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### Authors

Guo, Yuan  
Zhang, Zhiyong  
Wu, Hsiang-en  
et al.

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## Increased thrombospondin-4 after nerve injury mediates disruption of intracellular calcium signaling in primary sensory neurons

Yuan Guo<sup>1</sup>, Zhiyong Zhang<sup>1</sup>, Hsiang-en Wu<sup>1</sup>, Z. David Luo<sup>2,3</sup>, Quinn H. Hogan<sup>1</sup>, and Bin Pan<sup>1</sup>

<sup>1</sup>Department of Anesthesiology, Medical College of Wisconsin, 8701 Watertown Plank Road, Milwaukee, WI 53226

<sup>2</sup>Department of Anesthesiology & Perioperative Care, University of California Irvine, Irvine, CA 92697

<sup>3</sup>Department of Pharmacology, University of California Irvine, Irvine, CA 92697

### Abstract

Painful nerve injury disrupts Ca<sup>2+</sup> signaling in primary sensory neurons by elevating plasma membrane Ca<sup>2+</sup>-ATPase (PMCA) function and depressing sarco-endoplasmic reticulum Ca<sup>2+</sup>-ATPase (SERCA) function, which decreases endoplasmic reticulum (ER) Ca<sup>2+</sup> stores and stimulates store-operated Ca<sup>2+</sup> entry (SOCE). The extracellular matrix glycoprotein thrombospondin-4 (TSP4), which is increased after painful nerve injury, decreases Ca<sup>2+</sup> current (I<sub>Ca</sub>) through high-voltage-activated Ca<sup>2+</sup> channels and increases I<sub>Ca</sub> through low-voltage-activated Ca<sup>2+</sup> channels in dorsal root ganglion neurons, which are events similar to the effect of nerve injury. We therefore examined whether TSP4 plays a critical role in injury-induced disruption of intracellular Ca<sup>2+</sup> signaling. We found that TSP4 increases PMCA activity, inhibits SERCA, depletes ER Ca<sup>2+</sup> stores, and enhances store-operated Ca<sup>2+</sup> influx. Injury-induced changes of SERCA and PMCA function are attenuated in TSP4 knock-out mice. Effects of TSP4 on intracellular Ca<sup>2+</sup> signaling are attenuated in voltage-gated Ca<sup>2+</sup> channel  $\alpha_2\delta_1$  subunit (Ca<sub>v</sub> $\alpha_2\delta_1$ ) conditional knock-out mice and are also Protein Kinase C (PKC) signaling dependent. These findings suggest that TSP4 elevation may contribute to the pathogenesis of chronic pain following nerve injury by disrupting intracellular Ca<sup>2+</sup> signaling via interacting with the Ca<sub>v</sub> $\alpha_2\delta_1$  and the subsequent PKC signaling pathway. Controlling TSP4 mediated intracellular Ca<sup>2+</sup> signaling in peripheral sensory neurons may be a target for analgesic drug development for neuropathic pain.

**Corresponding author:** Bin Pan M.D., Ph.D., Department of Anesthesiology, Medical College of Wisconsin, 8701 Watertown Plank Road, Milwaukee, WI 53226, Phone: 414-955-7526, Fax: 414-955-6507, bpan@mcw.edu.

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## Keywords

Neuropathic pain; Intracellular calcium signaling; Thrombospondin-4; Plasma membrane  $\text{Ca}^{2+}$ -ATPase;  $\text{Ca}^{2+}$  stores and stimulates store-operated  $\text{Ca}^{2+}$  entry; Sarco-endoplasmic reticulum  $\text{Ca}^{2+}$ -ATPase

## 1. Introduction

Neuronal cytoplasmic  $\text{Ca}^{2+}$  is the essential regulator of numerous physiological functions, such as excitation, synaptic transmission, synaptic plasticity, and neuronal differentiation and survival (Catterall et al., 2013; Gemes et al., 2011; Paschen, 2001). When neurons are activated, free cytoplasmic  $\text{Ca}^{2+}$  concentration ( $[\text{Ca}^{2+}]_i$ ) increases due to  $\text{Ca}^{2+}$  influx through voltage-gated  $\text{Ca}^{2+}$  channels (VGCCs) or ligand-gated  $\text{Ca}^{2+}$  channels on the plasma membrane. This may be supplemented by release of  $\text{Ca}^{2+}$  from intracellular stores upon activation of ryanodine receptors (RyRs) and inositol 1,4,5-trisphosphate receptors ( $\text{IP}_3\text{R}$ ) on the membrane of endoplasmic reticulum (ER). In addition,  $\text{Ca}^{2+}$  can enter neurons through channels that are indirectly activated by the depletion of ER  $\text{Ca}^{2+}$  stores in the process known as store-operated  $\text{Ca}^{2+}$  entry (SOCE) (Gemes et al., 2011). After neuronal activation, elevated  $[\text{Ca}^{2+}]_i$  recovers back to resting levels by  $\text{Ca}^{2+}$  extrusion through the plasma membrane  $\text{Ca}^{2+}$ -ATPase (PMCA) (Duncan et al., 2013) and  $\text{Na}^+$ - $\text{Ca}^{2+}$  exchanger (NCX), and by  $\text{Ca}^{2+}$  sequestration into the ER by the sarco/endoplasmic reticulum  $\text{Ca}^{2+}$ -ATPase (SERCA) (Gemes et al., 2012). We have previously observed that, after nerve injury, sensory neurons of the dorsal root ganglia (DRG) develop decreased resting  $[\text{Ca}^{2+}]_i$  (Fuchs et al., 2005), depressed  $\text{Ca}^{2+}$  influx through VGCCs (Hogan et al., 2000), and decreased SERCA function (Duncan et al., 2013), as well as elevated  $\text{Ca}^{2+}$  clearance from the cytoplasm by PMCA function (Gemes et al., 2012) and enhanced  $\text{Ca}^{2+}$  influx through SOCE due to depletion of ER  $\text{Ca}^{2+}$  stores (Gemes et al., 2011). However, the pathological effector driving these events is undefined.

Thrombospondins (TSP) are a family of large oligomeric, extracellular matrix glycoproteins that mediate interactions between cells and interactions of cells with underlying matrix components (Adams, 2001; Risher and Eroglu, 2012). The TSP family consists of five members (TSP 1–5) divided into two subgroups (TSP1/2 vs. TSP3/4/5) according to their functional domains (Risher and Eroglu, 2012). TSP4 is expressed by astrocytes and neurons, in which it can promote neurite outgrowth and synaptogenesis (Arber and Caroni, 1995; Eroglu et al., 2009; Pan et al., 2015). Recently, TSP4 has been found to be a factor contributing to neuropathic pain (Crosby et al., 2014; Kim et al., 2012). Specifically, TSP4 gene and protein expression is elevated in DRGs after peripheral nerve injury, and intrathecal administration of TSP4 protein amplifies excitatory pre-synaptic transmission in the dorsal horn via promoting excitatory synaptogenesis by binding to its receptor, voltage-gated  $\text{Ca}^{2+}$   $\alpha_2\delta_1$  subunit ( $\text{Ca}_v\alpha_2\delta_1$ ) (Kim et al., 2012; Pan et al., 2015; Park et al., 2016). However, specific cellular and molecular mechanisms by which TSP4 acts are still unclear.

Various observations support the view that actions of TSP4 may contribute to the pathological effects of axotomy in sensory neurons. Specifically, application of TSP4 *in vitro* reproduces the *in vivo* effects of nerve injury on VGCCs (Hogan et al., 2000; Pan et al.,

2016). Additionally, expression of TSP4 protein is elevated in axotomized sensory neurons following peripheral nerve injury (Pan et al., 2015), which suggests that TSP4 may have an important role in regulating intracellular signaling. Here, we test our hypothesis that elevated TSP4 after nerve injury leads to depletion of ER  $\text{Ca}^{2+}$  stores and disordered function of PMCA, SERCA, and SOCE in primary sensory neurons.

## 2. Materials and methods

### 2.1. Animals

Male adult mice (129S1/SvImJ) and *TSP4* gene knock-out (KO) mice (B6.129P2-Thbs4tm1Dgen/J) were obtained from The Jackson Laboratory. The  $\text{Ca}_v\alpha_2\delta_1$  advillin conditional knock-out (CKO) mice were generated by crossing mice with Cre-recombinase expression in advillin positive cells to 129/sv background mice with exon 6 of the  $\text{Ca}_v\alpha_2\delta_1$  gene (MGI ID: 88295) floxed with loxP sites (Park et al., 2016).  $\text{CKO}^{\text{Adv-Cre}+/+}$  mice have conditionally knocked out  $\text{Ca}_v\alpha_2\delta_1$  in sensory neurons.  $\text{CKO}^{\text{Adv-Cre}-/-}$  littermates mice were used as wildtype (WT) control. Male Sprague Dawley rats weighing 170–200g used in some experiments were obtained from The Taconic Biosciences. All animals were housed in a room with a 12/12 h day/night cycle and free access to food and water. Male mice were used for initial studies and female mice were used to confirm major findings. All animal experiments were performed according to protocols approved by the Institutional Animal Care and Use Committee of the Medical College of Wisconsin (AUA00001809 for experiments on mice and AUA00002752 for experiments on rats). Those animals were maintained and used according to the NIH *Guide for the Care and Use of Laboratory Animals*, and in compliance with federal, state, and local laws. DRGs from mice and rats were rapidly harvested following deep isoflurane anesthesia and decapitation.

### 2.2. Neuron isolation and plating

DRGs from mice or rats were rapidly harvested following deep isoflurane anesthesia and decapitation. Ganglia were placed in a 35 mm dish containing  $\text{Ca}^{2+}/\text{Mg}^{2+}$ -free, cold HBBS (Life Technologies) and cut into four to six pieces that were incubated in 0.01% blendzyme 2 (Roche Diagnostics, Indianapolis, IN) for 26 min followed by incubation in 0.25% trypsin (Sigma Aldrich, St. Louis, MO) and 0.125% DNase (Sigma) for 30 min, both dissolved in Dulbecco's modified Eagle's medium (DMEM)/F12 with glutaMAX (Invitrogen, Carlsbad, CA). After exposure to 0.1% trypsin inhibitor and centrifugation, the pellet was gently triturated in culture medium containing Neural Basal Media A with B27 supplement (Invitrogen), 0.5 mM glutamine, 10 ng/ml nerve growth factor 7S (Alomone Labs, Jerusalem, Israel) and 0.02 mg/ml gentamicin (Invitrogen). Dissociated neurons were plated onto poly-L-lysine coated glass cover slips (DeutschesSpiegelglas, Carolina Biological Supply, Burlington, NC) and maintained at 37°C in humidified 95% air and 5%  $\text{CO}_2$  for 2 hours, and were studied no later than 8 hours after harvest.

### 2.3. Measurement of cytoplasmic $\text{Ca}^{2+}$ concentration

Measurement of  $[\text{Ca}^{2+}]_i$  was performed following our previously published protocols (Duncan et al., 2013; Gemes et al., 2012). In brief, unless otherwise specified, regular Tyrode's solution (in mM: NaCl 140, KCl 4,  $\text{CaCl}_2$  2,  $\text{MgCl}_2$  2, glucose 10, HEPES 10 with

an osmolarity of 297–300 mOsm and pH 7.40) was used to bathe the neurons. Stock solution of Fura-2-AM (Invitrogen) was dissolved in DMSO and subsequently diluted in the relevant bath solution such that final bath concentration of DMSO was 0.2% or less, which does not affect  $[Ca^{2+}]_i$  (Gemes et al., 2011). The 500  $\mu$ l recording chamber was superfused by gravity-driven flow at a rate of 3 ml/min. Agents were delivered by directed microperfusion controlled by a computerized valve system through a 500  $\mu$ m diameter hollow quartz fiber 300  $\mu$ m upstream from the neurons. This flow completely displaced the bath solution, and constant flow was maintained through this microperfusion pathway by delivery of bath solution when specific agents were not being administered. Dye imaging shows that solution changes were achieved within 200 ms.

