

# Opioid-Induced Hyperalgesic Priming in Single Nociceptors

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Clinical  $\mu$ -opioid receptor (MOR) agonists produce hyperalgesic priming, a form of maladaptive nociceptor neuroplasticity, resulting in pain chronification. We have established an *in vitro* model of opioid-induced hyperalgesic priming (OIHP), in male rats, to identify nociceptor populations involved and its maintenance mechanisms. OIHP was induced *in vivo* by systemic administration of fentanyl and confirmed by prolongation of prostaglandin  $E_2$  (PGE<sub>2</sub>) hyperalgesia. Intrathecal cordycepin, which reverses Type I priming, or the combination of Src and mitogen-activated protein kinase (MAPK) inhibitors, which reverses Type II priming, both partially attenuated OIHP. Parallel *in vitro* experiments were performed on small-diameter (<30  $\mu$ m) dorsal root ganglion (DRG) neurons, cultured from fentanyl-primed rats, and rats with OIHP treated with agents that reverse Type I or Type II priming. Enhancement of the sensitizing effect of a low concentration of PGE<sub>2</sub> (10 nM), another characteristic feature of priming, measured as reduction in action potential (AP) rheobase, was found in weakly isolectin B4 (IB4)-positive and IB4-negative (IB4<sup>-</sup>) neurons. In strongly IB4-positive (IB4<sup>+</sup>) neurons, only the response to a higher concentration of PGE<sub>2</sub> (100 nM) was enhanced. The sensitizing effect of 10 nM PGE<sub>2</sub> was attenuated in weakly IB4<sup>+</sup> and IB4<sup>-</sup> neurons cultured from rats whose OIHP was reversed *in vivo*. Thus, *in vivo* administration of fentanyl induces neuroplasticity in weakly IB4<sup>+</sup> and IB4<sup>-</sup> nociceptors that persists *in vitro* and has properties of Type I and Type II priming. The mechanism underlying the enhanced sensitizing effect of 100 nM PGE<sub>2</sub> in strongly IB4<sup>+</sup> nociceptors, not attenuated by inhibitors of Type I and Type II priming, remains to be elucidated.

**Key words:** excitability; fentanyl; isolectin B4; neuroplasticity; nociceptor; sensitization

## Significance Statement

Commonly used clinical opioid analgesics, such as fentanyl and morphine, can produce hyperalgesia and chronification of pain. To uncover the nociceptor population mediating opioid-induced hyperalgesic priming (OIHP), a model of pain chronification, and elucidate its underlying mechanism, at the cellular level, we established an *in vitro* model of OIHP. In dorsal root ganglion (DRG) neurons cultured from rats primed with fentanyl, robust nociceptor population-specific changes in sensitization by prostaglandin  $E_2$  (PGE<sub>2</sub>) were observed, when compared with nociceptors from opioid naive rats. In DRG neurons cultured from rats with OIHP, enhanced PGE<sub>2</sub>-induced sensitization was observed *in vitro*, with differences identified in non-peptidergic [strongly isolectin B4 (IB4)-positive] and peptidergic [weakly IB4-positive (IB4<sup>+</sup>) and IB4-negative (IB4<sup>-</sup>)] nociceptors.

## Introduction

Opioid analgesics remain among the most effective treatments for moderate-to-severe pain (Manchikanti et al., 2012; Cheung et al., 2014; Ballantyne, 2015; Bicket et al., 2019; Pisanu et al., 2019; Preuss et al., 2020). They can, however, produce serious adverse effects, including analgesic tolerance, addiction, opioid-induced hyperalgesia (OIHP), and pain chronification (Ossipov et al.,

2005; Chu et al., 2006; De Felice and Porreca, 2009; Roeckel et al., 2016; Mercadante et al., 2019), which limits their use and contributes to their abuse. Even when opioid analgesics are used briefly, in the perioperative period, they may lead to increased, postoperative pain and opioid requirements, up to a year later (Chia et al., 1999; van Gulik et al., 2012; Rashiq and Dick, 2014; Yildirim et al., 2014; Lyons et al., 2015; Chang et al., 2018; Rong et al., 2019), indicating that opioids can produce long-lasting alterations in pain processing, well beyond the duration of opioid exposure. We have established a preclinical model of the transition from acute to chronic pain, hyperalgesic priming, here referred to as priming. Priming is a form of long-lasting nociceptor neuroplasticity manifested by a left-shift in dose dependence and marked prolongation of mechanical hyperalgesia in response to pronociceptive mediators, prototypically prostaglandin  $E_2$

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(PGE<sub>2</sub>), but unchanged baseline nociceptive threshold (Aley et al., 2000; Parada et al., 2003, 2005; Reichling and Levine, 2009; Araldi et al., 2015, 2017a, 2018b,c, 2019; Ferrari et al., 2015, 2019; Khomula et al., 2017). Thus, while in control rats PGE<sub>2</sub>-induced hyperalgesia is of short duration (<4 h), in the primed state hyperalgesia is produced by lower concentrations of PGE<sub>2</sub> (Parada et al., 2003, 2005) and is still present, unattenuated, 4 h later (Aley et al., 2000).

We have recently shown that  $\mu$ -opioid receptor (MOR) agonists, including two commonly used clinical opioids, fentanyl (Araldi et al., 2018c; Khomula et al., 2019) and morphine (Araldi et al., 2019; Ferrari et al., 2019), as well as two biased MOR agonists, PZM21 and TRV130 (Araldi et al., 2018a), can produce both OIH and opioid-induced hyperalgesic priming (OIHP). We have recently established an *in vitro* model of OIH, neuroplasticity induced by *in vivo* administration of fentanyl that persists in cultured DRG neurons, differentially affecting peptidergic and non-peptidergic nociceptors (Khomula et al., 2019). However, despite a growing literature obtained from *in vivo* experiments, lack of an *in vitro* model of OIHP has made it difficult to study underlying mechanisms at the cellular level, in identified nociceptor populations.

Primary afferent nociceptors can be divided into three categories based on isolectin B4 (IB4)-binding status: strongly IB4-positive (IB4+), weakly IB4+, and IB4-negative (IB4-; Kashiba et al., 2001; Fang et al., 2006). IB4+ neurons depend on glial-derived neurotrophic factor (GDNF), and express Ret, whereas IB4- neurons depend on NGF for survival and express TrkA (Molliver et al., 1997; Bennett et al., 1998; Kashiba et al., 1998). Weakly IB4+ neurons also express TrkA (Kashiba et al., 2001; Fang et al., 2006). Nociceptors can also be classified into non-peptidergic and peptidergic (Bennett et al., 1996; Molliver et al., 1997; Woolf and Ma, 2007). Strongly IB4+ nociceptors belong to the non-peptidergic class, whereas weakly IB4+ and IB4- nociceptors are peptidergic (Kashiba et al., 2001; Fang et al., 2006). These classes differ in electrophysiological properties (Stucky and Lewin, 1999), physiological function (Snider and McMahon, 1998), and their role in OIHP (Araldi et al., 2015, 2018c, 2019; Ferrari et al., 2019).

To establish an *in vitro* model of OIHP we (1) produced OIHP *in vivo*, assessed at the nociceptor peripheral and central terminal as prolongation of PGE<sub>2</sub>-induced mechanical hyperalgesia, in rats treated with systemic fentanyl; (2) established, *in vivo*, a protocol to reverse OIHP and also evaluate whether systemic fentanyl-induced priming occurs in peptidergic and/or nonpeptidergic neurons; (3) employed, *in vitro*, concentration dependence of PGE<sub>2</sub>-induced sensitization, in populations of cultured DRG neurons (i.e., strongly and weakly IB4+, and IB4-) from rats primed *in vivo* with systemic fentanyl, to reveal *in vitro* correlates of OIHP at the single cell level; and (4) tested putative maintenance mechanisms of OIHP in nociceptor populations classified by their IB4-binding status (based on *in vitro* correlates, in cultured DRG neurons from animals in which OIHP was reversed *in vivo*).

## Materials and Methods

### Animals

Experiments were performed on 220–420 g male Sprague Dawley rats (Charles River Laboratories). Experimental animals were housed three per cage, under a 12/12 h light/dark cycle, in a temperature-controlled and humidity-controlled animal care facility at the University of California, San Francisco. Food and water were available *ad libitum*. Nociceptive testing was performed between 9 A.M. and 5 P.M. All experimental protocols approved by the Institutional Animal Care and Use

Committee at the University of California, San Francisco and adhered to the National Institutes of Health *Guide for the Care and Use of Laboratory Animals*. Effort was made to minimize number of animals used and their suffering.

### Nociceptive threshold testing

Mechanical nociceptive threshold was measured using an Ugo Basile Analgesymeter (Randall-Selitto paw-withdrawal device, Stoelting), which applies a linearly increasing mechanical force to the dorsum of a rat's hind paw, with a dome-shaped plinth, as previously described (Taiwo et al., 1989; Taiwo and Levine, 1989; Araldi et al., 2015, 2017a; Ferrari and Levine, 2015). Rats were placed in cylindrical acrylic restrainers that provide ventilation, and lateral ports that allow extension of their hind legs for the assessment of nociceptive threshold, while minimizing restraint stress. To acclimatize rats to the testing procedure, they were placed in restrainers for 40 min before starting training sessions (three consecutive days of training) and for 30 min before experimental manipulations. Nociceptive threshold was defined as the force, in grams, at which the rat withdrew its paw. Baseline nociceptive threshold for each individual rat was defined as the mean of three readings taken before the first experimental treatment. In each experiment, only one paw per rat was used. Each experiment was performed on a different group of rats (one paw/rat, six rats/group). The individuals performing behavioral experiments (D.A. and I.J.M.B.) were blinded to treatments.

### Drugs and routes of administration

The following compounds were used in these experiments: cordycepin 5'-triphosphate sodium salt (a protein translation inhibitor), fentanyl citrate salt (a MOR agonist), SU6656 (a Src family kinase inhibitor), U0126 [a MAPK/extracellular signal-regulated kinase [ERK] inhibitor], and PGE<sub>2</sub> (a direct-acting nociceptor-sensitizing agent), all of which were purchased from Sigma-Aldrich.

A stock solution of PGE<sub>2</sub> (1  $\mu$ g/ $\mu$ l) was prepared in 100% ethanol and additional dilutions made with physiological saline (0.9% NaCl), to prepare the final solution used in experiments (100 ng/5  $\mu$ l), which yielded a final ethanol concentration of 2%. In previous *in vivo* control experiments, we have shown that ethanol in this concentration had no effect on mechanical nociceptive threshold (Ferrari et al., 2016). Fentanyl and cordycepin were dissolved in saline. SU6656 and U0126 were dissolved in 100% dimethylsulfoxide (DMSO; Sigma-Aldrich) and further diluted in saline containing 2% Tween 80 (Sigma-Aldrich). The final concentration of DMSO and Tween 80 was ~2%.

Intradermal administration of PGE<sub>2</sub> (100 ng/5  $\mu$ l) was performed on the dorsum of the hindpaw using a 30-gauge hypodermic needle adapted to a 50  $\mu$ l Hamilton syringe by a segment of PE-10 polyethylene tubing (Becton Dickinson).

Intrathecal administration of PGE<sub>2</sub>, cordycepin, the combination of SU6656 and U0126, and the combination of cordycepin, SU6656, and U0126 was performed in rats briefly anesthetized with 2.5% isoflurane (Phoenix Pharmaceuticals) in 97.5% O<sub>2</sub>, using a 29-gauge hypodermic needle (300 unit/ $\mu$ l syringe; Walgreens Pharmacy) inserted into the subarachnoid space, between the L4 and L5 vertebrae. The maximum intrathecal volume administered was 20  $\mu$ l; cordycepin (4  $\mu$ g) was dissolved in saline and injected intrathecally in a volume of 20  $\mu$ l, when followed by intradermal PGE<sub>2</sub> (100 ng/5  $\mu$ l), and in a volume of 10  $\mu$ l when followed by intrathecal PGE<sub>2</sub> (400 ng/10  $\mu$ l; Araldi et al., 2018c). The combination of SU6656 (10  $\mu$ g) and U0126 (10  $\mu$ g) was injected intrathecally in a final volume of 20 or 10  $\mu$ l (these drugs were mixed in the syringe at the moment of injection), followed by PGE<sub>2</sub> injected intradermally (100 ng/5  $\mu$ l) or intrathecally (400 ng/10  $\mu$ l), respectively. The combination of all three inhibitors [cordycepin (4  $\mu$ g/6  $\mu$ l), SU6656 (10  $\mu$ g/6  $\mu$ l), and U0126 (10  $\mu$ g/6  $\mu$ l)] was injected intrathecally followed, 10 min later, by intradermal PGE<sub>2</sub> (100 ng/5  $\mu$ l). To verify whether the inhibitors reversed systemic fentanyl-induced priming permanently, PGE<sub>2</sub> was again injected intradermally (100 ng/5  $\mu$ l) or intrathecally (400 ng/20  $\mu$ l), 4, 14, and 28 days after the inhibitors. The intrathecal site of injection was confirmed by a sudden flick of the rat's tail, a reflex that is evoked by accessing the subarachnoid space followed by bolus intrathecal injection (Mestre et al., 1994).

Systemic (subcutaneous) administration of fentanyl was performed at the nape of the neck (Araldi et al., 2018c, 2019). Rats received an injection of fentanyl (30  $\mu\text{g}/\text{kg}$ , s.c.) diluted in saline and administered subcutaneously (100  $\mu\text{l}/100$  g body weight).

