



TRPA1 modulation by Sigma-1 receptor prevents oxaliplatin-induced painful peripheral neuropathy

Aida Marcotti,^{1,†,‡} Jorge Fernández-Trillo,^{1,†} Alejandro González,^{1,§} Marta Vizcaíno-Escoto,¹ Pablo Ros-Arlanzón,^{1,‡} Luz Romero,² José Miguel Vela,² Ana Gomis,¹ DFélix Viana¹ and Elvira de la Peña^{1,3}

[†]These authors contributed equally to this work.

Chemotherapy-induced peripheral neuropathy is a frequent, disabling side effect of anticancer drugs. Oxaliplatin, a platinum compound used in the treatment of advanced colorectal cancer, often leads to a form of chemotherapy-induced peripheral neuropathy characterized by mechanical and cold hypersensitivity. Current therapies for chemotherapy-induced peripheral neuropathy are ineffective, often leading to the cessation of treatment. Transient receptor potential ankyrin 1 (TRPA1) is a polymodal, non-selective cation-permeable channel expressed in nociceptors, activated by physical stimuli and cellular stress products. TRPA1 has been linked to the establishment of chemotherapy-induced peripheral neuropathy and other painful neuropathic conditions. Sigma-1 receptor is an endoplasmic reticulum chaperone known to modulate the function of many ion channels and receptors. Sigma-1 receptor antagonist, a highly selective antagonist of Sigma-1 receptor, has shown effectiveness in a phase II clinical trial for oxaliplatin chemotherapy-induced peripheral neuropathy. However, the mechanisms involved in the beneficial effects of Sigma-1 receptor antagonist are little understood. We combined biochemical and biophysical (i.e. intermolecular Förster resonance energy transfer) techniques to demonstrate the interaction between Sigma-1 receptor and human TRPA1. Pharmacological antagonism of Sigma-1R impaired the formation of this molecular complex and the trafficking of functional TRPA1 to the plasma membrane. Using patch-clamp electrophysiological recordings we found that antagonists of Sigma-1 receptor, including Sigma-1 receptor antagonist, exert a marked inhibition on plasma membrane expression and function of human TRPA1 channels. In TRPA1-expressing mouse sensory neurons, Sigma-1 receptor antagonists reduced inward currents and the firing of actions potentials in response to TRPA1 agonists. Finally, in a mouse experimental model of oxaliplatin neuropathy, systemic treatment with a Sigma-1 receptor antagonists prevented the development of painful symptoms by a mechanism involving TRPA1. In summary, the modulation of TRPA1 channels by Sigma-1 receptor antagonists suggests a new strategy for the prevention and treatment of chemotherapy-induced peripheral neuropathy and could inform the development of novel therapeutics for neuropathic pain.

- 1 Instituto de Neurociencias de Alicante, Universidad Miguel Hernández-CSIC, 03550 San Juan de Alicante, Spain
- 2 WeLab Barcelona, Parc Científic de Barcelona, 08028 Barcelona, Spain
- 3 Institute for Health and Biomedical Research (ISABIAL), 03010 Alicante, Spain
- ‡ Present address: Instituto de Farmacología Experimental de Córdoba (IFEC) CONICET, Haya de la Torre y Medina Allende, Ciudad Universitaria, Córdoba 5000, Argentina
- § Present address: Division of Molecular Neurobiology, Department of Medical Biochemistry and Biophysics, Karolinska Institutet, Stockholm, Sweden

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Present address: Institute for Health and Biomedical Research (ISABIAL), 03550 San Juan de Alicante, Spain

Correspondence to: Elvira de la Peña Instituto de Neurociencias de Alicante Universidad Miguel Hernández-CSIC 03550 San Juan de Alicante, Spain E-mail: elvirap@umh.es

Correspondence may also be addressed to: Felix Viana E-mail: felix.viana@umh.es

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Introduction

Chemotherapy-induced peripheral neuropathy (CIPN) is a frequent, disabling side effect of different anticancer treatments, including taxanes, vinca alkaloids, bortezomid and platinum analogues, affecting millions of patients worldwide (reviewed by Staff et al.,¹ Gordon-Williams and Farquhar-Smith² and Seretny et al.³). In the case of oxaliplatin, a platinum compound frequently used in the treatment of colorectal cancer, the neuropathy affects over 70% of treated patients3 and manifests itself with paraesthesias, dysesthesias and pain, preferentially in hands and feet.^{1,4} The severity of CIPN painful symptoms, including mechanical and cold hypersensitivity often lead to dose reduction or early cessation of treatment, compromising therapeutic effectiveness and survival.⁵ The chronicity of many cases has a long-term negative impact on patient's quality of life. There is no effective therapy to treat or prevent CIPN. The results of many clinical interventions have been negative or inconclusive⁶; current therapies are primarily geared towards the relief of pain symptoms.⁷

The mechanisms underlying painful CIPN are complex and poorly understood (reviewed by Starobova and Vetter,⁸ Ma et al.⁹ and Colvin¹⁰). Proposed mechanisms include mitochondrial dysfunction leading to oxidative stress in peripheral tissues, ^{11–13} neuroimmune alterations^{14,15} and ion channel dysregulation in sensory afferents.^{16–20} In particular, several studies implicated the ion channel transient receptor potential ankyrin 1 (TRPA1) in the painful symptomatology observed in several animal models of CIPN,²¹ including those caused by oxaliplatin administration.^{22–25} TRPA1 is a polymodal, non-selective cation channel expressed in a fraction of nociceptors,²⁶ where it senses noxious temperatures, mechanical forces and a broad array of exogenous and endogenous irritants, including reactive oxygen species (reviewed by Viana²⁷ and Talavera et al.²⁸). The role of TRPA1 as a molecular detector of mechanical and thermal stimuli is particularly evident in pathological conditions causing neuroinflammation or neuropathy.²⁹

Sigma-1 receptor (Sigma-1R) is a ligand-regulated chaperone residing at mitochondria-associated endoplasmic reticulum membranes, expressed in many tissues, including peripheral sensory neurons.³⁰⁻³² Sigma-1R is involved in multiple cellular functions, such as cellular bioenergetics, calcium homeostasis, reactive oxygen species generation, chromatin remodelling, inflammation and cell proliferation.³²⁻³⁶ Sigma-1R can translocate to other cellular compartments, such as the plasma membrane, regulating the expression and function of many ion channels.³⁷⁻³⁹ Previous studies linked Sigma-1R with pain modulation in animal models.^{40,41} Moreover, a recent phase II clinical trial established the efficacy of sigma 1 receptor antagonist (S1RA) (also known as E-52862), a highly selective Sigma-1R antagonist,⁴² in ameliorating oxaliplatin-induced peripheral neuropathy in patients treated for colorectal cancer.⁴³ Treatment with S1RA significantly reduced cold pain threshold temperature and cold-evoked pain intensity. However, the cellular and molecular mechanism involved in pain relief by Sigma-1R ligands are currently unknown.

We found that antagonism of Sigma-1R inhibits TRPA1 activity by a mechanism involving alterations in TRPA1-Sigma-1R complex formation, channel membrane trafficking and reduced expression at the plasma membrane of nociceptors. In an experimental model of oxaliplatin neuropathy, *in vivo* Sigma-1R antagonists reduced TRPA1 activity and prevented the painful mechanical and cold symptoms caused by the drug. This finding paves the way for innovative treatments of painful CIPN by targeting Sigma-1R.

Materials and methods

All experimental procedures abide to the Spanish Royal Decree 53/ 2013 and the European Community Council directive 2016/63/EU, regulating the use of animals in research. The Committee on Animal Research at the University Miguel Hernández approved all the animal procedures.

Animals

Studies were performed on young adult (2–4 months old) male and female C57BL/6J wild-type (WT) mice (https://www.jax.org/ strain/000664 Jackson Laboratory) and the transgenic mouse lines detailed next. Their genotype was confirmed by polymerase chain reaction. Mice were bred at the Universidad Miguel Hernández Animal Research Facilities (ES-119-002001) and kept behind a specific pathogen free barrier under 12/12-h light dark cycle with access to food and water *ad libitum*. TRPA1 knockout mice (TRPA1 KO), bred on a C57Bl/6J background, were kindly provided by Dr David Corey.⁴⁴ To identify TRPA1-expressing neurons, Trpa1 Cre mice (kindly provided by Dr Mark Hoon, NIH NIDCR) were crossed to the a tdTomato reporter line, Ai14 (https://www.jax.org/strain/007908, Jackson laboratory), both backcrossed to a C57Bl/6J background and named Trpa1 Cre-tdTomato for simplicity.

Oxaliplatin injection

Mice received three intraperitoneal (i.p.) injections of oxaliplatin (6 mg/kg) or vehicle (glucose 5% in dH_2O) blindly, on alternate days (Day 0, Day 2, Day 4) (Fig. 6A). Oxaliplatin (Tocris) was freshly dissolved in 5% glucose solution prepared in sterile distilled water. Behavioural tests (see next) were used to evaluate the development of neuropathic symptoms.

