Role of Decreased Sensory Neuron Membrane Calcium Currents in the Genesis of Neuropathic Pain

Quinn H. Hogan

The pathogenesis of neuropathic pain is incompletely understood and treatments are often inadequate. Cytoplasmic Ca\(^{2+}\) regulates numerous cellular processes in neurons. This review therefore examines the pathogenic contribution of altered inward Ca\(^{2+}\) flux (I\(_{Ca}\)) through voltage-gated Ca\(^{2+}\) channels in sensory neurons after peripheral nerve injury. We reviewed studies that recorded membrane currents through intracellular and patch-clamp techniques, as well as intracellular Ca\(^{2+}\) levels using fluorimetric indicators, and performed behavioral analysis of rodent nerve injury models. Following nerve injury by partial ligation, a response characterized by sustained lifting, shaking, and licking of the paw after sharp mechanical stimulation is a reliable indicator or neuropathic pain. Primary sensory neurons isolated from animals with this behavior show a decrease in high-voltage activated I\(_{Ca}\) by approximately one third. Low voltage-activated I\(_{Ca}\) is nearly eliminated by peripheral nerve injury. Loss of I\(_{Ca}\) leads to decreased activation of Ca\(^{2+}\)-activated K\(^{+}\) currents, which are also directly reduced in traumatized neurons. As a result of these changes in membrane currents, membrane voltage recordings show increased action potential duration and diminished afterhyperpolarization. Excitability is elevated, as indicated by resting membrane potential depolarization and a decreased current threshold for action potential initiation. Traumatized nociceptive neurons develop increased repetitive firing during sustained depolarization after axotomy. Concurrently, cytoplasmic Ca\(^{2+}\) transients are diminished. In conclusions, axotomized neurons, especially pain-conducting ones, develop instability and elevated excitability after peripheral injury. Treatment of neuronal I\(_{Ca}\) loss at the level of injury of the dorsal root ganglion may provide a novel therapeutic pathway.
Neuropathic pain, sensory neurons, and calcium

Nerve injury is a dominant or contributing factor in a wide variety of painful conditions, including persistent pain following thoracic, breast, and amputation surgery, radiculopathy from disc disease, nerve invasion by cancer, trauma, metabolic injury from ischemia or diabetes, infectious conditions such as herpes zoster and AIDS, and complex regional pain syndrome (CRPS) following minor injury. Neuropathic pain is often disabling because of its intensity, lancinating quality, and resistance to currently available treatments.

The pathogenic mechanisms generating hypersensitivity and spontaneous pain following injury of a peripheral nerve are anatomically distributed and complex. Altered neural structure and function have been identified at the peripheral tissues, the dorsal horn of the spinal cord (1-6), and the brain (7). In addition, the primary afferent neuron itself is an important site of changes generating pain. Increased neuronal excitability is evident at the injury site and also in the somata of the injured neurons (8,9). Persistence of membrane instability in the dorsal root ganglion (DRG) neurons isolated from injured animals (10,11) demonstrates that aberrant excitability is intrinsic to the soma of primary afferents. Prolonged afterdischarge following stimulation (5,12) and altered patterns of provoked activity (13) further characterize injured sensory neurons.

Major aspects of neuronal function are regulated by Ca\(^{2+}\), including neurotransmitter release, excitability, neuron growth, differentiation, and death (14,15), as well as the development of plasticity and gene expression (16). A similarly important role for Ca\(^{2+}\) has been established in processes of signaling pain, especially in facilitated pain states. Also, voltage-gated Ca\(^{2+}\) channels (VGCC) modulate pain in clinical and experimental settings (17-24). Due to the importance of this disease condition and the relatively recent attention to membrane Ca\(^{2+}\) currents (I\(_{\text{Ca}}\)) in its pathogenesis, this paper reviews recent studies examining the effects of nerve injury on directly measured I\(_{\text{Ca}}\) and cytoplasmic Ca\(^{2+}\) levels ([Ca\(^{2+}\)]\(_{c}\)) in primary afferent neurons.

