Nerve growth factor delivery by ultrasound-mediated nanobubble destruction as a treatment for acute spinal cord injury in rats

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Background: Spinal cord injuries (SCIs) can cause severe disability or death. Treatment options include surgical intervention, drug therapy, and stem cell transplantation. However, the efficacy of these methods for functional recovery remains unsatisfactory.

Purpose: This study was conducted to explore the effect of ultrasound (US)-mediated destruction of poly(lactic-co-glycolic acid) (PLGA) nanobubbles (NBs) expressing nerve growth factor (NGF) (NGF/PLGA NBs) on nerve regeneration in rats following SCI.

Materials and methods: Adult male Sprague Dawley rats were randomly divided into four treatment groups after Allen hit models of SCI were established. The groups were normal saline (NS) group, NGF and NBs group, NGF and US group, and NGF/PLGA NBs and US group. Histological changes after SCI were observed by hematoxylin and eosin staining. Neuron viability was determined by Nissl staining. Terminal deoxynucleotidyl transferase-mediated dUTP-biotin nick end labeling staining was used to examine cell apoptosis. NGF gene and protein expressions were detected by quantitative reverse transcription polymerase chain reaction and Western blotting. Green fluorescent protein expression in the spinal cord was examined using an inverted fluorescence microscope. The recovery of neural function was determined using the Basso, Beattie, and Bresnahan test.

Results: NGF therapy using US-mediated NGF/PLGA NBs destruction significantly increased NGF expression, attenuated histological injury, decreased neuron loss, inhibited neuronal apoptosis in injured spinal cords, and increased BBB scores in rats with SCI.

Conclusion: US-mediated NGF/PLGA NBs destruction effectively transfects the NGF gene into target tissues and has a significant effect on the injured spinal cord. The combination of US irradiation and gene therapy through NGF/PLGA NBs holds great promise for the future of nanomedicine and the development of noninvasive treatment options for SCI and other diseases.

Keywords: nanobubbles, nerve growth factor, spinal cord injury, apoptosis, gene therapy

Introduction

Spinal cord injuries (SCIs) can cause severe disability or death. The most common causes of SCI are motor vehicle accidents (38%), falls (22%), violence (13.5%), and sport/recreational accidents (9%).¹ In SCI, structural and functional damage of the spinal cord occurs by primary and secondary injury. This results in loss of movement and sensation and sphincter and autonomic nerve dysfunction below the damaged plane. These effects can damage the patient’s mental health and places an immense burden on society from a public health perspective.²–⁴

The principal treatments for SCI include surgical stabilization and decompression. Physiotherapy and neurotrophic rehabilitation attenuate further damage, relieve spinal
cord ischemia and hypoxia, save the damaged area, rescue necrotic neurons, and promote neuronal function.5-8 In recent years, gene therapy has provided an exciting new direction for research, and substantial progress has been made with improved vectors. Viral vectors have a high transfection efficiency, but their application is restricted because they are cytotoxic with high immunogenicity.9,10 Compared with viral vectors, nonviral vectors such as liposomes and polymeric nanospheres are safe and less toxic. However, their utilization is limited by instability and inefficiency.11-13 Therefore, the search for a safe, efficient, and highly specific transfection system for gene therapy of SCI has become a focus of interest for researchers. Poly(lactic-co-glycolic acid) (PLGA) nanobubbles (NBs) are nonviral vectors that possess unique advantages as gene carriers, such as targeting, slow release, and penetration.14-16 These properties make PLGA NBs ideal candidates for use as drug or gene carriers for the treatment of SCI.

Recently, there has been a rapid development in ultrasound (US)-mediated microbubble destruction (UTMD). UTMD improves the transfection efficiency of exogenous genes noninvasively by sonoporation of microbubbles via US irradiation. Many studies have shown that UTMD is a safe and effective method for transfection.17,18 To date, UTMD has been successfully used for gene transfection in diseases of the heart, lung, brain, uterus, pancreas, kidneys, retina, skin, and testes.19-27 To the best of our knowledge, very few studies have investigated the use of US-mediated gene therapy for SCI. Thus, it is still unknown whether US-mediated gene transfection represents an effective treatment strategy for SCI.