Systemic treatment with S1RA

S1RA, also known as E-52862 (40 mg/kg, ESTEVE Pharmaceuticals S.A.) or vehicle (sterile saline 0.9% NaCl, B-Braun) was injected intraperitoneally once a day for 11 consecutive days, starting 3 days before oxaliplatin injection (Figs 6G and 7C). Behavioural tests were used to evaluate antinociceptive effects.⁴⁵

Intradermal injection of AITC

Ten microlitres of 10 mM allyl isothiocyanate (AITC) or vehicle (ethanol) were blind injected subcutaneously into the plantar surface of the right hind paw. Nocifensive behaviour was videotaped and the time spent licking the injected paw was measured for 5 min.⁴⁶

Behavioural assays

Before testing, mice were habituated to the behavioural room for a minimum of 1 week and placed inside a transparent Plexiglass box for at least 1 h. The same investigator performed the scoring in all behavioural tests, which was blind in respect to the treatment or mouse strain. Both male and female mice were evaluated in all experimental groups, except in the experiments related to Fig. 6H, I and J; on this occasion the experiments were performed on male mice. The timeline of behavioural testing is diagrammed in the relevant figures. Each animal was evaluated for mechanical sensitivity and the acetone test before and after oxaliplatin treatment. The unilateral cold plate test was only performed after treatment, comparing results of drug- and vehicle-treated groups.

Mechanical threshold

Changes in mechanical withdrawal threshold were assessed with calibrated von Frey filaments (BIO-VF-M model from Bioseb-In vivo Research Instruments) applied to the plantar side of the hind paw, using the 'up and down' method described by Llorian-Salvador et al.⁴⁷ Mice were placed in transparent bottomless plastic boxes on a metal mesh (5 × 5 mm hole size) platform and von Frey filaments, size (force in g) = 2.44 (0.04), 2.83 (0.07), 3.22 (0.16), 3.61 (0.4), 4.08 (1), 4.56 (4), were applied to the plantar surface of one hind paw for up to 1 s. Measurements started with the 3.61 filament size, randomly starting with the left or right paw. The observation of a positive response (lifting, shaking or licking of the paw) was followed by the application of the immediate thinner filament or the immediate thicker one if the response was negative, a series of six measurements were obtained in each animal. The 50% response threshold was calculated using the following formula: 50% g threshold = $(10^{Xf+kd})/10000$; where Xf is the size value of the last von Frey filament applied, k, is the correction factor based on pattern of responses (Dixon's calibration table), d is the mean distance between size's filaments stimuli (here, 0.39). Reported values are the averages of left and right hind paws. Threshold values were assessed in mice 3 days before the first oxaliplatin or glucose injection as control baseline.

Acetone testing

To test acetone-evoked evaporative cooling-induced sensations, mice were covered with a round plastic chamber placed on a metal mesh platform. A drop (50 μ l) of acetone was applied onto the plantar surface of the hind paw, and the number of lifting, licking, biting, shaking and finger-opening events were counted over the next 60 s. Results obtained after acetone application to the left and right paw were averaged. A baseline value was obtained 3 days before the first oxaliplatin or glucose injection.

Unilateral cold plate

Mice were gently held with the hind paws resting on a flat surface at room temperature for 30 s. Thereafter, the plantar side of one paw was gently placed on a cold plate (Cold/hot plate, Bioseb-In vivo Research Instruments) set to 10°C for a maximum 30 s. Measurements of withdrawal latencies from the cooled surface of each hind paw were made separately at 2 min intervals, and the mean of both measures was considered.

Culture and transfection of cells lines

Human embryonic kidney -293 (HEK293) cells were obtained from the European Collection of Authenticated Cell Cultures. Cells grown at 37°C in Dulbecco's modified Eagle were medium (DMEM) containing foetal bovine serum (FBS) and 1% penicillin/streptomycin in a 5% CO₂ atmosphere. Cells were seeded in 24-well dishes at 2×10^5 cells/well and, 48 h after plating, were transfected with different plasmid DNA using Lipofectamine™ 2000 (Thermo Fisher Scientific); 1 µg total DNA and 2 µl Lipofectamine[™] per well were used, respectively. For transfection, LipofectamineTM was mixed with the DNA in 100 μ l of OptiMem (Thermo Fisher Scientific). Cells were trypsinized and replated on poly-L-lysine-coated round coverslips (12 mm diameter) at 80 000 cells/coverslip at 24-48 h post-transfection. Fluorescent protein (+) cells were selected for calcium or electrophysiological recordings. The different plasmid DNA used and their source are described in Supplementary Table 1.

In some experiments, hTRPA1-HEK293 cells stably expressing human TRPA1 channels were used. This cell line was obtained by geneticin resistance selection; HEK293 cells were transfected with hTRPA1-tGFP and cultured in DMEM containing of 10% FBS, 1% penicillin/streptomycin and 450 µg/ml Geneticin (G418).

HEK293 cells KO for Sigma-1R (S1R-KO), provided by ESTEVE Pharmaceuticals S.A., were generated with CRISPR–Cas9 technology. The absence of Sigma-1R protein was verified by western blot and radioligand binding assays using the selective ligand [3H]-(+) pentazocine. All cell lines were tested for mycoplasma and confirmed free.

Isolation and culture of mouse dorsal root ganglion neurons

Dorsal root ganglion (DRG) neurons from mice were dissociated as previously described.⁴⁸ Briefly, mice of either sex were anaesthetized with isoflourane and killed by cervical dislocation. The spinal cord was removed and 25–30 DRGs were extracted and washed in cold Hank's buffered saline solution (Invitrogen). Ganglia were incubated in 900 U/ml collagenase (type XI Sigma) and 5.46 U/ml dispase (Invitrogen) for 45 min at 37°C in 5% CO_2 . After that, they were gently triturated using a flame-polished glass pipette in calcium free culture medium Hank's buffered saline solution (Invitrogen) + 1% MEM-Vit (Invitrogen) + 10% FBS (Invitrogen) + 100 mg/ml penicillin/streptomycin (Invitrogen) and the resulting suspension was centrifuged. The pellet obtained was resuspended in culture medium with the following composition: MEM (Invitrogen) + 1% MEM-Vit (Invitrogen) + 10% FBS (Invitrogen) + 100 mg/ml penicillin/streptomycin (Invitrogen). Cells were plated on round glass coverslips treated with 0.01% poly-L-lysine (Sigma) and kept at 37°C, 5% CO_2 .

Ca²⁺ microfluorimetry

Ratiometric calcium imaging experiments were conducted with the fluorescent indicator Fura-2AM (Fura-2, Thermo Fisher Scientific). Cells were loaded with $5\,\mu\text{M}$ Fura-2 for 45 min and incubated at 37°C in a 5% CO₂ atmosphere. Recordings were performed in a lowvolume chamber continuously perfused. Bath solution contained (in mM) 140 NaCl, 3 KCl, 2.4 CaCl₂, 1.3 MgCl₂, 10 HEPES and 10 glucose adjusted to pH 7.4 with NaOH. Fluorescence measurements were made with a Leica inverted microscope (Leica DMI 3000B, Leica Microsystems). Fura-2 was excited using a xenon arc lamp (Sutter Instruments) at 340 and 380 nm wavelengths, switching between both with a computer-controlled filter wheel (Lambda 10-2; Sutter Instruments), and the emitted fluorescence was long-pass filtered at 510 nm. Images were acquired using an ORCA ER CCD camera (Hamamatsu Photonics K.K.) at a frequency of 0.33 Hz and analysed with Metafluor software (Molecular Devices). Cytosolic Ca²⁺ increases are presented as the ratio of the emission intensities after excitation at 340 and 380 nm (F₃₄₀/F₃₈₀). Measurements in DRG neurons were performed at $33 \pm 1^{\circ}$ C. Unless indicated otherwise, calcium measurements in cells lines were performed at $24 \pm 1^{\circ}$ C, controlled by a homemade temperature feedback device.

Electrophysiology

Current and voltage signals were recorded with a Multiclamp 700B or an Axoclamp 200B amplifier in conjunction with the pCLAMP 10.1 software suite and a Digidata 1322A digitizer (Molecular Devices). Intracellular solution for whole-cell recordings in HEK293 cells contained (in mM): 135 CsCl, 1 EGTA, 2 MgCl₂, 5 Na₂ATP 5, 0.1 Na-GTP 0.1 and 10 HEPES, adjusted to pH 7.2 with CsOH; 300 mOsm/l. DRG neurons were recorded using a pipette solution containing (in mM): 115 K-gluconate, 25 KCl, 9 NaCl, 1 MgCl₂, 0.2 EGTA, 10 HEPES, 3 K₂-ATP, 1 Na-GTP, pH 7.3 adjusted with NaOH; 300 mOsm/l. In both cases, bath solution contained (in mM): 140 NaCl, 3 KCl, 2.4 CaCl₂, 1.3 MgCl₂, 10 HEPES and 10 glucose adjusted to pH 7.4 with NaOH. Measurements in neurons were performed at 33 ± 1°C while measurements in cell lines were performed at room temperature ($24 \pm 1^{\circ}$ C). Patch electrodes were made from borosilicate glass using a P-97 Flaming/Brown micropipette puller (Sutter Instruments) and had resistances between 5-8 $M\Omega$ when filled with the different internal solutions. In voltage-clamp whole-cell configuration, series resistance (Rs) was compensated to >70%, cells were clamped at -60 mV and voltage ramps applied from -100 to 100 mV (0.33 mV/ms) at 0.33 Hz. Whole-cell currents were acquired at 10 kHz and filtered at 2 kHz. Neuronal recordings, acquired at 20 kHz and filtered at 2 kHz, were adjusted for a +13 mV pipette-liquid junction potential.

