Research Article

Changes in Cell Viability of Wounded Fibroblasts following Laser Irradiation in Broad-Spectrum or Infrared Light

Denise Hawkins and Heidi Abrahamse

Laser Research Group, Faculty of Health, University of Johannesburg, P.O. Box 17011, Doornfontein 2028, Johannesburg, South Africa

Received 29 June 2006; Revised 14 August 2006; Accepted 22 August 2006

Recommended by Frederik Claeyssens

Objective. This study aimed to establish if broad-spectrum or infrared (IR) light in combination with laser therapy can assist phototherapy to improve the cell function of wounded cells. Background. The effect of laser light may be partly or completely reduced by broad-spectrum light. Methods. Wounded human skin fibroblasts were irradiated with 5 J/cm² using a helium-neon laser, a diode laser, or an Nd:YAG laser in the dark, in the light, or in IR. Changes in cell viability were evaluated by cell morphology, ATP cell viability, LDH membrane integrity, and caspase 3/7 as an early marker of apoptosis. Results. Wounded cells exposed to 5 J/cm² using 632.8 nm in the dark or 830 nm in the light or 1064 nm in the dark showed an increase in ATP viability, an increase in cytokine expression, and a decrease in LDH cytotoxicity indicating that the metabolic activity of the wounded cells was stimulated. Conclusion. Wounded cells irradiated in IR light showed an undesirable thermal effect that was proportional to the duration of exposure.

Copyright © 2007 D. Hawkins and H. Abrahamse. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

1. INTRODUCTION

Light is considered part of the spectrum of electromagnetic energy. This spectrum includes wavelengths of energy that vary considerably in length from the very short waves in the ultraviolet end of the spectrum (gamma and X-rays) to the very long waves at the infrared extreme of the spectrum [1]. In the visible portion of the spectrum, light is composed of wavelengths that range from 400 nm to 800 nm. Tissue is generally more transparent to near-IR wavelengths (800–1200 nm) than to visible light [1].

Properties of laser light

(i) Laser light is collimated, meaning that the beam travels in a single direction without divergence over a long distance with a high intensity. Ordinary light scatters rather quickly and does not provide illumination over any real distance [1].

(ii) Laser light is monochromatic, with all waves having the same wavelength or frequency. By generating light of one specific wavelength, lasers can be more specific in their application. In medical lasers, the laser effect is dependent upon targeted tissue’s inherent sensitivity to a particular wavelength. Ordinary light is composed of many different wavelengths or colors [1].

(iii) Laser light is coherent, with all waves in phase. The peaks and troughs of each wave are synchronous in space. Normal light is composed of light traveling in many different directions and out of phase [1].

A laser, whether large or small, always includes the following parts: an energy source (power supply), a lasing or amplifying medium (solid, liquid, or gas), and a resonating cavity (mirrors). Neodinium:yttrium-aluminum-garnet (Nd:YAG) lasers have short wavelengths (1064 nm and 1440 nm) within the infrared (invisible) zone of the spectrum. Neodinium’s wavelength is relatively poorly absorbed by water (and thus most relevant to musculoskeletal tissues), and it therefore penetrates tissue for relatively greater-than-desired depths, estimated at 4–6 mm in a vascular tissue [1, 2]. When less is absorbed by target tissue, more laser energy is transmitted beyond the target. This unintended effect can lead to inadvertent heating of deeper tissues and clinically significant thermal damage. Reports of thermal necrosis caused by subchondral bone penetration soon followed the clinical introduction of the Nd:YAG laser and signified its rapid efficiency demise [1, 2].
Semiconductor lasers are small, potentially inexpensive, and much more efficient than traditional lasers. Diode lasers generate low powers generally below 1.0 W. The diode laser is an example of a semiconductor laser used in ophthalmology for photocoagulation. Fibers easily transmit diode laser radiation, which can be pulsed or continuous [2].

The helium-neon (HeNe) laser emits red laser light at a wavelength of 632.8 nm and is within the visible spectrum. HeNe lasers only operate in continuous-wave (CW) mode and no cooling system is required. The maximum power output (50–60 mW) is so weak that these are not useful as surgical lasers [2, 3]. The laser can be used as a visible guide beam that accompanies otherwise invisible treating laser beams. HeNe laser light can penetrate as far as 0.5–50 mm into freshly excised human skin and delivers the highest relative percentage of incident energy to a certain volume of tissue. A penetration depth of even some microns can be regarded as sufficient, because most of the relevant target cells of low-level laser irradiation, namely fibroblasts, keratinocytes, macrophages, and endothelial cells for the induction of wound healing, are located within the epidermis and upper dermis [4]. The HeNe laser has been reported to stimulate wound healing and cell growth [4] and is frequently used for biostimulation in pain management and for wound healing [2].

