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Low-level laser (light) therapy (LLLT) in skin: stimulating, healing, restoring

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Abstract

Low-level laser (light) therapy (LLLT) is a fast-growing technology used to treat a multitude of conditions that require stimulation of healing, relief of pain and inflammation, and restoration of function. Although the skin is the organ that is naturally exposed to light more than any other organ, it still responds well to red and near-infrared wavelengths. The photons are absorbed by mitochondrial chromophores in skin cells. Consequently electron transport, adenosine triphosphate (ATP) nitric oxide release, blood flow, reactive oxygen species increase and diverse signaling pathways get activated. Stem cells can be activated allowing increased tissue repair and healing. In dermatology, LLLT has beneficial effects on wrinkles, acne scars, hypertrophic scars, and healing of burns. LLLT can reduce UV damage both as a treatment and as a prophylaxis. In pigmentary disorders such as vitiligo, LLLT can increase pigmentation by stimulating melanocyte proliferation and reduce depigmentation by inhibiting autoimmunity. Inflammatory diseases such as psoriasis and acne can also benefit. The non-invasive nature and almost complete absence of side-effects encourages further testing in dermatology.

Keywords

Acne; Dermatology; Herpes; Laser; LLLT; Low level laser therapy; Phototherapy; Skin disease; Skin Rejuvenation; Pigmentation; Vitiligo

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Increasingly, non-invasive therapies for skin disease and skin rejuvenation are used, especially in Western countries where relatively high disposable incomes are combined with the desire for an ideal appearance fostered by societal pressures. Although the skin is the organ that is naturally most exposed to light, it still responds well to red and near-infrared wavelengths delivered at the correct parameters with therapeutic intent. Low-level laser therapy (LLLT) was discovered in the late 1960s, but only in recent times has it been widely applied in dermatology. The introduction of light emitting diode (LED) devices has reduced many of the concerns formerly associated with lasers, such as expense, safety concerns and the need for trained personnel to operate them. In fact, many LED devices are designed for home use and are widely sold on the internet. This review will cover the use of LLLT as possibly the ultimate non-invasive approach to treating the skin.

Low-Level Laser (Light) Therapy and Its Mechanism of Action

LLLT, phototherapy or photobiomodulation refers to the use of photons at a non-thermal irradiance to alter biological activity. LLLT uses either coherent light sources (lasers) or non-coherent light sources consisting of filtered lamps or light-emitting diodes (LED) or, on occasion, a combination of both. The main medical applications of LLLT are reducing pain and inflammation, augmenting tissue repair and promoting regeneration of different tissues and nerves, and preventing tissue damage in situations where it is likely to occur.^{1,2} In the last few decades, non-ablative laser therapies have been used increasingly for the aesthetic treatment of fine wrinkles, photoaged skin and scars, a process known as photorejuvenation (Table 1). More recently, this approach has also been used for inflammatory acne (Table 1).³ LLLT involves exposing cells or tissue to low-levels of red and near infrared (NIR) light. This process is referred to as 'low-level' because the energy or power densities employed are low compared to other forms of laser therapy such as ablation, cutting, and thermally coagulating tissue. Recently, medical treatment with LLLT at various intensities has been found to stimulate or inhibit an assortment of cellular processes.⁴

The mechanism associated with the cellular photobiostimulation by LLLT is not vet fully understood. From observation, it appears that LLLT has a wide range of effects at the molecular, cellular, and tissue levels. The basic biological mechanism behind the effects of LLLT is thought to be through absorption of red and NIR light by mitochondrial chromophores, in particular cytochrome c oxidase (CCO) which is contained in the respiratory chain located within the mitochondria,⁵⁻⁷ and perhaps also by photoacceptors in the plasma membrane of cells. Consequently a cascade of events occur in the mitochondria, leading to biostimulation of various processes (Figure 1).⁸ Absorption spectra obtained for CCO in different oxidation states were recorded and found to be very similar to the action spectra for biological responses to the light.⁵ It is hypothesized that this absorption of light energy may cause photodissociation of inhibitory nitric oxide from CCO⁹ leading to enhancement of enzyme activity,¹⁰ electron transport,¹¹ mitochondrial respiration and adenosine triphosphate (ATP) production (Figure 1).^{12–14} In turn, LLLT alters the cellular redox state which induces the activation of numerous intracellular signaling pathways, and alters the affinity of transcription factors concerned with cell proliferation, survival, tissue repair and regeneration (Figure 1).^{2,5,6,15,16}

