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Cellular/Molecular

Calcineurin Dysregulation Underlies Spinal Cord Injury-Induced K⁺ Channel Dysfunction in DRG Neurons

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Dysfunction of the fast-inactivating Kv3.4 potassium current in dorsal root ganglion (DRG) neurons contributes to the hyperexcitability associated with persistent pain induced by spinal cord injury (SCI). However, the underlying mechanism is not known. In light of our previous work demonstrating modulation of the Kv3.4 channel by phosphorylation, we investigated the role of the phosphatase calcineurin (CaN) using electrophysiological, molecular, and imaging approaches in adult female Sprague Dawley rats. Pharmacological inhibition of CaN in small-diameter DRG neurons slowed repolarization of the somatic action potential (AP) and attenuated the Kv3.4 current. Attenuated Kv3.4 currents also exhibited slowed inactivation. We observed similar effects on the recombinant Kv3.4 channel heterologously expressed in Chinese hamster ovary cells, supporting our findings in DRG neurons. Elucidating the molecular basis of these effects, mutation of four previously characterized serines within the Kv3.4 N-terminal inactivation domain eliminated the effects of CaN inhibition on the Kv3.4 current. SCI similarly induced concurrent Kv3.4 current attenuation and slowing of inactivation. Although there was little change in CaN expression and localization after injury, SCI induced upregulation of the native regulator of CaN 1 (RCAN1) in the DRG at the transcript and protein levels. Consistent with CaN inhibition resulting from RCAN1 upregulation, overexpression of RCAN1 in naive DRG neurons recapitulated the effects of pharmacological CaN inhibition on the Kv3.4 current and the AP. Overall, these results demonstrate a novel regulatory pathway that links CaN, RCAN1, and Kv3.4 in DRG neurons. Dysregulation of this pathway might underlie a peripheral mechanism of pain sensitization induced by SCI.

Key words: calcineurin; Kv3.4; pain; RCAN1; spinal cord injury

Significance Statement

Pain sensitization associated with spinal cord injury (SCI) involves poorly understood maladaptive modulation of neuronal excitability. Although central mechanisms have received significant attention, recent studies have identified peripheral nerve hyperexcitability as a driver of persistent pain signaling after SCI. However, the ion channels and signaling molecules responsible for this change in primary sensory neuron excitability are still not well defined. To address this problem, this study used complementary electrophysiological and molecular methods to determine how Kv3.4, a voltage-gated K⁺ channel robustly expressed in dorsal root ganglion neurons, becomes dysfunctional upon calcineurin (CaN) inhibition. The results strongly suggest that CaN inhibition underlies SCI-induced dysfunction of Kv3.4 and the associated excitability changes through upregulation of the native regulator of CaN 1 (RCAN1).

Introduction

Accumulating evidence suggests that changes in the peripheral nervous system play a major role in promoting pain sensitization after spinal cord injury (SCI) (Bedi et al., 2010; Yang et al., 2014). In neuropathic pain, ion channel dysfunction is associated with...
hyperexcitability in dorsal root ganglion (DRG) neurons (Dubin and Papatoupin, 2010; Gold and Gebhart, 2010; Hughes et al., 2012; Trimmer, 2014; Tsantoulas and McMahan, 2014; Walters, 2015). Whether analogous changes mediate SCI-induced pain sensitization, which does not involve direct injury of primary sensory neurons, remains to be determined. Here, we elucidated a novel mechanism involving dysregulation of the Kv3.4 channel induced by dysregulation of calcineurin (CaN) in DRG neurons after SCI.

Homotetrameric Kv3.4 channels are highly expressed and are major regulators of action potential (AP) polarization in small-diameter DRG neurons. Kv3.4 channel dysfunction in DRG neurons has been implicated in chronic pain syndromes (Chien et al., 2007; Duan et al., 2012; Li et al., 2014; Ritter et al., 2015a). Protein kinase C (PKC) modulates Kv3.4 by phosphorylating four serines within the N-terminal inactivation domain (NTID) (Covarrubias et al., 1994; Beck et al., 1998; Antz et al., 1999). This post-translational modification quickly switches the Kv3.4 current phenotype from fast-inactivating to slow/non-inactivating. However, the phosphatase that reverses it was not known. SCI induces attenuation of the Kv3.4 current and slowing of inactivation (Ritter et al., 2012; Ritter et al., 2015a). This suggests a homeostatic mechanism whereby slower inactivation strengthens the remaining Kv3.4 current as it strives to oppose SCI-induced hyperexcitability. In support of a dysregulated physiological role, severe siRNA-induced Kv3.4 knock-down in DRG neurons broadens the AP and eliminates acceleration of AP repolarization upon PKC activation (Ritter et al., 2012). To investigate molecules responsible for SCI-induced attenuation of the Kv3.4 current, we hypothesize that calcineurin (CaN) modulates Kv3.4 function. SCI-induced dysregulation of CaN would impair Kv3.4 function, promoting DRG hyperexcitability and the associated pain sensitization.

CaN is a Ca$^{2+}$-dependent serine-threonine protein phosphatase expressed in the nervous system, including the DRG (Goto et al., 1986; Strack et al., 1996). The immunosuppressants tacrolimus (FK506) and cyclosporin A (CsA) are CaN inhibitors responsible for CaN inhibitor-induced pain syndrome (CIPS) in subsets of transplant patients (Grotz et al., 2001; Fuji et al., 2006; Noda et al., 2008; Prommer, 2012; Azzi et al., 2013). Other studies found downregulation of CaN in the dorsal horn after rat sciatic nerve ligation (Miletic et al., 2002; Miletic et al., 2011; Miletic et al., 2013; Miletic et al., 2015). Intrathecal injection of CaN protein was sufficient to attenuate the resulting hyperalgesia (Miletic et al., 2013). Although these studies link reduced CaN activity and persistent pain syndromes, the downstream ion channel targets responsible for the underlying hyperexcitability are not known. A recent study reported upregulation of the endogenous regulator of CaN 1 (RCAN1) in rat spinal cord after SCI (Wang et al., 2016). RCAN1 upregulation typically inhibits CaN, which might affect CaN-dependent modulation of ion channels in DRG neurons (Davies et al., 2007; Liu et al., 2009; Li et al., 2011; Patel et al., 2015). However, until recently, there was no evidence for such a maladaptive change in the DRG.