Nerve growth factor (NGF) is one of the best-studied neurotrophic factors (NTFs) and has been shown to promote the growth, differentiation, survival, and synaptogenesis of central and peripheral neurons.28 In this study, we performed US-mediated destruction of NGF/PLGA NBs and investigated its ability to promote the growth, differentiation, and synaptogenesis of neurons for repair of damaged spinal cord tissues in rats after SCI.

Materials and methods
Sample preparation and assay kits
PLGA (50:50, molecular weight [MW] =12,000) and polyvinyl alcohol (PVA, MW =25,000) were purchased from Daigang Biomaterial Co., Ltd. The pCMV6-NGF-GFP plasmid was purchased from OriGene Technologies, Inc. and extracted using the E.Z.N.A. Plasmid Maxi Kit (D6942-01; Qingdao Superri Bio-equip Co., Ltd.). Primary anti-NGF antibody and secondary antibodies were purchased from ABclonal Biotechnology Co., Ltd. Real-time quantitative reverse transcription polymerase chain reaction (qRT-PCR) kits were purchased from Takara Bio Inc. (RR047A). Terminal deoxynucleotidyl transferase-mediated dUTP-biotin nick end labeling (TUNEL) Apoptosis Assay Kit was purchased from Roche (Roche Applied Science).

Preparation of NGF/PLGA NBs
NGF/PLGA NBs were fabricated using a double-emulsion process as reported previously.29 First, 1 mg/mL pCMV6-NGF-GFP plasmid was added to the organic phase of 2 mL CH₂Cl₂ containing 25 mg dissolved PLGA. This mixture was emulsified using an ultrasonic processor at 100 W for 60 s in an ice bath. Subsequently, 10 mL cold PVA solution (1% w/v) was poured into this initial emulsion, and the mixture was emulsified using an ultrasonic processor for 3 min at 75 W in an ice bath. The final solution was stirred for 3 h by a magnetic stirrer and centrifuged at 13,000×g for 5 min. The supernatant was discarded, and the precipitate was washed in deionized water. Centrifugation and washing were repeated three times. Finally, the washed precipitate was lyophilized for 48 h. The dried samples were harvested, filled with perfluorocarbon gas, and stored at −20°C until further use.

The mean diameter, polydispersity index (PDI), and zeta potential of the samples were analyzed by a Laser Particle Size Analyzer System (Zeta SIZER, Malvern). The morphological characterization of the NBs was performed using a light microscope (Olympus CKX41) and a scanning electron microscope (SEM; Hitachi S-340 ON). Transmission electron microscopy (TEM; Hitachi H-7600) was used to estimate the shape of the nanoparticles. The gene encapsulation efficiency was determined using an ultraviolet spectrophotometry method (Thermo NanoDrop 2000; Thermo Fisher Scientific). The following equation was used to calculate the gene encapsulation efficiency:

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\text{Encapsulation efficiency} = \left( \frac{W_1}{W_2} \times 100 \right)\% ,
\]

where \(W_1\) represents the gene amount in NGF/PLGA NBs and \(W_2\) represents the total amount of NGF used to prepare the NGF/PLGA NBs.29,30 Each experiment was performed in triplicate. We compared NGF encapsulation efficiency of five different NGF plasmid concentrations (0.1, 0.2, 0.4, 0.8, and 1 mg). In vitro sonication was performed to assess gene release behavior using a gene transfection instrument (Model UTG 1025; Institution of Ultrasound Imaging, Chongqing Medical University, Chongqing). Sonication parameters were set at 1.5 W for 60 s. The NGF/
PLGA NBs in 10 mL phosphate-buffered saline (PBS) were transferred to dialysis bags (molecular weight cutoff: 10,000 Da) and placed in a reservoir of 100 mL PBS after sonication. At appropriate intervals, 1 mL of the dialysate was extracted and used to determine the gene concentration by ultraviolet spectrophotometry. Consecutively, 1 mL of fresh PBS was added back to the reservoir to maintain a constant volume. The accumulative ratios of the released NGF were calculated.