Immunoprecipitation, cell surface biotinylation and western blot

For immunoprecipitation, HEK293 cells were transfected with hTRPA1-tGFP and mSigma-1R-EYFP fusion proteins. They were harvested, 48 h after transfection, at 800g for 10 min and washed with cold phosphate-buffered saline (PBS) twice. Pellets were lysed (Buffer lysis: 50 mM Tris-HCl, pH 7.5, 120 mM NaCl, 0.5 mM EDTA, 0.5% Nonidet P-40 with phosphatase and protease inhibitors) and centrifuged at 4°C. Supernatants were recovered, precleared and incubated with green fluorescent protein- (GFP-)Trap agarose beads (Chromotek) overnight at 4°C under gentle rotation. Note that GFP-Trap agarose beads do not recognize t-GFP (Chromotek specifications). Beads were washed three times with Tris-buffered saline containing protease and phosphatase inhibitors and finally eluted with $2\times$ sodium dodecyl sulphate (SDS) sample buffer (0.5 M Tris-HCl, pH 6.8, 10% SDS, 10% glycerol, 2-mercaptoethanol).

Biotinylation assays were performed using the Pierce Cell Surface Protein Isolation Kit (Thermo Fisher Scientific). Briefly, hTRPA1-HEK293 cells (stable cell line) were incubated with 25 µM S1RA or with 2 µl dimethyl sulphoxide (DMSO) as a control for 24 h. Then, cells were washed twice with ice-cold PBS pH 7.2 and incubated with sulfo-NHS-SS-biotin in PBS for 30 min at 4°C or with PBS as a negative control of biotinylation. Hereafter quenching free biotin, biotinylated and control cells were lysed in 300 µl of lysis buffer with a protease inhibitor cocktail (cOmplete Mini Roche, Sigma-Aldrich). Cell lysates were sonicated for 10 min at 4°C, and finally harvested at 10 000g for 15 min at 4°C. For each experimental condition, proteins were quantified (Pierce TM BCA Protein Assay Kit, Thermo Fisher Scientific) and the same amounts (200–300 µg) of protein were incubated with NeutrAvidin beads (pretreated with 1% BSA) for 1 h at room temperature, after collecting an aliquot as input control. Beads were washed with wash buffer and then proteins were eluted with 2× SDS sample buffer (62.5 mM Tris-HCl, pH 6.8, 1% SDS, 10% glycerol and 50 mM DTT) for immunoblot analysis

In immunoprecipitation and cell surface biotinylation assays, proteins were subjected to SDS–polyacrylamide gel electrophoresis and blotted onto Protran Nitrocellulose Membrane-Whatman (GE Healthcare Life Science). Membranes were blocked and incubated overnight at 4°C with the indicated primary antibody: anti-TRPA1 (1:500, NB110-40763 Novus), anti-GFP/EYFP (1:2000, A6455, Thermo Fisher Scientific), anti-tGFP (1:2000, TA150096 Origene), anti-GAPDH (1:5000 G9545, Sigma-Aldrich) and anti-Sigma-1R (1:100 sc-137075, Santa Cruz Biotechnology). Hereafter, membranes were washed and incubated with secondary antibody (Horseradish 1:2000 Sigma-Aldrich) and developed with ECLplus (Thermo Fisher Scientific). Image background subtraction and quantification analysis were performed with Fiji (ImageJ 1.53c, NIH).

Förster resonance energy transfer

HEK-293 cells were cotransfected with hTRPA1-tGFP and hSigma-1R-mCherry, and excited with 488 nm light to measure net Förster resonance energy transfer (FRET) between both fluorophores. Images for GFP and Cherry fluorescence were obtained using a Nikon TE-2000-U Eclipse fluorescence microscope (Nikon) using a ×60/numerical aperture1.49 oil immersion objective. The excitation light was generated by a Polychrome V monochromator (Thermo Fisher Scientific). The filter cube contained a dual band GFP/RFP filter set (51022-EGF/mRFP-1; Chroma Technology). GFP and Cherry emissions were simultaneously collected using a

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Dual-View Micro-Imager (Optical Insights) in which tGFP and Cherry wavelengths were separated through a 565dcxr dichroic mirror (Chroma). Images were acquired with an Andor iXon DU885K EMCCD camera (Oxford Instruments, UK). The camera and monochromator were controlled by FEI Live Acquisition software (Thermo Scientific). The same software, together with ImageJ, were used for image analysis.

Direct excitation of Cherry at 488 nm (Cherry crosstalk; $1.67 \pm 1.3\%$ of Cherry emission excited at 585 nm) and GFP emission in the FRET channel (GFP crosstalk; $1.89 \pm 3.9\%$ of GFP emission in the GFP channel) were determined for our experimental conditions. FRET was calculated with the following equation⁴⁹:

$$netFRET = \frac{I_{FRET} - I_{GFP} \times C_{GFP} - I_{Cherry} \times C_{Cherry}}{I_{GFP}}$$
(1)

where I_{FRET} is the fluorescence intensity in the Cherry channel after excitation at 488 nm, I_{GFP} the intensity in the GFP channel with 488 nm excitation, I_{Cherry} the intensity in the Cherry channel with 585 nm excitation, C_{GFP} the GFP crosstalk percentage in the Cherry channel and C_{Cherry} the Cherry crosstalk percentage at 488 nm excitation in the Cherry channel.

A short, membrane-targeted, tandem construct (RhoCG) expressing tGFP and mCherry followed by the Rho prenylation sequence was used as a FRET positive control.⁵⁰ The level of FRET obtained with RhoCG was taken as the maximum observable under our experimental conditions, and used as a reference (100%) to normalize FRET levels observed with other FRET pairs. In cotransfections, only cells with a mCherry-GFP fluorescence intensity ratio of at least 2:1 were used for analysis.

TIRF-fluorescence recovery after photobleaching microscopy experiments

HEK293 cells were transfected with hTRPA1-tGFP. GFP-labelled TRPA1 channels near the plasma membrane-cover slip interface were visualized at room temperature with total internal fluorescence (TIRF) microscopy.⁵¹ The setup consisted of a Nikon TE2000-U Eclipse microscope (Nikon) equipped with a ×60/numerical aperture 1.49 oil immersion ApoTIRF-objective (Nikon). A 491-nm laser light was produced by a Calypso100 laser emitter (Cobolt,) that was further filtered and power regulated by an acoustic optical tuneable filter (TillPhotonics-AOTF, Thermo Fisher, USA) and delivered to the rear illumination port of the microscope through a Till TIRF condenser (Thermo Fisher). The filter cube contained a Dual 488/561 HC BrightLine laser filter set (Semrock). Images were collected with an Andor iXon DU885K CCD camera (Oxford Instruments). Camera and laser system were controlled by FEI Live Acquisition software (Thermo Fisher), the same software was used for image analysis. The incidence angle of the laser was manually adjusted to obtain an evanescent field at the glasswater interface.

Reagents

Acute application to cells: Carvacrol (98%, 100 mM stock in DMSO, Sigma-Aldrich), allylisothiocyanate (AITC, 95% 100 mM stock in DMSO, Sigma-Aldrich), A967079 (5 mM stock in DMSO, Tocris). Cells were incubated with: S1RA (50 mM stock in DMSO, E-52862, ESTEVE Pharmaceuticals S.A.), BD1063 (50 mM stock in H₂O, ESTEVE Pharmaceuticals S.A.) and control solution (CS, 0.2% DMSO). All substances diluted in bath solution (in mM): 140 NaCl,

3 KCl, 2.4 CaCl $_2,\,1.3$ MgCl $_2,\,10$ HEPES and 10 glucose adjusted to pH 7.4 with NaOH.

For hind paw intradermal injections, 10 mM AITC in ethanol was used. For intraperitoneal injections, a solution of 40 mg/kg S1RA was prepared in sterile saline (0.9% NaCl, B-Braun). Oxaliplatin (Tocris), 5 mM stock in H_2O , was dissolved in 5% glucose and injected at final dose of 6 mg/kg.

Statistical analysis

Values are given as mean \pm standard error (SE). Statistical significance was estimated with the t-test, one-way ANOVA (followed by a post hoc test; Bonferroni or Dunnett) or Fisher's exact test, according with the data and specified in each experiment. A P-value <0.05 was considered statistically significant. Analysis was performed using GraphPad Prism v.5.0. Statistics for each experiment are described within the figure legends.

Data availability

All information necessary to evaluate the reported findings are included in the main text or the Supplementary material. Additional data or further methodological details will be provided by the authors upon request.

