Recent Advances of Low-Level Light Therapy: Fundamentals, Efficacy and Applications

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Submission: September 20, 2018; Published: September 28, 2018

Abstract

The mechanisms and fundamental of low-level light therapy (LLLT), or photobiomodulation are reviewed and compared with that of photodynamic therapy (PDT). However, LLLT combined with PS will enhance the efficacy, where the generation of singlet oxygen via the PS triplet state is more effective. The biomedical applications of LLLT including dermatology and plastic surgery, wound healings, nerve stimulation, dentistry, cancer therapy, and ophthalmic surgeries are reviewed, where lights (lasers or LED) spectra of UV (200-400) um, visible (400-700) nm, near-IR (700-2900) nm, and mid-IR (3-5) um are presented. Two important laws related to the fundamental and efficacy of the photo-biological systems: The Beer-Lambert law (BLL) and the Bunsen-Roscoe law (BRL) of reciprocity are discussed. LLLT efficacy may be improved by combined with photosensitizer-initiated reactive oxygen species (ROS) for light/heat sensitive ion channels.

Keywords: Lasers; Photobiomodulation; Photodynamic therapy; Low level light therapy

Introduction

Lights (lasers and LEDs) have been used for various medical procedures such as dermatology and cosmetic surgery, wound healings, nerve stimulation, dentistry ophthalmology applications for vision corrections and corneal deceases, and many other therapeutic procedures [1-3]. Combining the nanoparticles, diode lasers have been also used for cancer therapy, bio-sensing, bio-imaging, drug delivery, tissue engineering and diagnostics of cancer cell [4-7].

Lights in the near infrared (IR), wavelength of (750-1200) nm, have deeper tissue penetration depths than that of visible lasers. Therefore, near IR lasers are good candidates for procedures which need deep penetrations such as hair removal and nano-gold mediated cancer therapy [1]. On the other hand, visible lasers (430-680) nm with strong absorption in blood and color-dyes have been used for phototherapy of oral cancer, retina deceases and tattoo removal [2]. Mid-IR lasers (1.9-3.0) um and (9.3-10.6) um with strong absorption in water and tissue have been used for super surface procedures or ablative type procedures such as soft and hard tissue ablation. Other IR lasers (1.3-1.6) um have been used for so called minimally invasive procedures such as resurfacing due to their smaller tissue absorption than that of mid-IR lasers [1,2].

High power lights (in CW or pulsed mode) have been reviewed by Lin [1,6-9] for photothermal therapy (PTT) and photodynamic therapy (PDT), using the light spectra of UV (200-400) um, visible (400-700) nm, near-IR (700-2900) nm, and mid-IR (3-5) um having various penetration depths which define invasive and noninvasive procedures. In contrast to PTT (with high light intensity) and PDT (with photosensitizers), low level light therapy (LLLT), or photobiomodulation (PBM) has a much lower intensity between 5 to 500 mW/cm² and does not require photosensitizers (PS). However, LLLT combined with PS will enhance the efficacy, where the generation of singlet oxygen via the PS triplet state is more effective. In this article, we will review the recent progress of LLLT including its fundamental, mechanisms and applications. We will also discuss the efficacy of LLLT with PS and compare the differences and similarity between PDT and LLLT.

The fundamental

The basic definition and units: Lights are defined by their output "colors", i.e., wavelength (or spectrum) as follows:

- UVC (200-300nm), UVB (300-320nm), UVA (320-400) nm, visible (400-700) nm, near-IR (700-2900) nm, and mid-IR (3-5) um, where 1 um=1,000nm.

- The output of a light is defined by the following units:
  - Power (P) in mW (or W), 1W=1,000mW.
  - Intensity, or irradiance (I) in mW/cm² (power per unit area).
  - Energy (E) in mJ;
  - Fluence, or dose (F) in mJ/cm² (energy density).
  - Relations: E=It, P=I/area, F=E/area

- Laser-matter interaction: Light and tissue (or other media) interaction, in general, could be categorized into three processes:
  - (a) pure thermal, (b) non-thermal (or chemical), and (c) combined...
thermal and non-thermal effects. Lights can be reflected, absorbed, scattered or transparent to the matter. These processes are governed by not only the tissues (media) optical properties but also the laser parameters such as its wavelength, energy, intensity, pulse-width, repetition rate and the operation modes, continuous wave (CW) or pulsed mode. Various fiber structures for effective delivery of the laser energy to the treated areas are also critical in specific applications. Tissue penetration depth (d) of various lasers depends on their wavelength, or their tissue/water absorption coefficients (A), where A is inverse proportional to d (i.e. d=1/A). The absorption spectrum in water (or tissue) having 3 major peaks at 1.45um, 1.93um and 2.94nm, besides the high absorption of CO₂ lasers (9.6 to 10.6um).

