Comparison of the Effects of BMSC-derived Schwann Cells and Autologous Schwann Cells on Remyelination Using a Rat Sciatic Nerve Defect Model

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Abstract

Schwann cells (SCs) are primarily responsible for the formation of myelin sheaths, yet bone marrow mesenchymal stem cell (BMSC)-derived SCs are often used to replace autologous SCs and assist with the repair of peripheral nerve myelin sheaths. In this study, the effects of the two cell types on remyelination were compared during the repair of peripheral nerves. Methods: An acellular nerve scaffold was prepared using the extraction technique. Rat BMSCs and autologous SCs were extracted. BMSCs were induced to differentiate into BMSC-derived SCs (B-dSCs) in vitro. Seed cells (BMSCs, B-dSCs, and autologous SCs) were cocultured with nerve scaffolds (Sca) in vitro. Rats with severed sciatic nerves were used as the animal model. A composite scaffold was used to bridge the broken ends. After surgery, electrophysiology, cell tracking analyses (EdU labeling), immunofluorescence staining (myelin basic protein (MBP)), toluidine blue staining, and transmission electron microscopy were conducted to compare remyelination between the various groups and to evaluate the effects of the seed cells on myelination. One week after transplantation, only a small number of B-dSCs expressed MBP, which was far less than the proportion of MBP-expressing autologous SCs (P<0.01) but was higher than the proportion of BMSCs expressing MBP (P<0.05). Four weeks after surgery, the electrophysiology results (latency time, conductive velocity and amplitude) and various quantitative indicators of remyelination (thickness, distribution, and the number of myelinated fibers) showed that the Sca+B-dSC group was inferior to the Sca+autologous SC group (P<0.05) but was superior to the Sca+BMSC group (P<0.05). Conclusions: Within 4 weeks after surgery, the use of an acellular nerve scaffold combined with B-dSCs promotes remyelination to a certain extent, but the effect is significantly less than that of the scaffold combined with autologous SCs.

Key words: Schwann cells; remyelination; bone marrow mesenchymal stem cell; nerve regeneration; cell tracking

Introduction

Two prerequisites for functional recovery after peripheral nervous system (PNS) injury have been identified: axonal regeneration and remyelination [1-2]. After PNS injury, Wallerian degeneration occurs in the distal nerve, and the locally proliferating Schwann cells (SCs) subsequently remyelinate and modify the regenerated axon[3]. The myelin sheath protects the axons and promotes the rapid and effective transmission of nerve impulses. Eventual remyelination of the regenerated axon is a necessary step for the effective transmission of nerve signals and functional recovery.
SCs play the most important roles in axon remyelination, and their functions are complex and diverse[4]. SCs have recently been shown to exist in three states[5,6]. In the normal PNS, SCs play a role in forming (myelin states) and maintaining the integrity of the myelin sheath (nonmyelin states, also called Remak cells)[7]. After PNS injury, SCs in the above two states are immediately activated and dedifferentiated into repair Schwann cells[8], which nourish severed axons to promote axon regeneration. These cells act as scavengers to remove the disintegrated myelin debris, and more importantly, they form unique “Büngner bands” to guide the extension of the regenerating axon and establish connections with target sites. When axon regeneration is complete, repair SCs once again shift to a myelinating state, and Remak cells wrap the regenerated axon to complete the remyelination process[9]. For PNS lesions with a longer separation distance (a distance greater than 2 cm)[10], the proximal regenerated axon has difficulty crossing the lesion to reconnect with the distal axon structure because the injured area lacks the support of SCs and a neural basal lamina scaffold. Because the injured area lacks nerve innervation for a long period of time, the distal axon gradually disintegrates[11-13]. Therefore, SC transplantation into an injury lesion is an effective neural repair strategy[14]. However, many limitations in obtaining and expanding autologous SCs in vitro have been identified. Autologous SCs are not suitable for use as ideal seed cells. Recently, many research groups have turned their attention to the stem cell field to conduct in-depth research and have achieved some promising results[15].

Bone marrow mesenchymal stem cells (BMSCs) are widely used in the field of nerve regeneration. BMSCs have great application advantages in neural differentiation, neuroprotection, and neurotroph. In particular, BMSCs have recently been successfully and directionally induced to differentiate into BMSC-derived SCs (B-dSCs)[16,17], which not only express PNS glial lineage markers (i.e., S-100, O4, and glial fibrillary acidic protein (GFAP)) but also wrap the axon in a myelin sheath when cocultured with neurons[18]. Moreover, animal experiments have confirmed that these cells play certain roles in promoting neural regeneration in vivo in place of autologous SCs[19,20]. B-dSCs display a similar efficacy to autologous SCs in many respects[21], and we are interested in the extent to which B-dSCs substitute for autologous SCs in vivo. In this project, acellular nerve scaffolds were used as cell transplantation vectors, and rats with a severed sciatic nerve were used as a model to investigate whether these two seed cell populations exerted the same effects on promoting remyelination.