Although LLLT is now used to treat a wide variety of ailments, it remains somewhat controversial as a therapy for 2 principle reasons. First, there are uncertainties about the fundamental molecular and cellular mechanisms responsible for transducing signals from the photons incident on the cells to the biological effects that take place in the irradiated tissue. Second, there are significant variations in terms of dosimetry parameters: wavelength, irradiance or power density, pulse structure, coherence, polarization, energy, fluence, irradiation time, contact vs non-contact application, and repetition regimen. Lower dosimetric parameters can result in reduced effectiveness of the treatment and higher ones can lead to tissue damage.¹ This illustrates the concept of the biphasic dose response that has been reported to operate in LLLT ^{1,18,19}. Many of the published studies on LLLT include negative results. It is possibly because of an inappropriate choice of light source and dosage. It may also be due to inappropriate preparation of the patient's skin before application of LLLT, such as: lack of removal of makeup and oily debris, which can interfere with the penetration of the light source, and failure to account for skin pigmentation.¹⁷ Inappropriate maintenance of the LLLT equipment can reduce its performance and interfere with clinical results as well. It is important to consider that there is an optimal dose of light for any particular application.

Laser radiation or non-coherent light has a wavelength and radiant exposure dependent capability to alter cellular behavior in the absence of significant heating.²⁰ Phototherapy employs light with wavelengths between 390–1.100 nm and can be continuous wave or pulsed. In normal circumstances, it uses relatively low fluences (0.04–50 J/cm²) and power densities (< 100 mW/cm²).²¹ Wavelengths in the range of 390 nm to 600 nm are used to treat superficial tissue, and longer wavelengths in the range of 600nm to 1,100nm, which penetrate further, are used to treat deeper-seated tissues (Figure 2).⁴ Wavelengths in the range 700 nm to 750 nm have been found to have limited biochemical activity and are therefore not often used.¹ Various light sources used in LLLT include inert gas lasers and semiconductor laser diodes such as helium neon (HeNe; 633 nm), ruby (694 nm), argon (488 and 514 nm), krypton (521, 530, 568, 647 nm), gallium arsenide (GaAs; > 760 nm, with a common example of 904 nm), and gallium aluminum arsenide (GaAlAs: 612-870 nm).¹⁷ A wide range of LED semiconductors are available at lower wavelengths, whose medium contains the elements indium, phosphide and nitride. One question that has not yet been conclusively answered is whether there is any advantage to using coherent laser light over non-coherent LED light.²² While some medical practitioners treat deep tissue lesions using focused lasers in "points", in dermatology the use of LEDs is becoming increasingly common due to the relatively large areas of tissue that require irradiation.

LLLT for Skin Rejuvenation

Skin starts showing its first signs of aging in the late 20s to early 30s and it usually presents with wrinkles, dyspigmentation, telangiectasia, and loss of elasticity. Common histologic and molecular-level features are reduction in the amount of collagen, fragmentation of collagen fibers, elastotic degeneration of elastic fibers, upregulation of matrix metalloproteinases (MMPs), especially MMP-1 and MMP-2, dilated and tortuous dermal vessels, and atrophy and disorientation of the epidermis.^{23,24} Both chronological and

environmental influences are responsible for the aging process of skin; however photodamage seems to be one of the most important causes of these changes.

Several modalities have been developed in order to reverse the dermal and epidermal signs of photo- and chronological aging. The main concept of most of these modalities is removing the epidermis and inducing a controlled form of skin wounding in order to promote collagen biosynthesis and dermal matrix remodeling. The most commonly used interventions as of today are retinoic acid (a vitamin A derivative), dermabrasion, chemical peels, and ablative laser resurfacing with carbon dioxide (CO₂) or erbium: yttriumaluminum-garnet (Er:YAG) lasers or a combination of these wavelengths.^{25–27} However, these procedures require intensive post-treatment care, prolonged downtime and may lead to complications such as long-lasting erythema, pain, infection, bleedings, oozing, burns, hyper- or hypopigmentation and scarring.^{28,29} These limitations created a need for the development of alternative rejuvenation procedures that were safer, more effective, had fewer side effects and minimum postoperative care and downtime, which in turn led to the emergence of non-ablative rejuvenation technologies.^{30–32} Non-ablative skin rejuvenation aims to improve photoaged and aging skin without destroying the epidermis.^{31,32} Irregular pigmentation and telangiectasia can be treated with intense pulsed light sources (IPL), 532 nm potassium-titanyl-phosphate lasers (KTP), and high-dose 585/595 nm pulsed dye lasers (PDL)³³. Wrinkle reduction and skin tightening through thermal injury to the dermis (photothermolysis) can be achieved by other IPL sources (*ie*, low-dose 589/595 nm PDLs, 1064 & 1320 nm neodymium:yttrium-aluminum-garnet lasers, (Nd:YAG) 1450 nm diode lasers, and 1540 nm erbium fiber lasers).³³