In this study, we found that CaN inhibition prolonged the AP of small-diameter DRG neurons, which is associated with attenuation of the Kv3.4 current. Concomitantly, CaN inhibition slowed Kv3.4 fast inactivation. SCI induced changes in the Kv3.4 current of small-diameter DRG neurons that resemble those induced by CaN inhibition. Elucidating a mechanism, first, we found that modulation of Kv3.4 by CaN depends on four serines within the NTID of Kv3.4. Therefore, CaN is a phosphatase that counters PKC action on the NTID. Second, we determined that RCAN1 is upregulated in the DRG after SCI and that overexpression of RCAN1 in naive DRG neurons recapitulates the effects of CaN inhibition on the Kv3.4 current and the AP. We propose a pathway that links CaN and RCAN1 to functional modulation of the Kv3.4 channel in DRG neurons. Dysregulation of this pathway after SCI could be a major factor contributing to the peripheral hyperexcitability associated with SCI-induced persistent pain.

Materials and Methods

**DRG dissociation.** DRG neurons were harvested and dissociated from adult female Sprague Dawley rats (Taconic) as described previously (Ritter et al., 2012). After removal of the spinal cord, the cervical DRGs C3–C7 were harvested. Nerve fibers coming from the DRGs were trimmed and ganglia placed in Hank’s buffered saline solution (HBBS) + 10 mM HEPES. DRGs were then treated with collagenase (1.5 mg/ml in HBBS + 10 mM HEPES solution) for 30 min at 37°C, followed by 2 washes with HBBS + 10 mM HEPES. This was followed by an incubation with trypsin (1.0 mg/ml in HBBS + 10 mM HEPES) for 30 min at 37°C. DRGs were then placed in 1-15 Leibovitz medium supplemented with 10% fetal bovine serum, 38 mM glucose, 2 mM l-glutamine, 24 mM NaHCO3, and 2% penicillin–streptomycin and mechanically dissociated with a fire polished Pasteur pipette. Dissociated DRG neurons were plated on glass polyornithine-coated coverslips, incubated at 37°C, and used for electrophysiological experiments within 2–3 h after harvesting.

For embryonic cultures, DRGs were dissected out from embryonic day 18 (E18) rat embryos, washed once in HBBS, and incubated at 37°C with 0.25% trypsin for 10–15 min. Trypsin-treated DRGs were resuspended in L15 medium plus 10% horse serum and then mechanically triturated with a fire polished pipette. Dissociated rat DRG neurons (~7.5 × 10^5 cells) were transfected with various constructs by nucleofection (Lonza) using reagent P3 and the CU-133 program and then sets of ~30,000 cells were cultured on 18 mm glass coverslips coated with 10 μg/ml poly-β-lysine and 10 μg/ml laminin in F12 medium (with the N3 supplement, 40 μg/ml glucose, and 25 ng/ml NGF).

**Heterologous expression in mammalian cells.** Chinese hamster ovary (CHO) cells were grown on 10 cm tissue culture dishes in F12 medium containing 2 mM l-glutamine, 10% heat-inactivated fetal bovine serum, and 0.1% penicillin–streptomycin. Cells were kept in a humidified incubator at 37°C (5% CO2) and passed every ~4 d. After ~30 passages, cells were harvested. For heterologous expression, cells were cotransfected with Kv3.4 and enhanced green fluorescent protein (EGFP) cDNAs using Lipofectamine 2000 (Thermo Fisher Scientific) according to the manufacturer’s instructions. Transfections were performed in 60-mm plates and then plated onto 5 mm glass coverslips coated with polyornithine. Under epifluorescence conditions, EGFP expression allowed identification of successfully transfected cells.

**Transfection and staining of acutely dissociated DRG neurons.** DRG neurons were isolated and plated as described above. Two hours after plating on glass coverslips in a 35 mm dish, cells were transfected with RCAN1-Flag (gift from Jeffery Molkentin, University of Cincinnati; Addgene plasmid # 65413) and EGFP plasmids or with an EGFP plasmid alone. Transfections were done using Qiagen’s Effectene transfection reagent following the instructions provided by the manufacturer. Briefly, 150 μl of EC buffer was combined with 1 μg of DNA (1 μg of EGFP or 0.75 μg of RCAN1-Flag + 0.25 μg of EGFP) and 8 μl of the enhancer reagent. After a 5 min incubation, 25 μl of Transfectene reagent was added. Subsequently (10 min later), this solution was added to DRG neurons, which were then incubated at 37°C for 4–6 h. The medium was then swapped out and replaced every 5–15 h until DRGs were recorded 36–48 h after transfection. Cells not used for electrophysiology were fixed with 4% PFA for 10 min and proceeded to be blocked with PBS containing 0.2% Triton X-100 and 10% goat serum for 1 h at room temperature. Coverslips were then stained overnight at 4°C in PBS containing 0.1% Tween 20, 1% Triton X-100, and rabbit anti-Flag antibodies 1:1000 (Abcam catalog #ab1162 RRID:AB_298215). Coverslips were then washed with PBS and a coverslip was mounted on a slide using DAPI Prolong Gold Antifade Reagent.
Electrophysiological recordings. As reported previously, all electrophysiological experiments with acutely dissociated DRG neurons were performed at room temperature (20–24°C) (Ritter et al., 2012). A PIP5 micropipette puller (HEKA Instruments) was used to pull patch electrodes made from borosilicate thin-walled patch glass (Warner Instrument). Electrodes tip resistances ranged between 1 and 3 MΩ after fire polishing. The Multiclamp 700B amplifier (Molecular Devices) was used to amplify signals and a computer running Clampex 10.5 (Molecular Devices) was used for data acquisition and storage. Signals were digitized at 10 kHz (Digidata 1440; Molecular Devices) and OriginPro version 9.1 (Origin Laboratory) was used for data processing and analysis. The peak chord conductance (\( G_p \)) in cell-attached patches was calculated using the following relation:

\[
G_p = \frac{I_p}{(V_m - \text{rev})},
\]

where \( I_p \) is the peak current, \( V_m \) is the membrane potential estimated as stated above, and \( \text{rev} = -73 \) mV, as estimated previously (Ritter et al., 2012). To normalize the peak chord conductance (\( G_{pmax} \)), the maximum peak chord conductance (\( G_{pmax} \)) was calculated from the following best-fit fourth-order Boltzmann function:

\[
G_p(V_m) = \left[ G_{pmax} \left( 1 + \exp((V_m - V_{1/2})/k) \right) \right]^4,
\]

where \( V_{1/2} \) is the activation midpoint voltage of a single subunit and \( k \) is the slope factor. The midpoint voltage (\( V_{1/2} \)) of the \( G_p/V_m \) curve was calculated from the following equation:

\[
V_{1/2} = V_c + 1.67k
\]

The steady-state inactivation parameters were determined from the best-fit to the following simple Boltzmann function:

\[
I_p(V_m) = \left[ I_{max} \left( 1 + \exp((V_m - V_{1/2})/k) \right) \right],
\]

where \( V_{1/2} \) is the midpoint voltage of the curve and \( k \) is the slope factor. In some instances, the sum of two Boltzmann terms yielded a better fit. Therefore, the median voltage was used to describe the midpoint position. Calculation of the double Boltzmann curve along the voltage axis.