Neuropathic models and sensory testing

Examination of the pathogenesis of neuropathic pain has been accelerated by the introduction of rodent models of nerve injury that produce behavior indicative of spontaneous and inducible pain (25). A complete section of a nerve produces spontaneous pain but also an anesthetic limb. Partial injury retains a subset of afferent fibers and results in altered sensory function, including more intense pain during noxious stimulation (hyperalgesia) and pain after normally innocuous stimuli (allodynia). The first widely used model of incomplete peripheral nerve injury was chronic constriction injury (CCI, Figure 1), in which four ligatures of chromic gut suture produce axotomy (transection of the neuronal axon), ischemia, or inflammation (26-28). Within 10 days of injury, animals may demonstrate hyperalgesia and allodynia induced by mechanical and
thermal stimuli (29). These amplified behavioral responses are mediated by a subgroup of surviving afferent fibers, as demonstrated by similar findings after incomplete sciatic transection (30).

After partial nerve injury, axotomized sensory neurons develop substantial phenotypic shifts, including altered membrane channels and receptors, and new sensitivities to chemical stimulation by catecholamines, cytokines, bradykinin, and neurotrophins at both the injury site and proximally in the DRG (10,31). Surviving intact fibers are also exposed to abnormal conditions, due to production of inflammatory mediators from degeneration of disconnected fibers of axotomized neurons sharing the same nerve fascicles (32). The spinal nerve ligation (SNL) mode (33), in which the fifth lumbar (L5) and L6 spinal nerve components of the sciatic nerve are ligated but L4 remains intact, permits separate evaluation of the anatomically distinct axotomized neurons of the L5 DRG and the neighboring L4 neurons (Figure 1).

As in human clinical conditions, animal subjects show variability in the sensory and behavioral consequences of incomplete nerve injury. It is therefore important to distinguish those animals that successfully develop the desired phenotype. Since pain is “an unpleasant sensory and emotional experience” (34), the best we can do in animal experimentation is to record behavior and infer the experience. When a pin is applied to the footpad of a rat with only enough pressure to indent but not puncture the skin, the response is either a brief reflex withdrawal or a hyperalgesic reaction characterized by sustained lifting, shaking, and licking of the paw (35). As the latter response occurs only after true SNL but not sham exposure of the nerve alone, and only on the side ipsilateral to the injury (Figure 2), this may be accepted as an indication of a neuropathic pain state. Other commonly used measures, such as threshold testing for withdrawal from low intensity mechanical stimulation with von Frey fibers or from thermal stimuli, are altered after sham surgery without nerve section and also contralateral to the nerve section (35). Accordingly, we have adopted the complex and sustained hyperalgesia response as an indicator of neuropathic pain.

**IC\textsubscript{a} in injured sensory neurons**

DRG neurons express a variety of VGCCs. High voltage activated (HVA) currents are present in a variety of subtypes (L, N, P/Q, R) that are distinguished by their voltage dependency, kinetics, and pharmacology. The specific roles of these subtypes in DRG cells are not fully established, but currents through N and possibly P/Q channels initiate neurotransmitter release. Low voltage activated (LVA) currents (36), or T currents, inactivate rapidly during sustained depolarization but close (deactivate) slowly after repolarization of the membrane. Because of these features, T-currents account for up to 50% of Ca\textsuperscript{2+} entry (37-40). DRG neurons show definite heterogeneity with respect to HVA Ca\textsuperscript{2+} channels. L-type currents contribute substantially to total IC\textsubscript{a} in small neurons, N-type current is present in all sizes of neurons, and non-L/non-N current (presumptive P/Q and R) is prominent in large (≥40µm) and medium neurons (41). DRG so-
mata are particularly heterogeneous in their expression of T-currents, which are most evident in medium sized (30-40 μm) neurons (42-45).

The importance of \( I_{Ca} \) to the functioning of neurons makes it likely that altered \( I_{Ca} \) may contribute to functional abnormalities that accompany neuropathic pain. Also, analgesia from intrathecal administration of Ca\(^{2+}\)-channel blockers (17-24) raises the possibility that the primary disorder includes overexpression of \( I_{Ca} \). Our initial observations in neurons dissociated from hyperalgesic rats after CCI (46) show that peak \( I_{Ca} \) density is in fact diminished by injury (from 3.1 ± 0.3 to 2.2 ± 0.3 pS/pF in medium neurons, and from 3.9 ± 0.4 to 3.0 ± 0.4 pS/pF in large neurons) using standard patch clamp whole cell recording (47) (Figure 3). The medium-sized neuronal population includes cells that transmit both nociceptive and low threshold sensations, while the large neurons are specific for low threshold sensory modality.