LED which is a novel light source for non-thermal, non-ablative skin rejuvenation has been shown to be effective for improving wrinkles and skin laxity (Figure 3). $^{34-40}$ It is not a new phenomenon since the first reports of LLLT effects on increased collagen go back to 1987. Studies by Abergel et al. and Yu et al. reported an increase in production of pro-collagen, collagen, basic fibroblast growth factors (bFGF) and proliferation of fibroblasts after exposure to low-energy laser irradiation in vitro and in vivo animal models (Figure 4).^{41,42} Furthermore, LLLT was already known to increase microcirculation, vascular perfusion in the skin, alter platelet-derived growth factor (PDGF), transforming growth factor (TGF- β 1) and inhibit apoptosis (Figure 4).^{1,43,44} Lee et al. investigated the histologic and ultrastructural changes following a combination of 830 nm, 55 mW/cm², 66 J/cm² and 633 nm, 105 mW/cm², 126 J/cm² LED phototherapy and observed alteration in the status of MMPs and their tissue inhibitors (TIMPs).³³ Furthermore, mRNA levels of IL-1β, TNF-*a*, ICAM-1, and connexin 43 (Cx43) were increased following LED phototherapy whereas IL-6 levels were decreased (Figure 4)³³. Finally, an increase in the amount of collagen was demonstrated in the post-treatment specimens ³³. Pro-inflammatory cytokines IL-1β and TNF-a are thought to be recruited to heal the intentionally formed photothermally-mediated wounds associated with laser treatments, and this cascade of wound healing consequently contributes to new collagen synthesis.³³ LED therapy may induce this wound healing process through non-thermal and atraumatic induction of a subclinical 'quasi-wound', even without any actual thermal damage which could cause complications as in some other laser treatments.³³ TIMPs inhibit MMP activities, so another possible mechanism for the

increased collagen could be through the induction of TIMPs (Figure 4). When these observations are put together, it is possible that increased production of IL-1 β and TNF-a might have induced MMPs in the early response to LED therapy. This may clear the photodamaged collagen fragments to enable biosynthesis of new collagen fibers. Later on, an increase in the amount of TIMPs might protect the newly synthesized collagen from proteolytic degradation by MMPs.³³ Furthermore, increased expression of Cx43 may possibly enhance cell-to-cell communication between dermal components, especially the fibroblasts, and enhance the cellular responses to the photobiostimulation effects from LED treatment, in order to produce new collagen in a larger area which even includes the nonirradiated regions.³³ In a clinical study performed by Weiss et al., 300 patients received LED therapy (590 nm, 0.10 J/cm2) alone, and 600 patients received LED therapy in combination with a thermal-based photorejuvenation procedure. Among patients who received LED photorejuvenation alone, 90% reported that they observed a softening of skin texture and a reduction in roughness and fine lines ranging from a significant reduction to sometimes subtle changes.³⁶ Moreover, patients receiving a thermal photorejuvenation laser with or without additional LED photomodulation (n = 152) reported a prominent reduction in post-treatment erythema and an overall impression of increased efficacy with the additional LED treatment.^{36,45} This reduction in post-treatment erythema could be attributed to anti-inflammatory effects of LLLT.⁴⁰ Using different pulse sequence parameters, a multicenter clinical trial was conducted, with 90 patients receiving 8 LED treatments over 4 weeks.^{37,46–48} The outcome of this study showed very favorable results, with over 90% of patients improving by at least one Fitzpatrick photoaging category and 65% of patients demonstrating global improvement in facial texture, fine lines, background erythema, and pigmentation. The results peaked at 4 to 6 months following completion of 8 treatments. Markedly increased collagen in the papillary dermis and reduced MMP-1 were common findings. Barolet et al.'s study is also consistent with the previously mentioned studies. They used a 3-D model of tissue-engineered human reconstructed skin to investigate the potential of 660 nm, 50 mW/cm, 4 J/cm² LED in modulating collagen and MMP-1 and results showed upregulation of collagen and down-regulation MMP-1 in vitro.⁴⁰ A split-face, single-blinded clinical study was then carried out to assess the results of this light treatment on skin texture and appearance of individuals with aged/photoaged skin.⁴⁰ Following 12 LED treatments, profilometry quantification demonstrated that while more than 90% of individuals had a reduction in rhytid depth and surface roughness, 87% of the individuals reported that they have experienced a reduction in the Fitzpatrick wrinkling severity score.⁴⁰

