Schwann Cell Engraftment Into Injured Peripheral Nerve Prevents Changes in Action Potential Properties

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Yu, Kewei and Jeffery D. Kocsis. Schwann cell engraftment into injured peripheral nerve prevents changes in action potential properties. J Neurophysiol 94: 1519–1527, 2005; doi:10.1152/jn.00107.2005. Peripheral nerve injury results in changes in action potential waveform, ion channel organization, and firing properties of primary afferent neurons. It has been suggested that these changes are the result of reduction in basal trophic support from skin targets. Subcutaneous injections of Fluoro-Gold (FG) in the hind limb of the rat were used to identify cutaneous primary afferent neurons. Five days after FG injection, sciatic nerves were ligated and encapsulated in a silicon tube allowing neuroma formation. Green fluorescent protein (GFP)-expressing Schwann cells (SCs) were injected proximal to the cut end of the nerve. Thirteen to 22 days after injury and SC injection, the L2 and L4 dorsal root ganglia (DRG) were prepared for acute culture. Whole cell patch-clamp recordings in current clamp mode were obtained and action potential properties of medium-sized (34–45 μm) FG+ DRG neurons were characterized. In the neuroma group without cell transplantation, action potential duration and spike inflections were reduced as were the amplitude and duration of spike afterhyperpolarizations. These changes were not observed after transection by nerve crush where axons were allowed to regenerate to distal peripheral targets. In the transplantation group, GFP+/−SCs were extensively distributed throughout the neurona, and oriented longitudinally along axons proximal to the neuroma. Changes in action potential properties were attenuated in the GFP+/−SC group. Thus the engrafted SC procedure ameliorated the changes in action potential waveform of cutaneous primary afferents associated with target disconnection and neuroma formation.

INTRODUCTION

Axotomy results in changes in excitability of cutaneous afferent dorsal root ganglion (DRG) neurons and in action potential waveform (Dever and Wall 1990; Gurdu and Smith 1988; Oyelese et al. 1997). Medium-sized DRG neurons (likely Aβ neurons) projecting to skin can show both fast noninflected (A0) and inflected (A1) action potentials (Amir et al. 1999, 2002; Kim et al. 1998; Liu et al. 2002; Oyelese et al. 1997). Voltage-clamp studies indicate that within this neuronal population both fast TTX-sensitive (TTX-S) and slow TTX-resistant (TTX-R) sodium currents develop rapidly repriming sodium current in axotomized axons (Cummins and Waxman 1997). While much less work has been carried out on injury-induced changes in sodium channel subtypes of medium to large Aδ-Aβ DRG neurons, several reports indicate that after neuroma formation there is a reduction in slow TTX-R currents with a singularly fast TTX-S current predominating (Everill et al. 2001; Oyelese et al. 1997).

The mechanisms for these changes in sodium and potassium currents and in spike waveform are not fully understood. However, focal application of nerve growth factor (NGF) to cut sciatic nerve prevents the reduction in Aδ spikes (Oyelese et al. 1997) and the reduction in potassium currents in Aβ DRG neurons. Leffler et al. (2002) demonstrated that glial line-derived neurotrophic factor (GDNF) and NGF reverse changes in repriming of TTX-S currents after axotomy. Thus it is hypothesized that retrograde transport of trophic factors derived from target or peripheral nerve components such as Schwann cells (SCs) play an important role in stabilizing ion channel distribution and excitability of primary afferent neurons. We asked whether transplantation of SCs, which are known to produce neurotrophic factors, could stabilize action potential properties in a nerve ligation and encapsulation lesion model.

METHODS

Sciatic nerve injury and SC transplantation

Adult Sprague-Dawley (SD) rats of both sexes (150–250 g) were used (n = 69). The surgical procedure was in concordance with the recommendations of the International Association for the Study of Pain (IASP) and was approved by the Institutional Animal Care and Use Committee of Yale University. Sciatic nerve ligation and encapsulation was performed as previous described (Oyelese et al. 1995, 1997) with slight modifications for implanting SCs dissociated acutely from Green fluorescent protein (GFP)-expressing rats. Briefly, for the injury group (n = 19), the sciatic nerve on the left side was exposed in the middle thigh region of anesthetized (ketamine, 40 mg/kg; xylazine, 2.5 mg/kg ip) rats and was ligated and transected distal to the ligature; the proximal stump of sciatic nerve was inserted into a blind-ended silicone tube to prevent nerve regeneration (Fitzgerald et al. 1985; Oyelese et al. 1995, 1997). In the transplantation group (n = 21), four to six injections of a SC suspension (~30,000 cells/μl) were injected 3–5 mm proximal to the nerve crush and were ligated and transected distal to the ligature; the proximal stump of sciatic nerve was inserted into a blind-ended silicone tube to prevent nerve regeneration (Fitzgerald et al. 1985; Oyelese et al. 1995, 1997). In the transplantation group (n = 21), four to six injections of a SC suspension (~30,000 cells/μl) were injected 3–5 mm proximal to the