Cell cultures that are first irradiated with laser light and have consequently exhibited biological effects are then irradiated with broad-spectrum (nonmonochromatic and incoherent) light, subsequently have their laser-produced biological effects reduced to almost nothing [5]. Karu et al. [5] demonstrated stimulating biological effects in cell cultures from monochromatic incoherent light and indicated that there are more mechanisms at work than simply the excitation of polarization-sensitive chromophores [5]. The influence of ambient light may influence the outcome of both clinical and laboratory studies [5] and may well explain some of the variability in the results/effects reported in the literature.

Experiments following phototherapy have shown that immediate heat rise in target tissue is negligible (±1°C). Stadler et al. [6] used a mouse model to demonstrate that black skin (4.5 ± 0.15°C) exhibits a greater temperature change relative to white skin (0.3 ± 0.10°C), thus failure to consider this variable can result in thermal damage and unwanted thermal influences during phototherapy [6]. Greppi [7] investigated the use of an 810 nm wavelength diode laser for hair removal using low-energy settings and long pulse duration and concluded that diode lasers can be used on dark pigmented skin with positive outcomes [7]. Complications such as hypopigmentation, or hyperpigmentation do occur, but all are temporary and resolve within a few months. The 810 nm wavelength diode laser can safely and successfully treat all skin types, including dark pigmented skin [7]. Visible and infrared irradiations have a pronounced wound healing effect, which develops as a consequence of local and systemic light effects, but many aspects of their mechanism are still unclear [8].

Kubota [9] investigated the effect of defocused gallium-aluminium-arsenide (GaAlAs) diode laser therapy (830 nm, CW, 669 mW/cm², 6.3 J/cm² to 21 J/cm²) in the treatment of unresponsive skin ulcers and reported that 830 nm diode laser therapy was well tolerated, and was very effective in the treatment of compromised skin ulcers of different etiologies in a large range of patient ages [9]. Meffert et al. [10] studied the systemic effect of ultraviolet, visible, and infrared radiation in whole body irradiation using experimental conditions similar to those used in phototherapeutical practice [10]. The study concluded that there were no negative side effects after the application of large doses of visible or infrared radiation and considerable systemic responses were exclusively found with ultraviolet irradiation [10]. Belotsky et al. [11] studied the effect of visible-light irradiation of 5 J/cm² and 12 J/cm² applied to buffy coat leukocytes [11]. The study found an inhibitory effect dependent on the light intensity that was more pronounced at 12 J/cm².

Little information is available regarding the potential thermal effects of near-infrared photoradiation during low-level laser therapy (LLLT) [12]. Lanzafame et al. [12] studied the effect of temperature-controlled 830 nm LLLT on experimental pressure ulcers created in C57BL mice and reported that the group receiving laser therapy (830 nm 5 J/cm², CW) showed a maximum temperature change during therapy of 2.0 ± 0.64°C and complete wound closure at 18 ± 4 days while the group receiving incandescent light (5 J/cm²) showed a maximum temperature change during therapy of 3.54 ± 0.72°C and complete wound closure at 25 ± 6 days [12]. The results also showed that the group receiving incandescent light had a 36.3 ± 4.8% wound closure at day 14 while the group receiving laser therapy showed a significantly higher rate of wound closure at 75.4 ± 7.2% [12].

Maegawa et al. [13] studied the relationship between temperature increase and enhanced microcirculation during near-infrared irradiation and found a 0.8–1.0°C increase in temperature and enhanced microcirculation in exteriorized rat mesentery during irradiation at 830 nm with a 38 mW power output [13].

There are few studies that investigate the benefit or detriment of combining laser irradiation with broad-spectrum (visible white light) or infrared light (ExoTerra Heat Glo, 240 V, 100 W). This study aimed to establish if broad-spectrum or infrared light has any benefit when irradiating wounded human skin fibroblast with a dose of 5 J/cm² using an HeNe (632.8 nm, 18.8 mW) laser, a diode (830 nm, 65 mW) laser, or an Nd:YAG (1064 nm, 1 W). Wounded cells irradiated in IR light demonstrate an undesirable thermal effect that is proportional to the duration or length of exposure. Wounded cells responded with an increase in cell function and viability when exposed to 5 J/cm² using 632.8 nm in the dark, 830 nm in the light, or 1064 nm in the dark. Results indicate that 5 J/cm² using 632.8 nm in the dark or 830 nm in the light are the most effective doses, which may ultimately stimulate wounded cells to accelerate or improve the rate of wound healing.
2. MATERIALS AND METHODS