The CCI model does not allow a distinction between direct effects of axotomy and indirect inflammatory mechanisms. Examination of \( I_{Ca} \) specifically in axotomized neurons (L5 after SNL) (48) shows that current loss is present in all neuronal size groups, including the nociceptive small diameter population. Current loss is also evident in the adjacent L4 neurons, but only in the large cell category. These recordings were performed with sustained square wave currents, which lack the kinetic complexity of an actual action potential (AP). Additional recordings of the neuronal current response to voltage commands in the form of an AP (Figure 4) show comparable findings, which assure the pathophysiologic validity of the \( I_{Ca} \) loss.

LVA channels that generate the T-type current have an undefined role in sensory neurons, but may be substantially diminished by injury (49). The peak T-type \( I_{Ca} \) isolated by the elimination of other \( I_{Ca} \) with relevant toxins, is reduced by 60% after CCI, and total LVA Ca\(^{2+}\) influx is reduced by 80%. The mechanism is a depolarizing shift in the voltage dependence of activation and an increase in the rate of channel deactivation and inactivation.

Together, these findings indicate that nerve injury, particularly axotomy, results in a loss of \( I_{Ca} \) in primary sensory neurons that is present after different types of injury, affects the full range of neuron sizes, and includes both LVA and HVA current types. Although this was not our expected result, various other observations indicate a role of decreased \( I_{Ca} \) in generating pain. Norepinephrine, which produces pain when applied to injured nerves, reduces \( I_{Ca} \) and increases excitability of rat DRG somata injured by axotomy (50,51). Also, intrathecal Ca\(^{2+}\) administra-

---

**Figure 3.** Inward high-voltage activated Ca\(^{2+}\) currents measured by patch-clamp recording in dissociated sensory neurons decrease after chronic constriction injury. (A) Sample currents elicited by square-wave voltage commands (bottom) are decreased in an injured neuron (middle) compared to a neuron from a control animal (top). (B) Current-voltage plot of average data for medium sized neurons shows a loss of peak current in injured neurons from animals with neuropathic pain. From Hogan et al (46), with permission.

**Figure 4.** Inward low-voltage activated Ca\(^{2+}\) currents measured by patch-clamp recording in dissociated sensory neurons decrease after chronic constriction injury. (A) Current-voltage plot of average data for medium sized-neurons shows a loss of peak current in injured neurons compared to control neurons. (B) Presentation of a voltage command in the form of an action potential (top) produces current through low-voltage activated Ca\(^{2+}\) channels (bottom) that is substantially reduced in an injured neuron compared with a control neuron. From McCallum et al (49), with permission.
tion is antinociceptive (52). Agents that induce pain, including bradykinin and capsaicin, inhibit DRG ICa (53-55), although this is not necessarily their principal algesic mechanism. Axonal injury in other systems (56,57) depresses ICa. Some of our findings have been confirmed (58), although these authors failed to identify a loss in LVA currents.

**Biophysical response of intact neurons to injury**

Although patch-clamp recording is ideal for evaluation of the performance of a specific membrane channel, dialysis of the cytoplasm by the pipette solution alters the natural internal conditions of the cell and blockade of other currents precludes the natural interactions that dictate the generation of APs. To determine neuronal function in a setting that less disrupts function, we used the intracellular microelectrode technique in intact DRGs, which further avoids disruption by cell dissociation and permits characterization of the neurons by conduction velocity (CV).

Pronounced electrophysiological changes were seen only in L5 neurons following SNL (59). Both Aα/β (fast CV) and Aδ (slow CV) myelinated neuron types show increased AP duration, and decreased afterhyperpolarization (AHP) amplitude and duration (Figure 5A). The AHP duration in neurons with C fibers (CV<1.5m/s) shortens after axotomy. In contrast to the axotomized L5 neurons, neighboring L4 neurons develop no changes in AP duration or AHP dimensions. These parameters are of particular importance in considering mechanisms of elevated pain sensitivity, as a prolonged AP duration may release more neurotransmitter at the first synapse in the spinal cord dorsal horn (60), while reduced AHP dimensions results in increased burst firing and elevated maximal firing rate (61,62).

