Delivery of nitric oxide-releasing silica nanoparticles for in vivo revascularization and functional recovery after acute peripheral nerve crush injury

Abstract

Nitric oxide (NO) has been shown to promote revascularization and nerve regeneration after peripheral nerve injury. However, in vivo application of NO remains challenging due to the lack of stable carrier materials capable of storing large amounts of NO molecules and releasing them on a clinically meaningful time scale. Recently, a silica nanoparticle system capable of reversible NO storage and release at a controlled and sustained rate was introduced. In this study, NO-releasing silica nanoparticles (NO-SNs) were delivered to the peripheral nerves in rats after acute crush injury, mixed with natural hydrogel, to ensure the effective application of NO to the lesion. Microangiography using a polymer dye and immunohistochemical staining for the detection of CD34 (a marker for revascularization) results showed that NO-releasing silica nanoparticles increased revascularization at the crush site of the sciatic nerve. The sciatic functional index revealed that there was a significant improvement in sciatic nerve function in NO-treated animals. Histological and anatomical analyses showed that the number of myelinated axons in the crushed sciatic nerve and wet muscle weight excised from NO-treated rats were increased. Moreover, muscle function recovery was improved in rats treated with NO-SNs. Taken together, our results suggest that NO delivered to the injured sciatic nerve triggers enhanced revascularization at the lesion in the early phase after crushing injury, thereby promoting axonal regeneration and improving functional recovery.

Key Words: crush injury; nerve injury, nerve regeneration; nitric oxide; peripheral nerve; revascularization; silica nanoparticles

Introduction

Peripheral nerve injury (PNI) is primarily caused by motor vehicle accidents, industrial accidents, stretching or crushing after falls, stab or laceration injuries, gunshot injury, and fractures (Tapp et al., 2019). PNI results in functional impairment or permanent disability and is associated with substantial social and personal costs to the affected individual (Tapp et al., 2019; Bergmeister et al., 2020). As axonal regeneration and growth support of Schwann cells diminishes with increasing time after an injury and distance from the injured site, it is a race against time for regenerating axons to regain satisfactory function (Chan et al., 2014). Thus, the acceleration of axonal regeneration may allow partial or complete functional recovery after PNI.

Manipulation of the nitric oxide (NO) supply in the neural...
The regenerative process is one possible way to boost axonal regeneration after PNI. NO is a crucial bioregulatory molecule in degenerative and regenerative processes of peripheral nerves after PNI, for example, in Wallerian degeneration (Kellihoff et al., 2002a), vascular remodeling in angiogenesis (Kellihoff et al., 2002b), and re-establishment of neuromuscular junctions (Kellihoff et al., 2002b; Moreno-Lopez, 2010). During neural regeneration, the NO supply depends on three major types of nitric oxide synthase (NOS) isoforms: constitutive neuronal (nNOS), endothelial (eNOS), and inducible (iNOS) forms (Gonzalez-Hernandez and Rustioni, 1999a; Moreno-Lopez, 2010). NO produced by eNOS in the peripheral neurons enhances reinnervation and reorganization of blood vessels, helping to remove cell debris by activating the infiltration of inflammatory cells (Kellihoff et al., 2002a, b) and recruitment of other cells necessary for regeneration from the bloodstream (Gonzalez-Hernandez and Rustioni, 1999a; Kellihoff et al., 2002b; Moreno-Lopez, 2010). NO originating from the upregulation of nNOS/iNOS plays a critical role in Wallerian degeneration and axon regeneration (Levy et al., 2001; Kellihoff et al., 2002a; Zachodnie and Levy, 2005).

Despite its important biological roles, NO has not been widely used as a therapeutic material in PNI because of the lack of suitable vehicles to facilitate storage and controlled and prolonged delivery. The silica nanoparticle system has been reported to chemically store NO molecules (up to 1780 nmol/mg) and spontaneously release them at physiological temperature and pH under aqueous conditions over a physiologically significant time scale (up to 30 hours) (Shin et al., 2007; Shin and Schoenfisch, 2008). Previous studies have shown the anti-microbial and anti-tumor effects of NO-releasing silica nanoparticles (NO-SNs) (Hetrick et al., 2008, 2009; Stevens et al., 2010). However, as nerve regeneration involves a complex series of biological steps, it is essential to determine a suitable carrier material to deliver the NO supply only to the lesion, to prevent negative effects on the surrounding tissue.

