Standardization of techniques used in facial nerve section and facial movement evaluation in rats

Sumário

OBJETIVO: standardização da técnica para seccionamento da facial extra-temporal em ratos e criação de uma escala para avaliação dos movimentos faciais destes animais antes e após a cirurgia. Métodos: vinte ratos Wistar foram anestesiados com xilazina e ketamina e submetidos à seccionamento da facinal ao redor do orifício mastóideo. A constrição dos movimentos ocular e vibrissal de todas as ratas foram observados e uma escala de avaliação desses parâmetros foi criada. Resultados: foi encontrada a ação da faciana entre as margens tendineas de clavotrapezius e auricular. O tronco foi seccionado proximalmente ao seccionamento do mastóideo e os punhados suturados com 9-0 nylon. Uma escala de avaliação e graduação de movimentos faciais, independente de ocular e vibrissal, foi criada, juntamente com uma soma dos parâmetros, como um meio de avaliação de alopático facial. A consistência do movimento ocular e reflexo de fechamento ocular foi abandonada com uma avaliação e graduação de 1; a presença de contração muscular orbicular, sem reflexo de fechamento ocular, foi abandonada com uma avaliação e graduação de 2; 50% de fechamento ocular por reflexo de fechamento ocular, foi abandonada com uma avaliação e graduação de 3; 75% de fechamento ocular por reflexo de fechamento ocular, foi abandonada com uma avaliação e graduação de 3; 100% de fechamento ocular por reflexo de fechamento ocular, foi abandonada com uma avaliação e graduação de 4; a presença de fechamento ocular completo e reflexo de fechamento ocular, foi abandonada com uma avaliação e graduação de 5. Conclusão: A anatomia do ratão permite fácil acesso ao facial extra-temporal, permitindo sua seccionamento e standardização de sutura. Foi possível também estabelecer uma escala de avaliação e graduação de movimentos faciais do ratão com alopático facial baseada na observação clínica destes animais.

Keywords: evaluation, surgery, facial nerve, facial palsy, rats.
INTRODUCTION

Facial paralysis caused by facial nerve injury is a relatively common clinical condition. Its unilateral manifestation may cause alterations in facial symmetry, incomplete eye closure with or without ophthalmic repercussion, difficulty swallowing and in pronouncing some phonemes, amongst other problems. Therefore, this condition brings about relevant social and individual consequences, requiring great efforts in the attempt to understand the many factors related to facial nerve injury, its regeneration and repair, so as to reduce its consequences.

The facial nerve motor fibers, although they originate in the central nervous system, have great regenerative capacity, comparable to a peripheral motor nerve. The facial nerve has a debris and myelin clearance greater than that of the Central Nervous System (CNS) and, contrary to what happens to the CNS, the facial nerve has some sort of guidance for its regenerative process. The facial nerve also has better response capability to growth factors after injuries.

After injury, the axon is influenced by a number of events, among them we list: local paracrine factors, retrograde molecule transportation (cytokines and growth factors), calcium and sodium inflow, generation of free radicals, increase in the activity of phospholipases, generation of hyaluronic acid derivatives, genetic factors (balance between bcl2 and bax/bak).

Wallerian degeneration, which follows nerve injury, is influenced by these factors and is completed in approximately thirty days, causing the removal of the distal axon fragment. In the regeneration process, which starts in the first Ranvier node, close to the injury site and that has a growth rate of one to two millimeters per day, the Schwann cells play a fundamental role (they remove neural debris, secrete cytokines and growth factors); notwithstanding, the cells play a fundamental role (they remove neural debris, secrete cytokines and growth factors); notwithstanding, the facial nerve has some sort of guidance for its regenerative process. The facial nerve also has better response capability to growth factors after injuries.

Thus, the type of injury to which the nerve is submitted is of fundamental importance to the regeneration process and the approach to be used. Sunderland classified the nerve injury based on histopathological findings into: first grade – intact nerve structure (neurapraxia); second grade – ruptured axon with intact Schwann cells (axonotomy); third grade injury to the endoneurium; fourth grade – injury to the perineurium and fifth grade – complete rupture (neuromatosis). The first two injury grades usually allow full nerve recovery. However, when there is loss of nerve continuity, its direct repair has been advocated as the most effective way to specifically rehabilitate the facial nerve. When this is not possible, it is necessary to use nerve grafts.