Experimental animal models and drug injection
This study was conducted in strict accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals, and all the experimental protocols were approved by the animal ethics committee of Chongqing Medical University.

A modified Allen method was used to induce contusion injury. The tenth segment of the thoracic cord was exposed and injured by dropping a 10 g rod from a distance of 5 cm on to the spinal cord and letting the rod rest for 3 min on the lesion site. Adult male Sprague Dawley rats weighing 180–200 g were randomly separated into the following four groups and administered the following treatments: 1. Control group: 0.5 mL normal saline (NS) was injected via the tail vein (n=37). 2. NGF + NBs group: 0.5 mL NGF plasmid (50 µg) and pure PLGA NBs solution were injected via the tail vein (n=37). 3. NGF + US group: 0.5 mL NGF plasmid (50 µg) solution was injected via the tail vein and then subjected to US irradiation for 5 min (n=37). 4. NGF/PLGA NBs + US group: 0.5 mL NGF (50 µg)/PLGA NBs solution was injected via the tail vein and then subjected to US irradiation for 5 min (n=37).

All rats received the corresponding drug injections every 12 h and were then subjected to US irradiation (2.5 W/cm²) for 5 min. This was continued for 3 days, and then specimens were collected. All surgeries were performed under chloral hydrate anesthesia (10%, 0.3 mL/100 g), and all efforts were made to minimize animal suffering.

Following surgery, all rats were housed in separate cages to prevent them from attacking each other. Food and water were provided ad libitum. Intraperitoneal injections of penicillin (2,000 U/100 g) were administered twice a day for 3 days. Urination was stimulated in rats by massaging the bladder twice a day for 7 days. Dressings were kept dry and replaced every 3 days.

qRT-PCR
Total RNA was isolated from the spinal cord tissue using Trizol reagent (ComWin Biotech Co., Ltd., Beijing, China). RNA was reverse transcribed to complementary DNA (cDNA) using the PrimeScript™ RT reagent kit (Perfect Real Time kit) (TAKARA BIO INC). The cDNA samples were prepared for qRT-PCR using a SYBR Premix Ex Taq™ (Perfect Real Time kit) (TAKARA BIO INC). The NGF primer sequences were 5′-GGGGTGATGCTGGTGCTGAGTATG-3′ (sense) and 5′-GGTCTTATCTCACCACCCACA-3′ (antisense), and GAPDH (internal control) primer sequences were 5′-GGGGTGATGCTGGTGCTGAGTATG-3′ (sense) and 5′-CCGCTGCTCCACCACCTTTT-3′ (antisense). qRT-PCR was carried out in 96-well plates using an ABI 7500 Real-Time PCR System and 7500 System Software (Applied Biosystems, Alameda, CA, USA). Each qRT-PCR analysis was performed in triplicate. qRT-PCR data were analyzed by the 2^−ΔΔCT method.

Western blotting
Tissue lysates were prepared from samples frozen at −80°C. Samples were homogenized in RIPA lysis buffer to obtain protein lysates. The lysates were centrifuged at 10,000x g for 10 min at 4°C, and the supernatants were transferred into separate tubes. Protein concentrations were determined using the bicinchoninic acid assay (Beyotime Institute of Biotechnology, China). Equal amounts of protein were separated by electrophoresis on 5% sodium dodecyl sulfate polyacrylamide gels (SDS-PAGE) and then transferred onto polyvinylidene fluoride (PVDF) membranes. The membranes were incubated with a mouse anti-rat NGF monoclonal antibody at 4°C in 5% skim milk overnight. Then, membranes were incubated with a goat anti-mouse secondary immunoglobulin G (IgG) conjugated to horseradish peroxidase. Labeled protein bands were visualized using an electrochemiluminescence (ECL) Western blot substrate kit (Millipore, USA). NGF protein bands were normalized to β-actin, and all immunoblots were quantified with ImageJ software.

