Sympathetic Sprouting Near Sensory Neurons After Nerve Injury Occurs Preferentially on Spontaneously Active Cells and Is Reduced by Early Nerve Block

Wenrui Xie, Judith Ann Strong, Huiqing Li, and Jun-Ming Zhang

Department of Anesthesiology, University of Cincinnati College of Medicine, Cincinnati, Ohio

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Xi W, Strong JA, Li H, and Zhang J-M. Sympathetic sprouting near sensory neurons after nerve injury occurs preferentially on spontaneously active cells and is reduced by early nerve block. J Neurophysiol 97: 492–502, 2007. First published October 25, 2006; doi:10.1152/jn.00899.2006. Some chronic pain conditions are maintained or enhanced by sympathetic activity. In animal models of pathological pain, abnormal sprouting of sympathetic fibers around large- and medium-sized sensory neurons is observed in dorsal root ganglia (DRGs). Large- and medium-sized cells are also more likely to be spontaneously active, suggesting that sprouting may be related to neuron activity. We previously showed that sprouting could be reduced by systemic or locally applied lidocaine. In the complete sciatic nerve transection model in rats, spontaneous activity initially reduced by systemic or locally applied lidocaine. In the complete to neuron activity. We previously showed that sprouting could be reduced by systemic or locally applied lidocaine. In the complete sciatic nerve transection model in rats, spontaneous activity initially originates in the injury site; later, the DRGs become the major source of spontaneous activity. In this study, spontaneous activity reaching the DRG soma was reduced by early nerve blockade (local perfusion of spontaneous activity. In this study, spontaneous activity reaching the DRG soma was reduced by early nerve blockade (local perfusion of spontaneous activity in the injured nerve is critical for establishing the more long-lasting pathologies observed in the DRG. Individual spontaneously active neurons, labeled with fluorescent dye, were five to six times more likely than quiescent cells to be co-localized with sympathetic fibers, suggesting a highly localized correlation of activity and sprouting.

INTRODUCTION

In some chronic-pain patients, pain and hyperalgesia are maintained by efferent noradrenergic sympathetic activity and circulating catecholamines (sympathetically maintained pain, SMP) (Roberts 1986) and may be partly responsive to sympathetic blockade, whereas in others, the pain is sympathetically independent (Campbell et al. 1988). SMP may be a component of many painful conditions such as complex regional pain syndrome (CRPS), phantom pain, neuralgias, and herpes zoster. The lack of understanding of the neurophysiological mechanisms by which the sympathetic system invades the peripheral sensory system has hindered progress in the treatment of these painful conditions.

Clinical observations and animal studies have shown that coupling of the sympathetic nervous system and the sensitized sensory nervous system is important for development of SMP (Janig et al. 1996). An abnormally enhanced communication between these two systems may occur under a variety of pathologic conditions. For example, sympathetic stimulation may excite sensory neurons in animals with inflamed peripheral tissue or after peripheral nerve injury (Devor et al. 1994; Xie et al. 1995). Chemical or surgical sympathectomy relieves allodynia and hyperalgesia and improves chronic pain behavior in several animal models (Choi et al. 1994; Kim et al. 1993; Kinnman and Levine 1995; Malmberg and Basbaum 1998; Seltzer and Shir 1991). These observations suggest that increased activity of the sympathetic nervous system may contribute to the enhanced sensitivity to painful stimuli observed in chronic pain models, although the behavioral relevance of sympathetic activity in animal models of pain has been somewhat controversial (e.g., Habler 2000; Kim et al. 1999). Sympathetic-sensory coupling may occur either centrally or peripherally. The dorsal root ganglion (DRG) has been identified as one important site for peripheral sympathetic-sensory coupling (McLachlan et al. 1993). Within the normal DRG, sympathetic axons are only found accompanying blood vessels (Kummer et al. 1990). After peripheral nerve injury, sympathetic efferent fibers extensively sprout into both DRG and spinal nerves. Sprouting fibers sometimes form distinctive basketlike webs—sympathetic “baskets” or tyrosine hydroxylase (TH)-immunoreactivity (IR) rings wrapping around medium and large DRG neurons (Chung et al. 1996; Lee et al. 1998; McLachlan et al. 1993; Ramer and Bisby 1997). Pain induced by localized inflammation of the DRG or mechanical compression of the DRG in the absence of nerve injury can also be accompanied by sympathetic sprouting in the DRG (Chien et al. 2005; Xie et al. 2006). Most recently, sympathetic sprouting was observed in the glabrous skin after partial nerve injury (Grelik et al. 2005; Yen et al. 2006).

Abnormal spontaneous activity is also a common feature of many different animal pain models (Amir 1999; Govrin-Lippmann and Devor 1978; Hu and Xing 1998; Liu et al. 2000; Song et al. 1999; Study and Kral 1996; Xie et al. 2006). Large- and medium-diameter cells, which preferentially acquire sympathetic baskets, also show a much higher incidence of spontaneous activity in various neuropathic pain models. This suggested the possibility that spontaneous activity might be one cause of sympathetic sprouting. In support of this idea, we recently found (Zhang et al. 2004) that systemic application of the local anesthetic lidocaine could reduce sympathetic sprouting in two different pain models, the neumata model (sciatic...
axotomy) originally described by Wall (Wall et al. 1974) and the spinal nerve ligation model (Kim and Chung 1992). The inhibition of sprouting outlasted the duration of lidocaine application by >7 days. Sprouting could also be reduced by local application of lidocaine to the nerve trunk proximal to the injury site in the sciatic axotomy model. In the present study, we examined in more detail the possible connection between spontaneous activity and sympathetic sprouting in the sciatic axotomy model. We determined the effects on sympathetic sprouting of either enhancing the spontaneous activity originating at the injury site, or of preventing spontaneous activity from reaching the DRG by local nerve blockade with tetrodotoxin (TTX). We also conducted histological analyses on identified individual neurons, to determine whether spontaneously active cells were more likely to be the target of sympathetic sprouts.