After axotomy, both Aα/β low-threshold neurons and Aδ presumed nociceptive neurons show resting membrane potential depolarization and a decreased current threshold for AP initiation. Importantly, axotomized Aδ neurons develop increased repetitive firing during sustained depolarization after axotomy (Figure 5B), whereas Aα/β neurons do not. Thus, axotomized neurons, especially pain-conducting ones, develop instability and elevated excitability. Additionally, in the L5 ganglion after axotomy, a novel set of neurons (24% of total) have fast CVs characteristic of myelinated neurons, despite exhibiting long AP durations typical of slowly conducting C-type fibers (Figure 6). The histologic counterpart of these cells may have recently been identified by Hammond et al (63), who described the emergence exclusively in the L5 ganglion after SNL of a novel group of very small neurons that label with NS2 antibody, which identifies myelinated neurons. These cells might thus ex-
hibit the AP features of small neurons but have accelerated CV due to myelination. Overall, our findings indicate a clear pathogenic distinction between the substantially altered axotomized neurons and the less affected neighboring ones.

**Functional consequence of decreased I\(_{\text{Ca}}\)**

I\(_{\text{Ca}}\) is an inward current, so the direct effect of its loss should stabilize the membrane and shorten the AP. However, there are secondary effects through the operation of the Ca\(^{2+}\) admitted through the VGCCs upon Ca\(^{2+}\)-sensitive K\(^+\) channels, which generate I\(_{\text{K(Ca)}}\). These release K\(^+\) from the cell when the [Ca\(^{2+}\)]\(_{\text{c}}\) in their immediate vicinity cause them to enter the open state. In operation, they contribute to the repolarization to the AP and the generation of the AHP (64). The end result of I\(_{\text{Ca}}\) loss will thus be determined by the balance of the direct effect of decreased I\(_{\text{Ca}}\) and the indirect effect of lost stimulation of I\(_{\text{K(Ca)}}\). We examined the effect of selective I\(_{\text{Ca}}\) blockers on dissociated neurons to determine the combined effect of these two processes (48).

The form of a triggered AP was measured before and after ablation of I\(_{\text{Ca}}\) by combined application of toxins. Likewise, presentation of a voltage command in the form an AP was used to record the provoked currents with I\(_{\text{Ca}}\) intact and after ablation (Figure 7). We observed that I\(_{\text{Ca}}\) loss was associated with a prolongation of AP duration and loss of inward current during depolarization, but also loss of outward current during repolarization and the AHP. Thus, loss of I\(_{\text{Ca}}\) as occurs with injury, results in a substantial decrease in outward current. Others have similarly observed a predominant effect of Ca\(^{2+}\) entry in producing outward current (65).

Neuronal function is dictated by the complex interplay of the entire ensemble of membrane channels, which interact with each other through their effect on transmembrane voltage and on [Ca\(^{2+}\)]\(_{\text{c}}\). Using non-dissociated intact DRGs to limit artifact, we therefore examined the role of I\(_{\text{Ca}}\) by manipulating conditions to limit or en-
Our preliminary results confirm that suppressing ICa with cadmium application in the superfusate, Ca2+ withdrawal, or intracellular EDTA delivery decreases AHP dimensions and increases repetitive firing during sustained depolarization (Figure 8). Selective VGCC subtype toxins similarly reduced AHP dimensions and also prolonged the AP duration in the neurons of intact ganglia. Reciprocally, elevating Ca2+ entry with high bath Ca2+ concentrations or application of a Ca2+ ionophore increases the AHP and suppresses repetitive firing. These findings clearly indicate the excitatory consequences of decreased ICa in sensory neurons and support the loss of ICa as a mechanism causing increased excitability after injury.

A model of the role of ICa loss in nerve injury

K(Ca) channels help regulate neuronal excitability by hyperpolarizing the membrane after the AP and by decreasing membrane resistance, making the membrane less excitable in response to depolarizing currents (67). In the normal state (Figure 9), Ca2+ entry through VGCCs during the AP stimulates I(K,Ca) and assures a normal level of excitability. In most neurons, sustained depolarizations result in only a single AP. After injury, decreased ICa results in less IK(Ca) and therefore burst firing.

The frequency-encoded sensory signal is filtered at points of impedance mismatch along the axon, especially at the site in the DRG where the afferent neuron splits into the dorsal root fiber and the T-branch that leads to the neuronal soma (68,69). Since the ability of a neuron to conduct repetitive spikes through this particular site is regulated by Ca2+-sensitive processes (70), we hypothesized that injury and the loss of ICa would modulate spike conduction failure. We found (71) that axotomy causes low threshold neurons, identified by their lack of an inflection on the AP, to develop a prolonged refractory period during paired spike stimulation and a decreased maximal following frequency of tetanic bursts. In contrast, axotomy of nociceptive neu-
rons limited AHP amplification during an impulse train and increased the frequency at which these neurons can fire repetitively during tetanic stimulation. As a result, axotomy increases the ability of putative nociceptors to conduct high frequency trains of APs, whereas injury increases filtering of non-nociceptive afferent sensory traffic. This is important since high frequency bursts are transmitted with increased synaptic reliability while tonic discharge with the same average rate of firing may not successfully induce activity in the postsynaptic neuron (72). Also, burst discharge is particularly effective in producing dorsal horn neuronal plasticity (73), which may play a critical role supporting chronic pain states (74,75).