2.1. Cell culture procedure

Human skin fibroblast monolayer cultures (ATCC CRL1502 WS1) were grown in Eagle’s minimal essential medium with Earle’s balanced salt solution that was modified to contain 2 mM L-glutamine, 1.0 mM sodium pyruvate, 0.1 mM nonessential amino acids, 1% fungizone, and 1% penicillin-streptomycin and supplemented with 10% V/V foetal bovine serum (serum rich medium). The cultures were incubated at 37 °C with 5% CO₂ and 85% humidity [14]. Cells were trypsinized using a 0.25% (w/v) trypsin-0.03% EDTA solution in Hanks balanced salt solution (HBSS) and approximately 6.5 × 10⁵ cells (in 3 mL culture medium) were seeded in 3.4 cm diameter culture plates and incubated overnight to allow the cells to attach [15].

2.2. Laser specifications and exposure regime

Once the fibroblasts had attached, 2 mL of culture medium was removed and a wound was induced before the cells were irradiated. Normal and wounded fibroblasts were exposed to 5 J/cm² on day 1 and day 4. Since the HeNe laser has a power density between 2.07 mW/cm² and a spot size of 3.4 cm, the light is divergent and is not as harmful as a narrow parallel beam that allows the entire volume of intense light to be focused, or concentrated on one small area [16]. For the simulated wound environment, confluent monolayers were first scratched with a sterile pipette of 2 mm diameter and the plates were incubated at 37 °C for 30 minutes before they were irradiated [17–19]. Each scratch was irregular and the size of the wounds ranged from 1 to 2 mm in diameter [17–19].

Wounded cells were exposed to 5 J/cm² either in the dark or in broad-spectrum white light or in infrared light (Exo Terra Heat Glo, 240 V, 100 W, 17.5 cm above cells). Irradiations were performed with an HeNe laser (632.8 nm, 18.8 mW power output, 2.07 mW/cm² power density, and 3.4 cm diameter spot size or area 9.1 cm²), a semiconductor diode laser (830 nm, 54 mW power output, 5.95 mW/cm² power density, and 3.4 cm diameter spot size or area 9.1 cm²), and an Nd:YAG laser (1064 nm, 1 W power output, 12.73 mW/cm² power density, and 10 cm diameter spot size (to accommodate four culture dishes simultaneously) or area 78.5 cm²) using the following exposures:

(i) wounded unirradiated control cells (0 J/cm²);
(ii) wounded human skin fibroblasts exposed to 5 J/cm².

Using an average laser power density of 2.07 mW/cm², the duration of each exposure for the HeNe laser was calculated at 41 minutes and 40 seconds for 5 J/cm². Using an average laser power density of 5.95 mW/cm², the duration of each exposure for the 830 nm diode laser was calculated at 12 minutes and 45 seconds while using an average laser power density of 12.73 mW/cm² for the Nd:YAG, the duration of the exposure was 6 minutes and 30 seconds for 5 J/cm². The cellular response measurements (biological assays) were made 24 hours (day 5) after laser irradiation to observe the modifications or responses that occur with time [20].

2.3. Biological assays

Following laser irradiation, the fibroblasts were trypsinized from the 3.4 cm culture dishes and the cell suspension (1 × 10⁵ cells/100 μL) was used to assess the effect of laser irradiation on cell viability using cell morphology, the ATP luminescent viability assay, and caspase 3/7 activity. The culture medium was used to assess damage or additional stress caused by the irradiation using the LDH membrane integrity assay.

Changes in cell morphology

The control (unirradiated or 0 J/cm²) and wounded fibroblast behavior were observed using an inverted microscope. The number and intensity of colony formation, the haptotaxis (direction or orientation) of the edge fibroblasts, the number of fibroblasts present in the centre of the scratch, and chemotaxis-chemokinesis (movement or migration of cells across the central scratch) were evaluated to determine the activity of fibroblasts [21].

ATP cell viability assay

The CellTiter-Glo luminescent cell viability assay (Promega G7570) is based on the quantitation of ATP present, which signals the presence of metabolically active cells or viable cells [22]. An equal volume of reconstituted CellTiter-Glo reagent was added to 50 μL of cell suspension (1 × 10⁵ cells/100 μL). The contents were mixed on an orbital shaker for 2 minutes to induce cell lysis and were then incubated at room temperature for 10 minutes to stabilize the luminescent signal, which was recorded using a Berthold EG and Junior luminometer [22, 23].

LDH membrane integrity

The CytoTox 96 nonradioactive cytotoxicity assay (Promega G1780) measures lactate dehydrogenase (LDH), a stable cytosolic enzyme that is released upon cell lysis [24]. Hundred μL of culture medium were removed from each plate after each irradiation. Fifty μL of the culture medium were mixed with an equal volume of reconstituted substrate mix. The plate was covered with foil and incubated at room temperature for 30 minutes. Fifty μL of stop solution were added and the absorbance was read at 490 nm [24].

