Effects of low-level helium-neon laser on induced wound healing in rats

Os efeitos do laser hélio-neônio de baixa intensidade na cicatrização de lesões cutâneas induzidas em ratos

Viviane L. Busnardo¹, Maria L. P. Biondo-Simões²

Abstract

Objective: To evaluate the effects of low-level helium-neon (HeNe) laser on cutaneous wound healing in rats. Methods: Sixty Wistar rats were divided into control group and experimental group. A sutured longitudinal, dorsal-medial incision was made, with simple separate stitches. The experimental group was irradiated daily in three areas of the wound with HeNe laser (5mW maximum continuous power, 632.8 nm wavelength, 0.015 cm² laser beam area) for 36 seconds. The areas were evaluated on the third, seventh and fourteenth days postoperative. Histological sections were stained with hematoxylin-eosin to determine the type of inflammatory reaction according to the protocol by Vizzotto et al. (2003)* and with Picrosirius to identify types I and III collagen and the collagen maturation index (CMI). Immunohistochemical detection was employed with anti-CD45-LCA to identify the inflammatory cells. Results: Both groups had the same inflammatory pattern. The experimental group had fewer inflammatory cells at the three evaluation times (p<0.001) with faster reduction in the number of leukocytes. The experimental group had greater total collagen density on the third day (p=0.001), with more type III collagen (p=0.001) and more type I collagen (p=0.001). There was no significant difference in the CMI. Conclusion: Low-level laser irradiation does not change the quality of the inflammatory response, but it does reduce its intensity; it increases collagen deposition in the early stages of the healing process and does not interfere with collagen maturation.

Key words: wound healing; low-level laser therapy; helium-neon laser; collagen.


Resumo

Objetivo: Avaliar os efeitos do laser de baixa potência hélio e neônio (HeNe) na cicatrização de feridas cutâneas de ratos. Métodos: Sessenta ratos Wistar foram divididos em grupos controle e experimento. Utilizou-se ferida incisional, longitudinal, dorso-mediana, suturada com pontos separados simples. No grupo experimento, as feridas foram irradiadas diariamente com aparelho de laser de HeNe com potência contínua máxima de 5mW, comprimento de onda de 632,8 nm, visível com densidade de energia de 4J/cm², área de raio do laser de 0,015 cm², durante 36 segundos, em três pontos da lesão. As feridas foram avaliadas no 3º, no 7º e no 14º dia de pós-operatório. Cortes histológicos foram corados com hematoxilina-eosina (H&E) e avaliados segundo protocolo de Vizzotto et al. (2003)* para identificar o tipo de reação inflamatória e com Picrosirius para identificar os colágenos I e III e o índice de maturidade da cicatriz (IMaC). Utilizou-se imunoistoquímica com anti-CD45-LCA para o reconhecimento das células inflamatórias. Resultados: Ambos os grupos mostraram o mesmo padrão inflamatório. No grupo experimento, observaram-se menos células inflamatórias nos três tempos estudados (p<0,001), com diminuição mais rápida do número de leucócitos. Verificou-se que as do grupo experimento tinham maior densidade de colágeno total no 3º dia (p=0,001), com mais colágeno III (p=0,001) e mais colágeno I (p=0,001). Não houve diferença significativa no IMaC. Conclusão: A irradiação com laser de baixa intensidade não modifica a qualidade da reação inflamatória, mas diminui a intensidade dela; aumenta a deposição do colágeno no início do processo cicatricial e não interfere na maturação da cicatriz.

Palavras-chave: cicatrização de feridas; laserterapia de baixa intensidade; laser de gases Hélio e Neônio; colágeno.


