mas and angiotensin receptors and to address their functionality in neurons and microglia. Novel interactions were discovered by using resonance energy transfer techniques. The functionality of individual and interacting receptors was assayed by measuring levels of the second messengers cAMP and Ca²⁺ in transfected human embryonic kidney cells (HEK-293T) and primary cultures of striatal cells. Receptor complex expression was assayed by in situ proximity ligation assay. Functionality and expression were assayed in parallel in primary cultures of microglia treated or not with lipopolysaccharide and interferon-γ (IFN-γ). The proximity ligation assay was used to assess heteromer expression in parkinsonian and dyskinetic conditions. Complexes formed by Mas and the angiotensin AT₁ or AT₂ receptors were identified in both a heterologous expression system and in neural primary cultures. In the heterologous system, we showed that the three receptors—MasR, AT₁R, and AT₂R—can interact to form heterotrimers. The expression of receptor dimers (AT₁R-MasR or AT₂R-MasR) was higher in microglia than in neurons and was differentially affected upon microglial activation with lipopolysaccharide and IFN-γ. In all cases, agonist-induced signaling was reduced upon coactivation, and in some cases just by coexpression. Also, the blockade of signaling of two receptors in a complex by the action of a given (selective) receptor antagonist (cross-antagonism) was often observed. Differential expression of the complexes was observed in the striatum under parkinsonian conditions and especially in animals rendered dyskinetic by levodopa treatment. The negative modulation of calcium mobilization (mediated by AT₁R activation), the multiplicity of possibilities on RAS affecting the MAPK pathway, and the disbalanced expression of heteromers in dyskinesia yield new insight into the operation of the RAS system, how it becomes unbalanced, and how a disbalanced RAS can be rebalanced. Furthermore, RAS components in activated microglia warrant attention in drug-development approaches to address neurodegeneration.

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Key Words Parkinson · microglia · Mas receptor · GPCR · angiotensin · angiotensin AT₁ receptor · angiotensin AT₂ receptor

Introduction

The renin–angiotensin system (RAS) has been widely studied for its role in controlling blood pressure. Protein components of RAS are angiotensin-converting enzyme 1 (ACE1), which produces angiotensin II (Ang II); angiotensin-converting enzyme 2 (ACE2), which converts Ang II to angiotensin 1–7 (Ang(1-7)); the Ang II receptors AT₁R and AT₂R; and the Ang(1–7) receptor Mas. For several decades, the balance between the prooxidative, proinflammatory, and the antioxidative, anti-inflammatory RAS arms was thought to be centered in the functional equilibrium of AT₁R and AT₂R. Now, the role of the ACE2/Mas receptor function has been revealed as a crucial element of RAS balance and RAS function. The Mas receptor (MasR) was identified as a product of an oncogene and, because of its resemblance to the mitochondrial assembly gene from *Saccharomyces cerevisiae*, it was first known as Mas-related proto-oncogene. All RAS receptors belong to the superfamily of G protein-coupled receptors (GPCRs). A novel family of RAS-related receptors are named Mas-related GPCRs (Mrgprs); they respond to Ang (1–7) but their endogenous agonist is one of the newest members of the RAS system, alamandine [1–5]. ACE2 is considered the canonical cell surface receptor for SARS-CoV-2, the virus that causes COVID-19. In fact, ACE2 was serendipitously identified as the receptor for viruses of the SARS family [6–10].

RAS has a relevant role in controlling neurotransmission in both the central and peripheral nervous systems. Identification of Ang II receptors in neural cells of the substantia nigra and other regions within the basal ganglia has uncovered novel therapeutic approaches to address neurodegeneration in Parkinson’s disease (PD) and other synucleinopathies [11–18]. For obvious reasons, we must consider the brain RAS to understand the neurological manifestations of some patients infected with SARS-CoV-2. In fact, some cases of COVID-19 have neurological symptoms as diverse as encephalitis and seizures [19–21]. In some cases, the basal ganglia have been identified as mediating virus-induced central nervous system (CNS) alterations [13, 22, 23].

Although angiotensin I has little biological activity, ACE1 converts it into Ang II, the principal RAS effector, which activates AT₁R and AT₂R, whose actions in CNS development have been clearly delineated. However, in adults, the scenario is quite complex because AT₁R and AT₂R seemingly mediate opposite actions, whereas AT₁R activation usually leads to production of reactive oxygen species, activation of AT₂R counterbalances this noxious effect. In macrophages and microglia, AT₁R is thought to contribute to inflammation, whereas AT₂R attenuates inflammation and contributes to neuroprotection. These data reinforce the hypothesis that RAS balance is important in illnesses such as PD, which involves glial activation and neuroinflammation [14, 24–27]. MasR, whose endogenous agonist is Ang (1–7), adds complexity by mediating antioxidant and anti-inflammatory effects of AT₂R [24, 28]. In summary, RAS action in a given cell or brain circuit depends on peptide production by ACE1 and ACE2 and the expression of RAS receptors. Overall, the mechanisms underlying RAS balance in healthy conditions and RAS unbalance in aging [14] or disease remain unknown.

The physiological action of GPCRs is often mediated by heteromers; that is, by complexes having more than one receptor. Consensus is that receptor heteromers are novel functional units—their properties are different from those of the individually expressed receptors [29–31]. The first reported heteromer consisted of two GPCRs with the same endogenous agonist; namely, the μ-δ opioid receptor heteromer [32]. Further examples of heteromers formed by two receptors for the same endogenous ligand are adenosine A₁-A₂A and adenosine A₂A–A₂B receptors. In these cases, the presence and/or activation of one of the receptors blunts the response arising in the partner receptor, and structural information can explain how the trans blockade may happen [33–38]. Importantly, it has been reported that AT₁R and AT₂R and MasR and AT₂R form heteromers [39–41]. The aim of this study was to demonstrate the presence of Mas and AT₁/AT₂ receptor heterodimerization in neurons and microglia. The function of the MasR-AT₁R/AT₂R complexes in resting and activated microglia was investigated to complement our recent findings on AT₁R-AT₂R heteromer function in neural cells [42]. Importantly, we assessed the expression of heteroreceptor complexes in animal models of parkinsonian and dyskinnesia. We focused on the striatum because of its relevance in PD. Our results provide a holistic model to understand RAS action, especially under conditions in which microglia become activated.

Materials and Methods

Reagents

Lipopolysaccharide (LPS), interferon-γ (IFN-γ), Ang II, CGP-42112A, Ang [1–7], candesartan, PD123319, and A779 were purchased from Sigma-Aldrich (St. Louis, MO, USA). Forskolin was purchased from Tocris (Bristol, UK). The concentrations of ligands used for signaling assays were selected on the basis of the dose–response experiments shown in Supplementary Figure S1.
Cell Culture

Human embryonic kidney (HEK-293T) cells were grown in Dulbecco’s modified Eagle’s medium (DMEM; Gibco/Thermo Fisher Scientific, Paisley, UK) supplemented with 2 mM l-glutamine, 100 μg/mL sodium pyruvate, 100 U/mL penicillin/streptomycin, MEM Non-Essential Amino Acids Solution (1/100) and 5% (v/v) heat-inactivated fetal bovine serum (FBS) (all supplements were from Invitrogen/Thermo Fisher Scientific, Paisley, UK). Cells were maintained at 37 °C in a humid atmosphere of 5% CO₂.