Reagents. FK506, CsA, and the CaN auto-inhibitory peptide were purchased from Tocris Biosciences. Stock solutions of FK506 and CsA stocks were made in dimethyl sulfoxide (DMSO) with the final concentrations in the recording solution of 0.5–2%; the CaN auto-inhibitory peptide stocks were made in nuclease-free H2O. The rat Kv3.4 plasmid used for CHO cell transfection was a gift from Olaf Pongs (University of Hamburg, Germany).

Analysis of electrophysiological experiments. Clampfit version 10.5 (Molecular Devices) and OriginPro version 9.1 (Origin Laboratory) were used for data processing and analysis. The peak chord conductance (\( G_p \)) in cell-attached patches was calculated using the following relation:

\[
G_p = \frac{I_p}{(V_m - \text{rev})},
\]

where \( I_p \) is the peak current, \( V_m \) is the membrane potential estimated as stated above, and \( \text{rev} = -73 \) mV, as estimated previously (Ritter et al., 2012). To normalize the peak chord conductance (\( G_{pmax} \)), the maximum peak chord conductance (\( G_{pmax} \)) was calculated from the following best-fit fourth-order Boltzmann function:

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where \( V_{1/2} \) is the midpoint voltage of the curve and \( k \) is the slope factor. In some instances, the sum of two Boltzmann terms yielded a better fit. Therefore, the median voltage was used to describe the midpoint position.

Animals and SCI model. Adult female Sprague Dawley rats (200–250 g) were purchased from Taconic Farms and treated according to a protocol approved by the Thomas Jefferson University Institutional Animal Care and Use Committee. SCI and laminectomy surgeries were performed as described previously (Nicaise et al., 2012). Animals were anesthetized using a mixture of ketamine (95 mg/kg), xylazine (10 mg/kg), and acepromazine (0.075 mg/kg). A small incision was made in the skin using a mixture of ketamine (95 mg/kg), xylazine (10 mg/kg), and acepromazine (0.075 mg/kg). A small incision was made in the skin using a mixture of ketamine (95 mg/kg), xylazine (10 mg/kg), and acepromazine (0.075 mg/kg). Anesthesia was induced in the Infinite Horizons Impactor (Precision Systems and Instrumentation). A 1 mm impactor tip was centered over the exposed right side of the spinal cord, raised to height of 4 mm, and a 200 kilodyne unilateral contusion delivered (range, 195–205 kilodynes).
muscle and skin were closed in layers and animals were allowed to recover for 2 weeks. Laminectomy controls received all procedures except a contusion. Two weeks after injury, animals were killed by overdose of ketamine (380 mg/kg), xylazine (40 mg/kg), acepromazine (0.3 mg/kg), followed by thoracotomy. The spinal column was then extracted and C3–C7 DRG neurons ipsilateral to the SCI were extracted and subjected to the same dissociation protocol described above.

**RNA FISH.** Rats that had received either SCI or laminectomy underwent a thoracotomy followed by cardiac perfusion with 4% paraformaldehyde. C4 –C6 DRGs ipsilateral to the injury were then removed and sectioned at 30 μm on a cryostat before being mounted on glass slides. Sections were then stained with Quasar 570 tagged antisense oligonucleotides against rat RCAN1 mRNA. All reagents used for this procedure, including antisense probes, were obtained commercially (Custom Stellaris FISH, Biosearch Technologies). The simultaneous FISH (antisense RCAN1 probes) and immunofluorescence (anti-pan-cadherin antibodies) protocol used for the staining procedure was performed according to the manufacturer’s instructions.

**DAPI with Prolong Gold Antifade Reagent.** DAPI with Prolong Gold Antifade Reagent was used after the secondary antibody incubation, eliminating the need for sealing coverslips with clear nail polish. Z-stacks were obtained using a Nikon C1 Plus confocal microscope at the Bioimaging Facility of the Sidney Kimmel Cancer Center (NIH Cancer Center Core Grant 5 P30 CA-56036) and images precisely ½ way through each section were taken and analyzed using ImageJ. Puncta were counted within the bounds of individual neurons as shown by pan-cadherin staining.

**Immunofluorescence studies.** Animals were killed 2 weeks after surgery and transcardially perfused with 0.9% saline followed by 4% paraformaldehyde. DRGs from C4 –C6 ipsilateral to the unilateral spinal contusion were harvested and stored in 4% paraformaldehyde (24 h), followed by 0.1M phosphate buffer (24 h), and finally in 30% sucrose containing phosphate buffer (3 d). The tissue was then embedded in tissue freezing medium (Tissue-Tek) and 30 μm sections were cut. Sections were collected on glass slides and stored at −80°C.

For immunofluorescence experiments, we used an antigen retrieval kit from R&D Systems (catalog #CTS015). Sections were washed with TBS (3×/10 min) and placed in 1× retrieval buffer at 90°C for 10 min. Sections were then rinsed with TBS and incubated with 10% goat serum and 0.1% Tween 20 in TBS for 1 h at room temperature. Sections were then incubated with 2% goat serum and 0.1% Tween 20 in TBS containing phosphate buffer (3 d). The tissue was then embedded in tissue freezing medium (Tissue-Tek) and 30 μm sections were cut. Sections were collected on glass slides and stored at −80°C.

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**Figure 1.** Calcineurin inhibition alters the AP waveform in DRG neurons. A, APs before and after exposure to DMSO (left). B, Phase plane plots of the waveforms represented in (A). C, Comparison of the APD90 before and after a 5 min exposure to DMSO (n = 14; DF = 13; tstat = 1.93). D, Comparison of the maximum repolarization rate (Max. repol.) before and after a 5 min exposure to DMSO (n = 14; DF = 13; tstat = 1.11). E–H, Same as A–D except for the exposure to 10 μM FK506 (n = 15; DF(APD90) = 14; tstat(APD90) = 3.51; DF(Max. repol.) = 14; tstat(Max. repol.) = 7.21). I–L, Same as A–D except for the exposure to 50 μM intracellular CaN inhibitory peptide (n = 6 for K and n = 6 for L; DF(APD90) = 5; tstat(APD90) = 3.97; DF(Max. repol.) = 5; tstat(Max. repol.) = 4.75). p-values are indicated in the graphs (Student’s paired t test).
perature before a final wash. Coverslips were added along with Pro- long Gold Antifade Reagent with DAPI (Sigma-Aldrich). Confocal microscopy was performed at the Bioimaging Facility of the Sidney Kimmel Cancer Center (NIH Cancer Center Core Grant 5 P30 CA- 56036). Images were then compiled and an individual blinded to the conditions quantified the mean fluorescence of individual cells from each section using ImageJ.