**METHODS**

**Surgery**

All the surgical procedures were reviewed and approved by the University of Arkansas for Medical Sciences and University of Cincinnati Institutional Animal Care and Use Committee (IACUC).

**Surgical procedure for complete sciatic nerve transection**

Young female Sprague-Dawley rats weighing 80–100 g at the time of surgery were anesthetized with pentobarbital sodium (40 mg/kg ip). The right sciatic nerve was exposed at mid-thigh level, tightly ligated (3–0 silk suture), and transected 5 mm distally from the ligature as described previously (Wall et al. 1974). The incision was then closed in layers and a single dose of penicillin (8,000 IU) was given to all rats.

**Surgical procedure for implantation of osmotic minipumps for local delivery of chemicals to the nerve trunk**

At the time of sciatic nerve ligature/transaction, osmotic minipumps (model No. 2002; Durect, Cupertino, CA) preloaded with artificial cerebrospinal fluid (ACSF, in mM: 130 NaCl, 24 NaHCO3, 3.5 KCl, 1.25 NaH2PO4, 1.2 MgCl2, 1.2 CaCl2, and 10 dextrose, pH = 7.3), along with TTX (780 μM in ACSF), tetraethylammonium chloride (TEA; 10 mM in ACSF), or 4-aminopyridine (4-AP; 10 mM in ACSF; Sigma Chemical, St. Louis, MO) were implanted subcutaneously after 4-h incubation in saline at 37°C. The flow rate was 1 μl/h for 7 days (i.e., a total of 130 nmol of TTX or 1.7 μmol of TEA or 4-AP was delivered over the course of 7 days). To block the injured sciatic nerve, the minipump filled with TTX was attached to a piece of silicone tubing (50–60 mm in length, 0.64 mm ID ×1.19 mm OD; Dow Corning, Midland, MI) that was led to the sciatic nerve ~5–10 mm proximal to the ligature. The silicone tubing was slit over the final 10 mm. The two halves of the slit end of the tubing were placed surrounding the sciatic nerve and tightened together at the very end with 6–0 silk suture as previously described (Xie et al. 2005). The TTX pump was in place before the actual transection of the nerve, and hence the procedure included blockade of the injury discharge. In control experiments, the TTX pump was placed on the uninjured contralateral sciatic nerve to rule out the possibility of a systemic effect on sympathetic sprouting or abnormal activity by TTX. To enhance the firing of the injured axons, the injury site of the sciatic nerve was infused with osmotic pump preloaded with TEA or 4-AP solution through a fine PE tubing that was inserted into the sciatic nerve through the transected stump. The sciatic nerve and the inserted tubing were then tightly ligated with 3–0 silk suture at 5 mm proximal to the injury site.

**In vitro microelectrode intracellular recording**

At postoperative days (POD) 35–49, intracellular recording was performed on sensory neurons in whole DRG preparations isolated from normal and axotomized rats. As described in previous publications (Liu et al. 2002; Zhang et al. 1999), the ipsilateral L3 or L4 DRG was placed in the recording chamber and mounted on the stage of an upright microscope (BX50-W1, Olympus). A U-shaped stainless steel rod with three pieces of fine nylon filaments crossed from one side to the other was used to gently hold the ganglion in place within the recording chamber. The DRG was continuously perfused with oxygenated ACSF at a rate of 2–5 ml/min. The temperature was maintained at 36 ± 1°C by a temperature controller.

DRG cells were visualized under differential interference contrast (DIC). Images of the DRG neurons were clearly visible on a high-resolution video monitor fed by a cooled CCD camera (CCD300T, Dage MTI, Michigan City, IN). Intracellular, electrophysiological recordings were made from each cell with a microelectrode filled with 2 M potassium acetate (pH = 7.2). Satisfactory recordings were obtained with electrodes of 50–80 MΩ. Before electrode penetration, the DRG soma was visually classified by the diameter of its soma as small (<30 μm), medium (30–50 μm), or large (>50 μm). The electrophysiological data were collected with the use of single-electrode continuous current clamp (AxoClamp-2B, Axon Instruments, Union City, CA) and analyzed with pClamp 9 software (Axon Instruments).

In experiments to determine the incidence of spontaneous activity, individual DRG neurons were first impaled with a recording electrode. If spontaneous activity was absent during the first 60 s of the impaling, incremental currents (±4 nA) were then injected to ensure that action potentials could be evoked indicating a healthy cell. If any spontaneous activity was present, then we would wait for 3 min to ensure that the activity was not caused by penetrating the somata with the sharp electrode. For experiments investigating co-localization of sympathetic fibers with identified cells, after the initial recording was completed and the recording electrode removed, a second, higher-resistance, dye-filled electrode was used to impale the same neuron for intracellular injection of Lucifer yellow, a technique similar to that described by Lawson and her colleagues (Djouhri et al. 2003; Fang et al. 2002; Lawson et al. 1997). A successful injection was verified under the same microscope by briefly exposing the neuron to UV light with proper emission/excitation filter set. The whole DRG was then fixed for TH staining (see following text) to determine if the labeled neurons were surrounded by or co-localized with sympathetic fibers. In other experiments, some quiescent neurons were labeled using the same method.