Materials and Methods

Animals

Animals: Neonatal Sprague-Dawley (SD) rats aged between 3 and 5 days were used for BMSC isolation, and 88 SD rats (either sex, body weights 180-200 g) were used to prepare the animal models and for autologous SC isolation. All experimental animals were housed under standard conditions, and all protocols were approved by the Experimental Animal Administration and Ethics Committee of Sun Yat-Sen University.

Preparation and detection of the scaffold

Scaffold preparation

A 4.5-cm section of the sciatic nerve trunk was removed from an SD rat (used for other experiments) as the raw material for the preparation of the scaffold and was treated as follows: 1. rinsed with ddH2O for 2 h, 2. mechanical shaking in 125 mM sulfobetaine-10 (SB-10) in ddH2O for 12 h, 3. rinsed with PBS 3 times (5 min/rinse), 4. mechanical shaking in 0.14% Triton X-200 and 0.6 mM sulfobetaine-16 (SB-16) for 24 h, and 5. rinsed with PBS 3 times (5 min/rinse). Steps 2-3 were repeated, and then the material was treated with 4% sodium deoxycholate for 24 h. Next, 0.1% (v/v) peracetic acid (Adamas-Beta, Switzerland) diluted in ddH2O was used to sterilize the material for 3 h, followed by 3 washes in sterile PBS (10 min/rinse). The scaffold was stored in sterile PBS at 4°C until subsequent experiments. All procedures described above were performed at room temperature with shaking on a shaker (150 rpm/min). All chemicals were purchased from Sigma (St. Louis, MO, USA), unless indicated otherwise.

Determination of the structure and composition of the scaffold

Nerve grafts subjected to the decellularization treatment were embedded in Tissue-Tek® O.C.T. Compound (Sakura Finetek, USA). Frozen longitudinal and transverse sections were cut at a thickness of 5 µm using a freezing microtome (Leica, USA) and placed on poly-L-lysine-coated slides.

Composition

Immunostaining was performed to analyze the scaffold composition. Sections were incubated with a primary antibody (anti-laminin (LN); 1:1000, Sigma-Aldrich, St. Louis, MO, USA, or anti-fibronectin (FN); 1:400, Abcam, Cambridge, MA, USA) in a humidified chamber at 4°C overnight.
Sections were warmed to room temperature, incubated with a secondary antibody for 1 h in a humidified chamber at 37°C, and washed 3 times with 0.1 M PBS (pH 7.4) on a shaker (100 r/min). 4',6-Diamidino-2-phenylindole (DAPI; Sigma-Aldrich, St. Louis, MO, USA) was used for counterstaining, and the images were visualized using an Eclipse 80i (Nikon, Japan).

**Structure**

The sections prepared as described above were stained with hematoxylin and eosin (H&E) to confirm that the nuclei had been removed. The transmission electron microscopy detection method is described in detail below.

**Preparation of seed cells**

**Extraction and culture of BMSCs**

BMSCs were harvested from the bone marrow of neonatal rat femurs. This procedure is briefly described below. Neonatal rat femurs were isolated under aseptic conditions and rinsed three times with PBS. The bone cavity was repeatedly rinsed in the longitudinal direction with α-Minimum Essential Medium (α-MEM; Gibco, USA) containing 10% fetal bovine serum (FBS; Gibco, USA) and penicillin/streptomycin (50 U/ml and 50 mg/ml, respectively) 15-20 times. The rinse solution was collected, filtered through a 200-mesh sieve, and cultured under standard conditions (37°C). Twelve hours later, the medium was replaced with fresh medium (used above) to remove unattached cells. Then, the medium was replaced once every 3-4 days, and P3 cells were used for flow cytometry-based identification.

**Flow cytometry-based identification of BMSCs**

The concentration of the P3 BMSC suspension was adjusted to 5x10⁶ cells/ml, and the cells were fixed with 4% neutral paraformaldehyde for 30 min. Each EP tube contained 100 μl (approximately 5.0x10⁵ cells) of the suspension. The following antibodies were added to the tubes: CD90-fluorescein isothiocyanate (FITC) as a stem cell marker; CD29-phycoerythrin (PE) and CD44-PE (Serotec Oxford, UK) as mesenchymal lineage markers; and CD11b-PE, CD45-PE, CD106, and CD34-PE (Santa Cruz Biotechnology) as hematopoietic lineage markers. The tube contents were mixed well and incubated at 4°C in the dark for 30 min. Then, a BD FACSCalibur flow cytometer was used to analyze the cells. All antibodies used in the flow cytometry analyses were purchased from BioLegend, unless indicated otherwise.