Received: 21/08/2008 – Revised: 10/02/2009 – Accepted: 30/06/2009

¹ Graduate Program in Clinical Surgery, Pontifícia Universidade Católica do Paraná (PUCPR), Curitiba (PR), Brazil
² Department of Surgery, Universidade Federal do Paraná (UFPR), Curitiba (PR), Brazil
Correspondence to: Viviane L. Busnardo, Universidade Positivo, Rua Prof. Pedro Viriato Parigot de Souza, 5.300, CEP 81280-330, Curitiba (PR), Brazil, e-mail: vivianefucci@up.edu.br
Introduction

The process of tissue healing is very complex and involves several biological effects, such as vascular and cell changes, epithelial proliferation, fibroblast proliferation, synthesis and deposition of collagen, production of elastin and proteoglycans, revascularization and wound contraction. The incorporation of laser as a therapeutic tool in the biomedical field has been investigated since 1960 but, in spite of the numerous studies on the effects of laser therapy, it is difficult to justify physical variables such as: application technique, dosages, depth, modes and duration of exposure. It has been observed that photostimulation influences macrophage production of growth factors, which increases cell proliferation. In 1976, Mester et al. reported that low-level helium-neon (HeNe) laser could aid the healing of mechanical injuries. Since then, it was shown that this laser has several effects on live tissue, effects known as laser biostimulation.

Laser photobiomodulation has been increasingly used with the purpose of improving the quality of wound healing. The therapeutic effects of laser on the different biological types are broad and include trophic-regenerative, anti-inflammatory and analgesic effects. It has also been demonstrated that tissue regeneration becomes more effective when treated with low-level laser. There are reports that laser irradiation stimulates the release of fibroblast growth factor (FGF) and the replication of these cells. Irradiation with HeNe laser would accelerate the healing process, with a better weave of collagen fibers and greater collagen deposition, combined with faster reepithelialization and neovascularization.

In situations of deficient healing, such as ischemia, diabetes and pressure ulcers, irradiation with low-level laser could be an alternative for the recovery. The objective of the present study was to evaluate the effects of low-level helium-neon (HeNe) laser on the healing process of skin wounds in rats.

Methods

The presented study was conducted in accordance with Federal Law no. 6638 and the recommendations of the Colégio Brasileiro de Experimentação Animal (COBEA), an entity associated with the International Council for Laboratory Animal Science. This project was approved by the Animal Research Ethics Committee of Universidade Católica do Paraná (PUC-PR), under the protocol no. 181.06/CEUA-PUC-PR.

Sixty Wistar male rats (Rattus norvegicus albinus, Rodentia mammalia) from the vivarium of PUC-PR were used in the study. The mean weight was 397.84 ± 32.74 g and the age ranged from 100 to 150 days. Over the course of the study, the animals remained in the vivarium and were divided into groups of five per box. The room temperature was kept at 22±2°C with a light/dark cycle of 12 hours and ambient relative humidity and noise volume. All of the boxes were placed on shelves equally distant from the source of light. The animals had free access to water and specific food for the species (NUVILAB, NUVITAL®).

The animals were anesthetized with an intramuscular injection (0.1 ml/100g of weight) of ketamine (50 mg) and xylazine 2% (20 mg), the dorsal region was shaved and cleaned with povidone-iodine and the surgical field was delimited with a sterile fenestrated drape. Next, a longitudinal dorsal-medial incision was made through the skin and subcutaneous tissue, starting below an imaginary line that corresponds to the insertion of the front paws. The wound measured approximately four centimeters and it was closed with four simple separate stitches one centimeter apart using monofilament nylon thread 4.0. The laser was applied to the three areas between the stitches (Figure 1). After the surgery, the animals received a single intramuscular injection of diclofenac potassium (10 mg/kg) for analgesic purposes. After the recovery from the anesthesia, the rats were adequately marked and randomly distributed into two distinct groups, with 30 animals each: experimental group (EG) treated with laser and untreated control group (CG).