Herein, we describe a novel method for the application of NO-SNs in PNI in male rats (we used male mice to avoid the potential influence of the hormonal status or estrous cycles on regeneration in female mice. In addition, because traumatic peripheral nerve injury mostly occurs in male patients, translational research of nerve regeneration is usually performed with male animals), which involves mixing nanoparticles with a biocompatible natural hydrogel. Inspired by the controllable gelation behavior and biocompatibility of fibrin (Noori et al., 2017), this material was chosen as a carrier for NO-SNs so that the nanoparticles embedded in the glue-like fibrin gel may adhere to the damaged sciatic nerve of rats and release NO only at the site of injury. Histological analyses were performed to reveal revascularization and growth of myelinated axons. Furthermore, the physical performance of animals was recorded, because the ultimate aim of peripheral nerve regeneration is the recovery of functional performance, which is sufficient for the social comeback of the individual.

Materials and Methods

Synthesis of NO-SNs

The NO-SNs were synthesized and characterized as described previously (Shin and Schoenfisch, 2008; Hetrick et al., 2009). Briefly, an aminoalkoxysilane solution was prepared by dissolving 6.8 mM N-methylaminopropytrimethoxysilane (MAP3; Gelest, Tullytown, PA, USA) in a mixture of 16 mL ethanol (EtOH) and 4 mL TMAH (35 wt%) in deionized water (Milli-Q) for 2 minutes. The silane solution was then added to EtOH (22 mL) with an ammonia catalyst (6 M, 10% w/v) in water), and mixed vigorously for 30 minutes at 4°C. The precipitated nanoparticles were collected by centrifugation (2516 × g, 5 minutes), washed with EtOH several times, dried under ambient conditions for 1 hour, and stored in a sealed container at −20°C until use.

NO release kinetics of MAP3/NO nanoparticles was monitored in deoxygenated phosphate-buffered saline (PBS; 0.01 M, pH 7.4) at 37°C using a Sievers NOA 280i Chemiluminescence Nitric Oxide analyzer (Sievers Instruments, Boulder, CO, USA) (Shin et al., 2007; Shin and Schoenfisch, 2008). The instrument was calibrated with air passed through a zero filter (0 ppm (0.0001%) NO) and 45 ppm of NO standard gas (balance N₂). The morphology and size of the NO-SNs were analyzed using scanning electron microscopy.

Ethics statement

All animal experiments were conducted according to the protocol approved by the Institutional Animal Care and Use Committee of the Korea University College of Medicine (IACUC Approval No. 2016-0266) on November 30, 2017. This study was reported in accordance with the ARRIVE 2.0 guidelines (Animal Research: Reporting of In Vivo Experiments) (Percie du Sert et al., 2020).

Surgical procedure

Fifty-four male Lewis rats weighing 250–300 g were randomly divided into two groups of 27 each and received a standardized sciatic nerve crushing injury, as described in Figure 1. The rats were anesthetized using isoflurane inhalation; animals were sedated with 5% isoflurane in a plastic chamber and then moved on to the surgical field for a survival procedure maintained under 3% isoflurane administered via a mask placed over the mouth. The right limb and buttock were shaved, washed with 70% ethanol, and prepared with povidone iodine. The sciatic nerve was carefully exposed around the trifurcation site using gluteal splitting. All rats received sciatic nerve crushing injury (6 mm in length) on a site just proximal to the trifurcation, made by closing a new surgical needle holder (Integra Millex, Plainsboro, NJ, USA) to the third notch for 20 seconds (Sundem et al., 2016). Fibrin gel, a natural hydrogel, was subsequently applied around the crushed nerve, with and without NO-SNs. Only fibrin gel without NO-SNs was injected into animals of the control group: 1 µL of thrombin (T4648, Sigma-Aldrich, St. Louis, MO, USA; 50 U/mL in sterile water) was diluted with 37.5 µL of phosphate-buffered saline (PBS), and then, 12.5 µL of filtered fibrinogen solution (F8630, Sigma-Aldrich; 10 mg/mL in DPBS with Ca²⁺ and Mg²⁺) was added to the PBS-diluted thrombin solution just before the application of the gel to prevent pre-clotting. For the treatment of animals in the NO-treated group, NO-SNs were dissolved in PBS at a concentration 4-fold higher than the target concentration, and 37.5 µL of this solution was homogeneously mixed with 1 µL of thrombin. The same amount (12.5 µL without NO-SNs) and concentration of fibrinogen solution as used in the control group was added to the mixture of thrombin and NO-SNs. The final concentration of NO in the fibrin gel was 70 µM. After confirmation of clotting of the fibrin gel around the crush site, the skin was sutured with 4-0 Prolene, which was removed 14 days after surgery. All rats received a single intramuscular injection of ketoprofen (5 mg/kg) for immediate postoperative pain control and enrofloxacin (5 mg/kg) to prevent postoperative infection. After the surgery procedures, rats were returned to their cages and raised in an environment with a 12-hour light/dark cycle and provided with food and water ad libitum.