LLLT is essentially a process of photobiomodulation (PBM) without thermal effects, and its does not require photosensitizer (PS), where the light interaction or stimulate biological photoreceptors located inside human cells via electronic transfer from the donors to the acceptors (such as oxygen), augmenting the production of a trans-membrane electric and mechanical proton-H⁺ gradient. Greater mechanisms of LLLT (and PBM) will be discussed later.

The biomedical applications

LLLT uses lasers or LEDs typically having power density (or intensity) between 5 to 500mW/cm² and wavelengths of visible (400-700nm), near-infrared (NIR) 780 to 1100nm; can be a continuous wave (CW) or a pulsed light consisting of a relatively low dose (fluence) (0.5 to 50J/cm²). The wavelength range between 700 and 780nm was ineffective as it matches with the absorption spectrum of cytochrome c oxidase. Red (630-670nm) and NIR lights are the most effective due to their larger penetration depth and lower scattering and absorption by tissue chromospheres.

LLLT is also known for biostimulatory and biomodulatory effects in vivo and in vitro and shows not a thermal but photochemical effect by triggering biochemical changes within cells after the application of a lights. The result of LLLT on human tissues is a reduction of inflammation, pain relief, hair growth, adjuvant-induced arthritis, skin and nail psoriasis, breastfeeding, and accelerated tissue regeneration [10-16]. In dentistry, the use of LLLT is to promote wound-healing, dentine repairing process and accelerated tissue regeneration [10-16]. In dentistry, the use of LLLT is to promote wound-healing, dentine repairing process and accelerated tissue regeneration [10-16].

The efficacy and mechanisms

The mechanisms: The kinetics and mechanism of PDT has been reported by Lin [7-9], in which both oxygen-mediated (type-II) and non-oxygen-mediated (type-I) are presented. LLLT has been known for almost 50 years, since the first reported article using a ruby laser (694nm) reported by McGuff et al. [10] (in 1965) for human malignant tumors, and by Mester et al. [11] (in 1968) for hair growth in mice [11]. However, the complete mechanisms of LLLT action remain elusive and was reviewed recently by Ferrares et al. [17] and Freitas & Hamblin [18]. It was reported [12] that LLLT can augment the oxidative phosphorylation (OXPHOS) modifying the redox basal-status of the whole cell and in particular of mitochondria via electrons transfer from a donor to oxidizing agent, oxygen. The free energy of these reduction-oxidation (redox) reactions is spent to synthetize chemical energy of the adenosine triphosphate (ATP). The mechanism of LLLT is based on the absorption of a specific visible red and near-infrared wavelengths by biological photoreceptors located inside human cells, augmenting the production of a trans-membrane electric and mechanical proton-H⁺ gradient in mitochondria necessary for OXPHOS, and enhancing the activity of mitochondrial complexes IV in an exposure-response relationship [12].

Two major hypothesis about LLLT mechanisms were reported [18]: (i) the photons dissociate nitric oxide from the enzyme causing the increase in electron transport, mitochondrial membrane potential and ATP production, and (ii) ion channels activated by light allowing calcium to enter the cell. The light-activated signaling pathways including the increase of reactive oxygen species (ROS), cyclic adenosine triphosphate, NO and Ca²⁺, leading to activation of transcription factors and the increased expression of protein synthesis, cell migration and proliferation, anti-inflammatory signaling, anti-apoptotic proteins, antioxidant enzymes etc. The transcription factors include, such as, nuclear factor kappa B, kappa-B ligand, hypoxia inducible factor, protein kinase B, forkhead box protein M1, peroxisome proliferator-activated receptors, runt-related transcription factor. The growth factors include, such as, TNF, various interleukins, histamine, TGF-β, prostaglandins and eicosanoids, brain-derived neurotropic factor, vascular endothelial growth factor, hepatocyte growth factor, basic fibroblast growth factor (bFGF) and keratinocyte growth factor, heat shock protein [18].