### Administration of saporins

#### IB4-saporin

IB4-saporin, an IB4<sup>+</sup> nociceptor neurotoxin (Advanced Targeting Systems), was diluted in saline and a dose of 3.2  $\mu\text{g}$ , in a volume of 20  $\mu\text{l}$ , administered intrathecally, 14 d before systemic administration of fentanyl (30  $\mu\text{g}/\text{kg}$ ). The dose and timing of IB4-saporin administration were chosen based on previous reports from our group and others (Vulchanov et al., 2001; Nishiguchi et al., 2004; Joseph et al., 2008, 2010; Araldi et al., 2015, 2016a, 2017b, 2018c).

#### [Sar<sup>9</sup>, Met(O<sub>2</sub>)<sup>11</sup>]-substance P-saporin (SSP-saporin)

SSP-saporin, a neurotoxin specifically targeting neurons that express substance P (SP) receptors (Advanced Targeting Systems), was diluted in saline and a dose of 100 ng in a volume of 20  $\mu\text{l}$  administered intrathecally 14 d before rats received a systemic fentanyl (30  $\mu\text{g}/\text{kg}$ ). The addition of [Sar<sup>9</sup>, Met(O<sub>2</sub>)<sup>11</sup>] to the SP-conjugated to saporin makes the agent more stable and potent than when SP alone is bound to saporin. The dose and time point of the treatment (14 d before fentanyl) were based on previous studies (Wiley et al., 2007; Choi et al., 2012), where there was observed loss of intrinsic lumbar dorsal horn neurons expressing neurokinin 1 receptor (NK1R), a receptor for SP, in Laminae I–III but not in deeper laminae (Khasabov et al., 2002; Vierck et al., 2003; Wiley et al., 2007; Choi et al., 2012; Weisshaar and Winkelstein, 2014; Kras et al., 2015; Araldi et al., 2016b, 2017b, 2018b,c), thus disrupting signaling dependent on peptidergic nociceptors (Seki et al., 2005).

We used rats treated with unconjugated saporin as controls, since previous studies demonstrated that intrathecal administration of saporin alone has no effects on NK1R staining in the spinal cord (Hwang et al., 2003) and does not decrease IB4 labeling in Lamina II of the dorsal horn (Joseph et al., 2008). The control group received 3.2  $\mu\text{g}$  of unconjugated saporin, diluted in 20  $\mu\text{l}$  of saline, intrathecally. This dose (3.2  $\mu\text{g}$ ) is equal to the IB4-saporin dose and higher than SSP-saporin (100 ng).

To inject unconjugated saporin, IB4-saporin, or SSP-saporin, rats were briefly anesthetized with 2.5% isoflurane (Phoenix Pharmaceuticals) in 97.5% O<sub>2</sub>, and then a 29-gauge hypodermic needle (300 units/ $\mu\text{l}$  syringe, Walgreens Pharmacy) was inserted, on the midline, into the subarachnoid space, between the L4 and L5 vertebrae. Rats regained consciousness  $\sim$ 2 min after stopping anesthesia. There was no effect of unconjugated saporin, IB4-saporin, or SSP-saporin on mechanical nociceptive threshold per se ( $t_{(5)} = 0.6$ ;  $p = 0.59$ , for the unconjugated saporin-treated group,  $t_{(5)} = 1.3$ ;  $p = 0.24$ , for the SSP-saporin-treated group, and  $t_{(5)} = 0.5$ ;  $p = 0.66$ , for the IB4-saporin-treated group, when the mechanical nociceptive threshold is compared before and 14 d after intrathecal saporins; paired Student's  $t$  test).

### Culture of DRG neurons

Primary cultures of dorsal root ganglion (DRG) neurons were made from adult male Sprague Dawley rats (220–235 g), as described previously (Ferrari et al., 2016, 2018; Khomula et al., 2017; Araldi et al., 2018c). In brief, under isoflurane anesthesia, rats were decapitated, and the dorsum of the vertebral column surgically removed; L<sub>4</sub> and L<sub>5</sub> DRGs were rapidly extracted, bilaterally, chilled, and desheathed in HBSS, on ice. Ganglia were then treated with 0.25% collagenase type 4 (Worthington Biochemical Corporation) in HBSS for 18 min at 37°C, and then treated with 0.25% trypsin (Worthington Biochemical Corporation) in calcium-free and magnesium-free PBS (Invitrogen Life Technologies) for 6 min, followed by three washes, and then trituration in Neurobasal-A medium (Invitrogen Life Technologies) to produce a single-cell suspension. This suspension was centrifuged at 1000 RPM for 3 min and re-suspended in Neurobasal-A medium supplemented with 50 ng/ml nerve growth factor, 100 U/ml penicillin/streptomycin, B-27, GlutaMAX, and 10% fetal bovine serum (Invitrogen Life Technologies). Cells were then plated on cover slips and incubated at 37°C in 3.5% CO<sub>2</sub> for at least 24 h before use in experiments.

### In vitro patch-clamp electrophysiology

Cultured DRG neurons were used in *in vitro* electrophysiology experiments 24–96 h after DRG dissociation and plating. DRG from at least three rats (separate culture preparations) were used for each experimental series. Within the text,  $n$  refers to the number of neurons. Cells were identified as neurons by their double birefringent plasma membranes (Cohen et al., 1968; Landowne, 1993). While small, medium, and large sized DRG neurons were routinely observed in the same preparation, this study focused on cells with a soma diameter  $<30$   $\mu\text{m}$  (small DRG neurons), predominantly representing C-type nociceptors (Harper and Lawson, 1985; Gold et al., 1996b; Petruska et al., 2000, 2002; Woolf and Ma, 2007). After mounting a coverslip with cells in the recording chamber, the culture medium was replaced with Tyrode's solution containing 140 mM NaCl, 4 mM KCl, 2 mM MgCl<sub>2</sub>, 2 mM CaCl<sub>2</sub>, 10 mM glucose, and 10 mM HEPES and adjusted to pH 7.4 with NaOH; osmolarity is 310 mOsm/kg (Ferrari et al., 2016, 2018; Khomula et al., 2017; Araldi et al., 2018c). Tyrode's solution was used as external perfusion solution in the *in vitro* experiments. Drugs used *in vitro* were diluted to their final concentration in this solution just before application. The volume of the recording chamber is 150  $\mu\text{l}$ . The perfusion system is gravity-driven, flow rate of 0.5–1 ml/min. All experiments were performed at room temperature (20–23°C).

Whole-cell patch-clamp recordings, in current clamp mode, were made to assess changes in the excitability of cultured DRG neurons. Holding current was adjusted to maintain membrane potential at  $-70$  mV. Rheobase, the minimum magnitude of a current step needed to elicit an action potential (AP), was determined from a protocol using increasing square wave current pulses. To estimate rheobase, a series of current steps with 200-pA increments was applied until APs were elicited. The protocol was then adjusted to five to six pulses with step size of 5–10% of the rheobase estimate (two to three subthreshold stimuli and two to three with AP generation, to avoid overstimulation/desensitization; Ferrari et al., 2018; Khomula et al., 2019).

Recording electrodes were fabricated from borosilicate glass capillaries (0.84/1.5 mm i.d./o.d., Warner Instruments, LLC) using a Flaming/Brown P-87 puller (Sutter Instrument Co). Recording electrode resistance was  $\sim$ 3 M $\Omega$  after being filled with a solution containing the following: 130 mM KCl, 10 mM HEPES, 10 mM EGTA, 1 mM CaCl<sub>2</sub>, 5 mM MgATP, and 1 mM Na-GTP; pH 7.2 (adjusted with Tris-base), 300 mOsm (measured by Wescor Vapro 5520 osmometer, ELITech Group; Ferrari et al., 2018). Junction potential was not adjusted. Series resistance was below 20 M $\Omega$  at the end of recordings and was not compensated. Recordings were made with an Axon MultiClamp 700 B amplifier, filtered at 10 kHz, and sampled at 20 kHz using Axon Digidata 1550B controlled by pCLAMP 11 software (all from Molecular Devices LLC).

Drugs were applied at least 10 min after the establishment of whole cell configuration, at which time baseline current was stable.

### Identification of priming *in vitro*

We have previously shown that PGE<sub>2</sub> sensitizes small-diameter DRG neurons, putative nociceptors, *in vitro* (Gold et al., 1996b,c, 1998). Within 1–2 min, sensitization can be detected in “whole-cell” patched neurons, in current clamp mode, as a reduction of rheobase, the minimal sustained current required to generate an AP, for a saturating concentration of PGE<sub>2</sub> (1  $\mu\text{M}$ ), in both IB4<sup>-</sup> and IB4<sup>+</sup> small-diameter DRG neurons (Gold et al., 1996b; Ferrari et al., 2018). Since it is not practical to hold cells “clamped” for 4 h, to reveal priming, we relied on another *in vitro* feature of hyperalgesic priming, a more than one order magnitude leftward shift (to lower values) for the dose dependence of PGE<sub>2</sub>-induced hyperalgesia (Parada et al., 2005). Since only  $\sim$ 60% of cultured DRG neurons are responsive to PGE<sub>2</sub> (Gold et al., 1996b), it would not be possible to distinguish between responsive and nonresponsive neurons by their response to a low concentration of PGE<sub>2</sub>; response to a higher concentration is required. Previously, it has been shown that the dose dependence for the *in vitro* sensitizing effect of PGE<sub>2</sub> ranges from concentrations of 1 nM to 1  $\mu\text{M}$  (Gold et al., 1996b). Therefore, we chose two PGE<sub>2</sub> concentrations, 10 and 100 nM, to examine our hypothesis that priming *in vivo* induces a leftward shift of the dose response curve in primed DRG neurons.

### Fluorescence imaging of *in vitro* IB4 binding

The bright-field imaging system consisted of an inverted microscope (Eclipse TE-200, Nikon Instruments Inc) with epi-fluorescence using a xenon lamp (Lambda LS, Sutter Instruments Co) for excitation (Ferrari et al., 2018). Illumination was controlled by a Lambda 10–2 filter wheel and Lambda SC Smart Shutter controllers (Sutter Instruments Co); an Andor Clara Interline CCD camera (Andor Technology Ltd.) was used for high-resolution digital image acquisition. MetaFluor software (Molecular Devices LLC) provided computer interface and controlled the whole system as well as being used for image processing. A Plan Fluor objective (20× UV, NA 0.50; Nikon Instruments Inc) was used for both fluorescent and transmitted light imaging with phase contrast illumination.

Cells were incubated in Tyrode's solution supplemented with 10 μg/ml IB4 conjugated to Alexa Fluor 488 dye (Invitrogen Life Technologies) for 10–12 min, in the dark. After washout, fluorescent images were captured during the first 15 min of each experiment using a standard GFP filter set (Chroma Technology). Cells demonstrating bright fluorescence and a halo around the neuronal plasma membrane were recognized as IB4+ (strongly positive, if intensity above 40% of maximum for the selected field of view, and weakly positive if intensity in the range 20–40% of maximum), whereas those having intensity below 20% of maximum were considered as IB4– (Fang et al., 2006; Khomula et al., 2013, 2017, 2019; Ferrari et al., 2016).

### Data analysis

All data are presented as mean ± SEM of *n* independent observations. In all figures bars show mean and error bars show SEM. Statistical comparisons were made using GraphPad Prism 8.0 statistical software (GraphPad Software Inc); *p* < 0.05 was considered statistically significant.

In behavioral experiments, the dependent variable was mechanical paw-withdrawal threshold. Each data point was the mean of three readings. All magnitudes were expressed as percentage change from baseline. Average mechanical nociceptive threshold before systemic fentanyl was 143.4 ± 1.3 g and 4 d after was 142.8 ± 1.14 (*n* = 60 rats; *t*<sub>(59)</sub> = 0.48, *p* = 0.64, paired Student's *t* test).

In electrophysiological experiments, magnitude of the sensitizing effect of PGE<sub>2</sub> was expressed as percentage reduction in rheobase (relative to a baseline rheobase value recorded before PGE<sub>2</sub> administration). Only neurons sensitized above 10% by 10 min after PGE<sub>2</sub>, at the lower (10 nM) or higher (100 nM) concentration, applied to the same neuron, were considered PGE<sub>2</sub> responsive, and included in our analysis.

As described in detail in the figure legends, the following statistical tests were used: paired (Fig. 1) and unpaired (Fig. 5C) Student's *t* test, two-way repeated-measures ANOVA followed by Bonferroni's *post hoc* test (Figs. 1–3), one-way ANOVA followed by Tukey's *post hoc* test (Figs. 6D, 7D, 8), two-way ANOVA followed by Holm–Sidak's *post hoc* test (Figs. 5A,B,D, 6A–C, 7A–C).