Figure 1  | Flowchart describing the experimental procedures and animal assignment.

NO-SN: Nitric oxide-releasing silica nanoparticle.
Microangiography
The effect of NO delivery on revascularization at the crushed nerve was analyzed using microangiography and immunohistochemical staining of nerves. We performed microangiography (Giusti et al., 2016) to evaluate revascularization around the crush site of the nerve 3 weeks after crush. The sciatic nerves were anesthetized with an intravenous injection of alfalfaxone (1–1.5 mg) into the ventral tail vein, the rib cage was carefully opened to expose the heart, and a cannula was inserted into the aorta through a small incision in the left ventricle. A small incision was made in the right atrium for drainage, and the region was flushed with saline mixed with heparin until the liver turned pale. This was followed by flushing with 200 mL of 4% paraformaldehyde. Blue-colored microangiographic polymer dye (Microfil®, Flowtech, Carver, MA, USA) (40 mL) was injected through the cannula under physiological pressure (100 mmHg) using an infusion pump (Harvard Apparatus, Holliston, MA, USA). The specimen was stored overnight at 4°C for complete polymerization of the injected dye. CDs, which were harvested 3 days after nerve crush, were stained with toluidine blue (89640, Sigma-Aldrich). The image of the cross-sectioned tibial division of the sciatic nerve at the digital level of the crushing site was captured at 200× magnification, using a charged-coupled device camera (DP21, Olympus, Tokyo, Japan) attached to a light microscope (BX46, Olympus, Tokyo, Japan) and then incubated with rabbit anti-goat CD34 (AF4117, R&D System, Minneapolis, MN, USA, 1:100, AB_2074613) overnight at 4°C. Sections were washed with PBS and then incubated with rabbit anti-goat IgG (Al-5000, Vector Laboratories, Burlingame, CA, USA, 1:100, AB_2336125) for 1 hour at 37°C. Staining for CD34, a transmembrane protein in the endothelial cells of blood vessels, was used to evaluate the number of regenerated vessels at the crush site. The number of vessels, which are shown as brown-colored cells lining the luminal structure, was counted manually using a light microscope (BX46, Olympus, Tokyo, Japan) in the whole nerve section at a magnification of 200×.

Quantitative analysis of the myelinated axons, the N ratio, myelin thickness, and G-ratio, harvested nerves at 3 or 6 weeks after the injury were stained with toluidine blue. The nerves harvested at 3 or 6 weeks after injury were embedded in epoxy resin, cut into 1-µm sections, 2 mm distal to the crush site, and stained with toluidine blue (89640, Sigma-Aldrich). The image of the cross-sectioned tibial division of the sciatic nerve at the digital level of the crushing site was captured at 200× magnification, using a charged-coupled device camera (DP21, Olympus, Tokyo, Japan) attached to a light microscope (BX46, Olympus). The images were analyzed for the area in a semiautomatic fashion using ImageJ to determine the number of myelinated axons and the N ratio, which was calculated as the total myelinated axon area divided by the total nerve area (Tobin et al., 2014; Lee et al., 2016). Myelin thickness and G-ratio, which were calculated as axonal diameter divided by axoglial diameter, were measured using 30 randomly selected axons in each image (300 axons were analyzed per group).