Neurons plated on cover slips were exposed to Fura-2-AM (5  $\mu$ M) at room temperature in a solution that contained 2% bovine albumin to aid dispersion of the fluorophore. After 30 min, they were washed 3 times with regular Tyrode's solution, given 30 minutes for de-esterification, and then mounted in the recording chamber. Neurons were first examined under bright field illumination, and those showing signs of lysis, crenulation or superimposed glial cells were excluded. To determine  $[Ca^{2+}]_i$ , the fluorophore was excited alternately with 340 nm and 380 nm wavelength illumination (150 W Xenon, Lambda DG-4, Sutter, Novato, CA), and images were acquired at 510 nm using a cooled 12-bit digital camera (Coolsnapfx, Photometrics, Tucson, AZ) and inverted microscope (Diaphot 200, Nikon Instruments, Melville, NY) through a 20X objective. Recordings from each neuron were obtained as separate regions (MetaFluor, Molecular Devices, Downingtown, PA) at a rate of 3 Hz. After background subtraction, the fluorescence ratio R for individual neurons was determined as the intensity of emission during 340 nm excitation ( $I_{340}$ ) divided by  $I_{380}$ , on a pixel-by-pixel basis, and  $[Ca^{2+}]_i$  was estimated by the formula  $K_d \cdot \beta \cdot (R - R_{min}) / (R_{max} - R)$  where  $\beta = (I_{380max}) / (I_{380min})$ . Values of  $R_{min}$ ,  $R_{max}$  and  $\beta$  were determined by *in-situ* calibration and were 0.38, 8.49 and 9.54, and  $K_d$  was 224 nm. Only neurons with stable baseline R traces were further evaluated. Traces were analyzed using Axograph X 1.1 (Axograph Scientific, Sydney, Australia).

Depolarization-induced  $Ca^{2+}$  transients were generated by microperfusion application of  $K^+$  (50 mM) for 0.3 s. To characterize the rate of recovery of the activity-induced  $[Ca^{2+}]_i$  transient, the trace was fit by a mono-exponential curve ( $y = y_0 + k \times \exp(-x/\tau)$ ) from which the time constant ( $\tau$ ) was derived as a measure of the pace of recovery. To block PMCA function, HEPES was replaced by Trizma (10 mM) in regular Tyrode's with pH at 8.8. To block SERCA function, thapsigargin (TG, 1  $\mu$ M, Sigma-Aldrich, St. Louis, MO) was added into solutions. For quantifying SERCA and PMCA activity, we limited the participation of mitochondria by excluding traces that either had a transient peak above 400nM or showed a shoulder of sustained  $[Ca^{2+}]_i$  elevation during the descending limb of the activity-induced transient (Gemes et al., 2012). To induce the release of stored  $Ca^{2+}$  from the ER, caffeine (20mM) was added to regular Tyrode's solution. SOCE function was quantified as the amplitude of the transient induced by switching from  $Ca^{2+}$ -free bath solution to solution with 2 mM  $Ca^{2+}$ , either with or without TG (1  $\mu$ M). In all TSP4 treatment groups, neurons were incubated with TSP4 (60nM) for 4 hours, which is the dose and duration of application necessary to produce effects based on our previous electrophysiological experiments (Pan et al., 2016).

## 2.4. IB4 staining

After the measurement of cytoplasmic  $\text{Ca}^{2+}$  concentration, neurons were identified by incubating with isolectin B4 (IB4) conjugated with fluorescein isothiocyanate (10  $\mu\text{g}/\text{ml}$ , Sigma-Aldrich, St. Louis, MO) for 10 min and then washed with regular Tyrode's solution. IB4-positive (IB4pos) neurons were identified as those that retained a complete ring of FITC stain around the perimeter of the soma (Barabas et al., 2012).

## 2.5. Injury model

Fifteen mice were subjected to spinal nerve ligation (SNL), based on the original technique (Kim and Chung, 1992) with modifications (Gemes et al., 2012; Pan et al., 2015; Rigaud et al., 2008). Briefly, during anesthesia by inhalation of isoflurane (1.5–2.5% in oxygen), the right lumbar paravertebral region was exposed through a midline incision, and the fifth lumbar (L5) transverse process was removed to expose the L4 spinal nerve, which was ligated with 6-0 silk suture and severed approximately 1 mm distal to their respective DRGs. The wound was closed in layers and the skin stapled. Control mice received skin incision and closure only. DRGs were collected for experiments 15–21 days after the surgery.

## 2.6. Reagents

Fura-2 AM, thapsigargin and all other common chemicals were obtained from Sigma-Aldrich. YM 244769, CGP 37157, N-[2-[[3-(4-Bromophenyl)-2-propenyl]amino]ethyl]-5-isoquinolinesulfonamide dihydrochloride (H89), Calphostin C, Phorbol 12-myristate 13-acetate (PMA), (9R,10S,12S)-2,3,9,10,11,12-Hexahydro-10-hydroxy-9-methyl-1-oxo-9,12-epoxy-1H-diindolo[1,2,3-fg:3',2',1'-kl]pyrrolo[3,4-i][1,6]benzodiazocine-10-carboxylic acid, hexyl ester (KT5720) were obtained from Tocris Bioscience. Ru360 was obtained from EMD Millipore. TSP4 proteins were expressed and purified according to our previous report (Kim et al., 2012).

## 2.7. Data Analysis

DRG sensory neurons are a diverse population with various functional and molecular attributes. Sensory neuron somatic diameter is broadly associated with specific sensory modalities. Whereas small unmyelinated neurons generally carry nociceptive traffic, large myelinated neurons are activated by low threshold stimuli (Ma et al., 2003). Small-diameter sensory neurons are further divided neurochemically into IB4pos and IB4-negative (IB4neg) populations, which have distinct electrophysiological properties and sensitivities to noxious stimuli (Stucky and Lewin, 1999). By defining the small-diameter population of our recorded neurons from mice as having a diameter  $\leq 27\mu\text{m}$  (Barabas et al., 2012), we found that 98% of the IB4pos population were small (Fig. 1A) and 65% of small neurons were IB4pos, which is consistent with previous observations (Barabas et al., 2012; Dirajlal et al., 2003). Defining large neurons as diameter  $> 27\mu\text{m}$ , 4% of which were IB4pos. Those few IB4pos large neurons were ignored in further analysis. For experiments using neurons from rats, we defined the population of our recorded neurons with a diameter  $\leq 34\mu\text{m}$  as small, and neurons with a diameter  $> 34\mu\text{m}$  as large (Gemes et al., 2011).

Statistical analyses were performed with Excel (Microsoft, Redmond, WA), GraphPad Prism 6 (GraphPad Software, Inc, CA) and Sigmaplot 11.0 (Systat Software Inc, Chicago, IL).

Student's *t*-test, and one-way or two-way analysis of variance (ANOVA) with Bonferroni *post hoc* analysis were used to test significance of differences between groups. When our experiment question only focuses on TSP4 effects, analyses were performed by Student's *t*-test to test significance of the effect of TSP4 unless stated specifically. A *p* value less than 0.05 was considered significant. Data are reported as mean  $\pm$  SEM.

### 3. Results

#### 3.1. TSP4 reduces resting $[Ca^{2+}]_i$ of DRG neurons

We have previously found that resting  $[Ca^{2+}]_i$  in both large- and small-sized DRG neurons is decreased after spinal nerve ligation and section (Fuchs et al., 2005). We therefore tested whether TSP4 has similar effects as nerve injury on resting  $[Ca^{2+}]_i$ , which showed that TSP4 incubation (4 hours) minimally reduced the resting  $[Ca^{2+}]_i$  in large-sized neurons but not in small neurons (Fig. 1B).

#### 3.2. TSP4 slows recovery of depolarization-induced $Ca^{2+}$ transients

Following neuronal activation by brief application of  $K^+$ , the depolarization-induced  $Ca^{2+}$  transient returns to resting level in an exponential fashion (Fig. 2A) with a recovery  $\tau$  that is controlled by PMCA and SERCA (Duncan et al., 2013; Gemes et al., 2012). TSP4 significantly increased the recovery  $\tau$  of the depolarization-induced  $Ca^{2+}$  transient in IB4pos and IB4neg small-sized neurons, and also showed a trend towards increased recovery  $\tau$  in large-sized neurons ( $p = 0.08$ ) (Fig. 2A, B).

We and others (Gemes et al., 2012; Usachev et al., 2002) have previously shown that NCX contributes minimally to  $Ca^{2+}$  extrusion after neuronal activation in DRG sensory neurons. Here, we further confirmed this with blockade of NCX by applying NCX inhibitor YM 244769 (5  $\mu$ M), which did not change recovery  $\tau$  with or without TSP4 treatment (Fig. 2C, D). Additionally, we eliminated NCX function by replacing  $Na^+$  with N-methyl-d-glucamine (NMDG) in the Tyrode's solution, which also failed to change recovery  $\tau$  with or without TSP4 treatment (data not shown). In contrast, a study in which  $Li^+$  was used to replace  $Na^+$  showed decreased  $Ca^{2+}$  clearance following depolarization in a subpopulation of DRG neurons (Lu et al., 2006). We have confirmed similar results using  $Li^+$  Tyrode's solution in our preparations (data not shown).

While this may appear to be discordant with our findings with YM 244769 and NMDG substitution,  $Li^+$  is known to participate in  $Li^+/Ca^{2+}$  exchange (Palty et al., 2004), and can activate  $Na^+$ -activated  $K^+$  currents (Hess et al., 2007). Thus, we believe the most reliable findings indicate that NCX contributes minimally to  $Ca^{2+}$  transient recovery in sensory neurons, and is not a target of TSP4.

In order to clearly identify effects of TSP4 on SERCA and PMCA, we limited the participation of mitochondria by excluding traces that either had a transient peak above 400nM or showed a shoulder of sustained  $[Ca^{2+}]_i$  elevation during the descending limb of the activity-induced transient (Gemes et al., 2012). It has been reported that mitochondria might also contribute to  $Ca^{2+}$  buffering in sensory neurons even during small increases in  $[Ca^{2+}]_i$ . This was concluded in a study using the protonophore carbonyl cyanide m-

chlorophenyl hydrazine (CCCP) to block uptake and release of  $\text{Ca}^{2+}$  in mitochondria (Lu et al., 2006). However, the use of protonophores, including CCCP and carbonyl cyanide-p-trifluoromethoxyphenylhydrazone, also leads to robust hydrolysis of cytoplasmic ATP (Budd and Nicholls, 1996), which in turn limits PMCA function, making interpretation of these data difficult. Therefore, to further test whether mitochondria contribute to  $\text{Ca}^{2+}$  buffering under the conditions of our present study (transient peak less than 400 nM and without a shoulder), we used Ru360, a cell permeable and selective inhibitor of the mitochondrial  $\text{Ca}^{2+}$  uniporter, to block mitochondrial  $\text{Ca}^{2+}$  influx and CGP 37157 (a specific antagonist of mitochondrial  $\text{Na}^{+}$ - $\text{Ca}^{2+}$  exchanger) to block mitochondrial  $\text{Ca}^{2+}$  efflux. We found that application of Ru360 (10 $\mu\text{M}$ ) and CGP 37157 (10 $\mu\text{M}$ ) did not change recovery  $\tau$  with TSP4 treatment (Fig. 2E, F) or without TSP4 treatment (data not shown).

On the basis of these findings, our further examination of the delayed recovery of the depolarization-induced  $\text{Ca}^{2+}$  transient during TSP4 treatment focused on specific regulation of PMCA and SERCA to resolve the specific effects of these interacting  $\text{Ca}^{2+}$  buffering systems.