As mentioned in the “Introduction,” in this study, we focused on brain striatum. To prepare mice striatal primary microglial cultures, the brain was removed from C57BL/6 mice at 2 to 4 days of age. Microglial cells were isolated as described in [43] and grown in DMEM supplemented with 2 mM l-glutamine, 100 U/mL penicillin/streptomycin, MEM non-essential amino acids preparation (1/100), and 5% (v/v) heat-inactivated FBS. Briefly, striatum tissue was dissected, carefully stripped of its meninges, and digested with 0.25% trypsin for 20 min at 37 °C. Digestion was stopped by washing the tissue. A cell suspension was prepared by passing the cells through a 100-μm pore mesh. Glial cells were suspended in medium and seeded at a density of 1 × 10⁶ cells/mL in 6-well plates for cyclic adenyllic acid (cAMP) assays, in 12-well plates with coverslips for in situ proximity ligation assays (PLA), and in 96-well plates for mitogen-activated protein kinase (MAPK) experiments. Cultures were maintained at 37 °C in a humidified 5% CO₂ atmosphere, and unless otherwise stated, the medium was replaced once a week.

For neuronal primary cultures, the striatum from mouse embryos (E19) was removed and the neurons were isolated, as described by [44], and plated at a density of approximately 120,000 cells/cm². The cells were grown in a neurobasal medium supplemented with 2 mM L-glutamine, 100 U/mL penicillin/streptomycin, and 2% (v/v) B27 supplement (Gibco) in a 6-, 12-, or 96-well plate for 19 to 21 days. Cultures were maintained at 37 °C in a humidified 5% CO₂ atmosphere, and unless otherwise stated, the medium was replaced once every 4 to 5 days.

Immunodetection of specific markers (NeuN for neurons and CD-11b for microglia) showed that neuronal preparations contained >98% neurons and that microglia preparations contained at least 98% microglial cells [45, 46]. Contamination by astrocytes was negligible; in our hands, the passage of the suspension through a syringe disrupts astroglial cells, and in addition, the culture medium used does not favor astrocyte survival (checked in every culture).

PD Model Generation, Levodopa Treatment, and Dyskinesia Assessment

All experiments were carried out in accordance with EU directives (2010/63/EU and 86/609/CEE) and were approved by the ethical committee of the University of Santiago de Compostela. Similar to the approach elsewhere described [47], our experimental design using male Wistar rats aimed to obtain four groups of animals as described below. Animals were 8 weeks old at the beginning of the experimental procedure.

Details of model generation and the protocol of drug administration and behavioral analysis, performed by a blinded investigator, are given elsewhere [48, 49]. Surgery was performed on rats anesthetized with ketamine/xylazine (1% ketamine, 75 mg/kg, and 2% xylazine, 10 mg/kg). Lesions were produced in the right medial forebrain bundle to achieve a complete degeneration of the nigrostriatal pathway. The rats were injected with 12 μg of 6-hydroxydopamine (6-OH-DA) (to provide 8 μg of 6-hydroxy-DA free base; Sigma-Aldrich) in 4 μL of sterile saline containing 0.2% ascorbic acid. These were considered “lesioned” animals. Injection of the vehicle led to the generation of naïve (or non-lesioned) animals.

The 6-OH-DA hemilesoned rat is considered a PD model. Amphetamines-induced rotation was tested in a bank of 8 automated rotometer bowls (Rota-count 8, Columbus Instruments, Columbus, OH, USA) by monitoring full (360°) body turns in either direction. Right and left full body turns were recorded over 90 min following an injection of D-amphetamine (2.5 mg/kg i.p.) dissolved in saline. Rats that displayed more than 6 full body turns/min ipsilateral to the lesion were included in the study (this rate would correspond to >90% depletion of dopamine fibers in the striatum [50]).

Spontaneous use of forelimb can be measured by the cylinder test [51, 52]. Rats were placed individually in a glass cylinder (20 cm in diameter) and the number of left or right forepaw contacts was scored by an observer blinded to the animals’ identity and presented as left (impaired) touches as a percentage of total touches. A control animal would thus receive an unbiased score of 50%, whereas the lesion usually reduces the performance of the impaired paw to less than 20% of total wall contacts.

Of the lesioned animals displaying parkinsonism-like behavior according to the above described tests (18 in total), 12 were chronically treated with L-DOPA daily for 3 weeks. A mixture of L-DOPA methyl ester (6 mg/kg) plus benserazide (10 mg/kg) was administered subcutaneously. The treatment reliably induces dyskinetic movements in some rats. As described in a previous report [47], abnormal involuntary movements were evaluated according to the rat dyskinesia scale described in detail elsewhere [48, 53–56]. The severity of each abnormal involuntary movement (AIM) subtype (limb, orolingual, and axial) was assessed using scores from 0 to 4 (1 = occasional, present < 50% of the time; 2 = frequent, present > 50% of the time; 3 = continuous but interrupted by strong sensory stimuli; 4 = continuous, not interrupted by strong sensory stimuli). Rats were classified as “dyskinetic” if
they displayed a score ≥2 per monitoring period on at least two AIM subtypes. Animals classified as “non-dyskinetic” exhibited either no L-DOPA-induced abnormal involuntary movements or very mild/occasional ones [57]. Animals with low scores, either non-dyskinetic or dyskinetic, were excluded. In summary, four groups of animals were obtained: [1] non-lesioned; [2] lesioned, treated with vehicle; [3] lesioned and became dyskinetic when treated with L-DOPA; and [4] lesioned and did not become dyskinetic upon L-DOPA treatment. Tyrosine hydroxylase immunostaining was performed in every animal from sections taken postmortem [18, 49]; selected animals undergoing 6-OH-DABA treatment showed, in the lesioned hemisphere, >95% nigral dopaminergic denervation. Overall, 4 animals (those with better scores) were selected in each of the following 4 groups: naïve, lesioned, lesioned/L-DOPA dyskinetic, and lesioned/L-DOPA nondyskinetic. The PLA analysis (see below) was performed in different fields of striatal sections from each of the 16 selected animals. The striatum was delimited in sections using a bright field, and images were captured within delimitation coordinates.

**Fusion proteins**

Human cDNAs for AT1, AT2, Mas, and σ1 receptors cloned into pcDNA3.1 were amplified without their stop codons using sense and antisense primers harboring either BamHI and HindIII restriction sites to amplify AT1R and AT2R or with BamHI and EcoRI restriction sites to amplify Mas and σ1 receptors. Amplified fragments were then subcloned to be in frame with an enhanced yellow fluorescent protein (pEYFP-N1; Clontech, Heidelberg, Germany) or a Rluc (pRluc-N1; PerkinElmer, Wellesley, MA) on the C-terminal end of the receptor to produce AT1R/YFP, AT1R/Rluc, AT2R-YFP, Mas-YFP, Mas-Rluc, and σ1R-Rluc fusion proteins.

**Cell Transfection**

HEK-293T cells were transfected with correspondent cDNA by means of the poly-ethylenimine (PEI; Sigma-Aldrich) method, as previously described [58]. Briefly, the corresponding cDNA diluted in 150 mM NaCl was mixed with PEI (5.5 mM in nitrogen residues) also prepared in 150 mM NaCl for 10 min. The cDNA-PEI complexes were transferred to HEK-293T cells, which were incubated for 4 h in serum-starved medium. Then, the medium was replaced by fresh supplemented culture medium and cells were maintained at 37 °C in a humid atmosphere of 5% CO2. Forty-eight hours after transfection, cells were washed, detached, and re-suspended in the assay buffer.

