The earthworm *Dendrobaena veneta* (Annelida): A new experimental-organism for photobiomodulation and wound healing

Andrea Amaroli,1,2 Sara Ferrando,3 Marina Pozzolini,4 Lorenzo Gallus,2 Steven Parker,1 Stefano Benedicenti1

1Department of Surgical Sciences and Integrated Diagnostic, University of Genoa
2Department of Earth, Environmental and Life Sciences, Laboratory of New Model Organism (NeMoLAB), University of Genoa
3Department of Earth, Environmental and Life Sciences, Laboratory of Marine Biotechnology, University of Genoa, Italy

**Abstract**

Photobiomodulation (PBM) is a manipulation of cellular behavior using non-ablative low intensity light sources. This manipulation triggers a cascade of metabolic effects and physiological changes resulting in improved tissue repair, of benefit in the treatment of tissue injury, degenerative or autoimmune diseases. PBM has witnessed an exponential increase in both clinical instrument technology and applications. It is therefore of benefit to find reliable experimental models to test the burgeoning laser technology for medical applications. In our work, we proposed the earthworm *Dendrobaena veneta* for the study of non-ablative laser-light effects on wound healing. In our preliminary work, *D. veneta* has been shown to be positively affected by PBM. New tests using *D. veneta* were set up to evaluate the effectiveness of a chosen 808 nm-64 J/cm²–1W-CW laser therapy using the AB2799 hand-piece with flat-top bean profile, on the wound healing process of the earthworm. Effective outcome was assimilated through examining the macroscopic, histological, and molecular changes on the irradiated posterior-segment of excised-earthworms with respect to controls. Three successive treatments, one every 24 hours, were concluded as sufficient to promote the wound healing, by effects on muscular and blood vessel contraction, decrement of bacteria load, reduction of inflammatory processes and tissue degeneration. *D. veneta* was demonstrated to be a reliable experimental organism that meets well the 3Rs principles and the National Science Foundation statement. Through their genetic and evolutionary peculiarity, comparable to those of scientifically accredited models, *D. veneta* allows the effect of laser therapies by multidisciplinary methods, at various degree of complexity and costs to be investigated.