Green fluorescent protein (GFP) quantification by inverted fluorescence microscopy
To monitor the location and expression of NGF, a plasmid DNA encoding a GFP reporter gene was introduced using optimal transfection conditions. Spinal cord specimens were cut into frozen sections (10 µm), and GFP was detected by an inverted fluorescence microscope (Leica DMI4000B,
Germany. Semiquantitative analysis of GFP fluorescence intensity was performed using ImageJ software.

Hematoxylin and eosin (HE) and Nissl staining
Spinal cord tissues were immersed in 4% paraformaldehyde for 24 h, transferred to 70% ethanol, dehydrated through a serial alcohol gradient, embedded in paraffin wax blocks, and serially sectioned into 8 µm-thick segments. Tissue sections were dewaxed in xylene, rehydrated through decreasing concentrations of ethanol, and washed in distilled water for HE and Nissl staining. Stained cells were observed using a light microscope (Leica DM500 ICC50).

Detection of apoptotic cells by TUNEL assay
Neuronal apoptosis in the injured region of the spinal cord was analyzed using a TUNEL staining kit (Roche) according to the manufacturer’s instructions. Briefly, the tissue sections were dewaxed in xylene, rehydrated through decreasing concentrations of ethanol, washed in distilled water, treated with proteases, and incubated with the TUNEL reaction mixture. The nuclei were counterstained with 4′,6-diamidino-2-phenylindole (DAPI) (Roche). TUNEL-positive cells had a pyknotic nucleus with green fluorescence, indicative of apoptosis. Sections were imaged using a fluorescence microscope (Leica DMI4000B). The TUNEL reaction was also visualized by chromogenic staining with diaminobenzidine (DAB) (Roche). TUNEL-positive cells were stained brownish-yellow, and positive cells were counted. Sections were imaged using a microscope (Leica DM500 ICC50).

Basso, Beattie, and Bresnahan (BBB) locomotor rating scale
The recovery of neural function was assessed 7, 14, 21, and 28 days after SCI using the BBB locomotor rating scale.33 Rats were placed in an open field (80×130×30 cm) with a pasteboard-covered nonslippery floor. In each testing session, individual rats were observed for 3 min.

Statistical analysis
SPSS18.0 software (IBM Corporation, Armonk, NY, USA) was used for statistical analysis. All data were recorded as mean values ± standard deviation (SD). One-way analysis of variance was used to determine the statistical differences within groups at multiple time points. Student’s t-test was used to analyze differences between two groups at the same time point. A P-value of <0.05 was considered statistically significant.

Results
Characterization of NGF/PLGA NBs
The mean diameter of NGF/PLGA NBs was 215.3±55.29 nm. The PDI was 0.027 (Figure 1A), and the zeta potential was −11.3±5.65 mV (Figure 1B). SEM analysis revealed that NGF/PLGA NBs were highly dispersed and had a well-defined spherical morphology (Figure 1C). The plasmid solution was observed in the core of NBs by TEM (Figure 1D). The encapsulation efficiency of NGF/PLGA NBs was measured by the ultraviolet spectrophotometry method and was highest (34.33%±2.02%) when 0.4 mg of NGF DNA plasmid was used (Figure 1E). The release efficiency of NGF from NGF/PLGA NBs, from the accumulative ratios, was measured by ultraviolet spectrophotometry after low-frequency ultrasonic irradiation. Within 12 h after ultrasonic irradiation, 52.7% of NGF was released from NGF/PLGA NBs and 62.6% of NGF was released after 48 h (Figure 1F).

Verification of the animal model
A modified Allen method was used to induce contusion injury at the T10 segments (Figure 2A and B). Randomly selected rats from each group were analyzed by 7.0 T magnetic resonance imaging (MRI, Biospec 70/20USR; BRUKER, Germany) (Figure 2C and D), and specimens were collected for HE staining 1 day after surgery (Figure 2E). Pathological analysis verified the animal model, and 7.0 T MRI confirmed that SCI was accurately induced.

