

The present study assessed whether intact tibial nerves are successful unmyelinated fiber donors to the sectioned fibular nerves using SEN with a gap, using a rigid silicone tube interposed between the lateral portion of the intact tibial nerve (without an epi-perineurial window) and the distal stump of the fibular nerve. We report here that the side-to-end tubulization (SET) is a reliable reconstructive technique and can promote a remarkable plasticity of unmyelinated axons.

MATERIALS AND METHODS

Animals and Surgical Procedures

All experiments were conducted with approval from the Animal Experimentation Ethics Committee at the Medical School of Ribeirão Preto, University of São Paulo (protocol number 024/2010). Ten 7- to 8-week-old female Wistar rats (250–300 g) were used. Food and water were provided ad libitum. The animals were anesthetized using ketamine chloridrate (75 mg/kg) and xylazine (15 mg/kg), administered intraperitoneally. The right sciatic nerve and its branches were exposed, and the common fibular nerve was sectioned 3 mm from its origin. The proximal stump was then buried and sutured into the adjacent musculature. A silicone tube (6 mm in length \times 1.47 mm inner diameter \times 1.96 mm outer diameter; DeganiaSilicone Ltd., Israel) was attached to the adjacent lateral portion of the intact tibial nerve, and the distal stump of the common fibular nerve was sutured to the other end of the tube. All sutures were made with 10-0 nylon (Bioline) sutures. This procedure is referred to as SET (Fig. 1). The distal segment of the left common fibular nerve was used as normal (sham group). Fibular nerves were fixed in situ using 2% glutaraldehyde in 0.025 M, pH 7.4 sodium cacodylate buffer and dissected 70 days after tubulization. After removal of the nerves, the animals were euthanized using large quantities of the previously described anesthetics.

Histological Evaluation and Morphometry

The distal segment of the fibular nerve was sectioned 3 mm from the distal end of the tube, immersed in a 2% glutaraldehyde solution (48 h at 4°C), washed with sodium cacodylate buffer, postfixed in 1% osmium tetroxide, progressively dehydrated, and embedded in epoxy resin (Epon 812, Electron Microscopy Sciences, Hatfield, PA).

Samples were then cut into 0.5- μ m-thick sections using a microtome (MT 6000XL- RMC, Boeckeler Instruments Inc., Tucson, AZ), stained with toluidine blue, and mounted using Entellan (Sigma-Aldrich Co., Darmstadt, Germany) for analyses using light microscopy. Details of the methods have previously been published.²

Transmission Electron Microscopy Procedures

An ultramicrotome (Carl Zeiss, model G/214711) was used to cut the plastic embedded blocks into 80-nm-thick sections, which were placed on oval grids covered with 5% Formvar film (Formvar solution in ethylene dichloride, Electron Microscopy Sciences Inc., Hatfield, PA) and stained with 5% uranyl acetate and 0.5% lead citrate.

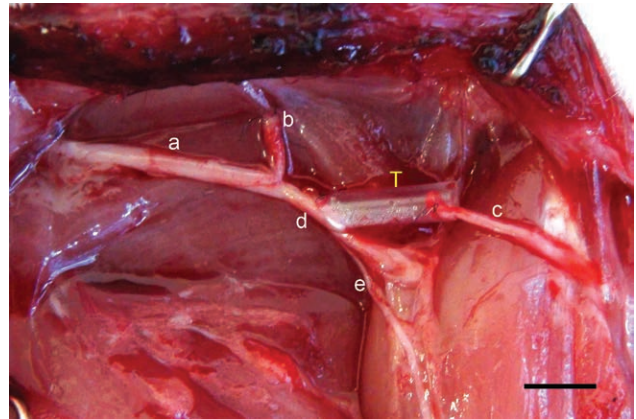


Fig. 1. SET in the right rat fibular nerve immediately after surgery, using a 6-mm silicone tube (T). A 5-mm segment was removed between the nerve stumps. (a) Sciatic nerve, (b) proximal stump of the common fibular nerve, (c) distal stump of the common fibular nerve, (d) tibial nerve, (e) sural nerve. Bar = 3 mm.