### 3.3 TSP4 accelerates PMCA function

PMCA is the principal means by which DRG neurons extrude  $\text{Ca}^{2+}$  from the cytoplasm to reduce  $[\text{Ca}^{2+}]_i$  (Gemes et al., 2012). PMCA function is elevated following nerve injury by SNL, which contributes to neuronal hyperexcitability (Gemes et al., 2012). To test the regulation of PMCA by TSP4, we blocked SERCA function with TG (1 $\mu\text{M}$ ) and measured recovery of  $[\text{Ca}^{2+}]_i$  following brief neuronal depolarization (Gemes et al., 2012). Blocking SERCA function with TG increased  $\tau$  in all neuronal types (Fig. 3). Two-way ANOVA analysis showed an interaction between effects of TSP4 and TG treatments in large-sized neurons ( $F_{1,92} = 21.35$ ,  $p < 0.001$ ; Fig. 3B left), small IB4neg neurons ( $F_{1,63} = 9.00$ ,  $p < 0.01$ ; Fig. 3C left) and small IB4pos neurons ( $F_{1,225} = 8.93$ ,  $p < 0.01$ ; Fig. 3D left). To determine whether there was a difference in  $\tau$  between groups treated with TG with respect to the TSP4-induced increase, data from TG-treated neurons were analyzed as a percent change from control. These data confirmed a diminished TG effect after TSP4 treatment in all neuronal groups (Fig. 3B right, C right, D right), which suggests elevated PMCA function by TSP4.

### 3.4 TSP4 inhibits SERCA function

SERCA, which resides on neuronal ER membrane, is a  $\text{Ca}^{2+}$  ATPase that reduces  $[\text{Ca}^{2+}]_i$  by pumping  $\text{Ca}^{2+}$  from the cytosol of the cell into the ER lumen. To functionally isolate SERCA, a bath Tyrode's solution with pH 8.8 was used to eliminate PMCA  $\text{Ca}^{2+}$  extrusion, which requires reciprocal  $\text{H}^{+}$  exchange (Duncan et al., 2013). Blocking PMCA function with pH 8.8 Tyrode's solution causes resting  $[\text{Ca}^{2+}]_i$  to increase due to unopposed SOCE function (Fig. 4A). This rise in  $[\text{Ca}^{2+}]_i$  was greater for TSP4-treated neurons in each neuronal group (Fig. 4B), which indicates that SOCE function is elevated by TSP4. To measure the effect of TSP4 on isolated SERCA function, the depolarization-induced  $\text{Ca}^{2+}$  transient was tested in pH 8.8 Tyrode's solution, which blocks PMCA function (Fig. 4C). Two-way ANOVA analysis showed an interaction between effects of TSP4 and pH 8.8 solution treatments in large-sized neurons ( $F_{1,103} = 10.96$ ,  $p < 0.001$ ; Fig. 4D left), small IB4neg neurons ( $F_{1,59} =$

8.29,  $p < 0.01$ ; Fig. 4E left) and small IB4pos neurons ( $F_{1,300} = 34.49$ ,  $p < 0.001$ ; Fig. 4F left). To determine whether there was a difference between groups treated with pH 8.8 solution with respect to the TSP4-induced increase in  $\tau$ , data from pH 8.8 solution-treated neurons were analyzed as a percent change from control. These data confirmed an increased effect of pH 8.8 solution after TSP4 treatment in all neuronal groups (Fig. 4D–F right), which suggests inhibited SERCA function by TSP4. This effect of TSP4 on SERCA, which acts to prolong the activity-induced  $[Ca^{2+}]_i$  transient, competes with the effect of TSP4 on PMCA, which acts to shorten the duration of the transient. Since the overall effect of TSP4 is to prolong the transient (Fig. 2), it appears that the TSP4 effect on SERCA is dominant. To determine if this effect of TSP4 on SERCA is influenced by sex of the animal subject, female mice were also tested. In them, TSP4 had similar effects on the recovery  $\tau$  of depolarization-induced  $Ca^{2+}$  transient during SERCA block with pH 8.8 Tyrode's solution in large-sized neurons (vehicle:  $3.73 \pm 0.56$  s,  $n = 10$ ; TSP4 treatment:  $7.65 \pm 0.77$  s,  $n = 9$ ;  $p < 0.001$ , t-test) and small-sized neurons (vehicle:  $6.67 \pm 0.92$  s,  $n = 27$ ; TSP4 treatment:  $16.67 \pm 1.9$  s,  $n = 23$ ;  $p < 0.001$ , t-test).

### 3.5 TSP4 reduces ER $Ca^{2+}$ stores

Since nerve injury depletes ER  $Ca^{2+}$  stores and SERCA's function is necessary to replenish ER  $Ca^{2+}$  stores (Rigaud et al., 2009), we next examined the effect of TSP4 on this parameter. The ryanodine receptor  $Ca^{2+}$  release channels (RyRs) are the major pathway of both constitutive and  $Ca^{2+}$ -induced  $Ca^{2+}$  release from ER in DRG neurons (Shmigol et al., 1995; Usachev et al., 1993). By activating the RyRs with caffeine (20mM), the resulting transient elevation of  $[Ca^{2+}]_i$  can be measured as a representation of  $Ca^{2+}$  transfer from the ER pool into the cytoplasm. The caffeine induced transient amplitude reflects the initial release from the ER while the duration of the transient is influenced by other  $Ca^{2+}$  regulatory processes including PMCA, SOCE and mitochondrial  $Ca^{2+}$  buffering. We therefore quantified transient amplitude as the best reflection of cytoplasmic  $Ca^{2+}$  load released from the ER. The caffeine-induced transient of large-sized neurons and IB4neg small-sized neurons was suppressed by TSP4, indicating reduced  $Ca^{2+}$  stores in ER (Fig. 5A, B). Interestingly, IB4pos small-sized neurons showed minimal response to caffeine. It has been previously reported that small-sized DRG neurons from mice have little or no response to caffeine application (Shmigol et al., 1994), whereas we have previously seen caffeine induced ER  $Ca^{2+}$  release in small-sized DRG neurons from rats that on average was even greater than in large-sized neurons (Rigaud et al., 2009). To confirm that these differences are species dependent, we again tested the response to caffeine in DRG neurons from rats. Consistent with our previous findings, caffeine induced larger transient in small-sized IB4pos (Fig. 5C). In contrast to mouse neurons, TSP4 had no effect on large-sized and IB4pos small-sized rat neurons, whereas TSP4 suppressed caffeine-releasable  $Ca^{2+}$  stores in IB4neg small neurons (Fig. 5C). These findings show that TSP4 treatment reduces ER  $Ca^{2+}$  stores in sensory neuron in a subtype-specific fashion, but the sensitivity of specific neuronal subtypes is species-dependent.

An alternative approach to measure intracellular  $Ca^{2+}$  stores is by TG application, which triggers a  $Ca^{2+}$  release transient by blocking SERCA and leaving constitutive  $Ca^{2+}$  release through RyRs unopposed. Under this condition, the  $Ca^{2+}$  release is a relatively slow process.

We quantified  $\text{Ca}^{2+}$  release from ER stores by measuring transient amplitude (Rigaud et al., 2009). This revealed TSP4 suppression of  $\text{Ca}^{2+}$  release from ER stores in all neuronal groups (Fig. 5D, E), including those IB4pos small neurons.

### 3.6 TSP4 increases SOCE by depletion of ER $\text{Ca}^{2+}$ stores

We have previously reported that nerve injury elevates SOCE function, which increases neuronal excitability (Gemes et al., 2011). This raises the question of whether TSP4 is involved in the regulation of SOCE. Here, we first examined the effect of TSP4 on constitutive SOCE in resting neurons. Resting  $[\text{Ca}^{2+}]_i$  is set by the opposing actions of PMCA and SOCE. Our findings above show that TSP4 activates PMCA, but the resting  $[\text{Ca}^{2+}]_i$  is affected only minimally and only in the large neuron group. This predicts that TSP4 administration may induce an enhanced level of SOCE. We tested this by incubating neurons with TG (1 $\mu\text{M}$ ) and allowing the  $\text{Ca}^{2+}$  release transient to recover (10 min). This showed that resting  $[\text{Ca}^{2+}]_i$  was elevated after TG, reflecting increased activity of SOCE in all neuronal subtypes and in the presence or absence of TSP4 (Fig. 5D, E). However, this effect of TG on resting  $[\text{Ca}^{2+}]_i$  was diminished in neurons incubated with TSP4 compared to controls (Fig. 5D, F). This suggests that in the presence of added TSP4, either SOCE is already relatively activated, causing TG application to produce proportionately less effect, or that maximum inducible SOCE is diminished by TSP4.

To resolve this, we next examined SOCE directly by eliminating SOCE with  $\text{Ca}^{2+}$ -free solution and measuring the increase of  $[\text{Ca}^{2+}]_i$  upon readdition of  $\text{Ca}^{2+}$  to the bath, which represents  $\text{Ca}^{2+}$  influx through SOCE. Bath  $\text{Ca}^{2+}$  withdrawal resulted in a fall of  $[\text{Ca}^{2+}]_i$  that was greater in TSP4 treated neurons in all neuronal groups (Fig. 6A, B).  $\text{Ca}^{2+}$  transients induced by readdition of  $\text{Ca}^{2+}$  were larger in neurons treated with TSP4 in all neuronal groups (Fig. 6A, C). These findings indicate that SOCE is activated in the presence of TSP4.

The influence of TSP4 upon SOCE may be indirect through its effect of depleting ER  $\text{Ca}^{2+}$  store level (Fig. 5B, C), which is the definitive factor that regulates SOCE. To identify whether TSP4 additionally interacts directly with the SOCE channel elements, we first depleted ER  $\text{Ca}^{2+}$  stores by exposing neurons to TG (1 $\mu\text{M}$ , 8min) in  $\text{Ca}^{2+}$ -free bath solution. Subsequent readdition of bath  $\text{Ca}^{2+}$  resulted in SOCE transients that were comparable in control and TSP4 groups (Fig. 6D, E), which indicates that TSP4 has no effect on fully activated SOCE channels. Taken together, these results indicate that TSP4 increases SOCE function indirectly through its depletion of ER  $\text{Ca}^{2+}$  stores, which is at least in part the result of TSP4 blocking SERCA function.

To summarize (Table 1), our findings indicate that TSP4 activates PMCA and suppresses SERCA, which depletes ER  $\text{Ca}^{2+}$  stores and thereby indirectly drives greater SOCE activity.

### 3.7 SERCA and PMCA dysfunction after nerve injury is dependent on TSP4

We next tested the role of TSP4 in generating injury-induced loss of SERCA function and gain of PMCA and SOCE function that we have previously observed (Duncan et al., 2013; Gemes et al., 2011; Gemes et al., 2012) by examining the effects of SNL in TSP4 KO mice, compared to that in littermate WT animals. For these experiments, only small-sized DRG neurons were studied, without further differentiation by IB4 binding. Two-way ANOVA

analysis showed an interaction between nerve injury and effects of TSP4 ( $F_{1,395} = 4.15$ ,  $p < 0.05$ ; Fig. 7A, B, C left). To determine whether there was a difference between different TSP4 genotype groups with respect to the nerve injury induced decrease in  $Ca^{2+}$  transient amplitude, data from SNL rats were analyzed as a percent change from control. These data confirmed that decreased  $[Ca^{2+}]_i$  transient amplitude after SNL was absent in DRG neurons from TSP4 KO mice (Fig. 7C right), which is consistent with our previous findings of TSP4-dependent loss of  $I_{Ca}$  after neuronal injury (McCallum et al., 2006; Pan et al., 2015). SNL increased the recovery  $\tau$  of depolarization-induced  $Ca^{2+}$  transients in neurons from WT mice, but this effect was absent in DRG neurons from TSP4 KO mice (two-way ANOVA,  $F_{1,395} = 10.79$ ,  $p < 0.01$ ; Fig. 7D left). SNL injury also specifically decreased SERCA function (two-way ANOVA,  $F_{1,205} = 8.59$ ,  $p < 0.01$ ; Fig. 8A, B left) and elevated PMCA function (two-way ANOVA,  $F_{1,205} = 3.96$ ,  $p < 0.05$ ; Fig. 8C left, D left), both of which were eliminated in neurons from TSP4 KO mice (Fig. 8C right, D right) when data from SNL mice were analyzed as a percent change from control. We infer from these results that TSP4 is a critical mediator of the effects of injury upon intracellular  $Ca^{2+}$  signaling. In the absence of injury,  $[Ca^{2+}]_i$  transient amplitude (Fig. 7C left) and SERCA function (Fig. 8B left) is also elevated in neurons from TSP4 KO mice compared with neurons from TSP4 WT mice, which indicates that constitutive TSP4 has a physiological role in maintaining SERCA function and intracellular  $Ca^{2+}$  homeostasis.