## Results

### Systemic fentanyl induces Type I and Type II priming at the nociceptor peripheral terminal

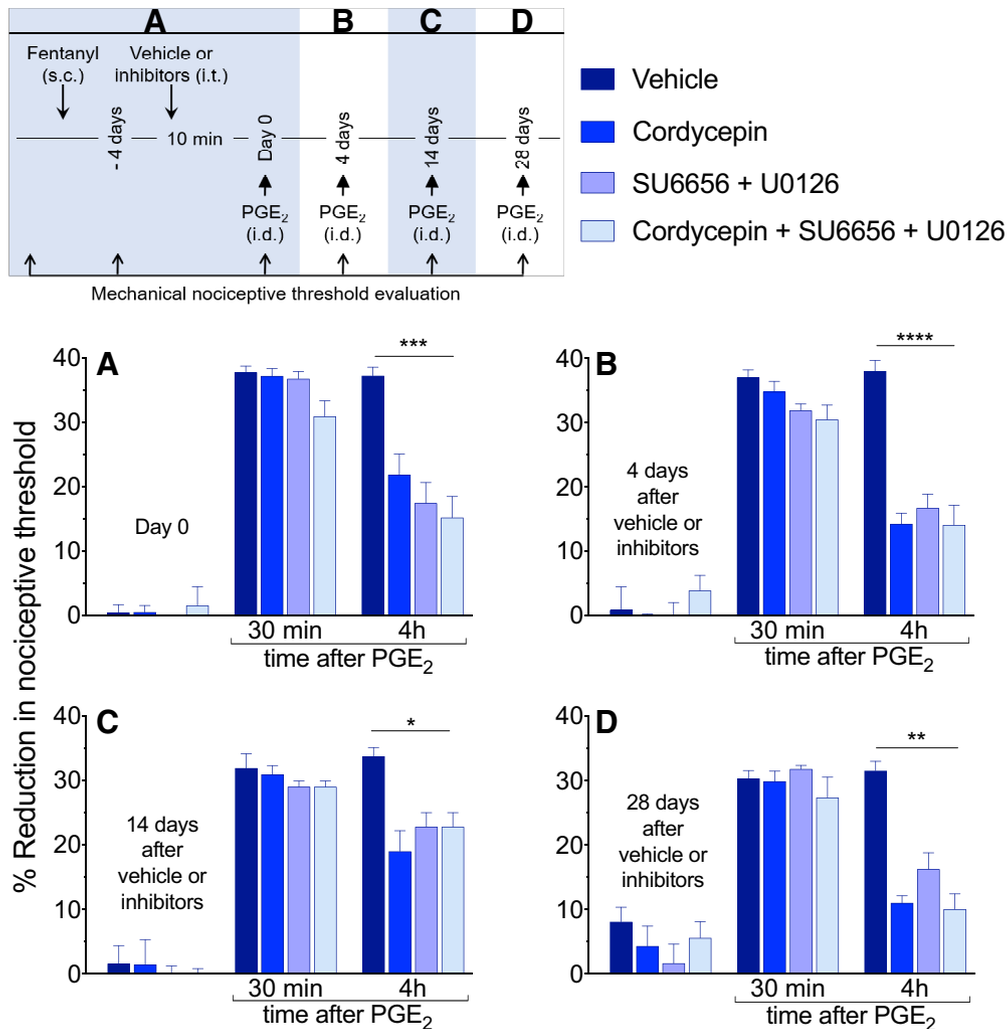
We recently demonstrated the presence of OIH and OIHP, *in vivo*, at the nociceptor peripheral terminal after a single systemic (subcutaneous) injection of an analgesic dose of fentanyl (30 μg/kg; Khomula et al., 2019). We also demonstrated that OIH, induced by systemic fentanyl, is attenuated by administration of cordycepin at the peripheral terminal (Khomula et al., 2019), indicative of Type I priming (Ferrari et al., 2013). More recently, we identified a second type of hyperalgesic priming, referred to as Type II (Araldi et al., 2015, 2018c). In distinction to the maintenance mechanisms for Type I priming (Ferrari et al., 2013), Type II is dependent on the combined activation of Src and MAP kinases (Araldi et al., 2017a). While we recently demonstrated that OIH induced by systemic fentanyl is reversed by cordycepin (Khomula et al., 2019), the maintenance mechanism of

OIHP has not been tested. Therefore, we evaluated, using the same *in vivo* model, whether hyperalgesic priming in the peripheral terminal is Type I (reversed by cordycepin) or Type II (reversed by the combination of Src and MAPK inhibitors).

Rats received a systemic injection of fentanyl (30 μg/kg, s.c.) followed, 4 d later by intrathecal injection of vehicle (20 μl), cordycepin (4 μg/20 μl), the combination of Src (SU6656, 10 μg/10 μl) and MAPK (U0126, 10 μg/10 μl) inhibitors, or a combination of all three drugs [cordycepin (4 μg/6 μl), SU6656 (10 μg/6 μl), and U0126 (10 μg/6 μl)]. Ten minutes later, PGE<sub>2</sub> (100 ng) was injected intradermally and mechanical nociceptive threshold evaluated a further 30 min and 4 h later. When measured 30 min after injection, PGE<sub>2</sub>-induced hyperalgesia in all groups. However, the prolongation of PGE<sub>2</sub> hyperalgesia, measured at the fourth hour, was attenuated in the groups treated with cordycepin, SU6656 + U0126 or the combination of all three inhibitors ( $F_{(3,20)} = 10.8$ , *p* = 0.0002, when the vehicle-treated and inhibitors-treated groups are compared at the fourth hour after intradermal PGE<sub>2</sub>; two-way ANOVA multiple comparisons test; Fig. 1A). To determine whether intrathecal administration of inhibitors was able to permanently attenuate priming induced by systemic fentanyl, the same groups of rats again received intradermal PGE<sub>2</sub> (100 ng), 4 d (Fig. 1B), 14 d (Fig. 1C), and 28 d (Fig. 1D) after vehicle or inhibitors. In the groups treated with cordycepin, the combination of SU6656 + U0126, and the combination of all three inhibitors, the prolongation of PGE<sub>2</sub>-induced hyperalgesia was attenuated, at all time points evaluated [4 d, (Fig. 1B)  $F_{(3,20)} = 23.6$ , *p* < 0.0001;  $F_{(3,20)} = 15.4$ , *p* < 0.0001; 14 d (Fig. 1C)  $F_{(3,20)} = 4.1$ , *p* = 0.02; and 28 d (Fig. 1D)  $F_{(3,20)} = 7.9$ , *p* = 0.0011, when the vehicle-treated group is compared with all the inhibitors-treated groups at the fourth hour after intradermal PGE<sub>2</sub>; two-way ANOVA multiple comparisons test]. These results provide confirmation that the reversal effects of cordycepin, and the combination of Src and MAPK inhibitors persisted for at least 28 d. In contrast, in the vehicle-treated group, priming did not reveal any attenuation out to day 32 after induction, as demonstrated by unattenuated magnitude of PGE<sub>2</sub>-induced hyperalgesia at 4 h. Altogether, these findings indicate that in the peripheral terminal of the nociceptor, maintenance of OIHP produced by a single systemic administration of fentanyl (30 μg/kg), shares mechanisms with both Type I and Type II priming. Of note, the combination of all three inhibitors (cordycepin + SU6656 + U0126) did not produce additional attenuation of PGE<sub>2</sub>-induced hyperalgesia, at 4 h, in systemic fentanyl-primed rats, when it was compared with the cordycepin-treated and SU6656 + U0126-treated groups.

### Priming induced by systemic fentanyl in the central terminal of the nociceptor is Type II

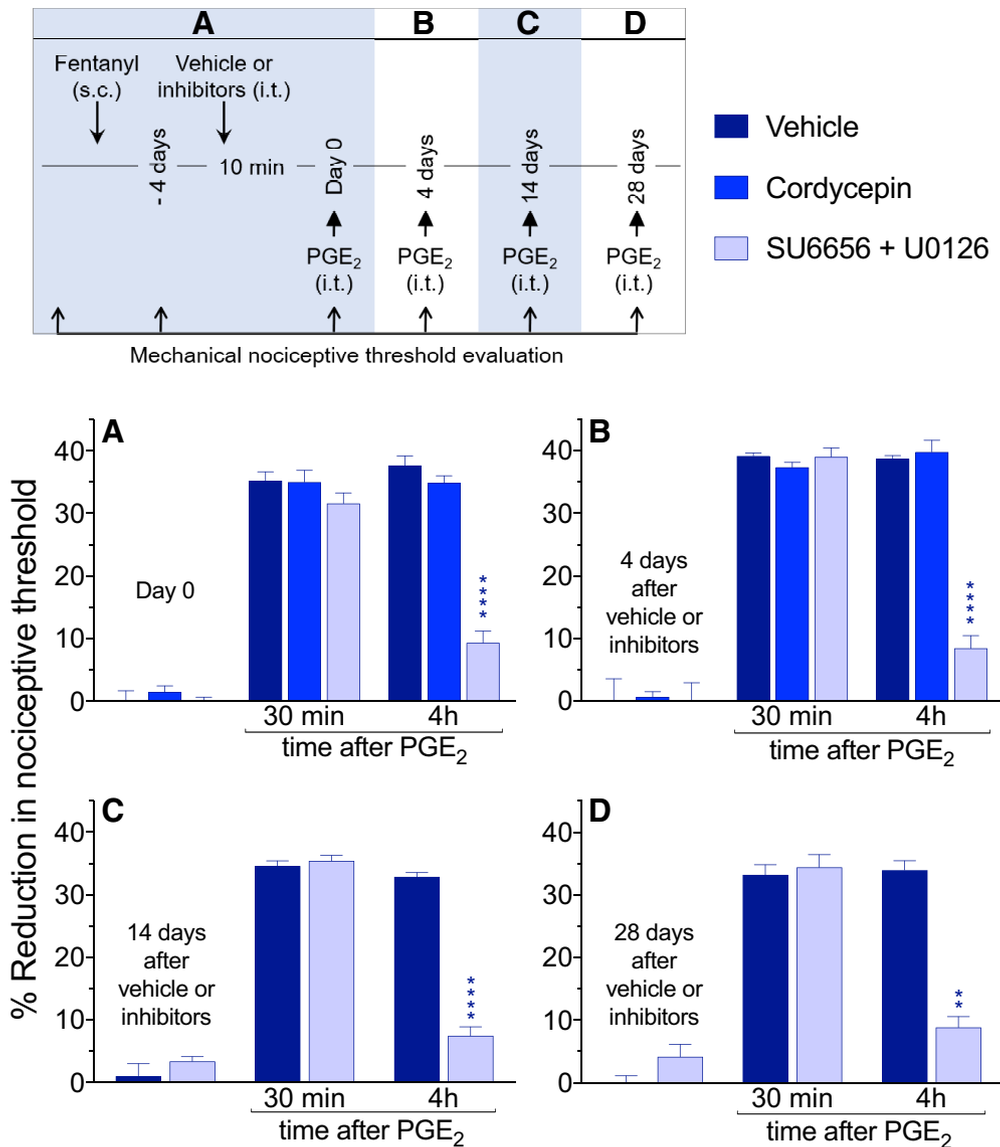
Systemic fentanyl at an analgesic dose (30 μg/kg) induces both Type I and Type II priming in the peripheral terminal of the nociceptor (Fig. 1). We next investigated the type of priming that systemic fentanyl induces in the central terminal. Rats received systemic fentanyl (30 μg/kg, s.c.) and 4 d later, vehicle (10 μl), cordycepin (4 μg/10 μl), or the combination of Src (SU6656, 10 μg/5 μl) and MAPK (U0126, 10 μg/5 μl) inhibitors was administered intrathecally, followed, 10 min later by PGE<sub>2</sub> (400 ng/10 μl), injected at the same site. Mechanical nociceptive threshold was evaluated 30 min and 4 h after intrathecal PGE<sub>2</sub>. Measured 30 min after its administration, intrathecal PGE<sub>2</sub> induced hyperalgesia in all groups. The prolongation of hyperalgesia induced by intrathecal PGE<sub>2</sub> was markedly attenuated in the group treated with the combination of SU6656 + U0126, but not in the vehicle-treated and cordycepin-treated groups ( $F_{(2,15)} = 49.9$ ,



**Figure 1.** Hyperalgesic priming in the peripheral terminal of the nociceptor, induced by systemic fentanyl, is attenuated by inhibitors of protein translation and a combination of a Src and MAP kinases. **A**, Male rats received a single systemic injection of an analgesic dose of fentanyl (30  $\mu$ g/kg, s.c.). Four days later, at which time the mechanical nociceptive threshold was not different from pre-fentanyl baseline (Day 0;  $t_{(23)} = 1.2$ ;  $p = 0.26$ , when the mechanical nociceptive threshold is compared before and 4 d after systemic fentanyl; paired Student's  $t$  test), vehicle (2% DMSO in saline, 20  $\mu$ l), cordycepin (4  $\mu$ g/20  $\mu$ l), the combination of Src kinase inhibitor (SU6656, 10  $\mu$ g/10  $\mu$ l) and MAPK inhibitor (U0126, 10  $\mu$ g/10  $\mu$ l) or the combination of all three inhibitors (cordycepin + SU6656 + U0126: 4  $\mu$ g/6  $\mu$ l, 10  $\mu$ g/6  $\mu$ l, and 10  $\mu$ g/6  $\mu$ l, respectively) was administered intrathecally followed, 10 min later, by PGE<sub>2</sub> (100 ng/5  $\mu$ l) injected intradermally, on the dorsum of the hind paw. Mechanical nociceptive threshold was evaluated 30 min and 4 h after intradermal PGE<sub>2</sub>. In all groups, PGE<sub>2</sub>-induced hyperalgesia was measured 30 min after its injection. However, the prolongation of PGE<sub>2</sub>-induced hyperalgesia at the fourth hour was markedly attenuated in the groups treated with cordycepin, SU6656 + U0126, and the combination of all three inhibitors ( $F_{(3,20)} = 10.8$ ,  $***p = 0.0002$ , when the vehicle-treated group is compared with the inhibitors-treated groups at the fourth hour after intradermal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test). **B**, Four days after intrathecal treatment with vehicle or inhibitors, at which time the mechanical nociceptive threshold was not different from the pre-vehicle/inhibitor baseline ( $t_{(5)} = 1.2$ ;  $p = 0.27$ , for the vehicle-treated group,  $t_{(5)} = 0.72$ ;  $p = 0.51$ , for the cordycepin-treated group,  $t_{(5)} = 0.44$ ;  $p = 0.68$ , for the combination of Src and MAPK inhibitors-treated group, and  $t_{(5)} = 1.6$ ;  $p = 0.17$ , for the group treated with the combination of all three inhibitors, when the mechanical nociceptive threshold is compared before and after intrathecal inhibitors; paired Student's  $t$  test), PGE<sub>2</sub> (100 ng/5  $\mu$ l) was again injected. In the groups previously treated with intrathecal cordycepin, SU6656 + U0126 and the combination of all three inhibitors, prolongation of PGE<sub>2</sub>-induced hyperalgesia was markedly attenuated ( $F_{(3,20)} = 23.6$ ,  $****p < 0.0001$ , when the hyperalgesia in the vehicle-treated and the inhibitors-treated groups is compared at the fourth hour after intradermal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test). Similarly, 14 d (**C**) and 28 d (**D**) after intrathecal injection of vehicle, cordycepin, SU6656 + U0126, or the combination of all three inhibitors, when PGE<sub>2</sub> (100 ng/5  $\mu$ l) was again injected intradermally, the prolongation of PGE<sub>2</sub>-induced hyperalgesia was still attenuated in all three groups treated with the inhibitors (14 d, **C**:  $F_{(3,20)} = 4.1$ ,  $*p = 0.02$ ; and 28 d, **D**:  $F_{(3,20)} = 7.9$ ,  $**p = 0.0011$ , when the vehicle-treated group is compared with the inhibitors-treated groups at the fourth hour after intradermal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test). These findings support the suggestion that systemic fentanyl induces both Type I and Type II priming, in the peripheral terminal of the nociceptor ( $n = 6$  paws/6 rats per group).