For quantification of differences in membrane staining, ImageJ was used to create fluorescence intensity plots by first tracing a straight line at a random angle through a 2D axis of each cell within a field of view by an individual blinded to the experimental conditions. The peak pan-cadherin mean fluorescence intensity (plasma membrane, two peaks) from a given Z-plane in each group was obtained and the mean fluorescence intensity of CaN was measured at the peaks. The CaN signal was then normalized to the peak pan-cadherin signal from that cell and the data were compared between groups (pan-cadherin signals did not differ between SCI and laminectomy groups). For quantification of cytoplasmic versus nuclear staining, borders were drawn around nuclei using the DAPI stain and around cells using the pan-cadherin stain. Mean fluorescence intensity was then gathered from either the nucleus or cytoplasm.

Figure 2. Attenuation of the Kv3.4 current and disruption of its inactivation profile upon inhibition of CaN in DRG neurons. A, Time course of Kv3.4 peak current (I_p) at 1 min intervals after DMSO wash-in (left). From individual patches, the data point at 1 min is the average I_p from five sweeps before exposure to DMSO and subsequent data points are normalized snapshots of the I_p at the indicated times. The I_p is normalized to the average I_p from the first five sweeps before exposure to DMSO. The arrow represents the approximate time point of compound administration. The graph depicts the mean ± SEM from multiple patches (n = 12). Representative Kv3.4 current traces at baseline and after exposure to DMSO (center) are shown. Scaled currents are also shown (right). Patches were held at −100 mV before a step to +30 mV for 1 s followed by a test pulse to +80 mV for 500 ms before returning to −100 mV. The start-to-start interval was 12 s. B, Same as A but with 1 μM FK506 (n = 11). C, Same as A but with 10 μM CsA (n = 11). The Student’s paired t test was used to evaluate statistically significant differences between the data at the 2 min time point and subsequent data points. *p < 0.05. D, E, Overlay of representative Kv3.4 currents and the corresponding best-fit exponential to the decay phase (blue line) before (left) and after (right) exposure to FK506 and CsA. F, Dot plot comparing fold change in the inactivation time constant upon exposure to DMSO (n = 14), FK506 (n = 10; DF = 1; χ² = 1.03), and CsA (n = 11; DF = 1; χ² = 4.86). The horizontal black line represents the median value. The p-value is indicated in the graph (Kruskal–Wallis ANOVA).
and a ratio was calculated. Graphs and corresponding statistical analyses were produced in Origin Pro version 9.1 (OriginLab).

Before DRG transfections, both the expression of the RCAN1-Flag construct coupled with a testing of the anti-RCAN1 antibody were done in HEK293T cells. Cells were either mock transfected or transfected with the RCAN1-Flag construct. Individual coverslips from both groups were then stained with the rabbit anti-RCAN1 antibodies (1:1000, gift from Dr. Rejji Kuruvilla, Johns Hopkins University) and rabbit anti-Flag antibodies 1:1000 (Abcam catalog #ab1162, RRID:AB_298215), respectively. Only those cells transfected with the RCAN1-Flag construct and not mock transfected showed immunoreactivity to either the anti-Flag antibody or the anti-RCAN1 antibody.

Statistical analysis. Student’s paired t test was used to evaluate paired datasets. The one-way ANOVA test was used to compare independent datasets if data passed tests for normality and equal variance (as determined by Levene’s test). Otherwise, the Wilcoxon–Mann–Whitney or Kruskal–Wallis ANOVA tests were used. The Kolmogorov–Smirnov test was used to compare cumulative frequency histograms. Details of the statistical analysis are provided in the corresponding figure legends.

Results
CaN inhibitors broaden the AP waveform in putative nociceptors
Previously, we found that Kv3.4 siRNA prolongs the AP and decreases the maximum rate of repolarization in small-diameter DRG neurons (Ritter et al., 2012). To determine whether a similar relation exists after inhibition of CaN, we first examined the effects of two well established CaN inhibitors (FK506 and the auto-regulatory peptide inhibitor) on the AP of small-diameter DRG neurons (21.5+/–0.4 μm, n = 35 somata; see Materials and Methods; Table 1). The vehicle (2% DMSO) did not affect AP amplitude, APD90, or the maximal rates of depolarization and repolarization (Fig. 1A–D). Consistent
with the broadening, the maximum rate of AP repolarization decreased to 77% of control (Fig. 1H, Table 1). In addition, the AP amplitude and maximum rate of AP depolarization were modestly reduced upon exposure to FK506 (Table 1). FK506 inhibits CaN indirectly through its binding to FKBP12 and might modestly reduced upon exposure to FK506 (Richter et al., 1995; Ahn et al., 2007; Sachewsky et al., 2014). To verify that the effects of FK506 on the AP waveform resulted from inhibition of CaN, we also examined the effects of the CaN auto-regulatory peptide, which inhibits the phosphatase directly (Hashimoto et al., 1990). This peptide, however, does not cross the cell membrane. Therefore, peptide freshly chewsky et al., 2014). This peptide, how-
 ever, does not cross the cell membrane. Therefore, peptide freshly dissolved in the intracellular solution at a final concentration of 50 μM was included in the patch pipette and delivered into the neuron upon establishing the whole-cell configuration. The first AP was then recorded 10 min later (“control”) and subsequent recordings were taken at intervals of 5 min (at least two more AP recordings after the initial 10 min control). The effects of the CaN auto-regulatory peptide on the AP waveform were qualitatively similar to those of FK506 (Fig. 1I–L, Table 1). It also significantly prolonged the AP (41% longer at APD90) and slowed the maximum rates of depolarization and repolarization (80% and 71% of control, respectively). In addition, it had a modest depolarizing effect on the afterhyperpolarization (ΔV = +4.8 mV). Both FK506 and the auto-regulatory peptide inhibitor had no effect on the resting membrane potential, input resistance, or AP threshold (Table 1). Overall, these results demonstrate that CaN inhibition modulates the AP waveform by consistently slowing the maximum rates of depolarization and repolarization, which broadens the AP. Focusing on the Kv3.4 channel as a major regulator of AP repolarization in small-diameter DRG neurons (Ritter et al., 2012), we investigated the effects of CaN inhibition on the Kv3.4 current.