### 3.8 TSP4 regulates $[Ca^{2+}]_i$ by binding with the $Ca_v\alpha_2\delta_1$ subunit

The VGCC  $\alpha_2\delta_1$  subunit is the binding site of both TSP4 (Eroglu et al., 2009; Park et al., 2016) and the analgesic gabapentin (GBP) (Hendrich et al., 2008; Pan et al., 2016) although they might work on different domains of  $Ca_v\alpha_2\delta_1$  subunit (Lana et al., 2016). This suggests that TSP4 may have its intracellular effects by binding with the  $Ca_v\alpha_2\delta_1$ . We therefore examined whether genetic deletion of the  $Ca_v\alpha_2\delta_1$  can block TSP4's effects on cytoplasmic  $Ca^{2+}$ . In sensory neurons dissociated from CKO<sup>Adv-Cre<sup>-/-</sup></sup> control mice, TSP4 increased the recovery  $\tau$  of the depolarization-induced  $Ca^{2+}$  transient ( $2.16 \pm 0.20$  s,  $n = 62$ ; Fig. 9A). This is somewhat faster than the recovery in wildtype animals in Fig. 2 ( $3.00 \pm 0.32$  s,  $n = 132$ ,  $p < 0.05$ ), which may be due to somewhat different backgrounds as they were obtained from different breeding colonies at different vendors and have different ability to clear intracellular  $Ca^{2+}$ . Two-way ANOVA analysis showed an interaction between  $Ca_v\alpha_2\delta_1$  knockout and effects of TSP4 on the recovery  $\tau$  of the depolarization-induced  $Ca^{2+}$  transient ( $F_{2,272} = 4.25$ ,  $p < 0.05$ ; Fig. 9B left). Further analysis of data from TSP4-treated neurons, using percent change from vehicle control, revealed that prolongation of the  $Ca^{2+}$  transient by TSP4 was eliminated (Fig. 9B right) in sensory neurons dissociated from the CKO<sup>Adv-Cre<sup>+/-</sup></sup> mice that lack  $Ca_v\alpha_2\delta_1$ , suggesting that TSP4 regulates SERCA by initially binding with the  $Ca_v\alpha_2\delta_1$  subunit.

### 3.9 Regulation of SERCA function by TSP4 is PKC signaling dependent

Both TSP4 and GBP can bind with the VGCC  $\alpha_2\delta_1$  subunit. GBP can potently block TSP4-induced formation of excitatory synapses (Dolphin, 2013; Eroglu et al., 2009; Hendrich et al., 2008). Additionally, GBP can modulate PKC signaling in the spinal cord dorsal horn (Yeh et al., 2011; Zhang et al., 2015) and regulate NMDA receptor function via PKC (Gu and Huang, 2001). Furthermore, activation of PKC can accelerate SERCA  $Ca^{2+}$  uptake into

ER (Usachev et al., 2006). We therefore examined whether PKC signaling is involved in the TSP4's regulation on SERCA function. Two-way ANOVA analysis showed an interaction between PKC signaling regulation and effects of TSP4 in regular solution ( $F_{3,204} = 9.37$ ,  $p < 0.001$ ; Fig. 10A left) and in pH8.8 solution ( $F_{3,204} = 3.60$ ,  $p < 0.05$ , Fig. 10B left), and *post hoc* testing revealed that incubation (4hr) with PKC inhibitors GF109203X (5  $\mu\text{M}$ ) or calphostin C (500 nM) increased the recovery  $\tau$  of brief depolarization-induced  $\text{Ca}^{2+}$  transients both in regular solution (Fig. 10A) and in pH8.8 solution (PMCA blocked, Fig. 10B), and both inhibitors blocked the effect of TSP4 on the recovery  $\tau$  of brief depolarization-induced  $\text{Ca}^{2+}$  transients (one-way ANOVA,  $F_{3,100} = 25.42$ ,  $p < 0.001$ ; Fig. 10A right) or TSP4 suppression of SERCA function (one-way ANOVA,  $F_{3,100} = 16.6$ ,  $p < 0.001$ ; Fig. 10B right). These observations suggest that TSP4 regulates SERCA via PKC signaling. To test this further, we examined the effect of PKC activation on TSP4's effects on SERCA function. We found that the PKC agonist PMA (1  $\mu\text{M}$ ) accelerated SERCA function and occluded the further action of TSP4 (Fig. 10 A, B). These findings together support a role of PKC in mediating the action of TSP4 on SERCA.

GBP has been reported to activate cAMP-dependent protein kinase A (PKA) signaling in various tissues (Lee et al., 2008; Takasu et al., 2008). In cardiomyocytes, SERCA function is regulated by PKA via phospholamban (Dutta et al., 2002; Kranias and Hajjar, 2012; Veglia and Cembran, 2013), which is not expressed in neurons (Plessers et al., 1991), although similar proteins are (Dou and Joseph, 1996). We therefore tested whether PKA signaling took part in TSP4's effects on SERCA function. Two-way ANOVA analysis showed an interaction between PKA signaling regulation and effects of TSP4 in regular solution ( $F_{2,146} = 3.96$ ,  $p < 0.05$ ; Fig. 10C left) and in pH8.8 solution ( $F_{2,146} = 3.37$ ,  $p < 0.05$ ; Fig. 10D left), while further testing revealed that PKA blockers H89 (10  $\mu\text{M}$ ) and KT5720 (1  $\mu\text{M}$ ) had no effect on TSP4 on the recovery  $\tau$  of brief depolarization-induced  $\text{Ca}^{2+}$  transients (one-way ANOVA,  $F_{2,75} = 0.95$ ,  $p = 0.39$ ; Fig. 10C right) or TSP4 suppression of SERCA function (one-way ANOVA,  $F_{2,75} = 0.62$ ,  $p = 0.54$ ; Fig. 10D right). From these findings, we conclude that the action of TSP4 on SERCA is not dependent on PKA signaling.

## 4 Discussion

The experiments reported here have investigated the effects of TSP4 on intracellular  $\text{Ca}^{2+}$  homeostasis regulation in primary sensory neurons. Our findings show that TSP4 accelerates the function of PMCA and inhibits the SERCA function, which results in reduced ER  $\text{Ca}^{2+}$  stores that in turn triggers increased SOCE function (Fig. 11). We further confirmed that increased TSP4 after nerve injury contributes to the altered PMCA and SERCA function through interacting with the  $\text{Ca}_v\alpha_2\delta_1$ .

DRG sensory neurons are a diverse population with different functions that can be characterized by neuronal size and neurochemical staining, such as IB4pos or IB4neg (Dirajlal et al., 2003; Ma et al., 2003; Stucky and Lewin, 1999). We found that IB4pos and IB4neg populations of small neurons differ in resting  $[\text{Ca}^{2+}]_i$  levels and response to caffeine application (Fig. 1, 5). Furthermore, the subpopulations of neurons show different responses to caffeine application mice and rats (Fig. 5). A possible explanation for this discrepancy is that neurons with different IB4 staining from different species have different expression

levels of RyRs isoforms, and thus different sensitivities to caffeine. All three isoforms (RyR1, RyR2 and RyR3) (Lanner et al., 2010) can be co-expressed in most of the tissues, but RyR1 is primarily expressed in skeletal muscle and RyR2 is in cardiac muscles, while RyR3 is mainly expressed in brain and DRG neurons (Lanner et al., 2010; Ooashi et al., 2005). These RyRs isoforms have different sensitivities to caffeine (Fessenden et al., 2000; Lanner et al., 2010), so further study may determine whether differential expression accounts for the distinct  $\text{Ca}^{2+}$  signaling features we have identified in DRG neuronal subgroups. Despite the differences in caffeine release of stores, our thapsigargin experiments nonetheless show that  $\text{Ca}^{2+}$  stores in IB4pos small neurons are present, and are depressed by TSP4, suggesting that TSP4 suppresses SERCA function in this neuronal subpopulation as well.

Nerve injury disrupts  $\text{Ca}^{2+}$  signaling in DRG neurons. Both inward  $\text{Ca}^{2+}$  current through VGCCs and resting  $[\text{Ca}^{2+}]_i$  are reduced after nerve trauma (Fuchs et al., 2005; Hogan et al., 2000), PMCA function is increased, and SERCA is impaired, culminating in depleted ER  $\text{Ca}^{2+}$  stores and reduced  $[\text{Ca}^{2+}]_i$  in the injured neurons (Duncan et al., 2013; Gemes et al., 2011). Resting  $[\text{Ca}^{2+}]_i$  level is regulated by  $\text{Ca}^{2+}$  influx through SOCE, uptake into ER by SERCA, expulsion by PMCA, and sequestration by mitochondria. Our previous studies of DRG neurons from SNL injury rats indicate that elevated PMCA function contributes to depressed resting  $[\text{Ca}^{2+}]_i$  (Duncan et al., 2013; Fuchs et al., 2005). In the present study, we found only minimally effects of TSP4 on resting  $[\text{Ca}^{2+}]_i$  in large sized neurons (Fig. 1), despite substantial influences on SERCA, PMCA and SOCE. One possible explanation is that TSP4 has offsetting effects, such that TSP4-induced SERCA depression and consequent SOCE enhancement increases resting  $[\text{Ca}^{2+}]_i$ , while TSP4-induced PMCA activation drives resting  $[\text{Ca}^{2+}]_i$  down, resulting in minimal or no change overall. Thus, factors other than TSP4 must account for the strong depression of resting  $[\text{Ca}^{2+}]_i$  after injury (Fuchs et al., 2005).

SERCA plays an important role in maintaining cellular homeostasis, including in sensory neurons (Gemes et al., 2011). For instance, blocking SERCA with thapsigargin is a trigger for ER stress through depletion of ER  $\text{Ca}^{2+}$  stores (Luciani et al., 2009). Our present study shows that TSP4 reduces SERCA function, depletes ER  $\text{Ca}^{2+}$  stores, and elevates SOCE, which are key features of ER stress. A consequence of ER stress is activation of extracellular signal-regulated kinase (ERK), JNK, and p38 MAPK, all of which are activated in sensory neurons after peripheral nerve injury (Jin et al., 2003; Zhuang et al., 2006), supporting the view that ER stress is a component of neuropathic pain. Furthermore, depleting ER  $\text{Ca}^{2+}$  stores increases sensory neuron excitability (Gemes et al., 2009). These observations together suggest that the TSP4 signaling pathway may account for ER stress in neuropathic pain. This is supported by our observation of enhanced TSP4 expression in painful nerve injury models (Pan et al., 2015).

Finally, ER stress in cardiac myocytes can itself trigger the generation of TSP4, perhaps as an ER-resident adaptive (Lynch et al., 2012). If this also occurs in sensory neurons, then a positive feedback loop could result in which increased expression of TSP4 produces ER stress, which then drives further production of TSP4.