LLLT for Acne

Pathogenesis of acne vulgaris has not yet been clarified, however current consensus is that it involves four main events: follicular hyperconification, increased sebum secretion effected by the androgenic hormone secretions, colonization of *Propionibacterium acnes* and inflammation.⁴⁹ *P. acnes* plays a key role by acting on triglycerides and releasing its cytokines which in turn trigger inflammatory reactions and alter infundibular keratinization.⁴⁹ Current treatments for acne vulgaris include topical and oral medications such as topical antibiotics, topical retinoids, benzoyl peroxide, alpha hydroxy acids, salicylic acid, or azaleic acid. In severe cases, systemic antibiotics such as tetracycline and

doxycycline, oral retinoids, and some hormones are indicated.⁵⁰ Medications work by counteracting microcomedone formation, sebum production, *P. acnes*, and inflammation.⁵⁰ Despite many options that are available for treatment of acne vulgaris, many patients still respond inadequately to treatment or experience some adverse effects.

Phototherapy (light, lasers, and photodynamic therapy) has been proposed as an alternative therapeutic modality to treat acne vulgaris and was proposed to have less side effects compared to other treatment options.⁵¹ Exposure of sunlight was reported to be highly effective for treatment of acne with efficacy up to 70%.⁵² The sunlight decreases the androgenic hormones in the sebaceous gland, but the unwanted effect of exposure to UVA and UVB limits sunlight for the treatment of acne. Recently, phototherapy with visible light (mainly blue light, red light or combination of both) started being used in the treatment of acne (Figure 3).⁵² One mechanism of action of phototherapy for acne is through the absorption of light (specifically blue light) by porphyrins that have been produced by *P*. acnes as a part of its normal metabolism, and that act as endogenous photosensitizers. ^{49,53} This process causes a photochemical reaction and forms reactive free radicals and Singlet oxygen species which in turn lead to bacterial destruction (Figure 5).^{49,53} Red light is known to penetrate deeper in tissues when compared to blue light.⁵⁰ It has been demonstrated that red light can affect the sebum secretion of sebaceous glands and change keratinocyte behavior.⁵⁴ Furthermore, red light might also exert its effects by modulating cytokines from macrophages and other cells, which in turn could reduce inflammation, 51,54

Several studies reported that LLLT in the red to near infrared spectral range (630–1000 nm) and nonthermal power (less than 200 mW) alone or in combination with other treatment modalities (mainly blue light), is effective for treatment of acne vulgaris.^{17,49,52,54,55} One of these studies demonstrated significant reduction in active acne lesions after 12 sessions of treatment using 630-nm red spectrum LLLT with a fluence of 12 J/cm² twice a week for 12 sessions in conjunction with 2% topical clindamycin; however the same study showed no significant effects when a 890 nm laser was used.⁵⁰ A few studies also showed that the combination of blue light and red light have synergistic effects in acne treatment.^{49,54–56} It is proposed that synergistic effects of mixed light is due to synergy between the antibacterial and anti-inflammatory effect of blue and red light respectively (Figure 5).^{49,56} It is also worth mentioning that in most studies improvement in inflammatory lesions were higher than the improvement in comedones.^{49,56}

LLLT for Photoprotection

It is widely accepted that the UV range (< 400 nm) exposure is responsible for almost all damaging photo-induced effects on human skin.^{57–59} Some proposed mechanisms for UV induced skin damage are collagen breakdown, formation of free radicals, inhibition of DNA repair, and inhibition of the immune system.^{57–59} Existing solutions to prevent UV induced damaging effects are based on minimizing the amount of UV irradiation that reaches the skin, which is achieved by either avoidance of sun exposure or by use of sunscreens. However sometimes sun avoidance might be hard to implement, especially for the people involved in outdoor occupations or leisure activities. On the other hand, the photoprotective efficacy of topical sunscreens have their own limitations as well, which include decreased

efficacy following water exposure or perspiration, spectral limitations, possible toxic effects of nanoparticles that are contained by most sunscreens,⁶⁰ user allergies, and compliance.