**Introduction**

A few years after the first working laser was invented, Endre Mester and co-workers,1 while applying low-level laser energy irradiation to the backs of shaved mice, noticed that the shaved hair grew back more quickly on the treated group than the untreated group. This manipulation of cellular behavior using low intensity light sources (non-ablative) was referred to as photobiomodulation (PBM). Conceptually, PBM and the application of photonic energy of specific individual wavelengths works on the principle of inducing a biological response through energy delivery and transfer into the tissue, modulating biological processes within that tissue and within the biological system of which that tissue is a part.2 Photonic energy incorporated within a molecular structure, may increase kinetic energy, activate or deactivate enzymes or alter physical or chemical properties of main macromolecules. Growth factor response within cells and tissue as a result of increased cellular energy and protein synthesis, together with a change in cell membrane permeability to Ca²⁺ uptake, may lead to cell proliferation.3–5 This may form part of a cascade of metabolic effects and physiological changes resulting in improved tissue repair associated with injuries, degenerative diseases or autoimmune diseases.6 However, a lot of confusion still reigns in this field since to date, the correct laser parameters and resultant applied energy density that effectively promote cell rescue without significant side effects, are still elusive. Limited evidence suggests that the energy can be applied with the same efficiency from cellular to organismic level.7 Indeed, if the photonic energy can have a photobiomodulatory positive effect on *in vitro* cells and tissues, that energy must reach target cell at the appropriate intensity to be effective *in vivo*. It is possible that this may be due to incomplete knowledge surrounding the biochemical mechanisms underlying the PBM positive effect. Additionally, there is the complex influence of choice among a large number of operating parameters such as wavelength of the applied electromagnetic energy, the absorbing chromophores within the target tissue, the energy density or fluence of the applied electromagnetic energy and the power density or irradiance of the applied electromagnetic energy.4,8 In addition, the application of PBM using “standard” (hitherto conventional) hand-piece is determined by its very features. Through beam divergence, the applied energy is distributed over an increasing area as the tip-to-tissue distance increases, dramatically affecting energy density at cellular level.8 Laser energy emitted from a standard hand-piece is not evenly distributed across its emitting surface. In almost all instances it has a Gaussian profile, so the density of the photons is significantly higher at the center of the beam and lower at its outskirts.8–9 In order to overcome these problems, PBM has witnessed an exponential increase in both clinical instrument technology and applications. It is therefore of benefit to find reliable experimental models to test the burgeoning laser technology for medical applications. Recently, a novel hand-piece (AB2799) with flat-top bean profile has been produced (LAMBDA Spa, Vicenza, Italy). This particular probe provides homogeneous irradiation and energy density, its beam is non-divergent and provides the same power density over any tip-to-tissue distance from contact to 100 cm (see product characteristics on http://www.lambdaspa.com/en/). In our previous papers,9,10,11 we...
proposed the unicellular eukaryotic organism *Paramecium primaurelia* as a model to test novel PBM therapies and laser technologies. Through these investigations with *P. primaurelia*, we showed that the higher fluence (64 J/cm²) using higher-power (1 W), continuous wave (CW) irradiation, with the AB2799, is able to induce PBM, affecting mitochondrial activity, increasing oxygen consumption, and ATP production, as well as modulating Ca²⁺ fluxes, Ca²⁺ stored release, nitric oxide production and cilia beating. In addition, the higher-fluence higher-power irradiation by flat-top hand-piece of gametes and embryos of the sea urchin *Paracentrotus lividus*, does not affect the DNA and does not generate aberration. In the current work, we moved from the unicellular to the multicellular level and we propose an animal experimental organism to test the photobiomodulatory effect of PBM on wound healing. An attempt to find more ethical and more human closed model organism, which meets well the 3Rs (Replacement, Reduction, Refinement) principles, has emerged through recent decades. The words “model organism”, “model species” or “model animal” are more and more used in different science fields. Some of these models have consequently been promoted for “human health”, while others have been ignored but without real motivation. The accredited models for the study of inflammatory process, wound healing and regeneration are both Vertebrates (*Mus musculus*, *Rattus norvegicus*, *Danio rerio*, *Xenopus laevis*) and Invertebrates (*Caenorhabditis elegans*, *Drosophila melanogaster*). However, the responses to inflammatory stress in the mouse model are not so easily correlated to that of humans. In addition, as reviewed by Katz, “Thomas Hunt Morgan, the father of *Drosophila* genetics, never once referred to *Drosophila* as a model organism in his iconic books. He cited the advantages of choosing the fly *Drosophila* as an experimental organism, but he never used the word “model”. After all, he was not studying a model of the mechanisms of heredity; he was studying the actual mechanism of heredity. Sydney Brenner did not refer to the worm *Caenorhabditis elegans* as a model organism in his classic work on its genetics”. Basically, in a scientific world where the future of medicine is the “Personalized Medicine”, the concept of “model for human health” is actually being questioned. Therefore, according to Katz in our work we proposed *D. veneta* as an “experimental organism” for the study of non-ablative laser light effects on the wound healing.

Earthworms, as annelids, are segmented worms and belong to the group of Clitellata. When mature, they have a visible set of swollen segments, the clitellum, specialized for reproduction. The anterior part of the body, from the mouth to the clitellum (in *D. veneta* roughly 30 segments), contains several different organs (e.g., ovaries, testes, multicellular glands, cerebral ganglia, nephridia, specialized regions of the alimentary canal such as pharynx, crop, and gizzard). Posteriorly to the clitellum, there are several tenths of segments that are anatomically more homogeneous, excluding the few, very caudal ones. In our preliminary work, *D. veneta* showed to be promising model, which was positively affected by PBM. In the present work, we focalized our attention to the wound healing in the posterior part of *D. veneta*, where all the segments share the following anatomy. The body wall is a muscular structure covered externally by a cuticle and epidermis and internally by the peritoneum. The anatomical structures in the coelomic cavity, running along the body length, are the gut, enveloped in a chloragogen layer, the major dorsal and ventral blood vessels, and the ventral nerve cord. Furthermore, a couple of nephridia are present laterally in each segment. In the fluid that fills the coelomic cavity are present free coelomocytes; they are a heterogeneous population of cells. The main cell types are amebocytes, deriving from the peritoneum, and eleocytes, deriving from the chloragogen layer. The body-wall muscle is formed by two layers of helical muscle fibers; they are elongated mononucleated cells, with spiral-ly-arranged sarcomeres and well developed, tubular, sarcoplasmic reticulum. When the body of the earthworm is transversally sectioned both the gut eversion and the formation of a coelomocytes-formed coelomic plug can be involved in wound closure. To reach our goal, we set up new tests on *D. veneta* to evaluate the effectiveness of the 808 nm-64 J/cm²-1 W laser therapy irradiated with the AB2799 hand-piece, on the wound healing process of that earthworm, by examining the macroscopic, histological, and molecular changes on the irradiated posterior-segment of excised-earthworms with respect to controls.