The digitalization was obtained using a transmission electron microscope (JEM-100CXII, JEOL Ltd., Tokyo, Japan) equipped with a digital camera (Hamamatsu ORCA-HR, model C4742-51- 12HR). Twenty nerves were assessed, 10 of each group. The cross-sectional fascicular area of the nerve was completely scanned, and images of the endoneurial content were obtained. Sequential photomicrographs were manually collected at 14,000 \times magnification while scanning occurred. We scanned means of 62% and 76% of the fascicle area from the sham group and the SET group, respectively. Each digital image obtained from the microscopic fields was 14.0 μ m wide \times 14.0 μ m high and 1,024 \times 1,024 pixels in tagged image file format. All images were analyzed. Owing to variability in the fascicular area,² we obtained a mean of 413 images per nerve.

Assessment of Unmyelinated Axons

Unmyelinated fibers and their axonal area were measured. The total number of axons, axonal density (axons/mm²), and the minimum axonal diameter (μ m) were measured using ImageJ 1.47 software (National Institutes of Health, Bethesda, MD), as described above. Unmyelinated axons were identified using established criteria,⁴³ as follows: (1) circular or oval profile; (2) surrounded by Schwann cell cytoplasm forming mesaxons; (3) axoplasm that was clearer than the cytoplasm of the respective Schwann cell; (4) clustering into “units,” which had a direct relationship with Schwann cells; and (5) basal lamina externally surrounding each fiber unit consisting of Schwann cells and axons.

Only regular sectioned fibers and axons with circular shapes were reported. Measurements of the internal fascicular area (mm²) were similar to those obtained in semithin sections; since semithin sections produced clearer images, we used them to measure the fascicular areas.²

Statistical Analyses

Student's *t* tests were used to compare frequencies and percentage minimum diameter data between groups, and

a χ^2 test was used to compare frequencies of fiber diameters between groups. All values are reported as the mean \pm SDs. Data were analyzed using SPSS v.17.0 (SPSS Inc., Chicago, Ill.) statistical software. Differences were considered significant when $P \leq 0.05$.

RESULTS

Seventy days after tubulization, connections between the nerve stumps occurred in all animals of the SET group, and the macroscopic aspects of the regenerated area appeared similar to normal fibular nerves (Fig. 2). In 5 of the 10 animals, the fascicular area of regenerated nerves was lower than that of normal nerves (Fig. 3A), whereas the other five animals had fascicular areas similar to normals. The perineurium, which was composed of one layer in the normal fibular nerve, had 5 layers in the SET group, as shown in Figure 3B. Myelinated and unmyelinated fibers, Schwann cell nuclei, endothelial cells, and mast cells were identified in all samples. However, these features were observed in greater numbers in regenerated nerves than in normals. The unmyelinated fibers were similar to normal nerve fibers, and usually consisted of a few axons, with considerable variability in both area and circularity. The unmyelinated axons had greater variation in their minimum diameters and there were a greater number of these axons in the SET group than in the sham group (Fig. 4A). Three nerves from the SET group showed endoneurial content predominantly populated by collagen fibers, with only a few myelinated fibers, unmyelinated fibers, and other cell types (Fig. 4B). However, the number of unmyelinated fibers was significantly greater in the SET group than in the distal contralateral fibular nerve (Table 1). Additionally, the variability among nerves was greater in the SET group. The median number of axons in normal nerves was 1,987.5, whereas in the SET group, the median was 1,349.5.

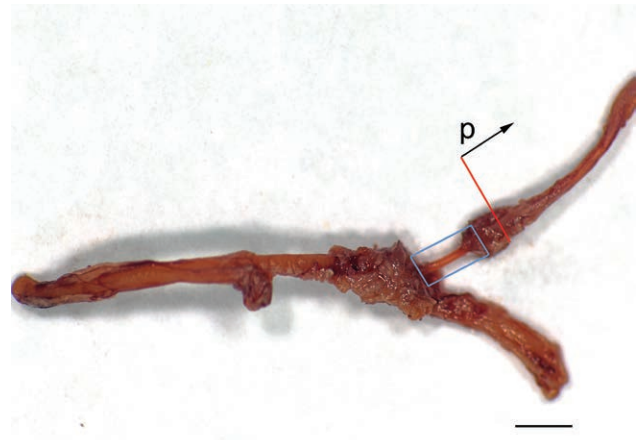


Fig. 2. Nerve segment from the right sciatic SET group, with branches. The blue rectangle represents the position of the silicone tube, with the regeneration cable inside. The red line represents the analysis site, which was 3 mm from the end of the silicone tube. The dashed arrow in black represents the nerve segment designed for proximal inclusion to the enclosing block (p). Bar = 5 mm.