Total NGF expression in vivo after transfection
NGF protein and mRNA expression were measured 7 and 14 days after SCI. The qRT-PCR data were analyzed by the 2−ΔΔCt method. NGF mRNA expression was significantly higher in the NGF/PLGA NBs + US group compared with other groups at 7 and 14 days after transfection (P<0.05; Figure 3A and B). Furthermore, expression was significantly different (P<0.05) between all pairs of groups, except between NS and NGF + NBs. NGF mRNA expression was highest in the NGF/PLGA NBs + US than all other groups (Figure 3A and B).

NGF protein expression was detected by Western blotting. A single band was observed for each group. NGF protein expression was highest in the NGF/PLGA NBs + US group 7 and 14 days after treatment, compared with the other groups (P<0.05; Figure 3C–F). NGF protein expression was higher in the NGF + US group than the NS and the NGF + NBs groups (P<0.05) in the early stages after treatment. This indicates that US irradiation alone can improve the gene transfection efficiency.
GFP expression

GFP expression was detected by fluorescence microscopy 7 and 14 days after transfection. GFP expression was higher in the NGF/PLGA NBs + US group than other groups after transfection ($P<0.05$; Figure 4A and B). The intensity of green fluorescence was highest in the NGF/PLGA NBs + US group, second highest in the NGF + US group, followed by the NGF group and NS group. GFP expression differed significantly ($P<0.05$) between all pairs of groups, except between the NS group and the NGF group. GFP expression...
Figure 2 MRI image analysis and pathological evaluation indicate the success of the SCI model.

Notes: (A) Exposed rat spinal cord (arrow). (B) The spinal cord was surgically exposed and hit with 10 g rod, leaving obvious contusions (arrow). (C) and (D) MR image of rat spinal cord shows contusion injury at the T10 segment. Sagittal image (C) and coronal image (D) show long T2 signals in the SCI site (arrows). (E) HE staining showing hemorrhage, disordered white matter nerve fibers, and formation of cavities in the SCI site (arrow).

Abbreviations: MRI, magnetic resonance imaging; SCI, spinal cord injury; HE, hematoxylin and eosin.

Figure 3 NGF therapy using US-mediated NGF/PLGA NBs destruction increases NGF expression after in vivo transfection.

Notes: Graphs showing the fold-change increase in NGF expression in the NGF/PLGA NBs + US group by qRT-PCR at 7 days (A) and 14 days (B) after transfection. NGF protein expression was higher after US-mediated NB destruction and NGF therapy at 7 days (C and E) and 14 days (D and F). *P<0.05 compared with other groups. **P<0.05 compared with NGF + US group.

Abbreviations: NS, normal saline; NGF, nerve growth factor; US, ultrasound; PLGA, poly(lactic-co-glycolic acid); NBs, nanobubbles; qRT-PCR, quantitative reverse transfection polymerase chain reaction; ns, not significant.
was highest in the NGF/PLGA NBs + US group at 7 and 14 days after transfection.

**General histopathology and Nissl staining**

Histopathological alterations caused by SCI were examined using HE staining 7 and 14 days after treatment. Hemorrhage was observed in the gray matter, and only a small number of neurons survived. Swollen axons and many vacuoles were visible in the remaining white matter in all groups after SCI. The extent of histopathological changes in the NGF/PLGA NBs + US group was attenuated compared with other groups, at day 7 and day 14. Some glial cell proliferation was detected and many neurons survived and showed normal morphology (Figure 5).

Nissl staining was used to verify neuron viability. Analysis of variance revealed a statistically significant difference in the number of viable neurons among the four groups 7 and 14 days after transfection. Furthermore, significantly fewer Nissl bodies were detected in the NS group and the NGF + NBs group than other groups ($P<0.05$), while significantly more Nissl bodies were observed in the NGF/PLGA NBs + US group compared with other groups ($P<0.01$). In addition, Nissl bodies in the NGF/PLGA NBs + US group were large and plump (Figure 6).

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**Figure 4** NGF therapy using US-mediated NGF/PLGA NBs destruction increased GFP expression.

**Notes:** (A) GFP expression 7 days after SCI ($\times 200$). GFP expression was the strongest in NGF/PLGA NBs + US group, second strongest in NGF + US group, followed by the NGF group, and no expression was found in NS group. (B) GFP expression 14 days after SCI ($\times 200$). GFP expression was highest in the NGF/PLGA NBs + US group.