**Materials and Methods**

**Dendrobaena veneta** (Phylum: Annelida, Class: Oligochaeta)

*D. veneta* (also sometimes called *Eisenia hortensis*) is a hermaphroditic robust earthworm that grows very rapidly and may tolerate much wider moisture ranges than many other species; it prefers temperature in the range of 15-25°C. Its life cycle can be completed in 100-150 days and 65 days is the average time to reach sexual maturity.

*D. veneta* can be easily bred in laboratory, in standard conditions, without addition of animal manure, by organic soil and vegetable or oatmeal as food.

**Experimental setting**

*D. veneta* were obtained from Redbug S.r.l (Milan, Italy), acclimatized in laboratory into organic soil at a temperature of 22±1°C and oatmeal fed. The experimental setting was performed as summarized in Table 1. Adult clitelled selected specimens with a similar length (~150 segments), weight (~1.5 g) and color were rinsed with water and anesthetized on an ice block for a few minutes. Under sterile laminar flow hood, 30 posterior segments were excised from the animals and non-irradiated (control) or irradiated with a 808 nm diode (Doctor Smile, LAMBDA Spa) using a hand-piece with flat-top beam profile (AB2799, Doctor Smile, LAMBDa Spa), with a fluence of 64 J/cm² and a power of 1 W in CW. Laser irradiation was performed with the hand-piece at 1 mm away from the samples, immediately after the excisions (0 h) and 24 h and 48 h thereafter. The irradiated and non-irradiated excised posterior segments were placed in individual sterile Petri dishes (Ø 19 cm), with 60 gr of organic soil and 50 mL of water. The samples were kept under this experimental condition for up to twelve-days and 20 mL of water was added every three days.

Simple organisms like the worms show a high-ability for wound healing and regeneration. Therefore, in our experiments, to point out the laser treatments’ effectiveness, we chose to excise the 30-posterior segments because the literature reported this condition induces the death of the 100% excised samples, within 15 days, as a consequence of the lack of healing.

The irradiated and non-irradiated samples were analyzed at different time intervals for the evaluation of bacteria load, macroscopic changes, histochemical and immunohistochemical analysis, expression of HSP70 and telomerase activity.

**Sample temperature analysis**

The temperature of the excised posterior segments was measured in 5 points of the earthworm section, before and after the irradiation by a Testo 826-T4 infrared thermometer with laser marking (Testo Spa, Milan, Italy).

**Bacterial load analysis**

The irradiated and non-irradiated excised posterior segments were immediately swabbed, under sterile laminar flow...
hood, on either Tryptic Soy Agar (15 gr/L agar, 15 gr/L peptone from pancreatic digest casein, 5 gr/L papaisoy peptone, 5 gr/L sodium chloride, pH 7.4) or in LB broth (10 gr/L tryptone, 5 gr yeast extract, 10 gr/L sodium chloride, pH 7.4). The Tryptic Soy Agar and the LB broth were kept in the incubator (Gallenkamp GmbH, UK) at 22°C (D. veneta’s breeding temperature), for 24 h. After 24 h the Tryptic Soy Agar was photographed and the images acquired by the Zeiss stereomicroscope with a Cellpad E (TiEsseLab s.r.l., Milan, Italy). The bacteria load was deduced by the measurement with ImageJ 1.33J software (National Institute of Health, Bethesda, MD, USA) of the bacterized area. In addition, the bacterized areas were carefully collected, suspended in LB broth and the bacteria loads were detected spectrophotometrically by a Jenway-6405-UV-Vis Spectrophotometer (Jenway, Felsted, Dunmow Essex, UK). The bacteria load of the experiments in LB broth was deduced spectrophotometrically by a Jenway-6405-UV-Vis Spectrophotometer, at 3, 6 and 24 h after the swab.