The  $\text{Ca}_v\alpha_2\delta_1$  has previously been identified as the major binding partner for TSP4 synaptogenesis (Eroglu et al., 2009; Park et al., 2016). TSP4 has weak binding with  $\text{Ca}_v\alpha_2\delta_1$  intracellularly and robust binding with membrane low density lipoprotein receptor-related protein 1 receptors in a tsA201 cell culture system (Lana et al., 2016). In HEK 293 cell culture system, TSP4 can also bind with intracellular stromal interaction molecule 1, which can bind with cartilage oligomeric matrix protein to regulate intracellular  $\text{Ca}^{2+}$  homeostasis (Duquette et al., 2014). Our data also suggest that binding of TSP4 to  $\text{Ca}_v\alpha_2\delta_1$  is required for the observed modulation of SERCA and PMCA. Like TSP4, expression of  $\text{Ca}_v\alpha_2\delta_1$  is also upregulated after nerve injury in neurons of the involved DRGs (Luo et al., 2001; Valder et al., 2003; Wang et al., 2002), and  $\text{Ca}_v\alpha_2\delta_1$  knock-out delays the onset of mechanical hypersensitivity after nerve injury (Patel et al., 2013). VGCCs, which are the main pathway for  $\text{Ca}^{2+}$  entry during neuronal activation, are composed of a pore-forming  $\alpha_1$  subunit as well as auxiliary  $\beta$ ,  $\alpha_2\delta$  and  $\gamma$  subunits (Catterall, 2000). Phosphorylation of the  $\text{Ca}_v\alpha_2\delta_1$  can modulate N-type VGCCs via an ERK-dependent mechanism (Martin et al., 2006). We previously found that TSP4 regulates  $I_{\text{Ca}}$  through VGCCs by binding with  $\text{Ca}_v\alpha_2\delta_1$  (Pan et al., 2016). Thus it appears that the  $\text{Ca}_v\alpha_2\delta_1$  is a key element for TSP4 signaling in general.

We have found that activation of sensory neuron PKC accelerates  $\text{Ca}^{2+}$  uptake into ER by accelerating SERCA function, consistent with a previous report (Usachev et al., 2006), and we additionally showed blocking PKC inhibits SERCA function. By binding with  $\alpha_2\delta_1$  calcium subunit, TSP4 modulates SERCA and PMCA through a PKC dependent pathway that links activation of  $\alpha_2\delta_1$   $\text{Ca}^{2+}$  subunit and inhibition of SERCA function. Additional evidence of this role of PKC comes from a report that shows GBP regulation of PKC signaling in the spinal cord of rats underlying neuropathic pain (Yeh et al., 2011).

Taken together, our findings show that TSP4 regulates  $\text{Ca}^{2+}$  homeostasis in primary sensory neurons by affecting  $\text{Ca}^{2+}$  influx and buffering, including SERCA and PMCA, via PKC signaling after binding to  $\text{Ca}_v\alpha_2\delta_1$ . Elevated TSP4 expression may be a key factor that disrupts  $\text{Ca}^{2+}$  homeostasis after neuronal injury, and may serve as a valuable target for treatment of neuropathic pain.

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## Abbreviations

<b>TSP4</b>	thrombospondin-4
<b><math>\text{Ca}_v\alpha_2\delta_1</math></b>	voltage-gated calcium channel $\alpha_2\delta_1$ subunit
<b><math>[\text{Ca}^{2+}]_i</math></b>	cytoplasmic $\text{Ca}^{2+}$ concentration
<b>DRG</b>	dorsal root ganglion
<b>VGCCs</b>	Voltage-gated calcium channels

<b>SERCA</b>	sarco-endoplasmic reticulum Ca <sup>2+</sup> -ATPase
<b>PMCA</b>	plasma membrane Ca <sup>2+</sup> -ATPase
<b>NCX</b>	Na <sup>+</sup> -Ca <sup>2+</sup> exchanger
<b>ER</b>	endoplasmic reticulum
<b>SOCE</b>	Ca <sup>2+</sup> stores and stimulates store-operated Ca <sup>2+</sup> entry
<b>RyRs</b>	ryanodine receptors
<b>IP<sub>3</sub>R</b>	inositol 1,4,5-trisphosphate receptors
<b>KO</b>	knock-out
<b>CKO</b>	conditional knock-out
<b>Wildtype</b>	WT
<b>SNL</b>	spinal nerve ligation
<b>PKC</b>	protein kinase C
<b>PKA</b>	protein kinase A
<b>IB4</b>	isolectin B4
<b>ERK</b>	Extracellular signal-regulated kinase
<b>GBP</b>	gabapentin

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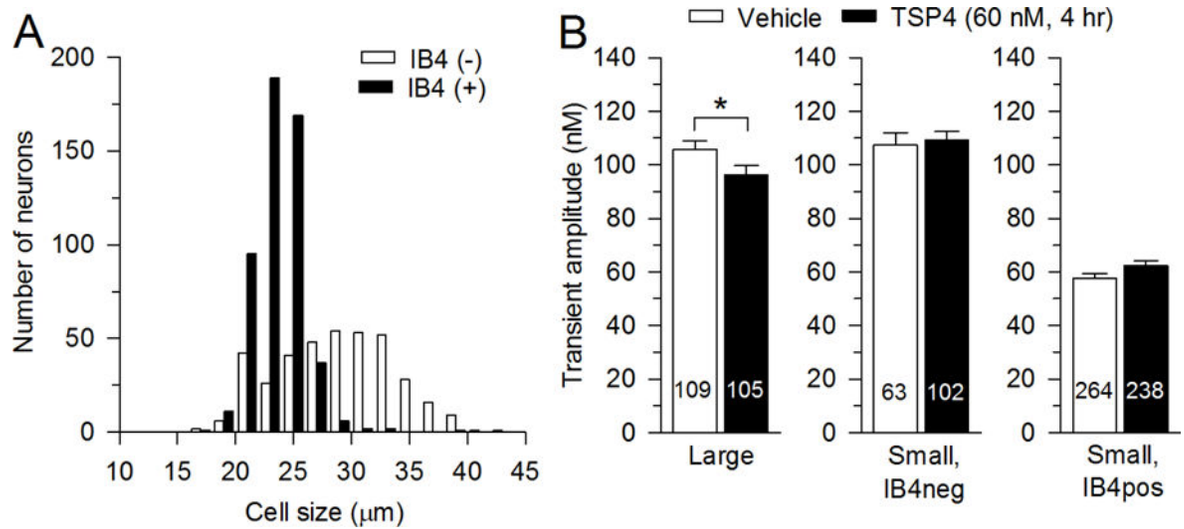
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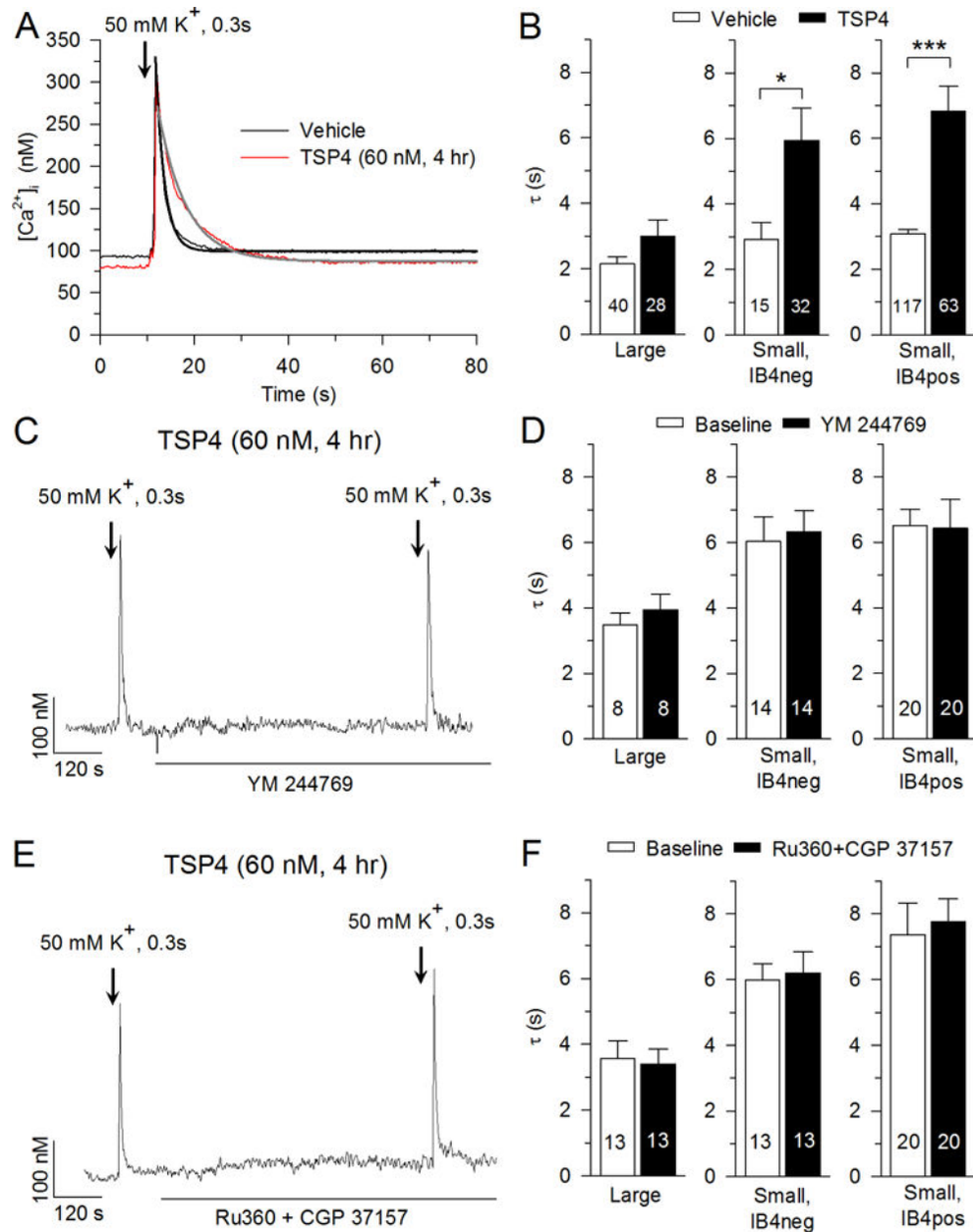
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### Highlights

- Thrombospondin-4 (TSP4) inhibits sarco-endoplasmic reticulum  $\text{Ca}^{2+}$ -ATPase (SERCA) function.
- Effects of TSP4 on SERCA function are attenuated in voltage-gated  $\text{Ca}^{2+}$  channel  $\text{Cav}\alpha_2\delta_1$  subunit knockout mice.
- Nerve injury-induced depression of SERCA function is attenuated in TSP4 knockout mice.
- Effects of TSP4 on SERCA function are dependent on protein kinase C signaling.



**Fig. 1.** Effects of TSP4 on resting  $[Ca^{2+}]_i$  of DRG neurons. **A**, Classification of neurons according to size and IB4 staining. **B**, Summary data show effects of TSP4 incubation on resting  $[Ca^{2+}]_i$  in neurons of different groups categorized by size and IB4 binding. The numbers in the bars represent the n for neurons in groups, and bars show mean  $\pm$  SEM, here and in subsequent figures. \*  $p < 0.05$ .



