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How to cite this article
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This original article is available in Indian Journal of Research in Homoeopathy: https://www.ijrh.org/journal/vol2/iss2/3
**FUNDAMENTAL RESEARCH**

**Effect of Low Level Laser Therapy and Calendula officinalis 3 CH on Wound Healing in Human Skin Fibroblasts**

Annelise Bresler (BA (WITS) M.Tech HOM), Denise Hawkins(M. Biomed), Radmila Razlog (M.Tech HOM) and Heidi Abrahamse (Prof)*.

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Keywords: wound healing, a comparison study of Calendula 3 CH and Laser phototherapy; Calendula, and wound healing; laser therapy, and wound healing.

**Introduction**

Skin fibroblasts are a vital component of the dermis; they not only produce and organize the extracellular matrix of the dermis, but they also communicate with each other and other cell types, playing a crucial role in regulating skin physiology. (1) Fibroblasts are a well-established system for in vitro analysis of fibroblast growth, migration and collagen metabolism in wound healing.(2,3,4)

The laser is an electro-optical device capable of efficiently transmitting energy in the form of an intense beam of light. The radiant energy of the laser beam can be transformed into heat energy through its interaction with tissues. The interaction is dependent on the wavelength of the laser and the properties of the tissue. Laser light is absorbed by water, and as most biologic tissues are composed mainly of water, the tissues are heated. (5) However, experiments following LLLT have shown that immediate increase of heat in the target tissue is negligible (+/-1°C). Low level lasers do not damage the hydrogen bonds since there is no heating effect. The effect is photochemical.(6) The basic tenet of LLLT is that it has a wavelength-dependent capability to alter cellular behavior in the absence of significant heating.(7)

It is the coherence of laser which is the most significant factor for skin penetration, thus allowing for a photochemical effect to occur.(8) When a laser beam is irradiated onto tissue, a local or topical primary response occurs at the cellular level; this is followed to some degree over a greater area by a systemic secondary response.

In the laboratory, Calendula extracts have been proven to be anti-edematous (9) antibacterial (10), antifungal (11) antiviral (12) and immuno-stimulating.(13) Calendula officinalis is a remedy seldom used internally.
Its major value lies in its local application to external wounds. (14) Topical application of Calendula officinalis markedly stimulates physiological regeneration and re-epithelization. (15) Dressing materials containing Calendula officinalis ointment applied to experimental animal wounds have also been found to enhance tissue repair. (16) Jouanny (17) advises its use externally as an antiseptic and cicatrizating agent. Phatak (18), Boericke (19) and Hering (20) state that Calendula officinalis promotes wound healing.

Interestingly enough, homeopathy has been explained to work in a similar manner to LLLT, which is based on its biostimulatory effect. The Arndt-Schulz law is used by both modalities to elucidate the underlying mechanisms involved. (21, 8) This law states that weak stimuli excite physiological activity while strong stimuli retard it. (8) It is evident from reviewing the literature that both modalities would argue that their treatment helps the body's own innate capacity to heal.

Allied health professionals regularly care for a variety of wounds to the skin, namely abrasions, burns, surgical incisions, and perhaps the most difficult to treat, ulcerations. (22) From acute wound management to augmentation of scar tissue remodelling, the clinician seeks to optimize wound care to promote healing. A comprehensive treatment plan for wounds including complementary and alternative therapies may improve the rate of wound healing. This may result in reduced hospitalization thus limiting treatment and nursing time and reducing complications. The patient will be in a position to return to active life sooner and in doing so the economy at large will benefit.

Materials and Methods

Cell Culture

Commercially available human skin fibroblast (HSF) cell lines (CRL1502 WSI) were obtained from the American Type Culture Collection (ATCC). Cells were grown in 25 ml of Eagle's minimal essential medium (EMEM) with Earle's balanced salt solution (BSS) that was modified to contain 2 mM L-glutamine, 1.0 mM sodium pyruvate, 0.1 mM non-essential amino acids, 1 % fungicide (BioWhittaker 17836-E) and 1 % penicillin-streptomycin. In addition, it was supplemented with 10 % foetal bovine serum (Delta Bioproducts 14-501Al). The cultures were incubated at 37°C with 5% CO2 and 85% humidity. The cells were examined by inverse light microscopy and subcultured according to standard procedures. (23)