**Macroscopic changes analysis**

The irradiated and non-irradiated excised posterior segments were examined by a Zeiss stereomicroscope, immediately after the irradiation or the non-irradiation (t0) and every 24 h from those. The images were acquired with a Cellpad E (TiEsseLab s.r.l.). The vitality of the segments was evaluated by prodding with a metal tip. The stimulation reactions were observed. No reaction was considered as synonymous with the death of the organism.

**Histochemical analysis**

The 24 h-, 48 h- and 72 h-irradiated and non-irradiated excised posterior segments were fixed in 4% paraformaldehyde in a 0.1 M phosphate buffered solution (PBS, pH 7.4) at 4°C, Paraplast (Bio-Optica Spa, Milan, Italy) embedded and cut into 5-μm-thick sections. Histological observations were performed using hematoxylin-eosin (H&E) and Masson’s trichrome. The Alexa 488-conjugated lectin wheat germ agglutinin (WGA; source: *Triticum vulgaris*, binding specificity: N-acetylglucosamine and N-acetylneuraminic acid, working dilution: 10 mg/mL – Termofisher Scientific, Waltham, MA, USA) was used. DNA pres-

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**Table 1. Description of the irradiation parameters and experimental design.**

| Irradiation parameters | Laser | Wavelength | Hand-piece | Electromagnetic wave | Power |
|------------------------|-------|------------|------------|----------------------|-------|
| Diode                  | 808 nm| Flat-top   | Continuous wave | 1 W |
| Irradiation time       | Fluence| Irradiance | Distance | Treatments |
| 60 sec                 | 64 J/cm²| 1 W/cm² | 1 mm | 0 h, 24 h, 48 h from excision |

| Experimental design | Organism | Segments number | Weight | Segment excised |
|---------------------|----------|-----------------|--------|-----------------|
|                     | *Dendrobaena veneta* | ~ 150 | ~ 1.5 gr | 30 |
ence was identified, when necessary, by 4',6-diamidino-2-phenylindole (DAPI) fluorescence staining. Sections were examined on a Leica DMRB light and epifluorescence microscope; images were acquired with a Leica CCD camera DFC420C (Leica, Heerbrugg, Switzerland).

HSP70 expression analysis

Immune responses to heat shock proteins (HSP) develop in virtually all inflammatory diseases and evidence on anti-inflammatory effects of HSP70 are shown in literature. Therefore, the amount of HSP70 mRNA (MF093577) in excised posterior segments of *D. veneta* 24 h- and 48 h-irradiated and non-irradiated was determined by qPCR using Chromo 4 apparatus (MJ Research, Waltham, MA, USA).

First total RNA was extracted from frozen liquid nitrogen tissue samples using RNeasy Mini Kit (QIAGEN, Milan, Italy) and RNase-Free DNase Set (QIAGEN) according to manufacturer’s instructions. Then worm cDNA was synthesized by iScript reverse transcriptase (Bio-RAD, Hercules, CA, USA) in a final volume of 20 μL using 500 ng of purified total RNA. Subsequently, each PCR reaction was performed in 20 μL containing: 1× master mix iQSYBR® Green (Bio-RAD), 0.2 μM of each primer, and 0.8 μL of synthesized cDNA or negative control. All samples were analyzed in triplicate. The reference gene used for the samples normalization was *D. veneta* 18S ribosomal RNA gene (KJ911964.1) whose primers sequence were (5′CCGTAACTCTGATGACTCTG) for forward and (5′CTTCCTTTGAGTGGTAGC 3′) for reverse while the HSP70 primers sequence were (5′ GGACAACAACCTCCTTGGAA 3′) for forward (5′ CGAAGGTTACCTCCTGTA 3′) for reverse. All primers were designed using the software Beacon Designer 7.0 (PREMIER Biosoft International, Palo Alto, CA, USA) and obtained from TibMolBiol (Genoa, Italy). The following thermal conditions were used: initial denaturation for 3 min, followed by 45 cycles with denaturation at 95°C for 15 s, annealing and elongation at 60°C for 30 s. The fluorescence was measured at the end of each elongation step. The next step was a slow heating (1°C/s) of the amplified product from 55°C to 92°C in order to generate a melting temperature curve. Data analyses were obtained using the DNA Engine Opticon® 3 Real-Time Detection System Software program (ver. 3.03) and, in order to detect the relative gene expression of HSP70, compared with a calibrator sample (uncut worm), the comparative threshold Ct method was used with the software GeneExpression Analysis for iCycleriQ Real-Time Detection System (Bio-Rad).