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ligation for a total of 2.4–3.6 × 10^6 cells. In eight animals, DMEM alone was injected as a sham control group. Nerve crush lesions were achieved in nine rats by compressing the sciatic nerve for 20–30 s with a pair of fine forceps; this transects the axons but allows regeneration to peripheral targets (Oyelese and Kocsis 1996). Age-matched unoperated rats were used as controls (n = 12).

**DRG neuron dissociation and culture**

Two to 3 wk after surgery, the animals were killed under pentobarbital sodium anesthesia (50 mg/kg ip), the left lumber ganglia L₄ and L₅ were excised and dissociated acutely with the use of previously described methods (Liu et al. 2002). Briefly, L₄ and L₅ ganglia were excised in ice-cold sterile calcium-free Krebs’ solution, minced, incubated in HBSS containing 1 mg/ml collagenase A (Boehringer-Mannheim, Indianapolis, IN) followed by a solution that consisted of 0.4 mg/ml collagenase D (Boehringer-Mannheim), 0.4 mg/ml bovine serum albumin (Sigma), and were gently triturated using a fire-polished Pasteur pipette and then distributed onto uncoated glass coverslips. Neurons were then kept in a 5% CO₂-95% O₂ incubator at 37°C. No antibiotic was added to the medium. The neurons were plated at low density (Fig. 1A), and recordings were obtained over the course of 8 h in culture before neurite extension occurred.

**Fluorescence tracer labeling of cutaneous and muscular afferents**

Cutaneous afferent DRG neurons were identified by retrograde labeling with Fluoro-Gold (Fluorochrome, Englewood, CO) as previously described (Liu et al. 2002; Oyelese and Kocsis 1996). Five days before rats were killed, 10–15 μl Fluoro-Gold (FG) solution (1% dissolved in sterile distilled water) was injected intradermally in the lateral region of the ankle. The labeled cells were identified in vitro in culture by yellow fluorescence emission when exposed ultraviolet light (Fig. 1B). Only neurons exhibiting a high degree of fluorescence were selected for study. Underlying muscle and tendon tissues were examined, and no FG was observed on these tissues but was confined to cutaneous tissues as previously reported (Liu et al. 2002). To eliminate errors that might arise from comparing electrophysiological properties that vary with neuronal size, an effort was made to select medium-sized cutaneous afferent neurons ranging from 34 to 45 μm for study. This neuronal size range corresponds to neurons giving rise to myelinated axon conducting in the Aβ and Aδ ranges (Harper and Lawson 1985) and was reported before to obtain waveform changes of action potential (AP) after axotomy (Oyelese et al. 1995).

**Preparation of donor SCs**

SCs were harvested from sciatic nerves of 4- to 8-wk-old SD transgenic rats expressing GFP [“green rat” CZ-004, SD-Tg(Act-EGFP)CZ-004Osborn; SLC, Shizuoka, Japan]. Sciatic nerves from desheathed, minced with a pair of scalpel blades, and incubated for 45 min in DMEM containing 10% fetal calf serum (Hyclone Laboratories, Logan, UT), 1 mg/ml Collagenase A (Boehringer-Mannheim), and 1 mg/ml Collagenase D (Boehringer-Mannheim), followed by a 15-min incubation in Ca²⁺-free CSS containing 0.5 mg/ml trypsin (Sigma), EDTA (Sigma), and a crystal of cysteine (Sigma). Then the nerve tissue was mechanically dissociated by trituration and washed with DMEM. The concentration of the cells suspension was counted and adjusted to about 30,000 cells/μl. Cryostat sections (12 μm) of nerve were incubated with antibodies against neurofilament (1:500; Sigma, St. Louis, MO). Sections were then washed in PBS (0.3% Triton X100 for 15 min) and incubated with appropriate secondary antibodies comprising goat anti-mouse IgG-Alexa Fluor 594 (1:1000; Thermo Scientific, Waltham, MA). Sections were then incubated with 10% normal swine serum (Dako, Carpentaria, CA) and 1:100 Alexa Fluor 488 goat anti-swine IgG (Invitrogen, Carlsbad, CA). Sections were then mounted with fluorescent mounting medium (Dako, CA) and coverslipped. Sections were then imaged using a fluorescence microscope (Olympus, Japan).