Each group was subdivided into three subgroups with the same number of animals to perform the time evaluation on postoperative days 3, 7 and 14. These subgroups were denominated: EG day 3, EG day 7 and EG day 14 and CG day 3, CG day 7 and CG day 14. For the treatment, the device laser He-Ne Plasmax IV, LHN 9709 (KLD Biossistemas®) was used. The wounds were treated with HeNe laser at energy density of 4 J/cm² over a 3 cm² area, resulting in a calculated use of the laser of 12 seconds per area of the wound. Thus, the HeNe laser was applied at the maximal continuous energy level of 5 mW, with wavelength of 632.8 nm and laser beam area of 0.015 cm².

Twenty-four hours after the incision procedure, the wounds of the EG animals received treatment with HeNe laser. One subgroup was irradiated for 3 days, another for 7 days and the last for 14 days. For the administration of the laser, the animals were sedated with intraperitoneal propophol (10 mg/Kg). The animals from the CG received the same sedation. After macroscopic analysis on the 3rd, 7th and 14th days, ten animals from each group were selected by draw and euthanized with an intraperitoneal injection of sodium thiopental (120 mg/Kg) in the left iliac fossa.

In the sampled animals, skin segments (5x4cm) containing the wound in the center were resected. About half a centimeter in the cranial and caudal extremities of the segment was rejected, and the three remaining centimeters of the wound were used for the study. The segments were fixed in buffered formalin at 10% for 24 hours and later submitted to routine...
histological procedure. Through the hematoxylin-eosin (H&E) staining, the general morphological evaluation of the wound was obtained, and the inflammatory pattern was recognized. Ten fields were viewed at 400x magnification, according to the guidelines described by Vizzotto et al.²⁶ (Table 1). For the cell count, the following scale was adopted: no cell=0; up to 50 cells=1; 50 to 100 cells=2 and more than 100 cells=3, positive for monomorphonuclear cells and negative for polymorphonuclear cells²⁶. After the attribution of the indices, their total was calculated so that each group of animals had a final score for classification into three phases of the inflammatory process²⁶ (Table 2).

The histological sections stained with Picrosirius (Picrosirius-red F3BA) under microscopy and polarized light allowed the identification of the collagen density in the wounds and the fractions of type I and III collagen. With this technique, the thicker and highly birefringent type I collagen fibers appear orange and red in color, and the finer, dispersed and less birefringent type III collagen fibers appear green in color²⁵. The images were captured on a Sony CCD 101/Trinitron system, digitized by an Oculus TCX capture board and analyzed by the Image Plus 4.5 for Windows application. For each slide, three fields were read at 200X magnification over the wound area, and the mean was calculated.

The collagen maturation index (CMI²⁷) was defined as the ratio of the percentage of type I collagen to the percentage of type III collagen. This index varies from zero (percentage of type I collagen=0 and percentage of type III collagen=100) to infinite (percentage of type I collagen=100 and percentage of type III collagen=0). Values higher than one indicate that the percentage of type I collagen is greater than the percentage of type III collagen and show the state of collagen maturation²⁸.

The tissue segments used for anti-CD45-LCA immunostaining (LCA – Leukocyte Common Antigen, Clone 136-4B5 IV WS - IgG1, FK-Biotec®) at 1:100 dilution were submitted to antigen retrieval in bain-marie in 10 mmol/L citrate buffer (pH 6.0) for further immunohistochemical processing. To guarantee the veracity of the staining, controls were made. For external positive control, a lymph node fragment was used. The dermis and the epithelium were used as internal negative control. For the analysis, a count was taken of the positive cells stained brown for LCA per field of magnification, in a total of ten fields.

The descriptive results obtained from the study were expressed as means and standard deviations. Two-way ANOVA was used to determine the effect of the group and of the evaluation day on the study variables. The normality condition of the variables was assessed by the Shapiro-Wilk test, and the homogeneity of the variances, by Levene’s test. In the cases in which there was a significant interaction between the factors, Student’s t test was used for intragroup analysis and ANOVA.