Results

Sigma-1R antagonists reduce calcium responses to TRPA1 agonists

To explore the modulatory action of Sigma-1R on TRPA1 channels, cultured HEK293 cells transiently transfected with human TRPA1 (hTRPA1-tGFP) were incubated with CS, 0.2% DMSO) or with different concentrations (25, 50 and 100 µM) of the highly selective Sigma-1R antagonist S1RA.42 Intracellular calcium responses to AITC, an electrophilic TRPA1 agonist, were monitored with the fluorescent calcium indicator Fura-2 (Fig. 1A and B). A 4-h incubation with S1RA caused a dose-dependent reduction in the fraction of responding cells and in the peak amplitude [half-maximum inhibitory concentration (IC₅₀) = $47 \pm 3 \mu$ M] (Fig. 1B–D). At the maximum concentration tested (100 µM) the amplitude reduction was (83%, and the number of cells responding to AITC decreased to 44%, compared to 95% responses in cells incubated in CS: Fig. 1E and F). Incubations, as short as 1 h with S1RA (100 μ M), produced a clear inhibition. Longer incubations (24 h) at lower concentrations (25 µM) also produced marked reductions in the response to AITC (Supplementary Fig. 1).

A similar inhibition in TRPA1 activity was observed after 4 h of incubation with BD1063 (100 µM), a structurally unrelated Sigma-1R antagonist (Fig. 1G). The amplitude was reduced to 36% of control and the percentage of responders was reduced to 45%. The neurosteroid progesterone has been shown to act as an antagonist.^{52,53} endogenous Sigma-1R Incubation of hTRPA1-expressing cells with progesterone also inhibited AITC-evoked responses (Supplementary Fig. 2). We also examined responses to carvacrol (100 µM), a non-electrophilic agonist of TRPA1 found in oregano. After incubating hTRPA1-tGFP-HEK293 cells with S1RA (100 μ M, 4 h), the fraction of responding cells was reduced from 90 to 57% and their mean amplitude was reduced to 26% of control (Fig. 1H).

In summary, three different Sigma-1R antagonists markedly reduced human TRPA1 responses to agonists with different chemical properties and activation mechanisms, showing that



Figure 1 Sigma-1R antagonists inhibit intracellular calcium responses in TRPA1 expressing cells. (A) Representative intracellular calcium responses monitored with Fura-2 in response to AITC (50 μ M) after incubation in CS, 4 h). Green trace is the average of 32 cells (grey traces) expressing hTRPA1-tGFP. (B) Same as in A but after incubation with S1RA (100 μ M, 4 h). The blue trace is the average of 30 cells (grey traces) expressing hTRPA1-tGFP. (C) Dose-response curve of S1RA effects. ***P<0.001 (one-way ANOVA in combination with Dunnett's post hoc test) CS n=37, S1RA 25 μ M n=68, 50 μ M n=90, 100 μ M n=64. (D) Percentage of responding cells to AITC after pre-incubation (4 h) with S1RA at different concentrations. ***P<0.001 (Fisher's exact test) versus CS. (E) Time-dependent inhibition of TRPA1 responses to AITC (50 μ M) by pre-incubation with S1RA (100 μ M, 4 h). ***P<0.001 (ANOVA in combination with Dunnett's post hoc test). CS n=37, 1 h n=21, 3 h n=77, 4 h n=64. (F) Effect of incubation time with n=255 versus CS n=255 ***P<0.001 (upaired t-test). (H) Graph summarizing the effect of pre-incubation with S1RA (100 μ M, 4 h) to responses when the cells were stimulated with carvacrol (100 μ M, n=21) versus CS n=45. ***P<0.001 (unpaired t-test).

the inhibitory actions of Sigma-1R antagonists on TRPA1 activity are independent of their structural properties or the TRPA1 mode of activation.

TRPA1 inhibition by Sigma-1R antagonists depends on the presence of Sigma-1R

Sigma-1R is broadly expressed in most tissues, including human kidney cells.³⁰ To confirm that Sigma-1R is the target of S1RA effect, we transfected hTRPA1-tGFP transiently in Sigma-1R KO HEK293 cells. As expected, these cells did not express Sigma-1R protein (Fig. 2C). Cells were pre-incubated for 4 h with S1RA (100 μ M) or CS and the calcium response to AITC (50 μ M) was tested. In cells lacking Sigma-1R, incubation with S1RA had no effect on the amplitude of AITC-evoked responses (Fig. 2A, top). In contrast, the transient expression of hSigma-1R-mCherry in silenced cells restored the strong inhibitory action of S1RA incubation on TRPA1 activity (Fig. 2A, bottom). In these cells, the peak amplitude of AITC-evoked responses was reduced 32% by S1RA compared to hSigma-1R-mCherry transfected cells incubated with CS (Fig. 2B).

These results indicate that Sigma-1R expression is necessary and sufficient for TRPA1 inhibition after S1RA incubation. They also suggest that these inhibitory effects are not exerted directly on TRPA1 channels.

The reduction of TRPA1 currents by S1RA depends on Sigma-1R

To investigate the influence of Sigma-1R on TRPA1 channels directly, whole-cell voltage-clamp recordings were performed from HEK293 cells stably expressing hTRPA1-tGFP. Cells were preincubated 4 h with CS or with different concentrations of S1RA. AITC-evoked inward currents were inhibited in a dose-dependent manner (IC₅₀ = $37 \pm 11 \mu$ M) when cells were pre-incubated with S1RA (Fig. 3A and B). Incubation with 100 µM S1RA, the highest dose tested, reduced AITC-evoked currents by 81% at -60 mV, compared to cells incubated in CS (Fig. 3C). AITC-activated inward currents at -60 mV (Fig. 3A) had a mean density of $52 \pm 11 \text{ pA/pF}$ (n = 10) in CS compared to 10 ± 3 pA/pF (n = 10) after S1RA 100 μ M. Of note, S1RA pre-incubation also modulated the kinetics of AITC-evoked inward currents (Fig. 3A). The time to peak increased to 65 ± 10 s (n = 18) after pre-incubation with 100 μ M S1RA, compared to a time to peak of 25 ± 2 s (n = 20) in cells incubated in CS, suggesting that S1RA slowed the macroscopic activation rate.



Figure 2 TRPA1 inhibition by Sigma-1R antagonists depends on Sigma-1R expression. (A) Top: Representative intracellular calcium responses to AITC (50 μ M) in Sigma-1R silenced HEK293 cells (S1R-KO) transfected with hTRPA1-tGFP. Cells were incubated with CS (4 h) or in S1RA (100 μ M, 4 h). Green and blue traces represent the averages of individual cells in grey. Bottom: The same experiment as in **A** but in hTRPA1-tGFP cells cotransfected with hSigma-1R-mCherry. (B) Summary histograms of mean \pm SE calcium responses to AITC in S1R-KO cells + hTRPA1-tGFP in control (green, n = 312), S1R-KO cells + hTRPA1-tGFP in S1RA (100 μ M, 4 h) (blue, n = 276), S1R-KO cells + hTRPA1-tGFP + hSigma-1R-mCherry in control (magenta, n = 118) and S1R-KO cells + hTRPA1-tGFP + hSigma-1R-mCherry in S1RA (100 μ M, 4 h) (pink, n = 144). ***P < 0.001 (ANOVA in combination with Bonferroni's post hoc test). (C) Western blot of HEK293 cells transiently transfected with hTRPA1-tGFP, WT HEK293 cells and S1R-KO HEK293 cells, demonstrating the expression of endogenous Sigma-1R in WT cells and the absence in S1R-KO cells.

Next, to demonstrate the specificity of S1RA actions, we examined the effect of S1RA pre-incubation on TRPA1 currents after silencing Sigma-1R (i.e. S1R-KO). In this case, incubation with S1RA had no effect on the amplitude of AITC-evoked currents (Fig. 3D and E).

These results confirm that Sigma-1R expression is necessary for the reduction in TRPA1 channel activity observed after S1RA pre-incubation.

S1RA impairs the formation of TRPA1–Sigma-1R complexes

Sigma-1R is a molecular chaperone known to interact directly with various ion channels, affecting their gating and their expression at the plasma membrane, reviewed by.⁵⁴ To probe the molecular interaction between Sigma-1R and TRPA1, we performed co-immunoprecipitation experiments, using HEK293 transiently transfected with hTRPA1-tGFP and mSigma-1R-EYFP fusion proteins. Cell extracts containing mSigma-1R-EYFP were purified with GFP beads and, as shown in Fig. 4A, TRPA1 was readily pulled down only in the presence of Sigma-1R (Fig. 4A, lane 1).

FRET experiments were performed to define the possible formation of a complex between Sigma-1R and TRPA1, and to study the effect of S1RA on this interaction. HEK293 cells cotransfected with hTRPA1-tGFP and hSigma-1R-mCherry were excited at 488 nm to measure sensitized emission FRET between both fluorophores (see the 'Materials and methods' section). As shown in Fig. 4B, cotransfection of both proteins produced a level of FRET that was $35.3 \pm 1.1\%$ of RhoCG FRET (n = 153) (the FRET positive control, see the 'Materials and methods' section). This value was significantly higher than that observed when hTRPA1-tGFP was cotransfected with free soluble mCherry and no specific interaction was expected (10.4 \pm 3.6% of RhoCG FRET, n = 199). These results indicate that the physical distance between TRPA1 and Sigma-1R is <10 nm, sufficiently close for making a protein–protein complex between them. Interestingly, when cotransfected cells were pre-incubated with S1RA (100 μ M, 4 h), the levels of FRET decreased significantly (28.6 \pm 1.1% of RhoCG FRET, n = 221), compared to the level obtained in cells treated with CS (P < 0.01, unpaired t-test), suggesting a reduced interaction.