In some experiments, in addition to determining spontaneous activity, we characterized the membrane properties and excitability more fully. Measured parameters were the threshold current (rhobase), action potential (AP) threshold, resting membrane potential (Vm), input resistance (Ri), and afterhyperpolarization (AHP) of the recorded DRG cell. The Vm was taken 3 min after a stable recording was first obtained. Depolarizing currents of 0.05–4 nA (100-ms duration) were delivered in increments of 0.05 nA until an AP was evoked. The threshold current was defined as the minimum current required to evoke an AP. The AP voltage threshold was defined as the first point on the rising phase of the spike at which the change in voltage exceeded 50 mV/msec. The duration of the AP was measured at the threshold voltage. The AP amplitude was measured between the peak and the AP threshold. The Ri for each cell was obtained from the slope of a steady-state I-V plot in response to a series of hyperpolarizing currents, 100-ms duration delivered in steps of 0.05 nA from 0.2 to −2 nA. The AHP amplitude was measured between the maximum hyperpolarization and the final plateau voltage, and the AHP duration was measured at the voltage half way between these two points.
Extracellular fiber recording

For some experiments (Table 2), spontaneous activity was determined with extracellular fiber recording instead of intracellular recording. The spontaneous activity of the dorsal root fibers was extracellularly recorded using in vitro microfilament dissection technique (Zhang et al. 1997). Briefly, the L₄ or L₅ DRG with attached dorsal root and initial segment of sciatic nerve was removed and maintained in oxygenated ACSF at 37°C in a recording chamber. The Govrin-Lippmann and Devor method (Govrin-Lippmann and Devor 1978) was used to determine the incidence of ongoing discharge in those dorsal root fibers in which conduction velocity was able to be measured by electrical stimulation of the dorsal root of L₄ and L₅. Briefly, each fiber bundle of approximately equal diameter (40–50 μm), the sciatic nerve was stimulated with a gradually increasing intensity of current (0.1–0.5 ms square wave pulses, 1–2 Hz) ≤10 mA, resulting in a gradual recruitment of A-fibers then C-fibers until the number of fibers saturated. The total number of different spontaneous action potential waveforms was counted and summed for all strands and divided by the total number of activatable fibers (in all strands) recruited by electrical stimulation of the sciatic nerve to obtain the incidence of ectopic discharge.

TH immunostaining of sympathetic fibers in sectioned and whole-mount DRGs

For measurement of sympathetic fiber density in sectioned DRGs (Fig. 4), rats were anesthetized with pentobarbital sodium (40 mg/kg ip) and fixed by perfusing 200–300 ml of Zamboni’s fixative through the left ventricle of the heart. The bilateral DRGs of L₄ and L₅ were removed, postfixed in the fixative for 2 h at 4°C, and embedded in gelatin overnight. The ganglia were horizontally sectioned with a Vibratome at a thickness of 40 μm.

A whole-mount DRG staining procedure was used to visualize the distribution of sympathetic fibers on the surface of the ganglion (Fig. 3). The whole-mount method was also used to visualize sympathetic fibers near individual neurons after intracellular recording and dye injection (Figs. 5 and 6) because only neurons on the ganglion surface are accessible to these procedures. For whole-mount DRG staining, the unsectioned DRG was fixed in 4% paraformaldehyde (0.1 M phosphate buffer, pH 7.4) for 2 h. The capsule was always removed before whole-mount staining or intracellular recording.

After blocking with 10% normal goat serum in PBS for 30 min, the tissue sections or whole DRGs were incubated in antibodies to TH (from Pel-Freeze, Rogers, AR) at a dilution of 1:1,000 for 24 h at 4°C, followed by the reaction with biotinylated secondary antibody and, finally, with Vector ABC reagent. The TH antibody is an affinity-purified polyclonal rabbit antibody; the antigen is purified denatured rat TH isolated from pheochromocytoma cells. Specificity is demonstrated by the ability to stain the noradrenergic and dopaminergic systems in rat brain with low background. Triton-X (0.3%) was used in all reaction solutions to enhance antibody penetration. Immunoreaction products were visualized by the dianimobenzidine method in the presence of H₂O₂ in 0.1 M phosphate buffer. Tissue sections were then mounted on gelatin-coated slides, air dried, dehydrated, and coverslipped for light-microscopic observation. Whole-mount DRGs were placed on slides coverslipped with anti-fade mounting medium.

Measurement of TH-IR fiber density in the DRG sections

Using ImagePro Plus software (Media Cybernetics, Silver Spring, MD), images from all sections of each DRG were captured under a light microscope (×20) equipped with a SPOT Insight colored digital camera (Diagnostic Instruments, Burlington, CA). Using the ImagePro program, TH-IR fibers in each image were traced, and a new image containing all the traced fibers was generated from the original image. After applying a “thinning filter”, all traced fibers were converted to single-pixel lines. The total number of pixels within each image was then counted and converted to fiber length (in μm). The numerical density of the TH-IR fiber within each image was obtained by dividing the total fiber length by the size of the measured area (in mm²). The density of TH-IR fibers for each DRG was calculated after all images from all sections were measured and counted.