**Immunocytochemistry**

HEK-293T cells were seeded on glass coverslips in 12-well plates. After 24 h of culture, cells were transfected with AT1R-YFP cDNA (1 μg) and Mas-Rluc cDNA (1 μg) or with AT2R-Rluc cDNA (1 μg) and Mas-Rluc cDNA (1 μg). Forty-eight hours later, cells were fixed in 4% paraformaldehyde for 15 min and washed twice with PBS containing 20 mM glycine before permeabilization with PBS-glycine containing 0.2% Triton X-100 (5 min incubation). A blocking solution consisting of PBS containing 1% bovine serum albumin was added (1 h). HEK-293T cells were then labeled with a mouse anti-Rluc antibody (1/100; Millipore, Darmstadt, Germany) and subsequently treated with Cy3-conjugated anti-mouse (1/200; Jackson ImmuResearch, West Grove, PA, USA; red) antibody (1 h each). The expression of Mas-YFP and AT2R-YFP was detected by their YFP fluorescence. Nuclei were stained with Hoechst (1/100 from 1 mg/mL stock; Sigma-Aldrich). Samples were washed several times and mounted with 30% Mowiol (Calbiochem, Merck, Darmstadt, Germany). Images were obtained in a Zeiss LSM 880 confocal microscope (Zeiss, Jena, Germany) with a × 63 oil objective.

**Bioluminescence Resonance Energy Transfer Assay**

HEK-293T cells were transiently cotransfected with a constant amount of cDNA encoding Mas-Rluc (0.75 μg) and with increasing amounts of cDNAs corresponding to AT1R-YFP (0.5 to 2.5 μg) or AT2R-YFP (0.5 to 3 μg). When indicated cDNA encoding for AT1R fused to Rluc was used. For negative control, HEK-293T cells were transiently cotransfected with a constant amount of cDNA encoding σ1R-Rluc (0.75 μg) and with increasing amounts of cDNA corresponding to AT2R-YFP (0.5 to 3 μg). To control the cell number, sample protein concentration was determined using a Bradford assay kit (Bio-Rad, Munich, Germany) using bovine serum albumin (BSA) dilutions as standards. To quantify fluorescent proteins, cells (20 μg of total protein) were distributed in 96-well microplates (black plates with transparent bottoms) and fluorescence was read in a Fluostar Optima Fluorimeter (BMG Labtech, Offenburg, Germany) equipped with a high-energy xenon flash lamp, using a 10-nm bandwidth excitation filter at 485 nm. For bioluminescence resonance energy transfer (BRET) measurements, the equivalent of 20 μg of total protein cell suspension was distributed in 96-well white microplates with white bottoms (Corning 3600; Corning Inc., Corning, NY, USA). BRET was determined 1 min after adding coelenterazine H (Molecular Probes, Eugene, OR), using a Mithras LB 940 reader (DLReady, Berthold Technologies, Bad Wildbad, Germany), which allows the integration of the signals detected in the short-wavelength filter at 485 nm and the long-wavelength filter at 530 nm. To quantify Mas-Rluc expression, luminescence readings were
obtained 10 min after the addition of 5 μM coelenterazine H. MilliBRET units (mBU) are defined as follows:

\[
mBU = \left( \frac{\lambda_{530}(\text{long-wavelength emission}) - \lambda_{485}(\text{short-wavelength emission})}{C_f} \right) \times 1000
\]

in which \(C_f\) corresponds to \([\text{(long-wavelength emission)}/(\text{short-wavelength emission})]\) for the Rluc construct expressed alone in the same experiment.

**Sequential BRET-FRET Assay**

HEK-293T cells were transiently cotransfected with a constant amount of cDNA for MasR-Rluc (0.6 μg) and for AT1R-GFP\(^2\) (1 μg) and increasing amounts of cDNA for AT2R-YFP (0.5 to 4.1 μg). For the negative control, HEK-293T cells were transiently cotransfected with a constant amount of σ1R-Rluc (0.3 μg) and AT1R-GFP\(^2\) (1 μg) and increasing amounts of cDNA for AT2R-YFP (0.5 to 4.1 μg). To control the cell number, sample protein concentration was determined using a Bradford assay kit (Bio-Rad) using BSA dilutions as standards and adjusted to 0.2 mg/mL. To quantify fluorescent proteins, cells (20 μg of total protein) were distributed in 96-well microplates (black plates with transparent bottoms) and fluorescence was read in a Fluostar Optima Fluorimeter (BMG Labtech) equipped with a high-energy xenon flash lamp, using a 10-nm bandwidth excitation filter with a 485-nm excitation filter for YFP or 410-nm excitation filter for GFP\(^\text{a}\). For SRET measurements, the equivalent of 20 μg of total protein cell suspension was distributed in 96-well microplates (black plates with transparent bottoms) and fluorescence was read in a Fluostar Optima Fluorimeter (BMG Labtech) equipped with a high-energy xenon flash lamp, using a 10-nm bandwidth excitation filter with a 485-nm excitation filter for YFP or 410-nm excitation filter for GFP\(^\text{a}\). For SRET measurements, the equivalent of 20 μg of total protein cell suspension was distributed in 96-well white microplates (Corning 3600). SRET was determined 30 s after the addition of Deepblue C (5 μM) (Molecular Probes), using a Mithras LB 940 reader (Berthold Technologies), which allows the integration of the signals detected in the short-wavelength filter at 400 nm and the long-wavelength filter at 530 nm. To quantify MasR-Rluc or σ1R-Rluc expression, luminescence readings were obtained 10 min after the addition of 5 μM coelenterazine H.

**Assessment of Cytoplasmic Calcium Ion Levels**

HEK-293T cells were cotransfected with the cDNAs for AT\(_1\) (1 μg) and/or AT\(_2\) (1 μg), and/or MasR (1 μg) and/or the GCaMP6 calcium sensor (1 μg) [59] by the use of PEI method, as described above. Forty-eight hours after transfection, HEK-293T cells plated in 6-well plates (black, clear-bottomed) were incubated with Mg\(^2+\)-free Locke’s buffer (154 mM NaCl, 5.6 mM KCl, 3.6 mM NaHCO\(_3\), 2.3 mM CaCl\(_2\), 5.6 mM glucose, 5 mM HEPES, 10 μM glycine, pH 7.4). Receptor antagonists were added 10 min before readings and receptor agonists were added a few seconds before readings. Fluorescence emission intensity of GCaMP6 was recorded at 515 nm upon excitation at 488 nm on an EnSpire Multimode Plate Reader (PerkinElmer, Waltham, MA, USA) for 150 s every 5 s at 100 flashes per well.

**Determination of cAMP Level**

The analysis of cAMP levels in primary neural cells or in transfected HEK-293T was performed using the Lance® Ultra cAMP kit (PerkinElmer). Two hours before the experiment, cells were placed in serum-starved DMEM. Cells growing in the medium containing 50 μM zardaverine were distributed in 384-well microplates (2000 HEK-293T cells or 4000 striatal neurons or microglial cells per well) and pretreated with the AT\(_1\)R, AT\(_2\)R, and MasR antagonists candesartan, PD123319, and A779, respectively, or with the vehicle at room temperature for 15 min, and then stimulated with the AT\(_1\)R, AT\(_2\)R, and MasR agonists Ang II, CGP-42112A, and Ang(1-7), respectively, for 15 min before adding 0.5 μM forskolin or vehicle for an additional 15 min. Homogeneous time-resolved fluorescence energy transfer (HTRF) measures were performed using the Lance® Ultra cAMP kit (PerkinElmer). Fluorescence at 665 nm was measured on a PHERAstar Flagship microplate reader equipped with an HTRF optical module (BMG Labtech). A standard curve for (cAMP) was obtained in each experiment.

