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The effects of low-level light emitting diode on the repair process of Achilles tendon therapy in rats

Casalechi · Nicolau · Casalechi · Silveira · Paula · Pacheco

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56	Abstract	<p>Thirty Wistar rats (350 ± 20 g) were subjected to total Achilles tendon tenotomy of the right fore limb. They were submitted to a daily dose of 20 J/cm² light emitting diode (LED) (640 ± 20 nm) therapy. The LED was applied punctually and transcutaneously to the lesioned region. The animals were separated into six groups, C1 and L1, C2 and L2, C3 and L3. The C groups were used for control and the L groups, treated for 7, 14 and 21 consecutive days, respectively. The animals were killed on the 7th, 14th and 21st days after surgery. After the animals had been killed, their tendons were extracted and dissected, fixed in formaldehyde at 10%, and sent for histological analysis by light microscopy in which the repair process was analysed. This study demonstrated that LED interfered in the repair process of the tendon tissue, reducing the number of fibroblasts in the initial periods and improving the quality of the repair in all periods studied.</p>	
57	Keywords separated by ' - '	Achilles tendon - Light emitting diode (LED) - Low-level laser therapy - Tissue repair	
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3 BRIEF REPORT

4 The effects of low-level light emitting diode on the repair 5 process of Achilles tendon therapy in rats

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13 **Abstract** Thirty Wistar rats (350 ± 20 g) were subjected to
14 total Achilles tendon tenotomy of the right fore limb. They
15 were submitted to a daily dose of 20 J/cm^2 light emitting
16 diode (LED) (640 ± 20 nm) therapy. The LED was applied
17 punctually and transcutaneously to the lesioned region. The
18 animals were separated into six groups, C1 and L1, C2 and
19 L2, C3 and L3. The C groups were used for control and the
20 L groups, treated for 7, 14 and 21 consecutive days,
21 respectively. The animals were killed on the 7th, 14th and
22 21st days after surgery. After the animals had been killed,
23 their tendons were extracted and dissected, fixed in
24 formaldehyde at 10%, and sent for histological analysis
25 by light microscopy in which the repair process was
26 analysed. This study demonstrated that LED interfered in
27 the repair process of the tendon tissue, reducing the number
28 of fibroblasts in the initial periods and improving the
29 quality of the repair in all periods studied.

30 **Keywords** Achilles tendon · Light emitting diode (LED) ·
31 Low-level laser therapy · Tissue repair

Introduction

Many scientific works have been published about tendon
lesions with the objective of optimizing tissue repair.
According to Goffi [1], surgeons worry about tendon
lesions in reconstructive surgeries, because, when there is
a suture, there may be adherence which hinders its full
recovery. Schmitt called attention to the use of physiother-
apeutic resources in speeding the tissue repair process in 1993
[2] in a study that showed the effects of laser radiation on
the regeneration of tendons in dogs. Nowadays, researchers
are still looking for the best way to restore the functions and
improve the repair in tendons. Most studies suggest that
coherent light (low-power laser) can start the modulation of
physiological processes.

The high cost of laser-emitting devices stimulates
research on the effects of alternative light sources such as
light emitting diode (LED). According to Soley et al. [3]
and Clark and colleagues [4], irradiation with non-coherent
light has a lower cost and can be as efficient as laser
radiation. Vinck et al. [5] obtained satisfactory results in
their study and suggest beneficial effects with LEDs on
different kinds of skin lesions.

The stimulant effects produced by low-power laser on
biological tissues were attributed to their coherence by
Boulton and Marshall [6]. The light transmitted by an LED,
unlike that from laser, is non-coherent. However, in more
recent studies, Pontinen [7] and Whelan et al. [8] state that
the light coherence is not responsible for the therapeutic
effects of low-power laser, because this property is lost in
the first layers of biological tissue. The growth of cellular
activity, both in division and synthesis, has been related to
the length and dosage and not specifically to the light
source [9]. Nowadays, LEDs are being commercially
introduced as alternatives to low-power laser therapies. It

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66 is known that the LED activity has an influence on the
67 regenerating process of wounds [5].