Histomorphometry analysis
The sciatic nerves were harvested at 3 days and 3 weeks after nerve crush. All nerves were fixed with 4% paraformaldehyde. The nerves harvested on day 3 after injury were embedded in paraffin, and 5-µm cross-sections were taken from the crush site. Immunohistochemical staining was performed using primary antibody goat anti-rat CD34 (AF4117, R&D system, Minneapolis, MN, USA, 1:100, AB_2074613) overnight at 4°C. Sections were washed with PBS and then incubated with rabbit anti-goat IgG (Al-5000, Vector Laboratories, Burlingame, CA, USA, 1:100, AB_2336125) for 1 hour at 37°C. Staining for CD34, a transmembrane protein in the endothelial cells of blood vessels, was used to evaluate the number of regenerated vessels at the crush site. The number of vessels, which are shown as brown-colored cells lining the luminal structure, was counted manually using a light microscope (BX46, Olympus, Tokyo, Japan) in the whole nerve section at a magnification of 200×.

Statistical analysis
All results are expressed as mean ± standard error of the mean (SEM). Mann-Whitney U test or Student’s t-test was used to detect statistical differences between two groups of vascular density and the number of vessels. One microangiographic specimen in the control group was excluded from the analysis because the specimen was improperly stained. Multiple unpaired t-test was used to detect statistical differences between two groups of SFI, muscle contraction force, muscle weight, and histologic analysis of nerve. Analysis of variance was used to detect statistical improvements in the SFI with time. Statistical analysis was performed using GraphPad Prism 9 (GraphPad Software, San Diego, CA, USA). Differences were considered significant at P-values < 0.05. A power analysis was performed based on the differences in SFI found between groups in previous publications (Elfar et al., 2008). Based on 35% difference detection and the desired power level of > 80%, we determined that a sample size of minimum five rats per group for each outcome is required (Elfar et al., 2008; Leem et al., 2017; Leem et al., 2020) had suggested that a group size of 3 or 4 would be sufficient for immunohistochemical studies. Biochemical and histological analyses were conducted blinded to the treatment, whereas in vivo microscopy and analysis were not performed blinded to the conditions of the experiments.

Results
Nitric oxide release from silica nanoparticles
The total amount of NO ([NO]) released from MAP3/NO nanoparticles was approximately 6.8 μM/mg, with a maximum NO flux ([NO]m) of 155,000 ppb/mg and a NO release half-life (t1/2) of 5 minutes. The diameter of MAP3/NO nanoparticles analyzed using scanning electron microscopy was approximately 120 nm (Figure 2), similar to the previously reported value (Shin et al., 2007).

Revascularization
The results of microangiography are presented in Figure 3. Based on microangiography evaluation, the regenerated endoneurial vessels passing through the crushing injury site were identified in specimens from all animals, except in one control specimen which was improperly stained and thus excluded from the analysis. Relative to the control group, more vessels stained with blue dye were found in the NO-treated group, indicating an increase in vascular density at the crushing site of the sciatic nerve was significantly greater in the NO-treated group than in the control group (P < 0.05).
The level of revascularization after treatment with NO-SNs. Microangiography was performed using a blue-colored polymer dye on day 3 after nerve crush injury to evaluate the level of revascularization after treatment with NO-SNs. The swollen and yellowish-colored area is the crush site of the sciatic nerve in the animal. There were more blue colored capillaries in the crush site in animals treated with NO-SNs (NO) compared with the control group (without NO-SNs), which suggest that the NO-SNs (MAP3/NO) promoted vascular proliferation in the crush site of rat sciatic nerve (n = 3/group); *P < 0.05 (Mann-Whitney U test). Data are expressed as the mean ± SEM. Group I: Crushed nerve; Group II: crushed nerve treated with NO-SNs. NO-SNs: Nitric oxide-releasing silica nanoparticles.

For direct identification of the newly formed vessels, immunohistochemical staining was performed targeting CD34, a marker for evaluating angiogenesis (Figure 4). The number of vessels stained with CD34 at the nerve sections in the control group is similar to the NO-treated group (P = 0.38).