Although surgical techniques have improved, the functional result related to the repair of full injuries to the facial nerve is still short of being satisfactory. Such fact has fostered many experimental studies in attempts to establish the best approach, and the best moment to approach an injury to this nerve.

Another line of research is to study the substances that act on facial nerve regeneration after injury, in an attempt to introduce a therapeutic alternative that may bring about better results, specially in paralysis caused by a full section of the facial nerve. Among these substances we have the neurotrophic factors, androgens and nimodipine.

With the progress we are seeing in experimental research, many animal models have been used (rats, mice, rabbits, cats). However, there is no experimental model standardized to study facial paralysis – behavioral evaluation, approach technique, facial nerve injury and regenerative process onset.

It is of great importance to establish a scale to assess facial movements and to standardize the surgical technique used to approach the facial nerve and the various stages of its regenerative process in order to better develop investigations on systemic drugs that may promote a better recovery for the facial nerve.

The goal of the present study is to standardize the technique of sectioning the facial nerve extratemporally in rats and establish a scale to evaluate facial movement in these animals before and after the sectioning of this cranial nerve.

MATERIALS AND METHODS

Twenty adult, male, Wistar rats, weighing between 250 and 300g, were used in this study. The animals were kept in a proper room, with light-dark cycles of 12 hours. Housed in proper cages, they had free access to food and water. All the procedures were carried out according to the recommendations from the Ethics Committee in Research of the São Paulo Hospital – Federal University of São Paulo (UNIFESP).

These rats were anesthetized by intraperitoneal injection of 2% Xylazine (0.5ml/kg) and 10% ketamine (0.9ml/kg). After that, the hair from the right side retroauricular region was removed. They were then placed in lateral decubitus and received an injection of xylocaine (it was repeated if necessary) in this region, and then we started the surgical procedure, which resulted in right side peripheral facial paralysis in these animals. The procedure was carried out by two surgeons, through the use of surgical microscopes DF-Vasconcellos M90, with a 200mm lens and 16x magnification. Surgical results were photographed with a 3.2 megapixels Sony Cybershot camera and were standardized.

All the animals were observed as to spontaneous and stimulated facial movements. This assessment was carried out by the same observer, and occurred before the...
surgery (to rule out facial movement alterations present prior to surgery that could compromise the later assessment), and in alternate days in the postoperative time, during 30 days. The parameters observed were: eye closure, blinking reflex, vibrissae movement and positioning. To do that, each rat was individually evaluated in a cardboard box (37x21x13cm) painted black inside (to better visualize the vibrissae), and as stimulation we used air inflation (a 20 ml syringe) on the animal face to trigger the blinking reflex, and hand clapping (three to four times) to cause vibrissae movement. The left side served as the control side (blinking reflex present with complete eye closure; normal and correctly positioned vibrissae movement). Through this observation we created a scale to assess facial movement in these animals.

RESULTS

With the animals properly positioned, the surgery started. After incision in the right retroauricular region (about three centimeters), the surgeons proceeded with the dissection by planes. The platysma muscle was superficially identified, and its cranial portion was sectioned. The tendinous border of the clavotrapezius muscle was found at the postero-superior portion of the incision and the facial nerve trunk was located close to this border, which was then pushed posteriorly in order to better show the nerve emergence. There was no need to cut this border. The nerve trunk was then followed antero-inferiorly down to its furcation, located medially to the external jugular vein and its branches. The facial nerve branches covered by fascia were not dissected.

The facial nerve trunk was then cut with the scalpel, proximal to its emergence, and then sutured with nylon 9.0 wire. The stumps were sutured with one stitch. In seven animals (35%), a small branch came off the main trunk close to the mastoid foramen. In four cases, we sectioned and sutured the branch and the trunk, however in the other three cases we managed to isolate the main trunk and section it alone.

After epineurium suture, we closed the skin with simple stitches made by nylon 4-0 wires, cut close to the incision (Pictures 1 to 4).

No animal was excluded from the study. Based on our observation after nerve cutting and suturing, it was possible to establish a scale to assess facial movements in these rats. Eye closure through the blinking reflex and vibrissae movements were observed and distributed separately in one scale.

Vibrissae movement and its anterior positioning, symmetrically in the control side, was established as a sign of complete functional recovery of this facial nerve branch in the operated side, and received a score of 5. Blinking reflex present with complete eye closure also received a score of 5 points. Therefore, the complete recovery of the right side of the face received a score of 10, same value assigned to facial movements before surgery (complete recovery was not seen in any animal for 30 days).