CaN inhibition attenuates the Kv3.4 current and slows its fast inactivation profile
To determine whether attenuation of the Kv3.4 current contributes to the modulation of the AP by CaN inhibition, we examined cell-attached patches from acutely dissociated small-diameter DRG neurons as described previously (see Materials and Methods). Under these conditions, we have previously observed robust macroscopic Kv3.4 currents exhibiting fast inactivation (Ritter et al., 2012). At intervals of 12 s (start to start), we evoked the current by a step depolarization to +80 mV after a 1 s conditioning pulse to −140 mV. The normalized test pulse peak current is compared in the figure (paired Student’s t-test; n = 18; DF = 17; t = 2.92 and 23; DF = 22; t = 1.53 patches, respectively). F, Combined data of the peak current change in A–E to compare the effects of CaN inhibitors and DMSO globally. The p-value is indicated in the graph (one-way ANOVA; n = 42 and 29, respectively; DF = 1; F = 18.01).

Figure 4. Attenuation of the total outward K+ current upon inhibition of CaN in DRG neurons and CHO cells. A–C, Paired scatter plot of the maximal peak outward K+ currents recorded before and 10 min after applying DMSO, CsA, or FK506 to DRG neurons. Currents were evoked by a test pulse to +80 mV after a 20 s conditioning pulse to −140 mV. The normalized test pulse peak current is compared in the figure (paired Student’s t-test; n = 10; DF = 9; t = 2.92 and 9; DF = 8; t = 2.46 patches, respectively). D, E, Paired scatter plot of the maximal peak outward K+ currents recorded before and 10 min after washing DMSO or FK506 onto CHO cells expressing Kv3.4. Currents were evoked by a test pulse to +70 mV after a 20 s conditioning pulse to −140 mV. The normalized test pulse peak current is compared in the figure (paired Student’s t-test; n = 18; DF = 17; t = 2.92 and 23; DF = 22; t = 1.53 patches, respectively). F, Combined data of the peak current change in A–E to compare the effects of CaN inhibitors and DMSO globally. The p-value is indicated in the graph (one-way ANOVA; n = 42 and 29, respectively; DF = 1; F = 18.01).
induced by recombinant Kv3.4 channels heterologously expressed in CHO cells, whereas the vehicle had little to no effect on the peak current or the time constant of inactivation (Fig. 3). The median fold change and range were as follows: DMSO 0.99 (0.21, 3.18) and FK506, 4.41 (0.58, 51.49).

As mentioned previously, the test pulse that evokes the Kv3.4 currents is delivered from a conditioning pulse to /H1100230 mV. Therefore, it was important to verify that the total outward K/H11001 current was reduced due to the attenuation of Kv3.4. The /Imax,a t a hyperpolarized conditioning voltage that is expected to recruit all available Kv3.4 channels (i.e., /H11002140 mV), was not affected by the vehicle, but was significantly reduced in the presence of FK506 or CsA in DRG neurons (/H1101180% of control; Fig. 4).

We confirmed this result with the recombinant Kv3.4 heterologously expressed in CHO cells (Fig. 4D, E).

Phosphorylation sites within the Kv3.4 NTID are responsible for Kv3.4 dysfunction induced by CaN inhibition
PKC-dependent phosphorylation at four sites (S8, S9, S15, and S21) within the Kv3.4 NTID, and the resulting conformational change, disrupt fast N-type inactivation of the Kv3.4 channel (Covarrubias et al., 1994; Beck et al., 1998; Antz et al., 1999). We investigated whether phosphorylation of these sites might also underlie Kv3.4 current attenuation in response to CaN inhibition. Phosphorylation and dephosphorylation of the NTID might occur under basal conditions. The balance between the basal activities of PKC and CaN would then determine the phosphorylation state of the NTID. Therefore, dephosphorylation of these sites by CaN would be necessary, not only to preserve the fast inactivating profile of the Kv3.4 current, but also to maintain a robust Kv3.4 current in DRG neurons. To investigate a novel role of the aforementioned phosphorylation sites, we tested the effect of CaN inhibition by FK506 on the current generated by the Kv3.4 channel with all four N-terminal serines mutated to alanine (Fig. 5).

The vehicle and FK506 results were indistinguishable. There was no significant change between the alanine mutant groups when using paired data within one group (i.e., alanine mutant patches exposed to FK506 over time) or when comparing datasets at dis-
distinct time points between the groups (i.e., DMSO vs FK506 exposed patches at a given time point). Furthermore, the changes in macroscopic inactivation of the mutant Kv3.4 current induced by exposure to either vehicle or FK506 were also generally small or negligible (Fig. 5D). In a few instances, upon exposure to either vehicle or FK506, spontaneous disruption of inactivation could have resulted from previously described cysteine oxidation in the absence of NTID phosphorylation sites (Fig. 5D) (Ruppersberg et al., 1991; Duprat et al., 1995). These results demonstrate that Kv3.4 NTID phosphorylation sites play a novel role in maintaining a robust basal level of the Kv3.4 current in a manner that depends on CaN activity.

SCI induces Kv3.4 dysfunction that mirrors the effects of CaN inhibition

To assess the role of CaN inhibition on the SCI-induced dysregulation of the DRG Kv3.4 current, we first examined the properties of the Kv3.4 current in cell-attached patches from small-diameter DRG neurons after either a laminectomy (sham surgery control) or a unilateral cervical contusion that results in persistent neuropathic pain-related behaviors (Materials and Methods; Fig. 6). As reported previously, we found that the Kv3.4 current undergoes modulation of fast inactivation under SCI conditions (Ritter et al., 2015a). Whereas the neurons from the laminectomy group mainly express robust fast inactivating outward currents, those from the SCI group displayed highly variable outward currents with more diverse phenotypes: fast inactivating, slow inactivating, and noninactivating (Fig. 6A–C). Patches with negligible outward currents were also observed in this group (Fig. 6D). To assess quantitatively the degree of inactivation of the currents, we measured the ratio of the peak outward current to the magnitude of the current at 75 ms, the time at which most of the fast inactivating Kv3.4 current under basal conditions would be inactivated ($I_{P}/I_{75}$; Fig. 6E). Demonstrating that modulated Kv3.4 currents in the SCI group have disrupted fast inactivation, we found that the $I_{P}/I_{75}$ was significantly reduced compared with the ratio from currents in the laminectomy group (Fig. 6E). Then, to determine whether the slowest inactivating currents were associated with small peak currents, we compared currents with $I_{P}/I_{75}$ at least 1 SD greater than the mean (currents dominated by fast inactivation) to those with $I_{P}/I_{75}$ at least 1 SD smaller than the mean (currents dominated...
SCI induces minimal changes on Kv3.4 biophysical properties. A, Normalized $G_p/G_{\text{max}}$ relation from laminectomy (black) and SCI (red) 2 weeks after surgery. Pulse protocol is as described in Figure 5 legend ($n = 29$ and 27 cells for laminectomy and SCI groups, respectively). Solid lines are best-fit fourth-order Boltzmann functions (Materials and Methods). B, Scatter plot of $V_{1/2}$ values extracted from individual $G_p - V$ relations 2 weeks after laminectomy and SCI. Horizontal bars indicate means. (one-way ANOVA; DF $= 1$; $F = 3.89$) ($V_{1/2}$ mean $\pm$ SEM; laminectomy $= +27.45 \pm 2.67$; SCI $= +20.25 \pm 2.49$). C, Steady-state inactivation curves 2 weeks after laminectomy (black) and SCI (red) ($n = 28$ and 33, respectively). The solid lines are best-fit Boltzmann functions.