It has recently been suggested that infrared (IR) exposure might have protective effects against UV-induced skin damage mainly by triggering protective/repair responses to UV irradiation. In the natural environment, visible and IR solar wavelengths predominate in the morning and UVB and UVA are maximal around noon which suggest that mammalians already possess a natural mechanism which, in reaction to morning IR radiation, prepares the skin for upcoming potentially damaging UV radiation at noon.⁶¹ However, opposing views also exist, such as Krutmann's study demonstrating IR-induced disturbance of the electron flow of the mitochondrial electron transport chain which leads to inadequate energy production in dermal fibroblasts.⁶² Schroeder's report is another example stating that IR alters the collagen equilibrium of the dermal extracellular matrix by leading to an increased expression of the collagen-degrading enzyme MMP-1, and by decreasing the de novo synthesis of the collagen itself.⁵⁹ As previously mentioned, the same light source may have opposite effects on the same tissue depending on the parameters used and these conflicting views are probably due to the biphasic effects of light.^{18,19}

Menezes et al. demonstrated that non-coherent near infrared radiation (NIR) (700-2,000 nm) generated a strong cellular defense against solar UV cytotoxicity in the absence of rising skin temperature and it was assumed to be a long-lasting (at least 24 hours) and cumulative phenomenon.⁶³ Following this study, Frank et al. proposed that IR irradiation prepares cells to resist UVB-induced damage by affecting the mitochondrial apoptotic pathway.⁶⁴ IR preirradiation of human fibroblasts was shown to inhibit UVB activation of caspase-9 and -3, partially release of cytochrome c and Smac/Diablo, decrease pro-apoptotic (ie, Bax) and increase anti-apoptotic proteins (ie, Bcl-2 or Bcl-xL).⁶⁴ The results suggested that IR inhibited UVB-induced apoptosis by modulating the Bcl2/Bax balance, pointing to a role of p53, a sensor of gene integrity involved in cell apoptosis and repair mechanisms. In a further study, Frank et al. studied more specifically the role of the p53 cell signaling pathway in the prevention of UVB toxicity.⁶⁴ The response to IR irradiation was shown to be p53 dependent which further suggests that IR irradiation prepares cells to resist and/or to repair further UVB-induced DNA damage. Finally, the IR induction of defense mechanisms was supported by Applegate et al. who reported that the protective protein, ferritin, normally involved in skin repair (scavenger of Fe²⁺ otherwise available for oxidative reactions) was induced by IR radiation.65

In an in vitro study, it was reported that an increase dermal fibroblast procollagen secretion reduces metalloproteinases (MMP) or collagenase production following non-thermal non-coherent deep red visible LED exposures (660 nm, sequential pulsing mode).⁴⁰ These results correlated with significant clinical improvement of rhytids in vivo.⁴⁰ In a subsequent in vivo pilot study, effect of this wavelength in 3 healthy subjects using a minimal erythemal dose (MED) method adapted from sunscreen SPF determination has been investigated.⁶¹ The results showed that LED therapy was effective, achieving a significant response in the reduction of the erythema induced by UVB.⁶¹ Following this pilot study a further investigation has been performed to find out in vivo aspects of this phenomenon. Effects of non-thermal, non-coherent 660 nm LED pulsed treatments in providing enhanced skin

resistance prior to upcoming UV damage was investigated in a group of subjects with normal fair skin and patients presenting polymorphous light eruption (PLE). Results suggested that LED based therapy prior to UV exposure provided significant dose-related protection against UVB induced erythema. A significant reduction in UVB induced erythema reaction was observed in at least one occasion in 85% of subjects as well as in the patients suffering from PLE. Furthermore, a sun protection factor SPF-15-like effect and a reduction in post-inflammatory hyperpigmentation were observed. An in vitro study by Yu et al. revealed that HeNe laser irradiation stimulated an increase in nerve growth factor (NGF) release from cultured keratinocytes and its gene expression.⁶⁶ NGF is a major paracrine maintenance factor for melanocyte survival in skin.⁶⁷ It was shown that NGF can protect melanocytes from UV-induced apoptosis by upregulating BCL-2 levels in the cells.⁶⁸ Therefore, an increase in NGF production induced by HeNe laser treatment may provide another explanation for the photoprotective effects of LLLT.