**Preparation of donor SCs**

**Preparation of donor SCs**

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**Fig. 1.** A: bright-field image (Hoffman Modulation Contrast Optics) of dorsal root ganglion (DRG) neurons isolated from L₄ and L₅. Note the lack of neurites. B: Fluoro-Gold retrograde labeled cutaneous afferent cell was identified by yellow fluorescence emission when exposed to ultraviolet light. C and D: parameters of action potentials (APs) recorded from medium size (34–45 μm) DRG neurons. Typical somatic APs of a cutaneous afferent DRG neuron evoked by either 1.2-nA, 30-ms (C) or 3.2-nA, 1-ms (D) step depolarizations. C: electrophysiological parameters measured include half-width duration (1), 90% rise time (RT) (2), 90% fall time (FT) (3), AP amplitude (4), AP overshoot (5), maximum slope of the AP rising phase (MSR) (6), and maximum slope of the AP falling phase (MSF) (7). D: AHP amplitude (1), AHP duration at 80% recovery (2) and AHP area at 80% recovery (3). Calibration bar corresponds to 100 μm.
Molecular Probes) in blocking solution for 4 h, washed in PBS, and mounted on slides.

Electrophysiological analysis

Whole cell patch-clamp recordings were obtained 1–8 h after plating to minimize morphological and physiological changes. Neurons plated on glass coverslips were placed in a recording chamber on the stage of an inverted microscope (Nikon) and continuously superfused with a modified Krebs’ solution (composition in mM) was 124 NaCl, 26 NaHCO3, 3 KCl, 1.3 NaH2PO4, 2 MgCl2, 2 CaCl2, and 10 dextrose, pH 7.4 (KOH); osmolarity, 305–315 mosM at room temperature (~22°C) bubbled continuously with 95% O2-5% CO2 at a flow rate of 0.5–1 ml/min. Only medium-sized neurons (34–45 μm diam) with positive FG staining were studied. Micropipettes were pulled from borosilicate glass (World Precision Instruments, Sarasota, FL) with a P97 micropipette puller (Sutter Instrument, San Rafael, CA) and fire polished with a microforge (Narisige, Tokyo, Japan). Electrode resistances ranged from 2 to 6 MΩ. The pipette solution contained (in mM) 140 KCl, 1 CaCl2, 11 EGTA, 2 Mg-ATP, and 10 HEPES, pH 7.3; osmolarity, 300–310 mosM. Recordings were sometimes contaminated by injected current (Blair 2002; Liu 2002) (Fig. 1A) and fire polished with a microforge (Narishige, Tokyo, Japan). The pipette solution contained (in mM) 140 KCl, 1 CaCl2, 11 EGTA, 2 Mg-ATP, and 10 HEPES, pH 7.3; osmolarity, 300–310 mosM. Recordings were included if they met all of the following criteria: >1 MΩ seal resistance prior to “break-through” with further suction; a resting membrane potential (Vr) of −55 mV or more negative; and >100 MΩ input resistance after whole cell mode was established. Once the gigaseal was established, the voltage-clamp mode was changed to current-clamp mode. The series resistance was balanced under fast current-clamp mode with an Axopatch-200B patch-clamp amplifier (Axon Instruments). The voltages were filtered at 5 kHz and acquired at 20 kHz using Clampex 8 software (Axon Instruments). The DigiData 1200B interface (Axon Instruments) was used for A-D conversion. Action potentials were elicited from Vr levels by delivering depolarizing step pulses of either 1- or 30-ms duration generated by Clampex 8.

Definitions of AP components

Parameters of somatic APs recorded from medium-sized (34–45 μm) cutaneous afferent DRG neurons (1.2 nA, 30-ms stimulation) were illustrated in Fig. 1C and defined as follows: AP half-width duration (ms) was measured as the width at half-maximal amplitude of APs (Fig. 1C, 1); 90% rise time (RT, ms) as 90% of the duration from start of current injecting at Vr to the peak of AP (Fig. 1C, 2); 90% fall time (FT, ms) as 90% of the time from peak of AP back to the baseline of Vr (Fig. 1C, 3); AP amplitude (mV) as the voltage value from the baseline at Vr to the positive peak of the spike (Fig. 1C, 4); AP overshoot (mV) as the voltage value from the baseline at 0 mV to the positive peak of the spike (Fig. 1C, 5); maximum slopes of the rising (MSR, dV/dt; Fig. 1C, 6) and falling (MFR, dV/dt; Fig. 1C, 7) phases of the AP.