Because CaN inhibition modulates the Kv3.4 current of DRG neurons in two major ways that mirror SCI-induced dysregulation of this current (smaller peak and reduced inactivation), we hypothesized that SCI might inhibit CaN activity in the DRG. Therefore, we investigated whether CaN expression had been affected by SCI. Immunofluorescence staining of the catalytic subunit, CaN $\alpha$, demonstrated localization at the plasma membrane, diffuse cytoplasmic staining, and punctate nuclear staining (Fig. 8A). Although the overall DRG immunofluorescence intensity per cell was 21% greater in the SCI group (laminectomy median and range (arbitrary units) $= 338$ (84, 700); SCI median and range (arbitrary units) $= 428$ (42, 853), $p = 5.6 \times 10^{-13}$), we observed no significant difference in plasma membrane expression, the nuclear to cytoplasmic ratio, or the number of nuclear puncta per cell per section (Fig. 8B–E). Therefore, SCI does not induce CaN A$\alpha$ downregulation or redistribution in DRG neurons. Based on the results above, we reasoned that, rather than changing CaN protein expression or localization, SCI may induce upregulation of endogenous modulators of CaN activity. Specifically, upregulation of RCAN1 in the DRG could inhibit CaN after SCI. RCAN1 is abundantly expressed in the nervous system and is upregulated in many neurological disorders, including...
Alzheimer’s disease, stroke, and Down syndrome (Cho et al., 2008; Patel et al., 2015; Kipanyula et al., 2016; Lee et al., 2016). After SCI, RCAN1 is also upregulated in the rat spinal cord (Wang et al., 2016). To test the role of RCAN1 in the SCI-induced dysregulated Kv3.4 function in the DRG, we first investigated the effect of unilateral cervical SCI on RCAN1 expression at the mRNA and protein levels in DRG sections ipsilateral to the injury using FISH and RCAN1 immunostaining, respectively (for antibody validation, see Materials and Methods). Whereas DRG satellite cells displayed negligible RCAN1 mRNA expression, we observed distinct mRNA puncta in the somata of individual DRG neurons throughout the tissue (Fig. 9). Two weeks after surgery, we found significantly increased RCAN1 mRNA levels in DRG neurons (56% increase; \( p = 0.008 \)) from rats in the SCI group compared with the laminectomy control group (Fig. 9A, C). Furthermore, we found a corresponding increase in RCAN1 immunoreactivity (27% increase; \( p = 3.9 \times 10^{-15} \)) in individual DRG neurons from the SCI group (Fig. 9B, D). These results demonstrate that SCI-induced RCAN1 upregulation could be responsible for Kv3.4 dysfunction through inhibition of CaN.

**RCAN1 overexpression in DRG neurons attenuates the Kv3.4 current, disrupts Kv3.4 inactivation, and alters AP trajectory**

To determine whether RCAN1 could be responsible for the SCI-induced attenuation of the Kv3.4 current in DRG neurons, we overexpressed a RCAN1-Flag construct in these neurons (for construct validation, see Materials and Methods). Once immunostaining confirmed the expression of RCAN1-Flag (Fig. 10A), we investigated Kv3.4 currents of DRG neurons transfected with EGFP alone or cotransfected with RCAN1-Flag and EGFP (Fig. 10B, C, respectively). Cumulative frequency histograms show that, compared with mock-transfected neurons (EGFP alone), RCAN1-Flag-expressing neurons displayed an excess of smaller Kv3.4 currents (Fig. 10E). In addition, the degree of inactivation (\( I_p/I_{75} \)) was also reduced significantly (Fig. 10D). These observa-
tions recapitulate the effects of CaN inhibitors (FK506 and CsA) and SCI on the Kv3.4 current (Figs. 2, 3). Therefore, there is a plausible link among SCI-induced RCAN1 upregulation, CaN inhibition, and Kv3.4 attenuation accompanied by disrupted fast inactivation.

We then investigated whether the extent of Kv3.4 current attenuation induced by RCAN1 overexpression is sufficient to recapitulate the effects of CaN inhibitors on the trajectory of the AP in DRG neurons (Fig. 1). We found that the APD₉₀ (median and ranges: EGFP, 0.93 ms (0.35, 1.45); EGFP + RCAN1-Flag, 0.97 ms (0.61, 1.08); n = 11 and 14 respectively; Kruskal–Wallis ANOVA; p = 0.38) and the maximum rate of repolarization of the APs (median and ranges: EGFP 34.01 (17.44, 118.58); EGFP + RCAN1-Flag, 32.69 (25.29, 58.42); n = 11 and 14, respectively; Kruskal–Wallis ANOVA; p = 0.44) under control and RCAN1 overexpression conditions were indistinguishable. Therefore, the extent to which acute RCAN1 overexpression in DRG neurons attenuates the Kv3.4 current is insufficient to change

Figure 9. SCI induces upregulation of RCAN1 at the mRNA and protein levels in DRG neurons. A, FISH analysis of RCAN1 mRNA expression (red) in DRGs from rats that received either SCI or laminectomy. DRG sections were counterstained with DAPI (blue) and pan-cadherin (green). B, Immunostaining of RCAN1 protein (green) counterstained with DAPI (blue) and pan-cadherin (red) in DRGs from rats that received either SCI or laminectomy. C, Quantification of RCAN1 mRNA puncta in DRG neurons from rats that received laminectomy or SCI (n = 95 and 121 neurons, respectively; DF = 1; \( \chi^2 = 7.0 \)). D, Quantification of RCAN1 immunofluorescence in DRG neurons from rats that received laminectomy or SCI (n = 264 and 260 neurons, respectively; DF = 1; \( \chi^2 = 61.73 \)). The p-values are indicated in the graphs (Kruskal–Wallis ANOVA; n = 3 and 3 SCI and laminectomy rats, respectively).
repolarization of the AP. However, the coefficients of variation of the APD$_{90}$ (EGFP = 0.41; RCAN1-Flag = 0.20) and the maximum rate of repolarization (EGFP = 0.65; RCAN1-Flag = 0.27) are decreased upon RCAN1-Flag expression, suggesting a reduced ability to regulate the AP.