This result suggests that the antagonistic effect of S1RA on TRPA1 activity could be due to an impairment in the assembly of TRPA1–Sigma-1R complexes or an alteration in the complex conformation.

S1RA incubation diminishes TRPA1 expression at the plasma membrane

So far, the results indicate that Sigma-1R interacts with TRPA1 channels and this interaction is affected by S1RA, leading to reduced TRPA1 activity. Next, we asked whether trafficking of TRPA1 channels to the plasma membrane was altered by S1RA incubation. To this end, we performed fluorescence recovery after photobleaching experiments in HEK293 cells transiently transfected with hTRPA1-tGFP, using TIRF excitation to restrict the photobleaching and recovery visualization to a thin layer of ~200 nm containing the plasma membrane. Sustained, high-power TIRF illumination for 120 s photobleached ~80% of fluorescence excited near the plasma membrane. Thereafter, fluorescence recovery due to hTRPA1-tGFP channel trafficking from non-bleached regions was monitored for 10 min. hTRPA1-tGFP was observed in the entire footprint of the cell as a dense mesh of punctate structures (Fig. 4C and Supplementary Fig. 3). After photobleaching, recovery of hTRPA1-tGFP takes place homogenously



Figure 3 Sigma-1R expression is necessary for the reduction of TRPA1 whole-cell currents by S1RA. (A) Representative time course of whole-cell currents, from the HEK293 hTRPA1-tGFP stable cell line, in response to the indicated repeated voltage ramp protocol (red) and AITC (50 μ M), HP = -60 mV, in CS and after S1RA pre-incubation (100 μ M, 4 h). Note the slow development of current and the reduced peak amplitude after S1RA incubation. (B) Average IV relationships from HEK293-hTRPA1-tGFP cells using the protocol shown in **A** recorded in control (*n* = 10) and S1RA at different concentrations: 10 μ M (*n* = 10), 30 μ M (*n* = 10) and 100 μ M (*n* = 10). Average of maximal response to AITC in each trace. (C) Dose-response curve of S1RA effects on HEK293 hTRPA1-tGFP cells in response to AITC (50 μ M). Normalized current density at -60 mV with respect to CS. ***P < 0.001 (ANOVA in combination with Dunnett's post hoc test). (D) Average IV relationships from WT HEK293 cells transiently transfected with hTRPA1-tGFP in control (*n* = 15), S1RA pre-incubation (100 μ M, 4 h, *n* = 12) and in Sigma-1R silenced cells transiently transfected with hTRPA1-tGFP (S1R-KO) recorded after pre-incubation in CS for 4 h (*n* = 12) or S1RA (100 μ M, 4 h, *n* = 18). Average of maximal response to AITC in each case. (E) Summary of individual and mean \pm SE current density at -60 and +60 mV for the different experimental conditions shown in D. **P < 0.01 (ANOVA in combination with Bonferroni's post hoc test). Results from transient transfections were obtained in seven independent experiments.

across the entire surface of the cell in contact with the coverslip, suggesting trafficking of the channel from the cytosol towards the plasma membrane and not by membrane lateral diffusion (Supplementary Fig. 3).⁵¹ As shown in Fig. 4C, the observed final level of fluorescence recovery after photobleaching after 600 s was significantly lower (blue trace, $16.6 \pm 1.5\%$ n = 22) in cells pretreated with S1RA (100 µM, 4 h) compared to cells incubated in CS (red trace, $20.7 \pm 1.2\%$ n = 31). Comparing the time constants of fluorescence recovery, no differences were found in recovery kinetics (data not shown). These results suggest that Sigma-1R facilitates TRPA1 trafficking to the plasma membrane, a process impaired by S1RA.

Additional evidence for the reduced expression of TRPA1 channels at the plasma membrane after S1RA incubation (100 μ M, 4 h) came from dose-response curves to the agonist AITC. As shown in Fig. 4D, S1RA pre-incubation produced a rightward shift in the dose-response curve [half-maximum effective concentration (EC₅₀) 3.61 \pm 0.03 μ M in control versus EC₅₀ 14.12 \pm 0.11 μ M after S1RA] and a significant reduction (~20%) in the maximal calcium response.

Finally, to establish more directly an effect of S1RA on plasma membrane expression of TRPA1, we used cell surface biotinylation techniques (see the 'Materials and methods' section). We evaluated TRPA1 plasma membrane and total protein levels in CS and after S1RA incubation (25 μ M–24 h) (Fig. 4E). Notably, total TRPA1 expression level was unaffected by S1RA treatment (Fig. 4E). In contrast, treatment produced a significant reduction in the avidin-bound fraction (i.e. biotinylated) of TRPA1. The level of TRPA1 at the surface, normalized to TRPA1 in the total protein extract, was reduced to about half in cells pretreated with S1RA (Fig. 4F).

Collectively, these results indicate that the observed decrease in TRPA1 channel activity after S1RA incubation is caused by a reduction in TRPA1 membrane expression without alteration in total TRPA1 levels.

S1RA incubation reduces endogenous TRPA1 activity in sensory neurons

To investigate the effects of Sigma-1R antagonists on TRPA1-mediated responses in nociceptors, DRGs from adult C57BL/ 6J mice were cultured and incubated with S1RA (100 μ M, 4 h) or CS. Representative traces of neurons responding to AITC in control and pre-incubated with S1RA are shown in Supplementary Fig. 4A. The amplitude of intracellular calcium responses to AITC (50 μ M) were significantly reduced from 0.46 \pm 0.01 in CS (n = 314) to 0.25 \pm 0.02 after S1RA treatment (n = 195). Treatment also reduced the percentage of



Figure 4 S1RA impairs the formation of TRPA1-Sigma-1R complexes and reduces TRPA1 expression at the plasma membrane. (A) Co-immunoprecipitation of TRPA1 and Sigma-1R-EYFP in cotransfected HEK293 cells. Input is one-tenth of the total protein used in each immunoprecipitation. TRPA1 was pulled down by Sigma-1R-EYFP (lane 1) using GFP-Trap (Chromotek), while it was not when Sigma-1R-EYFP was not present (lane 2). The figure is representative of four independent protein extractions. GFP-Trap agarose beads do not recognize tGFP. (B) Detection of net sensitized emission (FRET) levels between hTRPA1-GFP and hSigma-1R-mCherry in transfected HEK293 cells. Top: Representative image of the fluorescence signal of hTRPA1-tGFP+Sigma-1R-mCherry as well as the signal in the FRET channel (scale bar = 20 µM). Bottom: Summary histogram of FRET levels (as a percentage of RhoCG FRET) of TRPA1-tGFP+mCherry (negative control, green bar) and TRPA1-tGFP+Sigma1-mCherry in control conditions (black bar) and after S1RA incubation (100 μ M, 4 h) (blue bar). ^{##}P < 0.001 (ANOVA in combination with Bonferroni's post hoc test) test TRPA1-tGFP+soluble mCherry versus all groups, **P < 0.01 (unpaired t-test), S1RA untreated versus treated hTRPA1-tGFP+Sigma-1R-mCherry transfected cells. (C) Left: The time course of hTRPA1-tGFP TIRF fluorescence recovery after TIRF photobleaching, in control treated cells (red trace) and cells incubated with S1RA (100 µM, 4 h, blue trace). *P < 0.05 (unpaired t-test). Error bars represent SE. Representative TIRF images of HEK293 transfected with hTRPA1-GFP before (top), immediately after photobleaching (middle) and after 10 min recovery (bottom), scale bar = 10 µM. (D) Dose-response curves of AITC-evoked calcium responses in stable HEK293-hTRPA1-tGFP cells in CS (red) and pre-incubated with S1RA (100 µM, 4 h, blue). Values have been normalized to 500 µM AITC in CS. The lines represent fits to Hill functions $(Y = Y_{max}/(1 + (EC_{50}/X)^{nH}))$ with parameters $EC_{50} = 14.12 \pm 0.11 \mu$ M and $3.60 \pm 0.03 \mu$ M, $Y_{max} = 1.009$ \pm 0.002 and 0.797 \pm 0.002 and nH = 1.03 \pm 0.01 an 1.29 \pm 0.01 in control and S1RA (100 μ M, 4 h), respectively. (E) Western blot of total protein (TP) extracts from HEK293-hTRPA1-tGFP stable line incubated in CS for 24 h) or S1RA (25 µM, 24 h). A non-specific band at ~70 kDa is exposed with anti-GAPDH (1:5000 G9545, Sigma-Aldrich). Immunodetection values in S1RA and CS, normalized with respect to GAPDH, are similar (n = 5 independent extractions, P > 0.05, unpaired t-test). (F) Representative western blot from biotinylation assays of hTRPA1- tGFP cells. Total expression of TRPA1 was assessed by directly immunoblotting 15 µg of each cell lysate. TRPA1 cell surface (Plasma Membrane protein, PM) expression was calculated by expressing the intensities of the TRPA1 bands in the avidin-bound fraction as a fraction of those in the corresponding cell lysates (n = 3, *P < 0.05, unpaired t-test). GAPDH was used as a negative control blot for the presence of a soluble protein in the avidin bound fraction (ABF).