**Directionally induced differentiation of BMSCs into B-dSCs**

The induction method [18] is briefly described below. P3 generation BMSCs were collected, and the cell concentration was adjusted to 1x10⁶ cells/ml with mixed induction medium containing 1:1 (v/v) Dulbecco’s Modified Eagle’s Medium (DMEM)/F12 and Neurobasal medium (Invitrogen) supplemented with basic fibroblast growth factor (bFGF; 40 ng/ml; Peprotech), epidermal growth factor (20 ng/ml) (Peprotech), and B27 (2%, v/v, Invitrogen). Then, the cells were seeded onto ultralow attachment culture plates (Corning). Three weeks later, neurospheres were collected and seeded into poly-lysine-coated flasks. After the cells had attached to the wall, the medium was changed to a mixed medium composed of α-MEM (Gibco) supplemented with 10% FBS, forskolin (5 mM), platelet-derived growth factor (PDGF)-AA (5 ng/ml), bFGF (10 ng/ml), and β-hergululin (200 ng/ml). Cells were cultured for two additional weeks before the detection of the S-100, O4, and GFAP proteins. These B-dSCs were further cultured in basal medium (DMEM/F12 supplemented with 10% FBS).

**Isolation and identification of autologous Schwann cells**

One-half centimeter of the unilateral rat radial nerve was transected (the animal was sutured for the subsequent experiments) and digested with 0.05% collagenase. The digested mixture was repeatedly pipetted to collect the cell suspension, which was cultured in DMEM-F12 containing 10% fetal bovine serum (in a 37°C humidified incubator with 10% CO2). The mitotic inhibitors cytosine arabinoside (Ara-c) and G-418 were used to remove fibroblasts, and bFGF was used to promote the growth of Schwann cells. After the cells were confluent, the rapid digestion method with a low concentration of trypsin was used to collect the cells. The P3 generation of the cells was stained with the Schwann cell-specific antibody S-100 (1:800, Sigma, St. Louis, MO, USA), O4 (1:1000, Santa Cruz Biotechnology, USA), and GFAP (1:400, Sigma, St. Louis, MO, USA).

**Cell labeling**

EdU (5-ethynyl-2'-deoxyuridine) is a DNA thymine base pair analog that is incorporated into DNA when the cells undergo mitosis[22]. Therefore, EdU has been used to evaluate cell proliferation or to replace the traditional BrdU labeling method for cell tracking. The labeling method is described below. A total of 1x10⁶ cells were collected and seeded into a culture flask (25 cm²). After all cells had adhered to the flask wall, the Click-iTTM EdU (Molecular Probes;
Invitrogen) solvent was diluted with the culture medium to a working concentration of 20 μM, added to the culture flask, and incubated for 24 h. The medium was replaced with fresh medium. When the cells were traced, 4% paraformaldehyde was used to fix the specimen for 15 min; then, the cells were stained overnight with the detection medium (Alexa Fluor 594), and the cells were observed under a fluorescence microscope (Eclipse 80i, Nikon). The results were analyzed using Image-Pro Plus 6.0 software (Media Cybernetics, Inc., USA).

**Implantation and complex formation**

The cell complex formation method is briefly described below. The acellular nerve scaffold (approximately 2-cm long) prepared as described above was collected. A microsyringe was used to withdraw a 100-μl cell suspension (approximately 1x10⁶ cells), which was slowly injected into the scaffold. The surface of the scaffold was rinsed 3 times with PBS (3 min/rinse). Then, the nerve scaffold complexed with cells was placed into culture medium (DMEM/F12 supplemented with 10% FBS) and cultured for 7 days. Frozen sections were prepared for immunostaining (S-100), scanning electron microscopy (SEM), or animal experiments. For the SEM analysis, the specimen was fixed with glutaraldehyde for 2 h and postfixed with a 4% osmic acid solution for 2 h. After dehydration in a gradient of alcohol solutions, the specimens were soaked in isoamyl acetate overnight. Specimens were embedded in Poly/Bed 812 resin (Ted Pella, CA, USA) at 60°C for 48 h and then sectioned according to the different groups described in Table 1 according to the administration of different treatments. Sodium pentobarbital (30 mg/ml, 0.4 ml/100 g) was injected intraperitoneally, and the right sciatic nerve was exposed. The trunk of the sciatic nerve was flipped 180° in situ and sutured using the same method. In the sham group, the skin was cut, muscles were isolated, and the sciatic nerve was exposed but not severed. The animals were housed in a standard environment. All surgical procedures minimized the number of rats used and their suffering.