**Fig. 2.** Effects of TSP4 on the depolarization-induced  $\text{Ca}^{2+}$  transient. **A**, Sample traces show the influence of TSP4 on recovery of the  $\text{Ca}^{2+}$  transient induced by neuronal depolarization with KCl (50 mM  $\text{K}^+$  for 0.3 s). Black and gray lines show single exponential fit of the recovery of  $[\text{Ca}^{2+}]_i$ . **B**, Summary data show TSP4 decreased  $[\text{Ca}^{2+}]_i$  recovery following depolarization in small-sized neurons. **C**, Sample trace shows the effect of NCX inhibitor, YM 244769 (5  $\mu\text{M}$ ), on recovery of the  $\text{Ca}^{2+}$  transient induced by neuronal depolarization with KCl solution. **D**, Summary data show that YM 244769 did not change  $[\text{Ca}^{2+}]_i$  recovery following depolarization (paired  $t$ -test). **E**, Sample trace shows the effect of mitochondrial  $\text{Ca}^{2+}$  uniporter inhibitor, Ru360 (10  $\mu\text{M}$ ), and mitochondrial NCX inhibitor, CGP 37157 (10  $\mu\text{M}$ ), on recovery of the  $\text{Ca}^{2+}$  transient induced by neuronal depolarization with KCl

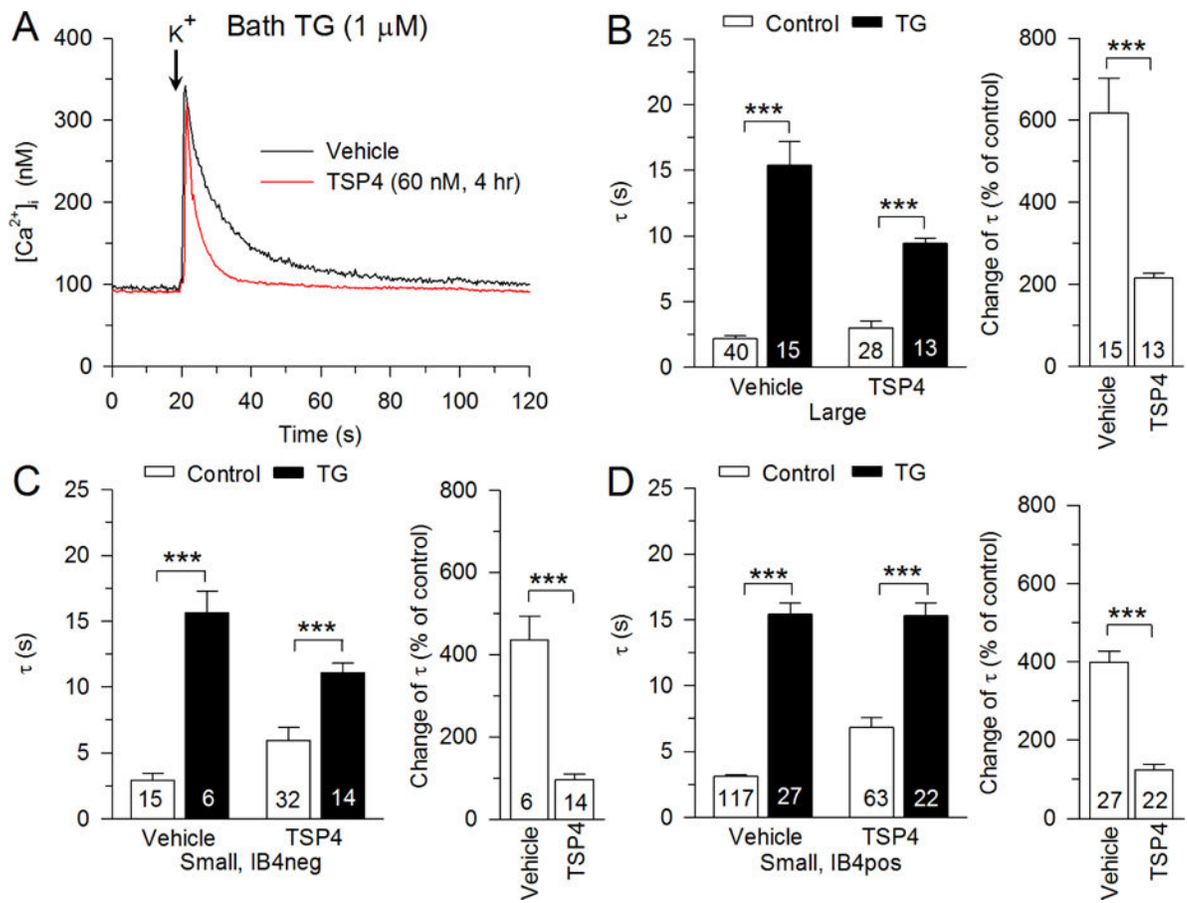
solution. *F*, Summary data show that Ru360 and CGP 37157 did not change  $[Ca^{2+}]_i$  recovery following depolarization (paired t-test). \*  $p < 0.05$ , \*\*\*  $p < 0.001$ .

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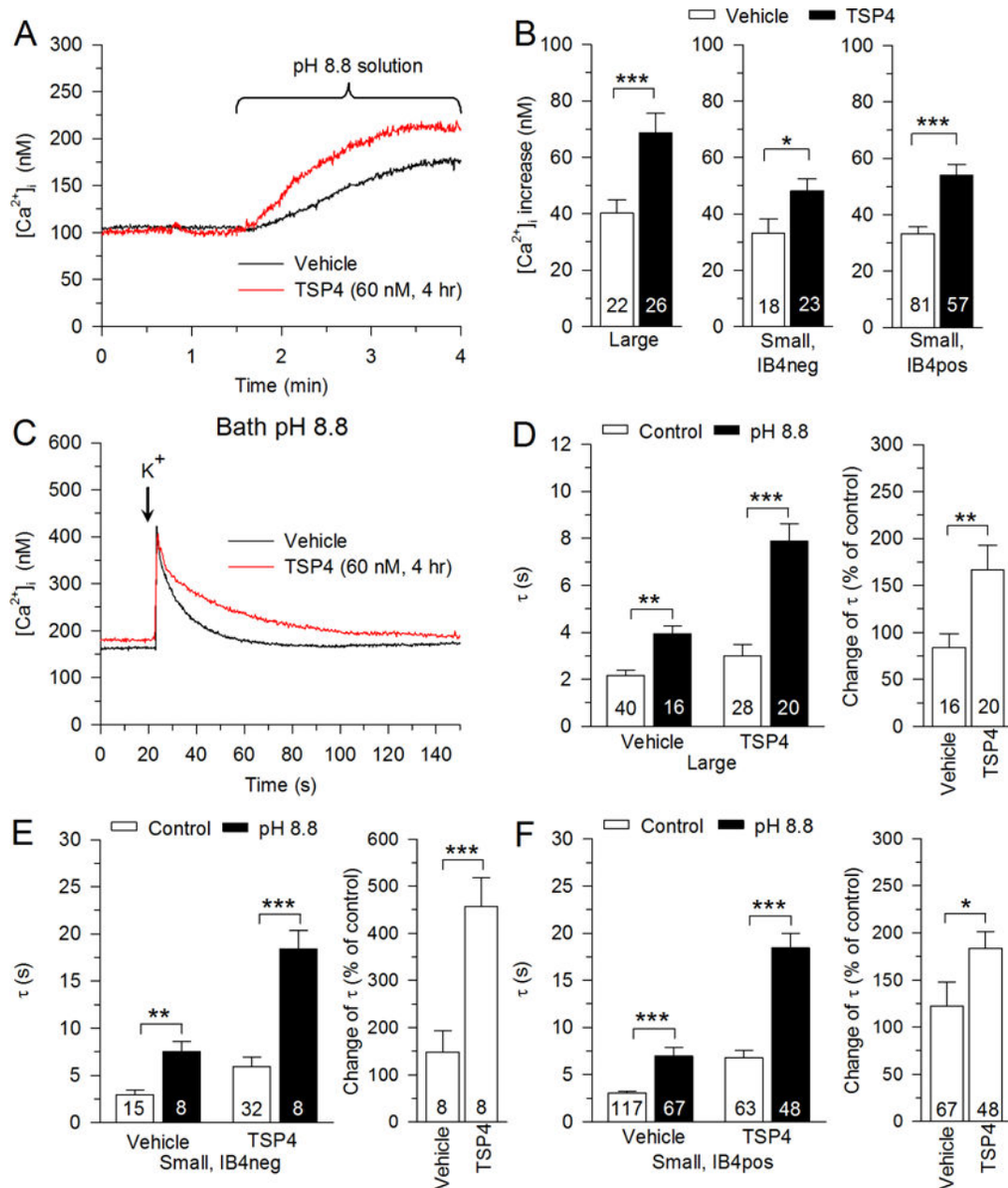
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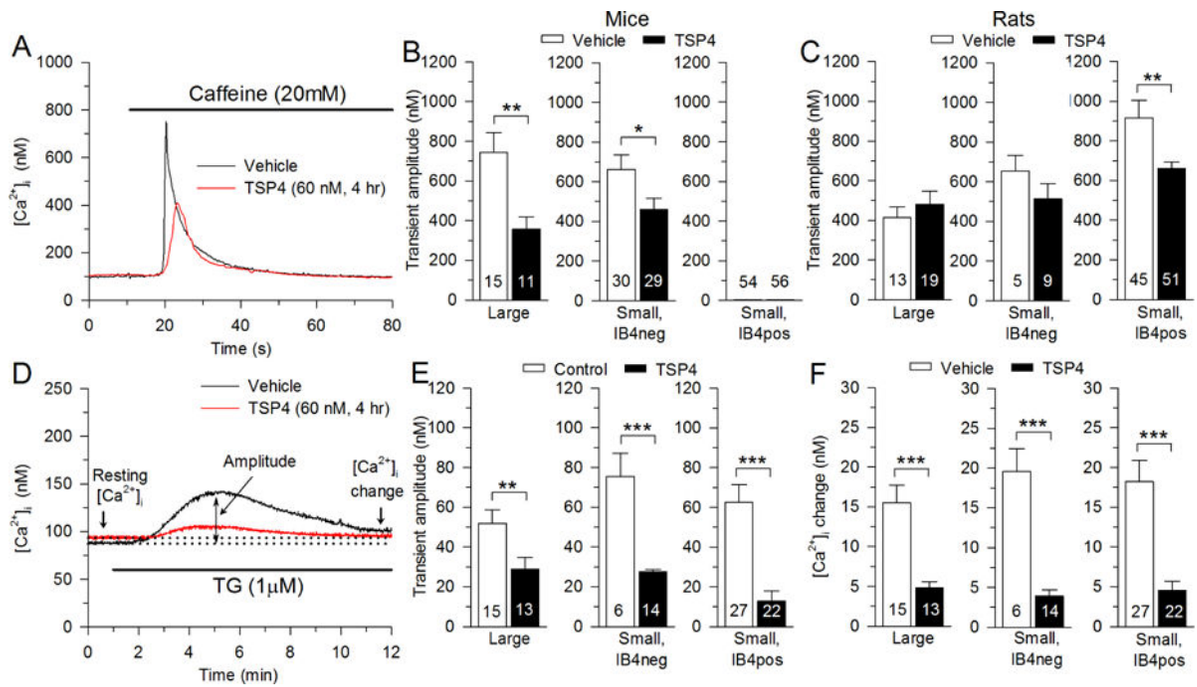
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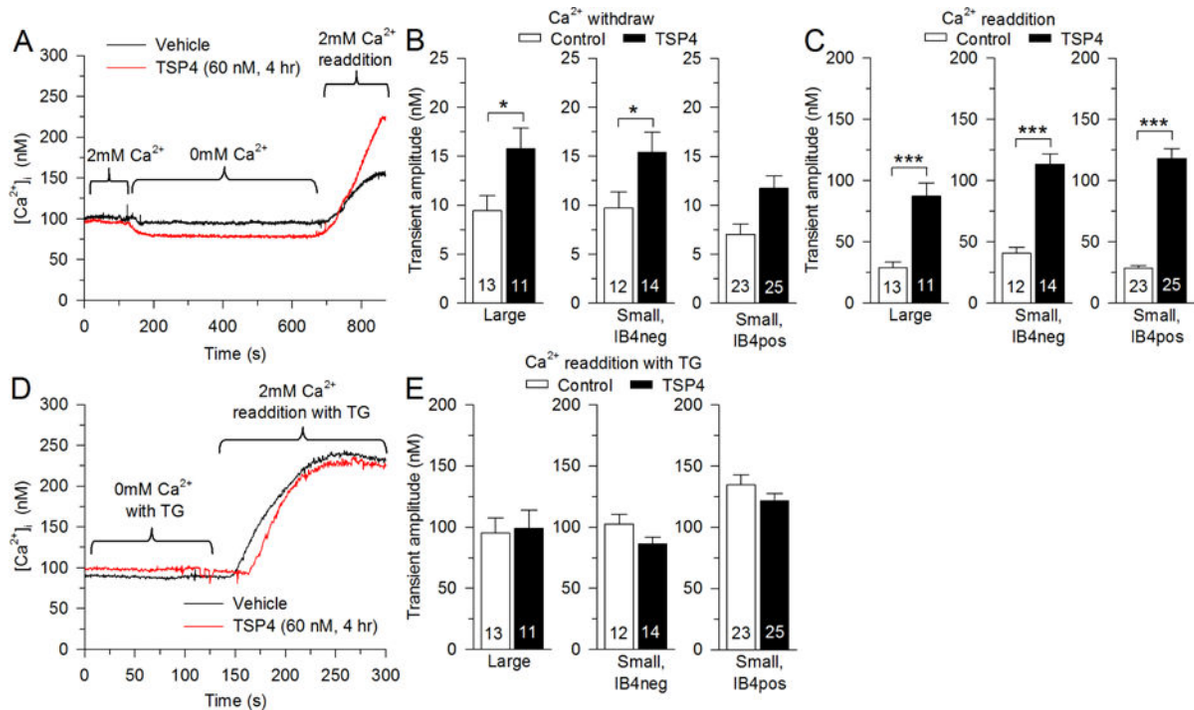
**Fig. 3.** Regulation of PMCA function by TSP4. **A**, Sample traces show the effect of TSP4 on PMCA function, revealed by measuring recovery of the depolarization-induced transient during inhibition of SERCA function with thapsigargin (TG, 1  $\mu$ M). Summary data show that effects of TSP4 on PMCA functions in large-sized neurons (**B**), small-sized IB4neg neurons (**C**), and small-sized IB4pos neurons (**D**). Control data are from Fig. 2. \*\*\*  $p < 0.001$ .