AITC responders from 47 to 32% (Supplementary Fig. 4A and B). In TRPA1-expressing DRG neurons, obtained from Trpa1 Cre-tdTomato mice and identified by their red fluorescence, incubation with S1RA (100 μ M, 4 h) produced a similar reduction in AITC-evoked responses (Supplementary Fig. 4C and D). Similar to our findings in TRPA1-expressing HEK293 cells, longer incubations (24 h) of DRG neurons with S1RA or BD1063 reduced the concentration of antagonists necessary to observe a significant reduction in AITC-evoked responses (Supplementary Fig. 5). In contrast to the effects of S1RA incubation, the acute application of a high concentration of the drug had no effect on the time course of AITC-evoked response in mouse DRG neurons (Supplementary Fig. 6).

The effect of S1RA incubation on the excitability of TRPA1(+) DRG neurons was examined with cell-attached recordings, a condition where the intracellular milieu is preserved. In CS, all tdTomato(+) neurons tested (8/8) responded to 100 μ M AITC, reaching a maximum average firing of 1.4 Hz (Fig. 5A and B). In contrast, after S1RA treatment about a half (5/11) failed to respond to AITC, or

decreased their firing rate: their average frequency did not exceed 0.6 Hz.

Activation of nociceptor endings by TRPA1 agonists produce depolarizing generator potentials and spike firing.⁵⁵ We simulated these conditions in current-clamp recordings obtained from cultured TRPA1(+) DRG neurons (Fig. 5C). Only minor differences were found in the electrophysiological properties of TRPA1(+) neurons after S1RA incubation when compared to CS (Supplementary Table 2). In contrast, the maximum depolarization evoked by AITC was strongly reduced after S1RA incubation, $5 \pm 1 \text{ mV}$ (n = 9) versus $12 \pm 1 \text{ mV}$ (n = 11) after incubation in CS (Fig. 5D). In CS, all (11/11) TRPA1(+) neurons fired AP after AITC application. In contrast, this number was reduced to 56% (5/9) after S1RA incubation (P < 0.05, Fisher's exact test). Moreover, the number of evoked action potentials were also reduced by S1RA incubation and their frequency during AITC application did not exceed 0.76 ± 0.10 Hz (average of n = 5) in the presence of S1RA while in control it reached 2.0 ± 0.3 Hz (average of n = 11) (Fig. 5D). Finally, electrophysiological



Figure 5 S1RA reduces TRPA1 activity in mouse DRG nociceptors. (A) TRPA1(+) neurons in cultured DRGs from Trpa1 Cre-tdTomato mice were identified by their red fluorescence. AITC (100 μ M) evoked action currents in 100% of neurons in CS (n = 8) and in 55% after S1RA incubation (6/11) (*P < 0.05, Fisher's exact test). Firing frequency during the 2 min of AITC application. ***P < 0.001 (unpaired t-test). (B) Representative cell-attached recordings of TRPA1-expressing DRG neurons (n = 5 mice). Responses to AITC (100 μ M) and high KCl (30 mM) after incubation with CS, 4 h) or S1RA (100 μ M, 4 h). Note the robust firing in CS. After S1RA incubation (100 μ M, 4 h), six neurons showed a weak response (S1R1-R, responder) while five did not fire action potentials (S1RA-NR, non-responder). Note that in neurons in which no response to AITC was observed, an impulse discharge could be evoked with 30 mM KCl. (C) Representative whole-cell current-clamp recordings of TRPA1(+) DRG neurons (n = 8 mice) showing the response to AITC (100 μ M) and KCl (30mM) after incubation with CS-4 h or S1RA (100 μ M, 4 h). After S1RA, some neurons (n = 8 mice) showing the response to AITC (100 μ M) and KCl (30mM) after incubation with CS-4 h or S1RA (100 μ M, 4 h). After S1RA, some neurons (n = 8 mice) showing the response to AITC (100 μ M) and KCl (30mM) after incubation with CS-4 h or S1RA (100 μ M, 4 h). After S1RA, some neurons fined AP (S1RA-R) while others showed a subthreshold depolarization but no AP firing (S1RA-NR). Both fired AP in response to high KCl. (D) Histograms summarizing the amplitude of AITC-evoked depolarization, number of action potentials and mean firing frequency of neurons: in CS (n = 11 out of 11) and after S1RA incubation (n = 5 of 9) responded to AITC. Mean ± SE **P < 0.001, *P < 0.05 (unpaired t-test). (E) Left: Representative whole-cell voltage-clamp recordings (n = 2 mice) of TRPA1(+) neurons (V_{hold} = -73 mV). AITC-evoked inward current in CS for 4 h incubated neurons (-9.6 ± 1.4

recordings in whole-cell voltage-clamp mode showed a large reduction in the amplitude of AITC-evoked inward currents after S1RA incubation (Fig. 5E).

These results confirm that S1RA causes a substantial decrease in the activation of TRPA1-expressing nociceptors to a TRPA1 agonist, without significant alteration in their intrinsic excitability.

Systemic S1RA injections reduce pain-related behaviours to intradermal TRPA1 agonists

Considering that Sigma-1R antagonists mitigate nociceptive responses in different experimental pain models, ^{41,42,56} we examined the possible role of TRPA1 as an antinociceptive drug target of Sigma-1R. To this end, we quantified responses of WT and TRPA1 KO mice to intradermal injections of AITC (10 µl at 10 mM) in their hind paws.⁴⁶ One group of animals received vehicle (saline i.p.) and the other S1RA (40 mg/kg i.p.), 24-h before blind evaluation of painrelated behaviour (i.e. duration of paw licking after AITC injection). The results show that systemic S1RA injection produces a substantial reduction in pain-related responses to intradermal AITC in WT mice (Supplementary Fig. 7B). Responses to AITC were strongly diminished in TRPA1 KO mice compared to WT, and no further reduction was observed in these animals when pretreated with S1RA (Supplementary Fig. 7B). These findings demonstrate that the modulation of pain-related behaviours produced by S1RA after AITC application was dependent on TRPA1 activation.

TRPA1 channels participate in painful responses to oxaliplatin

Systemic oxaliplatin administration causes a painful peripheral neuropathy in patients and rodent models, characterized by mechanical and cold hypersensitivity in their extremities.^{1,8} To induce the neuropathy, mice received three injections of oxaliplatin (6 mg/kg i.p.) on alternate days (Fig. 6A), while the control group received its vehicle (glucose 5% in dH₂O). Mechanical and cold sensitivity were monitored in mice before and after oxaliplatin treatment (see the 'Materials and methods' section).

Systemic oxaliplatin treatment led to a marked hypersensitivity to mild (acetone application) (Fig. 6B) and noxious (10°C) cold stimuli (Fig. 6D).^{24,57} Notably, this thermal hypersensitivity was abolished in TRPA1 KO mice (Fig. 6C). Interestingly, baseline responses to mild cold were not altered in TRPA1 KO mice compared with WT, suggesting that other mechanisms, e.g. TRPM8, mediate innocuous cold sensitivity (Fig. 6B and C). In contrast, TRPA1 KO mice showed longer latencies in the noxious cold test, suggesting a role of TRPA1 in physiological noxious cold responses (Fig. 6D).^{44,58}

Mechanical sensitivity was evaluated with calibrated von Frey hairs at baseline and 3 days after the end of oxaliplatin or vehicle treatment. We noticed that vehicle injections tended to reduce mechanical thresholds (Fig. 6E). This should be noted because it introduces a confounding factor when evaluating the effects of oxaliplatin. This reduction may reflect a higher stress level caused by the repeated injections required in this protocol. Oxaliplatin caused a marked mechanical hypersensitivity in WT mice (n=11): their mean threshold was reduced from 0.64 ± 0.11 g at baseline to 0.10 ± 0.04 g after oxaliplatin (Fig. 6E). In contrast, the injection of vehicle resulted in a moderate, not significant, decrease in mechanical threshold compared to baseline (in average 0.98 ± 0.16 g versus 0.67 ± 0.11 g, n=9). The effects of oxaliplatin on mechanical sensitivity in TRPA1 KO mice were clearly attenuated when compared to WT mice. As shown in Fig. 6F, TRPA1 KO animals

treated with oxaliplatin showed a milder mechanical hypersensitivity when compared to WT animals, and this hypersensitivity was nearly identical to the one observed after vehicle injection, suggesting that it is mainly caused by the handling or repeated injections.

These results indicate that activation of TRPA1-expressing nociceptors plays a central role in the cold and mechanical hypersensitivity caused by systemic oxaliplatin treatment.