Experimental Procedure

The HSF cells were subcultured from the 75 cm² flask to 3.3 cm diameter culture plates. Approximately 6 x 10^5 cells in 3 ml culture medium were seeded in six sterile 3.3 cm diameter culture plates and incubated overnight to allow the cells to attach. The next day (Day 1) the plates were taken out of the incubator and 1 ml of culture medium was removed from each plate and

<table>
<thead>
<tr>
<th>Plate</th>
<th>Treatment</th>
<th>No wound/wound</th>
<th>Control/Test</th>
<th>Key</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>No Treatment</td>
<td>No wound</td>
<td>Control</td>
<td>-W -C -I</td>
</tr>
<tr>
<td>2</td>
<td>No treatment</td>
<td>Wound</td>
<td>Control</td>
<td>+W -C -I</td>
</tr>
<tr>
<td>3</td>
<td><em>Calendula officinalis</em> 3 CH in 5% ethanol, 10 µl.</td>
<td>Wound</td>
<td>Test</td>
<td>+W +C -I</td>
</tr>
<tr>
<td>4</td>
<td>Irradiation with 632.8 nm Helium Neon (HeNe) laser, 5 J/cm² at calculated dosage.</td>
<td>Wound</td>
<td>Test</td>
<td>+W -C +I</td>
</tr>
<tr>
<td>5</td>
<td><em>Calendula officinalis</em> 3 CH in 5% ethanol, 10 µl plus irradiation simultaneously.</td>
<td>Wound</td>
<td>Test</td>
<td>+W +C +I</td>
</tr>
<tr>
<td>6</td>
<td>Irradiation with 632.8 nm HeNe laser, 5 J/cm² at calculated dosage thereafter the plate was placed in the incubator for 4 min and subsequently the 10 µl of <em>Calendula officinalis</em> 3 CH in 5% ethanol was added.</td>
<td>Wound</td>
<td>Test</td>
<td>+W +I +C</td>
</tr>
</tbody>
</table>

Table 1: Experimental Procedure
discarded. Five of the six plates were wounded. To simulate a wound, mechanical trauma was induced by means of a central scratch with a 1 ml sterile Pasteur pipette.(24,25,26). Each scratch was irregular and the wounds ranged from 1-2 mm in diameter. The plates were incubated at 37°C for 30 min before each treatment regime.

Treatment was applied on two consecutive days (Day 1 and Day 2). Cells were visualised 24 hours (Day 2) and 48 hours (Day 3) after wound induction with an inverted light microscope. Cellular responses were measured on Day 3. Four of the plates received treatment while the remaining two served as controls. The experimental procedure is summarized in Table 1. The procedure was repeated six times (n=6).

Laser Irradiation

Laser irradiation was carried out with a Helium Neon (HeNe) laser, which has a wavelength of 632.8 nm, a power output of 18.8 mW and a power density of 2.07 mW/cm². The dose delivered was 5J/cm² (40 min 15 s) for a 3.3 cm spot size. Cell culture dishes were placed under the laser beam and irradiated with the culture dish lid on at room temperature in the dark on a dark surface.

Calendula officinalis 3 CH

A preliminary study established that cells which received 10 l l of Calendula officinalis 3 CH dispensed in 5% ethanol showed optimal signs of cell migration with minimal cell death. A volume of 10 l l of Calendula officinalis 3 CH was added to the culture medium for specified plates on Day 1 and Day 2 of the experiment.

Cellular responses

Cell morphology

Both wound contraction and re-epithelialization from the margins of the wound play an important role in wound closure.(27,28,29) Wound closure has been studied both in vivo and in vitro.(3,30,31) Two digital photos were taken of each wound on Day 2 and Day 3. Photos were evaluated for changes in migration (wound closure), haptotaxis (change in orientation of edge fibroblasts) and the number of fibroblasts present in the central scratch.(26)

ATP Cell Viability Assay

The CellTiter-Glo® (Promega G7571) luminescent cell viability assay determines the number of viable cells in culture based on the quantitation of ATP present, which signals the presence of metabolically active cells.(32) An equal volume of reconstituted CellTiter-Glo® reagent was added to 50 l l of cell suspension (x10⁵/100 l l).