The mean internal fascicular areas, number of axons, minimum axonal diameters, and axonal densities of the sham and SET groups are given in Table 1. The greater density of unmyelinated fibers in the SET group is explained by reduced fascicular area and increased variability in the number of unmyelinated axons. Figure 5 presents the frequency distribution for minimum axonal diameter in both groups. The axon diameters of the normal fibular nerve ranged from 0.2 to 2.8 μm . The minimum diameter of axons in the SET group ranged from 0.2 to 2.6 μm . There were more unmyelinated axons between 1.0 and 1.2 μm in the sham group, whereas there were more axons with minimum diameters between 0.4 and 0.6 μm in the SET group (Fig. 5).

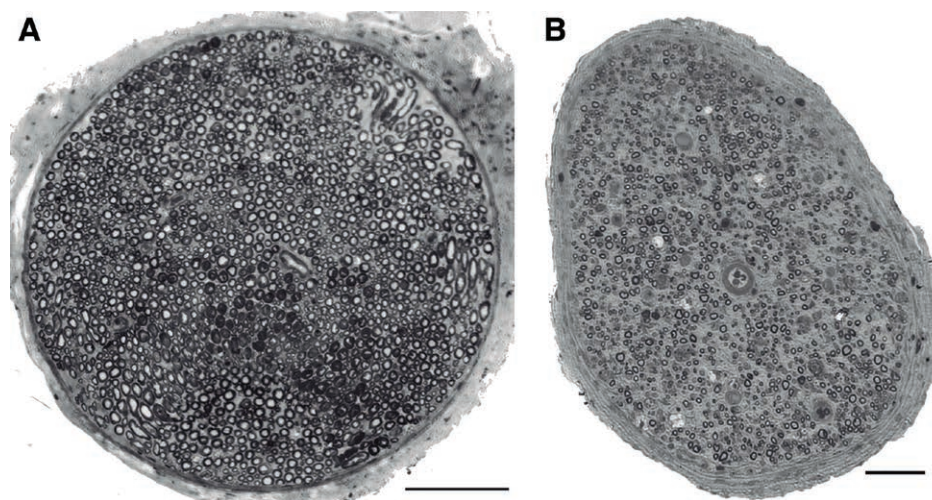


Fig. 3. Photomicrographs of the fascicular areas of the distal segments from normal common fibular nerves and from partially reconstructed nerves 70 days after surgery. Note the difference in the size of the fascicular area between groups. Panel B shows a perineurium constituted of 5 layers, which is in contrast to 1 layer in the normal fibular nerve. A, Normal distal fibular nerve (sham group). B, Reconstructed distal fibular nerve 70 days after SET group, at the same level as the image in A. Semithin sections (0.5 μm). Toluidine blue. Bar = 100 μm .