Apoptosis

The caspase-Glo 3/7 (Promega G8090) assay is a luminescent assay that measures caspases −3 and −7 activity. Apoptosis can be induced by stress from growth factor withdrawal or irradiation and is mediated by a cascade of highly specific proteases known as caspases [25]. Caspases 3, 6, and 7 are effector caspases that cleave cellular substrates and
precipitate apoptotic death [26]. The addition of caspase-Glo 3/7 reagent results in cell lysis, followed by caspase cleavage of the substrate and generation of a luminescent signal produced by luciferase. Twenty-five μL of the cell suspension (1 × 10^5 cells/100 μL) were added to 25 μL of caspase-Glo 3/7 reagent and the tube was mixed and incubated at room temperature for 3 hours. The luminescent signal was recorded which is directly proportional to the amount of caspase activity present [25].

Statistical analysis

Each experiment was repeated on different populations of wounded fibroblast cells between passage 13–31. Each biological assay was performed in duplicate and the average of the two results was used to obtain a final sample number of n = 4. The results were recorded for statistical analysis using SigmaPlot 8.0 and the mean, standard deviation, standard error, and significant percentage change between the un-irradiated control (0 J/cm²) and the irradiated cells were calculated for each data group and graphically represented. The Student t test and one-way analysis of variance (ANOVA) were used to analyze the difference firstly between the un-irradiated control (0 J/cm²) and the irradiated cells and secondly between the different data groups. Statistical significance was accepted at the 0.05 level (95% confidence interval). The ± value or error bars in the figures indicate standard error of the mean (SEM).

3. RESULTS

Fibroblast behavior was observed using an inverted microscope and changes in cell morphology were digitally recorded daily. Normal human skin fibroblasts are long slender cells that grow in monolayer sheets –×40 magnification. For the simulated wound environment, confluent monolayers were first scratched with a sterile pipette of 2 mm diameter and the plates were incubated at 37°C for 30 minutes before they were irradiated. A clear central scratch with a distinct wound margin (wm) could be observed directly after inducing a mechanical scratch [17–19]. Morphological changes that were assessed included haptotaxis (h) or change in orientation of the edge fibroblasts, chemotaxis-chemokinesis or migration (m) of cells across the central scratch, colony formation along the wound margin, and number of fibroblasts present in the central scratch (f) –×10 magnification. Wounded irradiated cells showed evidence of haptotaxis and migration of fibroblasts across the central scratch in an attempt to close the wound (Figure 1).

ATP cell viability assay

The ATP luminescent assay was used to determine the effect of laser irradiation on the viability of wounded fibroblasts. The measurements were made 24 hours post-laser irradiation (Figure 2). Wounded cells exposed to 5 J/cm² using 632.8 nm in the light showed a decrease in the ATP viability when compared to wounded cells exposed to 5 J/cm² in the dark (P = 0.022) indicating that cells irradiated with 5 J/cm² in broad-spectrum (nonmonochromatic and incoherent) light exhibit a decrease in biological response to the laser light when compared to cells irradiated with the same dose in the dark [5]. This suggests that there are more mechanisms at work than simply the excitation of polarization-sensitive chromophores [5].

Wounded cells exposed to 5 J/cm² using 632.8 nm in the dark, light, and IR showed a decrease in ATP viability of −3.789% (P = 0.587), −12.892% (P = 0.192), and −24.972% (P = 0.078), respectively, when compared to the wounded unirradiated control. Wounded cells exposed to 5 J/cm² using 830 nm in the dark, light, and IR showed a decrease in ATP viability of −5.341% (P = 0.626), −2.694% (P = 0.785), and −12.354% (P = 0.417), respectively, when compared to the wounded unirradiated control. Wounded cells exposed to 5 J/cm² using 830 nm in IR showed a decrease in the ATP viability when compared to wounded cells exposed to 5 J/cm² in the light (P = 0.053). Wounded cells exposed to 5 J/cm² using 632.8 nm in the light showed a decrease in ATP viability when compared to wounded cells exposed to 5 J/cm² using 830 nm in the light (P = 0.002). Wounded cells exposed to 5 J/cm² using 830 nm in the dark showed a decrease in ATP viability when compared to wounded cells exposed to 5 J/cm² using 632.8 nm in the dark (P = 0.048) indicating that the biological response is dependent on the wavelength. The results indicate that both the wavelength (632.8 nm, 830 nm, or 1064 nm) and irradiation conditions (light, dark, or IR) contribute to different biological responses of wounded cells.