$p < 0.0001$ , when the combination of SU6656 + U0126-treated group is compared with the vehicle-treated and the cordycepin-treated groups at the fourth hour after intrathecal PGE<sub>2</sub>; two-way ANOVA multiple comparisons test; Fig. 2A). To verify whether intrathecal treatment with inhibitors permanently reversed priming induced by systemic fentanyl in the central terminal of the nociceptor, PGE<sub>2</sub> (400 ng) was again injected 4 d

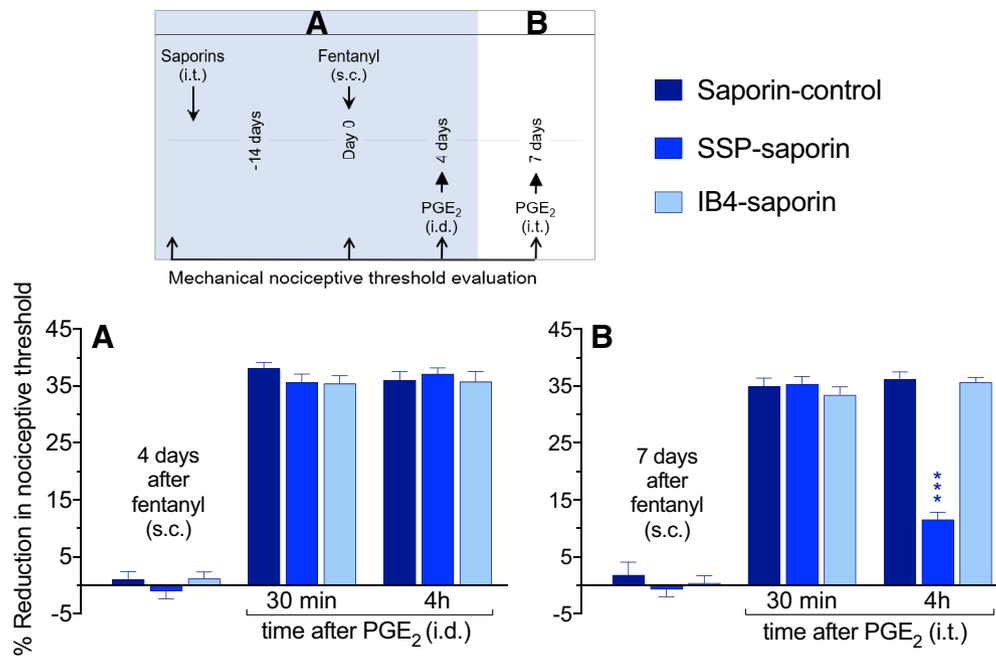
(Fig. 2B), 14 d (Fig. 2C), and 28 d (Fig. 2D) after vehicle and inhibitors. The prolongation of PGE<sub>2</sub>-induced hyperalgesia was still inhibited in the group treated with the combination of SU6656 + U0126 but not in the vehicle-treated and cordycepin-treated groups ( $F_{(2,15)} = 41.6$ ,  $p < 0.0001$ , when the combination of SU6656 + U0126-treated group was compared with the vehicle-treated and the cordycepin-treated groups at the fourth hour



**Figure 2.** Systemic fentanyl induces Type II priming in nociceptor central terminals. **A**, Male rats were treated with systemic fentanyl (30  $\mu\text{g}/\text{kg}$ , s.c.). Four days later, when mechanical nociceptive threshold was not different from pre-fentanyl baseline ( $t_{(17)} = 0.18$ ,  $p = 0.86$  when the mechanical nociceptive threshold was compared before and 4 d after systemic fentanyl; paired Student's  $t$  test), vehicle (2% DMSO in saline, 20  $\mu\text{l}$ ), cordycepin (4  $\mu\text{g}/10$   $\mu\text{l}$ ), or the combination of Src inhibitor (SU6656, 10  $\mu\text{g}/5$   $\mu\text{l}$ ) and MAPK inhibitor (U0126, 10  $\mu\text{g}/5$   $\mu\text{l}$ ) were administered intrathecally followed, 10 min later, by PGE<sub>2</sub> (400 ng/10  $\mu\text{l}$ ) administered at the same site, intrathecally. Mechanical nociceptive threshold was evaluated 30 min and 4 h after intrathecal PGE<sub>2</sub>. PGE<sub>2</sub>-induced hyperalgesia was present in all groups, 30 min after its injection. When the prolongation of PGE<sub>2</sub>-induced hyperalgesia was evaluated at the fourth hour after its injection, in the group treated with the combination of Src and MAPK inhibitors it was markedly attenuated ( $F_{(2,15)} = 47.9$ ,  $****p < 0.0001$ , when the combination of Src and MAPK inhibitors-treated group is compared with the vehicle- and cordycepin-treated groups at the fourth hour after intrathecal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test); however, no attenuation was observed in the vehicle-treated and cordycepin-treated groups. **B**, Four days after intrathecal vehicle and inhibitors, when the mechanical nociceptive threshold was not different from pre-vehicle/inhibitor baseline ( $t_{(5)} = 0.07$ ;  $p = 0.95$ , for the vehicle-treated group,  $t_{(5)} = 0.81$ ;  $p = 0.46$ , for the cordycepin-treated group, and  $t_{(5)} = 0.08$ ;  $p = 0.94$ , for the combination of Src and MAPK inhibitors-treated group, when the mechanical nociceptive threshold is compared before and after intrathecal inhibitors; paired Student's  $t$  test), PGE<sub>2</sub> (400 ng/20  $\mu\text{l}$ ) was again injected intrathecally and mechanical nociceptive threshold evaluated 30 min and 4 h later. In the group treated with the combination of Src and MAPK inhibitors the prolongation of PGE<sub>2</sub>-induced hyperalgesia was still attenuated ( $F_{(2,15)} = 41.6$ ,  $****p < 0.0001$ , when the combination of Src and MAPK inhibitors-treated group was compared with the vehicle-treated and cordycepin-treated groups, at the fourth hour after intrathecal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test). To verify that the reversal of priming in the central terminal of the nociceptor was permanent, intrathecal PGE<sub>2</sub> was again injected 14 d (**C**) and 28 d (**D**) after intrathecal vehicle or inhibitors. Of note, the cordycepin-treated group was not tested at the next two time points, since it did not show any attenuation in the prolongation of PGE<sub>2</sub>-induced hyperalgesia in the previous experiments. **C**, **D**, In the group of rats treated with the combination of Src and MAPK inhibitors, the prolongation of intrathecal PGE<sub>2</sub>-induced hyperalgesia was still markedly attenuated at 14 d (**C**;  $F_{(1,10)} = 56.4$ ,  $****p < 0.0001$ ) and 28 d (**D**;  $F_{(1,10)} = 14.3$ ,  $***p = 0.004$ , when the vehicle-treated and the combination of Src and MAPK inhibitors-treated groups were compared at the fourth hour after intrathecal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test) after intrathecal vehicle and inhibitors. These findings indicate that at the central terminal of the nociceptor systemic fentanyl induces Type II priming ( $n = 6$  paws/6 rats per group).

after intrathecal PGE<sub>2</sub>; two-way ANOVA multiple comparisons test; Fig. 2B). Since treatment with cordycepin did not affect the prolongation of PGE<sub>2</sub>-induced hyperalgesia in systemic fentanyl-primed rats, this group was not tested at the follow-up time points. In the group of rats treated with the combination

of SU6656 + U0126, the prolongation of PGE<sub>2</sub>-induced hyperalgesia was still inhibited at days 14 and 28 after its administration [14 d (Fig. 2C)  $F_{(1,10)} = 56.4$ ,  $p < 0.0001$ ; and 28 d (Fig. 2D)  $F_{(1,10)} = 14.3$ ,  $p = 0.0036$ , when the vehicle-treated group is compared with the combination of SU6656 + U0126-treated group



**Figure 3.** Role of IB4+ and IB4– nociceptors in priming induced at the peripheral and central nociceptor terminal by systemic fentanyl. Rats received an intrathecal injection of control saporin (3.2 μg/20 μl), SSP-saporin (100 ng/20 μl), or IB4-saporin (3.2 μg/20 μl). Fourteen days later, at which time the mechanical nociceptive threshold was not different from pre-saporin baselines [ $t_{(5)} = 0.59$ ;  $p = 0.58$ , for the saporin (control)-treated group,  $t_{(5)} = 1.3$ ;  $p = 0.24$ , for the SSP-saporin-treated group, and  $t_{(5)} = 0.47$ ;  $p = 0.66$ , for the IB4-saporin-treated group, when the mechanical nociceptive threshold is compared before and 14 d after treatments; paired Student’s  $t$  test], fentanyl (30 μg/kg) was injected subcutaneously. **A**, Four days later, at which time mechanical nociceptive threshold was not different from pre-fentanyl baseline [ $t_{(5)} = 0.25$ ;  $p = 0.81$ , for the saporin (control)-treated group,  $t_{(5)} = 0.52$ ;  $p = 0.62$ , for the SSP-saporin-treated group, and  $t_{(5)} = 0.061$ ;  $p = 0.95$ , for the IB4-saporin-treated group, when the mechanical nociceptive threshold was compared before and 4 d after systemic fentanyl; paired Student’s  $t$  test] PGE<sub>2</sub> (100 ng/5 μl) was injected intradermally and mechanical nociceptive threshold evaluated 30 min and 4 h later. Hyperalgesia was present in all saporin-treated groups measured 30 min after its injection. Also, prolonged hyperalgesia induced by intradermal PGE<sub>2</sub> was present in all groups treated with saporins ( $F_{(2,15)} = 0.69$ ,  $p = 0.52$ , when all saporin-treated groups are compared at the fourth hour after intradermal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test), indicating that in the peripheral terminal of the nociceptor systemic fentanyl does not require IB4+ and SP peptidergic nociceptors to develop priming. **B**, Seven days after systemic fentanyl and 3 d after intradermal PGE<sub>2</sub>, rats were tested for priming in the central terminal of the nociceptor. PGE<sub>2</sub> (400 ng/20 μl) was injected intrathecally and mechanical nociceptive threshold evaluated 30 min and 4 h later. Intrathecal PGE<sub>2</sub>-induced hyperalgesia in all saporin-treated groups, measured 30 min after its administration. However, the prolongation of PGE<sub>2</sub>-induced hyperalgesia at the fourth hour was markedly attenuated in the group treated with SSP-saporin, but not in the saporin-control-treated and IB4-saporin-treated groups [ $F_{(2,15)} = 16.0$ ,  $***p = 0.0002$ , when the SSP-saporin-treated group is compared with the saporin (control)-treated and the IB4-saporin-treated groups at the fourth hour after intrathecal PGE<sub>2</sub>; two-way repeated-measures ANOVA followed by Bonferroni *post hoc* test]. These findings support the suggestion that at the central terminal, systemic fentanyl requires IB4– peptidergic nociceptors to induce priming ( $n = 6$  paws/6 rats per group).