LLLT for Herpes Virus Lesions

One of the most common virus infections is caused by herpes simplex virus (HSV). HSV is chronic and lasts one's entire life. The exposure of the host to several kinds of physical or emotional stresses such as fever, exposure to UV light, and immune suppression causes virus reactivation and migration through sensory nerves to skin and mucosa, localizing particularly on the basal epithelium of the lips and the perioral area.⁶⁹ Up to 60% of sufferers will experience a prodromic stage, after which the lesions develop through stages of erythema, papule, vesicle, ulcer and crust, until healing is achieved. It is accompanied by pain, burning, itching, or tingling at the site where the blisters form. Immune responses to HSV infection involve: macrophages, Langerhans cells, natural killer cells, lymphocyte-mediated delayed-type hypersensitivity, and cytotoxicity.⁷⁰

While several anti-viral drugs such as acyclovir and valacyclovir are used to control recurrent herpes outbreaks, only limited reduction in the lesions' healing time has been observed.⁶⁹ Furthermore, development of drug-resistant HSV strains is of increasing significance especially in immunocompromised patients.⁷⁰ Therefore, new therapy modalities that can shorten the recurrent episodes and cause prominent reduction of related pain and inflammation are necessary.

LLLT has been suggested as an alternative to current medications for accelerated healing, reducing symptoms and influencing the length of the recurrence period.^{69,71,72} Among 50 patients with recurrent perioral herpes simplex infection (at least once per month for more than 6 months), when LLLT (690 nm, 80 mW/cm², 48 J/cm²) was applied daily for 2 weeks during recurrence-free periods it was shown to decrease the frequency of herpes labialis episodes.⁷³ In another study with similar irradiation parameters (647 nm, 50 mW/cm², 4.5 J/cm²), investigators achieved a significant prolongation of remission intervals from 30 to 73 days in patients with recurrent herpes simplex infection.⁷⁴ Interestingly, patients with labial herpes infection showed better results than those with genital infection. However, irradiation did not effect established HSV latency in a murine model.⁷⁵

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Even though mechanism of action is still not clear, an indirect effect of LLLT on cellular and humoral components of the immune system involved in antiviral responses rather than a direct virus-inactivating effect was proposed.⁷⁶ Inoue et al. investigated tuberculin reactions at the bilateral sites of the backs of sensitized guinea pigs. They applied a single low-power laser irradiation dose at a fluence of 3.6 J per cm²on one side and compared it to contralateral non-irradiated sites.⁷⁷ Interestingly, following irradiation, tuberculin reaction was suppressed not only at the irradiated site but also at the contralateral nonirradiated site. It is worth to note that this phenomenon was observed when mononuclear cells were dominant in the perivascular cellular infiltration. Based on their results, they have suggested a possible systemic inhibitory effect of LLLT on delayed hypersensitivity reactions.⁷⁷ Activation and proliferation of lymphocytes^{78–81} and macrophages⁸² as well as the synthesis and expression of cytokines^{83,84} following low intensities of red and NIR light have been reported by several investigators. The question of whether these effects of LLLT have any influence on HSV infection remains to be determined.

LLLT for Vitiligo

Vitiligo is an acquired pigmentary disorder characterized by depigmentation of the skin and hair. The underlying mechanism of how the functional melanocytes disappear from the involved skin is still under investigation. However, findings suggest that regardless of the pathogenetic mechanism involved, keratinocytes, fibroblasts, melanoblasts and melanocytes may all be involved in both the depigmentation and also the repigmentation processes of vitiligo.^{66,85–89} Therefore, stimulation of these epidermal and dermal cells may be a possible treatment option. Due to the obscure pathogenesis of the disease, treatment of vitiligo has generally been unsatisfactory. Current existing therapies that induce varying degrees of repigmentation in patients with vitiligo are topical corticosteroids, phototherapy, and photochemotherapy (PUVA).⁸⁹ In 1982, a group of investigators found that low energy laser irradiation had effects on defective biosynthesis of catecholamine in certain dermatological conditions including scleroderma and vitiligo.^{90,91} Later on, one of the investigators from the same group reported that after 6-8 months of treating 18 vitiglio patients with lowenergy HeNe laser (632 nm, 25 mW/cm²) therapy, marked repigmentation was observed in 64% of the patients and some follicular repigmentation was observed in the remaining 34%.⁹¹ Since then, LLLT has been suggested as an alternative effective treatment option for patients with vitiligo.66,88,89