The contents were mixed for 3 minutes by hand to induce cell lysis. It was then incubated at room temperature for 10 minutes to stabilize the luminescent signal and the luminescence was recorded using a Junior EG&G Berthold Luminometer.

Trypan Blue Exclusion Test

Trypan blue is a stain recommended for use in estimating the proportion of viable cells in a population.(33,34) A vital dye such as trypan blue can enter a damaged cell membrane and stain the cytoplasm acting as an indicator of non-viable cells.(23) Live (viable) cells do not take up the dye and dead (non-viable) cells do. For the trypan exclusion test, 10 l l of the cell suspension was mixed with 10 l l of 0.4% Trypan blue (Sigma T8154). This solution was left at room temperature for 5 minutes and loaded into the haemocytometer-counting chamber. The total number of cells per millilitre, the total number of unstained (viable) and the total number of stained (non-viable) cells were counted using a light microscope. The viable cells were expressed as a percentage and calculated as the number of viable cells/total number x 100.

Lactate Dehydrogenase (LDH) Membrane Integrity Assay

The CytoTox 96® (Promega G1781) assay indirectly measures the lactate dehydrogenase activity present in the cytoplasm of intact cells. Cell quantitation, therefore, can occur only if the cells are lysed to release the LDH present in the cell. The CytoTox 96® measures this stable cytosolic enzyme that is released upon cell lysis. Culture medium was removed from each plate on Day 3. 50 l l of culture medium was mixed with an equal volume of reconstituted substrate mix in a 96-well plate. The plate was covered with foil and incubated at room temperature for 30 minutes, protected from light. 50 l l of stop solution was added and the absorbance read at 490 nm.

Results

Changes in Cell Morphology

The morphological changes observed after treatment were recorded on Day 3 (Figure 1). The HSF cells that received no wound showed a normal state of biologic activity where normal growth was observed, highlighting the sustainability of the cultured environment to support continued existence and growth. Fibroblast activity was normal but also showed some non-viable cells (A). Wounded fibroblasts that received no treatment showed fibroblast migration into the central scratch (CS) area. The morphological changes of the wounded untreated fibroblasts served as the control against which the individual treatments were compared (B).

Indian Journal of Research in Homoeopathy
Vol. 2, No. 2, April-June 2008
Figure 1. Changes in Cell Morphology
HSF cells 48 h after initiation of double treatment protocol. Unwounded HSF cells receiving no treatment demonstrate normal cell growth (A). Wounded HSF cells receiving no treatment show slight migration of fibroblasts across the wound margin (B). Wounded HSF cells treated with Calendula officinalis 3 CH show considerable migration of cells across the WM with extensive fibroblast presence in the central scratch (C). Wounded HSF cells treated with laser irradiation show migration and haptotaxis of cells across the WM. This photo has been enlarged 40 X to show haptotaxis in detail (D). Wounded HSF cells treated with Calendula officinalis 3 CH followed by laser irradiation show minimal fibroblast migration across the WM (E). However, wounded HSF cells receiving laser irradiation followed by the administration of Calendula officinalis 3 CH demonstrate haptotaxis at the WM (F).
Fibroblasts treated with Calendula officinalis 3 CH showed migration (M) of fibroblasts across the (wound margin) WM into the CS area. A significant amount of fibroblasts (F) can be seen in the CS (C). Although Figure 1 (D) has been taken under the 10 X magnification, it was enlarged to show haptotaxis (change in the orientation of the edge fibroblasts) in detail. Wounded HSF cells treated with laser irradiation show haptotaxis at the wound margin. The cytoplasmic processes of the fibroblasts can be seen to change their orientation at the WM as it invades the CS area (D).

Slight infiltration of fibroblasts into the CS area can be observed for cells treated with Calendula officinalis 3 CH followed by laser irradiation on Day 3. Fibroblasts present in the CS were indicated with an arrow (E). Haptotaxis was visible for cells treated with laser irradiation followed by administration of Calendula officinalis 3 CH at the wound margin. Fragments of the plate, which came loose during the wound induction process, are encircled (F).

**Figure 2.** Cell viability was measured using ATP luminescence and Trypan blue stain. ATP luminescence (bars) and Trypan blue (scatter line) showed no statistical significant difference between groups. The p - value indicated in each bar represents the statistical difference between (W - C - I) and each treatment group. (n = 6).