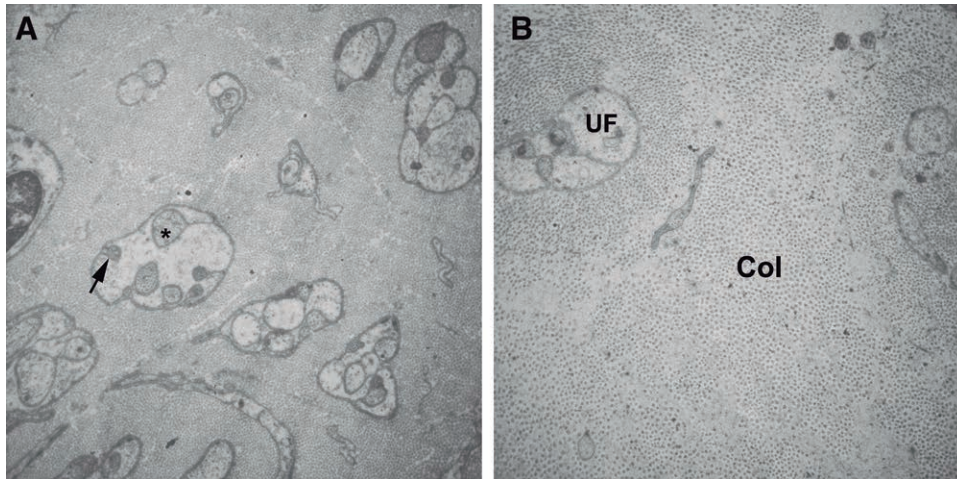


Fig. 4. Electron micrographs of cross-sections of distal segments from 2 common fibular nerves from the SET group. A, There are a large number of unmyelinated fibers; however, Schwann cell units are associated with one or few axons. Most axons have small diameters compared with the sham group. However, the variability in axonal diameter was often high within the same unit (arrow = unmyelinated axon of smaller caliber; * = unmyelinated axon of larger caliber). B, Predominance of collagen fibers (Col) involving an unmyelinated fiber (UF) with 3 unmyelinated axons of small diameters. Transmission electron microscopy—14,000x. Bar = 2 μm.

Table 1. Morphometric Parameters

Group	Nerve Area (mm ²)	No. Axons	Minimum Axonal Diameter (μm)	Density of Axons (Axons/mm ²)
Sham (n = 10)	0.13 ± 0.02	1,882 ± 270.9	0.968 ± 0.10*	13,935.0 ± 1,875.8
SET (n = 10)	0.11 ± 0.04	2,012 ± 1,060.8	0.648 ± 0.08	18,733.3 ± 5,668.6*

Data are presented as mean ± SD, except for number of axons.

*Indicates a significant difference between the sham group and experimental group (SET; $P \leq 0.05$; Student's *t* test).

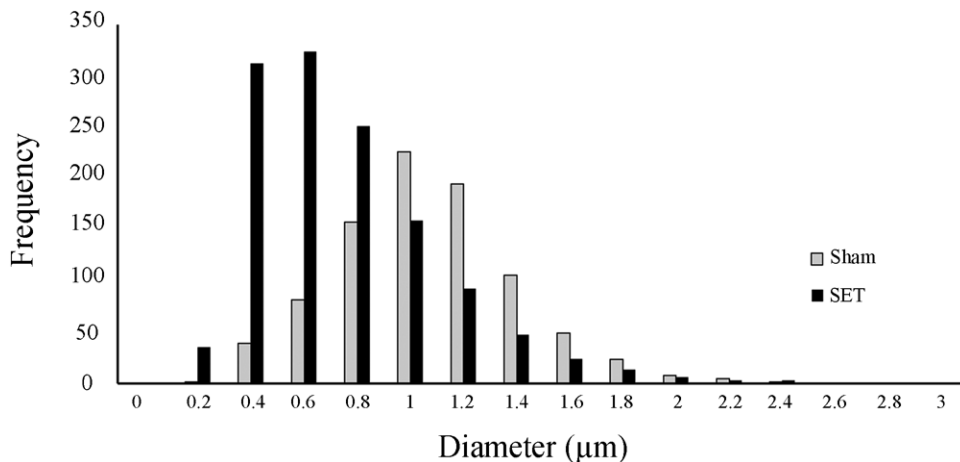


Fig. 5. Histogram of axon numbers in the left normal distal fibular nerve (normal group = sham) compared with the right distal fibular nerve reconstructed using SET. There is a shift to the left, indicating reduced diameters of unmyelinated fibers in the SET group, particularly in the 0.4–0.6 μm diameter range.