Systemic S1RA treatment prevents mechanical and cold hypersensitivity in oxaliplatin-treated mice

Since administration of oxaliplatin and other chemotherapeutics follows a planned regime,⁵⁹ we asked whether previous administration of a Sigma-1R antagonist could prevent or attenuate the painful symptoms produced by systemic oxaliplatin. The treatment protocol consisted of daily dosing of systemic S1RA (40 mg/ kg i.p., see the 'Materials and methods' section), starting 3 days before the first oxaliplatin injection (6 mg/kg), which was injected in three alternate days for 3 days (Fig. 6G). In the control group, the animals received vehicle (saline, i.p.) instead of S1RA and identical oxaliplatin treatment. Behavioural evaluation was performed before the start (baseline, BS) and after the last oxaliplatin injection, the same day or 1-5 days after the last S1RA injection (Fig. 6G), by a researcher blind to treatment. Consistent with our previous results, 7 days after the start of oxaliplatin treatment, control mice (i.e. treated with saline) developed a marked hypersensitivity to mild cold (i.e. acetone) (Fig. 6H). This hypersensitivity was greatly reduced in animals treated with S1RA (Fig. 6H). Similarly, responses to noxious cold were also substantially reduced in animals receiving S1RA instead of saline (Fig. 6I). Notably, this antiallodynic effect persisted even 3 days after the last S1RA injection (Fig. 6G), well passed the plasma clearance of the drug in mice.^{42,45} The effects of S1RA on attenuating mechanical allodynia following oxaliplatin treatment were also substantial (Fig. 6J). In this case, saline-treated animals showed a mechanical threshold of 0.32 ± 0.1 g (n = 12), compared to 0.69 ± 0.12 g (n = 12) after S1RA treatment (P < 0.05, unpaired t-test).

Collectively, these findings suggest that TRPA1 channels play a major role in the development of cold and mechanical hypersensitivity produced by oxaliplatin and these painful symptoms can be attenuated by a Sigma-1R antagonist applied systemically.

S1RA treatment normalizes TRPA1 responses in cultured DRG neurons from oxaliplatin-treated mice

Sigma-1R shows a very broad distribution throughout the nervous system.³¹ The cellular locus of S1RA antiallodynic effects is still unclear, with evidence for central and peripheral sites of action.⁴² Having established the antiallodynic effect that S1RA plays in oxaliplatin-induced neuropathic symptoms, and the dependence of these symptoms on TRPA1, we tested the modulatory role of Sigma-1R on sensitized peripheral neurons.

First, we investigated responses to AITC in short-term cultured DRG neurons prepared from oxaliplatin or vehicle-treated WT mice, and evaluated the effects of incubation with S1RA (100 μ M, 4 h) or CS on these responses. We selected animals that showed a marked behavioural phenotype (i.e. mechanical and cold allodynia) to oxaliplatin treatment. Cold hypersensitivity was confirmed by the unilateral cold plate test (10°C) the same day of neuronal isolation. Two hours after harvesting and plating, neurons were incubated 4 h with S1RA or CS and loaded with Fura-2, during the last



Figure 6 Systemic S1RA administration reduces TRPA1-dependent cold and mechanical hypersensitivity in oxaliplatin-treated mice. (A) Time line of experimental protocol. Oxaliplatin was injected three times on alternate days. Oxaliplatin effects to mild cold (acetone) or mechanical stimulation (von Frey) were evaluated 3 days before and 7 days after first injection. Noxious cold was evaluated 8-12 days after the first oxaliplatin injection. (B) Responses to acetone in WT mice before (baseline, BS) and after vehicle (green, n = 5), and before and after oxaliplatin (orange, n = 5) ***P< 0.001, n.s. P > 0.05 (unpaired t-test). Paired t-test when responses of the same mice were compared (lines connecting symbols). (C) Cold (acetone) sensitivity in TRPA1 KO mice before (BS) and after vehicle (purple-green, n=5), and before (BS) and after oxaliplatin (purple-orange, n=7) injection. n.s. P > 0.05 (unpaired t-test). Paired t-test when the same mice were compared. Oxaliplatin did not induce cold allodynia in TRPA1 KO mice. (D) Evaluation of noxious cold sensitivity by unilateral cold plate test after vehicle (WT green, n = 7, TRPA1 KO purple-green, n = 5) or oxaliplatin injection (WT orange n = 11, TRPA1 KO purple-orange n=7) ***P < 0.001, n.s. P > 0.05 (one-way ANOVA in combination with Bonferroni's post hoc test). Oxaliplatin-induced cold hyperalgesia in WT but not in TRPA1 KO mice. (E) Mechanical withdrawal thresholds, in WT mice before and after vehicle injection (green, n = 9), before and after oxaliplatin injection (orange, n = 11). ***P < 0.001, n.s. P > 0.05 (unpaired t-test). Paired t-test when the same mice were compared. Oxaliplatin induced a clear mechanical allodynia. (F) Mechanical withdrawal thresholds in TRPA1 KO mice before and after vehicle injection (purple-green, n = 5), and before and after oxaliplatin injection (purple-orange, n = 7), *P < 0.05, n.s. P > 0.05 (unpaired t-test). Paired t-test when the same mice were compared. Oxaliplatin induces less mechanical allodynia in TRPA1 KO than in WT mice. (G) Timeline of experimental protocol to evaluate preventive effects of S1RA. Mice were evaluated 3 days before and 7-12 after first oxaliplatin (OXA) injection. (H) Mild cold (acetone) sensitivity before and after or oxaliplatin (OXA) injections. One group received saline combined with OXA (orange-pink, n = 12), and the other S1RA combined with OXA (orange-blue, n=12) **P < 0.01, *P < 0.05 (paired t-test). (I) Unilateral cold plate test in saline + OXA injected mice (orange-pink, n=12), and S1RA + OXA injected mice (orange-blue, n = 12). *P < 0.05 (unpaired t-test). (J) Mechanical sensitivity before (black) and after saline + OXA injections (orange-pink, n = 12) or S1RA + OXA (orange-blue, n = 12) ***P < 0.001, *P < 0.05 (paired t-test).

45 min of incubation (see the 'Materials and methods' section). In sensory neurons obtained from oxaliplatin-treated mice, the average amplitude of AITC-evoked calcium responses was larger (0.55 ± 0.04 , n = 118) than in neurons obtained from vehicle-injected mice (0.38 ± 0.25 , n = 149) (Fig. 7A and B). Furthermore, incubation of neurons obtained from oxaliplatin-injected mice with S1RA (100μ M, 4 h), restored AITC-evoked responses to values similar to those obtained from mice treated with vehicle (0.35 ± 0.02 , n = 172) (Fig. 7A and B). These results show that acute S1RA treatment restores sensitized TRPA1 responses in DRG neurons obtained from oxaliplatin-treated animals to normal values.

In vivo S1RA treatment prevents TRPA1 sensitization in cultured DRG neurons from oxaliplatin-treated mice

Finally, to evaluate the preventive effect of systemic (i.p.) S1RA on oxaliplatin-mediated sensitization of TRPA1, we used the protocol shown in Fig. 7C. The effects of *in vivo* S1RA administration were compared to the injection of saline. Response to TRPA1 agonists were examined in short-term cultures 1–3 days after the last S1RA injection. Consistent with the behavioural results shown previously, the amplitude of AITC-mediated responses in cultured DRG neurons was reduced in animals treated *in vivo* with S1RA (Fig. 7C and D). These effects persisted for 72 h after systemic S1RA treatment.

These results suggest that the preventive effect of systemic S1RA in the development of the oxaliplatin-induced neuropathy involves the modulation of TRPA1 channels in nociceptors.

Discussion

CIPN is a frequent, insidious side effect of anticancer therapies. Without a full mechanistic understanding of its causes, treatments have been generally unsuccessful.^{2,5} The salient contribution of this study is the demonstration that TRPA1 is downmodulated by antagonism of Sigma-1R, a multifunctional chaperone, and this inhibition reduces the painful side effects of oxaliplatin in an experimental model of CIPN neuropathy. We established that antagonism of Sigma-1R impairs the formation of a molecular complex between TRPA1 and Sigma-1R, and



Figure 7 In vivo and in vitro S1RA treatment normalizes TRPA1 responses in DRG neurons from oxaliplatin-treated mice. (A) Oxaliplatin (OXA) was injected three times on alternate days. DRG neurons were cultured and studied by Ca²⁺ imaging at Days 8–12, shortly after mice were evaluated by unilateral cold plate test. Calcium responses in DRG neurons from mice that received vehicle (*left*) and OXA injections (*middle*). *Right*: Responses from mice that received of the condition are shown in grey. Coloured traces represent averages of the respective individual traces. (B) Histograms summarizing the sensitization of the AITC response by OXA and the inhibition by SIRA. Vehicle/CS n = 149, OXA/CS n = 118, OXA/SIRA n = 172, ***P < 0.001, n.s. P > 0.05 (one-way ANOVA in combination with Bonferroni's post hoc test). (C) Mice received saline via SIRA injections starting 3 days before the first OXA injection until Day 7. DRG neurons were cultured and studied by Ca²⁺ imaging at Days 8–10, right after mice were evaluated by unilateral cold plate test. Representative traces of calcium elevations evoked by AITC (50 µM) and SIRA + OXA (*right*, average in blue). (D) Histograms summarizing the preventive effect of SIRA in the sensitization of AITC responses by OXA. Saline/ OXA (n = 797), SIRA/OXA (n = 1208), ***P < 0.001 (unpaired t-test).

the trafficking of functional TRPA1 channels to the plasma membrane, leading to a reduction in the excitability of nociceptor neurons sensitized by oxaliplatin.