| Treatments                        | Groups (n=10 animals per group; unless indicated otherwise) |
|-----------------------------------|------------------------------------------------------------|
| Scaffold+B-dSCs                   | Sca+B-dSC group                                           |
| Scaffold+BMSCs                    | Sca+BMSC group                                            |
| Scaffold+autologous SCs           | Sca+autologous SC group                                   |
| B-dSC suspension                  | B-dSC group                                               |
| BMSC suspension                   | BMSC group                                                |
| autologous SC suspension          | Autologous SC group                                        |
| Scaffold                            | Sca group                                                 |
| Autologous nerve                  | Control group                                             |
| Just exposed the sciatic nerve    | Sham group (n=4 animals)                                   |
| Saline                             | Untreated group (n=4 animals)                              |

**Table 1. Heading**

**Surgical methods**

Eighty-eight SD rats were randomly divided into the 9 groups described in Table 1 according to the administration of different treatments. Sodium pentobarbital (30 mg/ml, 0.4 ml/100 g) was injected intraperitoneally, and the right sciatic nerve was exposed. The trunk of the sciatic nerve was completely severed at 0.5 cm proximal to the branch of the tibial nerve and peroneal nerve to cause an approximately 1.5-cm-long nerve defect. For the Sca+B-dSC, Sca+BMSC, Sca+autologous SC and Sca groups, the 1.5-cm nerve scaffold complexed with cells was used to bridge the two ends of the defect region using the adventitia suture method. For the B-dSC, BMSC and autologous SC groups, a 100-μl cell suspension (approximately 1x10⁶ cells) was injected into the lesion site. For the untreated injured group, 100 μl of saline was used as a control. For the autologous nerve group, the severed 1.5-cm long sciatic nerve was flipped 180° in situ and sutured using the same method. In the sham group, the skin was cut, muscles were isolated, and the sciatic nerve was exposed but not severed. The animals were housed in a standard environment. All surgical procedures minimized the number of rats used and their suffering.

**Nerve function test**

Four weeks after surgery, all groups of animals were subjected to a sciatic nerve electrophysiological examination using the RM6240 multichannel electrophysiological signal acquisition and analysis system. Briefly, the rats were anesthetized via an intraperitoneal injection of phenobarbital sodium (45 mg/ml). The surgical field was reopened to expose and dissect the repaired sciatic nerve. The examination of the function of the motor nerves was conducted using the procedure described below. The recording electrode was placed in the triceps surae, and the stimulating electrode was placed 0.5 cm from the proximal end of the sciatic nerve graft. The stimulation current, time, and frequency parameters were 1-2 mA, 0.1 ms, and 1.0 Hz, respectively. The amplitude (AP), latency (LT), and conduction velocity (CV) of the nerve impulse potential were recorded.

**Evaluation of remyelination**

At the 4th week postoperation, the transplanted grafts and the corresponding distal nerves located 2 cm from the graft were isolated from all animals to be tested. The surface fat and basal lamina tissue were removed to perform the experiments listed below.

**TEM and toluidine blue staining**

Specimens were washed with PBS three times (5 min/wash) and fixed with a 2.5% (m/v) glutaraldehyde solution for 2 h, followed by 1% osmic acid for 90 min. After dehydration in a gradient of alcohol solutions, the specimens were placed in 1% uranyl acetate overnight. Specimens were embedded in Poly/Bed 812 resin (Ted Pella, CA, USA) at 60°C for 48 h and then sectioned according to the different
experimental designs (i.e., 0.5-μm-thick tissue sections were stained with toluidine blue, and the myelin sheath was observed under a light microscope (Eclipse 80i, Nikon). For TEM, ultrathin sections (80-100 nm) were prepared and stained with lead citrate and uranyl acetate. Sections were observed under a transmission electron microscope (FEI TECNAI SPIRIT G²). Previous research methods were used as references for the analysis of relevant data. Three tissue sections per group were randomly imaged using a Nikon Eclipse 80i microscope equipped with a high-resolution color digital camera (Digital Sight US-U2, Nikon). The results were analyzed using the Image-Pro Plus software (Media Cybernetics, Inc.). The following methods were used to quantify the results of the cell tracing analysis: 10 high-power fields were randomly selected, and the software was used to calculate the numbers of DAPI+ cells, EdU+ cells, and EdU+MBP+ cells. The following method was used to perform a semiquantitative analysis of the levels of the NF-200 and S100 proteins: ten high-power fields were randomly selected, and the expression levels were calculated by dividing the positively stained area for each protein by the area of the total field of view.

Statistical analysis

All experiments were repeated at least three times, unless indicated otherwise. Data are presented as means±SD and were analyzed using the Statistical Program for Social Science (SPSS) software version 19.0. Statistical significance was estimated using one-way analysis of variance (ANOVA) with the Tukey-Kramer post hoc analysis. P values<0.05 were considered statistically significant.

