Post-Operative Time Effects after Sciatic Nerve Crush on the Number of Alpha Motoneurons, Using a Stereological Counting Method (Disector)

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ABSTRACT

There are extensive evidences that show axonal processes of the nervous system (peripheral and/or central) may be degenerated after nerve injuries. Wallerian degeneration and chromatolysis are the most conspicuous phenomena that occur in response to injuries. In this research, the effects of post-operative time following sciatic nerve crush on the number of spinal motoneurons were investigated. Twelve adult male Wistar rats, whose left sciatic nerves were highly compressed for 30 s, assigned to experimental groups 1 and 2 (n = 6). After 3 and 8 weeks post-operative (in groups 1 and 2 respectively) the lumbar segments of spinal cord were sampled, processed, sectioned serially and stained with toluidine blue (pH 4.65). By using stereological quantitative technique (physical disector), the number of alpha motoneurons in the right and the left ventral horns of spinal cord were counted and compared with each other. Statistical analyses showed a remarkable reduction in the number of alpha motoneurons in the left side (experimental or operated) when compared with the right side (control or unoperated) both in 3 and 8 weeks post-operative groups. This reduction may be due to the blockade of retrograde axonal transport. Iran. Biomed. J. 4: 45-49, 2000

Keywords: Peripheral nerves, Neuronal degeneration, Alpha motoneurons, Stereology, Chromatolysis

INTRODUCTION

A few days after axotomy, changes occur in the cell bodies of the most types of neurons. This phenomenon is referred to as chromatolysis [1]. If the neuron successfully regenerates its axon and restores the connections with other cells in the nervous system, the cell body usually returns to its former appearance [2]. Failure to contact a new target cell leads to atrophy and death [3]. The studies on neural development have identified several neurotrophic factors that are released by the targets of neurons and retrogradly transport to neuronal cell body [2, 4]. These factors are necessary and important for neuronal survival and growth [4-6].

To demonstrate the importance of axonal retrograde transport of neurotrophins, it has been shown that exogenous NGF enhances survival of axotomized neurons in adults and crushed neurons in neonatal rats [7]. Recently, it has been demonstrated that retrograde transport of neurotrophins in intact axons can be abolished by application of drugs such as colchicine [8-10] and neural lesions such as compression [11, 12]. In rodents after peripheral axotomy, some motoneurons survive and undergo typical reactive changes classic for chromatolysis, while other motoneurons undergo changes that lead to cell death. It was also reported that particularly immature rat motoneurons are sensitive to axotomy, with a greater than 80% loss of neurons occurring within 1 week after nerve injuries in newborn rats [13]. The purpose of the present research is to study the post-operative time effects after sciatic nerve compression on the rate of neuronal death. To evaluate this, the numbers of alpha motoneurons in spinal cord segments, L4 to L6 were estimated.

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MATERIALS AND METHODS

Animal model and experimental design. Twelve inbred young male Wistar rats weighing 250-300 g (Serum Production Institute, Hesarak, Karaj) were housed in pairs in plastic cages in the animal house and given rat chow and water ad libitum. Animals were coded and their ears were marked at the time of surgery and randomly assigned to 1 and 2 experimental groups (n = 6). All assessments were made in a blinded fashion and decoded at the end of study.

All the surgical procedures were performed with aseptic techniques. For axotomy, the animals were anesthetized under interperitoneal injection of 0.2 ml of a mixture (1:2) of 10% ketamin (Bayer, Germany) and 2% xylazine (Boxtel, Holland) and the left sciatic nerve was exposed through a gluteal muscle splitting incision. At this location, the nerve trunk was crushed for 30 seconds period between prongs of #5 clamp forceps. On the right side, a control operation was performed which exposed the sciatic nerve but did not disturb it. The muscle and skin were then closed with 14 mm stainless steel sutures. At the selected post-operative times (group 1, 3 weeks and group 2, 8 weeks), rats were anesthetized and intracardially perfused with cold (20°C) 10% formaldehyde (Holland ˚, Serum Production Institute, Hesarak, Karaj) were housed in pairs in plastic cages in the animal house and given rat chow and water ad libitum. Animals were coded and their ears were marked at the time of surgery and randomly assigned to 1 and 2 experimental groups (n = 6). All assessments were made in a blinded fashion and decoded at the end of study.

The spinal blocks were placed in the same fixative for post sampling fixation overnight and then processed and embedded in paraflin. The blocks were sectioned serially at 7 micrometer. A uniform random sampling scheme was employed so that about 10 sections from each block were sampled [14]. Sections were stained with toluidine blue with special buffer of acetic acid 1N (1 ml), sodium acetate 1N (1 ml) and distilled water (98 ml) pH 4.65. After permanent mounting, the number of motoneurons in the left and the right sides of ventrolateral regions of ventral horn of the spinal cord (L4 to L6) were determined, using stereological counting technique; physical disector.