68 The aim of the study was to investigate the effect of
69 LED (640±20 nm) therapy on tissue regeneration of the
70 Achilles tendon of rats through the quantitative evaluation
71 of the number of fibroblasts and the quality of tissue repair.
72 This study would allow evaluation of the effect of LED
73 therapy on the tissue repair process of the tendon, aiming at
74 a low-cost alternative to low-level laser therapy (LLLT) in
75 clinical treatments.

76 **Materials and methods**

77 Experimental groups

78 Thirty albino male Wistar rats (approximately 350±20 g,
79 3 months old) were used in this study. They were kept in
80 the Research and Development Institute of the University
81 of Vale do Paraíba in appropriate standard polyethylene
82 cages, in random groups of five animals per cage. They
83 underwent a period of adaptation for 5 days in a room with
84 constant temperature and humidity (24°C e 60%), natural
85 light, water and food ad libitum.

86 The 30 animals were randomly separated into six groups,
87 C1, L1, C2, L2, C3 and L3. Table 1 shows the groups and
88 the respective days on which they were killed.

89 Surgical procedure

90 A total tenotomy of the medium region of the right Achilles
91 tendon, between the tendon insertion and the myotendon
92 articulation, was made in all animals. For this procedure all
93 animals were subcutaneously pre-treated with atropine
94 (analgesic); the dosage was 0.04 ml/100 g body weight.
95 After this, there was an interval of 15 min before the
96 anaesthetic procedure [11]. The anaesthetic drug was given
97 in a single intramuscular injection of cetamine chlori-
98 drate 10%, 10 ml (Syntec, 0.1 ml/100 g body weight) and
99 xylazine chloridrate 2%-10 ml (Syntec, 0.1 ml/100 g) [12],
100 injected with a 1 ml insulin syringe for each animal. The
101 skin of the right limb was shaved and scrubbed with a 2%

t1.1 **Table 1** Number of animals, groups, treatments and days on which they were killed

t1.2	Group (n=5)	Therapy	Day killed
t1.3	C1	Control	7th day after surgery
t1.4	L1	LED	
t1.5	C2	Control	14th day after surgery
t1.6	L2	LED	
t1.7	C3	Control	21th day after surgery
t1.8	L3	LED	

iodine alcohol solution. The tendon was exposed through a 102
1 cm incision and transversally tenotomized in the medial 103
region. The skin was closed with 6.0 polyester monofila- 104
ment (Prolene®) and disinfected with 2% iodine alcohol 105
[11, 12]. 106

After surgery the animals received a single intramuscular 107
injection of broad-spectrum antibiotics (Fort Dodge®, 108
0.02 ml/100 g body weight). 109

LED therapy 110

The equipment used in the study was an LED (640± 111
20 nm), Red Star LED® model (100 mW) with a 0.5 cm² 112
area in direct contact with the right limb of the animal, on 113
the lesion area. The application time was 120 s, with a final 114
dose of 20 J/cm² at one point on the injured area. Before 115
the beginning of the experiments, the LED equipment was 116
checked with a power checker (13PEM001/J, Mellers 117
Griot, Netherlands). 118

The therapeutic procedure was begun 1 h after the 119
surgery and was repeated every 24 h. All animals were 120
treated the same way. For the procedure, the animals were 121
positioned on a table in ventral decubitus, and manually 122
immobilized. The LED was used on their hind limbs, 123
directly on the injury, at a 90° angle. The LED pen was 124
protected by plastic film after each application. 125

Experiment 126

The animals were killed on the 7th, 14th and 21st days after 127
surgery in groups C1 and L1, C2 and L2, C3 and L3, 128
respectively. Before the rats were killed, the same sedation 129
procedure was used as in the surgery, after which an intra- 130
cardiac injection of sodium thiopental (Cristalia) (1 ml/ 131
100 g body weight) was given. 132

The tendons were removed by dissection, from the 133
calcaneal insertion to the myotendon articulation. They 134
were fixed in 10% formaldehyde and sent for histological 135
processing. 136