**Extracellular Signal-Regulated Kinases 1/2 Phosphorylation**

To determine ERK1/2 phosphorylation, 40,000 HEK-293T cells expressing MasR and either AT\(_1\)R or AT\(_2\)R, or 50,000 striatal neurons, or microglial primary cultures were plated in transparent Deltalab 96-well microplates. Two hours before the experiment, the medium was substituted with serum-starved DMEM. Cells were treated or not for 10 min with the selective antagonists (300 nM candesartan, 1 μM PD123319, or 500 nM A779) followed by 7-min treatment with the selective agonists (Ang II, CGP-42112A, or Ang(1-7)). Cells were then washed twice with cold PBS before the addition of lysis buffer (15-min treatment). Ten microliters of each supernatant was placed in white ProxiPlate 384-well microplates, and ERK1/2 phosphorylation was determined using an AlphaScreen SureFire kit (PerkinElmer) following the instructions of the supplier and using an EnSpire Multimode Plate Reader (PerkinElmer).

**Dynamic Mass-Redistribution Label-Free Assays**

Cell signaling was explored using an EnSpire Multimode Plate Reader (PerkinElmer) by using label-free technology. Cellular cytoskeleton redistribution movement induced upon receptor activation was detected by illuminating the underside of the plate with polychromatic light and measured as changes in the wavelength of the reflected monochromatic light, which
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is a sensitive function of the index of refraction. The magnitude of this wavelength shift (in picometers) is directly proportional to the amount of dynamic mass-redistribution (DMR). To determine the label-free DMR signal, 10,000 HEK-293T cells cotransfected with cDNAs for AT1R (1 μg) or AT2R (1 μg) and MasR or 10,000 striatal neuronal or microglial primary cultures were plated on transparent 384-well fibronectin-coated microplates to obtain 70 to 80% confluent monolayers, and kept in the incubator for 24 h. Before the assay, cells were washed twice with assay buffer (Hanks’ balanced salt solution with 20 mM HEPES, pH 7.15, 0.1% dimethyl sulfoxide) and incubated in the reader for 2 h in 30 μL/well of assay buffer at 24 °C. Then, the sensor plate was scanned and a baseline optical signature was recorded for 10 min before adding 10 μL of test antagonist (candesartan, PD123319, or A779) dissolved in assay buffer, followed by the addition of 10 μL of selective agonist (Ang II, CGP-42112A, or Ang(1-7)) also dissolved in assay buffer. The DMR responses induced by the agonist were monitored for a minimum of 3600 s.

Proximity Ligation Assay

Detection in natural sources of clusters formed by AT1 and Mas receptors or AT2 and Mas receptors was addressed in primary cultures of microglia or striatal neurons and in brain sections. Cells grown on glass coverslips were fixed in 4% paraformaldehyde for 15 min, washed twice with PBS containing 20 mM glycine to quench the aldehyde groups, permeabilized with the same buffer containing 0.05% Triton X-100 for 5 to 15 min, and washed with PBS. After a 1-h incubation at 37 °C with the blocking solution in a preheated humidity chamber, samples were incubated overnight at 4 °C with a mixture of a rabbit monoclonal anti-AT1R antibody (1/100, rabbit anti-AT1R monoclonal antibody produced by Calbiochem, Merck). To detect red dots corresponding to AT1-MasHets or AT2-MasHets, samples were observed in a Zeiss LSM 880 confocal microscope equipped with an apochromatic 63× oil-immersion objective, and 405- and 561-nm laser lines. For each field of view, a stack of 2 channels (one per staining) and 3 Z-planes with a step size of 1 μm were acquired. Andy’s algorithm, a specific ImageJ macro (National Institutes of Health, Bethesda, MD, USA) macro for reproducible and high-throughput quantification of the total PLA foci dots and total nuclei, was used for data analysis [60].

The specificity of antibodies against angiotensin receptors has been questioned, even though different laboratories have reported excellent performance of different antibodies see [58–61]. All antibodies used in this study were monoclonal; however, to check the specificity of the antibodies used, we performed experiments in HEK-293T cells, either non-transfected or expressing AT1R or AT2R. Signal was negligible in AT2R-expressing cells and non-transfected cells when the anti-AT1R antibody was used and negligible in AT1R-expressing cells and non-transfected cells when the anti-AT2R antibody was used (Supplementary Figure S2). These results confirm the antibody specificity as previously reported [62].

Data Analysis

All data were obtained from at least five independent experiments and are expressed as the mean ± standard error of the mean (SEM). GraphPad Prism 8 software (GraphPad Inc., La Jolla, CA, USA) was used for data fitting and statistical analysis. One-way ANOVA and post hoc Bonferroni’s test were used when comparing multiple values. When a pair of values was compared, Student’s t test was used. Significant differences were considered when the p value was < 0.05.

Results

The Mas Receptor Physically Interacts with AT1 or AT2 Receptors

Colocalization of MasR with AT1R or AT2R was first addressed using a heterologous expression system. Immunocytochemical assays performed in HEK-293T cells transfected with cDNAs encoding MasR fused to YFP and AT1 fused to Rluc (Fig. 1A–C) indicated a marked degree of colocalization (yellow in Fig. 1C). A lower degree of colocalization was observed after immunocytochemical assays performed in HEK-293T cells transfected with cDNAs encoding MasR fused to Rluc and AT2R fused to YFP (Fig. 1D–F). In those experiments, red fluorescence was due to a secondary Cy3-conjugated antibody, whereas green fluorescence was due to YFP (see “Materials and Methods”); colocalization can be observed by yellow in the merged image. It should be noted that the images are taken near the glass...
surface, that is, most of the labeling was due to proteins in the cell membrane proximal to the slide.

Immunocytochemistry assays do not demonstrate physical interactions because they occur within distances of 200 nm. Thus, we addressed potential interactions using BRET in HEK-293T cells expressing a constant amount of MasR-Rluc and increasing amounts of AT1R-YFP. The saturable BRET curve (BRET max 60 ± 2 mBU; BRET 50 13 ± 4 mBU) indicates specific interactions between Mas and AT1 receptors (Fig. 1G). A saturable curve was obtained when a similar experiment was carried out in HEK-293T cells expressing a constant amount of MasR-Rluc and increasing amounts of AT2R-YFP (BRET max 32 ± 1; BRET 50 7 ± 2), indicating that the two receptors do physically interact in living cells (Fig. 1H), confirming the results previously reported in HEK-293T cells using fluorescence resonance energy transfer and cross-correlation spectroscopy [40]. In contrast, experiments in cells coexpressing σ1R-Rluc and AT2-YFP showed a linear nonspecific signal (negative control in Fig. 1H).

We addressed the potential formation of trimers by taking advantage of the sequential resonance energy transfer (SRET; [63]) technique using cells expressing MasR-Rluc, AT1R-GFP, and AT2R-YFP. The saturable curve, which was not detected in the negative control (σ1R-Rluc, AT1R-GFP, and AT2R-YFP), indicated that the three RAS receptors interacted in a heterologous system, forming trimers (Fig. 2A). Trimer formation led to a structural rearrangement; in fact, increased expression of MasR led to a higher BRET signal due to the AT1R-Rluc and AT2R-YFP pair. These results are consistent with the reduced distance between Rluc fused to AT1R and
YFP fused to AT$_2$R when forming receptor heterotrimers with MasR (Fig. 2B).