The ‘disector’ principle [15] was used to determine the numbers of motoneurons in each section. From each section and its adjacent neighbor four photos were taken, two from each section; one from right side and the other one from the left side of ventrolateral region of the spinal ventral horn with a final magnification of 100 (Fig. 1). A two-dimeniontal unbiased counting frame [16] was overlaid in a uniform, random manner onto regions of any two photos taken from left sides of both sections. Those cells were nuclei selected by the frame on the reference plane but not appearing on the adjacent "look-up" frame section were deemed to have their ‘tops’ in the volume described by the product of the area of the counting frame and the distance between sections [15]. A similar operation was done on photos of the right sides. These nuclei were counted (Q) to provide the numerical density of cells (Nv) in the ventral horn of spinal cord according to the following equation:

\[
N_v = \frac{\sum Q - \sum (\text{disector}) \times h \times a_{\text{frame}}}{\sum (\text{disector}) \times h \times a_{\text{frame}}}
\]

Where \(\sum Q\) is the sum of counted neurons, \(\Sigma\) (disector) is the number of disectors, \(h\) is the depth of the disector equal to the section thickness (7 micron) and \(a_{\text{frame}}\) is the scaled area of the disector frame. An example of morphometric calculation is as follow: In this research the area of the frame was 25 \(\times\) 25 = 625 mm\(^2\), and the linear magnification was 4 \(\times\) 10 \(\times\) 2.5 = 100. To calculate the \(N_v\) of motoneurons in the right side of spinal cord (Table 1, rat number 1) we should do as follows. A total of 22 neurons were counted in 16 disectors and the depth of the disector (h) was 7 \(\times\) 10\(^3\) mm, and the real area corresponding to each disector frame, at a linear magnification of \(\times\) 100, was:

\[
\frac{625}{(100)_2^2} = 6.25 \times 10^{-2} \text{mm}^2
\]

and the numerical density (\(N_v\)) of motoneurons of spinal cord was calculated according to the above equation:

\[
N_v = \frac{22}{16 \times 7 \times 10^{-3} \times 6.25 \times 10^{-2}} = 3142/\text{mm}^3
\]

The ratio of numerical density of neurons in operated/unoperated sides of spinal cord was then used as an index of neuronal death. All quantitative data were analysed by using ANOVA and \(t\)-test.

RESULTS

The results of the morphometric analyses of numerical density of alpha motoneurons in the ventral horns of spinal cord are shown in Table 1.
After sciatic nerve crush, the spinal ventral horn motoneuron counts show a decline in number. Reduction of cell number appeared complete at 3 weeks post-operative with no additional loss noted at 8 weeks (1489 and 1438 respectively). Although in operated sides of spinal cord of both groups the numerical density is decreased and remained constant but in unoperated sides the numerical density of alpha motoneurons is higher in group 2 (4353 vs 3441). Statistical analyses show that the reduction in the alpha motoneurons in treated sides of spinal cord of both groups, when compared with unoperated sides is significant ($p < 0.0001$) (Table 1, Fig. 2). Comparison of numerical density of motoneurons among operated sides in both groups showed no significant differences but when unoperated sides were compared, there was a significant difference between neuronal density ($p < 0.01$).

Fig. 1. Photomicrographs of toluidine blue stained transverse section from L4-L6 spinal cord following unilateral crushing of sciatic nerve, 3 weeks post-operation. Note the decrease in the numbers of motoneurones in the operated (left) side.

Fig. 2. The comparison of numerical density of motoneurons of spinal ventral horn in operated and unoperated sides, in 3 and 8 weeks post-operation time.

**DISCUSSION**

It has been well known that the survival and functional maintenance of neurons are clearly dependent upon retrogradely transport of neurotrophins [17, 18]. Peripheral nerve transaction or crush, blockades axonal transport and therefore might produce chromatolysis and cell death. In this research, we investigated the post-operative time effects following sciatic nerve crush on the numerical density of alpha motoneurons. The present results, are based on using method which provides absolute estimates of cell number rather than ratios [15, 19, 20]. This method has been rapidly gaining acceptance, because not only it is the most efficient quantification method, but also unbiased and simple to employ.

Statistical analyses of data indicate that, in comparison with unoperated side of spinal cord, the numerical density of alpha motoneurons in the operated side significantly decreases. However, there are no remarkable differences between group 1 and 2, which indicates that post-operative time has no effect on neuronal loss. It is possible that, the site of compression was very close to the spinal cord (gluteal region) and therefore after a few days following axotomy alpha motoneurons undergo degeneration. It has been shown that particularly immature rat motoneurons are sensitive to axotomy, with a greater than 80% loss of neurons occurring within 1 week after nerve injuries in newborn rats [10]. The magnitude of motoneuronal death observed in this research is consistent with that of previously reported [13]. Because sciatic nerve is a mixed nerve, it seems very likely that after compression efferent and also afferent pathways are damaged [21-23]. Therefore, it is possible that,
in addition to motoneurons, some interneurons which receive inputs from afferent pathways undergo degeneration. Since motoneurons of unoperated side also receive inputs, at least partly, via crossed interneurons, the transneuronal degeneration might affect cell bodies of motoneurons of unoperated side. By passing time, the remaining interneurons try to re-innervate further motoneurons. If this is the case, the appereance of motoneuronal cell bodies of the unoperated side becomes normal and therefore falls into counting. This explanation may be the cause of elevation of numerical density in unoperated side of group 2 (4353 vs 3441). The measurement we have used here allows an unbiased assessment of numerical density, and the result suggests that, although degeneration processes are completed as early as week 3 after compression, a partial recovery in numerical density of motoneurons in unoperated side may be possible.

ACKNOWLEDGMENT

This work was supported by the grant #3-2/5272 from Office of Research Affairs of Ferdowsi Mashhad University. The authors wish to thank Dr. A. R. Fazel, Mashhad University of Medical Sciences, for kindly reviewing the manuscript and M. Khaksar and M. Sabzehbin for their technical assistance and animal care.

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