The complete lack of blinking reflex without ocular closure scored 1 point, and the lack of vibrissae movements with posterior positioning scored 1 point as well. Thus, complete facial paralysis scored 2 and was seen in all rats that underwent the surgical procedure.

A light vibrissae tremor on the anterior position, scored 2 points, tremor with the vibrissae positioned more posteriorly scored 3 and the normal vibrissae movement keeping its posterior position scored 4 points.

The lack of blinking reflex (no ocular closure) and the contraction of the orbicular muscle scored 2 points; the closure of half the ocular cleft (50%) with blinking...
most are nerve sectioning and nerve crushing. However, there is no established pattern for facial nerve surgical approaches and for the observation of the subsequent facial paralysis in these animals. This standardization could bring great uniformity to the studies with certain animals and would certainly facilitate result comparisons.

A large number of studies have used rats as experimental models because they are easy to acquire and they tolerate well the facial paralysis, even when bilateral, without presenting problems in the immediate postoperative time\textsuperscript{13}.

The rats used in the present investigation were the Wistar which, together with the Sprague-Dawley, are the most frequently used.

The animals were anesthetized by intraperitoneal administration...
reflex present scored 3 points; and the almost total closure (75%) scored 4 points.

Charts 1 and 2 show the scales created:
The animal assessment through this scale may be exemplified in the pictures that follow. (Pictures 5 to 7).

**DISCUSSION**

The human face may be considered a mirror of our feelings, since it is through the movement of its muscles that we express different types of feelings in different degrees. It also has a fundamental participation in social communication because it is integrated to the speech process.

**Chart 1. Vibrissae observation scale.**

| Score | Movement | Position |
|-------|----------|---------|
| 1     | No movement | posterior |
| 2     | Light tremor  | posterior |
| 3     | Greater tremor | posterior |
| 4     | Normal movement | posterior |
| 5     | Normal movement | anterior |

**Chart 2. Scale of eye closing and blinking reflex observation.**

| Score | Movement |
|-------|----------|
| 1     | No movement |
| 2     | Contraction/no closure |
| 3     | 50% closure |
| 4     | 75% closure |
| 5     | Complete closure |

Facial nerve integrity, responsible for the innervation of the facial muscles, bears a relevant role in the social and individual context. Important physiological functions depend on this integrity, such as tearing and ocular protection, taste (anterior 2/3 of the tongue), food intake (mouth orbicular muscle takes part in the initial process), salivation, among others. Many are the causes of this dysfunction, inflammatory processes and injuries are among the most common.

Because of the relevant impact its dysfunction may cause, the facial nerve has been studied in the most varied ways, and experimental studies play a key role in this process.

Many animals have been used as experimental models, among them we list: rabbits, pigs, rats, mice and cats. Different types of facial nerve injuries are described in facial paralysis investigations, and the two that appear the injection of 2% Xylazine (0.5ml/kg) and 10% ketamine (0.9ml/kg). Such combination is often used to anesthetize both rats and other animals used in experimental studies.

The retroauricular incision used in this study allowed a clear field of vision to explore the facial nerve, and it is frequently considered a reference in papers that describe this approach in rats in details and in other animals. Another very much used incision is the intraauricular.

The facial nerve may be approached in its intratemporal pathway, notwithstanding; we chose to approach it in its extratemporal pathway, close to its emergence through the mastoid foramen, because it is the most commonly used way in studies that cause facial paralysis in rats.

In order to find the facial nerve emerging through the mastoid foramen, we sectioned the platysma muscle and used the auricular cartilage and the tendon border of the clavotrapezius muscle as reference points. This way, we located the main trunk of the facial nerve, between the auricular cartilage, anteriorly, and the clavotrapezius posteriorly. We followed this trunk until we found its furcation into minor branches, in order to confirm that structure was really the facial nerve, since we did not use intraoperative stimulation on the nerve. The use of intraoperative stimulation on the facial nerve has been described only in a handful of studies.

Having found the nerve trunk, we sectioned it close to the mastoid foramen. However, in seven animals the posterior cervical branch, probably, diverged from the main trunk still close to the mastoid foramen, was difficult to isolate and section. In four cases we sectioned the trunk and the branch, and in the other three, we were able to isolate the main branch and cut it. We did not find references in these regards in the literature. After sectioning the trunk, the stumps were sutured with a single epineurium stitch, using a nylon 9-0 wire because it is technically easier. Although the stump suturing was been carried out by Fernandez et al. (1992 and 1995) with a nylon 10-0 wire and by Guntinas-Lichius et al. (2001 and 2005) with nylon 11-0 wire, Yian et al. (2001) in his studies with rabbits used the nylon 8-0 suture wire.