**Table 1.** Effect of PGE<sub>2</sub> on small DRG neurons of different IB4-binding classes from control animals

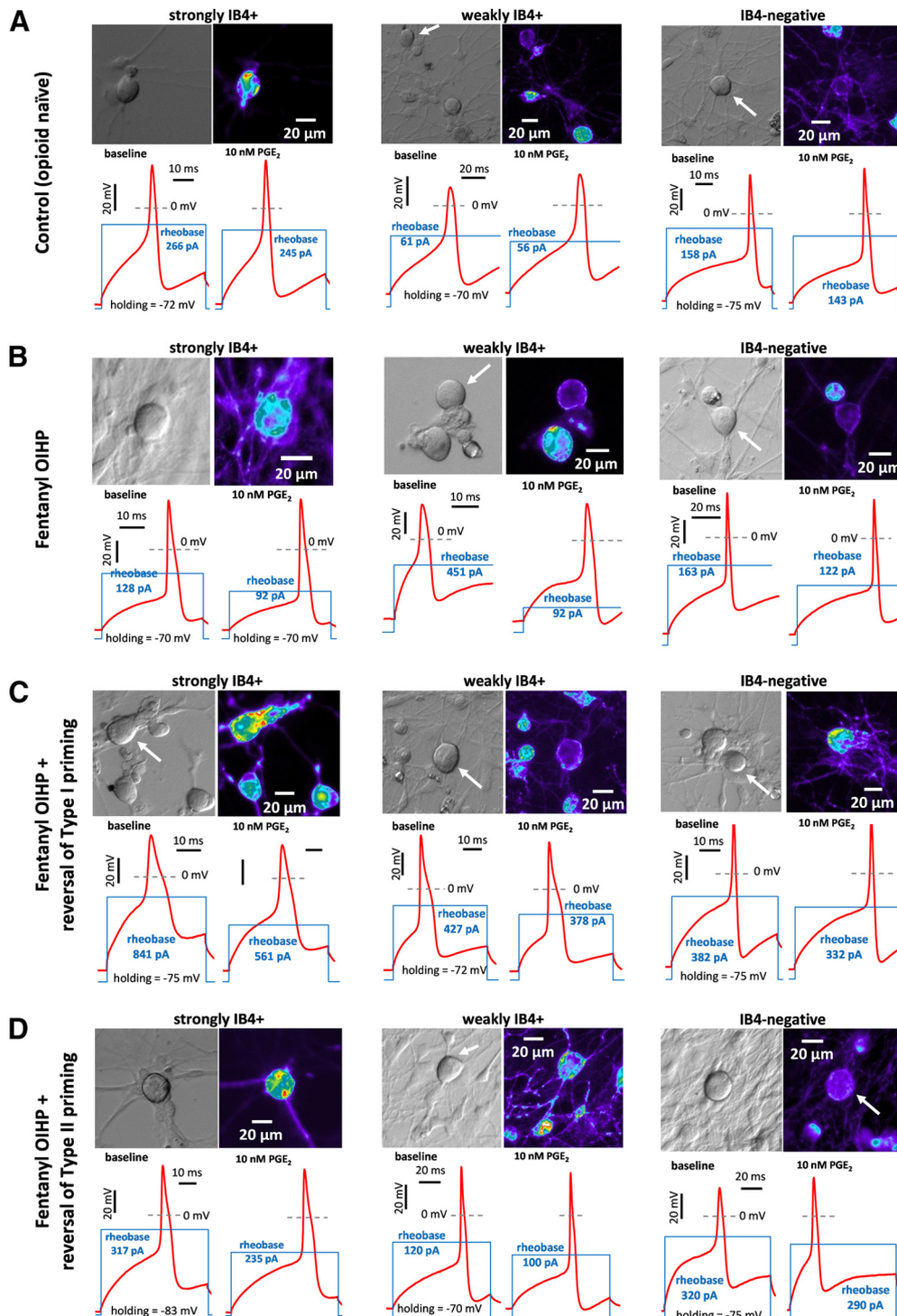
|  | Strongly IB4+ | Weakly IB4+  | IB4–         |
|--|---------------|--------------|--------------|
| Number of cells tested with PGE <sub>2</sub>                             | 10            | 14           | 9            |
| % of neurons responsive to PGE <sub>2</sub> [number of responsive cells] | 60%           | 71%          | 67%          |
|  | [6 cells]     | [10 cells]   | [6 cells]    |
| % reduction in rheobase induced by 10 nM PGE <sub>2</sub>                | 13 ± 5        | 12 ± 4       | 9 ± 4        |
| % reduction in rheobase induced by 100 nM PGE <sub>2</sub>               | 23 ± 3        | 24 ± 4       | 27 ± 7       |
| Baseline rheobase, pA  | 384 ± 56      | 313 ± 45     | 294 ± 47     |
| (including neurons not tested with PGE <sub>2</sub> )                    | ( $n = 20$ )  | ( $n = 23$ ) | ( $n = 20$ ) |

at the fourth hour after intrathecal PGE<sub>2</sub>; two-way ANOVA multiple comparisons test]. These data support the suggestion that systemic fentanyl induces Type II but not Type I priming in nociceptor central terminals, which is in accordance with our previous findings (Araldi et al., 2018c), while Type I priming only occurs at the peripheral terminal.

**Involvement of peptidergic nociceptors in priming at the central terminal**

Maintenance of Type I and Type II priming can occur in different classes of nociceptors. For example, Type I, induced by inflammatory mediators, occurs in IB4+ nonpeptidergic nociceptors (Joseph and Levine, 2010), while Type II priming, induced by intradermal DAMGO, was prevented by intrathecal administration

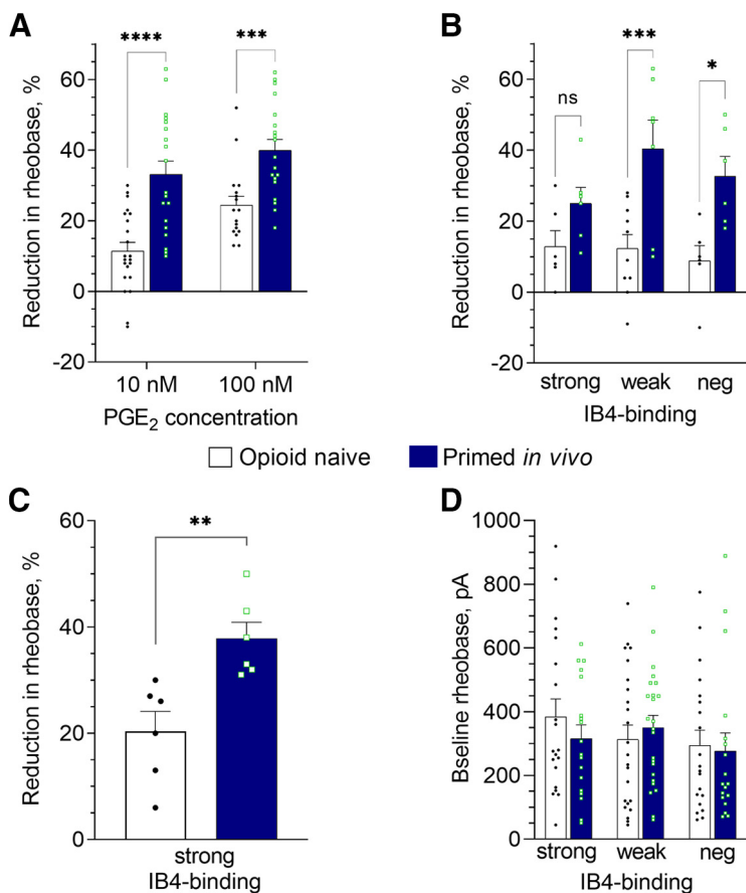
of SSP-saporin (Araldi et al., 2018b), known to disrupt signaling pathway of peptidergic (mostly IB4–) nociceptors (Seki et al., 2005). In the present experiments, we evaluated the role of these two nociceptor populations in priming induced by systemic fentanyl, in the peripheral and central nociceptor terminals. Saporin-control, IB4-saporin [destroys IB4+ (mostly nonpeptidergic) nociceptors], or SSP-saporin was injected intrathecally and, 14 d later, systemic fentanyl (30 μg/kg) was administered subcutaneously. Four days later, PGE<sub>2</sub> was injected intradermally and the mechanical nociceptive threshold evaluated 30 min and 4 h after its injection. Prolongation of PGE<sub>2</sub>-induced hyperalgesia was present in all saporin-treated groups [ $F_{(2,15)} = 0.69$ ,  $p = 0.52$ , when the control (saporin), the SSP-saporin-treated and the IB4-saporin-treated groups are compared at the fourth hour after intradermal PGE<sub>2</sub>; two-way repeated-measures



**Figure 4.** Illustration of PGE<sub>2</sub>-induced sensitization in small DRG neurons *in vitro*. Rows (A–D) represent groups of DRG neurons cultured from rats with different conditions: control (opioid naive, A), fentanyl-primed (B), and fentanyl-primed with reversal of Type I (C) and of Type II (D) priming, as defined in Figures 5–7. Strongly IB4+, weakly IB4+, and IB4– small DRG neurons, sensitized by PGE<sub>2</sub>, are depicted in each group. For each depicted neuron, its image in transmitted light (DIC contrast; upper left), fluorescent image of IB4-labeling (pseudocolor; upper right), and two electrophysiological recordings (below) are shown. The recordings, made in current clamp mode, show APs (red line) induced by current step of rheobase magnitude, before (lower left) and 10 min after (lower right) administration of PGE<sub>2</sub> (10 nM). Stimulation profile is shown as blue line, in the same scale in left and right recordings (the scale differs between neurons). Note, the reduction in rheobase after administration of PGE<sub>2</sub>, which was used to quantify magnitude of sensitization (further analyzed in Figs. 5–7), is larger in fentanyl-primed group for each IB4-binding status (B) compared with opioid naive group (A); attenuation of the sensitizing effect of PGE<sub>2</sub> was observed in both reversal groups (C, D) compared with the primed group (B), in weakly IB4+ and IB4–, but not in strongly IB4+ neurons.

ANOVA; Fig. 3A], indicating that fentanyl-induced priming in the nociceptor peripheral terminal is not dependent on SP peptidergic or IB4+ nociceptors, compatible with the role of a different class of nociceptors (Cavanaugh et al., 2011; Araldi et al., 2018c). Seven

days after systemic fentanyl and 3 d after intradermal PGE<sub>2</sub>, priming was tested in the central terminal. Rats received PGE<sub>2</sub> (400 ng) intrathecally and mechanical nociceptive threshold was evaluated 30 min and 4 h later. In the group treated with SSP-saporin, but not



**Figure 5.** Fentanyl-induced priming *in vivo* is associated with enhanced PGE<sub>2</sub>-induced sensitization of small DRG neurons, *in vitro*. Rats were primed by the systemic administration of fentanyl (30 μg/kg, s.c.) 8 d before preparing neuronal cultures, to parallel our *in vivo* protocol: 4 d before (injection of fentanyl) and 4 d after administering of vehicle or reversal agents. Patch-clamp electrophysiology recordings were made in small-diameter DRG neurons from fentanyl-primed and control (opioid naive) groups (depicted in all panels by the white and dark blue bars, correspondingly), after 24 h in culture. In **A–C**, bars show pooled magnitudes of decrease in rheobase, relative to preadministration baseline (measured before the first application of PGE<sub>2</sub>) and 5 min after application (10 and 100 nM); 10 nM PGE<sub>2</sub> was applied first, and then, 10 min later, its concentration was increased to 100 nM (cumulative concentration dependence). Symbols show individual values. Only those neurons with a change in rheobase of not <10%, for the higher or lower concentration, were considered for analysis. Some neurons were not tested with the higher concentration because of loss of the patch (but only used if they showed at least 10% reduction in rheobase for the lower concentration), therefore, regular ANOVA was used for analysis. **A**, Reduction of rheobase in response to 10 and 100 nM PGE<sub>2</sub>, analyzed regardless of IB4-binding status. In DRG neurons from primed rats the effect of both concentrations of PGE<sub>2</sub> was significantly greater than in controls [two-way ANOVA: effect of PGE<sub>2</sub> concentration,  $p = 0.0015$ ,  $F_{(1,73)} = 10.8$ ; effect of condition:  $p < 0.0001$ ,  $F_{(1,73)} = 38.2$ ; not significant (ns) interaction:  $p = 0.32$ ,  $F_{(1,73)} = 1.0$ ; Holm–Sidak’s *post hoc*  $t_{(73)} = 5.3$ , \*\*\*adjusted  $p < 0.0001$  for 10 nM;  $t_{(73)} = 3.6$ , \*\*\*adjusted  $p = 0.0007$  for 100 nM]. Number of cells in control group:  $n = 22$  for 10 nM and  $n = 17$  for 100 nM; in primed group:  $n = 19$  for both 10 and 100 nM. **B**, Reduction of rheobase in response to PGE<sub>2</sub> (10 nM), in neurons separated by IB4-binding intensity into: strongly IB4+ (“strong”), weakly IB4+ (“weak”), and IB4– classes (“neg”). Effect of priming was statistically significant (two-way ANOVA: effect of condition  $F_{(1,35)} = 23.0$ ,  $p < 0.0001$ ; effect of IB4-binding status  $F_{(2,35)} = 1.1$ ,  $p = 0.34$ ; interaction  $F_{(2,35)} = 1.2$ ,  $p = 0.33$ ). Holm–Sidak’s *post hoc* test revealed a statistically significant increase in the effect of PGE<sub>2</sub> in weakly IB4+ ( $t_{(35)} = 4.1$ , \*\*\*adjusted  $p = 0.0008$ ) and IB4– ( $t_{(35)} = 2.9$ , \*adjusted  $p = 0.011$ ) classes, indicating a leftward shift in the concentration–response curve for PGE<sub>2</sub> in these neurons. In strongly IB4+ neurons, the effect of priming was not statistically significant ( $t_{(35)} = 1.5$ , adjusted  $p = 0.14$ ). Number of cells (strong/weak/neg) in control group: 6/10/6, in primed group: 6/7/6. **C**, Reduction of rheobase in response to 100 nM PGE<sub>2</sub> in strongly IB4+ neurons from the primed group of rats was significantly enhanced compared with the control group ( $n = 6$  per group; two-tailed unpaired Student’s *t* test:  $t_{(10)} = 3.6$ , \*\* $p = 0.005$ ), indicating a different change in this neuronal population associated with *in vivo* priming. **D**, Baseline rheobase (before administration of PGE<sub>2</sub>) in three neuronal classes with different IB4-binding status. PGE<sub>2</sub>-sensitive and PGE<sub>2</sub>-insensitive neurons were included in the analysis of baseline rheobase. Differences between primed and control groups as well as between classes were not statistically significant [two-way ANOVA: effect of IB4-binding status, not significant (ns),  $p = 0.92$ ,  $F_{(2,117)} = 0.40$ ; effect of condition: ns,  $p = 0.68$ ,  $F_{(1,117)} = 0.17$ ; interaction: ns,  $p = 0.53$ ,  $F_{(2,117)} = 0.64$ ]. Number of cells (strong/weak/neg) in control group: 20/23/20, in primed group: 18/24/18.

the saporin (control)-treated and IB4-saporin-treated groups, prolongation of PGE<sub>2</sub>-induced hyperalgesia was markedly attenuated [ $F_{(2,15)} = 16.0$ ,  $p = 0.0002$ , when the saporin (control), the SSP-saporin-treated and the IB4-saporin-treated groups are compared at

the fourth hour after intrathecal PGE<sub>2</sub>; two-way repeated-measures ANOVA; Fig. 3B], indicating that priming induced by systemic fentanyl, in the central terminal of the nociceptor, is dependent on peptidergic nociceptors. These data support the suggestion that more than one population of nociceptors is involved in OIHP induced by systemic fentanyl.

