The Effect of Chick Embryo Cerebro-Spinal Fluid in Microwave Irradiated Collagen Guide Channel on Sciatic Nerve Regeneration in Rat

Abstract

The aim of this research was to evaluate the effect of chick embryo cerebrospinal fluid (e-CSF) across the irradiated collagen guide channel on sciatic nerve regeneration in comparison with autograft. Forty adult male Sprague-Dawley rats (250-300g) were randomized into (1) collagen channel + E-CSF, (2) collagen channel + normal saline (NS), (3) autograft (AG), and (4) sham surgery groups. The left sciatic nerve was exposed and a 10 mm nerve segment was cut and removed. In the collagen groups, the proximal and distal cuts ends of sciatic nerve were telescoped into the nerve guides and E-CSF or NS injected into collagen conduit. To the autograft group, the 10 mm nerve segment was turned backwards and used an autologous nerve graft. All animals were evaluated by sciatic functional index (SFI) and electrophysiology testing. The improvements in SFI since the beginning of the last evaluation in experimental groups were measured. On days 49 and 60 post-operation, the mean SFI of the collagen + E-CSF group was significantly greater than the autograft group (P< 0.05). On day 90, the mean nerve conduction velocity (NCV) of the collagen + CSF group was greater than autograft group (P< 0.05). These findings showed that RPS in collagen nerve guide channel effectively enhances nerve regeneration and promotes functional recovery in injured sciatic nerve of rats.

Keywords: Embryo cerebro-spinal fluid; Collagen conduit; Nerve regeneration

Introduction

Peripheral nerve injuries lead to significant loss of neuron function, and the first choice for treating severe nerve defects involves implantation of autografts to bridge the defect in the injured nerve site. However that is limited by graft availability, secondary deformities, potential differences in tissue structure and size [1].

Novel approaches attempt to provide an alternative to autologous nerve graft. Nerve guidance conduit (NGC) have been designed to bridge the gap and provides guidance for sprouting axon and provide a place for diffusion of neurotrophic factors and endogenous/transplanted cells at the lesion site [2]. It is believed that an ideal nerve conduit should provide not only structural support for damaged nerve but also the neurotropic and neurotrophic support for axonal regrowth [3]. Collagen biomaterials were found to be highly biocompatible, absorbable and able to promote Schwann cell adhesion, proliferation and migration [4]. Collagen nerve conduits have been shown to be effective in rat and monkey models of nerve repair and are FDA and EU-approved for clinical application [5]. Lohmeyer et al. (2007) used collagen conduits to repair sensory nerve avulsions in 11 patients and observed good improvement in nerve recovery. Numerous experiments indicated that, after nerve injury, cell adhesion molecules and neurotrophic factor played very important roles in nerve regeneration, cellular viability, growth, activity and functional recovery [6]. Among the neurotrophic factors, nerve growth factor (NGF) plays a critical role. Supplementing the nerve site with NGF has been shown to enhance proliferation and promote nerve repair following injury [7].

A number of investigators reported Embryonic CSF (E-CSF) contains high concentrations of NGF, transforming growth factor-α (TGFα), transforming growth factor-β (TGF-β), epidermal growth factor (EGF), and neurotrophin-3 (NT-3) [8]. Early experiments on rat embryos have shown E-CSF supports viability and proliferation of cortical cells in vitro [9]. Gato et al. [10] were cultured mesencephalic neuroectoderm for 24 hr with defined culture medium supplemented with chick E-CSF, show that E-CSF promotes neuroepithelial stem cell survival, and induces proliferation and neurogenesis in mesencephalic explants. Martin et al. [11] identified several isoform of FGF-2 in CSF during early stages of development in chick embryos and demonstrated that E-CSF has an effect on the regulation of neuroepithelial cell behavior, including cell proliferation and neurogenesis [11]. In the present study, our purpose was to evaluate the effect of collagen tube containing chick embryo cerebrospinal fluid on repair of a 10 mm sciatic nerve defect in rats.
Material and Methods

Animals

Forty adult male Sprague-Dawley rats (250-300g) were randomized into:

(i) Collagen channel + E-CSF
(ii) Collagen channel + normal saline (NS)
(iii) Autograft (AG)
(iv) Sham surgery groups.

The Tabriz University of Medical Sciences Ethical Committee approved all experiments.

Preparation of chick embryo cerebro-spinal fluid

Fertile chick white Leghorn eggs were incubated at 38±1°C in a humidified atmosphere. CSF carefully aspirated under a microscope from incubated chick embryos in day E17 by using a 20 µL pulled tip glass microcapillary pipette, which was held steadily inside the brain ventricles. The maximum amount of CSF collected from each embryo was 15 µL. E-CSF was collected from 30 chick embryos according to development stage based on Hamburger and Hamilton (1951). CSF samples were kept at 4 °C during collection to minimize protein degradation. The aliquots were centrifuged at 15,000 rpm at 4°C for 10 minutes to remove any contaminating cells [8]. None of the samples used in this experiment had any visible sign of contaminating red blood cells.