A direct effect of injury upon Ca²⁺-activated K⁺ channels

Altered function of other ionic membrane channels contributes to the disordered membrane biophysics observed after nerve injury, including substantial changes in voltage-gated Na⁺ and K⁺ channels (76,77). It is therefore possible that the post-injury shift in membrane function includes alteration of K(Ca), not just reduced Ca²⁺ stimulation of the channels. We tested this by maximally stimulating dissociated neurons during patch-clamp recording with high intracellular [Ca²⁺]c and identifying currents sensitive to selective I_K(Ca) blockers (78). This revealed I_K(Ca) with components sensitive to apamin, clotrimazole, and iberiotoxin, indicative of SK, IK, and BK subtypes of I_K(Ca). SNL decreases total I_K(Ca) in axotomized (L5) neurons, but increases total I_K(Ca) in adjacent (L4) DRG neurons. All I_K(Ca) subtypes are decreased by axotomy, but iberiotoxin-sensitive and clotrimazole-sensitive current densities are increased in adjacent L4 neurons after SNL. Thus, the injury has divergent effects on axotomized neurons and adjacent intact neurons, and direct effects on the K(Ca) channel amplify the action of I_Ca loss.

Effect of injury on intracellular Ca²⁺ signaling

The major signal downstream from AP-induced inward Ca²⁺ flux is the critical second messenger [Ca²⁺]c (79,80). AP trains trigger an elevation of [Ca²⁺]c (the Ca²⁺ transient) in the primary afferent neuron that persists seconds to minutes after the membrane activity (81), and thereby provides an integrative and memory process very early in the anatomic pathway of somatic sensory signaling. Virtually all aspects of neuronal function are controlled by the Ca²⁺ regulatory pathway, including synaptic transmission, enzymatic activity, membrane currents, cellular energetics, gene expression, cell differentiation, and death (16,82). All these events have been implicated in the reaction to nerve trauma but the effect of peripheral nerve injury on intracellular Ca²⁺ regulation has not been previously examined.

Intracellular Ca²⁺ level is controlled by balanced interactions of Ca²⁺ storage, release, and extrusion. Most of the Ca²⁺ that enters through VGCC or receptor-operated channels is buffered in the cytoplasm. Free Ca²⁺ in peripheral sensory neurons is tightly regulated to approximately 100 nM, or about 20 000-fold less than the extracellular concentration. However, after SNL injury (Figure 10), axotomized neurons show a depressed resting level for both small nociceptive

![Figure 10](image-url). Injury reduces resting level of cytoplasmic Ca²⁺ ([Ca²⁺]c). Spinal nerve ligation injury reduces [Ca²⁺]c in axotomized fifth lumbar (L5) neurons of both sizes, but only in large neurons from the adjacent L4 population. Small numbers in the bars indicate number of neurons. From Fuchs et al (83), with permission.
and large non-nociceptive categories (83). Only non-nociceptive neurons develop a depressed resting \([Ca^{2+}]_c\), in the adjacent L4 population. Decreased \([Ca^{2+}]_c\) may precipitate cell loss, including programmed cell death by apoptosis (84-86), as has been noted after axotomy (87,88). Resting \([Ca^{2+}]_c\) also regulates receptor-triggered calcium signaling (89-91).