Histological technique 137

After fixation, the tendons were dehydrated and embedded 138
in paraffin, followed by microtomy in a semi-automatic 139
revolving microtome to produce sections 5 µm thick, eight 140
sections per animal. Four sections were stained with 141
haematoxylin and eosin (HE) and four with Masson 142
trichrome. 143

Morphometry 144

A Nikon® optical binocular microscope, model YS100, 145
was used for the morphometry, with a Zeiss® ocular with 146

147 millimetre reticulated integrator. The analysis was done in
 148 12 microscopic fields, equivalent to a 150 μm^2 area, with
 149 histological sections dyed with HE and Masson trichrome.
 150 The area chosen for analysis in all samples was the most
 151 proximal to the edges of the tenotomized regenerating
 152 tissue. The numbers of fibroblasts and gradation of the
 153 repair were evaluated in the tissue of this area.

154 The tissue regeneration in the tenotomized area was
 155 graded as follows:

- 156 • Group ⁺. Absence of regeneration. Evidence of hyper-
 157 cellularity associated with the presence of delicate and
 158 rare collagen fibrils with no specific orientation in
 159 relation to the edges of the tenotomized area. Abundant
 160 amorphous fundamental substance. Diffuse infiltrate of
 161 chronic inflammatory and phagocytic cells.
- 162 • Group ⁺⁺. Initial regeneration. Evidence of hyper-
 163 cellularity associated with the presence of delicate and
 164 thick collagen fibres, already presenting indications of
 165 orientation (angulation of up to 25°) in relation to the
 166 the edges of the tenotomized area. Decrease in the
 167 presence amorphous fundamental substance. Chronic
 168 inflammatory and phagocytic cells.
- 169 • Group ⁺⁺⁺. Intermediate regeneration. Cellularity near
 170 normal, associated with mature collagen fibres, posi-
 171 tioned in orientated clusters in relation to the edges of
 172 the tenotomized area (25° to 10° angle). Occasional
 173 presence of inflammatory cells and absence of phago-
 174 cytic cells.
- 175 • Group ⁺⁺⁺⁺. Complete regeneration. Normal cellularity
 176 associated with clusters of mature collagen of fibres in
 177 parallel orientation (180°) in relation to the edges of the
 178 tenotomized. Absence of inflammatory cells and ab-
 179 sence of phagocytic cells.

180 **Statistical analysis**

181 We evaluated the data for coefficient variance and sample
 182 distribution to determine the statistical test, considering a
 183 statistical significance level of 5% ($P < 0.05$) [13]. The
 184 numbers of fibroblasts in the treated and untreated groups
 185 were subjected to analysis of variance (ANOVA), with the
 186 Bonferroni post-test. The level of significance was 5% ($P <$
 187 0.05). The program used was GraphPad Prism[®], version
 188 2.0

189 **Results**

190 **Qualitative analyses of tissue repair**

191 In the histopathological analyses of group C1 the most
 192 observed gradation of remodelled collagen fibres was in the

193 absence of the regeneration phase and, in rare cases, in the
 194 initial regeneration phase. The C1 group presented hyper-
 195 cellularity associated with the presence of delicate and rare
 196 collagen fibrils with no specific orientation in relation to the
 197 edge of the wound. There was evidence of an amorphous
 198 fundamental substance and a diffuse infiltrate of chronic
 199 and phagocytic inflammatory cells (Fig. 1a).

200 Group L1 presented a remodelled gradation of the
 201 collagen fibres, especially in the initial remodelling.
 202 Evidence of hypercellularity was observed, in association
 203 with delicate and sparse collagen fibres, but with signs of
 204 orientation in relation to the edge of the wound. The
 205 presence of an amorphous fundamental substance and a
 206 diffuse infiltrate of chronic and phagocytic inflammatory
 207 cells diminished (Fig. 1b).

208 In group C2 the gradation of collagen fibre remodelling
 209 was mostly observed in the initial remodelling phase.
 210 Evidence of hypercellularity was observed, in association
 211 with delicate and rare collagen fibres, with signs of
 212 orientation in relation to the edge of the wound (Fig. 2a).