Wounded cells exposed to 5 J/cm² using 1064 nm in the dark, light, and IR showed a decrease in ATP viability of −7.778% (P = 0.801), −10.436% (P = 0.520), and −4.121% (P = 0.673), respectively, when compared to the wounded unirradiated control. Wounded cells exposed to 5 J/cm² using 1064 nm in the light showed a decrease in ATP viability when compared to wounded cells exposed to 5 J/cm² using 1064 nm in IR (P = 0.071). Wounded cells exposed to 5 J/cm² using 1064 nm in the dark showed a decrease in ATP viability when compared to wounded cells exposed to 5 J/cm² using 1064 nm in IR (P = 0.128).

Wounded cells exposed to 5 J/cm² using 1064 nm in the light showed a decrease in ATP viability when compared to wounded cells exposed to 5 J/cm² using 830 nm in the light (P = 0.050). Wounded cells exposed to 5 J/cm² using 830 nm in IR showed a decrease in ATP viability when compared to wounded cells exposed to 5 J/cm² using 1064 nm in IR (P = 0.074). Wounded cells exposed to 5 J/cm² using 632.8 nm in IR showed a decrease in ATP viability when compared to wounded cells exposed to 5 J/cm² using 1064 nm in IR (P = 0.018).

LDH membrane integrity assay

Changes in cell membrane integrity were measured using LDH membrane integrity assay since LDH is a stable cytosolic enzyme that is released upon membrane damage [24]. Wounded cells exposed to 5 J/cm² using 632.8 nm in IR
Fibroblast behavior was observed using an inverted microscope and changes in cell morphology were digitally recorded daily. Normal human skin fibroblasts are long slender cells that grow in monolayer sheets –×40 magnification (a). For the simulated wound environment, confluent monolayers were first scratched with a sterile pipette of 2 mm diameter and the plates were incubated at 37°C for 30 minutes before they were irradiated (b). A clear central scratch with a distinct wound margin (wm) could be observed directly after inducing a mechanical scratch (c). Morphological changes that were assessed included haptotaxis (h) or change in orientation of the edge fibroblasts, chemotaxis-chemokinesis or migration (m) of cells across the central scratch, colony formation along the wound margin (wm), and number of fibroblasts (f) present in the central scratch –×10 magnification (d).

showed an increase in cytotoxicity when compared to the unirradiated control \((P = 0.010)\) and when compared to wounded cells exposed to 5 J/cm² using 632.8 nm in the dark \((P = 0.950)\) and in the light \((P = 0.476)\). The result supports the ATP viability, which suggests that cells irradiated in broad-spectrum light exhibit a decrease in biological response (Figure 2).

Wounded cells exposed to 5 J/cm² using 830 nm in IR showed an increase in cytotoxicity when compared to wounded cells exposed to 5 J/cm² using 830 nm in the dark \((P = 0.008)\) and in the light \((P = 0.0003)\). The result corresponds to the ATP viability result, which suggests that cells irradiated in IR light exhibit a decrease in biological response. Wounded cells exposed to 5 J/cm² using 632.8 nm in IR showed an increase in cytotoxicity when compared to wounded cells exposed to 5 J/cm² using 830 nm in IR \((P = 0.082)\) indicating that the extended irradiation time results in an undesirable thermal effect that causes an increase in cytotoxicity and decrease in cell viability. The result show that there is a statistical difference between cells irradiated with 632.8 nm and 830 nm in IR indicating that either the wavelength or that the irradiation time influences the total cellular response.

Wounded cells exposed to 5 J/cm² using 1064 nm in IR showed an increase in cytotoxicity when compared to wounded cells exposed to 5 J/cm² using 632.8 nm in the dark \((P = 0.082)\) and in the light \((P = 0.109)\). Wounded cells exposed to 5 J/cm² using 632.8 nm in the dark \((P = 0.074)\) and when compared to wounded cells exposed to 5 J/cm² using 830 nm in the dark \((P = 0.0003)\). Wounded cells exposed to 5 J/cm² using 1064 nm in the dark showed a decrease in cytotoxicity when compared to wounded cells exposed to 5 J/cm² using 1064 nm in the dark \((P = 0.063)\). There was no statistical difference between wounded cells exposed to 5 J/cm² using 632.8 nm and 830 nm in the light \((P = 0.141)\) and between wounded cells exposed to 5 J/cm² using 1064 nm and 632.8 nm in the light \((P = 0.975)\),
however there is a difference between wounded cells exposed to 5 J/cm² using 1064 nm and 830 nm in the light ($P = 0.013$). Results show that there was no statistical difference between wounded cells irradiated with 5 J/cm² using 632.8 nm or 830 nm in the dark or the light, however wounded cells exposed to 5 J/cm² using 1064 nm in the dark show an increase in the cytotoxicity when compared to cells irradiated in the light ($P = 0.0004$).