Functional Characterization of AT$_1$R-MasR Heteromeric Complexes in HEK-293T Cells

Upon identification of AT$_1$-MasHets and AT$_2$-MasHets in cotransfected HEK-293T cells, we characterized the functionality of these complexes. It is well established that the AT$_1$ receptor couples to $G_q$ proteins, increasing inositol trisphosphate (IP$_3$) and diacylglycerol (DAG) levels by breaking phosphatidylinositol 4,5-bisphosphate (PIP$_2$) and subsequently releasing calcium (II) ion from endoplasmic reticulum channels, but it also couples to $G_i$ proteins, inhibiting adenylate cyclase and decreasing cAMP levels. In contrast, the MasR signal transduction pathway has not yet been fully characterized. Calcium measurements were first addressed in HEK-293T cells transfected with cDNAs for AT$_1$R-Rluc (0.5 $\mu$g), a constant amount of cDNA for AT$_2$R-YFP (3 $\mu$g), and increasing amounts of cDNA for MasR (0 to 1 $\mu$g). Values are the mean ± SEM of 5 independent experiments performed in triplicate. One-way ANOVA followed by Bonferroni’s multiple comparison post hoc tests were used for statistical analysis.

Fig. 2 All three RAS receptors interact in transfected HEK-293T cells. A Sequential resonance energy transfer (SRET$^2$) assays were performed in HEK-293T cells transfected with a constant amount of cDNA for MasR-Rluc (0.6 $\mu$g) or AT$_1$R-Rluc (0.3 $\mu$g) (as negative control), a constant amount of cDNA for AT$_2$R-GFP$^2$ (1 $\mu$g), and increasing amounts of cDNA for AT$_1$R-YFP (0.5 to 4.1 $\mu$g). Values are the mean ± SEM of 8 independent experiments performed in duplicate. B BRET assays were performed in HEK-293T cells transfected with a constant amount of cDNA for AT$_2$R-Rluc (0.5 $\mu$g), a constant amount of cDNA for AT$_1$R-YFP (3 $\mu$g), and increasing amounts of cDNA for MasR (0 to 1 $\mu$g). Values are the mean ± SEM of 5 independent experiments performed in triplicate. One-way ANOVA followed by Bonferroni’s multiple comparison post hoc tests were used for statistical analysis. C Schematic representation of the SRET$^2$ assay. *p < 0.05, **p < 0.001 versus absence of MasR.

Fig. 3 Functional analysis of AT$_1$R-MasR and AT$_2$R-MasR complexes in HEK-293T cells. HEK-293T cells were transfected with the cDNAs for MasR (1 $\mu$g) and for either AT$_1$R (1 $\mu$g) or AT$_2$R (1 $\mu$g) and, in assays of Ca$^{2+}$ determination, with the cDNA for an engineered calcium sensor, GCaMP6 (1 $\mu$g). Transfected cells were pretreated with the selective antagonists (300 nM candesartan for AT$_1$R, 1 $\mu$M PD123319 for AT$_2$R, and 500 nM A779 for MasR) and subsequently treated with selective agonists (100 nM Ang II for AT$_1$R, 300 nM CGP-42112A for AT$_2$R, and 500 nM Ang [1–7] for MasR). Thereafter, cytosolic calcium increases (A, B), intracellular cAMP levels (C, D), ERK1/2 phosphorylation (E, F), and the time-dependent DMR signal (G, H) were determined. Values are the mean ± SEM of 6 independent experiments performed in triplicate. One-way ANOVA followed by Bonferroni’s multiple comparison post hoc test were used for statistical analysis. *p < 0.05, **p < 0.01, ***p < 0.001 versus forskolin treatment in cAMP determinations or versus vehicle in pERK and DMR assays.
for AT₁ and Mas receptors (1 μg of cDNA each) and with the cDNA for the GCaMP6 calcium sensor (1 μg of cDNA). After treating cells with the AT₁R agonist Ang II (100 nM), a characteristic curve of cytoplasmic transient [Ca²⁺] increase was recorded. This ion mobilization was completely blocked in cells pretreated with the AT₁R...
antagonist candesartan (300 nM; Fig. 3A). Interestingly, when cells were pretreated with the MasR antagonist A779 (500 nM) followed by Ang II stimulation, we observed a marked increase in the calcium mobilization signal. Thus, MasR blockade potentiated AT1R-mediated signaling in the AT1-MasHet context. The MasR agonist Ang(1-7) (250 nM) induced no effect, indicating that the MasR receptor does not couple with Gq.

Because of the controversy existing around the pathways engaged by MasR activation, we assayed whether the receptor agonist could affect the cytoplasmic levels of cAMP. As can be observed in Fig. 3C, activation of MasR in HEK-293T cells coexpressing AT1 and Mas receptors (1 μg of cDNA each) did not have any effect on either basal or forskolin-induced cAMP levels, indicating that MasR does not couple with either Gi or Gs. Similar results were obtained in cells only expressing MasR (Supplementary Figure S3). Furthermore, Ang II stimulation induced no effect in cells expressing AT1-MasHets (Fig. 3C).

We next addressed activation of the MAPK pathway, which is linked to the function of many GPCRs. When HEK-293T cells expressing AT1-MasHets were treated with Ang II or Ang(1-7), observed an increase in ERK1/2 phosphorylation. However, when cells were simultaneously stimulated with both agonists, the signal was reduced (Fig. 3E). This phenomenon—in which the signal in a combined treatment is lower than the sum of individual activation—is known as negative cross-talk and it can be used to detect AT1-MasHets in native tissues. Furthermore, when cells were pretreated with selective antagonists, we observed that A779 treatment blocked not only the MasR-induced signal but also the AT1R-induced signal. Similarly, candesartan blocked both AT1R-induced and MasR-induced signals (Fig. 3E).

Finally, when AT1-MasR complex signaling was assayed by using label-free DMR, a technique that detects cytoskeletal rearrangements upon receptor activation, negative cross-talk was detected when cells were costimulated; cross-antagonism was unidirectional because it was only detected with candesartan (Fig. 3G).

### Functional Characterization of AT1R-MasR Heteromeric Complexes in HEK-293T Cells

After confirming that MasR and AT1R arrange into a functional unit with novel properties, we proceeded to analyze the possibility of a similar scenario for AT2R. The assays were similar to those described in the previous section. On the one hand, HEK-293T cells expressing AT2-MasHet and the GCaMP6 calcium sensor did not respond to the MasR agonist Ang [1–7] (250 nM) or to the AT2R agonist CGP-42112A (300 nM). These results fit with a lack of Gq coupling (Fig. 3B). On the other hand, assays to determine cAMP levels showed that CGP-42112A decreased the forskolin-induced effect by approximately 20%, whereas Ang [1–7] produced no effect (Fig. 3D). Similar results were obtained in cells expressing either AT2R (CGP-42112A) or MasR (Ang [1–7]) (Supplementary Figure S3). However, MasR activation partially blocked the AT2R signal. Interestingly, when HEK-293T cells expressing AT2-MasHets were pretreated with the MasR antagonist A779 followed by AT2R activation, potentiation of AT2R-mediated signaling was detected. Thus, these results show that MasR stimulation blocked AT2R-induced signaling, whereas, remarkably, MasR blockade potentiated AT1R functionality. To further characterize signaling in AT1-MasHet-expressing cells, ERK1/2 phosphorylation and DMR were determined. Equivalent results were found in both assays, namely negative cross-talk when receptors were simultaneously activated (Fig. 3F, H) and partial cross-antagonism in MAPK phosphorylation when cells were pretreated with selective antagonists. This partial cross-antagonism was not detectable in DMR assays (Fig. 3H). MasR blockade did not potentiate AT2R functionality in MAPK activation or DMR assays.

