Mandibular Branch of the Facial Nerve in Wistar Rats: New Experimental Model to Assess Facial Nerve Regeneration

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Abstract

Introduction The ideal animal model for nerve regeneration studies is the object of controversy, because all models described by the literature have advantages and disadvantages.

Objective To describe the histologic and functional patterns of the mandibular branch of the facial nerve of Wistar rats to create a new experimental model of facial nerve regeneration.

Methods Forty-two male rats were submitted to a nerve conduction test of the mandibular branch to obtain the compound muscle action potential. Twelve of these rats had the mandibular branch surgically removed and submitted to histologic analysis (number, partial density, and axonal diameter) of the proximal and distal segments.

Results There was no statistically significant difference in the functional and histologic variables studied.

Conclusion These new histologic and functional standards of the mandibular branch of the facial nerve of rats establish an objective, easy, and greatly reproducible model for future facial nerve regeneration studies.

Introduction

To study the processes that affect facial nerve degeneration and regeneration, several experimental models were created. Large animals such as pigs have a long vertical (mastoid) segment of the facial nerve, great for surgical procedures but difficult to use because of their handling, high maintenance costs, and small resistance to infections,1 besides having a high incidence of anesthetic complications.2 Medium to small mammals such as cats, rabbits, and guinea pigs are easier to handle and cost less compared with large animals,2 have facial nerves that anatomically resemble the human facial nerve,2,3 and have well-known reactions to posttraumatic microbiological exposure.4 However, such animals are relatively difficult to handle and maintain, are more prone to infections, and are less tolerant to bilateral facial paralysis.1,2 At the same time, small animals such as rats are being widely employed in experimental research for their advantages, such as easy caretaking and handling, high resistance to infections and interventions, low-cost and convenient housing, besides having well-described growth, anatomic, and metabolic parameters.1,2,5,6 Regarding facial nerve studies, rats endure well bilateral facial paralysis and exhibit similar anatomic nerves as primates.5

Keywords
► facial nerve
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One of the most used experimental models of peripheral nerve regeneration involves the sciatic nerve of rats, mainly because of its easy exposure and long extension. However, this nerve divides itself after leaving the sacrum, possibly altering the results of functional or histologic investigations. Furthermore, authors such as Aldskogius et al., Labelle and Allen, McCoy and Boyle, Spector et al., and Saz and Miller refer that, because of its long path inside a bony canal, the facial nerve has remarkable differences in regeneration and therefore cannot be compared with the sciatic nerve.

Electrophysiologic studies have the purpose of quantifying the function, and, therefore, investigating the pathologies of the peripheral nervous system, especially of the sensitive and motor neurons, the neuromuscular junction, and the motor unit. Hence, they are a fundamental part of experimental models that aim to evaluate nerve regeneration. Methods of subjective and noninvasive evaluation of the facial nerve have been created, such as the “walking track,” related to the sciatic nerve. Facial mimics observation scales, precise videotaping of facial movements, and muscle fibrillation sometimes associated to scales are examples of widely used methods of functional facial nerve evaluation. However, all these methods are subjective and, thus, altered when performed by distinct examiners or in the presence of postsurgical scars or edema.

Nerve morphometric studies, which include counting of axons and of axonal density and diameter, are considered by many authors an objective method of evaluating nerve regeneration. Qualitative analysis, although being a subjective method, can be associated to quantitative analysis to determine the quality of regenerated nerve tissue, especially in a time-dependent manner, adding precision to the experimental results.

Therefore, this experimental study aims to describe the qualitative and quantitative histologic patterns and motor neuroconduction patterns by electroneuromyography of the mandibular branch of the facial nerve of Wistar rats, to create a new experimental model for facial nerve regeneration studies, that is both objective and thorough.

Materials and Methods

All experiments were conducted according to the Guide for the Care and Use of Laboratory Animals, after approval by the institution’s ethics committee under protocol number 0291/07.

Functional evaluation included electroneuromyography of the mandibular branch of the facial nerve of rats. Forty-two male Wistar rats, weighing 250 to 300 g, with body temperatures ranging from 34.5 to 36.5°C, were anesthetized by an intraperitoneal injection of 10% chloral hydrate at 400 mg/kg and subjected to neurophysiologic evaluation by a motor neuroconduction test of the mandibular branch of the facial nerve to obtain the compound muscle action potential (CMAP) for posterior analysis of its latency, amplitude, and duration.

The motor neuroconduction test was performed according to method previously described by Salomone et al and consists of obtaining CMAPs by supramaximal stimulation of the mandibular branch, identified by transparency, through the skin at a distance of 20 mm from the labial commissure. The recording electrodes are two subdermal needle electrodes, separated by 5 mm and located 2 mm from the lower lip and labial commissure.

To prevent the influence of anatomic variations on electrical signals, all animals presenting branches of the mandibular nerve to the upper lip or branches of the buccal nerve to the lower lip or branches that connected the two were excluded.