**Abbreviations:** NGF, nerve growth factor; US, ultrasound; PLGA, poly(lactic-co-glycolic acid); NBs, nanobubbles; GFP, green fluorescent protein; SCI, spinal cord injury; NS, normal saline; ns, not significant.
Neuronal apoptosis was measured in the injured region of spinal cord using a TUNEL assay 14 days after transfection. The TUNEL-DAPI assay showed reduced apoptosis in the NGF/PLGA NBs + US group (Figure 7A). There were many TUNEL-positive cells in the spinal cord of rats in the NS group and NGF+NBs group. The number of TUNEL-positive cells decreased slightly in the US + NGF group. These findings indicate that US-mediated NB destruction combined with NGF treatment significantly reduced the number of apoptotic cells (P<0.01; Figure 7B). This suggests that NGF therapy through US-mediated NB destruction protects neurons effectively by inhibiting apoptosis.

**BBB locomotor rating scale**

Recovery of neural function in the hindlimb was assessed 7, 14, 21, and 28 days after treatment. All rats had normal limb function and a BBB score of 21 before SCI. Immediately after the operation, the BBB score reduced to 0 in all rats, indicating that an SCI model was effectively established. BBB scores were significantly higher following 7 days of NGF/PLGA NBs + US treatment. Recovery continued to improve more in rats treated with NGF/PLGA NBs + US compared with other groups, as shown by higher BBB scores (P<0.05; Figure 8).

**Discussion**

The clinical treatment of SCI poses difficult problems. In SCI, the force of the primary and secondary injury destroys neural tissues, which may lead to permanent loss of function. When the spinal cord is injured, cells and molecules associated with scar formation are activated to form a glial scar, which can limit axon regeneration. Although many promising molecular strategies have emerged to reduce secondary injury and promote axonal regrowth, there is still no effective cure and functional recovery remains limited. In recent years, the use of therapeutic exogenous genes to promote neuronal regeneration has gained much interest, and clinical trials have been initiated. Nevertheless, the safety and continuous expression of these exogenous genes must be addressed before they can be implemented in clinical practice.

NGF is one of the best studied NTFs. Previous studies have shown that NGF protects neurons from damage and promotes regeneration in the central nervous system. The purpose of gene therapy is to rectify abnormal protein expression at the genetic level. Introducing exogenous NGF to correct decreased NGF levels represents a promising treatment option for central nervous system disorders. The mechanism behind the NTF-like activity of NGF is well understood; however, its therapeutic efficacy is not satisfactory. This may be attributed to the blood–brain barrier, which prevents most drugs or genes from entering the brain. Several studies have demonstrated that low-power focused US irradiation following the intravenous administration of an US contrast agent can be used to induce reproducible and consistent focal opening of the blood–brain barrier without any permanent damage to brain tissue. UTMD facilitates the transfer of extracellular molecules
into cells via sonoporation\textsuperscript{42} and is a desirable physical gene transfection method. US can be used to monitor and crush microbubbles in specific tissues at specific times to achieve the accurate targeting required for gene therapy. Most studies have indicated that, under the right conditions, US does not destroy the exogenous gene but enhances its transfection efficiency.\textsuperscript{43} To date, limited research has been conducted into gene therapy of SCI by US-mediated NB destruction.

Shimamura et al\textsuperscript{44} intrathecally injected naked luciferase DNA with Optison through the 4/5th lumbar intervertebral space and applied sonication directly on the thoracic dural sac by removing the dorsal part of the 9–10th thoracic vertebra. This approach enhanced the expression of luciferase in the meningeal cells of the insonated region. Takahashi et al\textsuperscript{17} expressed transgenes in meningeal cells by intrathecal injection of luciferase DNA with Optison through the lower lumbar

**Figure 6** NGF therapy using US-mediated NGF/PLGA NBs destruction decreases neuron loss.

*Notes*: Nissl staining of neurons in different groups at 7 days (A, ×400) and 14 days (B, ×400) after treatment. Neurons in the NS group and the NGF + US group showed lighter staining. More Nissl bodies were detected in the NGF/PLGA NBs + US group, and these Nissl bodies were large and plump. *P*<0.05, compared with other groups. **P**<0.05, compared with the NBs + US group.