Telomerase activity analysis

The telomerase activity is typically correlated to cellular proliferation during wound healing. Therefore, the 24 h-, 48 h-, 72 h- and 96 h-irradiated and non-irradiated excised posterior segments were frozen in liquid nitrogen and their proliferative status was detected by qPCR using the “Quantitative Telomerase Detection kit” (Allied Biotech, Inc., Benicia, CA, USA), following the manufacturer’s instruction. The amount of extracted protein used in each sample was 1 μg. To correlate the Ct obtained in the different samples, a standard curve was prepared with a 1:5 serial dilution of an oligonucleotide (TSR) supplied by the kit and the enzymatic activity was finally expressed as attomoles per micrometer protein. Specifically, values for 24 h, 48 h, 72 h and 96 h-irradiated samples were compared to 24 h-, 48 h-, 72 h- and 96 h non-irradiated samples ones.

Statistical analysis

The statistical analysis was performed using a one-way ANOVA followed by the Tukey-Kramer multi-comparison test (GraphPadInStat 3) to discriminate statistically significant results. High significance level: P<0.001 (**), significance level: P<0.01 (**), significance level: P<0.05 (*), not-significant level: P>0.05. Each experi-

![Figure 1. Effect of the laser irradiation on the temperature of *Dendrobaena veneta* excised side. The temperature was measured before (control) and after (irradiated) the laser irradiation.](https://example.com/figure1)

![Figure 2. Bacterial load analysis in non-irradiated (control) and irradiated excised side of *Dendrobaena veneta*. A = bacterial colonies grown for 24 h on Tryptic Soy Agar. B= image analysis of the area of bacterial colonies grown for 24 h on Tryptic Soy Agar. C= spectrophotometric analysis of the bacterial colonies grown for 24 h on Tryptic Soy Agar. D= spectrophotometric analysis of bacterial colonies grown for 3 h, 6 h and 24 h in LB broth. O.D.= optical density. No-star symbol= P>0.05; * = P<0.05; **= P<0.01; ***= P<0.001.](https://example.com/figure2)
ment was carried out by 3 replications and 5 repetitions.

Results

Evaluation of thermic increment

The temperature of the excised posterior segments was 22.0°C±0.2 before the irradiation and 22.5°C±0.4 afterwards (Figure 1). The difference between the measures did not constitute statistical significance (P>0.05, data not shown).

Evaluation of bacteria load analysis

The analysis of the excised posterior segments swabs showed that the bacterized area on the Tryptic Soy Agar was reduced in the irradiated samples (~41%) compared to the non-irradiated posterior segments (P<0.001) (Figure 2A, B). The spectrophotometric analyses of the bacterized area emphasizes these differences, quantifying the bacteria load of the irradiated sample as 52% less than the non-irradiated sample (P<0.001) (Figure 2C). In addition, the spectrophotometric analysis of the LB bacterized broth, 3 h, 6 h and 24 h after the swab, showed that the irradiation reduced the LB bacteria load by about 45%, 40% and 62%, respectively (P<0.001) (Figure 2D).

Evaluation of survival rate

The reaction to the external stimulus showed that, 6 days after the excision, 100% of both irradiated and non-irradiated excised posterior segments were alive (P>0.05) (Figure 3), with the irradiated excised posterior segments appearing more vital and dynamic. 9 days and 12 days after the excision, 100% of the irradiated samples were healthy, while only 75% and 50% respectively, of the non-irradiated samples appeared vital (P<0.001) (Figure 3). Finally, 16 days after the excision, all the non-irradiated segments succumbed, whereas 100% of the irradiated segments survived (P<0.001) (Figure 3).

Evaluation of macroscopic changes

Immediately after the 30 posterior segment excision the analysis by stereomicroscopy revealed the evagination of the alimentary tract and the major blood vessel. The coelom cavity was open and the coelomic fluid emerged as well as the blood. The body wall was loose and the wound appearance revealed a circular section of diameter similar to the uncut body diameter (Figure 4 A,B). After the first laser treatment, performed 2 minutes after the excision (0 h), the body exhibited a dorsal-ventral thinning and the wound diameter reduced. The body contraction induced the internalization of the alimentary tract, which disappeared from the wound site and the major blood vessel was not visible. The coelomic fluid and the blood flow ceased (Figure 4B’).