**Apoptosis**

The caspase 3/7 luminescent assay was used to identify if the irradiation caused additional stress which may induce higher levels of apoptosis (Figure 3). Wounded cells exposed to 5 J/cm² using 632.8 nm in IR showed an increase in caspase 3/7 activity when compared to the wounded unirradiated control ($P = 0.364$) and when compared to wounded cells exposed to 5 J/cm² using 632.8 nm in the dark ($P = 0.0007$) or in the light ($P = 0.00003$) indicating additional stress caused by IR light. Wounded cells exposed to 5 J/cm² using 632.8 nm in light showed decrease in caspase 3/7 activity when compared to the wounded un-irradiated control ($P = 0.433$) and when compared to cells exposed to 5 J/cm² using 632.8 nm in the dark ($P = 0.069$).

Wounded cells exposed to 5 J/cm² using 830 nm in IR showed an increase in caspase 3/7 activity when compared to wounded cells exposed to 5 J/cm² using 830 nm in the dark ($P = 0.084$) or in the light ($P = 0.052$) indicating additional stress caused by IR light. Wounded cells exposed to 5 J/cm² using 830 nm in light showed decrease in caspase 3/7 activity when compared to the wounded unirradiated control ($P = 0.880$) and when compared to wounded cells exposed to 5 J/cm² using 632.8 nm in the dark ($P = 0.931$). Wounded cells exposed to 5 J/cm² using 1064 nm in IR showed a decrease in caspase 3/7 activity when compared to the wounded unirradiated control ($P = 0.852$) and when compared to wounded cells exposed to 5 J/cm² using 1064 nm in the dark ($P = 0.979$) or in the light ($P = 0.406$) indicating a limited change in cell stress that may be related to the duration (6 minutes and 30 seconds) of the exposure to laser light and IR. Wounded cells exposed to 5 J/cm² using 1064 nm in the light showed an increase in caspase 3/7 activity when compared to wounded cells exposed to 5 J/cm² using 632.8 nm in the light ($P = 0.0179$) indicating that the wavelength and treatment condition (in the light) may influence the biological response of the cells.

Wounded cells exposed to 5 J/cm² using 830 nm in IR showed an increase in caspase 3/7 activity when compared to wounded cells exposed to 5 J/cm² using 1064 nm in IR ($P = 0.0038$). Wounded cells exposed to 5 J/cm² using 1064 nm in IR showed a decrease in caspase 3/7 activity when compared to wounded cells exposed to 5 J/cm² using 632.8 nm in IR.
was detected using caspase 3/7 luminescence. Wounded cells exposed to 5 J/cm² using 830 nm in IR showed an increase in caspase 3/7 activity when compared to the wounded unirradiated control (P = 0.001), however there was no change in activity between wounded cells exposed to 5 J/cm² using 830 nm in IR and wounded cells exposed to 5 J/cm² using 632.8 nm in IR (P = 0.687).

4. DISCUSSION

Karu [27] stated that the laser effect depends on the radiation wavelength, dose, and intensity, as well as on the cell culture conditions [27]. There are biological limits to the effects of LLLT: the proliferation of fast-growing cells cannot be stimulated, or not all cellular functions can be activated. Not all cells in tissues or cellular cultures will respond to irradiation in exactly the same way. The total response from the cells may represent the average rather than the true value, which has little meaning in the clinic but is important in a cell study. Many experiments have been performed in vitro and the reaction seen or not seen in an in vitro experiment reflects the effect of laser therapy on a single isolated cell [28]. In the clinic, there is no single-cell effect, instead a very complex and multipath cascade of processes that influences every cell in the body at each particular moment.

Karu et al. [5] has shown that the effect of laser light may be partly or completely reduced by broad-spectrum light [5]. When wounded or scratched, cell monolayers respond to the disruption of cell-cell contacts with an increased concentration of growth factors at the wound margin and by healing the wound through a combination of proliferation and migration [29–31]. These processes reflect the behavior of individual cells as well as the properties of the cell sheet as a surrogate tissue.