---

**Table 1.** Methods for quantification of histological findings in sections stained with hematoxylin-eosin (H&E).

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Intensity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acute</td>
</tr>
<tr>
<td>Polymorphonuclear cells</td>
<td>-3</td>
</tr>
<tr>
<td>Edema</td>
<td>-3</td>
</tr>
<tr>
<td>Congestion</td>
<td>-3</td>
</tr>
<tr>
<td>Monomorphonuclear cells</td>
<td>3</td>
</tr>
<tr>
<td>Granulation Tissue</td>
<td>3</td>
</tr>
<tr>
<td>Fibrosis</td>
<td>3</td>
</tr>
</tbody>
</table>

---

**Table 2.** Characteristics of the phases of inflammatory process according to the final score.

<table>
<thead>
<tr>
<th>Inflammatory Process</th>
<th>Final Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute</td>
<td>-9 to -3</td>
</tr>
<tr>
<td>Subacute</td>
<td>-2.9 to 3</td>
</tr>
<tr>
<td>Chronic</td>
<td>3.1 to 9</td>
</tr>
</tbody>
</table>

---

Figure 1. Appearance of the wound at the end of surgery, showing the laser application areas.
was used for within-day analysis. For the multiple comparisons, the Least Significant Difference (LSD) test was used. For the analysis of the CMI tables, Fischer’s test was used. The level of significance adopted was 0.05.

Results

In the histological sections of the wounds on day 3 for both groups, the inflammation showed characteristics of an acute process evidenced by negative scores. For the between-group comparisons, on day 7 the inflammation had changed and had characteristics of a subacute process, and on day 14, it had characteristics of a chronic process evidenced by positive scores (Figures 2 and 3). With regard to the analysis of type I percentage of collagen, there was no significant difference between the groups in the three moments analyzed (p=0.527; Figure 4A). There was higher type III collagen density in the wounds of EG day 3 (p=0.001). However, in the between-group comparisons of day 7 and 14, the differences were not significant (Figure 4B). Furthermore, the between-group comparisons of the total collagen showed higher density in the wounds of EG day 3 (p=0.001), without significant differences in the comparisons of day 7 and 14 (Figure 4C).

Regarding the CMI, greater maturation was found in the wounds of CG day 3 when compared to EG day 3 (p=0.026; Figure 4D). However, in the dichotomized analysis separating the number of wounds with indices up to 1 and those with indices higher than 1, there was no difference in frequency between the groups. A difference between groups was found on day 3 (p=0.474), day 7 (p=1) and day 14 (p=1) (Figure 5). The immunohistochemical process showed a lower count of inflammatory cells for the EG group when compared to the CG group on the three assessment moments (p<0.001), with an earlier decrease in their number in the EG group (Figures 6 and 7).

Discussion

Wound healing can be improved when the main events that enable it are stimulated, e.g. nutrition, cell proliferation and control of inflammation and infection. In several studies with animal models, low-level laser has been identified as an adjuvant for healing because it improves revascularization, generation of energy in form of adenosine triphosphate (ATP) for the repair cells, fibroblast proliferation and inhibition of the chemical mediators of inflammation. Given the main factors involved in the healing process, the present study aimed to identify the influence of laser on the general aspects of this process. The factors for analysis were chosen to observe the resolution of the inflammatory process and the collagen synthesis. The results confirm that the low-level laser with 4 J/cm² energy density promotes type III collagen deposition on postoperative day 3.

Carvalho et al. conducted a morphometric analysis of the percentage of collagen fibers by color density of the wounds of 48 rats. The authors used HeNe laser (4 J/cm²) for 36 seconds in the EG and assessed the results on postoperative days 3, 7 and 14 and found a significant increase in the percentage of collagen from day 3 to day 7. When day 14 was compared to day 7, a deceleration in the percentage of collagen was observed. The authors attributed this deceleration to the state of maturation of these wounds because of reports of decrease in cell proliferation and fibroblast number and size in the remodeling/maturation.