213 The frequently observed gradation of remodelled colla-
 214 gen fibres in group L2 was in the intermediate remodelling
 215 phase. The group presented near normal cellularity, with

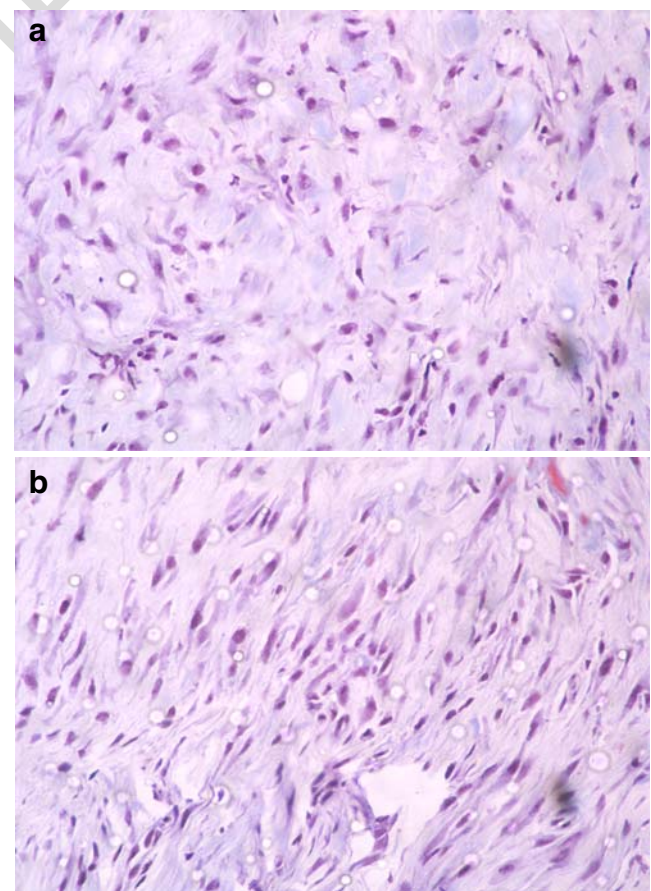


Fig. 1 Microscopy of calcaneal tendon of rats killed on the 7th day after surgery. **a** Group C1; **b** group L1. Masson trichrome, $\times 400$

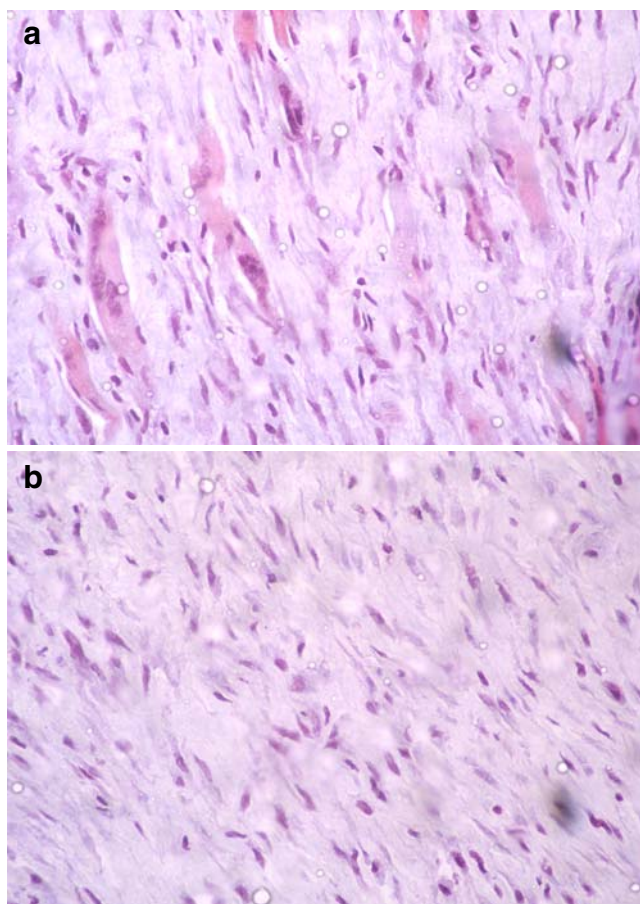


Fig. 2 Microscopy of calcaneal tendon of rats killed on the 14th day after surgery. **a** Group C2; **b** group L2. Masson trichrome, $\times 400$