### AT1R-MasR and AT2R-MasR Heteromeric Complexes in Neuronal Primary cultures

Parkinson’s disease is characterized by neuronal death and neuroinflammation, mainly affecting the indirect pathway of the basal ganglia, where angiotensin receptors are expressed. Thus, we isolated primary cultures of brain striatum to look for expression of angiotensin and Mas receptor complexes.

We first identified AT1-MasHets and AT2-MasHets in primary neurons by in situ PLA. Clusters of receptor pairs were identifiable as red fluorescent dots (Fig. 4A–D); 1.44 red dots/cell were counted using primary antibodies against AT1R and MasR, and 0.63 red dots/cell were counted using primary antibodies against AT2R and MasR. The non-specific signal was equivalent to 0.17 red dots/cell in the negative control.

Once expression of heteromers was demonstrated, we addressed their functionality in striatal neurons. Receptors in primary cultures of striatal neurons were treated with selective antagonists (candesartan for AT1R, PD123319 for AT2R, and A779 for MasR) and activated with agonists (Ang II for AT1R, CGP-42112A for AT2R, and Ang(1-7) for MasR), and cAMP levels and MAPK activation were analyzed.

In the cAMP assays, we observed that only AT2R activation produced a significant decrease in forskolin-induced cAMP levels (Fig. 4E–G). This effect was partially counteracted by MasR activation and thus congruent with the data obtained in the heterologous expression system. In addition, pretreatment with MasR antagonist potentiated AT2R signaling.
Finally, except for AT$_2$R in cells expressing AT$_2$-MasHets, individual activation of receptors led to ERK1/2 phosphorylation. Furthermore, AT$_2$R activation blunted the effect of MasR activation. Antagonists allowed us to identify unidirectional cross-antagonism, that is, the MasR antagonist blocked the AT$_1$R-induced but not the AT$_2$R-induced effect, whereas angiotensin receptor antagonists did not affect the link of MasR and the MAPK pathway (Fig. 4H, I). Compared with results in transfected HEK-293T cells, the cross-antagonism of AT$_1$ and AT$_2$ receptors over MasR was not observed in neurons.

**Fig. 4** AT$_1$-MasHet and AT$_2$-MasHet expression and function in primary cultures of striatal neurons. A–C Expression of AT$_1$-MasHets and AT$_2$-MasHets heteromers was determined by proximity ligation assay (PLA), which was performed using specific primary antibodies against AT$_1$, AT$_2$, and Mas receptors. Confocal microscopy images (stacks of 4 consecutive planes) show heteroreceptor complexes; nuclei are Hoechst-stained (blue). Scale bar: 20 μm. D Bar graph showing the percentage of AT$_1$R-MasR and AT$_2$R-MasR clusters as red dots/cell compared with the negative control (*p < 0.05, **p < 0.01; Student’s t test vs the negative control condition). For cAMP (E–G) or ERK1/2 phosphorylation (H, I), cells were pretreated with selective receptor antagonists (300 nM candesartan for AT$_1$R, 1 μM PD123319 for AT$_2$R, and 500 nM A779 for MasR) and subsequently treated with selective agonists (100 nM Ang II for AT$_1$R, 300 nM CGP-42112A for AT$_2$R, and 250 nM Ang(1-7) for MasR). Values are the mean ± SEM of 5 independent experiments performed in triplicate. One-way ANOVA followed by Bonferroni’s multiple comparison post hoc tests were used for statistical analysis. *p < 0.05, **p < 0.01, ***p < 0.001 versus forskolin treatment in cAMP determinations or versus vehicle treatment (basal) in pERK determinations.

**AT$_2$R-MasR Heteromeric Complex Expression in Microglia Treated or Not with LPS and IFN-γ**

In pathological conditions, microglia migrate to the injury site, releasing pro- and anti-inflammatory factors, and becoming key actors in regulating the neurodegenerative/neuroprotective balance [64]. First, expression of the AT$_2$R-MasR and AT$_1$R-MasR heteromeric complexes was detected by PLA in primary microglia, both resting and activated. Microglia were activated by treating the cells for 48 h with 1 μM LPS and 200 U/mL IFN-γ. As can be observed in Fig. 5, 3.8 and 4.4 red dots/cell were
counted for AT1-MasHets (Fig. 5A) and AT2-MasHets (Fig. 5D), respectively, in resting microglia; in the negative control using the MasR primary antibody, only 0.2 red dots/cells were counted (Fig. 5C). These results indicate an expression level of AT1-MasHets and AT2-MasHets that seems markedly higher in resting microglia than in striatal neurons (see previous section). Interestingly, when the same experiment was performed in activated microglia, a significant decrease was observed in expression of the AT2R-MasR complex (3.1 red dots/cell) (Fig. 5E) but not in expression of the AT1R-MasR complex (4.2 red dots/cell) (Fig. 5B).

**AT1R-MasR Complexes Show Negative Cross-talk in cAMP and MAPK Signaling Pathways in Microglia Treated or Not with LPS and IFN-γ**

Signaling outputs were used to look for any differential functionality of heteromers in resting versus activated microglial cells. As observed in Fig. 6A, B, neither AT1R nor MasR activation resulted in any cAMP level alteration in resting microglia. Also, activation of each receptor led to MAPK activation (Fig. 6C). Moreover, simultaneous coactivation of resting microglia with the two agonists induced negative cross-talk, whereas experiments with antagonists showed cross-antagonism. Interestingly, when primary cultures were treated for 48 h with 1 mM LPS and 200 U/mL IFN-γ, Ang II led to a significant G i-mediated effect, indicating that the blockade of AT1R that occurred when forming complexes with MasR disappeared in activated microglia. This effect was potentiated when receptors were coactivated and cross-antagonism was not detected (Fig. 6D). By analyzing ERK1/2 phosphorylation data, we observed a similar effect of the MasR agonist in both resting and activated microglia. However, as in resting microglia, negative cross-talk and cross-antagonism were observed (Fig. 6E). To summarize, in activated microglia, the AT1-MasHets seem to undergo structural reorganization favoring the action of Ang II on G i-coupled AT1R.

**AT2R-MasR Complexes Show Negative Cross-talk in cAMP and MAPK Signaling Pathways in Microglia Treated or Not with LPS and IFN-γ**

As in the previous section, we observed differential effects in resting and activated microglia when the AT2R-MasR couple was analyzed with a functional perspective. The cAMP data indicated a non-significant effect upon AT2R or MasR activation in resting cells (Fig. 7A). Again, in resting microglia, a blockade of G i-mediated AT2R function was detected when AT2R was coexpressed with MasR. In resting cells, only MasR was linked to MAPK pathway activation but with...
marked negative cross-talk when the AT2R agonist was present and with a cross-antagonism effect (Fig. 7B). These results are similar to those obtained in the heterologous expression system.

In microglia activated for 48 h with 1 mM LPS and 200 U/mL IFN-γ, we observed that the AT2R agonist CGP-42112A induced a significant decrease in forskolin-induced cAMP levels (Fig. 7C). Activation of microglia provoked structural alterations in AT2-MasHets because cross-antagonism and negative cross-talk were not found; that is, there was no blockade by MasR over AT2R. In terms of ERK1/2 phosphorylation, the results were similar to those found in resting cells: only MasR was linked to MAPK pathway activation, with marked negative cross-talk and cross-antagonism (Fig. 7D).