A complex system of \(Ca^{2+}\) sequestration, release, and extrusion regulates \([Ca^{2+}]_c\) and shapes the \(Ca^{2+}\) transient that follows neuronal activation. \(Ca^{2+}\) is pumped out of the cell by plasma membrane \(Ca^{2+}\)-ATPases (PMCAs) and is sequestered into endoplasmic reticulum (ER) stores by sarcoplasmic-endoplasmic reticulum \(Ca^{2+}\)-ATPase (SERCA) channels. The ER also functions as a source that releases \(Ca^{2+}\) into the cytoplasm through channels sensitive to inositol 1,4,5-triphosphate (IP3) and others sensitive to the plant alkaloid ryanodine, thereby known as ryanodine receptors (RyRs). Cytoplasmic accumulation of \(Ca^{2+}\) activates RyRs as well as IP3 channels, accounting for the phenomenon of \(Ca^{2+}\)-induced \(Ca^{2+}\) release (CICR). Depletion of the ER \(Ca^{2+}\) pool activates a voltage-independent plasma membrane channel (store-operated \(Ca^{2+}\) channel, SOCC) that conducts an inward \(Ca^{2+}\) flux, a process termed capacitative \(Ca^{2+}\) entry (CCE), which serves to refill \(Ca^{2+}\) stores (92). Our initial data (93) demonstrates that the activity-induced \(Ca^{2+}\) transient is markedly altered in injured neurons (Figure 11). Specifically, after axotomy, the transient duration and amplitude in nociceptive neurons are significantly diminished, whereas amplitude is increased in axotomized non-nociceptors. In the adjacent L4 neurons, only the transient amplitude is diminished only in nociceptors. These effects are likely attributable to combined effects of decreased \(Ca^{2+}\) load entering the neurons and injury-related disruption of the processes regulating \([Ca^{2+}]_c\).

What causes pain after peripheral nerve injury?

While the processes that produce hyperalgesia after nerve injury are diverse, Figure 12 attempts to explain the potential contributions that follow loss of \(ICa\) in primary sensory neurons. Increased burst firing from decreased AHP inflicts greater nociceptive traffic on the secondary neurons of the dorsal horn, where prolonged AP duration results in greater excitatory neurotransmitter release. Although axotomized neurons (L5 ganglion after SNL) are disconnected from sensory fields, they are nonetheless activated through direct mechanical stimulation during movement, depolarization by circulating and local algogens and inflammatory mediators, and sympathetic activity (8,10,31,94,95). Furthermore, the process of cross-excitation spreads activity among adjacent neurons in the DRG (96). This is critical since most clinical nerve injuries are partial. Afferent traffic from these sources is effectively
amplified by the reduced signal filtering at the T-branch of injured neurons. The intense bursts of activity along this L5 pathway sensitize the spinal dorsal horn to input along intact (eg, SNL L4) pathways (1,73), so that natural stimuli are perceived as more intense. Other processes that do not involve reduced ICa in the axotomized neurons include irritation of intact fibers by inflammation induced by adjacent degenerating neuron segments distal to the axotomy, altered channel and receptor expression in the intact fibers, and anatomic changes in connectivity in the dorsal horn.

Relief of pain in animal models by intrathecal administration of ICa blockers (18,97) may appear to contradict our model and findings. However, the function of Ca2+ signaling in different tissues is distinct, and this analgesia is explained by a blockade of neurotransmitter release in the spinal cord, which is not applicable at peripheral sites. VGCC blockers applied at the SNL injury site have no analgesic effect (97), although topical ICa blockers do decrease pain behavior in other models (98) that have a large inflammatory component. Interpretation of gene knockout studies (99-101) is severely limited by the simultaneous effects of current loss at multiple sites that have divergent roles for ICa.

**DRG is an undeveloped site for chronic pain treatment**

There is a great need for more effective treatment of neuropathic pain that follows peripheral nerve injury. Although critical changes leading to chronic pain reside in the DRG, no treatments for chronic pain after peripheral nerve injury have been devised that target the DRG. Drugs that elevate \([\text{Ca}^{2+}]_o\) for instance Ca2+ ionophores, might be administered directly to the DRG, optimizing drug potency at the effector site while minimizing undesirable CNS and systemic effects. Further, stable genetic transfer via viral vectors may allow regulated expression of VGCC. Clinical methods for administering drugs directly to the selected DRGs are well established (102). Therefore, it can be hoped that better understanding of the peripheral pathophysiology of neuropathic pain might ultimately translate into pain therapy through targeted delivery of drugs or genes to selected DRGs.

**Acknowledgment**

This work was supported in part by grant No. NS-42150 from the National Institutes of Health, Bethesda, Maryland, USA.

**References**

Hogan: Calcium in Neuropathic Pain

6:207:25


38. Livingstone DM, White G. Post-natal development of burst firing behavior and the low-threshold transient calcium current examined using freshly isolated neurons from...


60 Sabatini BL, Regehr WG. Control of neurotransmitter release by presynaptic waveform at the granule cell to Purkinje cell synapse. J Neurosci. 1997;17:3425-35.


Hogan: Calcium in Neuropathic Pain