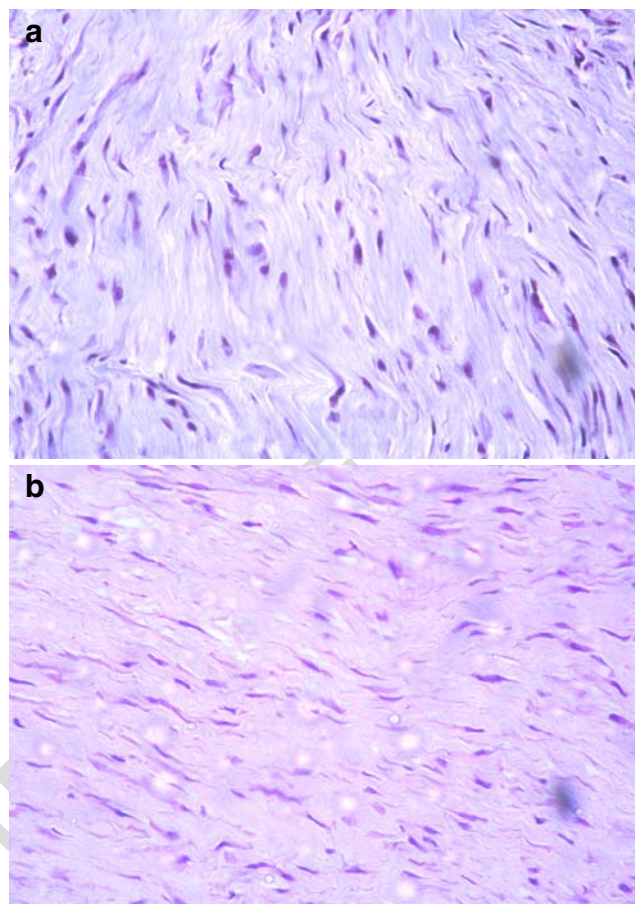


Fig. 3 Microscopy of treated calcaneal tendon of rats killed on the 21st day after surgery. **a** Group C3; **b** group L3. Masson trichrome, $\times 400$

216 mature collagen fibres disposed in clusters aligned with the
217 edge of the wound. Occasional chronic inflammatory cells
218 were observed (Fig. 2b).

219 Group C3 presented a remodeling gradation of collagen
220 fibers especially in the intermediate remodeling phase and in
221 rare cases in the complete remodeling phase. Cellularity near
222 normal was observed with the presence of clusters of mature
223 collagen fibers aligned with the edge of lesion area (Fig. 3a).

224 In group L3 the gradation of the collagen fibers presented
225 a complete remodelling phase. Evidence of normal cellular-
226 ity was observed, in association with clusters of mature
227 collagen fibers aligned with the edge of lesion area (Fig. 3b).

228 Figure 4 shows the comparison of results of the
229 gradation of tendon tissue remodelling between the control
230 and LED groups in the three different phases observed.

231 Quantitative analysis

232 ANOVA was performed between groups, and all groups
233 (control and treated) presented significant statistical differ-
234 ences in relation to the numbers of fibroblasts with the
235 passage of time: C1 and C2 ($P < 0.01$), C1 and C3 ($P <$

0.001), L1 and L3 ($P < 0.05$). In inter-group evaluations a
236 significant statistical difference was observed between
237 groups C1 and L1 ($P < 0.01$). In Table 2 the numbers of
238 fibroblasts in the control and LED animals at the 7th, 14th
239