Short injection of current (3.2 nA, 1 ms) were applied for measure the parameters of afterhyperpolarization (AHP) so that it will not be contaminated by injected current (Blair 2002; Liu 2002) (Fig. 1D). AHP amplitude (mV) was measured from the baseline of Vr to the negative peak of AHP (Fig. 1D, 1); AHP duration at 80% recovery (ms) was measured at the baseline at 20% AHP amplitude (Fig. 1D, 2), which is also the baseline for measuring AHP area at 80% recovery (mV·ms; Fig. 1D, 3).

Finally, the current (nA) and voltage threshold (mV) for evoking a single spike using 30-ms depolarizing pulses, as well as the utilization time (from start of current injecting to the start of AP spiking), were measured. The neurons were further characterized as having an inflected waveform (A5) if digital differentiation (Clampfit 8) indicated two peaks on the falling phase of the spike (Fig. 3A) or noninflected waveform (A0) if there was only one peak (Fig. 3B).

Statistical analysis

Electrophysiological data were processed by using Origin 6.1, Clampfit 8 (Axon Instruments) and Excel (Microsoft). Data are presented as means ± SE. Statistical evaluations were based on two-tailed t-test, χ2 test (Excel, Origin; criterion, P < 0.05).

RESULTS

Passive membrane properties of identified medium-sized cutaneous afferent DRG neurons

Whole cell current-clamp recordings were obtained from 140 dissociated medium-sized cutaneous afferent neurons. Five experimental groups were studied: control (n = 28), ligation (n = 23), nerve crush (n = 32), ligation with SC transplantation (n = 35), and sham injection (n = 22). A comparison of general membrane properties was performed to compare resting potential (Vr), input capacitance (Cap) and input resistance (Rc) among the five groups. These properties were similar for the five experimental groups (Table 1). Vr was measured in all neurons studied as soon as adequate access was attained, and it was relatively steady in the whole cell current-clamp mode. These data are presented in Table 1.

SC engraftment into the injury site prevents injury-induced changes in AP waveform

The voltage traces in Fig. 2, A–E, show APs recorded from the five experimental groups. The AP duration measured at half-maximal amplitude (AP half-width duration) was 1.75 ± 0.26 ms for the control neurons (Fig. 2, A and F; Table 2). In the ligation group the AP half-width duration was reduced to 0.94 ± 0.12 ms (Fig. 2, B and F; Table 2). In the nerve crush group where axons are completely axotomized but allowed to regenerate through the distal nerve stump, AP duration did not change (1.79 ± 0.38 ms; Fig. 2, C and F; Table 2). APs studied 2 wk after transplantation of SCs into the ligated nerve indicated that the AP narrowing observed in the ligated group did not occur in the transplant group (1.55 ± 0.22 ms; Fig. 2, D and F; Table 2). However, AP narrowing did occur in the sham group (0.83 ± 0.13 ms; Fig. 2, E and F; Table 2). The rise and fall times (RT and FT, respectively) were decreased, and the corresponding maximum slopes (MSR and MFR, respectively)

| Table 1. DRG membrane properties of normal, injured, and Schwann cell engrafted nerves |
|----------------------------------|---------------|---------------|---------------|---------------|---------------|
| Cell Membrane Properties        | Control       | Ligation      | Nerve crush   | SC Graft      | Sham Graft    |
| Size, μm                        | 39.2 ± 0.4 (28) | 39.0 ± 0.4 (23) | 38.3 ± 0.5 (32) | 37.3 ± 0.4 (35) | 38.6 ± 0.4 (22) |
| Cap, pF                         | 58.6 ± 3.1 (28) | 60.3 ± 2.4 (23) | 59.5 ± 2.3 (32) | 64.0 ± 3.0 (35) | 67.9 ± 2.4 (22) |
| Rc, MOΩ                         | 272.1 ± 22.3 (28) | 245.7 ± 15.5 (23) | 224.2 ± 19.5 (32) | 256.5 ± 29.4 (35) | 222.1 ± 4.7 (22) |
| Vr, mV                          | −67.0 ± 1.1 (28) | −65.5 ± 0.8 (23) | −63.9 ± 0.7 (32) | −64.2 ± 0.7 (35) | −64.8 ± 4.7 (22) |

Values are means ± SE with n in parentheses.
increased in the ligated and sham groups but not in the SC transplant group (Table 2). Thus the AP narrowing and the increased RT and FT in the ligation group were prevented by SC transplantation into the cut and ligated nerve end.