Histologic evaluation included qualitative and quantitative study of the mandibular branch of the facial nerve of rats. Twelve rats, randomly chosen after electroneuromyography, were subjected to dissection, in situ fixation, and removal of the mandibular branch of the facial nerve, according to method described by Costa et al. This method consists of removing 3 mm of the previously fixated nerve by two transversal incisions, made 25 and 28 mm from the labial commissure. Therefore, a nerve segment with two ends—distal (D) and proximal (P)—is produced. After removal, nerve segments were treated with a 2% osmium solution, dehydrated with ethanol, infiltrated with propylene oxide, and embedded in Epoxi resin (Burlington, Vermont, United States) until polymerization. After these steps, P and D segments were cut into smaller ones of 0.5 µm in width and stained with 1% toluidine blue.

For nerve documentation, slides with P and D segments were photographed (Nikon Coolpix E 955, Nikon Corporation, Tokyo, Japan) with an optical microscope using 50× magnification (one field to view the entire nerve) and 400× magnification (three fields of 3,000 µm², comprising a total area of 9,000 µm²). Afterward, images were transferred to a portable computer and analyzed by SigmaScan Pro 5.0 (Statistical Package for Social Sciences [SPSS] Science, Chicago, Illinois, United States). Photographs obtained with the 50× magnification were used for evaluation of the qualitative aspects of nerve segments, namely, quantity of fascicles and axonal arrangement (morphometry and distribution of myelinated axons).

Quantitative evaluation of P and D segments was performed with three images obtained with the 400× magnification and involved analyzing the total number of myelinated axons in a 9,000-µm² nerve area, the partial density of myelinated axons (partial quantity of myelinated axons/partial area [9,000 µm²]), and myelin maturation by measuring the smallest outer nerve diameter (including the myelin sheath) in the 9,000-µm² area.

Statistical Analysis

The Wilcoxon signed rank test from the SPSS software, version 21.0, was used to assess differences (p < 0.05) in values of CMAP latency, amplitude, duration, area, and intensity and in total axon number, axonal diameter, and axonal density of P and D segments obtained from each side.

Results

Table 1 compares means, medians, standard deviations, and percentiles of amplitude, latency, duration, and intensity of...
CMAPs obtained from each side. There were no statistically significant differences between sides.

Qualitative histologic evaluation of P and D segments showed no noticeable differences. All segments had two to three well-defined fascicles (Fig. 1A, B), with normal axons, exhibiting similar diameter and homogeneous distribution (Fig. 1C, D). The segment’s perineurium exhibited condensed loose tissue and the epineurium was visibly preserved (Fig. 1A–D).

Table 1 compares means, medians, standard deviations, and percentiles of axon number, axonal density, and medium axonal diameter of the P and D segments. Again, there were no statistically significant differences observed.

Discussion

Wistar rats were chosen for this new model to evaluate facial nerve regeneration not only because of the advantages they...
present as small animals (described in the introduction) but also because they have a very long and purely motor extra-temporal segment of the facial nerve, which is easy to access and divides itself similarly in all rodents. Another advantage is that, in Wistar rats, the facial nerve passes below and not inside the parotid gland, and its peripheral branches lay above the superficial facial fascia, simplifying exposure and dissection.2,7,20 Additionally, the facial nerve fibers do not vary according to animal weight or age.2–20 Male rats were chosen because of the proven effect that female hormones have on nerve regeneration.21

The mandibular branch was chosen because of some unique qualities: (1) easy access compared with other branches such as the posterior auricular, cervical, temporal, and zygomatic branches2,23–25; (2) longer path (10 to 12 mm) before splitting compared with the facial nerve trunk (6 mm) and to its posterior cervical, cervical, buccal, auriculotemporal, zygomatic, and temporal branches, allowing surgical manipulation and graft alignment with reliable results2–5,8,9, (3) less complex anatomy compared with the temporal and zygomatic branches2–22; (4) visualization of the nerve through the skin, previous to surgical intervention, avoiding operations on animals with anatomic variations, such as branches to the upper lip, other additional branches, or anastomosis with the buccal branch of the facial nerve, which could jeopardize the results of electrophysiologic tests5; (5) homogeneous histology2,19,23; (6) highly reproducible nerve conduction tests5; and (7) resemblance to the human facial nerve, including the presence of nerve fascicles2,23,25.

To create an objective method of functional evaluation of the facial nerve of rats, several methods were described. Mattox and Felix were pioneers in stimulating the facial nerve in multiple segments (trunk, buccal and mandibular branches) and registered answers with subcutaneous electrodes located on the upper and lower lips of rats.2 Several years later, Byrne et al stimulated the facial nerve trunk and registered the CMAPs using electrodes located bilaterally and symmetrically near the whiskers.23 Posteriorly, Shi et al evaluated only the buccal branch of the facial nerve, inserting it through a silicone tube, stimulating the proximal segment and registering the answer with microneedles in the distal segment to obtain the nerve’s action potential.24 However, besides being invasive, all the methods above failed to establish an exact place for the stimulus, which can cause distortion of distance-dependent results, such as latency and duration of CMAPs.25,26