*Abbreviations*: NGF, nerve growth factor; US, ultrasound; PLGA, poly(lactic-co-glycolic acid); NBs, nanobubbles; NS, normal saline; ns, not significant.
intervertebral space, together with transcutaneous insonation at the same lumbar region. With this approach, the target genes adhered to the microbubble and were not wrapped or carried in the microbubble. Thus, the stability of the DNA and Optison mixture is relatively poor. In contrast, we expressed NGF in the spinal cord by direct intravascular injection of NGF/PLGA NBs through the tail vein, together with gene transfection treatment at the SCI region. Our intravenous injection method is more useful clinically. Additionally, the gene transfection instrument (Model UTG 1025, Institution of Ultrasound Imaging, Chongqing Medical University) was developed and patented by our institute. Interestingly, transgenes were consistently expressed for as long as 2 weeks in our study, which is longer than previous reports. Furthermore, previous studies only demonstrated the feasibility of US microbubble-mediated plasmid DNA transfer into the target level of the normal spinal cord and did not examine the injured spinal cords or recovery of neural function.

Foreign genes can be transported into target cells by viral and nonviral vectors. Although viral vectors exhibit
a high transfection efficiency, their use is limited by high cytotoxicity and potent immunogenicity. Compared with viral vectors, nonviral vectors such as liposomes and polymeric nanospheres are safe and efficient. PLGA NBs exhibit unique advantages, such as targeting, slow release, and penetrability. These characteristics make PLGA NBs one of the most suitable materials for drug- or gene-based treatment of central nervous system diseases.

In this study, we used PLGA NBs to carry NGF by US into the injured spinal cord to treat SCI. SCI was established in rats using a classical method first reported by Allen. The histological features and the changes in neural function were similar to those seen in previous studies, indicating successful establishment of our SCI model. BBB scores for rats that obtained NGF therapy through US-mediated NB destruction rats were significantly higher at 7, 14, 21, and 28 days post-injury. This anti-apoptotic effect of NGF is consistent with previous studies and may improve recovery following SCI.

Functional recovery is an essential factor for successful SCI treatment. In this study, locomotor function was assessed in rats after SCI. NGF therapy through US-mediated NB destruction improved the BBB scores, but there were some complications, which impaired recovery of neurological function. These included death, urinary tract infection, incision infection, and self-destructive behavior. Ten rats died in the four groups. Incisions became infected in four rats and were cured by debridement and suturing. Six rats had urinary tract infections with pyuria, and two rats eventually died of these infections. Overall, seven rats had to be euthanized because of self-destructive behavior. Additionally, a few rats without complications showed poor functional recovery after NGF/PLGA NBs + US treatment. Poor functional recovery manifested as low BBB scores and a long convalescence and may have been caused by intraoperative nerve injury and denervation after surgery. NGF therapy through US-mediated NB destruction significantly improved the recovery of locomotor function recovery in rats. However, our follow-up was only short term, and long-term observation of neural functional recovery must be performed in the future.

There are several limitations that must be considered when interpreting these findings. All rats underwent laminectomy to expose the spinal cord to permit irradiation by US. However, it is difficult for ultrasonic waves to reach the bone-protected spinal cord in patients who do not receive decompression surgery. The development of a new percutaneous transforaminal endoscopic associated with a focused US system will improve future studies of gene therapy in SCI. Furthermore, low-frequency US waves used in this study may have released NGF into other organs, causing supernumerary biological effects. However, despite these limitations, our findings justify further study into the combination of low-frequency focused US and gene therapy associated with microbubbles for SCI treatment.

Conclusion
We used PLGA NBs to deliver NGF under US irradiation following SCI. This approach improved SCI therapy. Our findings indicate that US-mediated NB destruction facilitates...
gene transfer and may represent a novel gene delivery method for targeted SCI therapy.

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Disclosure

The authors report no conflicts of interest in this work.

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