Determination of co-localization of TH-IR fibers and individual DRG neurons in whole-mount ganglia

After individual neurons were identified by intracellular recording as spontaneously active (or quiescent, in separate experiments) and injected with Lucifer yellow as described in the preceding text, the whole DRG was fixed and stained for TH as described in the preceding text. Whole-mount preparations were observed with an upright microscope (Olympus BX50-WI). Dye-injected cells were scored as being co-localized with sympathetic sprouts if TH-positive fibers were observed to end anywhere on the cell surface. Fibers passing completely across the neuron but not ending there were not scored as co-localized (e.g., see Fig. 5A). It was also noted whether cells had more extensive sprouting in the form of rings or basket formations.

Data analysis

All data are expressed as means ± SE. The proportions of spontaneously active fibers or neurons in different experimental groups, and of neurons with sympathetic fibers around them, were compared using Fisher’s exact test. Fiber density data were analyzed using ANOVA with Tukey’s post hoc test. Average values for electrophysiological parameters were compared using Student’s t-test.

RESULTS

Development of hyperexcitability and spontaneous activity of DRG neurons after sciatic nerve transection depends on ectopic discharge from the injury site

EARLY NERVE BLOCKADE REDUCED SPONTANEOUS ACTIVITY IN DRG NEURONS AFTER SCIATIC AXOTOMY. Between POD 35 and 49 days, the incidence of spontaneous activity was measured using whole-DRG intracellular recording in the following groups of DRG neurons: 2,390 neurons from rats with axotomy but without any treatment (n = 21), 915 axotomy with early nerve blockade (TTX perfusion of the nerve proximal to the injury site for the 1st 7 days POD, n = 9), 404 axotomy with early ACSF perfusion (n = 8), and 782 normal (n = 10). The incidence of spontaneous activity in normal DRG was around 5% in both large (21/415)- and medium (10/200)-sized neurons. Sciatic nerve transection dramatically increased the incidence to 11.5% (156/1352; P < 0.001) in medium-sized neurons, whereas, early nerve blockade significantly reversed this abnormal high incidence of spontaneous activity in DRG neurons, returning it to normal levels of 3.3% (17/520; P = 0.18 vs. normal) in large-sized neurons and toward normal levels of 10.4% (26/249; P = 0.04 vs. normal) in medium-sized neurons. Perfusion with ACSF (vehicle) instead of TTX was indistinguishable from axotomy with no perfusion (P = 0.45, large cells; P = 0.30, medium cells). The effects of TTX perfusion were unlikely to be due to systemic effects because perfusion of the (uninjured) contralateral nerve with the same concentration of TTX and at the same low pump rate (1 μl/h) as was used on the
injured nerve was ineffective at reducing spontaneous activity, which remained at 19% in large-diameter cells and 34% in medium-diameter cells, not significantly lower than the axotomy perfused with ACSF group (large cells, $P < 0.45$; medium cells, $P < 0.30$). The incidence of spontaneous activity in C-cells after axotomy was already so low (5/439, 1.1%; $P < 0.60$ vs. normal) that it was not possible to measure statistically significant decreases after early nerve blockade (Fig. 1).

The particular patterns of spontaneous activity were also compared between neurons in the different experimental groups. As shown in Fig. 2, most of the increase in spontaneous activity after axotomy was due to increased incidence of cells with either a bursting or irregular activity, and both these types of activity were reduced by nerve blockade. The percentage of tonically firing cells was much less sensitive to axotomy or TTX.

EARLY NERVE BLOCKADE REDUCED EXCITABILITY IN DRG NEURONS AFTER SCIATIC NERVE TRANSECTION. In some experiments, excitability parameters were measured in addition to determining incidence of spontaneous activity. In addition to a high incidence of spontaneous activity, sciatic nerve transection also caused abnormal depolarization of the resting membrane potential and reduction (hyperpolarization) of the AP threshold, decreased rheobase, decreased amplitude of AHP, and increased AP and AHP durations in both large- and medium-sized DRG neurons. These parameters are summarized in Table 1. Only depolarized resting $V_{m}$, hyperpolarized AP threshold, and decreased rheobase were observed in small-sized neurons after sciatic nerve transection. More remarkably, instead of increasing, AP duration was significantly decreased in small-sized neurons.

Early nerve blockade inhibited the abnormal hyperpolarization of AP threshold in both large- and medium-sized neurons and increased rheobase significantly in large and medium-sized neurons. There was no clear effect observed on other abnormal parameters after early nerve blockade (Table 1). Also the electrophysiological parameters of DRG neurons after sciatic nerve transection were not affected by ACSF perfusion (Table 1). These results suggest initial neuronal blockade at the time of nerve injury may prevent the subsequent development of spontaneous activity and hyperexcitability in medium- and large-sized DRG neurons after sciatic nerve transection, even when measured 4–5 wk after the end of the nerve blockade period.