**SC engraftment into the nerve injury site prevents injury-induced changes in voltage threshold and AP utilization time**

AP threshold was measured by application of step depolarizations. The voltage threshold for spike initiation was 30.3 ± 1.7 mV for the control group and 34.9 ± 1.1 mV for the ligation group. Voltage threshold was also lowered in the sham group (37.0 ± 1.3 mV) but not in the crush (37.2 ± 1.4 mV) and SC transplant group (31.7 ± 1.3 mV). These data are summarized in Fig. 2F and Table 2. Moreover, the utilization time (UT) for spike initiation was decreased from 1.43 ± 0.06 ms (control) to 1.22 ± 0.05 and 1.23 ± 0.05 ms in the ligated and sham groups, respectively. The SC transplant group (1.37 ± 0.08 ms) and the nerve crush group (1.36 ± 0.08 ms) did not show changes in UT (data summary in Fig. 2G and Table 2). These changes in voltage threshold, together with decreased utilization time indicate that DRG neurons of the ligated group can be more readily activated and that SC transplantation prevents these changes.

**Effects of nerve injury and SC engraftment on AP inflections**

The APs of the medium-sized cutaneous afferent DRG neurons were characterized as inflected (A1) or noninflected (A0) based on the presence or absence of an inflection on the falling phase of the AP (Fig. 3A, top). The AP inflections were evident as two peaks in waveforms obtained after

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**Fig. 2.** Typical somatic action potentials recorded from cutaneous afferent DRG neurons in the control (A), nerve ligation (B), nerve crush (C), Schwann cell transplantation (D), and nerve ligation and vehicle-injection groups (E). Comparison of the AP duration, utilization time, and voltage threshold between the experimental groups are shown in F–H, respectively. (*P < 0.05, **P < 0.01, for comparisons with normal group; ^P < 0.05, for comparisons with ligation group; !!P < 0.01, for comparisons between Schwann cell (SC) grafting and sham group)
digital differentiation (dV/dt, clampfit 8.2; Fig. 3B, bottom). More than half of the control APs (57.1%, 16/28) displayed varying degrees of inflection on the falling phase of the AP (Fig. 3C, Table 2). In the ligation group, 17.4% (4/23) of the APs were inflected, and 46.9% inflected APs were observed in the crush group (15/32). In the SC graft group, 37.1% (13/35) of the APs were inflected as compared 13.6% (3/22) in the sham group. These data are summarized in Fig. 3C and in Table 2.

**Effects of nerve injury and SC engraftment on AP AHP**

Representative APs and AHPs recorded from DRG neurons for the five experimental groups are shown in Fig. 4, A–E. Comparison measurements were obtained for AHP amplitude (Fig. 4F), AHP duration (80% recovery; Fig. 4G) and the area of the AHP (Fig. 4H). AHP amplitude, duration, and area were reduced in the ligation, crush, and sham groups. These parameters of the AHP were not changed in the SC transplantation group (Fig. 4, F–H, and Table 3). Thus changes observed in the

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**TABLE 2. DRG action potential properties of normal, injured, and Schwann cell engrafted nerves**