Preparation of collagen tube

Type-I Collagens were extracted from rat-tail tendons by modification of the previously published method [12]. In brief, rat-tail teased out from 3-month-old rats and washed in ethanol 70 °C for 45 min. Tendons were excised from rat-tails, washed twice with PBS, and then soaked in distilled water and 4 mmol/L acetic acid. After stir for 7 d at 4 °C. The mixture was decanted twice with PBS, and then soaked in distilled water and 4 mmol/L acetic acid. After stir for 7 d at 4 °C. The mixture was decanted twice with PBS, and then soaked in distilled water and 4 mmol/L acetic acid. After stir for 7 d at 4 °C. The mixture was decanted twice with PBS, and then soaked in distilled water and 4 mmol/L acetic acid. After stir for 7 d at 4 °C. The mixture was decanted twice with PBS, and then soaked in distilled water and 4 mmol/L acetic acid. After stir for 7 d at 4 °C. The mixture was decanted twice with PBS, and then soaked in distilled water and 4 mmol/L acetic acid.

The supernatant, containing the collagen-I that was stored in 0.25 M Tris-HCl containing 0.15 M NaCl at pH 7.4 for 30 minute [15] and irradiated with microwave irradiation (MWI) (30 s each with intermittent cooling time of 2 min) [15]. Tubes were then exhaustively washed with double-distilled water and dried under sterile conditions.

Surgical procedure

Animals were anesthetized with intraperitoneal injection of Ketamine (90 mg/kg) and Xylazine (10 mg/kg). The left hind limb was disinfected, shaved and prepped. The left sciatic nerve was exposed through a 3-4 cm longitudinal gluteal muscle splitting incision on the lateral side of the left thigh. The sciatic nerve was exposed and dissected free from the underlying muscle bed. In the sham operation, the entire left sciatic nerve was exposed and carefully sutured. A 10-mm sciatic nerve segment was then transected at the mid tight level and removed. In the collagen group, nerve conduit, both the proximal and distal stumps of the sciatic nerve, were inserted into the lumen ends of the nerve guides and fixed in place with a single 10-0 nylon epineurial suture. After placement lumen in the collagen+, E-CSF and collagen +NS groups were filled with e-CSF and normal slain. In the Autograft group, 10 mm nerve segment was turned backwards and used as an autologous nerve graft. On both proximal and distal stumps, single 10-0 nylon epineurial sutures and small gauge needle were used. The muscle border was approximated with 4-0 Dexon sutures and the skin incision were closed with 3-0 nylon. The entire experiment was performed under aseptic conditions using under operating microscope. After surgery, animal was observed and its body temperature maintained with a heating pad until it awoke and were placed in individual cages with free access to food and water ad libitum and a cycle of 12h light/12 hr dark.

Functional tests

To determine the sciatic functional index (SFI), walking track analysis was taken on the day prior to and the 7th, 21st, 35th, 49th, 60th and 90th days after the operation. Black non-toxic water-soluble ink was applied to the plantar surface of the hind feet to dip all anatomical regions, and the rats walked across a clean sheet of white paper track and leave footprints. The footprints were measured and SFI calculated on both operated and unoperated limbs according to the formula developed by Bain [16].

Formula: $SFI = -38.3 \left(\frac{(EPL-NPL)}{NPL}\right) + 109.5 \left(\frac{(EPS-NTS)}{NTS}\right) + 13.3 \left(\frac{EIPNIT}{NIT}\right) - 8.8$

SFI 0 and 100 indicate normal function and complete transaction of the sciatic nerve.

Electrophysiological study

At days, 28 and 90-post operation, the rats in each group were anesthetized with an intraperitoneal injection of a mixture of Ketamine (90 mg/kg) and Xylazine (10mg/kg). The animals were prepared for surgery and their body temperature was maintained between 36.5-37°C using a temperature thermostatic pad. The left sciatic nerve (operated side) was re-exposed through a longitudinal muscle splitting incision in the mid-thigh at the pervious surgical site. The electrophysiological study was performed by using Narco bio-system (USA). The stimulating electrode pair were placed 20 mm apart on each side of the epineurial sutures, and a recording electrode was inserted into
the gastrocnemius muscle. The difference in electromyography latency, amplitude and distance between proximal and distal stimulation sites was measured to calculate conduction velocity.

Statistical analysis

Statistical analyses were processed using the statistical SPSS software (version 16.0) and presented as means ± standard error of mean. Result were analyzed using one-way ANOVA were computed with 95% confidence intervals using. A post-hoc study was carried out to examine any significant differences between the groups. Statistically significant differences were considered at P<0.05.