Results

Analysis of the scaffold properties

In terms of structure, after acellular extraction, the nerve tissue structure was basically complete, the nuclei had been completely removed (Figure 1A), the basal lamina scaffold wall was complete, and no significant collapse of the lumen was observed (Figure 1C). The tube wall was approximately 0.5-1-μm-thick, and the tube diameter was generally 4-10 μm. The SCs formed a laminar myelin sheath structure, and the axon structure in the basal lamina scaffold had completely disappeared (Figure 1D). The interior of the scaffold showed a wavy mesh-like structure; these tubular structures guide and limit the direction of axon extension.

In terms of composition, FN and LN immunostaining (Figure 1B-C) showed that the scaffold itself contained abundant FN and LN. These protein components play important roles in seed cell adhesion and nerve regeneration.

Cell culture and identification

Culture and identification of bone marrow mesenchymal cells

After the primary cells were isolated and purified, P3 cells were collected. After 24 h of adherent culture, all cells showed a spindle shape under the light microscope. After 5 additional days of culture, the cells showed whirlpool-like morphology (Figure 2A). Similar to previous studies[23], flow cytometry revealed that greater than 98% of the cells expressed the mesenchymal lineage markers CD29, CD44, and CD90, and no more than 1% of the cells expressed the hematopoietic lineage markers CD34, CD11b, CD45, and CD106 (Figure 2).

Induction, differentiation, and identification of BMSCs

The induced and differentiated B-dSCs showed an SC-like morphology and extended processes within 5-7 days (Figure 3B). According to the immunofluorescence staining, 91.3±5.4% (n=6) of cells were positive for S-100, O4, and GFAP during this period (Figure 3D-F).

Figure 1. Basic properties of a nerve scaffold. H&E staining shows that the nuclei in the scaffold have been completely removed. The composition test showed that the scaffold was enriched in LN (cross-section) and FN (longitudinal section). TEM image showing that the myelin sheath and axon components disappeared and that the neural basal lamina scaffold remained intact, with no significant collapse of the scaffold wall. Scale bars=100 μm in A, B and C.
Figure 2. BMSC identification. Cultured P3 BMSCs express high levels of CD90, CD44, and CD29 and low levels of CD34, CD45, CD106 and CD11b.

Figure 3. Detection of the properties of B-dSCs. Compared with normal BMSCs (A), the image in B shows induced B-dSCs with a disordered cell arrangement. The cell body is spindle-shaped with slender processes and has a morphology similar to SCs (C). Most of the B-dSCs expressed the SC cell-specific markers S-100 (D), O4 (E), and GFAP (F). Scale bar=50 µm.
Figure 4. Composite scaffold. The cells and the scaffold were cocultured in vitro for 48 h. A-C SEM images showing that all seed cells (A for B-dSCs; B for BMSCs; and C for SCs) adhere to and survive on the scaffold surface. After 7 days of coculture, immunostaining (D and E) shows stable expression of S-100 of B-dSCs and SCs cultured on the scaffold. The nuclear counterstain shows that these two cell types show a “Büngner band”-like linear arrangement on the scaffold. Scale bar=50 µm.

Figure 5. Electrophysiological test results. Comparison of the electrophysiological test results between groups of animals that received the scaffold transplant at four weeks after surgery. *P<0.05, **P<0.01, one-way ANOVA with Tukey’s post hoc test.

SC culture

The isolated and cultured autologous SCs exhibited a typical spindle-shape structure (Figure 3C).

Cell labeling

After coculture with EdU for 24 h, the numbers of EdU+ cells among the three cell types (BMSCs, autologous SCs, and B-dSCs) were 94.5%, 93.9%, and 97.6%, respectively (Figure 6A-C).

Cell complex formation

After the seed cells were complexed with the nerve scaffold for 7 days, SEM images of the three groups of specimens showed that most of the cells aggregated and adhered to the surface of the internal pore walls of the scaffold (Figure 4A-C). Based on the immunostaining results, cells in the B-dSC and the autologous SC groups continued to express S-100 (Figure 4E-F). Thus, the acellular scaffold had better cell compatibility and represented a potentially suitable cell delivery system for the three types of cells to repair the sciatic nerve defect. Notably, these two groups of cells showed a linear arrangement under the guidance of the physical scaffold structure (see the red dotted lines in Figure 4E-F).

Restoration of nerve function

With the exception of the sham group, the posterior limbs of all animals in other groups became completely paralyzed after the nerve was severed, and motor function was completely lost. During the
4-week observation period after surgery, all animals developed ulcers or lost toes, whereas these phenomena did not occur in the control group and the sham group. Instead, only the loss of a toenail and foot pad tissue atrophy were observed in the control group.