Motor function of rat hind limbs The SFI for animals in the control group improved from 2 weeks to 5 weeks after injury (from 2 weeks to 4 weeks after injury, P < 0.0001; from 4 weeks to 5 weeks after injury, P < 0.01) and the SFI for animals in the NO-treated group improved from 2 weeks to 4 weeks after injury (P < 0.0001). From 2 weeks to 4 weeks after injury, the SFI of animals in the NO-treated group exhibited a significant functional improvement compared to that in the animals of the control group (P < 0.05; Figure 5C). In summary, after crush injury, NO-SN treatment accelerated both revascularizations (3 days after injury, P < 0.01, Figure 3) and recovery of sciatic nerve function (from 2 weeks to 4 weeks after injury, P < 0.05; Figure 5). However, the SFIs of the NO-treated and control groups were identical at 5 weeks after injury (Figure 5C).

Figure 5 | Walking track analysis for the calculation of the SFI. (A) Custom-made corridor for rats to walk straight. A dark cage was installed on the opposite side to guide the rat walk in the front direction. (B) Representative photographs of the footprint of a rat after crushing injury of sciatic nerve showing parameters for the calculation of the SFI. Compared with a normal paw, the injured side shows decreased intermediate toe spread (IT, blue) and toe spread (TS, black) and increased paw length (PL, red) at 2 weeks. Serial measurements at 4 and 6 weeks show gradual improvements, proving the recovery of motor function. (C) The SFI for animals in both groups improved from 2 weeks to 5 weeks after injury. Based on the SFI change, NO-SNs promoted functional recovery until 4 weeks after nerve crush injury. Improvements in sciatic function in group II was faster than in group I by 4 weeks of injury. n = 20/group (3–5 weeks) and 10/group (4–6 weeks). Data are expressed as the mean ± SEM. *P < 0.05 (multiple unpaired t-test). Group I: Crushed nerve/control; Group II: crushed nerve treated with NO-SNs. NO-SNs: Nitric oxide-releasing silica nanoparticles; SFI: sciatic functional index.

Muscle function and muscle atrophy As presented in Figure 6, NO-SNs enhanced the isometric tetanic muscle force of the TA at 3 and 6 weeks after injury, and a significant effect of NO-SNs was apparent, with a mean intergroup difference at 6 weeks after injury (P < 0.05).

The wet weight of calf muscle (gastrocnemius and soleus) in animals of the NO-treated group was higher compared with that in the control group at 3 and 6 weeks after injury. However, a statistically significant difference between the two groups was observed only at 3 weeks (P < 0.05), but not at 6 weeks (P = 0.1; Figure 7) after injury.

Nerve regeneration The number of myelinated axons and the N ratio increased over time in both groups (Figure 8B and C). The number of myelinated axons was significantly higher in animals treated with NO-SNs compared with the untreated animals 3 weeks after injury (P < 0.05; Figure 8B). However, this intergroup difference was not significant at 6 weeks after injury. The N ratio was higher in animals treated with NO-SNs compared with the untreated animals at 3 and 6 weeks after injury, but there was no significant difference between the two groups (Figure 8C). NO-SNs treated animals had significantly thicker myelin at 3 and 6 weeks after injury (P < 0.01; Figure 8D) and significantly lower G-ratio at 3 weeks after injury compared with the untreated animals (P < 0.0001; Figure 8E).

Discussion Our results demonstrate that NO-SNs (MAP3/NO) enhance revascularization and nerve regeneration, and improve the motor function in a rat model of sciatic nerve injury.

NO is a pivotal neuromediator in revascularization of peripheral nerves and vasodilation for cell recruitment after PNI (Keilhoff et al., 2002b; Zochodne et al., 2005). Endogenous NO levels increase as soon as 2 days after PNI, and are modulated by eNOS overexpression in the vasa nervosum of the distal stump (Gonzalez-Hernandez and Rustioni, 1999a). Endogenous NO generated by eNOS plays a...
NO-SNs prevent denervation-induced muscle atrophy.

The values of the muscle weight are presented as percentages of the contralateral side. The isometric tetanic force of Group I was higher than in group I at 3 and 6 weeks after injury. The statistically significant difference between the two groups was observed only at 6 weeks after injury (n = 10/group; *P < 0.05 (multiple unpaired t-test)). Group I: Crushed nerve; Group II: crushed nerve treated with NO-SNs. NO-SNs: Nitric oxide-releasing silica nanoparticles; ns: not significant.