**Cell Viability**

The mean cell viability measured by ATP luminescence over six consecutive experiments shows cell viability was highest for wounded cells which received Calendula officinalis 3 CH as treatment (+W +C - I) (Figure 2). No statistically significant change was calculated between wounded cells receiving no treatment (+W -C - I) and HSF cells receiving Calendula officinalis 3 CH (+W +C - I) as treatment (P = 0.116).

Unwounded cells receiving no treatment represent the normal reference state of metabolic activity and measure the second highest cell viability (W -C - I). Wounded cells receiving no treatment (+W -C - I) show a decrease in cell viability when compared to the unwounded cells receiving no treatment (-W -C - I) (P = 0.111). Morphological differences between the unwounded cells (-W -C - I) and wounded cells receiving no treatment (+W -C - I) confirm that a wound environment was successfully induced.
Wounded HSF cells which received laser irradiation (+W-C+1) and wounded HSF cells which were irradiated followed by the administration of Calendula officinalis 3 CH (+W +1 +C) measured higher ATP luminescence when compared to wounded cells which received no treatment (+W-C-I). Wounded HSF cells, which received Calendula officinalis 3 CH followed by laser irradiation (+W +C +1) showed the lowest cell viability. No statistically significant difference was calculated between the treatment group which measured the lowest ATP viability (+W +C +1) and between the group which measured the highest ATP viability (+W +C -I) (P = 0.078).

**Trypan Blue Exclusion Test**

The scatter line in Figure 2 represents the percentage of viable cells for each procedure. Viability ranged between 90% and 96% and is read on the right Y-axis. HSF cells treated with Calendula officinalis 3 CH (+W +C -I) and the unwounded cells receiving no treatment (-W-C-I) calculated the highest percentage of viable cells, which corresponds with the results obtained for the ATP luminescence assay measuring cell viability.

This consistency between the two measuring tools for cell viability, namely ATP luminescence and Trypan blue was not established for the following two treatment groups: laser irradiation (+W -C +1) and irradiation followed by Calendula officinalis 3 CH (+W +1 +C). Viability percentages were lower for these when compared to wounded cells receiving no treatment (+W -C -I). This stands in contrast to the ATP cell viability measurements where both laser irradiated (+W -C +1) cells and irradiated cells followed by administration of Calendula officinalis 3 CH (+W +1 +C) measured higher cell viability compared to wounded cells receiving no treatment (+W -C -I).

**LDH Membrane Integrity Assay**

Lactate dehydrogenase (LDH) is released by dying cells and therefore gives a quantitative measurement of the non-viable population. LDH activity was lowest for both unwounded cells receiving no treatment (-W-C-I) and wounded cells receiving irradiation followed by the administration of Calendula officinalis 3 CH (+W +1 +C). Thus, the unwounded cells and wounded cells receiving

![Graph showing LDH membrane integrity assay](image)

**Figure 3.** Cytotoxicity was measured by means of a LDH membrane integrity assay. A statistical significant increase in cytotoxicity was measured for untreated HSF cells receiving no wound and those receiving a wound (P = 0.03). Furthermore, a statistically significant decrease in cytotoxicity was measured for wounded HSF cells receiving no treatment and wounded HSF cells treated with irradiation followed by the administration of Calendula officinalis 3 CH (P = 0.025). (n = 6, *P = less than or equal 0.05).
laser irradiation followed by the administration of Calendula officinalis 3 CH (+W +1 +C) were the least cytotoxic (Figure 3).

The wounded HSF cells which received laser irradiation (+W -C +1) as treatment achieved the highest LDH activity and were consequently identified as the most cytotoxic treatment group. The wounded cells that received no treatment (+W -C -I) showed the second highest LDH activity, closely followed by the fibroblasts which received Calendula officinalis 3 CH (+W +C -I). Administration of Calendula officinalis 3 CH followed by laser irradiation (+W +C +1) showed a cytotoxicity greater than wounded cells receiving laser irradiation followed by the administration of Calendula officinalis 3 CH (+W +1 +C) indicating that laser irradiation may adversely affect the activity of Calendula.