#### Enhanced PGE<sub>2</sub>-induced sensitization of sensory neurons cultured from fentanyl-primed rats

To further understand the populations of nociceptors in which priming occurs, we translated our *in vivo* to an *in vitro* priming model by examining the effect of systemic fentanyl on PGE<sub>2</sub>-induced sensitization of cultured DRG neurons. Since it is not practical to hold cells clamped for 4 h, to reveal priming, we relied on another *in vivo* feature of hyperalgesic priming, a more than one order magnitude leftward shift (to lower values) for the dose dependence of PGE<sub>2</sub>-induced hyperalgesia (Parada et al., 2005). Since only ~60% of cultured DRG neurons are responsive to PGE<sub>2</sub> (Gold et al., 1996b), it would not be possible to distinguish between responsive and nonresponsive neurons by their response to a low concentration of PGE<sub>2</sub>; response to a higher concentration is required. Therefore, we chose two PGE<sub>2</sub> concentrations, 10 and 100 nM, to examine our hypothesis that priming induces a leftward shift of the dose–response curve in primed DRG neurons.

In cultures of DRG neurons derived from opioid naive animals, corresponding magnitudes of sensitization were  $11 \pm 2\%$  ( $n = 22$ ) for 10 nM PGE<sub>2</sub> and  $24 \pm 2\%$  ( $n = 17$ ) for 100 nM PGE<sub>2</sub>; 67% of small DRG neurons (soma diameter <30 μm; see Materials and Methods) examined (22 out of 33) were responsive to PGE<sub>2</sub>. Average baseline rheobase of examined neurons was  $310 \pm 41$  pA ( $n = 33$ ). In DRG neurons from control animals, no significant difference was found between mean rheobase values of small DRG neurons responsive ( $314 \pm 55$  pA) and non-responsive ( $282 \pm 81$  pA) to PGE<sub>2</sub> (unpaired *t* test,  $t_{(31)} = 0.3$ ). Considering that DRG neurons in different IB4-binding classes have different electrophysiological properties (Stucky and Lewin, 1999) and are differentially

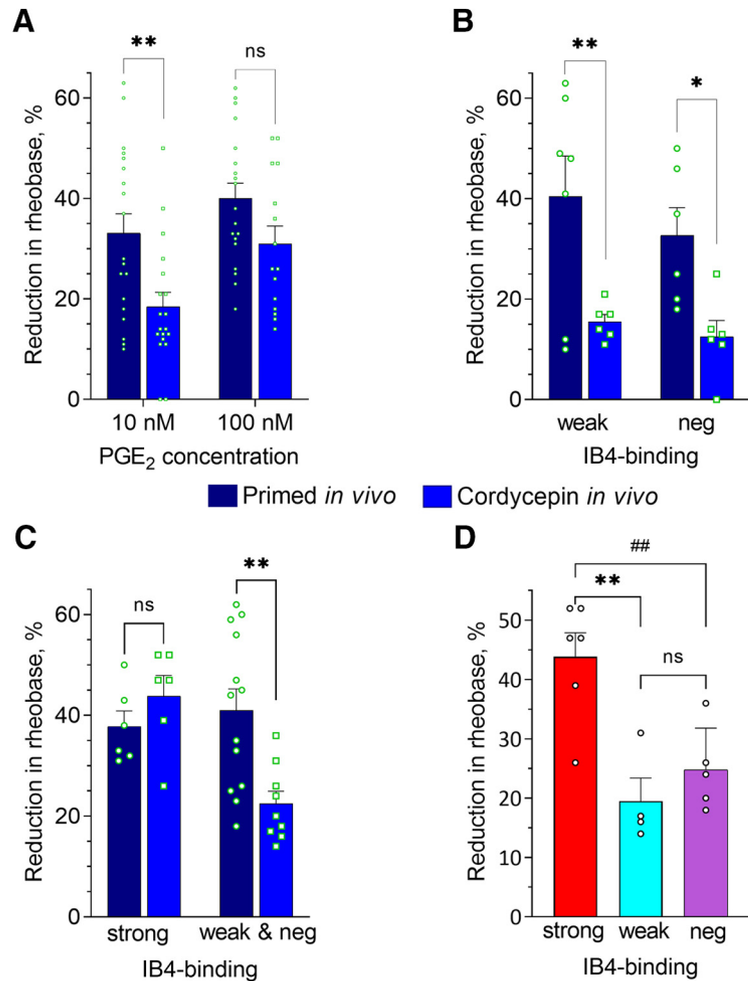
ent IB4-binding classes have different electrophysiological properties (Stucky and Lewin, 1999) and are differentially

affected by exposure to fentanyl (Khomula et al., 2019), we analyzed their response to PGE<sub>2</sub>, separately (Table 1). Fractions of neurons responsive to PGE<sub>2</sub> and magnitude of sensitization by 10 nM PGE<sub>2</sub> were similar under control conditions; ANOVA did not reveal a significant difference between baseline rheobase ( $p = 0.42$ ,  $F_{(2,60)} = 0.89$ ).

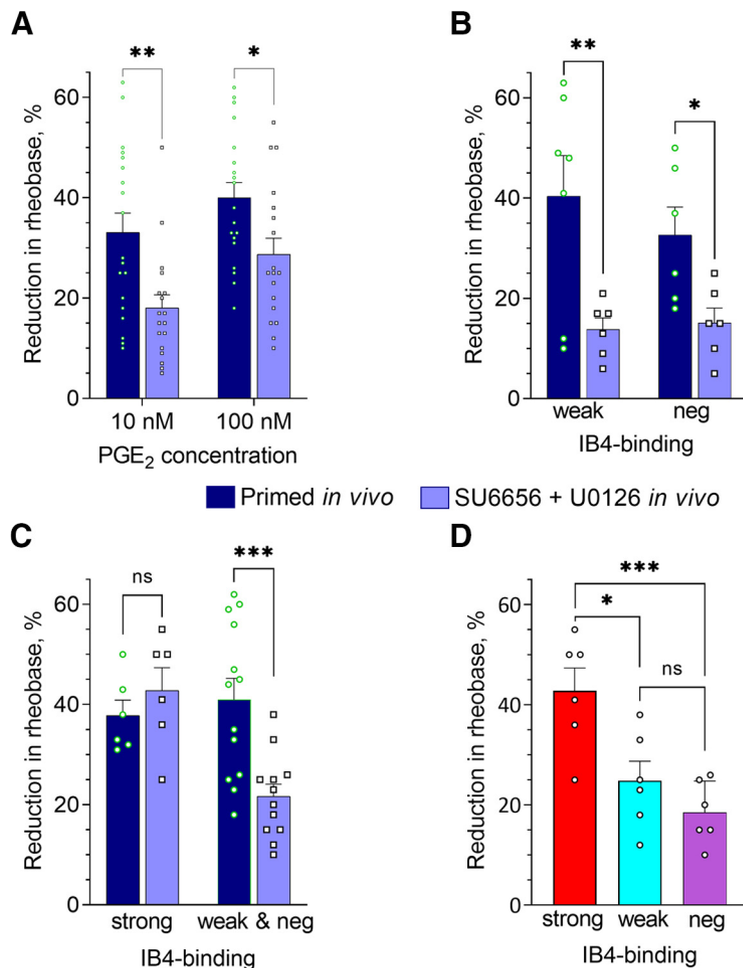
In cultured small-diameter DRG neurons derived from animals primed with fentanyl *in vivo* (primed group) reduction of rheobase induced by both concentrations of PGE<sub>2</sub> (10 and 100 nM) was significantly greater than in neurons from control animals (two-way ANOVA: effect of condition:  $p < 0.0001$ ,  $F_{(1,73)} = 38.2$ ; Holm–Sidak’s *post hoc*:  $t_{(73)} = 5.3$ , adjusted  $p < 0.0001$  for 10 nM;  $t_{(73)} = 3.6$ , adjusted  $p = 0.0007$  for 100 nM; Figs. 4A,B, 5A). And, when neurons were separated according to their IB4-binding status, and analyzed, the effect of 10 nM PGE<sub>2</sub> (Fig. 5B), our primary endpoint, a significant effect of priming was confirmed (two-way ANOVA: effect of condition  $F_{(1,35)} = 23.0$ ,  $p < 0.0001$ ). Holm–Sidak’s *post hoc* test revealed a significant increase in the effect of PGE<sub>2</sub> in weakly IB4+ ( $t_{(35)} = 4.1$ , adjusted  $p = 0.0008$ ) and IB4- ( $t_{(35)} = 2.9$ , adjusted  $p = 0.011$ ) classes. In strongly IB4+ neurons, the difference between primed and control groups was not statistically significant for 10 nM PGE<sub>2</sub> ( $t_{(35)} = 1.5$ , adjusted  $p = 0.14$ ). However, for these neurons we found a significant increase in sensitization by the higher concentration (100 nM) of PGE<sub>2</sub> (two-tailed unpaired Student’s *t* test:  $t_{(10)} = 3.6$ ,  $p = 0.005$ ; Fig. 5C).

Of note, we did not see a significant effect of priming on rheobase of neurons of any IB4-binding status [two-way ANOVA: effect of IB4-binding status, not significant (ns),  $p = 0.92$ ,  $F_{(2,117)} = 0.40$ ; effect of condition: ns,  $p = 0.68$ ,  $F_{(1,117)} = 0.17$ ; interaction: ns,  $p = 0.53$ ,  $F_{(2,117)} = 0.64$ ; Fig. 5D], or when the three classes were combined (control vs priming, two-tailed unpaired Student’s *t* test:  $t_{(122)} = 0.34$ ,  $p = 0.73$ ). This is in agreement with the results of our current (Fig. 1) and previous (Ferrari et al., 2013; Hendrich et al., 2013; Araldi et al., 2015, 2018c; Khomula et al., 2019) behavioral studies, which demonstrated no effect of priming on baseline mechanical nociceptive threshold.

These findings confirm our initial hypothesis that priming-like neuroplasticity could be observed *in vitro*, in nociceptors cultured from primed rats, and identify nociceptor populations in which this occurs. While all three nociceptor populations were affected by fentanyl, leftward shift in dose dependence of PGE<sub>2</sub>-induced sensitization was only observed in weakly IB4+ and IB4-