Critical role in revascularization, remodeling of vessels, hyperemia, vasodilatation, and inflammatory cell infiltration (Rudic et al., 1998; Gonzalez-Hernandez and Rustioni, 1999a; Kelihoff et al., 2002b). A 2-day delay in revascularization was observed in eNOS knockout (KO) mice relative to wild-type mice after nerve transection and autogenous nerve grafting (Kelihoff et al., 2002b). NO-mediated vasodilatation helps to remove cell debris, recruit inflammatory cells, such as macrophages from the bloodstream, and support the metabolic processes for regenerating axons and other cellular elements (Kelihoff et al., 2002a; Moreno-Lopez, 2010).

NO promotes revascularization following PNI, but also plays a critical role in Wallerian degeneration, axonal regrowth, and functional recovery after PNI (Gonzalez-Hernandez and Rustioni, 1999a, b; Kelihoff et al., 2002a; Kikuchi et al., 2018). Functional recovery is impaired after nerve transection and repair in nNOS KO mice because of delayed Wallerian degeneration, increased uncontrolled axonal sprouting, and delayed nerve regeneration (Kelihoff et al., 2002a).

nNOS is a major factor in axonal sprouting and synaptogenesis in nNOS KO mice compared with wild-type mice after nerve transection (Gonzalez-Hernandez and Rustioni, 1999a, b). Levy et al. (2001) made similar observations about Wallerian degeneration, axonal regrowth, and functional recovery in INOS KO mice. Thus, the lack of NO-mediated stimulation in nNOS or INOS KO mice might lead to delayed structural changes in nerves and functional recovery. Kelihoff et al. suggested that manipulation of the NO supply might offer therapeutic options for the treatment of PNI (Kelihoff et al., 2002a).

In this context, it is worth noting that in the present study, locally delivered exogenous NO supply might enhance the revascularization process and functional recovery in the case of PNI. Functional recovery following traumatic or iatrogenic peripheral nerve injury is unpredictable. These injuries require many complex microsurgical procedures, such as neurorrhaphy, nerve grafting, and nerve transfer. Regardless of the procedure, a patient undertakes, rapid revascularization of damaged nerves or grafts is mandatory to maintain or restore viable Schwann cells and to improve the rate and quality of nerve regeneration (Bunge, 1994; Best et al., 1999; Mompeo et al., 2003; Kannan et al., 2005; Sekiguchi et al., 2012).

NO is a crucial signaling molecule associated with several physiological processes, including angiogenesis, inhibition of thrombus formation, wound healing, and neurotransmission; therefore, several studies have focused on NO-based therapies (Carpenter and Schoenfisch, 2012; Riccio and Schoenfisch, 2012; Seabra et al., 2015). Low-molecular-weight NO donors, including sodium nitroprusside (Seabra et al., 2015), nitroglycerin, NO-aspirin (Song et al., 2018), S-nitrosothiols (Seabra et al., 2004), organic nitrates (Riccio and Schoenfisch, 2012), and N-diazemiumdilates (Keefner, 2011) have been developed for the chemical storage and delivery of NO. However, these NO donors have some disadvantages...
as therapeutic tools, such as untargeted delivery, evolving tolerance for some drugs, toxicity, limited NO payload, and uncontrolled NO release (Carpenter and Schoenfisch, 2012; Riccio and Schoenfisch, 2012; Seabra et al., 2015). Thus, previous studies on NO-based therapies have mainly focused on the synthesis of controlled and sustained NO-releasing materials, and these materials have now been widely applied in biomedical fields, including cardiovascular devices, treatment of sexual dysfunction, wound healing, and antibacterial and antitumor treatments (Hetrick et al., 2008; Stevens et al., 2010; Seabra et al., 2004, 2015; Nichols et al., 2012). Silica-based materials have been widely used in the biomedical field, owing to their stability, excellent biocompatibility, straightforward synthesis, and easy customization of the size, morphology, and composition (Riccio and Schoenfisch, 2012). However, to the best of our knowledge, locally delivered exogenous NO stimulation, including that caused by the NO-SNs used in this study, has not been investigated in all animal models of PNI.