Expression of Heteroreceptor Complexes in Parkinsonian and Dyskinetic Animals

In situ PLA is instrumental in detecting receptor heteromer clusters in natural sources. We performed PLA in striatal sections from the 6-OH-dopamine-based PD model.

Cytosolic cAMP levels (A, B, D) and ERK1/2 phosphorylation (C, E) were subsequently determined. Values are the mean ± SEM of 5 independent experiments performed in triplicate. One-way ANOVA followed by Bonferroni’s multiple comparison post hoc test were used for statistical analysis. *p < 0.05, **p < 0.01, ***p < 0.001 versus basal treatment (A) or forskolin treatment (B, D) in cAMP measurements or versus vehicle in pERK measurements (C, E).

Assays were performed in sections from non-lesioned and lesioned hemispheres in three animal groups (see “Materials and Methods”): untreated, levodopa-treated non-dyskinetic, and levodopa-treated dyskinetic. Both AT1R-MasR and AT2R-MasR clusters were expressed but at relatively low levels (Fig. 8) and their expression was markedly enhanced in the lesioned hemisphere (Fig. 8). Levodopa treatment per se did not lead to significant modification of expression in parkinsonian animals; however, in dyskinetic animals (i.e., animals that developed dyskinesias upon treatment with the drug), expression was modified. Importantly, the expression in dyskinetic animals (with respect to non-dyskinetic animals) was lower in the case of the AT2R-MasR heteromer but significantly higher in the case of AT1R-MasR (Fig. 8).

Discussion

At the beginning of the twenty-first century, research on GPCRs uncovered a novel property resulting from the
formation of heteroreceptor complexes. Receptor–receptor interactions lead to new functional units whose properties are different from those displayed by the individually acting receptors. Interestingly, consideration of receptor heteromers was first suggested for neurodegenerative conditions such as those occurring in PD. Heteromers containing adenosine and cannabinoid receptors are among those relevant to striatal function [65–71]. The occurrence of angiotensin receptors in

Fig. 7 AT$_2$-MasHet functionality in primary cultures of microglia. A–D Microglia in primary cultures were incubated for 48 h with 1 μM LPS and 200 U/mL IFN-γ (C, D) or vehicle (A, B). Microglial cells were pretreated with selective receptor antagonists (300 nM candesartan for AT$_1$R, 1 μM PD123319 for AT$_2$R, and 500 nM A779 for MasR) and subsequently treated with selective agonists (100 nM Ang II for AT$_1$R, 300 nM CGP-42112A for AT$_2$R, and 250 nM Ang [1–7] for MasR). Cytosolic cAMP levels (A, C) and ERK1/2 phosphorylation (B, D) were subsequently determined. Values are the mean ± SEM of 5 independent experiments performed in triplicate. One-way ANOVA followed by Bonferroni’s multiple comparison post hoc test were used for statistical analysis. *p < 0.05, **p < 0.01, ***p < 0.001 versus forskolin treatment in cAMP measurements (A, C) or versus vehicle in pERK assays (B, D).

Fig. 8 AT$_2$-MasHet and AT$_2$-MasHet complex expression in the striatum of lesioned and levodopa-treated animals. Heteroreceptor expression was assessed by the in situ proximity ligation assay (PLA) in the striatum of the different groups of animals treated as described in Methods. Graphs in the right represent the quantification of the red labeling in the different groups calculated using Andy’s algorithm. *p < 0.05, **p < 0.01, ***p < 0.001 versus non-lesioned animals.
the nigra and the striatum was followed by the identification of complexes formed by AT1R and AT2R [39, 42]. In this paper, we showed that the Mas receptor forms heteroreceptor complexes with AT1R and AT2R and that such complexes are relevant for better understanding the role of RAS in neuroinflammation.

The RAS imbalance has been well studied in relation to blood pressure alterations, which are exacerbated by aging [14, 72, 73]. However, alterations in RAS balance upon aging affects almost any physiological process. RAS regulates functions in the kidneys, lungs, and brain and thus is key for maintaining homeostasis both in the periphery and in the CNS. Unbalanced RAS upon aging is seen as an opportunity to intervene using ligands of RAS receptors to combat neurological diseases whose main risk factor is age. Parkinson’s and Alzheimer’s diseases are the most prevalent aging-related diseases affecting the CNS [74–76]. It is tempting to speculate that RAS is center stage in understanding the greater severity of COVID-19 in older patients and the variety of symptoms ranging from asymptomatic to serious lung, kidney, immunological, and/or neurological manifestations [19, 21, 22, 77].

A review in 2009 questioned whether results associated with the RAS system were due to cross-talk or to heteromerization [78]. Both AT1R-AT2R [39, 79] and AT2R-MasR interactions had previously been reported [4, 40]. In 2018, it was suggested that MasR and AT2R were “joining forces” to counteract deleterious actions on blood pressure mediated by AT1R and that AT2-MasHets could help explain the functionality of AT1R [41]. A scenario of RAS receptors interacting with each other raised numerous questions, particularly to understand why receptors mediating opposing effects would “join forces.” To approach the issue, two lines of inquiry are necessary: [1] assessing the expression levels of the different components of the system in targeted cells/tissues, heteromers included; and [2] deciphering the properties of the units resulting from receptor–receptor interactions. The functionality studies reported here show differential behaviors depending on the signaling pathway and inter-receptor interactions that lead to negative cross-talk, even reaching full blockade and/or antagonism. Such counterbalancing effects occurred for both AT1-MasHets and AT2-MasHets. Cross-antagonism was reported for AT2-MasHet expressed in mouse astrocytes using the mRNA level of the CX3C chemokine receptor-1 as a read-out [40]. Apart from cross-antagonism, which is often a characteristic of heteromer expression, a novel feature that is also explained by inter-receptor interaction is that the antagonist of MasR increased the calcium peak elicited by the AT1R agonist [31, 80, 81]. The properties of these heteromers must be considered to understand the efficacy of antihypertensive medication targeting RAS and for repurposing such approved drugs.

Unbalanced RAS associated with Parkinson’s and Alzheimer’s diseases opens new therapeutic perspectives. Pharmacological manipulation of RAS components has potential in PD [82]; it is suggested that antagonists already used to treat hypertension and able to cross the blood–brain barrier may be repurposed to treat PD [83]. On the other hand, the system has been extensively characterized in relation to nigral neurodegeneration [27, 72, 83–85]. On the other hand, there are confounding factors in assessing the risk of PD in individuals taking antihypertensive medication. This is due to the multiplicity of therapeutic choices but also to the fact that some drugs targeting RAS enter the brain (e.g., candesartan) [86], whereas others cannot cross the blood–brain barrier. Therefore, we cannot assess with certainty whether chronic administration of RAS-related antihypertensives is neuroprotective or whether they can delay neurodegeneration once the disease shows clinical symptoms. Selecting the most appropriate cell to be targeted is also important. It is feasible to target activated microglia to prevent neurodegeneration if the disease has a neuroinflammatory component. In addition, MasR is key for microglia-driven development of the retinal vasculature [87]. The results presented here and those related to expression of AT1R-AT2R and AT2R-MasR heteromers in striatal cells [42] suggest that MasR-related therapies to delay progression of PD should consider RAS components in microglia, with the ultimate goal of attenuating inflammation or skewing microglia toward the M2 neuroprotective phenotype [64]. RAS is well positioned to be targeted to polarization of microglia activated upon neuroinflammation [82]. In contrast to results in transfected HEK-293T cells and activated microglia, we did not observe cross-antagonism of AT1 and AT2 receptors over MasR in neurons. This, together with the lesser expression of MasR-containing heteromers, suggests that MasR functionality in neurons is not efficiently regulated by Ang II acting on AT1 or AT2 receptors.