**Electrophysiological test results**

The myelin sheath plays a role in insulation and improving the nerve conduction velocity. In the electrophysiological tests, CV and CMAP are effective parameters that reflect the function of the myelin sheath. The results from the electrophysiological tests of the groups at four weeks after the surgery (n=3) are provided below, except for the groups treated with cells alone (including BMSC, B-dSC, and autologous SC groups) and the untreated group, from which we were not able to record electrical signals because nerve regeneration did not occur in the lesion site. And atrophy of the distal nerve was observed. Rats in the Sca+B-dSC, Sca+BMSC, Sca+autologous SC and Sca groups all displayed worse recoveries than rats in the control group (CV: 115±15 m/s, LT: 3.3±0.4 ms, P<0.05; AP: 2.91±0.85 mV, P<0.05). However, the rats in the Sca+B-dSC group (CV: 28±10 m/s; LT: 7.2±1.3 ms; AP: 0.69±0.22 mV) displayed an even lower CV and CMAP than the rats in the Sca+autologous SC group (CV: 59±6 m/s; LT: 5.1±0.8 ms, P<0.05; AP: 1.51±0.47 mV, P<0.01), but greater values than the rats in the Sca+BMSC group (CV: 16±3 m/s, P<0.05; LT: 8.5±0.7 ms; AP: 0.98±0.21 mV, P<0.01) and Sca group (CV: 9±5 m/s, P<0.05; LT: 10.3±0.5 ms; AP: 0.63±0.22 mV, P<0.01).

**Cell tracing and myelin-associated protein detection**

On the 7th postoperative day (n=4), none of the transplanted cells in groups treated with cells alone (BMSC, B-dSC, and autologous SC groups) had survived (data not shown). Only 7% of the transplanted cells survived in the Sca+autologous SC group, which was less than the survival rate obtained in the Sca+B-dSC (12%) and Sca+BMSC groups (14%). However, most (85%) of these cells (Edu+) expressed MBP, suggesting that most of these cells were capable of forming a myelin sheath in vivo. For the Sca+B-dSC and Sca+BMSC groups, MBP+Edu+ double-positive

![Figure 6](http://www.ijbs.com)
cells accounted for only 15% and 2% of the total cells, respectively, values that were significantly lower than the percentage obtained for the Sca+autologous SC group (P<0.01) (Figure 6D-F and M). On the 14th postoperative day, the EdU signal was not detected in these three groups of animals that received transplanted cells (data not shown). However, the quantitative results from the S-100 and NF-200 immunofluorescence staining indicated higher levels of expression of these two proteins in the Sca+autologous SC group than in the Sca+B-dSC and Sca+BMSC groups (Figure 6J-L and N), indicating better regeneration of axons and the myelin sheath in the Sca+autologous SC group than in the other two groups at this time point (P<0.05).

Remyelination

The proportion of myelinated nerves and the depth and depth distribution of the myelin sheaths are powerful indicators of remyelination. During the 4th postoperative week (n=4), lesion sites were re-exposed, and groups transplanted with cells alone (BMSC, B-dSC, and autologous SC groups) and the untreated group did not display obvious nerve regeneration in the defect area, defect areas exceeding 1.5 cm in length were observed between the stumps of the nerve, and the distal nerve tissue exhibited obvious shrinkage. Therefore, the remyelination evaluation was mainly aimed at the groups in which the scaffold was transplanted (Sca+B-dSC, Sca+BMSC, Sca+autologous SC, and Sca groups), as well as the control group.

The toluidine blue and TEM data (Figure 7) indicated that 45% of the nerve fibers in the grafts of the Sca+B-dSC group were myelinated, a value that was significantly lower than the proportion in the Sca+autologous SC group (70%; P<0.01), did not differ from the proportion of the Sca+BMSC group (43%) (P>0.05) and was slightly higher than the proportion of the Sca group (31%) (P>0.05). However, the values obtained for these four groups were less than the values obtained for the control group (P>0.05). The average thickness of the myelin sheath in the Sca+autologous SC group was 0.73±0.19 μm, which was significantly higher than the thickness of 0.42±0.05 μm observed in the Sca+B-dSC group, 0.19±0.07 μm in the Sca+BMSC group and 0.21±0.02 μm in the Sca group. The G-ratio was 0.64±0.04 in the control group but only 0.88±0.06 in the Sca+B-dSC group, which was higher than the 0.73±0.09 ratio (P<0.05) obtained for the Sca+autologous SC group. No significant difference was detected compared to the 0.91±0.03 ratio (P>0.05) obtained for the Sca+BMSC group and 0.87±0.09 in the Sca group (P>0.05). Based on the analysis of the thickness of the myelin sheath, the distribution of myelin sheaths with a thickness greater than 0.6 μm was as follows: graft 74% and distal nerve 20% in the Sca+autologous SC group; graft 20% and distal nerve 5% in the Sca+B-dSC group; graft 10% and distal nerve 2% in the Sca+BMSC group; graft 71% and distal nerve 29% in the Sca group and graft 80% and distal nerve 73% in the control group. The toluidine blue and TEM results showed that the numbers of axons in the central region of the graft and in the distal nerves in the animals of the Sca+B-dSC, Sca+autologous SC, and Sca+BMSC groups were 487±89 and 276±66, 560±41 and 327±37, and 521±33 and 232±59, respectively. No significant differences were observed among the three groups. However, the values of the Sca+BMSC groups were significantly higher than the Sca group (256±19 and 188±35, respectively; P<0.05).