We always observed underlying membrane oscillations in spontaneously active neurons with bursting or irregular firing patterns, although this could not be observed in high-frequency tonically firing cells. Similar observations have been reported by others (e.g., Amir et al. 2005). Subthreshold membrane potential oscillations that were not sufficient to trigger APs were also observed in some neurons that were not spontaneously active. Like spontaneous activity, the incidence of such oscillations (observed 4–10 wk after nerve injury) was increased after sciatic nerve transection and reduced by early nerve blockade. Of the nonspontaneously active neurons, subthreshold oscillations were observed in 3.5% (20/577) of cells from normal rats. This increased to 8.5% (107/1263) after axotomy or 11.0% (38/346) after axotomy with ACSF perfusion of the nerve (both significantly different from control; $P < 0.0001$, but not different from one another, $P = 0.17$). Nerve blockade with TTX during the first week after injury reduced
Comparison of fiber density on the dorsal surface of the DRG (Fig. 3).

TH-IR fibers sprouted vigorously and invaded the center portion of the ganglion. None could be seen in the ventral side of the DRG. Small-diameter cells or "dark cells," as previously described (Lawson 1979; Price and Mudge 1983; Tandrup et al. 2000). Fibers accompanying the vascular processes and some were from the small TH-IR dopaminergic neurons. None could be seen in the center region of the dorsal surface of the ganglion. Most fibers accompanied the vascular processes and some were from the small TH-IR dopaminergic neurons or "dark cells," as previously described (Lawson 1979; Price and Mudge 1983; Tandrup et al. 2000). Fibers accompanying the vascular processes were from large bundles of TH-IR branches. The density of TH-IR fibers on the ventral side of the DRG was higher compared with the fiber density on the dorsal surface.

In DRG examined after nerve transection (n = 21), TH-IR fibers sprouted vigorously and invaded the center portion of the dorsal surface of the DRG (Fig. 3C). In some cases, the sprouted fibers formed ring or basket structures around large- and medium-sized DRG neurons. The overall density of the fibers increased significantly compared with normal DRG. The TH-IR dark cells disappeared completely in the axotomized DRG (Fig. 3, C and D). Similar changes occurred on both sides of the ganglion.

DENSITY OF SYMPATHETIC FIBERS IN DRG SECTIONS AFTER SCIATIC NERVE TRANSECTION WAS DECREASED BY EARLY NERVE BLOCKADE AND INCREASED BY POTASSIUM CHANNEL BLOCKADE. A great deal of evidence indicates that peripheral nerve injury increases

TABLE 1. Effect of early nerve blockade on electrical parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Normal</th>
<th>Axotomy</th>
<th>Axotomy + ACSF</th>
<th>Axotomy + TTX</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>42 cells, 5 animals</td>
<td>35 cells, 8 animals</td>
<td>54 cells, 7 animals</td>
<td>41 cells, 6 animals</td>
</tr>
<tr>
<td>V_m, mV</td>
<td>−58.1 ± 1.70</td>
<td>−51.2 ± 1.83*</td>
<td>−52.4 ± 0.98*</td>
<td>−52.02 ± 1.23*</td>
</tr>
<tr>
<td>Rheobase, nA</td>
<td>1.06 ± 0.03</td>
<td>0.78 ± 0.06**</td>
<td>0.73 ± 0.04**</td>
<td>0.72 ± 0.04**</td>
</tr>
<tr>
<td>AP threshold, mV</td>
<td>−17.62 ± 1.70</td>
<td>−23.1 ± 1.66*</td>
<td>−24.5 ± 1.04*</td>
<td>−25.6 ± 1.24***</td>
</tr>
<tr>
<td>AP duration, ms</td>
<td>3.0 ± 0.34</td>
<td>2.9 ± 0.25</td>
<td>2.80 ± 0.14</td>
<td>2.47 ± 0.11</td>
</tr>
<tr>
<td>AHP amplitude, mV</td>
<td>51.7 ± 1.18</td>
<td>48.3 ± 2.24</td>
<td>50.9 ± 1.12</td>
<td>50.0 ± 1.41</td>
</tr>
<tr>
<td>AHP duration, ms</td>
<td>6.0 ± 0.55</td>
<td>5.3 ± 0.52</td>
<td>4.4 ± 0.27*</td>
<td>3.45 ± 0.22*</td>
</tr>
<tr>
<td>R_m, MI</td>
<td>17.4 ± 0.93</td>
<td>16 ± 1.08</td>
<td>17.5 ± 0.50</td>
<td>17.06 ± 0.73</td>
</tr>
</tbody>
</table>

Cell diameters were obtained from microscopic observations made during intracellular recording. Values are means ± SE. AP, action potential; AHP, afterhyperpolarization; R_m, input resistance; ACSF, artificial cerebrospinal fluid. Compared with normal value, ***P < 0.0001; **P < 0.001; *P < 0.05.

Compared with axotomy and axotomy + ACSF group. §P < 0.05.

the percentage of oscillating cells to 5.7% (50/871), significantly lower than the axotomy group but not significantly different from the normal group. These oscillations were observed 2–9 wk after injury (longer times were not tested).

Relationship between sympathetic nerve sprouting and spontaneously active neurons

DISTRIBUTION OF TH-POSITIVE SYMPATHETIC FIBERS ON THE SURFACE OF DRGS BEFORE AND AFTER SCIATIC NERVE TRANSECTION. As shown in Fig. 3, in normal rats (n = 10), some TH-IR sympathetic fibers could be seen on the dorsal (Fig. 3, top) and ventral (Fig. 3, bottom) surface of the whole-mount DRG. But viewed in these orientations, the fibers were mainly located around the edge of the ganglion. None could be seen in the center region of the dorsal surface of the ganglion. Most fibers accompanied the vascular processes and some were from the small TH-IR dopaminergic neurons or "dark cells," as previously described (Lawson 1979; Price and Mudge 1983; Tandrup et al. 2000). Fibers accompanying the vascular processes were from large bundles of TH-IR branches. The density of TH-IR fibers on the ventral side of the DRG was higher compared with the fiber density on the dorsal surface.