<table>
<thead>
<tr>
<th>AP Properties</th>
<th>Control</th>
<th>Ligation</th>
<th>Nerve Crush</th>
<th>SC Graft</th>
<th>Sham Graft</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current threshold, nA</td>
<td>0.53 ± 0.05 (28)</td>
<td>0.54 ± 0.04 (23)</td>
<td>0.67 ± 0.05 (32)</td>
<td>0.64 ± 0.04 (35)</td>
<td>0.54 ± 0.06 (22)</td>
</tr>
<tr>
<td>Voltage threshold, mV</td>
<td>−30.3 ± 1.7 (28)</td>
<td>−34.9 ± 1.1 (23)*</td>
<td>−32.2 ± 1.4 (32)</td>
<td>−31.7 ± 1.3 (35)</td>
<td>−37.0 ± 1.3 (22)**</td>
</tr>
<tr>
<td>UT, ms</td>
<td>1.43 ± 0.06 (28)</td>
<td>1.22 ± 0.05 (23)*</td>
<td>1.36 ± 0.08 (32)</td>
<td>1.37 ± 0.08 (35)</td>
<td>1.23 ± 0.05 (22)*</td>
</tr>
<tr>
<td>Amplitude, mV</td>
<td>113.3 ± 1.9 (28)</td>
<td>112.2 ± 2.1 (23)</td>
<td>109.4 ± 1.4 (32)</td>
<td>111.0 ± 1.6 (35)</td>
<td>111.0 ± 1.3 (22)</td>
</tr>
<tr>
<td>Overshot amplitude, mV</td>
<td>46.6 ± 1.7 (28)</td>
<td>46.2 ± 1.6 (23)</td>
<td>44.1 ± 1.4 (32)</td>
<td>45.8 ± 1.3 (35)</td>
<td>44.6 ± 1.4 (22)</td>
</tr>
<tr>
<td>Half-width duration, ms</td>
<td>1.75 ± 0.26 (28)</td>
<td>0.94 ± 0.12 (23)**</td>
<td>1.79 ± 0.38 (32)^</td>
<td>1.55 ± 0.22 (35)^</td>
<td>0.83 ± 0.13 (22)***</td>
</tr>
<tr>
<td>90% RT, ms</td>
<td>1.38 ± 0.08 (28)</td>
<td>1.04 ± 0.05 (23)**</td>
<td>1.18 ± 0.08 (32)</td>
<td>1.23 ± 0.10 (35)</td>
<td>0.99 ± 0.05 (22)**</td>
</tr>
<tr>
<td>90% FT, ms</td>
<td>2.03 ± 0.31 (28)</td>
<td>1.01 ± 0.14 (23)*</td>
<td>2.86 ± 0.96 (32)</td>
<td>1.86 ± 0.31 (35)</td>
<td>0.88 ± 0.16 (22)!</td>
</tr>
<tr>
<td>MSR, V/s</td>
<td>254.1 ± 25.0 (28)</td>
<td>334.3 ± 23.1 (23)*</td>
<td>314.2 ± 24.2 (32)</td>
<td>313.8 ± 21.8 (35)</td>
<td>363.4 ± 18.9 (22)**</td>
</tr>
<tr>
<td>MSF, V/s</td>
<td>−115.2 ± 13.4 (28)</td>
<td>−167.5 ± 12.9 (23)**</td>
<td>−133.3 ± 14.4 (32)</td>
<td>−134.3 ± 13.4 (35)</td>
<td>−194.5 ± 14.6 (22)***</td>
</tr>
<tr>
<td>Inflected, %</td>
<td>57.1 (16/28)</td>
<td>17.4 (4/23)**</td>
<td>46.9 (15/32)^</td>
<td>37.1 (13/35)</td>
<td>13.6 (3/22)**</td>
</tr>
</tbody>
</table>

Values, except for percentages, are means ± SE with n in parentheses. UT, utilization time; RT and FT, rise and fall time; MSR and MSF, rise and fall maximum slope. *P < 0.05, **P < 0.01, comparisons with normal DRG neurons. \^P < 0.05, \^P < 0.01, comparisons with ligation group. !P < 0.05, !!P < 0.01, comparisons between Schwann cell (SC) graft and sham group.

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![Fig. 3. APs (top) with (A) or without (B) an inflection on the falling phase of the spike were recorded from cutaneous afferent DRG neurons evoked by 30-ms pulses ranging from −0.3 to 1.8 nA in 0.2-nA steps. Bottom: differentiated signals (digital differentiation, Clampfit 8.2). C: inflection rates (%) among the 5 experimental groups (**P < 0.01, for comparisons with normal group; ***P < 0.05, for comparisons with ligation group). Note that the number of inflected spikes is reduced in the ligation and sham group.](http://jn.physiology.org/ by 10.220.33.4.on June 22 2017)
AHP after nerve ligation were reduced in the SC transplantation group.

**Distribution and morphology of engrafted GFP⁺-SCs into ligated sciatic nerve**

The GFP-expressing SCs integrated into the cut nerve end and distributed longitudinally. Figure 5A is a longitudinal frozen section of the sciatic nerve and neuroma (right) 2 wk after nerve ligation and SC transplantation. GFP-expressing cell profiles can be seen in the neuroma head (Fig. 5B) and are distributed longitudinally in the proximal nerve trunk (Fig. 5, A and C). In neurofilament immunostained sections, the relationship to GFP-SCs and axons can be identified. In the neuroma head, the engrafted GFP-SCs distributed as spheroidal and