The ATP cell viability results show that wounded cells exposed to 5 J/cm² using 632.8 nm in IR showed a decrease in ATP viability indicating an increase in cellular damage possibly caused by an increase in culture temperature since the cells were irradiated for a longer period of 41 minutes and 40 seconds when compared to 830 nm (12 minutes and 45 seconds) and 1064 nm (6 minutes and 30 seconds). Wounded cells exposed to 5 J/cm² using 830 nm in IR showed a decrease in ATP viability supporting the evidence that IR may cause thermal damage or unwanted thermal influences during LLLT. Wounded cells exposed to 5 J/cm² using 1064 nm in the light and dark showed a decrease in ATP viability when compared to cells irradiated in IR indicating that the IR light may support the 1064 nm near-infrared laser light or that the cells are incubated for a minimal period of 6 minutes and 30 seconds limiting any undesirable thermal effects. Results show that there was a statistical difference between wounded cells exposed to 5 J/cm² using 632.8 nm, 830 nm, and 1064 nm in the light indicating that the cellular response is dependent on the wavelength. The similar response was demonstrated for wounded cells exposed to 5 J/cm² in the dark and IR supporting the evidence that the dose, treatment condition, and wavelength influence the average response of the cells (n = 4).
the dark, therefore it appears that the response is influenced by a change in temperature, dose, and wavelength.

The LDH cytotoxicity showed that wounded cells had an increase in membrane damage after they were exposed to 5 J/cm² using 632.8 nm and 830 nm in IR. The results correspond to the decrease in ATP cell viability that was observed indicating that IR causes thermal damage. The results show that wounded cells exposed to 5 J/cm² using 1064 nm in IR did not result in a significant change in the cytotoxicity when compared to wounded cells exposed to 5 J/cm² in the light. This result corresponds to an increase in the ATP viability for 5 J/cm² using 1064 nm in IR indicating that IR light may support and assist laser irradiation provided the duration of the irradiation is short since longer durations appear to cause undesirable thermal damage. The LDH results show that wounded cells exposed to 5 J/cm² using 632.8 nm and 830 nm in the light are not statistically different from cells exposed in the dark, however there is a difference between cells exposed to 5 J/cm² using 1064 nm in the light or dark supporting the evidence that ambient or broad-spectrum light reduces the biological effect and indicates that there are more mechanisms at work than simply the excitation of polarization-sensitive chromophores [5].

Results show that wounded cells exposed to 5 J/cm² using 632.8 nm in the light or IR have an increase in cytotoxicity when compared to wounded cells exposed to 5 J/cm² using 830 nm in the light or IR supporting the evidence that the wavelength is an important parameter that can influence the cellular response to laser irradiation. Wounded cells exposed to 5 J/cm² using 1064 nm in the dark or light show a difference in LDH cytotoxicity when compared to wounded cells exposed to 5 J/cm² using 830 nm in the dark or light supporting the evidence that the response depends on the wavelength. Results show that wounded cells exposed to 5 J/cm² using 632.8 nm and 830 nm in IR have an increase in cytotoxicity when compared to wounded cells exposed to 5 J/cm² using 1064 nm in IR suggesting that the increase in irradiation time of 41 minutes and 40 seconds for 632.8 nm and 12 minutes and 45 seconds for 830 nm compared to 6 minutes and 30 seconds for 1064 nm may result in thermal damage or unwanted thermal influences from the IR light. Wounded cells exposed to 5 J/cm² using 1064 nm in IR showed a similar biological response to wounded cells exposed to 5 J/cm² using 632.8 nm in the dark, however there was an increase in LDH cytotoxicity for 1064 nm which may indicate more damage to the cells. Wounded cells exposed to 5 J/cm² using 830 nm in light showed a similar biological response to wounded cells exposed to 5 J/cm² using 632.8 nm in the dark indicating an alternative wavelength and treatment condition that results in a similar response.

Wounded cells exposed to 5 J/cm² using 632.8 nm or 830 nm in the light showed a decrease in caspase 3/7 activity which did not correspond with the LDH cytotoxicity results. Wounded cells exposed to 5 J/cm² using 1064 nm in the light showed an increase in caspase 3/7 activity that corresponds to the ATP viability and LDH cytotoxicity. These results indicate that ambient or broad-spectrum light changes the biological response of wounded cells to laser irradiation. The caspase 3/7 activity increased for wounded cells that were exposed to 5 J/cm² using 632.8 nm (41 minutes and 40 seconds) or 830 nm (12 minutes and 45 seconds) in IR light when compared to the activity for 1064 nm (6 minutes and 30 seconds). The results suggest that IR light in combination with laser irradiation may only be of benefit if the duration of the exposure is minimal. The results indicate that the undesirable thermal effect of IR light is dependent on and directly proportional to the duration or length of exposure.