In DRG examined after nerve transection (n = 21), TH-IR fibers sprouted vigorously and invaded the center portion of the dorsal surface of the DRG (Fig. 3C). In some cases, the sprouted fibers formed ring or basket structures around large- and medium-sized DRG neurons. The overall density of the fibers increased significantly compared with normal DRG. The TH-IR dark cells disappeared completely in the axotomized DRG (Fig. 3, C and D). Similar changes occurred on both sides of the ganglion.

FIG. 3. Anti-tyrosine hydroxylase (TH) immunostaining of sympathetic fibers in whole-mount DRG. Sympathetic nerve fibers could be observed on the dorsal (top) and ventral (bottom) surfaces of both normal (A and B) and axotomized (C and D) DRGs, however, the axotomized DRGs had much higher fiber density on both sides of the ganglion. Scale bar = 200 μm.
the sprouting of sympathetic nerve in DRG after nerve transection. To determine whether ectopic firing in the DRG is related to the abnormal growth of sympathetic nerve fibers, we compared the density of sympathetic fiber sprouting between axotomized DRGs in which we experimentally increased or decreased the incidence of spontaneous activity reaching the DRG cell body. The density of sympathetic nerve fibers was counted in DRG sections by TH immunostaining. Early nerve blockade during the first 7-days post sciatic nerve transection \((n = 5)\) significantly \((P < 0.001, \text{ANOVA})\) decreased the sprouting of sympathetic nerve in DRG as measured between POD 35 and 49, reducing it to a level just slightly higher than that seen in uninjured animals (Fig. 4A). Conversely, injured peripheral nerve fibers are sensitive to \(K^+\) channel blockers such as TEA and 4-AP, which can evoke robust increases in spontaneous firing at the nerve injured site. To further confirm the relationship between spontaneous activity and sympathetic nerve sprouting, TEA \((n = 3)\) or 4-AP \((n = 3)\) filled osmotic pumps were used to perfuse the injury site for 7 days to increase ectopic firing at the site of sciatic nerve transection. Perfusion with ACSF was used as a control \((n = 3)\). Consistent with previous studies (Devor 1983; Xie et al. 1993), dorsal root fiber recording on POD 7 indicated that spontaneous activity from DRG neurons was significantly increased after TEA or 4-AP perfusion by 1.5-fold for both treatments (Table 2). The two treatments did not differ from each other significantly \((P = 0.21\) for \(L_4; 0.054\) for \(L_5\)). In addition, this increased ectopic discharge also resulted in a robustly higher density of sympathetic nerve fiber in the DRGs, which were collected and stained on POD 14 (Fig. 4B). These results indicate that sympathetic nerve sprouting around DRG cell bodies following axotomy is directly affected by the spontaneous activity evoked by the injury.

CO-LOCALIZATION BETWEEN SYMPATHETIC NERVE SPROUTING AND INDIVIDUAL ECTOPIC FIRING NEURONS. The preceding data clearly suggest that spontaneous activity in DRG directly correlates with the sprouting of sympathetic nerve after nerve transection. We further studied the relationship between ectopic firing and sympathetic nerve growth in DRG at the level of individual neurons. After sciatic nerve transection \((n = 21)\), we observed much higher density of sympathetic nerve fibers on the surface of DRG than on the normal ones \((n = 10; \text{Fig. 3})\). During intracellular recording, individual ectopically active or (in separate experiments) quiescent neurons were labeled with injection of the fluorescent dye Lucifer yellow, followed by whole DRG TH immunohistochemical staining to detect sympathetic fibers. This allowed us to determine whether the sympathetic sprouting occurred preferentially around ectopically firing neurons. Two hundred and sixty-one (261) ectopically firing neurons in 42 DRGs were successfully labeled, and sympathetic nerve fibers ended on 54\% (142) of them (Fig. 5).

In 43 of these cells, the sympathetic fiber endings took the form of a ring-like structure and in 12, a basket structure. In the

### TABLE 2. Effect of \(K^+\) channel blockers on spontaneous activity

<table>
<thead>
<tr>
<th></th>
<th>Axotomy + ACSF</th>
<th>Axotomy + TEA</th>
<th>Axotomy + 4-AP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of animals</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Incidence of Spontaneous activity, %</td>
<td>21.1 ± 0.5</td>
<td>31.5 ± 1.0</td>
<td>31.8 ± 2.9</td>
</tr>
<tr>
<td>(P) value vs. ACSF</td>
<td>—</td>
<td>0.0001</td>
<td>0.0001</td>
</tr>
<tr>
<td>Total number of fibers recorded</td>
<td>503</td>
<td>555</td>
<td>576</td>
</tr>
</tbody>
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Incidence of spontaneous activity determined on post-operative day (POD) 7 after perfusion of the injured nerve with \(K^+\) blockers TEA (10 mM) or 4-aminopyridine (4-AP; 10 mM), or with vehicle (ACSF). The pump was still in place at the time spontaneous activity was measured. SEs are based on comparison between different animals. \(P\) values are based on Fisher’s exact tests on the combined data. There was no significant difference between the TEA and 4-AP groups.
remaining cells, the co-localization took the form of simple fiber endings on the surface of the cell. We observed that sympathetic nerve also formed basket-like structures on 19 unlabeled neurons, which were just adjacent to 19 of the labeled (spontaneously active) neurons. We noticed that spontaneously firing neurons, usually concentrated in several spots on the surface of DRG, suggesting that cross-talk may exist between spontaneously active DRG neurons. In separate experiments, we also randomly labeled 124 neurons without spontaneous activity in 17 DRGs after axotomy. Only nine of them had sympathetic nerve fibers ending on the surface; none of these took the form of ring or barked structures. Hence growth of sympathetic fibers occurred preferentially around spontaneously active DRG cells (Figs. 5 and 6).