Pugliese et al. studied the effects of low-level laser (GaAlAs laser at different energy densities) on the standardized skin wounds of 62 Wistar rats. In the morphometric analysis, the authors observed a greater expression of the collagen fibers, however, without statistical significance. The best results were found in the groups irradiated with 4 J/cm². In the evaluation of the inflammatory response, the irradiated wounds had an earlier reduction in edema, congestion and inflammatory cells.
Regarding the improvement in healing, the authors attributed to the laser a significant increase in the collagen deposition that can happen by induction of the cell proliferation or even by an increase in protein synthesis and release, with the added possibility of both processes occurring simultaneously.

Araújo et al.\textsuperscript{19} found more activated fibroblasts, more collagen and more incorporation of 3H-proline in the wounds of rats treated with low-level laser and evaluated on days 8, 15 and 22 post-wounding. Carvalho et al.\textsuperscript{21} reported that this effect can be observed in the wounds of diabetic rats, showing that low-level laser therapy could be useful in adverse situations. Medrado et al.\textsuperscript{20}, however, observed a slight increase in collagen density.

Figure 4. Mean percentages of the areas of the histological sections examined and represented by collagen in the two groups at each evaluation and the collagen maturation index.

Figure 5. Photomicrograph of the histological sections of the wound areas of the EG and CG, stained with Sirius Red under a polarized light, (200x magnification) on the third-day evaluation.

Figure 6. Photomicrographs of histological sections showing the leukocytes marked by anti-CD45 (*) on the third-day evaluation (400x magnification).

Figure 7. Mean number of inflammatory cells (per site) marked with anti-CD45 in ten sites.
Collagen maturation was evaluated using the CMI. The Picrosirius-stained sections allowed the assessment of the type of collagen and to quantify each type in the samples. Several authors have compared different staining methods for collagen analysis and concluded that the Picrosirius pigment is the most selective and the easiest to use and interpret. It is also specific for the study of tissue collagen because the amount of adhered pigment is proportional to the amount of protein, allowing the quantification of the protein.

Despite a higher density of type III collagen on day 3, the results of the CMI in the dichotomized analysis showed that the collagen maturation was similar throughout the duration of the study. In fact, the type I collagen density tended to be greater in the EG, but it was not statistically significant. A larger sample could clarify this question by confirming or rejecting this tendency. The same can be stated about the wounds of the EG, which had a tendency for higher but non-significant maturation indices on assessment days 7 and 14. On day 14, a deceleration in the healing process was observed, a finding reported in a previous study and that may be explained by the decrease in the proliferation of cellular elements in this phase.

Although many clinical studies have been favorable to the use of conventional laser, the mechanism by which it accelerates the process of healing has not been fully elucidated. There are still some hypotheses to be tested, e.g. whether laser enhances the transcription of ribonucleic acid (RNA) while reducing the inflammatory phase and anticipating fibroplasia, and whether the laser-induced increase in oxygen and ATP is determined by an early angiogenesis.

To ensure the effects of low-level laser therapy on the healing process, there is still a need for consensus on the standards for the physical variables: application times and techniques, energy densities, output powers and wavelengths. The comparison of the results of several authors has been hindered by their use of different methodologies. The use of HeNe laser (632.8nm), applied with different densities can lead to different cellular responses, and this may preclude comparisons. Hawkins and Abrahamse showed that the use of low-level HeNe laser stimulated interleukin-6 expression of the interleukin-6, cell proliferation and migration in diabetic subjects, thus improving the healing process. Silveira, Streck and Pinho demonstrated that low-level laser can activate mitochondrial enzymes, such as cytochrome c oxidase, and improve wound healing.

Irradiation with low-level HeNe laser in the conditions of this experiment did not change the quality of the inflammatory response, but decreased its intensity. Collagen deposition increased at the beginning of the healing process and the maturation of the wound was not affected.

Conclusion

Irradiation with low-level HeNe laser in the conditions of this experiment did not change the quality of the inflammatory response, but decreased its intensity. Collagen deposition increased at the beginning of the healing process and the maturation of the wound was not affected.
Effects of low-level HeNe laser on wound healing

References