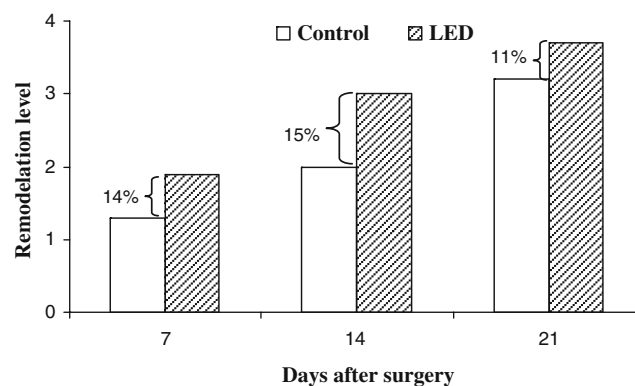


Fig. 4 Comparison of the intensity of remodelling between the control and LED groups in the three different phases observed. The numbers 1, 2, 3 and 4 represent tissue regeneration grades +, ++, +++ and +++, respectively. The difference between the control and LED is shown in percentages

t2.1 **Table 2** Means, standard deviations, and ANOVA results of the numbers of fibroblasts found on the 7th, 14th and 21st days in the control and LED groups

t2.3	Day	Control	LED
t2.4	7th day	201.5±6.3 *	167.6±5.3***
t2.5	14th day	166.5±11.5	160.1±1.4
t2.6	21 st day	161.6±1.2 **	138.8±1.4 **

t2.7 $P < 0.05 = *$ (7th day vs 14th day), $**$ (7th day vs 21 st day), $***$ (control vs LED)

240 and 21st days are shown. The lowest number of fibroblasts
 241 was found on the 21st day, with the lowest mean value in the
 242 LED group (L3). On the 14th day intermediate values were
 243 observed; the lowest mean value was in the LED group (L2).
 244 The group killed on the 7th day presented the highest
 245 number of fibroblasts; the lowest mean value was in the LED
 246 group (L1).

247 **Discussion**

248 This study was the result of the limited research into the use
 249 of LEDs in tendon healing and the need for alternative
 250 therapies to help in the process. It considered the success of
 251 photobiomodulation in tendon wounds with light sources,
 252 coherent or not, demonstrated by previous studies [11].
 253 However, this is a controversial area, because of conflicting
 254 results, justifying the need to identify the best parameters to
 255 be used in clinical practice [14–18].

256 Studies have demonstrated the effects of LED irradiation
 257 similar to the ones obtained with laser on cells [5].
 258 However, the differences between the in vitro and in vivo
 259 systems must be heeded, for the presence of the circulatory
 260 and lymphatic systems, among other aspects, can lead to
 261 modifications in the light–cell interaction. For this reason,
 262 we carried out this study in vivo, using young male rats to
 263 avoid interferences such as menopause and age [19]. The
 264 calcaneal tendon, known as the Achilles tendon, was
 265 selected for its easy access, because it is near the skin and
 266 is large, which makes it easy to undergo surgical techniques
 267 [11, 12]. The irradiation followed the recommended
 268 procedure. It was punctual, with a beam covering the
 269 whole injured area. The pen was positioned at 90° to the
 270 tendon’s longitudinal axis, in contact with the animal’s skin
 271 (transcutaneously). According to these authors, these con-
 272 ditions allow the deposited energy to penetrate the tissue
 273 with less loss by specular reflection [12, 20].

274 The tendon tissue has poor regenerative capacity,
 275 because of its scarce blood circulation, oxygenation and
 276 nutrition, which are very important in tissue repair, for the

adequate evolution of regeneration [21, 22]. Thus, the use
 of non-coherent light as an auxiliary in tendon repair was a
 relevant factor in this study because this kind of light
 source is similar to coherent light sources such as the laser,
 but at a lower cost [3, 4, 7, 9, 23].

Tissue remodelling demonstrated that there was superior
 healing in the groups treated with LED, resulting in a more
 mature tissue than in the other groups in this experiment.
 Our results agreed with those obtained by Faria [11] in a
 similar study with LEDs (4 J/cm²).

The average obtained in the tissue remodelling analysis
 in the group treated with the LED therapy, on the 7th day
 after surgery, was very similar to the that obtained in the
 control group on the 14th day after surgery. On the other
 hand, the group treated with the LED therapy, on the 14th
 day after surgery presented an average similar to that of the
 control group on the 21st day after surgery. In the groups on
 the 21st day after surgery the average of the LED group
 almost reached the maximum remodelling previously
 established, with a higher quality than the average of the
 control group. The results suggested that the remodelling
 quality of the groups treated with LEDs might have been
 achieved more quickly than in the groups that did not
 undergo the treatment, confirming previous studies that
 stated that LLLT would optimize tissue healing in tendons
 [11, 12, 16, 24, 25].