Twenty-four hours after the first laser treatment, in the irradiated samples the wound seemed to close and the body wall contracted (Figure 4D), while the non-irradiated samples remained similar to the previous day.

Forty-eight hours after the excision and 24 h from the second treatment, the irradiated samples wound continued to recover and its color appeared uniform with the rest of the body (Figure 4 F). Conversely, the alimentary tract of the non-irradiated sample remained evaginated, although the body wall was more contracted and both the coelomic fluid and blood were absent (Figure 4E).

Seventy-two hours after the excision and 24 h from the third treatment, the wound of irradiated samples regained shape and pigmentation in similar form to the rest of the body (Figure 4H), while the non-irradiated wound remained open with the alimentary tract visible (Figure 4G).

Lastly, 144 h (6 days) after the excision...
and 96 h from the third and last treatment, the irradiated posterior excised segments gave an appearance similar to the posterior side of a non-excised earthworm (Figure 4L), whereas the non-irradiated samples showed evidence of gangrene (Figure 4I).

**Histology - general observations**

Twenty-four hours after the dissection, in the control specimens, the body wall muscles were not contracted, the alimentary tract lumen was in contact with the external environment and the alimentary tract wall was everted, forming a wide protrusion (Figure 5A). The blood vessels close to the wound were enlarged (Figure 6A).

Twenty-four hours after the dissection and the first laser treatment, the body wall muscles were contracted and the alimentary tract wall was non-everted (Figure 5B). The blood vessels close to the wound revealed a reduced lumen compared to the control (Figure 6B).

Forty-eight hours after the dissection, in control specimens, the body wall muscles were partially contracted and formed an easily-visible bulge close to the wound site. The alimentary tract lumen was open to the external environment, and the alimentary tract wall was still everted (Figure 5C), as well as the blood vessels being still enlarged (Figure 6C).

Forty-eight hours after the dissection and the second laser treatments, the edges of the wound appeared near-apposed and the alimentary tract lumen withdrawn into the body cavity (Figure 5D). The blood vessels were still reduced in size (Figure 6D).

Seventy-two hours after the dissection, in control specimens, the alimentary tract was no longer everted and in some cases it appeared withdrawn into the body cavity (Figure 5E).

Seventy-two hours after the dissection and the third laser treatments, the body wall margins had closed and epidermis covered the wound site (Figure 5F).

**Histology - body wall muscles**

The body wall muscles in control specimens presented a progressive degeneration, from 24 h to 72 h after the dissection (Figure 7 A,C,E); in particular, gaps appeared among the muscle fibers, with a more abundant presence of collagen. The histology of body wall muscles in laser-treated specimens did not show degeneration (Figure 7 B,D,F).

**Histology - WGA labeling**

The WGA lectin highlights three different structures in the histological sections of *D. veneta*: the cytoplasm of scattered cells in the peritoneum, which lines the coelom (Figure 8A); cytoplasm of cells among the muscle fibers in the body wall (Figure 8 B-F); granules in the cytoplasm of epidermal cells (Figure 8F).

The scattered WGA-labeled cells in the peritoneum were always present and did not change their aspect and distribution in different specimens. Alternatively, the WGA-labeled cells among the muscle fibers changed their distribution over time and different treatments. In control specimens, they became visible 24 h after the dissection (Figure 8B) and increased their presence at 48 h (Figure 8D) and 72 h (Figure 8F). Also, a continuity between WGA-labeled cells of peritoneum and WGA-labeled cells in the muscle tissue was observed in some slides. In the laser-treated specimens the WGA-labeled cells in the muscle tissue were almost absent after 24 h (Figure 8C), 48 h (Figure 8E) and 72 h (and three laser treatments) from the dissection (data not shown). The WGA-labeled granules in epidermal cells were observed only in control specimens and they were most abundant at 72 h after the dissection (Figure 8F).
Evaluation of telomerase activity

The telomerase activity increased, compared to the uncut (P<0.001), after the excision in both the irradiated and the non-irradiated posterior segments (P<0.05) and it remained constant up to 72 h from the excision (Figure 9). However, at 96 h the level of telomerase activity was maintained in the irradiated sample, while there was a dramatic decrease in the non-irradiated samples (P<0.001) (Figure 9).