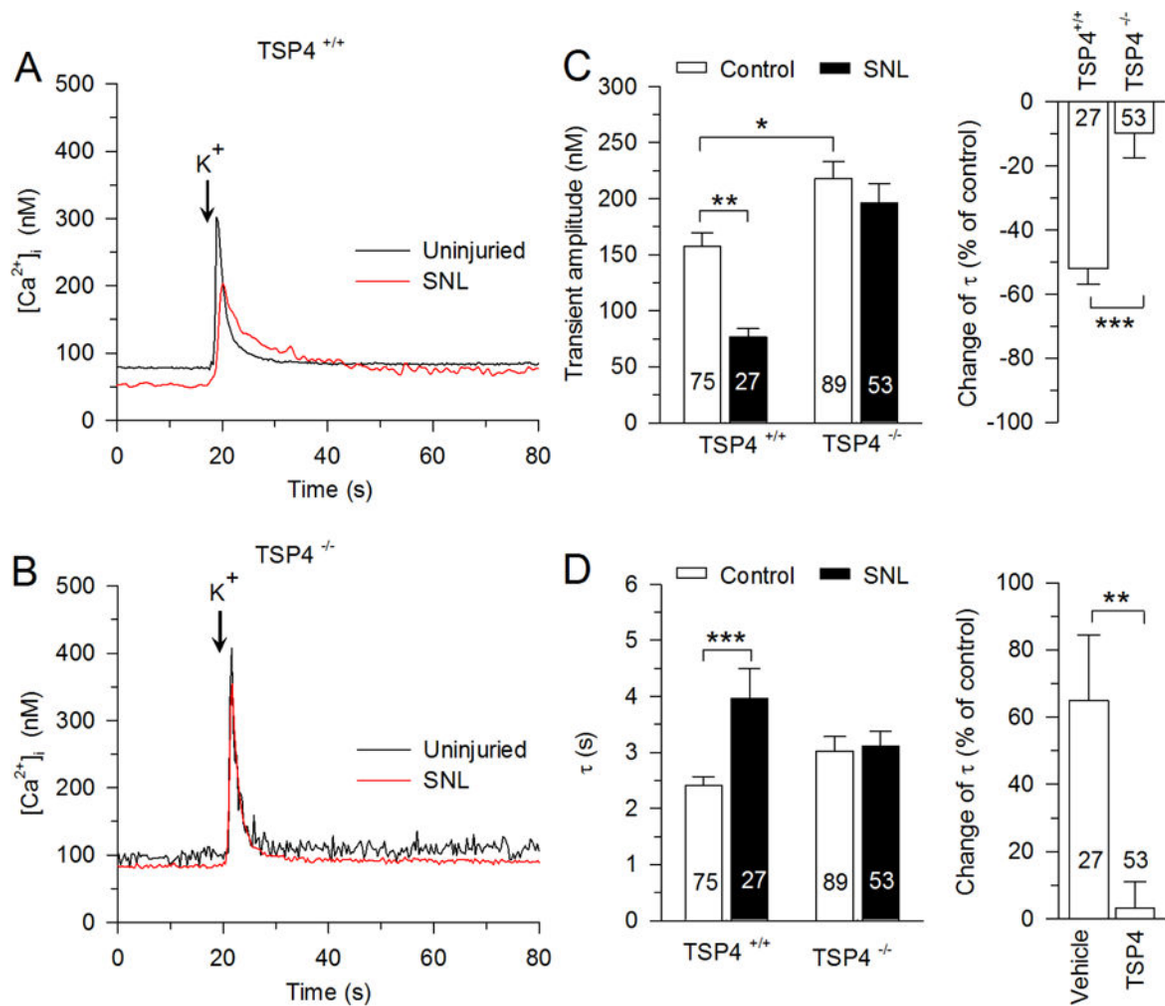
**Fig. 4.** Regulation of SERCA function by TSP4. **A**, Sample traces show the effect of inhibition of PMCA function with pH 8.8 Tyrode's solution on neurons with and without TSP4 treatment. **B**, Summary data show effects of TSP4 on resting [Ca<sup>2+</sup>]<sub>i</sub> in pH 8.8 Tyrode's solution. **C**, Sample traces show the effect of TSP4 on SERCA function, revealed by measuring recovery of the depolarization-induced transient recovery during inhibition of PMCA function with pH 8.8 Tyrode's solution. Summary data show that TSP4 decreased SERCA functions in large-sized neurons (**D**), small-sized IB4neg neurons (**E**), and small-sized IB4pos neurons (**F**). Control data are from Fig. 2. \* p < 0.05, \*\* p < 0.01, \*\*\* p < 0.001.



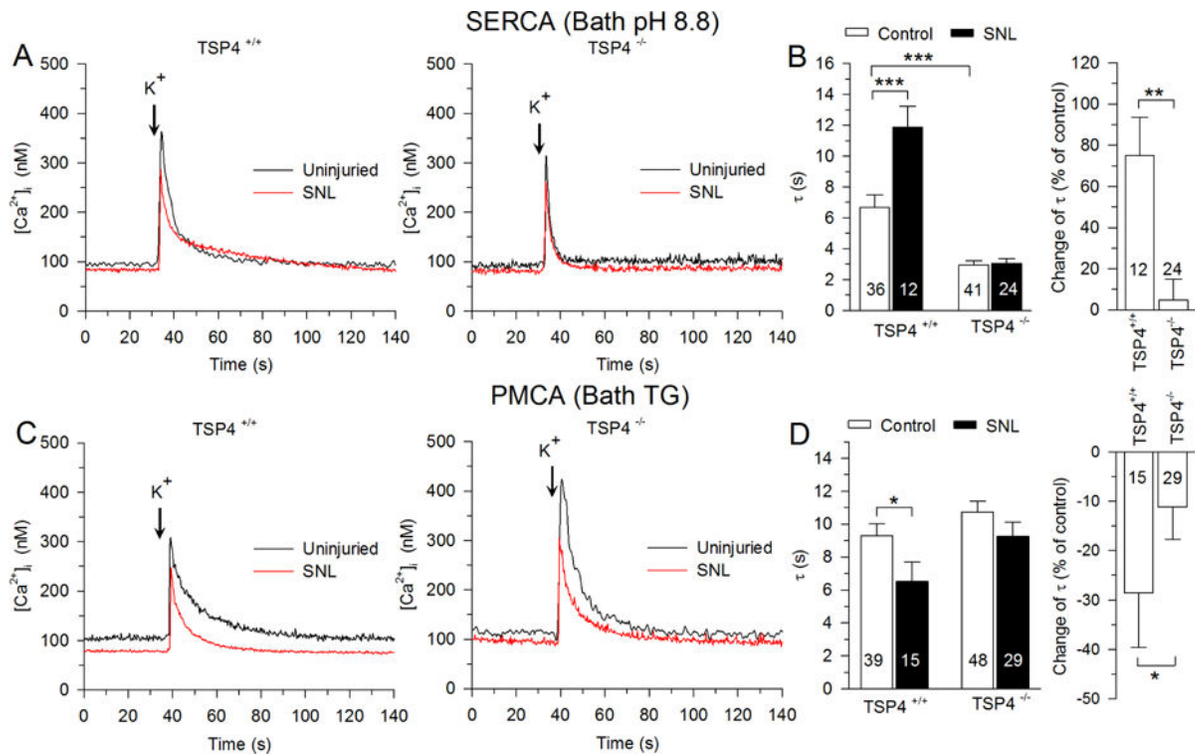
**Fig. 5.** Regulation of ER  $\text{Ca}^{2+}$  store by TSP4. **A**, Sample traces show the effect of TSP4 on the  $\text{Ca}^{2+}$  transient induced by release of ER stores by caffeine (20 mM). **B**, Summary data show reduced releasable  $\text{Ca}^{2+}$  stores, measured as  $\text{Ca}^{2+}$  transient amplitude, in mouse neurons incubated in TSP4. **C**, Summary data show TSP4 effects in rat neurons. **D**, Sample traces show the effect of TSP4 on the  $\text{Ca}^{2+}$  transient induced by the release of stores by the SERCA blocker TG, 1  $\mu\text{M}$ . **E**, Summary data show TSP4 suppresses the TG-induced  $\text{Ca}^{2+}$  transient amplitude. **F**, Summary data show TSP4 suppresses the TG-induced elevation of resting  $[\text{Ca}^{2+}]_i$ . \*  $p < 0.05$ , \*\*  $p < 0.01$ , \*\*\*  $p < 0.001$ .