### TABLE 3. DRG afterhyperpolarization properties

<table>
<thead>
<tr>
<th>AHP Properties</th>
<th>Control</th>
<th>Ligation</th>
<th>Nerve Crush</th>
<th>SC Graft</th>
<th>Sham Graft</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amplitude, mV</td>
<td>-17.5 ± 1.1 (27)</td>
<td>-13.1 ± 1.3 (23)⁺</td>
<td>-16.7 ± 1.0 (32) ^=</td>
<td>-15.9 ± 1.0 (33)</td>
<td>-12.4 ± 1.4 (22)⁺⁺</td>
</tr>
<tr>
<td>80% recovery duration, ms</td>
<td>51.5 ± 8.2 (27)</td>
<td>28.1 ± 7.5 (23)⁺⁺</td>
<td>28.4 ± 4.3 (32)⁺⁺</td>
<td>40.9 ± 8.2 (33)</td>
<td>17.1 ± 2.7 (22)⁺⁺⁺</td>
</tr>
<tr>
<td>80% recovery area, mV*ms</td>
<td>-204.3 ± 38.0 (27)</td>
<td>-122.0 ± 27.3 (23)⁺⁺⁺</td>
<td>-165.4 ± 25.2 (32)⁺⁺⁺</td>
<td>-216.7 ± 44.3 (33)</td>
<td>-88.0 ± 16.9 (22)⁺⁺⁺⁺</td>
</tr>
</tbody>
</table>

Values are means ± SE with n in parentheses. *P < 0.05; **P < 0.01, comparisons with normal DRG neurons. ^=P < 0.05, comparisons with ligation group. !P < 0.05, !!P < 0.01, comparisons between SC graft and sham group.

**FIG. 4.** Slow sweep action potentials with afterhyperpolarization (AHP) recorded from the 5 experimental groups (A–E). Comparison of AHP amplitude, duration and area are shown in F–H, respectively. (*P < 0.05, **P < 0.01, for comparisons with normal group; ^=P < 0.05, for comparisons with ligation group; !P < 0.05, !!P < 0.01, for comparisons between SCs grafting and sham group)
fusiform cells with short thin processes that intermingled with axons but did not form prominent longitudinal association with axons (Fig. 5D). However, at more proximal regions of the nerve prominent longitudinal processes were formed by the GFP-SCs which ran in parallel to the axons (Fig. 5E).

**DISCUSSION**

After sciatic nerve section, the transected axons die back for a few millimeters over the course of the first few days after injury, and then regenerate toward distal targets or in the case of nerve ligation, toward the ligation (Ramon y Cajal 1928). Just proximal to the site of ligation a bulbous enlargement or neuroma forms. The neuroma is a complex mass containing numerous nonmyelinated axonal sprouts that grow circuitously within the neuroma. Much work indicates that the neuroma can be the site of ectopic impulse generation and a source of neuropathic pain (for reviews, see Devor 1994; Kocsis and Devor 2000). Moreover, a number of changes on the cell bodies of cutaneous afferents occur after neuroma formation including reorganization of sodium (Cummins et al. 1997; Everill et al. 2001) and potassium (Everill and Kocsis 1999) channels and GABA_A receptors (Oyelese et al. 1995, Oyelese and Kocsis 1996). Changes in sodium (Dib-Hajj et al. 1998) and potassium (Everill and Kocsis 2000) channels have been reduced by focal application NGF to the cut nerve end. Although SCs are known to be a source of a variety of neurotro-
phic factors (Carroll et al. 1997; Curtis et al. 1994; Hall et al. 1997; Hoke et al. 2002; Lee et al. 1995; Terenghi 1999), and neurotrophin production by the engrafted cells might contribute to our observations, we have no direct data to support this view. The results of the present study indicate that transplantation of SCs ameliorated injury-induced changes in AP waveform of medium-sized cutaneous afferents. This neuronal size population likely includes Aβ and possibly Aδ neurons but clearly excludes C-type neurons.

**SCs engrafted into the proximal region of a ligated sciatic nerve survive and migrate**

GFP-expressing SCs transplanted into transected sciatic nerve survived within the nerve. The donor cells were observed both within the head of the neuroma near the ligature and along the proximal nerve trunk. Many cells within the head of the neuroma did not appear to be longitudinally associated with axons but rather were spheroidal or bipolar spindle-shaped cells with thin elongated processes. In contrast, many of the proximal cells were distributed longitudinally in association with axons. GFP+SCs were observed several millimeters proximal to the neuroma head. It is likely that the donor SCs were able to compete with endogenous SCs in the proximal nerve stump to longitudinally associate with axons, and that they maintained a spheroidal phenotype within the neuroma head. Moreover, the GFP-SCs survived and integrated within the nerve without immunosuppression, indicating that cell rejection did not occur. However, we cannot rule out the possibility of an immune response by the engraftment that could possibly contribute to the observed changes in AP properties.