Evaluation of HSP70 expression

The HSP70 protein expression increased approximately 10 times in the non-irradiated sample (control), compared to the uncut earthworm, whereas imperceptible, but statistically significant, change was recorded in the irradiated earthworm (P<0.001) (Figure 10). At 48 h from the excision the non-irradiated earthworm exhibited continued increase in HSP70 expression (P<0.001) and the irradiated sample retained a protein expression similar to the uncut earthworm (P>0.05) (Figure 10).

Discussion

The ability to respond to injury and to repair tissue is a fundamental property of multicellular organisms. In the moments after an injury occurs, various intracellular and intercellular pathways must be activated and coordinated, if tissue integrity and homeostasis are to be restored.19 If this response is successful and the injury does not result in the demise of the organism, these processes must be shut down in a precise sequence in the ensuing days as recovery progresses. Notwithstanding differences between varying organisms, the three classic stages of wound repair, inflammation, new tissue formation and remodeling, are observed in almost all eukaryotic organisms.37,7 In our work, the effect of the treatment with 808 nm diode laser, 64 J/cm2, 1 W(CW), on the wound healing of excised posterior segments of D. veneta are classified in three steps: immediate effects (0 h after irradiation), short-term effects (24 h, 48 h and 72 h after irradiation) and long-term effects (72 h over irradiation).

Immediately after treatment the irradiated D. veneta showed evident muscular contraction at the wound site, pointed out by the dorsal-ventral thinning of the body and the internalization of the alimentary tract, which disappeared from the wound site only in irradiated earthworms. Oligochaeta show striate transverse and helical musculature arrangement,27 which, like the vertebrate striate musculature, contracts by changes in calcium fluxes and variation of intracellular calcium concentration.28 Amaroli et al.14 proved the effect of the 808nm diode laser (64 J/cm², 1 W) therapy on P. primaurelia calcium fluxes and Sharmizaev35 showed marked spasmyolytic properties of laser irradiation, likely due to calcium signaling in the adjacent muscle’s cells. Therefore, our data suggest an effect of the laser on the cellular muscle calcium fluxes, which result in the D. veneta striate musculature contraction.

In the moments after irradiation, the PBM decreases the bacteria concentration on the top of excised posterior segments, as revealed by our microbiological analysis. Literature shows that PBM may modify growth of wound bacteria, which could affect wound healing.36 In particular, fluences from 4.2 J/cm² up to 50 J/cm² (CW) and laser light wavelengths in the range of 380-810 nm can excite endogenous porphyrins and kill the related pigmented-bacteria.36,37

We do not know the qualitative effect of our therapy on the earthworm excised posterior segments bacteria populations, however, the quantitative effect is clear and its role in the inflammatory process should not be underestimated. The short-term effects support the observed immediate effects. The histological analysis of samples fixed 24 h after the first irradiation or non-irradiation, highlights the everted intestine of the controls, which is folded back to expose the inside epithelium. The alimentary tract eversion and the formation of a coelomic plug are the two classical ways to rapidly close the wound, as described in the earthworm Eisenia fetida.25 Amaroli et al.24 showed that also D. veneta is able to produce the coelomic plug and this plug is thinner in the excised earthworm exposed to 808 nm diode laser, 64 J/cm², 1 W therapy, than in the control, as a demonstrable consequence of a decreased inflammatory process.

In our work, in contrast to the control, the irradiated excised tails did not close the wound through alimentary tract eversion, but by a vigorous muscular contraction. The apposition of the irradiated excised tail body wall flaps provides evidence of the consequence of this contraction. In addition, by the Masson’s trichrome stain, a new tissue mass is identified. This mass is collagen-free, while the collagen is evident in the flaps existing tissue. The differences between irradiated and non-irradiated (con-
trol) are also evident, for the major blood vessels close to the excised side. In fact, their diameter is smaller in the irradiated than in the non-irradiated histological sample and the contour edge of blood vessel lumen is more regular in the irradiated than in the non-irradiated histological sample and the contour edge of blood vessel lumen. The endothelial cells of the earthworm are properly considered mioendothelial cells, whose contraction ability increase with the increment of blood vessel lumen. The earthworm blood vessel contraction relies on the myofilaments, which contract by variations of intracellular calcium concentration. It is not unreasonable therefore to assume, as with the major blood vessels, that our laser irradiation promotes a contraction, with a mechanism similar to that hypothesized for the muscular tissue.