And finally we sutured the skin with simple nylon 4-0 stitches. Fernandez et al. (1995) used metal clips to close the skin, while Bento & Miniti (1989) in their study with cats, used catgut 2-0 wire to close the skin in a single layer; and Mersa et al. (2000) used separate nylon 4-0 stitches to close the rabbit skins. This final step of the surgical procedure also varies and is also not very often mentioned in the literature.

Observing the behavior of the animals involved in the facial paralysis study has been used as a parameter to assess facial nerve regeneration. Studies with rabbits...
hamsters\textsuperscript{10} or rats\textsuperscript{2,14,20} observed eye closure, blinking reflex and vibrissae movement and positioning. In non-operated animals, the blinking reflex should be present as symmetrical and complete eye closure, as well as vibrissae movement, which should be positioned anteriorly. Thus, in animals with facial paralysis, there are characteristic signs that include the loss of the blinking reflex and no vibrissae movement, which remain posteriorly positioned\textsuperscript{10,20,21}. By observing these signs, distinct facial paralysis scales have been described.

Yian et al. (2001)\textsuperscript{13} developed a scale to observe upper lip and eyelid movements in rabbits (who underwent cutting of facial nerve branches), with and without stimuli (forehead and nose touch). This scale assigned values between 0 and 3, 0 was assigned to the lack of movement and 3 to normal movement; 1 and 2 points were given to mild and moderate movement, respectively. Byers et al. and 3 to normal movement; 1 and 2 points were given to between 0 and 3, 0 was assigned to the lack of movement (forehead and nose touch). This scale assigned values cutting of facial nerve branches), with and without stimuli (facial movement in a symmetrical fashion. Haldock et al. (1996)\textsuperscript{20} assessed only the vibrissae function, their normal movement and position. In their study, Mersa et al. (2000)\textsuperscript{2} observed eye closure and the blinking reflex, and they built a scale with values between 0 and 2; and 0.5 was assigned only to the contraction of the orbicular muscle contraction. In their study with cats, Bento & Miniti (1989)\textsuperscript{12} studied behavior, paying special attention to eye closure and the blinking reflex, and the presence or absence of facial movement in a symmetrical fashion. Haldock et al. (2005)\textsuperscript{18} created a scale with values between 0 and 1, but contrary to these studies, they applied it separately to assess vibrissae movement and eye closure. They justified this separation based on eye and vibrissae function recovery, which occurred at distinct times.

We have then created an observational scale in order to assess vibrissae position and movement, and another, with the same variation, to assess eye closure and blinking, because we noticed the asymmetrical recovery of facial function as mentioned by Haldock et al. (2005)\textsuperscript{18}. The parameters assessed were similar to the ones mentioned: lack of movement, vibrissae tremor, normal movement and their posterior or anterior positioning; contraction of eye orbicular muscles, complete and incomplete eye closure. The scale has five scores (between 1 and 5), and also a separate score for the parameters. It includes a summation of points for each distinct parameter, similar to the Facial Gradation System presented by Ross et al.\textsuperscript{19} in 1996 to clinically assess facial recovery in human beings. Thus, it is possible to have an idea of the recovery for one isolate parameter and for the whole face of the animal, and also to compare animals from different groups (if the study involves both a study group and a control group), in a more accurate fashion.

Therefore, the rat allows the observation of its eyes and vibrissae movements and also the creation of an independent scale that allows a better analysis of these parameters and of facial movements as a whole. The facial nerve approach, in its extratemporal pathway, with cutting and suturing, is also a procedure that may be standardized in the rat, because this animal has an accessible facial nerve trunk and with such variability that does not compromise such approach.