LLLT can interact with cells, leading to changes at the molecular, cellular and tissue levels. The tissue mechanism of LLLT include: muscle recovery via the increase of creatine kinase, traumatic brain injury treatment via the newly formed neuroprogenitor cells travelling to the injured region of the cortex, and pain reduction via laser induced analgesia, where a conduction block of central and peripheral nerve fibers and to the release of endorphins. The effects of LLLT on hair growth are already well known, but the exact mechanism still needs clarifying [12,18].

Two possible mechanisms have been proposed by Avci et al. [15]: (i) the small amount of heat produced could induce follicular stem cells to proliferate and differentiate, due to the increased level of heat shock proteins; (ii) the release of certain factors affects the cell cycle and induces angiogenesis. Moreover, LLLT stimulates proliferation and differentiation of osteoblasts, leading to an increased bone formation, accompanied by an increase in the activity of alkaline phosphatase and in osteocalcin expression, as reported by Fujimoto et al. [19]. In contrast to LLLT using low fluence of light, high-fluence low-power laser irradiation (HFLP) activates the mitochondrial apoptosis pathway, altering the cell cycle, inhibiting cell proliferation and even causing cell death. Wu et al. [20] found that HFLP, using light at 633nm and 120J/cm², could
The role of light dose has been reported by Lin [30], Alkin et al. [31] & Dhiran et al. [32]. Effort to minimize these complications, various modified PDT protocols have been explored involving reduced verteporfin dosage, laser fluence, or a combination of both. Higher intensity while maintaining the similar efficacy. The linear BRL has limitation as that of BLL. A generalized, time dependent Beer-Lambert law (BLL), the non-BRL optimal conditions, and mathematical formulas (scaling laws) have been discussed by Lin [26-28]. Similar to the synergic efficacy of PDT/PTT [6], far infrared light (3 to 50um) can cause the selective absorption of biological lipid bilayer membranes to enhance the overall LLLT efficacy, via the synergic activation of light/heat sensitive ion channels. Figure 1 shows the PDT efficacy versus light dose (E) and photosensitizer (PS) concentration, C0. Limitation of BRL is shown by the nonlinear law of Lin [28], which demonstrate a higher threshold of [EC0] than that of linear BRL.

The efficacy of LLLT combined with PS has the similar efficacy as that of PDT and may be described by a scaling law of a S-function proportional to the light intensity (I) by a power law, I^n, with 0.3<n<1.2, depending on the tissues (media) optical properties and the kinetics involved. Clinical data are required to fit the scaling laws. Therefore, variation of exposure time and irradiance may account for conflicting results in the literature. Düzgören et al. [29] recently report the optimal combination of light intensity and dose for maximum efficacy. However, they do not present enough data to show the actual value of n in the scaling law. Lin [27] predicts the n value should be approximately 0.5 to 0.8 for most LLLT procedures. Figure 2 shows an optimal dose E* related to the Arndt-Schulz law having an optimal therapeutic window [26,27]. However, without PS, LLLT via the light-activated signaling pathway of reactive oxygen species (ROS), leading to activation of transcription factors is less effective due to the relatively lower rate of generation of ROS. Therefore, LLLT efficacy may be improved by combined with PTT (using light-induced temperature) and PS-initiated ROS for light/heat sensitive ion channels.
Half-fluence, half-dose, half-fluence and micropulse, 1/3 dose, and minimal-fluence protocols have all demonstrated some degree of treatment effect. Half-dose verteporfin and half-fluence treatments are the two most described modified protocols [31,32].

**Conclusion**

We have presented the limitation of the linear BRL and BLL. A generalized, time dependent BLL provides the non-BRL optimal conditions. The optimal dose is related to the Arndt-Schulz law having an optimal therapeutic window. Without PS, LLLT via the light-activated signaling pathway of reactive oxygen species (ROS), leading to activation of transcription factors is less effective due to the relatively lower rate of generation of ROS. Therefore, LLLT efficacy may be improved by combination with PTT (using light-induced temperature) and PS-initiated ROS for light/heat sensitive ion channels.

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DOI: 10.31031/RMES.2018.06.000645