Segmental-type vitiligo is associated with a dysfunction of the sympathetic nerves in the affected skin and it is relatively resistant to conventional therapies.⁶⁶ Based on the previous reports stating that HeNe laser irradiation leads to improvement in nerve injury^{92–94} and LLLT induces repigmentation responses,^{95,96} it was proposed that the HeNe laser might be a potential treatment modality for treatment of segmental type vitiligo. ⁶⁶ When the HeNe laser light was administered locally (3 J/cm², 1.0 mW, 632.8 nm), marked perilesional and perifollicular repigmentation (> 50%) was observed in 60% of patients with successive treatments. Both NGF and bFGF stimulate melanocyte migration and deficiencies of these mediators may participate in the development of vitiligo.^{86,97,98} In the same study, when cultured keratinocytes and fibroblasts were irradiated with 0.5–1.5 J per cm² HeNe laser, significant increase in bFGF release both from keratinocytes and fibroblasts as well as a

significant increase in NGF release from keratinocytes was reported. ⁶⁶ Additionally, the medium from HeNe laser irradiated keratinocytes stimulated [3H]thymidine uptake and proliferation of cultured melanocytes. Another study by Lan et al. demonstrated that the HeNe laser (632.8 nm, 1 J/cm² and 10 mW) stimulates melanocyte proliferation through enhanced $\alpha 2\beta 1$ integrin expression⁸⁸ and induces melanocyte growth through upregulation of the expression of phosphorylated cyclic-AMP response element binding protein (CREB) which is an important regulator of melanocyte growth.⁸⁸ ECM molecules are also important elements of the pigmentation process due to their regulatory roles for physiological functions of pigment cells including morphology, migration, tyrosinase activity and proliferation.^{99–101} Type IV collagen is present in the basement membrane and is known to have an intricate relationship with melanocytes in the epidermis such as increasing melanocyte mobility.⁸⁹ Following, HeNe irradiation, the attachment of melanocytes to type IV collagen was found to be significantly enhanced which also indicated modulation of melanocyte physiological function by HeNe laser irradiation.⁸⁸ Furthermore, among various ECM proteins found in the dermis, fibronectin was shown to have significant effects on both differentiation and migration of cultured melanoblasts and melanocytes.^{102,103} In 1983. Gibson et al. demonstrated that the physical distribution of fibronectin in vivo was closely associated with the migration path undertaken by melanoblasts during the repigmentation process of vitiligo.¹⁰⁴ Based on Lan at al.'s findings, an immature melanoblast cell line (NCCmelb4) showed significant decrease in the attachment to fibronectin following HeNe laser treatment while the attachment of a more differentiated melanoblast cell line (NCCmelan5) to fibronectin increased about 20% following 1 J/cm2, 10 mW HeNe laser treatment.⁸⁹ Lastly, expression of integrin a5b1 which mediate locomotion of pigment cells was found to be enhanced on NCCmelb4 cells.89

LLLT for Producing Depigmentation

Most studies carried out for vitiligo show the stimulatory effects of LLLT on pigmentation; however in a previously mentioned study, while testing effects of blue and red laser for acne treatment, an interesting and unexpected result was found for the first time.⁴⁹ Combining both blue (415 +–5 nm, irradiance 40 mW/cm2, 48 J/cm²) and red (633 +- 6 nm, 80 mW/cm², 96 J/cm²) light produced an overall decrease in the melanin level. Instrumental measurement results showed that melanin level increased by 6.7 (the median of differences between the melanin level before and after one treatment session) after blue light irradiation without a statistical significance (P > .1), whereas it decreased by 15.5 with statistical significance (P < .005) after red light irradiation. This finding may have some relationship with the laser's brightening effect of the skin tone, which 14 out of 24 patients spontaneously reported after the treatment period. However as of today, no other studies investigated or reported similar decrease in melanin levels following red light irradiation. Considering that different parameters are used for vitiligo and acne treatment, different effects of red light on the same tissue might be due to the biphasic effects of LLLT.^{18,19}

LLLT for Hypertrophic Scars and Keloids

Hypertrophic scars and keloids are benign skin tumors that usually form following surgery, trauma, or acne and are difficult to eradicate. Fibroblastic proliferation and excess collagen