**Figure 6.** Effect of *in vivo* treatment of fentanyl-primed rats with the reversal agent for Type I priming on PGE<sub>2</sub>-induced sensitization of small DRG neurons. Rats were primed by the systemic administration of fentanyl (30 μg/kg, s.c.) 8 d before preparing neuronal cultures (4 d before intrathecal cordycepin, followed by an additional 4 d before culture preparation). Recordings were made in small-diameter DRG neurons from fentanyl-primed animals either treated (reversal group) or not treated (primed group) with cordycepin (depicted in A–C by the blue and dark blue bars, correspondingly) after 24 h in culture. In A–D, bars show pooled magnitudes of decrease in rheobase, relative to baseline, after PGE<sub>2</sub> application (10 and 100 nM), measured and analyzed in the same way as described in Figure 5 and Materials and Methods. Symbols show individual values. In A–C, values for primed group were repeated from Figure 5A–C for the purpose of comparison. **A**, Reduction of rheobase in response to 10 and 100 nM PGE<sub>2</sub>, analyzed regardless of IB4-binding status. In neurons from primed rats the effect of 10 nM but not 100 nM PGE<sub>2</sub> was significantly greater than in the reversal group [two-way ANOVA; effect of condition:  $F_{(1,68)} = 12.5$ ,  $p = 0.0007$ ; Holm–Sidak’s *post hoc*:  $t_{(68)} = 3.2$ , \*\*adjusted  $p = 0.004$  for 10 nM;  $t_{(68)} = 1.8$ , adjusted  $p = 0.07$ , not significant (ns), for 100 nM]. Number of cells in primed group:  $n = 19$  for both 10 and 100 nM; in reversal group:  $n = 19$  for 10 nM and  $n = 15$  for 100 nM. **B**, Reduction of rheobase in response to 10 nM PGE<sub>2</sub> in weakly IB4+ (“weak”) and IB4- (“neg”) neurons. Attenuation of PGE<sub>2</sub>-induced sensitization after *in vivo* cordycepin was statistically significant in both neuronal populations (two-way ANOVA: effect of condition  $F_{(1,21)} = 16.3$ ,  $p = 0.0006$ ; Holm–Sidak’s *post hoc*:  $t_{(21)} = 3.2$ , \*\*adjusted  $p = 0.008$  for weakly IB4+;  $t_{(21)} = 2.5$ , \*adjusted  $p = 0.02$  for IB4-). Number of cells (weak/neg) in reversal group: 6/6, in primed group: 7/6. **C**, Reduction of rheobase in response to 100 nM PGE<sub>2</sub> in strongly IB4+ (“strong”) and merged weakly IB4+ and IB4- (“weak and neg”) neurons from primed and reversal groups. Two-way ANOVA revealed statistically significant interaction ( $F_{(1,30)} = 8.5$ ,  $p = 0.007$ ), indicating differential effects on the three different neuronal classes. Indeed, statistically significant attenuation in reversal compared with primed group occurred in weakly IB4+ and IB4- but not in strongly IB4+ neurons (Holm–Sidak’s *post hoc*:  $t_{(30)} = 0.9$ , adjusted  $p = 0.38$  for strong, not significant (ns);  $t_{(30)} = 3.7$ , \*\*adjusted  $p = 0.002$  for weak and neg). Number of cells (strong/weak and neg): 6/13 in primed group, 6/9 in reversal group. **D**, Reduction of rheobase in response to 100 nM PGE<sub>2</sub> in strongly IB4+ (strong), weakly IB4+ (weak), and IB4- (neg) neurons from reversal group. Values in strongly IB4+ class were significantly greater than in weakly IB4+ and IB4- neurons [one-way ANOVA:  $F_{(2,12)} = 11.8$ , \*\* $p = 0.002$ ; Tukey’s *post hoc*:  $q_{(12)} = 6.3$ , adjusted  $p = 0.002$  for strongly IB4+ vs weakly IB4+;  $q_{(12)} = 5.2$ , ##adjusted  $p = 0.008$  for strongly IB4+ vs IB4-;  $q_{(12)} = 1.3$ , adjusted  $p = 0.63$ , not significant (ns), for weakly IB4+ vs IB4-]. Number of cells: 6 strong, 4 weak, 5 neg.



**Figure 7.** Effect of *in vivo* treatment of fentanyl-primed rats with reversal agents for Type II priming on PGE<sub>2</sub>-induced sensitization of small DRG neurons. Rats were primed by the systemic administration of fentanyl (30 μg/kg, s.c.) 8 d before preparing neuronal cultures (4 d before reversal agents, followed by 4 d before culture preparation). Recordings were made in small-diameter DRG neurons from fentanyl-primed animals treated intrathecally with the combination of a Src and MAPK inhibitor (SU6656 + U0126; Type II reversal group; light blue bars) and not treated (primed group; dark blue bars). In **A–D**, bars show pooled magnitudes of decrease in rheobase after PGE<sub>2</sub> application (10 and/or 100 nM), relative to baseline (measured before the first application), measured and analyzed in the same way as described in Figure 6, while symbols show individual values. In **A–C**, values for primed group were repeated from Figure 5A–C for the purpose of comparison. **A**, Reduction of rheobase in response to 10 and 100 nM PGE<sub>2</sub>, analyzed regardless of IB4-binding status. In the primed group the effect of 10 and 100 nM PGE<sub>2</sub> was significantly greater than in the Type II reversal group (two-way ANOVA; effect of condition:  $F_{(1,70)} = 16.6$ ,  $p = 0.0001$ ; Holm–Sidak’s *post hoc*:  $t_{(70)} = 3.3$ , \*\*adjusted  $p = 0.003$  for 10 nM;  $t_{(70)} = 2.5$ , \*adjusted  $p = 0.02$  for 100 nM). Number of cells in primed group:  $n = 19$  for both 10 and 100 nM; in Type II reversal group:  $n = 18$  for both 10 and 100 nM. **B**, Reduction of rheobase in response to 10 nM PGE<sub>2</sub> in weakly IB4+ (“weak”) and IB4– classes (“neg”) neurons. Attenuation of PGE<sub>2</sub>-induced sensitization after *in vivo* administration of the combination of a Src and MAPK inhibitor was statistically significant in both neuronal subpopulations (two-way ANOVA: effect of condition  $F_{(1,21)} = 15.5$ ,  $p = 0.0008$ ; Holm–Sidak’s *post hoc*:  $t_{(21)} = 3.4$ , \*\*adjusted  $p = 0.005$  for weakly IB4+;  $t_{(21)} = 2.2$ , \*adjusted  $p = 0.04$  for IB4–). Number of cells (weak/neg) in Type II reversal group: 6/6, in primed group: 7/6. **C**, Reduction of rheobase in response to 100 nM PGE<sub>2</sub> in strongly IB4+ (“strong”) and merged weakly IB4+ and IB4– (“weak and neg”) neurons from primed and reversal groups. Two-way ANOVA revealed statistically significant interaction ( $F_{(1,33)} = 8.8$ ,  $p = 0.006$ ), indicating differential effects on different neuronal classes. Indeed, statistically significant attenuation in reversal group compared with the primed group occurred in weakly IB4+ and IB4– but not in strongly IB4+ neurons [Holm–Sidak’s *post hoc*:  $t_{(33)} = 0.74$ , adjusted  $p = 0.46$  for strong, not significant (ns);  $t_{(33)} = 4.1$ , \*\*\*adjusted  $p = 0.0005$  for weak and neg]. Number of cells (strong/weak and neg): 6/13 in primed group, 6/12 in Type II reversal group. **D**, Reduction of rheobase in response to 100 nM PGE<sub>2</sub> in strongly IB4+ (strong), weakly IB4+ (weak), and IB4– (neg) neurons from reversal group. Reduction of rheobase in the strongly IB4+ class were significantly greater than in weakly IB4+ and IB4– neurons [one-way ANOVA:  $F_{(2,15)} = 11.2$ ,  $p = 0.0010$ ; Tukey’s *post hoc*:  $q_{(15)} = 4.8$ , \*adjusted  $p = 0.011$  for strongly IB4+ vs weakly IB4+;  $q_{(15)} = 6.5$ , \*\*\*adjusted  $p = 0.0010$  for strongly IB4+ vs IB4–;  $q_{(15)} = 1.7$ , adjusted  $p = 0.48$ , not significant (ns), for weakly IB4+ vs IB4–]. Number of cells: 6 strong, 6 weak, 6 neg.

populations of putative nociceptors. However, enhanced sensitization by the high dose of PGE<sub>2</sub> was found in strongly IB4+ nociceptors. Together, our behavioral and *in vitro* findings support the suggestion that fentanyl-induced priming develops in the nociceptor.

### Intrathecal cordycepin attenuates priming-induced enhancement of PGE<sub>2</sub> sensitization of nociceptors, *in vitro*

Using our *in vitro* model of fentanyl-induced priming, we determined whether administration of cordycepin, which partially reversed priming *in vivo*, has a similar effect on priming-like neuroplasticity in cultured DRG neurons. We prepared cultures of DRG neurons derived from animals that were primed *in vivo* with fentanyl and 4 d later received intrathecal cordycepin (cordycepin group), used in our behavior experiments to attenuate Type I priming (Fig. 1). DRG neuron cultures were prepared 4 d after administering cordycepin, a time at which attenuation of priming was observed in behavioral experiments (Fig. 1). Small-diameter DRG neurons cultured from the cordycepin-treated group received PGE<sub>2</sub> (10 nM) and then, 10 min later, PGE<sub>2</sub> 100 nM, and magnitude of the reduction in rheobase induced was compared with that in DRG neurons from primed rats (Figs. 4B,C, 6).

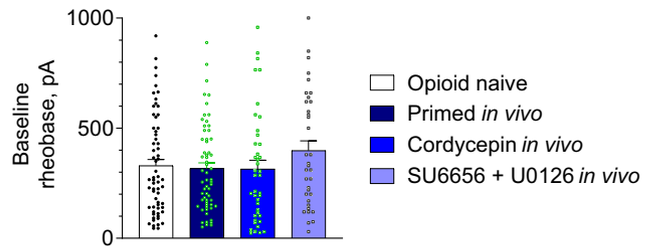
First, we compared the effects of PGE<sub>2</sub>, 10 and 100 nM, on small DRG neurons, regardless of IB4-binding status (Fig. 6A). We found a significant attenuation of the sensitizing effect of PGE<sub>2</sub> in the cordycepin-treated group compared with the primed group, for 10 nM but not for 100 nM PGE<sub>2</sub> (two-way ANOVA; effect of condition:  $F_{(1,68)} = 12.5$ ,  $p = 0.0007$ ; Holm–Sidak’s *post hoc*:  $t_{(68)} = 3.2$ , adjusted  $p = 0.004$  for 10 nM;  $t_{(68)} = 1.8$ , adjusted  $p = 0.07$ , for 100 nM). This finding supports the suggestion that *in vivo* administration of cordycepin attenuates priming in nociceptors, *in vitro*, similar to our *in vivo* observation.

To assess possible IB4-binding status-specific changes induced by *in vivo* administration of cordycepin, we took into account our finding of an enhanced effect of 10 nM PGE<sub>2</sub> on weakly IB4+ and IB4– populations, and the enhanced effect of only 100 nM PGE<sub>2</sub> on strongly IB4+ neurons. We found significant attenuation of the sensitizing effect of 10 nM PGE<sub>2</sub> in weakly IB4+ and IB4– neurons after cordycepin was administered *in vivo* to primed animals (two-way ANOVA; effect of condition:  $F_{(1,21)} = 16.3$ ,  $p = 0.0006$ ; Holm–Sidak’s *post hoc*:  $t_{(21)} = 3.2$ , adjusted  $p = 0.008$  for weakly IB4+;  $t_{(21)} = 2.5$ , adjusted  $p = 0.02$  for IB4–; Fig. 6B). In contrast, the enhanced sensitizing effect of 100 nM PGE<sub>2</sub> in

strongly IB4+ neurons was not significantly different between neurons from primed and cordycepin-treated primed rats (two-way ANOVA: statistically significant effect of interaction:  $F_{(1,30)} = 8.5$ ,  $p = 0.007$ ; Holm–Sidak's *post hoc*:  $t_{(30)} = 0.9$ , adjusted  $p = 0.38$  for strongly IB4+;  $t_{(30)} = 3.7$ , adjusted  $p = 0.002$  for weakly IB4+ and IB4– combined; Fig. 6C). In strongly IB4+ neurons, from cordycepin treated rats, the sensitizing effect of 100 nM PGE<sub>2</sub> was significantly greater than in weakly IB4+ and IB4– neurons (one-way ANOVA:  $F_{(2,12)} = 11.8$ ,  $p = 0.002$ ; Tukey's *post hoc*:  $q_{(12)} = 6.3$ , adjusted  $p = 0.002$  for strongly IB4+ vs weakly IB4+;  $q_{(12)} = 5.2$ , adjusted  $p = 0.008$  for strongly IB4+ vs IB4–;  $q_{(12)} = 1.3$ , adjusted  $p = 0.63$  for weakly IB4+ vs IB4–; Fig. 6D). These results support the suggestion that maintenance mechanisms of enhanced sensitization by PGE<sub>2</sub> are dependent on Type I priming in weakly IB4+ and IB4– nociceptor populations.

### Intrathecal Src plus MAPK inhibitors attenuates priming-induced enhancement of PGE<sub>2</sub> sensitization of nociceptors, *in vitro*

Using our *in vitro* model of fentanyl-induced priming, we next determined whether *in vivo* administration of agents that reverse Type II priming (Fig. 1) also attenuates enhanced PGE<sub>2</sub>-induced sensitization in DRG neurons cultured from fentanyl-treated animals, and which nociceptor populations were affected. Animals were primed with fentanyl, *in vivo*, and 4 d later received the combination of Src and MAPK inhibitors (intrathecally; Fig. 1A). DRGs were harvested for culturing 4 d later. Reduction of rheobase induced by 10 and 100 nM PGE<sub>2</sub> in small-diameter DRG neurons from these cultures were compared with corresponding values in cultures from the primed group of animals (Figs. 4B,D, 7). We found a significant attenuation of the sensitizing effect of PGE<sub>2</sub> in this reversal group compared with the primed group, when neurons were analyzed regardless of IB4-binding status (two-way ANOVA; effect of condition:  $F_{(1,70)} = 16.6$ ,  $p = 0.0001$ ; Holm–Sidak's *post hoc*:  $t_{(70)} = 3.3$ , adjusted  $p = 0.003$  for 10 nM;  $t_{(70)} = 2.5$ , adjusted  $p = 0.02$  for 100 nM; Fig. 7A). When IB4-binding status was considered, we found significant attenuation of the sensitizing effect of PGE<sub>2</sub> (10 nM), in weakly IB4+ and IB4– neurons (two-way ANOVA; effect of condition:  $F_{(1,21)} = 15.5$ ,  $p = 0.0008$ ; Holm–Sidak's *post hoc*:  $t_{(21)} = 3.4$ , adjusted  $p = 0.005$  for weakly IB4+;  $t_{(21)} = 2.2$ , adjusted  $p = 0.04$  for IB4–; Fig. 7B). The sensitizing effect of PGE<sub>2</sub> (100 nM) in strongly IB4+ neurons was not significantly different between reversal and primed groups (two-way ANOVA; statistically significant effect of interaction:  $F_{(1,33)} = 8.8$ ,  $p = 0.006$ ; Holm–Sidak's *post hoc*:  $t_{(33)} = 0.74$ , adjusted  $p = 0.46$  for strongly IB4+;  $t_{(33)} = 4.1$ , adjusted  $p = 0.0005$  for weakly IB4+ and IB4– combined; Fig. 7C). However, the sensitizing effect of 100 nM PGE<sub>2</sub> in strongly IB4+ neurons was significantly greater than in weakly IB4+ and IB4– neurons (one-way ANOVA:  $F_{(2,15)} = 11.2$ ,  $p = 0.0010$ ; Tukey's *post hoc*:  $q_{(15)} = 4.8$ , adjusted  $p = 0.011$  for strongly IB4+ vs weakly IB4+;  $q_{(15)} = 6.5$ , adjusted  $p = 0.0010$  for strongly IB4+ vs IB4–;  $q_{(15)} = 1.7$ , adjusted  $p = 0.48$  for weakly IB4+ vs IB4–; Fig. 7D). These results, along with the results of the previous section (Fig. 6), confirm a significant functional difference between strongly IB4+ nociceptors versus weakly IB4+ and IB4– nociceptors in how they were affected by systemic fentanyl, *in vivo*. While in both reversal groups, PGE<sub>2</sub>-induced sensitization in strongly IB4+ neurons remained unattenuated, compared with the primed group, in weakly IB4+ and IB4– neurons, it was significantly attenuated. Overall, the *in vitro* effect of combined *in vivo* inhibition of Src and MAPK on all three classes was