The results indicate that wounded cells exposed to 5 J/cm² using 1064 nm do not show statistical variations in cell stress when irradiated in the dark, in the light, or in IR, however there are statistical variations in ATP viability and LDH cytotoxicity. The results show that there are statistical variations between the different wavelengths when wounded cells are exposed to 5 J/cm² using 632.8 nm, 830 nm, and 1064 nm in the light and IR, however there are no statistical variations between the different wavelengths when wounded cells are exposed in the dark. The result suggests that the wavelength and treatment conditions are important parameters that influence the biological response. The results also support the evidence that broad-spectrum light changes the biological response [5]. Karu and Kolyakov [32] reported that the wavelength range important for phototherapy (600–860 nm) has four active regions, but peak positions are not exactly the same for all spectra. The peak positions are between 613.5 and 623.5 nm in the red maximum and two near-infrared maxima have peak positions in the range 750.7–772.3 nm and 812.5–846 nm, respectively [32]. This may explain why broad-spectrum light changes the biological response since it may influence the peak active region of each wavelength.

The results using the ATP viability, LDH cytotoxicity, and caspase 3/7 activity show the following: 5 J/cm² using 632.8 nm in the dark is more effective than 830 nm in the dark; 632.8 nm in the dark is more effective than 1064 nm in the dark, and 1064 nm in the dark is more effective than 830 nm. Results also show that 5 J/cm² using 830 nm in the light is more effective than 632.8 nm in the light but not as effective as 632.8 nm in the dark; 632.8 nm in the light is more effective than 1064 nm in the light but not as effective as 632.8 nm in the dark, and exposure to 830 nm in the light is more effective than 1064 nm in the light. Therefore the results indicate that 632.8 nm in the dark is the most effective.

5. CONCLUSION

Wounded cells respond with an increase in ATP viability and a decrease in LDH cytotoxicity when exposed to 5 J/cm² using 632.8 nm in the dark or 830 nm in the light. Results showed that only the increase in LDH cytotoxicity differentiated wounded cells exposed to 5 J/cm² using 1064 nm in IR from cells exposed to 1064 nm in the dark indicating that IR light can support laser irradiation with 1064 nm provided that the duration of exposure is minimal to limit the undesirable thermal effects.

Wounded cells exposed to 5 J/cm² using 632.8 nm in the light result in a decrease in caspase 3/7 activity when
compared to cells exposed in the dark. The results support findings from Hawkins and Abrahamse [33], which reported that 5 J/cm² using 632.8 nm stimulates mitochondrial activity, cell proliferation, and migration to accelerate wound closure [33]. However, wounded cells exposed to 5 J/cm² using 632.8 nm in IR show an increase in caspase 3/7, increase in cytotoxicity, and decrease in viability when compared to cells exposed in the dark or in the light indicating an undesirable thermal effect of IR light. Wounded cells respond to 5 J/cm² using 632.8 nm or 1064 nm in the dark with a decrease in LDH cytotoxicity and increase in ATP viability when compared to cells exposed in the dark indicating that broad-spectrum light decreases the biological response of cells to laser light. Wounded cells exposed to 5 J/cm² using 632.8 nm in IR show a statistical decrease in ATP viability demonstrating that IR light has a detrimental effect and does not assist phototherapy. Results show that the effect of IR light with 830 nm or 1064 nm laser light is dependent on the duration of the exposure.

Results from the study may be useful in defining laser parameters for clinical applications where clinicians may be able to use broad-spectrum or IR light in combination with phototherapy to improve the laser effect for specific applications. This in vitro study has shown that the effect of IR light is determined by the duration of the laser irradiation as may only be beneficial in combination with short laser exposures to avoid an undesirable thermal effect. It should be noted that this study concentrates on the effect of IR and broad-spectrum light in combination with phototherapy on cell culture and that the effect in an animal or human wound model may be different and may be dependent on other contributing factors such as blood flow, cytokine expression, inflammatory response, and endorphins (pain relief) to minimize the undesirable thermal effect. With further research, the use of IR light in combination with phototherapy may be useful in clinical applications such as pain relief and musculoskeletal injuries. The results support published findings by Enwemeka et al. [26] that revealed the positive effects of various wavelengths of laser light on tissue repair, with 632.8 nm having the highest treatment effect and 780 nm the least and concluded that phototherapy is a highly effective therapeutic armamentarium for tissue repair and pain relief [26].

To conclude, the results show that wounded cells responded optimally to 5 J/cm² using 632.8 nm in the dark, 830 nm in the light, and 1064 nm in the dark by maintaining cell viability, reducing cytotoxicity, stimulating the release of cytokines and growth factors to ultimately stimulate cell proliferation and cell growth. The results indicate that the exposures identified stimulate the metabolic activity of wounded cells, which may ultimately accelerate or improve the rate of wound healing.

ACKNOWLEDGMENTS

This project is jointly funded by the University of Johannesburg, National Laser Centre (NLC), National Research Foundation (NRF), and Medical Research Council (MRC).

REFERENCES