In the light of this result, we resorted to rat embryonic DRG neurons in culture to optimize the overexpression of RCAN1 and the effects on the Kv3.4 current. Compared with those from adults, embryonic DRG neurons allow improved transfection efficiency and survival after nucleofection. Within ~16 h after nucleofection, we detected a 1.87-fold increase in RCAN1-Flag expression (Fig. 11A). These neurons also expressed a robust high-voltage-activating A-type current with biophysical properties similar to those of DRG neurons from pups and adults (Fig. 11B). Moreover, as shown previously for DRG neurons from pups, Kv3.4 siRNAs nearly abolished the embryonic Kv3.4 current (Ritter et al., 2012). However, compared with the results from adult DRG neurons, RCAN1 overexpression in embryonic DRG neurons had a greater effect on the Kv3.4 current (Fig. 11B, C). Cumulative frequency histograms revealed that, compared with mock-transfected neurons (EGFP alone), RCAN1-Flag expressing neurons displayed a significant excess of smaller
Kv3.4 currents (Fig. 11E). Although these currents were too small to derive inactivation time constants reliably, there were instances of small currents displaying severely disrupted inactivation (Fig. 11D). We then compared the APs from mock-transfected neurons against those from RCAN1-Flag-transfected neurons (Fig. 12). As found with CaN inhibitors, RCAN1 overexpression in embryonic DRG neurons prolonged the APD_{90} (23.4% longer than control; \( p = 0.037 \)) and slowed the maximum rate of repolarization (83.9% of control; \( p = 0.033 \)). These changes are consistent with previously reported effects of Kv3.4 siRNA on the AP of DRG neurons (Ritter et al., 2012) and could be the result of optimizing the attenuation of the Kv3.4 current in embryonic DRG neurons (Fig. 11). In addition, as observed with CaN inhibitors (Table 1), RCAN1 overexpression decreased AP amplitude modestly (control: 100.02 ± 1.94 mV, RCAN1: 92.15 ± 2.32 mV, \( p = 0.011 \)) and slowed the maximum rate of depolarization (control: 59.97 ± 4.82 ms/mV, RCAN1: 44.27 ± 3.78 ms/mV, \( p = 0.014 \)).
Calcineurin-Dependent Modulation of Kv3.4

Discussion

The ionic and molecular bases of peripheral mechanisms of SCI-induced persistent pain remain poorly understood. We have proposed previously that dysfunction of the Kv3.4 channel is a major determining factor of DRG hyperexcitability and the resulting persistent pathological pain in a rat model of cervical SCI (Ritter et al., 2015a, b). In a search for the signaling molecules and the mechanism responsible for SCI-induced Kv3.4 dysfunction, we investigated CaN, which has been implicated previously in pain syndromes associated with immunosuppressants and peripheral nerve lesions (Grotz et al., 2001; Miletic et al., 2002; Prommer, 2012; Miletic et al., 2015). The results demonstrate the following: (1) exposure of small-diameter DRG neurons to distinct CaN inhibitors prolongs the AP and decreases the maximum AP repolarization rate; (2) similar treatments induce attenuation of the Kv3.4 current and disrupt the current’s fast inactivation profile in DRG neurons and a heterologous expression system; (3) attenuation of both peak currents and inactivation is eliminated by mutating NTID serines to alanine; (4) SCI induces Kv3.4 current alterations that mimic the effects of CaN inhibitors on this current; (5) SCI also induces upregulation of RCAN1 in DRG neurons, which has been shown to inhibit CaN activity; (6) overexpression of RCAN1 in naive DRG neurons is sufficient to induce Kv3.4 current alterations with features that recapitulate the effects of CaN inhibitors; and (7) overexpression of RCAN1 in naive DRG neurons is also sufficient to prolong the AP and slow the maximum AP repolarization rate. Altogether, these observations suggest that CaN is a key player in the phosphorylation-dependent modulation of the Kv3.4 channel NTID, which is disregulated upon SCI. Moreover, novel SCI-induced RCAN1 upregulation in the DRG could be the primary alteration that drives Kv3.4 dysfunction and its effects on pain signaling.

Novel role of NTID phosphorylation sites on the modulation of Kv3.4 functional expression by CaN

PKC activation eliminates Kv3.4 fast N-type inactivation and, consequently, it initially increases the peak current (Covarrubias et al., 1994). The underlying mechanism of this modulation involving phosphorylation of four NTID serines is well established (Covarrubias et al., 1994; Beck et al., 1998; Antz et al., 1999) and it is not limited to heterologous expression systems. It exists in small-diameter DRG neurons, where it helps to shape the repolarization phase of the action potential (Ritter et al., 2012) and thus may regulate excitatory synaptic transmission in nociceptive areas of the spinal cord dorsal horn. Showing that CaN inhibition also disrupts fast Kv3.4 inactivation in this study has two major implications. First, it suggests that PKC and CaN represent the kinase and phosphatase that determine the phosphorylation status of the Kv3.4 NTID. Second, it suggests the presence of basal PKC activity and that the role of CaN under basal conditions is to maintain the NTID of Kv3.4 mostly dephosphorylated. Therefore, the Kv3.4 current would remain the fast inactivating A-type. Here, we determined a link between phosphorylation-dependent elimination of fast Kv3.4 inactivation and Kv3.4 current attenuation induced by CaN inhibitors. If PKC is activated further under conditions that favor NTID phosphorylation (i.e., CaN inhibition), then the Kv3.4 current would exhibit time-dependent modulation in two phases. During the early phase, the peak Kv3.4 current increases due to elimination of fast inactivation and persistent phosphorylation during a second late phase would induce slow current attenuation. Elucidating a mechanism, we discovered that the PKC phosphorylation sites within the NTID (S8, S9, S15, and S21) are also involved in the late attenuation of the Kv3.4 current induced by CaN inhibition. Such a novel role suggests that phosphorylation of the NTID might additionally modulate functional surface expression (trafficking and protein stability) and/or conductance. Although more complex scenarios are possible (e.g., involving ancillary subunits), the consistency of the electrophysiological effects with various CaN inhibitors in DRG neurons and confirmation of the modulations in a heterologous system help to establish CaN as a major modulator of the expression and function of the Kv3.4 channel through its effects on phosphorylation sites in the NTID.