**Figure 8.** Priming by fentanyl *in vivo* as well as *in vivo* reversal of priming does not alter baseline rheobase of small DRG neurons *in vitro*. Bars show pooled baseline rheobase (before administration of PGE<sub>2</sub>) in the whole neuronal population (regardless of IB4-binding status) in control, primed, and two reversal groups, as defined in Figures 5–7. Symbols show individual values. PGE<sub>2</sub>-sensitive and PGE<sub>2</sub>-insensitive neurons were included in the analysis. Difference between groups was not statistically significant (one-way ANOVA:  $p = 0.26$ ,  $F_{(3,196)} = 1.33$ ). Number of cells: 63 in control, 61 in primed, 41 in Type I, and 35 in Type II reversal groups.

qualitatively similar to that found for *in vivo* cordycepin (Fig. 6). Of note, enhanced sensitization by PGE<sub>2</sub> in both weakly IB4+ and IB4– nociceptor populations shared maintenance mechanisms with both Type I and Type II priming.

Our analysis did not reveal a significant difference in baseline rheobase between DRG neurons from control, primed groups of rats, and groups of primed rats treated with reversal agents for Type I and Type II priming, correspondingly (one-way ANOVA:  $p = 0.26$ ,  $F_{(3,196)} = 1.33$ ; Fig. 8). Thus, baseline excitability of nociceptors is not affected by priming or by reversal of priming, in agreement with our current (Fig. 1) and previous behavioral studies (Aley et al., 2000; Parada et al., 2005; Reichling and Levine, 2009; Ferrari et al., 2013; Hendrich et al., 2013; Araldi et al., 2015, 2018c, 2019; Khomula et al., 2017), where baseline mechanical nociceptive threshold remained unchanged.

In summary, we have demonstrated, *in vivo*, the existence of long-lasting nociceptor neuroplasticity, induced by systemic administration of an analgesic dose of fentanyl. This neuroplasticity, which involves changes in nociceptor function, is responsible for both hyperalgesic priming and, as previously shown, for OIH (Khomula et al., 2017). Here, for the first time, we have created an *in vitro* model for studying OIHP. Using this model, we provide evidence for specific changes in electrophysiological properties of nociceptors, in their response to PGE<sub>2</sub>, which correlates with the presence of hyperalgesic priming *in vivo* and reversal (partial permanent attenuation) by cordycepin or the combination of Src and MAPK inhibitors. The combination of behavioral and *in vitro* cellular approaches allowed us to provide evidence of different roles for classes of nociceptors in hyperalgesic priming induced *in vivo* by systemic fentanyl. Additionally, our *in vitro* results identified a novel form of neuroplasticity in strongly IB4+ nociceptors, characterized by increased change in rheobase induced by the high but not low concentration of PGE<sub>2</sub>, which is not reversed by either cordycepin or the combination of Src and MAPK inhibitors. Studies are ongoing to elucidate the *in vivo* correlate of this novel form of nociceptor neuroplasticity. We suggest that the residual PGE<sub>2</sub>-induced hyperalgesia (~50%), which was still present at the peripheral nociceptor terminal at the fourth hour, in rats treated with inhibitors of both Type I and Type II priming (Fig. 1), is mediated by neuroplasticity in strongly IB4+ nociceptors.

## Discussion

Opioid analgesics, especially of the fentanyl class, can produce a hypersensitivity to noxious stimuli (OIH) and long-lasting pain

exacerbation (chronification; Zyllicz and Twycross, 2008; Silverman, 2009; Yi and Pryzbylowski, 2015). Since the action of MOR agonists on nociceptors plays an important role in their pronociceptive effects (Scherrer et al., 2009; Corder et al., 2017; Streicher and Bilsky, 2017), we focused on cellular mechanisms of OIHP, using *in vitro* patch-clamp electrophysiology.

Three forms of OIHP have been reported. Type I is reversed by cordycepin (Ferrari et al., 2013, 2019; Araldi et al., 2018c), Type II by the combination of a Src and MAPK inhibitor (Araldi et al., 2017a, 2018c), and a third type, produced by biased MOR agonists, is neither Type I nor Type II (Araldi et al., 2018a). In the case of fentanyl, different nociceptor populations and distinct intracellular mechanisms are involved in OIHP induced in the setting of different routes of administration and doses (i.e., subanalgesic and analgesic; Araldi et al., 2018c). Our present *in vivo* findings show that an analgesic dose of fentanyl induces priming in the peripheral terminal of the nociceptor, sharing maintenance mechanisms with both Type I and Type II priming.

Consistent with our *in vivo* findings, we observed enhanced PGE<sub>2</sub>-induced sensitization (reduction of rheobase) in small-diameter DRG neurons cultured from fentanyl-primed rats, confirming our hypothesis that priming-like neuroplasticity can be observed *in vitro*. Enhanced sensitization by the lower concentration of PGE<sub>2</sub>, which was our primary endpoint, was only observed in weakly IB4+ and IB4- neurons. In contrast, in strongly IB4+ neurons we found a significant increase in sensitization by the higher but not the lower concentration of PGE<sub>2</sub>. Of note, we did not observe an effect of priming on pre-PGE<sub>2</sub> rheobase, paralleling the lack of change in baseline nociceptive threshold in primed animals, in behavioral studies (Parada et al., 2005; Araldi et al., 2018c; Ferrari et al., 2019). Our finding of priming-like neuroplasticity in cultured DRG neurons, which have lost their terminals in peripheral tissues and spinal dorsal horn, during culture preparation, support the suggestion that the priming state in nociceptors also exists in the soma.

A significant attenuation of the enhanced sensitizing effect of the low concentration (10 nM) of PGE<sub>2</sub> was observed in weakly IB4+ and IB4- DRG neurons (both peptidergic) cultured from rats whose fentanyl-induced priming was reversed *in vivo* by agents that reverse Type I (cordycepin) or Type II (the combination of a Src and a MAPK inhibitors) priming. These findings support the suggestion that both Type I and Type II priming develop, in the same or different nociceptor populations, identified *in vitro* as weakly IB4+ and IB4- (peptidergic) nociceptors. However, underlying intracellular mechanisms, allowing coexistence of both Type I and Type II priming in the same neuron, still need to be investigated. One might suggest a convergent signaling event, downstream, affected by Type I and Type II reversal agents. For instance, protein translation in DRG neurons requires ERK signaling, and Src activation and PKA/PKC activation may also lead to ERK activation (Hu and Gereau, 2003; Kawasaki et al., 2004; Uttam et al., 2018), supporting the suggestion that Type II reversal agents might also inhibit protein translation (required for Type I priming) downstream of ERK signaling. However, Type II priming at the central terminal was not sensitive to cordycepin (Fig. 2; Araldi et al., 2018c). Similarly, Type I priming is insensitive to agents that reverse Type II priming (Araldi et al., 2018c; Ferrari et al., 2019). Together, these findings support the suggestion that there is no overlap between mechanisms of Type I and Type II priming.

Alternatively, it is possible that different neuronal subpopulations, within weakly IB4+ and IB4- peptidergic nociceptors

mediate Type I and Type II priming. In this case, however, one would expect an additive attenuating effect of reversal agents of Type I and Type II priming *in vivo* (as they would act on different neurons), which we did not observe (Fig. 1). The prominent attenuating effect of SSP-saporin on fentanyl-induced priming, only at the nociceptor central terminal (Fig. 3), indicates that different populations of nociceptors mediate priming, and in particular Type II priming, at the central and peripheral terminal. The fact that OIHP, in the nociceptor peripheral terminal, was not affected by either IB4-saporin or SSP-saporin treatment could be attributed to the involvement of a different class of nociceptors, negative for both nonpeptidergic and peptidergic markers, as suggested in our previous study in which intrathecal fentanyl-induced priming in the nociceptor peripheral terminal was not prevented by treatment with IB4-saporin or SSP-saporin (Araldi et al., 2018c). Of note, nonpeptidergic IB4- neurons account for ~20% of TRPV1<sup>+</sup> neurons, as shown in mice (Cavanaugh et al., 2011).

The enhanced sensitizing effect of the higher concentration of PGE<sub>2</sub>, in strongly IB4+ (non-peptidergic) neurons was not attenuated by drugs that reverse Type I or Type II priming, identifying a novel form of neuroplasticity in these nociceptors. It is possible that neuroplasticity in this class of nociceptors contributes to the remaining PGE<sub>2</sub>-induced hyperalgesia, at the fourth hour, in rats treated with a combination of drugs that reverse both Type I and Type II priming that did not attenuate priming any more than inhibition of either Type I or Type II priming alone.

Although we use only mechanical stimulation to reveal enhanced and prolonged hyperalgesia, we do not expect that priming has selectivity to stimulus modality. Indeed, our *in vitro* findings suggest that priming is associated with enhanced reduction of threshold for electrical excitability in response to PGE<sub>2</sub>. Thus, any depolarizing stimulus capable of activating a primed nociceptor would require lower stimulus magnitude to trigger AP generation after sensitization.

Observed changes in rheobase implicate changes in ion channel function. It is very likely that the function of channels mediating the effects of PGE<sub>2</sub> are affected, including voltage-gated sodium, potassium and calcium channels (Gold et al., 1996a, 1998; Kopp and Cicha, 1999; Gu et al., 2008; Fan et al., 2011; for review, see Kawabata, 2011). We do not, however, think that priming directly affects properties of ion channels involved in AP generation, rather it alters the PGE<sub>2</sub> signaling pathway (Aley et al., 2000; Khasar et al., 2008; for review, see Reichling and Levine, 2009; Reichling et al., 2013).

Once sensitized, any part of a primed nociceptor may contribute to a prolongation of hyperalgesia and in this way contribute to pain chronification. For instance, tissue damage, like a surgical incision or wound, is associated with production of sensitizing agents, resulting in hyperalgesia. In the case of priming, when nociceptors remain sensitized for a much longer time after administration of sensitizing agents, pain would be expected to be prolonged as well. Increased and prolonged nociceptor activity in the presence of a stimulus that leads to activation of the nociceptor likely will facilitate central sensitization. Of note, since priming by itself does not alter pain sensitivity, it is unlikely to lead to central sensitization.

Along with central sensitization, hyperalgesic priming is recognized as an integral part of the “pain memory” paradigm (Ji et al., 2003; Price and Inyang, 2015) and serves to illuminate a critical role of the nociceptor in pain chronification. Studies have

demonstrated the progressive nature of chronic pain (Price and Inyang, 2015), as well as OIH, which has been tightly linked to OIHP in our studies (Khomula et al., 2019). Use of opioids for analgesia in a clinical setting, from the perspective of our OIHP model may be viewed as the triggering event for induction of a pain memory, thus providing important clinical parallels that can help explain mechanisms and suggest potential pathways to resolution of neuroplasticity that drives chronicity.

In conclusion, our *in vitro* results indicate that fentanyl induces neuroplasticity in nociceptors of all three IB4-binding categories; neuroplasticity in weakly IB4+ and IB4- (peptidergic) nociceptors likely contributes to Type I and Type II OIHP *in vivo*, which differs from our previous finding for OIH induced by fentanyl, where only a role of weakly IB4+ nociceptors was observed (Khomula et al., 2019). Of note, strongly IB4+ nociceptors, which develop a novel type of neuroplasticity in fentanyl-primed animals, also revealed robust fentanyl-induced sensitization in cultures from opioid naive animals, which does not account for fentanyl induced OIH, since it did not become greater in cultures from primed animals. In contrast, weakly IB4+ and IB4- nociceptors were sensitized by fentanyl in cultures from *in vivo* primed animals, sensitization that was reversed by cordycepin, but only in weakly IB4+ neurons (Khomula et al., 2019). On the other hand, their enhanced PGE<sub>2</sub>-induced sensitization, attributed to priming, was attenuated by cordycepin and the combination of Src and MAPK inhibitors, in both peptidergic classes. Together, these findings indicate that fentanyl produces different effects in peptidergic and non-peptidergic neuronal classes; they also reveal differences between neuroplasticity in weakly IB4+ and IB4- nociceptors, dissected by the effect of intrathecal cordycepin on their sensitization by fentanyl, *in vitro*.

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