To determine how general the co-localization phenomenon might be, we conducted a similar experiment in neurons from DRG that had been subjected to inflammation in the absence of nerve injury. DRG were inflamed by local deposition of a small drop of the immune activator zymosan (n/H11005/H17). As shown previously (Xie et al. 2006), this treatment causes sympathetic sprouting, spontaneous activity, and mechanical allodynia and hyperalgesia. In co-localization experiments such as those shown in Fig. 5, conducted 15–35 days after localized inflammation of the DRG, 38% (35/93) of SA neurons were contacted by sympathetic fibers compared with only 11% (5/47) of quiescent neurons (P = 0.0007).

**REDDUCING SYMPATHETIC SPROUTING IN DRG AFTER SCIATIC NERVE TRANSECTION BY EARLY NERVE BLOCKADE DOES NOT CHANGE THE CORRELATION BETWEEN SPROUTING AND INDIVIDUAL SPONTANEOUSLY FIRING NEURONS.** As we have described in the preceding text, perfusing TTX onto the neuroma during the first 7 days after sciatic axotomy decreased but did not eliminate spontaneous activity in the DRG and also decreased but did not eliminate sympathetic sprouting. To determine whether the preferential sprouting around spontaneously active neurons was preserved even when both phenomena were reduced by early nerve blockade, we repeated the preceding experiment in DRG in which the neuroma was perfused with TTX for the first 7 days after nerve transection (n = 9). Recordings were made on POD days 14–45 (i.e., 7–37 days after the end of nerve blockade). Among 36 labeled ectopically firing neurons in 17 DRGs, 18 of them (50%) had adjacent sympathetic nerve sprouts. In three neurons, the length of sympathetic nerve was longer than the half of the cell diameter. We also randomly labeled 76 neurons without spontaneous activity in eight DRGs. Twenty of them (26%) had adjacent sympathetic nerve fibers ending on the cell surface (Fig. 6, P = 0.02). This result indicated that the correlation between sympathetic nerve and ectopic firing neurons was not reduced by decreasing spontaneous activity even though the overall incidence of both sprouting and activity was reduced. Using ACSF to replace TTX and perfuse transected sciatic nerve also did not change the relationship between sympathetic nerve and neurons with ectopic firing in the DRG (n = 8). This result further indicates the specific correlation between ectopic firing neuron and sympathetic nerve sprouting. This relationship is independent of the absolute incidence of spontaneous activity or the absolute density of sympathetic nerve fibers.

**DISCUSSION**

This study examined relationships between spontaneous activity and sympathetic sprouting around DRG cells in the complete sciatic nerve transection model of neuropathic pain. The primary new findings are that short-term (1 wk) blockade of the nerve proximal to the neuroma leads to reduction in
The effects on sympathetic sprouting, hyperexcitability, and spontaneous activity of TTX perfusion near the nerve injury site are likely to be due to its local impulse blocking effects on the injured nerve because in our previous study, a similar effect on spontaneous activity that occurred in DRG cell bodies. Simply preventing this activity from reaching the DRG during the first week after nerve injury led to increased sympathetic sprouting and that sympathetic sprouts were much more likely to terminate near spontaneously active rather than quiescent neurons.

The effects on sympathetic sprouting, hyperexcitability, and spontaneous activity of TTX perfusion near the nerve injury site are likely to be due to its local impulse blocking effects of drug, lidocaine, for local nerve blockade (Zhang et al. 2004); TTX perfusion of the contralateral uninjured nerve did not reduce spontaneous activity in the ipsilateral DRG, arguing strongly against a systemic effect; and perfusion of the injured nerve with chemically distinct drugs that increased spontaneous activity had opposing effects on the density of sympathetic sprouting.