Further parameters to consider are the expression of ACE1 and ACE2 and the possibility that these enzymes also interact with RAS receptors, thus affecting the local concentration of agonists of RAS receptors. In relation to COVID-19 management, it has been demonstrated that ACE2 interacts with AT1R [88]. Combining these results with the results presented here, we cannot rule out the occurrence of a functional unit constituted by an enzyme, two receptors, and the corresponding G-coupled proteins. The functionality of equivalent macromolecular complexes and the 3-dimensional structure underlying the particular functional features has been demonstrated for adenosine A1–A2A and A2A–A2B receptors ([33–38] and data in preparation). Both the allosteric modulation of ACE2 activity that results from the interaction with SARS-CoV viruses and the effect of viral infection on the surface expression of RAS components, heteromers included, need to be addressed.

Mechanistically, the two most relevant findings are the MasR-mediated regulation of cytosolic calcium mobilization triggered by AT1R agonists and the differential link between RAS and MAPK activation, depending on which heteromers are expressed in a given cell type and which
agonist(s) is (are) affecting RAS receptors. Very few mechanisms lead to radical changes in the activation of the MAPK pathway. However, we present here data that show that one of the properties of RAS receptor heteromers is the possibility to engage or not this relevant pathway, depending on the overall RAS balance. Several years ago, we reported that histamine H3 receptor activation could not lead to ERK phosphorylation unless it formed a heteromer with the dopamine D1 receptor. D1-H3 receptor heteromers are unique devices directing histaminergic and dopaminergic signaling toward the MAPK pathway in a G1-dependent but G2-independent manner [89]. Remarkably, MasR in activated microglia is linked to MAPK pathway activation with marked negative cross-talk and cross-antagonism. In a complementary recent study, we showed that AT1-AT2 receptor heteromers are expressed in microglia where [1] they are upregulated in both parkinsonian conditions and in L-DOPA-induced dyskinesias, and [2] their activation is seemingly neuroprotective. The data shown in Fig. 8 suggest that expression of heteroreceptor complexes formed by MasR and either AT1R or AT2R is higher in the striatum of the lesioned hemisphere in the rat 6-OH-DA-based PD model. The marked increase in the case of AT1R-MasR heteromers in dyskinetic animals open new therapeutic avenues for the assessment of this well-known side effect of antiparkinsonian medication. In microglia, a differential pharmacological trend, when Ang II receptor heteromers are compared with AT1-MasHets or AT2-MasHets, is cross-antagonism, which is not found in the former but in the latter. This is promising from a therapeutic point of view, because cross-antagonism in AT1R-AT2R heteromers would lead to a dead end in terms of neuroprotection; instead, the antagonist of one Ang II receptor releases the brake on activation of the partner Ang II receptor within the AT1R-AT2R heteromer [42].

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Data Availability The data that support the findings of this study are available from the corresponding author upon reasonable request.

Compliance with Ethical Standards

Conflict of Interest The authors declare that they have no conflict of interest.

Statement of Ethics Under the current legislation, obtaining protocol approval is not needed if animals are sacrificed to obtain a specific tissue. Animal handling, sacrifice and further experiments were conducted according to the guidelines set in Directive 2010/63/EU of the European Parliament and the Council of the European Union that is enforced in Spain by National and Regional organisms; also the 3R rule (replace, refine, reduce) for animal experimentation was taken into account.

References

1. Villela DC, Passos-Silva DG, Santos RAS, Alamandine: A new member of the angiotensin family. Vol. 23, Current Opinion in Nephrology and Hypertension. Curr Opin Nephrol Hypertens; 2014. p. 130–4.
2. Souza LL, Duchene J, Todiras M, Azevedo LCP, Costa-Neto CM, Alenina N, et al. Receptor mas protects mice against hypothermia and mortality induced by endotoxemia. Shock. 2014;41(4):331–6.
3. Santos RAS, Simoes e Silva AC, Maric C, Silva DMR, Machado RP, de Buhr I, et al. Angiotensin(1-7) is an endogenous ligand for the G protein-coupled receptor Mas. Proc Natl Acad Sci U S A [Internet]. 2003;100(14):8258–63. Available from: http://www.pnas.org/content/100/14/8258.full
4. Villela D, Leonhardt J, Patel N, Joseph J, Kirsch S, Hallberg A, et al. Angiotensin type 2 receptor (AT 2 R) and receptor Mas: a complex liaison. Clin Sci [Internet]. 2015;128(4):227–34. Available from: http://www.clinsci.org/content/128/4/227. abstract?5Ch, http://clinscilookup/10.1042/CS2015015
5. De Carvalho Santuchi M, Dutra MF, Vago JP, Lima KM, Galvão I, De Souza-Neto FP, et al. Angiotensin-(1-7) and Alamandine Promote Anti-inflammatory Response in Macrophages in Vitro and in Vivo. Mediators Inflamm [Internet]. 2019 [cited 2020 Apr 29];2019(1):2401081. Available from: https://doi.org/10.1155/2019/2401081
6. Clarke NE, Turner AJ. Angiotensin-Converting Enzyme 2: The First Decade. Int J Hypertens [Internet]. 2012;2012:1–12. Available from: http://www.hindawi.com/journals/ijhy/2012/ 307315/
7. Wan Y, Shang J, Graham R, Baric RS, Li F. Receptor Recognition by the Novel Coronavirus from Wuhan: an Analysis Based on Decade-Long Structural Studies of SARS Coronavirus. J Virol. 2020 Jan 29;94(7).
8. Shang J, Ye G, Shi K, Wan Y, Luo C, Aihara H, et al. Structural basis of receptor recognition by SARS-CoV-2. Nature. 2020 Mar 30;581(7807):221–4.
9. Kuhn JH, Li W, Choe H, Farzan M. Angiotensin-converting enzyme 2: A functional receptor for SARS coronavirus [Internet]. Vol. 61, Cellular and Molecular Life Sciences. 2004 [cited 2020 May 26]. p. 2738–43. Available from: https://www.nature.com/articles/nature02145
10. Li W, Moore MJ, Vasilieva N, Sui J, Wong SK, Berne MA, et al. Angiotensin-converting enzyme 2 is a functional receptor for the SARS coronavirus. Nature. 2003 Nov 27;426(6965):450–4.
11. Valenzuela R, Barroso-Chinea P, Villar-Cheda B, Joglar B, Muñoz A, Lancelio JL, et al. Location of Prorenin Receptors in Primate Substantia Nigra: Effects on Dopaminergic Cell Death. J Neuropathol Exp Neurol [Internet]. 2010 Nov [cited 2019 Sep 1];69(11):1130–42. Available from: http://www.ncbi.nlm.nih.gov/pubmed/20940627
12. Jarrott B, Williams SJ. Chronic Brain Inflammation: The Neurochemical Basis for Drugs to Reduce Inflammation. Neurochem Res [Internet]. 2016 Mar 16 [cited 2019 Oct 4];41(3): 523–33. Available from: http://www.ncbi.nlm.nih.gov/pubmed/26177578
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