DISCUSSION

The unmyelinated fibers in this study traversed the 5-mm gap created by SET and reached the distal segment of the fibular nerve. A literature search did not reveal previous experimental studies on unmyelinated fibers with SEN, either with or without tubulization from the normal

tibial nerve to the fibular stump of a previously divided fibular nerve, which is the most common experimental model for SEN. We conducted tubulization using a rigid silicon tube with the intention that future studies can build on our work and observe the actions of nerve growth promoters. Although both rigid and nonrigid tubes have been

used in previous studies, they have not previously been used in studies of unmyelinated fibers with SEN.^{33,34,37}

The study evidence a difference between fascicular areas in half of the SET group that was lower than in the sham group. At this time, 70 days post surgery, data describing nerves area in current developing at axonal phase, referring to the regenerating process not completed yet.⁴⁴ Data about perineurial layers difference between groups support this report, supposing that at the final regeneration process, the perineurial layers will be condensed in a unique multilayered cellular membrane.⁴⁵

Although some previous studies have demonstrated that a perineurial window induces greater collateral fibers growth,^{46,47} we chose not to use an epi-perineurial window, due to work of Viterbo et al.²⁴ not revealing any differences between end-to-side neurorrhaphies with and without perineurial window concerning the morphological features. Our findings demonstrate profuse growth of unmyelinated fibers, which were often more numerous than the average number of fibers in normal nerves. Future studies are needed to compare the growth of sensory and motor fibers this tubulization experimental model.

Unmyelinated fibers in the fibular nerve are generally classified as 73% afferent (sensory) and 27% efferent (sympathetic).⁴⁸ The majority of quantitative studies of unmyelinated fibers in peripheral nerves are conducted on the sural nerve.

Unmyelinated fibers or C fibers carry thermal and pain sensations; therefore, the sural nerve is commonly studied because 92% of its axons conduct sensory information.⁶ A previous SEN study used the sural nerve as the donor nerve at the distal stump of the fibular nerve and evaluated myelinated and unmyelinated fibers. After 56 days, there were <800 axons observed 4mm from the neurorrhaphy site in the fibular nerve.³⁹ This SEN procedure was conducted without tubulization of the sural-fibular nerves; therefore, more profuse growth of unmyelinated fibers was expected. In the present study, we found a mean of 2,011 unmyelinated axons (median = 1,349.5) in the distal end of the tubulized fibular nerve. Despite the prominent axonal growth in our research, even applying a tubular scaffold on a gap comparing with the study with SEN above cited, we can't assert that tibial nerve are better donor nerve than sural nerve, because the experimental models approaches are different.

We studied adult rats (130 days) because the sciatic nerve has more axons during development than in adult rats. Axons are eliminated during the maturation process.⁴⁹ Furthermore, by using the tibial nerve as a source of unmyelinated fibers, we avoided the possibility of transitory sprouts that might experience further elimination, because axon growth appears to be stable in the tibial nerve. After axotomy or nerve crush injury in adult rats, the number of unmyelinated axons transiently increases before returning to normal numbers.⁵⁰ A previous experiment by Kovacic found that coaptating the rat sural nerve as a lateral donor to the fibular nerve resulted in stabilization of unmyelinated fiber growth at 2–4 months after surgery.³⁹ This leads us to suppose that our research time (70th day) contemplates the same situation.

The majority of the SEN studies in man contemplate the motor function recovery.^{14–20,51,52} Although some studies have evaluated sensory recovery after SEN,^{53–57} the thermal, tactile, and pain sensations have not been systematically evaluated and few studies applied appropriate analysis that refers to those kinds of findings.^{58–60}

Lack of those modalities of sensations can be potential causes of pressure ulcers and accidental injuries, as occur in some predominantly sensory neuropathies, such as leprosy and hereditary sensory neuropathies. So, it is recommended in the clinical setting, not only a clinical evaluation but a quantitative sensory test evaluation of those sensations of the skin supposedly reinnervated after nerve regeneration surgical procedure.

CONCLUSIONS

The present study demonstrates plasticity of unmyelinated axons with SET, using the tibial nerve as a donor. Long-term studies are needed to study changes in thermal and pain sensation that result from fibular nerve sectioning and to evaluate the stability of the unmyelinated fiber growth accomplished by SET. It is possible and even probable that in SEN without tubulization, even greater fiber growth occurs. The effect of nerve growth enhancers should also be studied, and SET is an appropriate model for these studies.

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