**Key role of early activity**

It is well known that sciatic nerve transection induces abnormal spontaneous activity, originally arising from the neuroma. Later, spontaneous activity can originate in the DRG, and even cells not showing spontaneous activity may develop hyperexcitability (Amir 1999; Babbedge et al. 1996; Burchiel 1984a,b; DeSantis and Duckworth 1982; Govrin-Lippmann and Devor 1978; Kajander et al. 1992; Kirk 1970; Michaels 2000; Wall and Devor 1983). Indeed, spontaneous activity is a key feature in many different neuropathic pain models (see introduction). Such activity is prominent in medium-diameter cells, many of which are nociceptors, as well as in large-diameter cells, which are not normally nociceptors. Several mechanisms have been proposed to account for the apparent contribution of spontaneous activity in large-diameter cells to pathological pain (e.g., see (Abdulla et al. 2003)). Our data suggest that the initial early period of ectopic discharges from the injury site plays a key role in setting up the later, more prolonged excitability changes, and in generation of spontaneous activity that occur in DRG cell bodies. Simply preventing this activity from reaching the DRG during the first week after injury results in a substantial reduction in spontaneous activity and hyperexcitability measured in the DRG cell bodies and dorsal roots 5–6 wk after the nerve blockade has ended. Other studies have also suggested that blocking the initial period of spontaneous activity in various nerve injury models can have profound, long-lasting effects. Our previous work (Xie et al. 2005) with two different localized blockade methods, in two different partial injury neuropathic pain models, also provided evidence for the critical importance of spontaneous activity during the first week after nerve injury. Such blockade could lead to reduction in mechanical and thermal pain behaviors that long outlasted the duration of the blockade. Temporarily blocking spontaneous activity reduces or eliminates spontaneous pain, hyperalgesia, and allodynia in a variety of pain models, using methods to suppress spontaneous activity that vary widely in their specific targets (Boucher et al. 2000; Chaplan et al. 2003; Lai et al. 2002; Lyu et al. 2000; Seltzer et al. 1991; Xiao and Bennett 1995; Yoon et al. 1996), however, see (Suter et al. 2003).

In a previous study (Zhang et al. 2004), we showed that lidocaine, either systemic or locally applied to the injury site, could reduce sympathetic sprouting in the sciatic nerve transection model. The present study builds on those findings in showing that experimentally increasing activity with local perfusion of K\(^+\) channel blockers can increase sprouting; that a different local blocker (e.g., TTX) has the same effect as lidocaine and that individual spontaneously active neurons are preferentially targeted by sympathetic sprouts. A surprising finding in the Zhang et al. paper was that the reduction in sympathetic sprouting outlasted the duration of the blockade. Interestingly, prolonged effects of treatments of short-acting local anesthetics are also reported in clinical examples of sympathetically maintained pain. These findings, along with previous studies that emphasize the key role of the early period of spontaneous activity, support the notion of preemptive analgesia. Attempts to prevent development of chronic pain by providing analgesia early (e.g., immediately before or just after a surgery and before chronic pain has developed) have been conflicting and often disappointing (Kelly et al. 2001; Moiniche et al. 2002). However, many such clinical studies have used blockade periods that were much shorter than the 7-day blockade used in this study. In addition, this and previous studies suggest that analgesics that acted primarily at the level of the spinal cord would not be able to prevent the prolonged alterations in DRG properties...
that occur in animal models of pathological pain. Hence after the analgesic was removed, establishment of central sensitization by the abnormal DRG cells might still occur.

Relationship between spontaneous activity and sympathetic sprouting

The observation that sympathetic sprouting in this model occurs preferentially around large- and medium-diameter neurons, which are also most likely to develop high-frequency spontaneous activity, provided the initial suggestion that sprouting might be related to spontaneous activity. This study demonstrated a correlation between sprouting and spontaneous activity at the level of individual neurons. Our study does not directly address mechanism and causality. The observation that spontaneous activity appears very early after injury, well before sprouting can be observed, as well as our ability to affect sprouting by manipulating spontaneous activity suggest that if there is a causal relationship, it is that activity somehow induces sprouting around individual neurons. However, our data do not address this issue directly, and we cannot eliminate an indirect or a bidirectional relationship between sprouting and activity.

Many mechanistic studies of sympathetic sprouting into the DRG have focused on possible roles of neurotrophic factors and/or cytokines. Some of these theories focus on the possible role of the satellite glia cells that surround DRG neurons rather than on the neurons themselves [e.g., (Ramer et al. 1999; Walsh and Kawaja 1998; Zhou et al. 1999)], and it is worth noting that structural studies indicate that the sprouting fibers predominantly contact the satellite glia sheath rather than the DRG neurons (Shinder et al. 1999). The present findings are not inconsistent with a role of neurotrophic factors or cytokines in sprouting. There are a number of mechanisms by which abnormal activity in the DRG cell bodies might result in local elevation of trophic factors or cytokines. For example, activity might evoke release of substances from neurons that either enhance sympathetic sprouting directly, or stimulate nearby glial cells to produce substances such as NGF that then enhance sprouting. Many studies support the connection between neuronal activity and release of trophic factors (Castren et al. 1992; Ernfors et al. 1991; Gall and Isackson 1989; Gall et al. 1991a,b; Isackson et al. 1991; Kim et al. 1994; Lu et al. 1991; Patterson et al. 1992; Thoenen 1991; Zafra et al. 1990, 1992), as well as mechanisms by which neuronal activity is communicated to surrounding glial cells (Gunzel and Schlie 2000; Murphy et al. 1993; Schmidt et al. 1999). The latter include direct linking of neuronal and glial membrane potentials (Lohr and Deitmer 1999; Newman and Zahn 1998; Rouach et al. 2000), stimulation of calcium waves in glia by glutamate (Dani et al. 1992; Venance et al. 1995), and release of fractalkine by activated neurons (Chao et al. 1995; Kyrkanides et al. 1999; Watkins and Maier 2002; Winkelstein et al. 2001). Our observation that spontaneously active neurons tended to be found in clusters would be consistent with a role for highly localized factors. Highly localized effects of neuronal activity on sympathetic sprouting are also suggested by the finding that spontaneously active cells are more likely to have sympathetic fibers nearby even after early nerve blockade has greatly reduced the overall incidence of both activity and sprouting.

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