A statistically significant increase in cytotoxicity was calculated between unwounded and wounded cells, which received no treatment (P = 0.03) showing that a wound environment was success fully induced. Furthermore, a statistically significant decrease in cytotoxicity was measured between wounded cells receiving no treatment (+W -C -I) and wounded HSF cells receiving laser irradiation followed by the administration of Calendula officinalis 3 CH (+W +1 +C) (P = 0.025).

Discussion

The stimulatory effects of LLLT at the cellular and molecular levels have been shown in many studies (36), including in our own laboratory.(37) Laser light affects the mitochondrial respiratory chain by changing the electrical potential of cell membranes and, consequently, their selective permeability for sodium, potassium and calcium ions, or by increasing the activity of certain enzymes such as cytchrome oxidase and adenosine triphosphate.(38) It also increases DNA synthesis,(39) collagen and pro-collagen production, (40,41) and may increase the cell proliferation (42) or alter locomotor characteristics of cells. (43)

Although studies in both laboratory conditions and clinical settings have been numerous, the biochemical reactions induced by low level laser irradiation are still poorly understood.

Cordova et al.,(44) conducted a study with the aim to investigate the relationship between the beneficial properties of Calendula officinalis and its antioxidant properties. The butanolic fraction (BF) was studied because it is non-cytotoxic and is rich in a variety of bioactive metabolites including flavonoids and terpenoids. The results obtained suggest the BF of Calendula officinalis possesses a significant free radical scavenging and antioxidant activity and that the proposed therapeutic efficacy of this plant could be due, in part, to these properties.

After analysis of the results there was an indication that Calendula officinalis 3 CH can facilitate wound healing on wounded HSF cells in vitro in the following two respects - it accelerates wound closure and increases cell viability. Although no statistically significant difference was measured between the control group and the cells treated with Calendula officinalis 3 CH, the cell viability and cytotoxicity findings are encouraging. The lack of decrease in cell viability or increase in cytotoxicity despite wound induction and Calendula officinalis application may be indicative of the beneficial effect that this treatment may have on wounded HSF cells in vitro. Furthermore, minor changes evident on a cellular level may be more significant at a systemic level.

Laser treatment on its own showed increased ATP and a decrease in cytotoxicity when compared to wounded fibroblasts without treatment. Furthermore, cell migration for wounded fibroblasts treated with laser irradiation showed an increase in migration when compared to wounded cells without treatment. In conclusion, the cellular responses (migration and cell viability) of wounded fibroblasts appeared to be stimulated by laser irradiation when the responses were compared to wounded cells receiving no treatment. Wounded cells which received no treatment (+W -C -I) showed a decrease in trypan blue, decrease in ATP cell viability and increase in LDH cytotoxicity when compared to unwounded untreated cells (+W -C -I) confirming that a central scratch successfully induced a wound environment.

Wounded HSF cells treated with Calendula officinalis 3 CH followed by laser irradiation proved to be the least favourable treatment. It seemed to lower cytotoxicity and early cell death. A suggested explanation would be that the laser irradiation might alter the chemical characteristics/bonds of Calendula officinalis 3 CH dispensed in 5% ethanol consequently having a damaging effect on the cells and leading to lower cell viability.

Laser irradiation followed by the administration of Calendula officinalis 3 CH proved to be the better combination treatment procedure. It showed superior results with regard to cell morphology, cell viability and cytotoxicity when compared to the Calendula officinalis 3 CH followed by irradiation treatment group. Laser irradiation followed by Calendula officinalis 3 CH demonstrated a normalising effect on wounded HSF cells. Results were comparable to unwounded cells thus designating this treatment procedure to be the most favourable.
Conclusion

From this study cellular effects of LLLT and Calendula officinalis 3 CH on normal and wounded skin fibroblasts could be detected. Morphological changes indicate that a double treatment application of Calendula officinalis 3 CH increase wound closure on wounded HSF cells in vitro. Laser irradiation too showed morphological signs of increased wound closure for HSF cells after a double exposure. However, a synergistic relationship, based on morphology, was not established when combining the two treatments.

Although only a few statistically significant differences were obtained between the measurement tools and any of the treatment protocols, it is important and worth noting that small differences on the cellular level might have a significant influence at the systemic level. Changes experienced at the cellular level might be amplified at the systemic level because of cell-cell interactions that may elicit a cascade of events.

References