Results

The absence of rat mortality and wound infection postoperatively demonstrated that no wound infections or autophagy occurred. After operation operative procedures were completed, several animal were noted to have ulceration in experimental limbs but these healed without complication. Wounds healed properly, indicating that the operations were well tolerated by the animals. After implantation to repair the defects of the Rat's sciatic nerves, the tubes were completely resorbed by 12 weeks, leaving no residue. Figure 1 shows neurofunctional recovery was assessed by sciatic function index from the day prior to surgery, and then on days 7th, 21st, 35th, 49nd, 60th and 90th post-operation. Prior to surgery, SFI values were near zero for all groups. The mean SFI decreased to -100 after the nerve transaction due to the complete loss of sciatic nerve function in all animals. The SFI improved from the first to the last evaluation in collagen + E-CSF and autograft groups, since it increased from -90.26 ± 3.6 and -90.53 ± 2.55 during the 7th postoperative day to -32.95 ± 5.5 and -47.44 ± 3.21 during the 90th post-operation day, respectively. On the days 49 and 60 post-operation SFI analyses showed that neurofunctional recovery was better in the autograft and collagen +E-CSF than collagen + NS group (P< 0.05) (Figure 1).

Table 1: Nerve conduction velocity (NCV) and maximum Amplitude (AMP) comparisons in each group after operation.

| Group        | NCV (m/s) | Amp (mv) |
|--------------|-----------|----------|
| Collagen +E-CSF | 39.7*     | 5.72     |
| Autograft     | 30.05*    | 4.76     |
| Sham Surgery  | 49.6*     | 9.7      |
| Collagen +NS  | 21        | 3.6      |

*P<0.05, the difference sham surgery and Collagen +E-CSF /Autograft were significant. The difference between Collagen +E-CSF and Autograft groups were not significant (P>0.05, One-Way-ANOVA). Results are presented as mean ± SEM.

NCV: Nerve Conduction Velocity; AMP: Amplitude; E-CSF: Embryo Cerebro-Spinal Fluid

Discussion

Results from this study have clearly demonstrated the E-CSF in collagen guide channel significantly enhances peripheral nerve regeneration in vivo. Attempts to enhance nerve regeneration and promote functional recover of injured nerves is still one of the most challenging issues and a problem to surgeons [17]. Artificial nerve conduit to bridge the gap between severed nerve stumps is widely accepted as a useful alternative that provides structural support and a favorable microenvironment for axonal regeneration [18]. The great advantage of collagen is very resorbable, and Archibald et al. [19,20] described excellent regeneration following median nerve repair in monkeys as by its structure also offers biochemical support for nerve regeneration, as it has been showed that collagen is one of extra-cellular matrix (ESM) factors that has been shown to play a significant role in the process of nerve regeneration [6]. Collagen nerve conduits degradation speed can be controlled by controlled by manipulating the thickness of the tube and the temperature for thermal dehydration. This condition is essential to permit application of the collagen tube in nerve gaps of different length [19]. Chamberlain et al. [21] used silicon tubes filled with collagen and show improved regeneration over a 10 mm rat sciatic nerve gap compared to empty silicone controls. Our results demonstrate that both collagen and autograft repairs promote nerve regeneration. In this respect, neuromas had not been observed in collagen tube repaired nerves in which regeneration was successful. However, physical nerve guidance by them may not be sufficient to foster optimal nerve regeneration and functional recovery [3]. The neurotrophins NGF, NT-3 and BDNF show a well-defined and selective beneficial effect on the survival and phenotypic expression of primary sensory neurons in dorsal root ganglia and of motoneurons in spinal cord following to axotomy [22,23]. The remarkable finding of this study was the accelerating effect of chick E-CSF administration on axonal regeneration in rats. Chick E-CSF can be obtained in large quantities and is inexpensive, sterilized, and easily stored. Moreover, the fact that the protein fraction of the CSF is more complex in embryos than in adults, and that it is detected at higher concentrations...
NGF is an important neurotrophin that has a role in the survival and proliferation of the cells of the developing nervous system [27] while IGF-I plays an important role in the regulation of mammalian growth [28]. TGF-β1 regulates the differentiation of neuronal, immune, mesenchymal and epithelial cell types [29]. The importance of TGF-β1 signaling has been demonstrated in vascular morphogenesis [30]. During peripheral nerve regeneration, TGF-β1 up-regulates beta fibroblast growth factor expression in the anterior horn motoneurons of the spinal cord [31]. VEGF administration has also been shown to support and enhance the growth of regenerating nerve fibers, probably through a combination of endogenous, neurotrophic and Neuroprotective effects [32-34]. SFI is a simple and repeatable method for evaluating the functional condition of sciatic nerve injuries. After transaction, the SFI value decreased significantly, and NCV could not be detected at first month after surgery. At 90 days after operation, the mean SFI between Collagen +E-CSF and Collagen +NS was significant. The SFI value and NCV of group Collagen +NS was significant. The SFI value and NCV of group Collagen +E-CSF were to similar to that of group autograft. These finding supports that idea that the combination of SFI with electrophysiological assessment is more comprehensive than electrophysiological method alone.

Conclusion
In conclusion, the present study shows that compared to normal saline, treatment with chick E-CSF in collagen guide channel significantly promote peripheral nerve regeneration.

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