Another important topic to discuss regarding functional evaluation methods by electroneuromyography is the intensity of the stimulus. Amplitude is directly proportional to the number and synchrony of intact axonal fibers that the stimulus depolarizes.7,28,29 Therefore, below normal amplitude may occur because of the reduction of number or synchrony of preserved fibers or because of a low stimulus. Hence, it is necessary to establish a standard of stimulus intensity. In this sense, the supramaximal stimulus, which is the minimum intensity necessary to depolarize all the nerve fibers plus 20%,5–7,10–25,27–29 is ideal.1,28,29 Mattox and Felix2 did not describe the intensity of the stimulus used. Shi et al used a constant current of 0.5 mA,24 and Byrne et al found a supramaximal stimulus of 1.5 mA,23 different from the value that our group obtained in a series of experiments.5–7,10–30

Anatomical variations also demand caution. As previously described, the mandibular branch of the facial nerve of Wistar rats can give branches to the upper lip, anastomosing the mandibular and buccal branches.2,23–25 This fact was confirmed by Mattox and Felix when they stimulated the buccal and mandibular branches separately and detected synchronous answers in electrodes located in the upper and lower lips.2 However, the same authors demonstrated that the CMAPs produced by the mandibular branch (lower lip) have a broader trace but lower amplitudes when compared with those of the buccal branch (upper lip). Visualization of the nerve before surgery, through the skin, helps exclude animals with anatomical variations.

Our results show that there were no statistically significant differences in values of amplitude, latency, duration, and intensity of CMAPs when left and right sides were compared. The values obtained for these variables were compatible with those standardized by Salomone et al5 and endorsed by Costa et al,30 demonstrating that this objective evaluation method is highly reproducible.

In regard to the histologic studies, authors disagree about the best quantitative morphometric parameter to evaluate nerve regeneration. According to Hubbard31 and Vasconcelos and Gay-Escoda,32 the total axon number is the most important criterion. For Miyamoto,33 axonal density is the most

| Axonal Segment | n | Mean value | Standard derivation | Lower limit | Higher limit | 25th percentile | Median | 75th percentile | p value \( ^* \) |
|----------------|---|------------|---------------------|------------|-------------|----------------|--------|----------------|----------|
| Number         | Proximal | 12 | 1,662 | 131.03 | 1,336 | 1,847 | 1,590.75 | 1,669.5 | 1,753 | 0.666 |
| Density (number/\( \mu m^2 \)) | Proximal | 12 | 0.18 | 0.01 | 0.15 | 0.21 | 0.18 | 0.19 | 0.19 | 0.666 |
| Diameter (\( \mu m \)) | Proximal | 12 | 4.5 | 0.21 | 4.2 | 5.01 | 4.36 | 4.48 | 4.53 | 0.937 |

\( ^* \)Wilcoxon signed rank test.
important variable. Meanwhile, Donovan, Duncan, Fernand and Young, and Williams and Wendell-Smith defend that nerve fiber diameter is the most reliable parameter to compare experimental groups and individuals in the same experimental group. May and Schaitkin support that both axonal diameter and the width and length of the myelin sheath are the most relevant factors regarding histologic nerve recovery.

According to Harrison, the analysis of morphometric characteristics of myelinic fibers can be accomplished with software, simplifying counting and also yielding precise and reliable results. Furthermore, it is unnecessary to evaluate the morphometric characteristics of the entire nerve, because the results obtained in 6% of the total area are as precise as those of the whole area of myelinic fibers of a nerve.

In this study, the partial area analyzed (9,000 µm²) represents almost 90% of the total area of the mandibular branch of Wistar rats. The morphometric histologic data (total axon number, axonal density, and medium axon diameter in a partial area of 9,000 µm²) were analyzed by two independent and blind examiners, using the SigmaScan Pro 5.0 software (SPSS Science), and the final data represent the medium value of that obtained from the two examiners. The fibers that had centers out of the microscope’s field were excluded to avoid the edge effect and any possible resultant sampling errors.

Mattox and Felix described that normal mandibular branches of the facial nerve of Wistar rats contain between three and five fascicles. Our results show that the number of fascicles varied between two and three, up to their insertion in the lower lip muscle. However, the results obtained in the quantitative and qualitative histologic studies (number of axons, axonal density, and diameter) of the P and D segments were similar to those described by Mattox and Felix, Costa et al, and Salomone et al, indicating that the mandibular branch of the facial nerve of Wistar rats varies very little between sides and animals, besides being more similar to the human facial nerve when compared with the facial nerve trunk, qualifying as an adequate model for the study of facial nerve regeneration.

On the other hand, according to Ochi et al, the results obtained with histologic methods of neural regeneration, despite being qualitative, quantitative, or both, must be analyzed with caution, because the increase in the size or quantity of axons may not correspond to functional improvement. Regarding this, the authors note that species with many myelinated small axons or few myelinated large axons behave similarly during regeneration. Grosheva et al are even more extreme, claiming that axonal quantity, density, and diameter have no importance if axonal growth is not directed toward the motor end plate.

Conclusions
With histologic (qualitative and quantitative) and functional (nerve conduction studies) standardization of the mandibular branch of the facial nerve of rats, we obtained a novel experimental model, which is objective, easy, and highly reproducible, for future facial nerve regeneration studies.

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