**Fig. 6.**

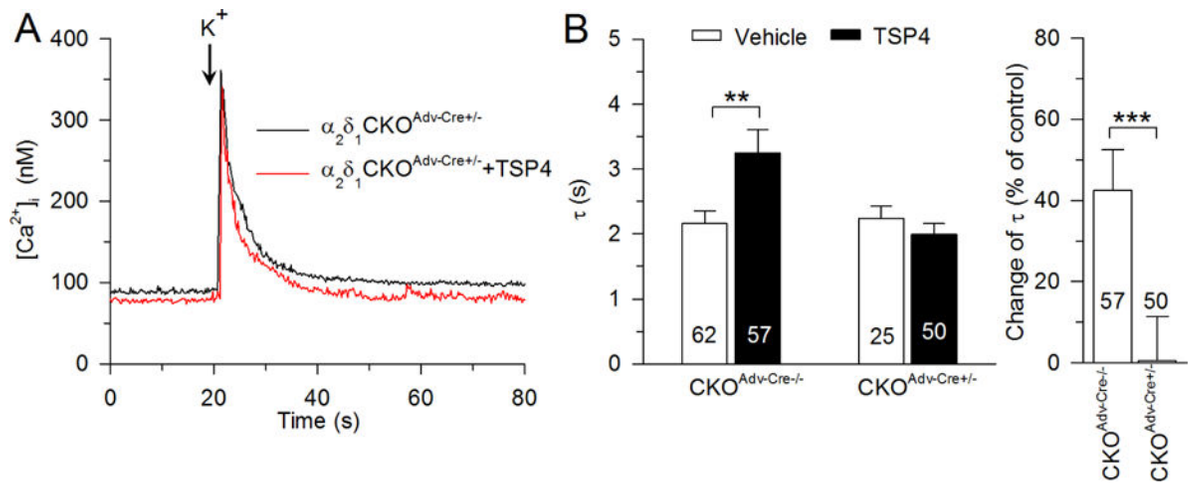
Regulation of SOCE by TSP4. **A**, Sample traces show the effect of TSP4 on SOCE. **B**, TSP4 amplifies the  $[Ca^{2+}]_i$  decrease during  $Ca^{2+}$ -free solution, which blocks SOCE. **C**, TSP4 incubation amplifies the  $[Ca^{2+}]_i$  rise during bath  $Ca^{2+}$  readdition. **D**, **E**, TSP4 has no effect on the  $[Ca^{2+}]_i$  rise during bath  $Ca^{2+}$  readdition when SOCE is fully activated by SERCA blockade with TG (1  $\mu$ M). \*  $p < 0.05$ , \*\*\*  $p < 0.001$ .

**Fig. 7.**

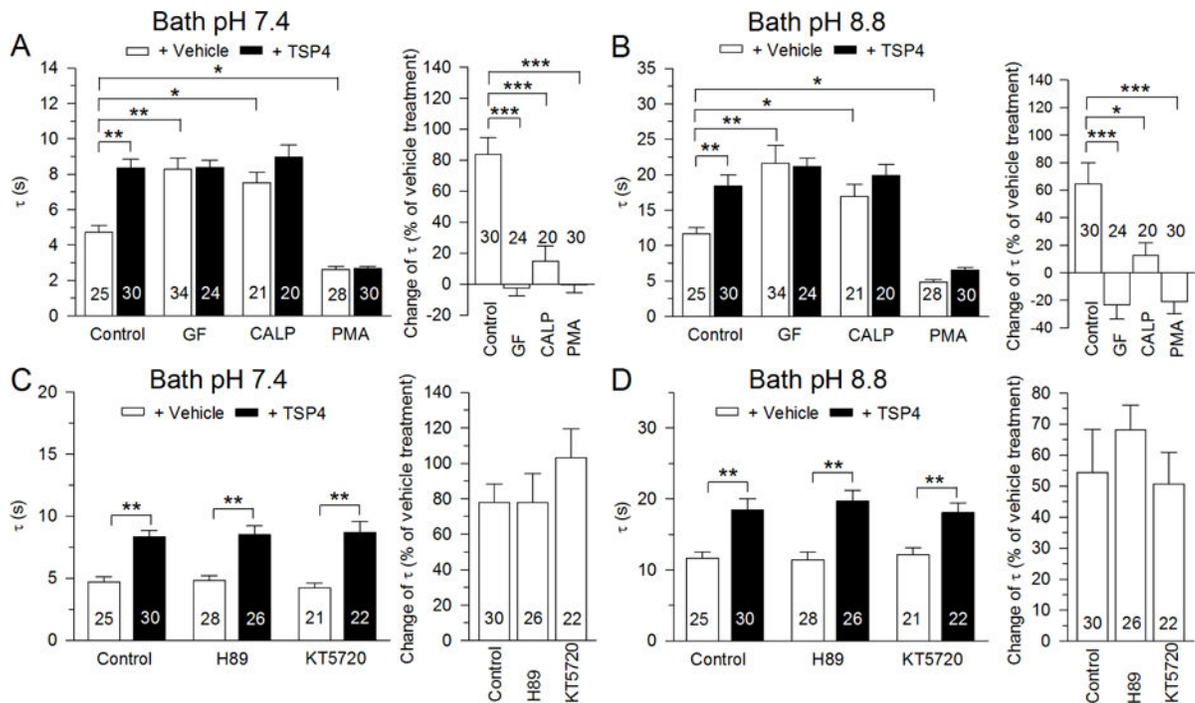
The influence of nerve injury (spinal nerve ligation, SNL) on the depolarization-induced Ca<sup>2+</sup> transient depends on TSP4. **A**, SNL slows the recovery of the Ca<sup>2+</sup> transient in small-sized neurons from WT mice. **B**, SNL has no effect on Ca<sup>2+</sup> transient recovery in small-sized neurons from TSP4 KO mice. **C**, Summary data show that the transient amplitude is reduced after SNL in WT mice (left), but this effect of injury is absent in TSP4 KO mice (right). **D**, Summary data show that the transient recovery  $\tau$  is reduced after SNL in WT mice (left), but this effect of injury is absent in TSP4 KO mice (right). \*  $p < 0.05$ , \*\*  $p < 0.01$ , \*\*\*  $p < 0.001$ .



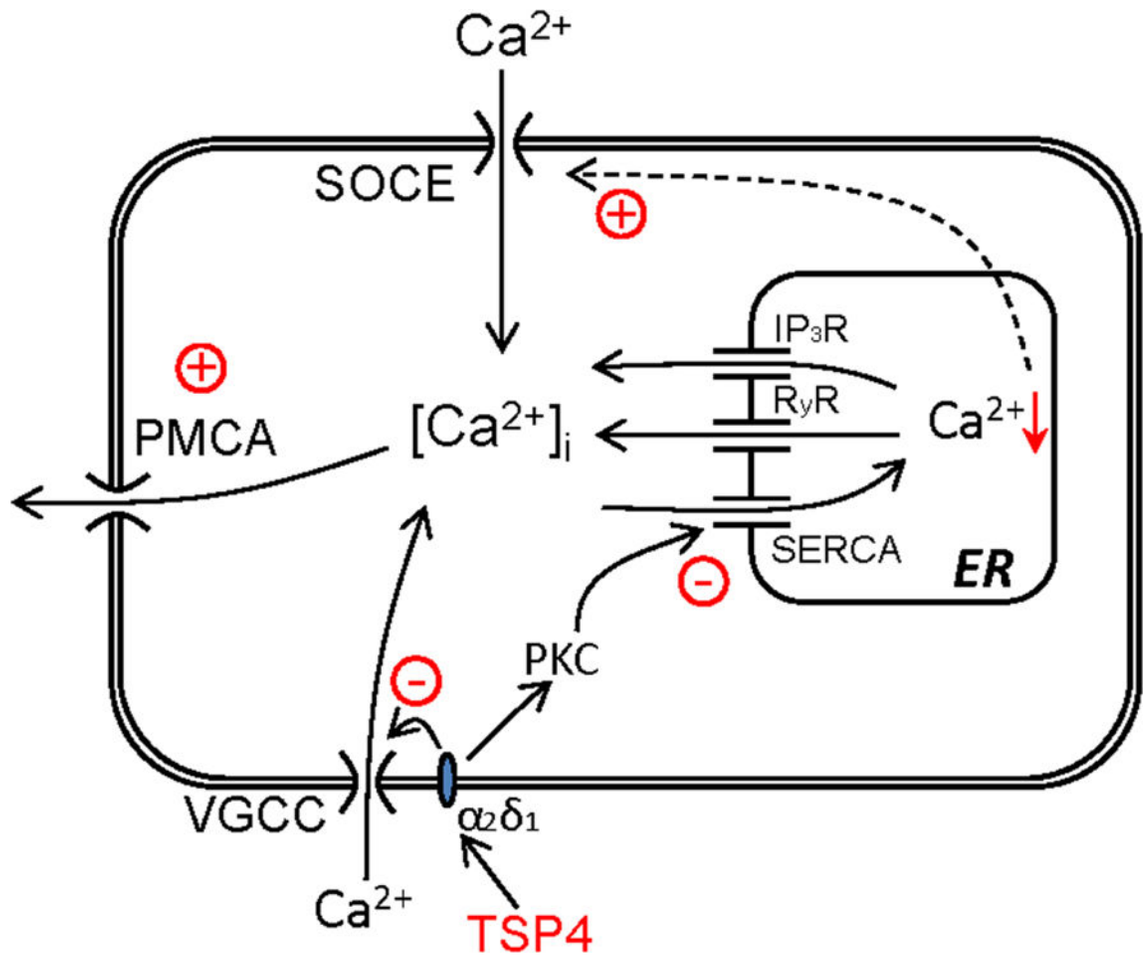
**Fig. 8.** Injury-induced inhibition of SERCA function and elevation of PMCA function depends on TSP4. Sample traces (A.) and summary data (B, left) show that SERCA function was reduced by injury (SNL) in small-sized neurons from WT mice. Similarly, sample traces (C.) and summary data (D, left) show that PMCA function is enhanced by injury. In TSP4 KO animals, these effects of injury are absent for both SERCA (B, right) and PMCA (D, left). \*  $p < 0.05$ , \*\*  $p < 0.01$ , \*\*\*  $p < 0.001$ .



**Fig. 9.** The  $Ca_v\alpha_2\delta_1$  is required for TSP4's regulation of recovery of the depolarization-induced  $Ca^{2+}$  transient. Sample traces (**A**) and summary data (**B**) show that TSP4-induced increase of  $[Ca^{2+}]_i$ ; the recovery  $\tau$  was eliminated in small-sized DRG neurons from  $Ca_v\alpha_2\delta_1$  conditional KO mice. \*\*  $p < 0.01$ , \*\*\*  $p < 0.001$ .

**Fig. 10.**

TSP4's regulation of depolarization-induced  $\text{Ca}^{2+}$  transient is PKC-dependent. The recovery  $\tau$  is prolonged and the effect of TSP4 is eliminated by PKC inhibitors GF 109203X (GF, 5  $\mu\text{M}$ ) and Calphostin C (CALP, 500nM) in both normal bath solution (A, left) and after isolating SERCA function by bath pH 8.8 (B, left) in small-sized DRG neurons. The PKC activator (PMA, 1  $\mu\text{M}$ ) enhances transient recovery and eliminates the effect of TSP4 in both normal bath solution (A, right) and after isolating SERCA function by bath pH 8.8 (B, right). PKA blockers H89 (10  $\mu\text{M}$ ) and KT5720 (1  $\mu\text{M}$ ) had no effect on TSP4 in both normal bath solution (C) or after isolating SERCA function by bath pH 8.8 (D) in small-sized DRG neurons. \*  $p < 0.05$ , \*\*  $p < 0.01$ , \*\*\*  $p < 0.001$ .



**Fig. 11.** Model of TSP4 regulation of  $[Ca^{2+}]_i$  signaling. TSP4 first binds with the  $Ca_v\alpha_2\delta_1$  and possibly with activated PKC. Activated PKC in turn decreases SERCA function, which indirectly increases SOCE function by depleting calcium stores in ER. Additionally, TSP4 amplifies PMCA activity, but to a lesser extent than the effect on SERCA, such that  $[Ca^{2+}]_i$  is minimally altered.

**Table 1**

Summary of TSP4's effects (Direction of arrow represents upregulation or downregulation).

	Large-sized Neurons	Small-sized Neurons	
		IB4neg	IB4pos
Resting $[Ca^{2+}]_i$	↓	–	–
Recovery	–	↓	↓
PMCA	↑	↑	–
SERCA	↓	↓	↓
ER $Ca^{2+}$ stores	↓	↓	–
Resting SOCE	↑	↑	↑
Maximal SOCE	–	–	–

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