**REFERENCES**

1. Choi D, Dunn IT. Facial nerve repair and regeneration: an overview of basic principles for neurosurgeons. Acta Neurochir (Wien) 2001;143:107-14.
2. Mersa B, Tiangco DA, Terzis JK. Efficacy of the “baby-sitter” procedure after prolonged denervation. J Reconstr Microsurg 2000;16(1):27-35.
3. Fernandez E, Pallini R, Marchese E, Lauretti L, Bozzini B, Sbriccoli A. Reconstruction of peripherial nerves: the phenomenon of bilateral reinnervation of muscles originally innervated by unilateral motor-neurons. Neurosurgery 1992;30:364-9.
4. Guntinas-Lichius O, Neiss WF, Schulte E, Stennert E. Quantitative image analysis of the chromatolysis in rat facial and hypoglossal motoneurons following axotomy with and without reinnervation. Cell Tissue Res 1996;265:537-41.
5. Barrs DM. Facial nerve trauma: optimal timing for repair. Laryngoscope 1991;101:835-48.
6. Chen YS, Hsu CJ, Liu TC, Yanagihara N, Murakami S. Histological rearrangement in the facial nerve and central nuclei following immediate and delayed hypoglossal-facial nerve anastomosis. Acta Otolaryngol 2000;120:551-6.
7. Guntinas-Lichius O, Efflenberger K, Angelov DN, Klein J, Streppel M, Stennert E, Neiss WF. Delayed rat facial nerve repair leads to accelerated and enhanced muscle reinnervation with reduced collateral axonal sprouting during a definite denervation period using a cross-anastomosis paradigm. Exp Neurol 2000;162:98-111.
8. Jergovic D, Stal P, Lidman D, Lindvall B, Hildebrand C. Changes in a rat facial muscle after facial nerve injury and repair. Muscle Nerve 2001;24:1202-12.
9. Kohnura E, Takamichi Y, Toshiyuki Y, Toshiyuki F, Norimasa K, Akhi-ko S et al. BDNF atelocollagen mini-pellet accelerates facial nerve regeneration. Brain Res 1999;849:255-8.
10. Jones KJ. Recovery from facial paralysis following crush injury of facial nerve in hamsters: differential effects of gender and androgen exposure. Exp Neurol 1993;121:133-8.
11. Mattson P, Aldskogius H, Sevensson M. Nimodipine-induced improved survival rate of facial motor neurons following intracranial transection of the facial nerve in the adult rat. J Neurosur 1999;90:760-5.
12. Bento RF, Miniti A. Comparison between fibrin tissue adhesive, epineural suture and natural union in intratemporal facial nerve of cats. Acta Otolaryngol Suppl 1989;465:1-36.
13. Mattos DE, Felix H. Surgical anatomy of the rat facial nerve. Am J Otol 1987;8(1):43-7.
14. Byers JM, Clark BF, Thompson GC. Effect of pulsed electromagnetic stimulation on facial nerve regeneration. Arch Otolaryngol Head and Neck Surg 1998;124:383-9.
15. Yian CH, Paniello RC, Spector G. Inhibition of motor nerve regeneration in a rabbit facial nerve model. Laryngoscope 2001;111:786-91.
16. Guntinas-Lichius O, Irintchev A, Streppel M, Lenzen M, Grosheva M, Wewetzer K et al. Factors limiting motor recovery after facial nerve transection in the rat: combined structural and functional analyses. Eur J Neurosci 2005;21:391-402.
17. Semba K, Egger D. The facial “motor” nerve of the rat: control of
vibrissal movement and examination of motor and sensory components. J Comp Neurol 1986;247:144-58.

18. Hadlock TA, Heaton J, Cheney M, Mackinnon SE. Functional recovery after facial and sciatic nerve crush injury in the rat. Arch Facial Plast Surg 2005;7:17-20.

19. Terrel GS, Terzis JK. An experimental model to study the blink reflex. J Reconst Microsurg 1994;10(3):175-83.

20. Gilad VH, Tetzlaff G, Rabey JM, Gilad GM. Accelerated recovery following polyamines and aminoguanidine treatment after facial nerve injury in rats. Brain Res 1996;724:141-4.

21. Guntinas-Lichius O, Angelov DN, Tomov TL, Dearn J, Neiss WF, Wewetzer K. Transplantation of olfactory ensheathing cells stimulates the collateral sprouting from axotomized adult rat facial motoneurons. Exp Neurol 2001;172:70-80.

22. Fernandez E, Pallini R, Marchese E, Lauretti L, Marca FL. Quantitative, morphological, and somatotopic nuclear changes after facial nerve regeneration in adult rats: a possible challenge to the “no new neurons” dogma. Neurosurgery 1995;37(3):456-63.

23. Ross BG, Fradet G, Nedzwelski JM. Development of a sensitive clinical facial grading system. Otolaryngol Head Neck Surg 1996;114:880-6.