Our histological analysis shows also other evident differences between irradiated and non-irradiated excised tails. Twenty-four hours after the excision, and ever more evident in the following days the muscular fibers progressively became distinct from each other, with the consequent increase in the connective tissue amongst them. This change is clear when observing the longitudinal cross-sections of circle musculature, just below the epidermis, as well as with the longitudinal musculature. Conversely, the irradiated sample had a compact muscular wall, with thin trabecular connective among the fibers. It is known that muscular atrophy is a phenomenon occurring in the organism with muscle tissue as consequence of senescence, inactivity or pathological causes and in mammalian species a correlation between inflammatory processes and muscle atrophy is documented. The HSP70 is one of the most conserved proteins of “animal kingdom” and its involvement in the inflammatory process is well documented. In fact, HSP70 is involved in the restoration of cell homeostasis, in the immunoregulation and in the phagocytosis of cellular residue, to promote cell proliferation and wound healing. Our data show for the first time the expression of HSP70 in D. veneta. In the control samples its expression was time-dependent and highest at 48 h (+1984% respect to the non-excised earthworm). The irradiated samples showed increase in HSP70-expression at 24 h but the increment was very small (+60% compared to the non-excised earthworm) and at 48 h, after the third irradiation the HSP70-levels were similar to that in the non-excised earthworm. Therefore, a reduction in the inflammation through PBM without a thermal-shock is evident.

This assertion is supported by our previ-
They could be evidence of a virus infection, as reported in *Drosophila melanogaster*, in which it induces the HSP70 increment. However, moving beyond these speculations, it is clear that this was present only in the epidermis of the control, progressively destined to gangrene, but not in the irradiated-epidermis. If the first step in wound healing is the control of the inflammation processes to restore homeostasis, the new tissue formation by new cell proliferation is the second. The histochemical analysis as little as at 24 h showed new tissue formation in the irradiated excised earthworm, but our evaluation of the telomerase activity pointed out an interesting cell behavior. The excise earthworm had an increment of telomerase activity, compared to the non-excised earthworm. Up to 72 h, this increment was similar for both irradiated and non-irradiated *D. veneta*. At 96 h it decreased drastically in the control, while it remained at high level in the irradiated samples. Actually, the high-level telomerase activity observed at 96 h in the irradiated organisms cannot be a direct effect of the irradiation (the last irradiation was at 48 h). Therefore, if as reported in literature, the telomerase activity is typically correlated to cellular proliferation during the wound healing of mammalian tissue, we can speculate that the inflammatory processes and the tissue degeneration of non-irradiated excised *D. veneta* deplete the cell energy and its proliferative potential within the fourth-day. Conversely, the laser irradiation would allow a more equilibrated employment of cell energy, by the mitigating effect on both inflammation and tissue degeneration and a likely effect on the energetic cellular metabolism, in according to our previous works.

A foreseeable consequence therefore emerges when all data is considered: the control specimens become infected, become gangrenous and die, while the irradiated samples complete the wound healing and tissue remodeling and after twelve-days they are alive.

Lastly, our results are stressed by the evidence that invertebrate species are as diverse as flies and worms and share a lot of genes and molecular pathways with humans. More than 90% of the domains that can be identified in human proteins are present in fruit fly and worm proteins, while 61% of fruit fly proteins and 43% of worm proteins have high sequence similarities to predicted human proteins. Furthermore, the gene conservation of earthworms is closer to humans than worms such as the well-known animal model *Caenorhabditis elegans*. In fact, the bilaterian metazoan showing bilateral symmetry, are divided into two clades: Deuterostomes and...
In conclusion, the 808 nm diode laser therapy, using a fluence of 64 J/cm² at 1 W (CW) with a flat-top hand-piece, proved to induce positive PBM. This was sustained in the transition from cell to the animal model. Three successive treatments, one every 24 h are sufficient to promote wound healing, through effects on muscular and blood vessel contraction, decrease in bacteria load, reduction of inflammatory processes and tissue degeneration. D. veneta was demonstrated to be a reliable experimental organism that satisfies the request of public opinion in the field of bioethics, the 3Rs principles and the National Science Foundation statement “…. All model organisms share a few common traits: They’re inexpensive, easy to care for, grow quickly and are relatively simple creatures”.46

In acknowledgement of their genetic and evolutionary peculiarity, not inferior to those of scientifically accredited models, D. veneta allows investigation of the effect of laser therapies by multidisciplinary methods, at various degree of complexity and costs.

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