Discussion

A nerve scaffold is a necessary prerequisite for the repair of defective nerves. If no bridging scaffold is available, the ability of regenerated axons to cross the injured area is hindered for long-distance nerve defects[24]. The nerve scaffold provides a good microenvironment for nerve regeneration, in which SCs adhere to and migrate along the scaffold surface [25]. The spatial structure of the scaffold provides natural physical guidance for the SCs, enabling them to eventually form a unique cell alignment structure (Büngner band), which is the most important condition for inducing axonal extension[26]. After the cells were seeded on the surface of the scaffold and cocultured for 7 days, the cells (B-dSCs and autologous SCs in our experiment) showed an ordered cell alignment under the guidance of the natural scaffold structure (see Figure 4D-E). We postulate that this cell structure plays an important role in the subsequent regeneration of nerve axons. A simple acellular nerve scaffold has been reported to promote a certain degree of myelin regeneration (see our data in the results section), but this approach still has many problems compared with SCs embedded in a scaffold [27]. Therefore, supplementation with exogenous SCs is necessary. The scaffold itself provides only a substrate for the adhesion and migration of cells, and when the number of SCs is insufficient to assist in the regeneration of the nerve axons, the efficacy of nerve scaffold is reduced. Simple transplantation of cells may promote regeneration under conditions in which the continuity of nerve is intact; however, for long-distance nerve defects, locally injected cells did not increase regeneration in the absence of a scaffold (on the 7th day after the operation, locally injected cells were unable to survive without a scaffold). As expected, in this experiment, the isolated cell transplantation groups and the
untreated injury group did not reconstruct the continuous nerve by the 4th week after surgery. Therefore, the role of the scaffold is very important.

We postulate that scaffold implantation is a prerequisite for the evaluation of exogenous cellular function in long-distance nerve defect models.
The regeneration and extension of axons are inseparable from the effects of SCs[28-31]. When the nerves are severed, the injured area lacks a sufficient number of SCs to support axon regeneration. Strategies that rely solely on the entry of proliferating SCs at the proximal stump into the scaffold and the subsequent induction of nerve regeneration is obviously unrealistic. Thus, exogenous SCs are an important tool to assist in nerve regeneration[4]. BMSCs have the ability to differentiate into ectodermal neuronal cells[32]. Monolayers of B-dSCs have recently been shown express their specific markers (S-100, O4, and GFAP)[33], and differentiated cells have the ability to perform the basic functions of SCs, such as participating in the ATP-induced intracellular Ca2+ signaling pathway (ATP-purinergic signaling) [34], the secretion of multiple neurotrophic factors [34], inducing the extension of neuronal axons, and wrapping new axons with myelin sheaths[18]. These characteristics confirm that B-dSCs are more effective than BMSCs at promoting nerve regeneration and are ideal substitutes for SCs[36,37]. In this experiment, we used the scaffold to simulate the 3-D environment in vivo and found that both B-dSCs and SCs continued to express the S-100 protein when the seed cells were complexed with the scaffold for 7 days. Based on this finding, the physical environment created by the scaffold does not change the basic characteristics of B-dSCs. Thus, the scaffold can be used as a vehicle for in vivo cell delivery. However, the role of SCs is extremely complex in the in vivo environment. Ideal seed cells should perform the same functions as genuine SCs. What are the actual functions of B-dSCs in vivo?