In an alternate scenario, slowing of inactivation could have resulted from a Kv3.4-independent, CaN inhibitor-induced, delayed rectifier current. To test this possibility directly, we examined DRG neurons treated with Kv3.4 siRNA. Transfection of E18 DRG neurons with Kv3.4 siRNA nearly abolished the Kv3.4 current and treating these neurons with 1 μM FK506 induced no novel slow inactivating current. Therefore, in addition to current attenuation, CaN inhibition also induces the slowing of Kv3.4 inactivation as explained above.

Novel role of RCAN1 in maladaptive modulations induced by SCI in the DRG

SCI also induces simultaneous disruption of inactivation and current attenuation of the Kv3.4 current in small-diameter DRG neurons, which prompted us to investigate a possible role of CaN downregulation in the pathophysiology of SCI. However, we found that the expression of CaN Aα throughout the DRG was not downregulated by SCI. Instead, we discovered that the regu-
latory protein RCAN1 is upregulated in the cervical DRGs from SCI rats at both the mRNA and protein levels. RCAN1 is an essential endogenous regulator of CaN that is highly expressed in mammalian excitable tissues and inhibits CaN activity through a direct interaction (Hoeffer et al., 2007; Liu et al., 2009; Li et al., 2011; Shin et al., 2011). This inhibitory action is typically associated with upregulation or overexpression of RCAN1. Several studies have observed inhibition of NFAT translocation to the nucleus after RCAN1 upregulation, which confirms its negative impact on CaN activity (Davies et al., 2007; Liu et al., 2009; Li et al., 2011). In light of the available RCAN1 literature and our discovery, it is remarkable that overexpression of RCAN1 in DRG neurons is sufficient to induce a robust dual modulation of Kv3.4 channels that closely recapitulates the effects of pharmacological CaN inhibitors (in DRG neurons and CHO cells) and the effects of SCI. Therefore, given that SCI induces RCAN1 upregulation in DRG neurons, we can conclude that RCAN1 is a significant novel player in the pathophysiology of SCI. Through inhibition of CaN, RCAN1 upregulation might then ultimately be responsible for the attenuation of the Kv3.4 current with disrupted inactivation in small-diameter DRG neurons. Our previous work revealed that such a negative effect can be a major contributing factor to the hyperexcitability of small-diameter DRG neurons that is a basis of neuropathic pain induced by SCI (Ritter et al., 2015a). A recent study reported RCAN1 upregulation in the spinal cord dorsal horn after SCI (Wang et al., 2016). However, it was not clear whether the reported upregulation occurred in spinal neurons or synaptic terminals from primary sensory inputs in the dorsal horn. In addition, this study did not investigate the effects of RCAN1 upregulation on molecular targets that could alter neuronal excitability. Nevertheless, it supports independently a significant role of RCAN1 in the pathophysiology of SCI.

Pharmacological inhibition of CaN broadens the AP and slows its maximal rate of repolarization in DRG neurons, resembling the effects of Kv3.4 siRNA (Ritter et al., 2012). However, we found that overexpression of RCAN1 in adult neurons does not produce these effects. Rather, the effects of RCAN1-Flag overexpression resemble previously reported effects of SCI on Kv3.4 currents and DRG APs. We found that AP shape remained unchanged after SCI upon partial inhibition of the Kv3.4 current accompanied by slowing of inactivation (Ritter et al., 2015a). Contrary to what is normally observed under naive conditions, there was no AP shortening in response to PKC activation because, after incomplete attenuation, the remaining Kv3.4 current after SCI is already slow inactivating and cannot undergo further modulation of inactivation by PKC (Ritter et al., 2015a). Therefore, the magnitude of Kv3.4 current attenuation might ultimately determine whether there is broadening of the AP. Supporting this explanation, we found that optimized overexpression of RCAN1 in embryonic DRG neurons not only produces very robust attenuation of the Kv3.4 current exhibiting disrupted inactivation, it also prolongs the AP and slows the maximum rate of repolarization. Nucleofection of embryonic DRG neurons resulted in an 87% increase in RCAN1 expression, which is significantly greater than a more modest increase induced by SCI (27%). We do not know whether the SCI-induced RCAN1 upregulation alone is sufficient to generate the observed SCI-induced alterations in the Kv3.4 current and whether other factors are also involved. Most likely, SCI, like other injuries, induces multiple changes that could also affect Kv3.4 function. For instance, other pain injury models suggest changes that implicate PKCs in the DRG and spinal cord (Malmberg et al., 1997; Hucho and Levine, 2007). In our current working model, we propose that concerted PKC activation and CaN inhibition (with the latter resulting from RCAN1 upregulation) might lead to the level of Kv3.4 phosphorylation responsible for the observed changes after SCI. This is a likely scenario because promoting CaN inhibition experimentally upon RCAN1 overexpression alone seems able to match the SCI-induced Kv3.4 phosphorylation status, which could be responsible for the functional alterations. A possible contribution of novel factors, however, remains open.

**Additional effects of CaN inhibition in the DRG**

In addition to prolonging the AP and slowing its maximum repolarization rate, we found that CaN inhibition (induced by inhibitors or RCAN1 overexpression) decreased the AP amplitude modestly and slowed the maximum depolarization rate (Table 1). Decreased AP amplitude might negatively affect high-voltage-activating Kv channels, which could also contribute to broadening of the AP upon CaN inhibition. However, these effects are not observed upon knocking down Kv3.4 in DRG neurons with siRNA, as we reported previously (Ritter et al., 2012). Therefore, we suggest that CaN might additionally modulate voltage-gated Na+ channels, a possibility that is not surprising given the widespread roles of CaN in excitable and nonexcitable tissues (Yakel, 1997; Rusnak and Mertz, 2000; Li et al., 2011).

**Concluding remarks**

The cellular and molecular bases of intractable pain in SCI and CIPS patients are poorly understood. The Kv3.4 channel is a key regulator of AP repolarization in putative DRG nociceptors. Here, we identified Kv3.4, CaN, and RCAN1 as key players in a putative peripheral mechanism of pain sensitization involving novel dysregulation of CaN signaling in DRG neurons. In particular, the results strongly suggest that RCAN1 is implicated in this mechanism through its downstream effects on the modulation of the Kv3.4 channel by CaN. In addition, we discovered that Kv3.4 NTID phosphorylation sites that modulate inactivation are involved in a novel mechanism that modulates Kv3.4 functional expression. These findings suggest tantalizing novel interventions to treat intractable pain more effectively in SCI and CIPS patients.

**References**


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