An acceleration in the formation of fibrils, collagen
 fibres, simulating regeneration of the muscle tissue and a
 great decrease in the inflammatory response in tenotomized
 tendons treated with LLLT [26, 27] improves the healing
 quality of the tissue.

The repair process in tendon tissue happens in three
 overlapping phases: inflammatory, proliferative and remod-
 elling [1, 21, 28, 29]. Our study demonstrated the presence
 of a larger number of fibroblasts in both groups killed on
 the 7th day after surgery (control and treated). It is known
 that until the 7th day an injury is in the inflammatory phase
 in which extrinsic regeneration happens and there is a
 proliferation of blood capillaries and fibroblasts that
 migrate to the injured area [21, 28, 30]. In the proliferation
 and remodelling phases intrinsic regeneration occurs in
 which there is fibroblast synthesis to produce collagen [1,
 29, 31], which could explain the gradually diminishing
 number of fibroblasts in the groups killed on the 14th and
 21st days after surgery.

Some specific studies have found that LLLT can
 stimulate or inhibit the various regenerating phases accord-
 ing to the dosage. The objective of our study was also to
 see whether this affirmation is also applicable to LED
 therapy. This could not be confirmed, because our results
 demonstrated that the LED therapy produced an accelera-
 tion of the healing process after total tenotomy. This was
 the expected effect of the electromagnetic radiation in the

330 red region, diffused inside the injured tissues [23]. Among
 331 the groups studied the biggest difference found in relation
 332 to the number of fibroblasts appeared when we compared
 333 group C1 with group L1, where the rats had been killed on
 334 the 7th day after surgery. The group stimulated with LED
 335 showed significant diminishing in the number of fibro-
 336 blasts, which disagrees with some authors who described an
 337 increase as an effect of LLLT [5, 11, 32]. Important factors
 338 are mentioned as possible causes for this increase in the
 339 number of fibroblasts in the initial tissue repair phase,
 340 among which the dosage stands out. Probably, the results
 341 obtained were related to the high dosage used in our study
 342 (20 J/cm²), pointing to new research in this direction.

343 Houreld and Abrahamse [33] observed the stimulatory
 344 effect of LLLT on cells that had received a 5 J/cm² dosage;
 345 meanwhile, cells that had received a 16 J/cm² dosage
 346 presented an inhibitory effect on the proliferation and
 347 fibroblast activity. Likewise, the results found in our work
 348 showed a significant increase in the numbers of fibroblasts
 349 after the rats had been irradiated with a 20 J/cm² dosage.
 350 This significant difference in the numbers of fibroblasts was
 351 only present in the control groups and the group treated in
 352 the initial phase of regeneration (C1 and L1), which might
 353 suggest little influence of the therapy in later phases. This
 354 would eliminate the need for LED treatment after 7 days.
 355 This proliferative aspect might be considered negative if the
 356 remodelling had not been superior in the irradiated groups
 357 in relation to the control.

358 It was observed that the effect of the LED therapy in the
 359 regenerating process of rats' Achilles tendons optimized the
 360 remodelling quality of the tendon tissue in all groups
 361 treated, presenting a great difference in all groups treated.
 362 There was a great difference in relation to the control
 363 groups and a significant difference between the days on
 364 which the rats had been killed, with a gradual decrease in
 365 the numbers of fibroblasts in both groups and between the
 366 groups killed on the 7th day after surgery.

367 To summarize, with the use of LED therapy in the
 368 parameters tested, the quality of the regenerated tissue was
 369 superior to that in the control group in all experimental
 370 phases, with significantly lower cellularity on the 7 days
 371 for the irradiated group. This information points to an
 372 increase in activity in collagen synthesis, probably related
 373 to the reduction of the inflammatory process in the first
 374 moments of regeneration. This would attest to the
 375 superiority of the quality of remodelling in the treated
 376 groups.

377 **References**

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