Role of TRPA1 in CIPN

Multiple mechanisms are thought to be involved in CIPN. In particular, dysregulation of different ion channels, including sodium, potassium and TRP channels have been implicated in the painful symptoms^{16–19,60,61} Our behavioural studies in TRPA1 KO mice defined a key role of this ion channel in the development of mechanical and cold hypersensitivity in oxaliplatin CIPN. Our findings are in line with previous pharmacological and genetic studies in mice and rats.^{22,24,62} Recent *in vivo* imaging studies of mouse DRG neurons found that oxaliplatin treatment unmasked a population of silent peptidergic nociceptors with altered excitability to cold and noxious pinch.²⁰ The molecular identity of the cold sensor in this subpopulation of sensory neurons was not established in the study and further work is needed.

A recurrent theme in various postulated mechanisms for CIPN is the alteration in the redox-status of peripheral sensory neurons, linked to excessive generation of reactive oxygen species by damaged mitochondria^{63–65}. This is very relevant to our findings because TRPA1 is activated by cellular stress, including reactive oxygen species (reviewed by Sakaguchi and Mori⁶⁶). In particularoxaliplatin activates TRPA1 by a glutathione-sensitive mechanism,^{21,23} and cytosol acidification of DRG sensory neurons.⁶⁷ Other *in vitro* studies showed that oxaliplatin can enhance cold sensitivity of human TRPA1 by a mechanism implicating inhibition of prolyl hydroxylase and oxidation of intracellular cysteines.^{25,68}

Current clinical guidelines recommend the use of duloxetine in the treatment of CIPN.⁶ This antidepressant drug showed modest analgesic efficacy in a randomized, double-blind, placebocontrolled trial for CIPN.⁶⁹ It is worth mentioning that a recent study found that duloxetine inhibits TRPA1 activity, and this inhibition had neuroprotective effects on delayed neuropathy caused by exposure to organophosphates, reactive compounds present in many insecticides, herbicides and nerve agents.⁷⁰ Thus, a deeper understanding of the mechanisms involved in the beneficial effects of duloxetine in CIPN, including the possible role of TRPA1 channels, may open new avenues for treatment.

Sigma-1R is necessary for S1RA effects on TRPA1

Using a KO cell line for Sigma-1R, we demonstrated that the receptor is essential for the inhibitory effect of S1RA on TRPA1. Moreover, reintroduction of Sigma-1R rescued the effect. In addition, acute application of S1RA did not reduce AITC-evoked responses. Altogether, this indicates that the actions of S1RA are not mediated by direct inhibition or block of the channel by the antagonist. This demonstration is important because some previous studies found that Sigma-1R ligands can inhibit other TRP channels (e.g. TRPC5 and TRPM3) by mechanisms independent of Sigma-1R.⁷¹ These findings also suggest that the actions of S1RA are not mediated by Sigma-2R. Despite their names, these two proteins show no sequence homology and derive from unrelated genes.⁷² Consistent with this view, S1RA is highly selective for Sigma-1R over Sigma-2R.42 The inhibitory effects of S1RA on TRPA1 required much higher concentrations than the estimates suggested from binding affinities of S1RA to Sigma-1R receptors.⁴² This discordance between affinities obtained in radioligand binding assays and IC₅₀s in functional studies has been noted previously for S1RA and other

potent Sigma-1R antagonists like BD1063,⁷³ that normally require micromolar concentrations to be effective.^{37,74,75} Multiple reasons could contribute to this discrepancy, including the fact that functional studies are performed on cells with intact membranes, and the complex, multistep process, taking place between drug–receptor interaction and biological effect. The crystal structure of the human Sigma-1 receptor revealed that the ligand-binding cavity is very hydrophobic and completely occluded from solvent.⁷⁶ These structural constraints account for the very slow ligand-binding kinetics characteristic of Sigma-1 receptors and help explain the discrepancies noted previously.

In calcium imaging and electrophysiological recordings, we established that Sigma-1R reduces TRPA1 activity in nociceptive neurons. In a recent study, Ortiz-Rentería and collaborators described the inhibition of TRPV1, another nociceptive channel, by Sigma-1R antagonists.³⁷ In addition, the authors found a reduced inhibition in cells treated with siRNA, suggestive of a role of Sigma-1R in the inhibitory action. Since TRPV1 and TRPA1 colocalize in a fraction of nociceptors,²⁶ the inhibitory effects of Sigma-1R antagonists may reflect shared signalling pathways downstream of both receptors.

Molecular interaction between Sigma-1R and TRPA1

We found that Sigma-1R antagonists reduce the plasma membrane expression of functional TRPA1 channels. The FRET experiments demonstrated that Sigma-1R and TRPA1 are close enough to form a molecular complex at the plasma membrane, a result corroborated by immunoprecipitation. We also show that FRET efficiency decreases after incubation with S1RA, indicating structural rearrangements in the TRPA1-Sigma-1R complex. Previous studies, using intramolecular FRET, showed that TRPA1 tetramers undergo notable conformational changes in the presence of agonists (e.g. calcium).77 We also show that S1RA incubation reduces biotinylated (i.e. plasma membrane TRPA1 levels) without affecting total TRPA1 expression, and impairs the repopulation of TRPA1 channels at the plasma membrane. We interpret this result as further evidence that the main effect of S1RA is on subcellular redistribution (i.e. assembly, folding or trafficking) of the channel, rather than direct interaction with the channel pore. Previous studies have shown that Sigma-1R can affect cellular excitability by modulating the traffic of various ion channels (reviewed by Soriani and Kourrich³⁵). Indeed, the same interpretation to ours was suggested for similar results obtained in a previous study examining the interaction between Sigma-1R and Kv1.2 channels in the nucleus accumbens.³⁸

We have not elucidated the subcellular location of the interaction between Sigma-1R and TRPA1. An obvious candidate is the endoplasmic reticulum-associated membrane, rich in Sigma-1R.³⁴ Another possible locus of modulation is the alteration in the subcellular localization of TRPA1. Both, Sigma-1R^{78–80} and TRPA1⁸¹ reside in membrane lipid rafts, possibly facilitating their interaction. Moreover, Sigma-1R silencing shifted these channels from lipid nanodomains to non-raft fractions.⁸⁰ In the case of TRPA1, disruption of lipid rafts reduced their activity.⁸¹ Future studies should address the interesting possibility that cholesterol or other lipids participate in the interaction between Sigma1-R and TRPA1.

Ankyrin repeats are structural motifs found in many proteins, including TRP channels.⁸² These repeats are associated with protein–protein interactions and have been postulated to act as mechanosensitive springs.^{82–84} TRPA1 function is modulated through its long ankyrin repeat-rich amino-terminal domain.⁸⁵ Previous work also demonstrated that Sigma-1R can form a complex with ankyrin B and inositol 14,5-trisphosphate receptors at the endoplasmic reticulum membrane, and this trimer was regulated by Sigma-1R agonists.⁸⁶ Future mutagenesis studies can address the relevance of TRPA1 ankyrin repeats in the interaction between both proteins.

Sigma-1R can exist in various oligomeric forms, with antagonists favouring higher order oligomerizations and agonists having the opposite effect.⁸⁷ The incubation with S1RA increases the proportion of Sigma-1R multimers,⁷⁴ possibly altering the interaction with TRPA1, as suggested by the reduced FRET signal that we observed between both proteins. This may lead to a misfolded TRPA1, interfering with the forward trafficking and reducing its functional expression at the plasma membrane without alteration in total TRPA1 protein levels.

Sigma-1R and CIPN

Chemotherapy administration to cancer patients is a planned procedure, allowing for the design of preventive strategies for minimizing neurotoxic side effects. In a clinical trial of patients treated with oxaliplatin for colorectal cancer, S1RA showed a protective effect on neuropathic symptoms.43 Moreover, in rats, co-administration of S1RA and oxaliplatin prevented the development of cold allodynia.45 We demonstrated similar effects with the anticipated systemic injection of S1RA in oxaliplatintreated mice. The reduction in mechanical and cold hypersensitivity extended 3 days past the last S1RA administration, suggesting a long-lasting effect on TRPA1 activity, well beyond the plasma half-life of the drug.⁴² This interpretation is reinforced by the demonstration of in vitro effects on TRPA1 activity, several days after the in vivo application of S1RA. However, ligand association and dissociation at the Sigma-1R is very slow,⁸⁸ possibly explaining the long-lasting effects we observed. Monitoring the reduction in neuropathic symptoms over longer times after ending the oxaliplatin treatment will be useful in defining the potential use of S1RA or other Sigma-1R antagonists in preventing CIPN in cancer patients. Future studies, examining the effects of Sigma-1R antagonists on human nociceptors derived from induced pluripotent stem cells (iPSCs) should provide further insight into the mechanisms of CIPN.

TRPA1 is expressed in many barrier tissues, including the skin, gut and airways (reviewed by Viana²⁷). Because Sigma-1R are also widely expressed in the same tissues,³⁰ the potential for functional interactions between TRPA1 and Sigma-1R beyond the one described here is a distinct possibility worth exploring, and potentially relevant for treatment of other diseases like chronic itch, asthma and inflammatory bowel disease.

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Competing interests

J.M.V. and L.R. are former full-time employees of ESTEVE Pharmaceuticals S.A., which partially financed the study. All other authors report no competing interests.

Supplementary material

Supplementary material is available at Brain online.

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