Our data showed that fibrin gel containing NO-SNs increased the capillary density at the crushing site of the sciatic nerve at 3 days after injury, and preserved the wet muscle weight and myelinated axons by 3 weeks of injury. We postulated that rapid revascularization stimulated by NO-SNs promoted axonal regeneration, preserved the myelin sheath, and prevented denervation-induced muscle atrophy in the early stages, which resulted in significantly higher SFI from 2 to 4 weeks after the injury in the group treated with NO-SNs. We hypothesize that NO-SNs supply NO, primarily important for restoring or maintaining Schwann cells, influencing their activity, and promoting successful nerve regeneration (Bunge, 1994; Best et al., 1999; Mompeo et al., 2003; Kannan et al., 2005; Sekiguchi et al., 2012). However, there is a discrepancy between the beneficial effects of NO and motor functional recovery. NO-SN releases the majority of NO for up to 30 hours after injury, which is a crucial time period in revascularization and Wallerian degeneration. Successful peripheral axonal regeneration is associated with rapid and efficient Wallerian degeneration. NO-SN promotes revascularization and the rapid recruitment of cells, including macrophages, and mediates Wallerian degeneration regardless of revascularization (Rudic et al., 1998; Giusti et al., 2016; Lee et al., 2016). We speculate that NO-SN treatment creates a favorable environment for regenerating axons for a certain duration, which does not contribute to functional recovery. After a certain time period, axons regenerate to the end-organ (muscle), which might contribute to functional recovery.

Unlike functional improvement in the early phase, there was no difference in the number of myelinated axons and wet muscle weight at 6 weeks after injury, or SFI at 5 and 6 weeks after injury in both groups. We speculate that rapid revascularization stimulated by NO-SNs may have little influence on long-term functional outcomes after nerve damage. Similar observations have been reported by other investigators (Kelihoff et al., 2002b; Giusti et al., 2016; Lee et al., 2016). Vascular endothelial growth factor promotes early revascularization in autografts, but does not improve functional motor recovery in the long term (Giusti et al., 2016; Lee et al., 2016). Although revascularization was delayed after nerve transection in eNOS KO mice, functional recovery of eNOS KO mice was similar to that in wild-type mice (Kelihoff et al., 2002b). However, our data showed that fibrin gel containing NO-SNs improved the TA contraction force at 6 weeks after injury, despite the similar number of myelinated axons and muscle weight at 6 weeks after injury in both groups. This can be explained by the fact that NO is required for the reconnection and maturation of the neuromuscular junction (Sunico et al., 2005). It is possible that the ability of the rats to undergo high and rapid intrinsic nerve regeneration (Kaplan et al., 2012) made the effect of NO less effective. Therefore, further investigations with animal models of low intrinsic recovery levels (e.g., medium/large animal models (Wang et al., 2005) or rat models using allograft (Amniattalab and Mohammadi, 2017), xenograft (Heberbrand et al., 1997), or artificial materials (Zhao et al., 2017)) are necessary to examine the possible effects and benefits of NO-releasing nanomaterials in long-term functional recovery.

The limitation of this study is that the control group should have been treated with fibrin gels containing silica nanoparticles for a stricter comparison, although it is widely accepted that mesoporous silica nanoparticles are biocompatible (Stein et al., 2000; Sayari et al., 2001; Jafari et al., 2019). SFI should have been measured from the same cohort of rats until 6 weeks, i.e. 10 rats per group after excluding 10 sacrificed rats at 3 weeks. In addition, the NO concentration (70 μM) was drawn from previous in vitro results (Jeon et al., 2019) multiplied by an arbitrary factor. However, for precise NO dose control, the diffusion length of NO in the gel (≈100 μm from the source (Wang et al., 2017)) should be quantitatively considered. Recent developments in spatially controlled NO delivery (Park et al., 2020) would enable the quantitative study of the optimized dose of NO in nerve regeneration.

**Conclusion**

Our results suggest that NO-SNs (MAP3/NO) enhance the revascularization of rat sciatic nerves in the early stages after crushing injury. Exogenous nitric oxide delivered by nanomaterials enhanced axonal regeneration and functional recovery in the early phase, as supported by morphometric analysis and sciatic functional assessment. Based on our findings, we suggest a practical method to therapeutically use exogenous nitric oxide. Future studies will focus on determining the optimal dose of NO-SNs using animal models with low intrinsic recovery levels to confirm the benefits of NO-SNs in promoting revascularization for peripheral nerve regeneration.

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**Availability of data and materials**: All data generated or analyzed during this study are included in this published article and its supplementary information files.

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