**Changes in DRG AP properties after nerve ligation and neuroma formation**

A number of changes in spike waveform have been reported after peripheral nerve injury including spike broadening (Kim et al. 1998; Stebbing et al. 1999) and narrowing (Oyelese et al. 1995; 1997). Action potential broadening was reported in nerve injury models where axons were transected but not encapsulated to prevent outgrowth in peripheral tissues (Kim et al. 1998; Stebbing et al. 1999). We found that if the nerve was ligated and encapsulated to prevent regeneration and reinnervation of target, the APs of identified medium-sized cutaneous afferents were narrowed and the inflection on the falling phase of the AP was eliminated or reduced as studied 2 wk after injury. Indeed, previous studies indicate that at 2 wk after nerve encapsulation that a kinetically slow Na+ current is reduced and that a singular fast Na+ current predominates in medium-sized cutaneous afferents (Oyelese et al. 1997). Moreover, the AHP amplitude and duration were both reduced 2 wk after nerve ligation. It is interesting in this regard that both sustained (K+ current) and a transient K+ (A current) currents of DRG neurons are reduced to about one-half in this injury model (Everill and Kocsis 1999); this may account for the changes in AHP properties.

**SCI transplantation reduces injury-induced changes in AP properties**

Changes in AP waveform, sodium (Oyelese et al. 1997), and potassium currents (Everill and Kocsis 2000) of DRG neurons induced by nerve ligation and encapsulation are reduced by direct application of NGF to the nerve injury site. It was hypothesized that target-derived NGF plays a role in regulating sodium and potassium channel organization on cutaneous afferent fibers after target disconnection (Dib-Hajj et al. 1998; Everill and Kocsis 2000; Oyelese et al. 1997). SCs are known to play a critical role in modulating sodium channel isoform expression in spinal sensory neuron either in vivo (Rasband et al. 2003) or in vitro (Hinson et al. 1997). The altered sodium channel isoform organization due to the lack of normal neuroglia contact may contribute to the pathophysiological changes in nerve injury or demyelinating diseases. Introduction of exogenous SCs into the nerve-injury site reduced AP narrowing, the reduction of the inflected APs, and changes in the AHP. One possibility to account for these observations is that the engrafted SCs provided neurotrophic support that could be transported retrogradely to the DRG cell body to substitute for the loss of skin-derived neurotrophins. It is interesting that the changes in spike waveform were reduced in the nerve crush model where the axons are completely transected but allowed to regenerate through the SC-enriched distal nerve segment and to peripheral targets. Both NGF and p75 receptor are upregulated on the SCs (Heumann et al. 1987; Taniuchi et al. 1988) of the distal segment and may provide sufficient trophic support during the regeneration process to ameliorate ion channel changes associated with changes in AP waveform. As mentioned, there are a number of potential neurotrophins besides NGF including neurotrophin-3 (NT-3), brain-derived neurotrophic factor (BDNF), glial line-derived neurotrophic factor (GDNF), ciliary neurotrophic factor (CNTF), leukemia inhibitory factor (LIF), and neurotrophins (Carroll et al. 1997; Curtis et al. 1994; Hall et al. 1997; Hoke et al. 2002; Lee et al. 1995; Terenghi 1999) that could contribute to this effect; we present no direct data implicating a particular neurotrophin. We hypothesize that transplantation of cultured SCs into the nerve-injury site provides an additional cellular neurotrophin source that may extend neurotrophin availability and thus limit changes in ion channel expression and changes in spike waveform. Clearly, additional biochemical studies will be necessary to directly address this hypothesis.

In the uninjured animal, cutaneous afferents associate with beta-keratinocytes in skin; beta-keratinocytes are a known source of at least one neurotrophin, NGF (Yiangou et al. 2002). The beta-keratinocyte may provide an ambient level of neurotrophin to maintain and stabilize appropriate Na+ and K+ channel organization of cutaneous afferents. In the case of neuroma formation, such an ambient source of neurotrophin may not be adequate to compensate for the lack of target-derived neurotrophic factor to maintain cell survival or normal function (Heumann et al. 1987; Johnson et al. 1985; Terenghi 1999), thus leading to altered ion channel expression and altered spike waveform (Oleyese et al. 1995, 1996). After nerve ligation, the endogenous SCs will require time to dissociate from the degenerating axon segments before they upregulate their expression of neurotrophins. The engrafted SCs may be primed by the dissociation protocol to more rapidly express neurotrophins and their receptors compared with the endogenous SCs in the proximal segment of transected nerve. The engrafted SCs may provide a transient source of neurotrophin during a critical period after nerve ligation before resident SCs are capable of doing so. The results of this study indicate that focal implantation of SCs into encapsulated blind end neuroma
reduces injury-induced changes in primary cutaneous afferent AP waveform. This suggests that a cellular therapeutic approach may be considered in stabilizing AP waveform after traumatic nerve injury.

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