We conducted in vivo experiments to answer these questions. In the present study, animals in the autologous nerve transplantation group showed only mild foot pad and toenail atrophy during the 4-week observation period. During the first four weeks after the operation, all animals in the other groups (except for the sham group) showed different proportions of ulcers or toe loss. Neurotrophic substances are synthesized by the soma of neurons and are transported through axoplasm flow to the nerve endings for release[38]. This process does not affect the conduction of nerve impulses. Even if the conduction of nerve impulses is abnormal (e.g., due to the myelin sheath), the axoplasm flow will not stop, and the tissue under its innervation will not undergo metabolic changes and atrophy[39]. Therefore, we preliminarily suggested that the nerve repair in the Sca+B-dSC, Sca+autologous SC, and Sca+BMSC groups did not reach the same efficacy as autologous nerve transplantation. We subsequently performed electrophysiological examinations to further explore the evidence of remyelination and repair in the Sca+B-dSC, Sca+autologous SC, and Sca+BMSC groups. In the electrophysiological analysis, the CV, LT, and AMP were significantly different in the Sca+B-dSC or Sca+BMSC group compared with the Sca+autologous SC group. SCs are the cell type responsible for the formation of myelin, and the quality of the resulting myelin sheath (i.e., the number of myelinated nerve fibers, the thickness of the myelin sheath, and the distance between the nodes of Ranvier) is a critically important factor that affects the electrophysiological outcome[40,41]. Therefore, we speculate that B-dSCs may not be able to perform all functions of autologous SCs, particularly the formation of the myelin sheath, and thus, a difference may exist between the two cell types.

We analyzed the differences in the promotion of myelin sheath formation by the seed cells in each group using cell tracing and morphology to verify this hypothesis. The thickness of the myelin sheath, the proportion of myelinated fibers, and the myelin area ratio were reduced in the Sca+B-dSC group compared with the Sca+autologous SC group. In particular, regarding the distribution of the myelin sheath, approximately 70% of myelin sheaths had a thickness >0.6 μm in the Sca+autologous SC group, whereas this value was less than 25% in the Sca+B-dSC group. We investigated whether these results were caused by differences in the survival rates of the seed cells. We tried to analyze the results using the cell tracing technique to exclude this possibility. On the 7th postoperative day, the tracing results showed that more transplanted BMSCs and B-dSCs survived than autologous SCs. However, based on the ratio of MBP+EdU+ cells, a significantly greater proportion of cells formed myelin sheaths in the Sca+autologous SC group than in the Sca+B-dSC group. These data most likely confirmed that a large difference existed in the abilities of B-dSCs and autologous SCs to form the myelin sheath. Most B-dSCs do not function like SCs in vivo, as expected. We speculated that a possible explanation was that some of the B-dSCs failed to be converted to the myelinating states that effectively wrap the axon and form the myelin sheath, thus increasing the number of unmyelinated fibers[42]. The nonmyelinating states are important conditions for maintaining the integrity of myelin[43]. According to the electron microscopy results, the myelin sheath structures formed by the B-dSCs, autologous SCs, and BMSCs were intact. We postulate that some B-dSCs are converted to nonmyelinating states in vivo, but in-depth in vivo studies are needed to clarify this speculation. The survival and maintenance of related Schwann cell properties in the repair state are the keys to successful nerve regeneration[5]. The so-called...
Büngner band structure is actually formed by Schwann cells in the repair state[44]. Both the Sca+B-dSC and the Sca+autologous SC groups formed cell alignments under the guidance of the physical structure of the scaffolds and formed a Büngner band-like structure to induce axon regeneration and extension[6]. We hypothesized that the seed cells of the Sca+B-dSC and Sca+autologous SC groups served to stabilize the Schwann cells in the repair state. The use of acellular nerve allografts (ANGs) combined with B-dSCs or BMSCs results in less axon regeneration than ANGs containing SCs[34,45]. In the present study, the analysis of the axon regeneration revealed lower levels of NF-200 and S-100 immunostaining in the Sca+B-dSC group during the second postoperative w than in the Sca+autologous SC group, although the expression levels were higher than those in the Sca+BMSC group. However, 4 weeks after the operation, the analysis of the number of regenerated axons showed significant differences among the Sca+B-dSC, Sca+autologous SC, and Sca+BMSC groups. Based on this result, we speculated that autologous SCs formed a Büngner band on the scaffold in the early stage after the operation and were induced axon extension to a greater extent than the other two cell types. Regarding the results obtained in the later stage, one possible explanation is that B-dSCs or BMSCs may promote nerve regeneration through other pathways[46-48], such as the secretion of certain neurotrophic factors, local regulation of immunity, or the induction of angiogenesis, and the comprehensive effects of these alternative pathways may partially compensate for some of the defects of B-dSCs or BMSCs in promoting remyelination. Based on the above analysis, we postulate that B-dSCs may play a certain role in the myelination of the severed nerves but do not completely replace autologous SCs. However, the relevant mechanisms require further exploration.

Conclusions

B-dSCs are a type of commonly used seed cell for nerve repair. When combined with an acellular nerve scaffold, they promote nerve regeneration to a certain extent. However, significant differences were observed compared with autologous SCs in vivo. In the first 4 postoperative weeks, B-dSCs do not achieve the same effect on myelination as autologous SCs but are better than BMSCs. The underlying mechanism requires further exploration.

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Competing Interests

The authors have declared that no competing interest exists.

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