deposits are the 2 main characteristics¹⁰⁵ and imbalance between rates of collagen biosynthesis and degradation superimposed on the individual's genetic predisposition have been implicated in their pathogenesis. A wide range of surgical (eg, cryotherapy, excision), non-surgical (e.g., pharmacological, mechanical pressure, silicone gel dressings), and laser therapies (CO2, pulsed dye, fractional ablative, and non-ablative lasers) have been tested with variable success, however until now, an optimal treatment of these lesions remains to be found.^{106–108} It has recently been proposed that poor regulation of interleukin (IL)-6 signaling pathways and transforming growth factor beta-I (TGF-BI) expression have a significant role in this process and thus inhibition of the IL-6 pathway and/or TGF-BI could be a potential therapeutic target.^{106,107,109–111} Based on the reports demonstrating the effects of LLLT on decreasing IL-6 mRNA levels,³³ modulation of PDGF, TGF-β, interleukins such as IL-13 and IL-15, MMPs, which are all also associated with abnormal wound healing,^{112,113} it was proposed to be an alternative therapy to existing treatment options. The use of LLLT as a prophylactic method to alter the wound healing process to avoid or attenuate the formation of hypertrophic scars or keloids has been investigated by Barolet and Boucher in 3 cases studies, where following scar revision by surgery or CO_2 laser ablation on bilateral areas, a single scar was treated daily by the patient at home with NIR - LED 805 nm at 30 mW/cm² and 27 J/cm².¹¹² The first patient had pre-auricular linear keloids bilaterally post-face lift procedure and surgical scar revision/excision had been performed. The second patient had hypertrophic scars on the chest bilaterally post-acne when the CO_2 laser was used for resurfacing. The third patient had hypertrophic scars on the back bilaterally post-excision and again the CO₂ laser was used for resurfacing. As a result, significant improvements on the NIR-LED treated vs the control scar were seen in all efficacy measures and moreover no significant treatment-related adverse effects were reported.112

LLLT for Burns

In a clinical study by Weiss et al. 10 patients received LED treatment (590 nm) for acute sunburn using a once-or twice-daily treatment regimen for 3 days, treating only half of the affected anatomic area.³⁶ Decreased symptoms of burning, redness, swelling, and peeling were reported. One patient received LED treatment twice daily for 3 days only on half of his back, and other half was left untreated.³⁶ When compared with the untreated side, decreased MMP-1 was demonstrated on the LED-treated side through immunofluorescence staining. Moreover, RT-PCR gene expression analysis showed a significant decrease in MMP-1 gene expression on the LED-treated side at both 4 and 24 hours post–UV injury compared with the untreated side. Other significant changes were also noted with LED treatment related to inflammation and dermal matrix composition 4 days post–ultraviolet (UV) exposure.³⁶

One of the main complications of receiving laser treatment is burns which may be devastating for the patient. LED was suggested as a treatment modality for facilitating faster healing. A group of 9 patients who had a variety of second-degree burns from nonablative laser devices were given LED therapy once a day for 1 week and according to both the patient and the physician, healing occurred 50% faster.³⁶ Also the same investigators conducted a pilot study, where one forearm was injured by a CO₂ laser using a computer pattern generator to deliver the identical treatment to both test sites. Both sites received daily

dressing changes using a non-stick dressing and Polysporin ointment, but one site also received additional LED treatment.³⁶ As a result, when compared to the untreated control site, accelerated reepithelialization was observed in the LED-treated site ³⁶.

LLLT for Psoriasis

More recently LLLT has been considered for treatment of plaque psoriasis. A recent preliminary study investigated the efficacy of a combination of 830 nm (near infrared) and 630 nm (visible red light) to treat recalcitrant psoriasis using LED irradiation. All patients with psoriasis resistant to conventional therapy were enrolled and were treated sequentially with 830 nm and 630 nm wavelengths in 2 20-min sessions with 48 hours between session for 4 or 5 weeks. The results showed no adverse side effects and a resolution of psoriasis.¹¹⁴ The limitation of this study was the small number of patients enrolled, however the results observed encourage future investigations for use of LLLT in treating psoriasis.

Conclusion

LLLT appears to have a wide range of applications of use in dermatology, especially in indications where stimulation of healing, reduction of inflammation, reduction of cell death and skin rejuvenation are required. The application of LLLT to disorders of pigmentation may work both ways by producing both repigmentation of vitiligo, and depigmentation of hyperpigmented lesions depending on the dosimetric parameters. The introduction of LED array-based devices has simplified the application to large areas of skin. There is no agreement as yet on several important parameters particularly whether red, NIR, or a combination of both wavelengths is optimal for any particular application. There is a credibility gap that needs to be overcome before LLLT is routinely applied in every dermatologist's office.

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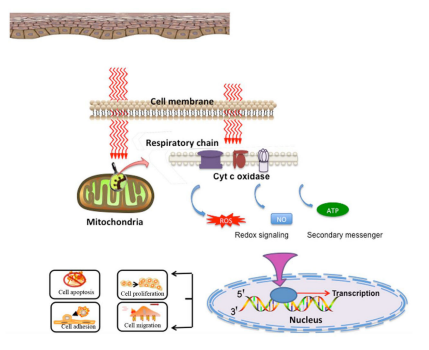
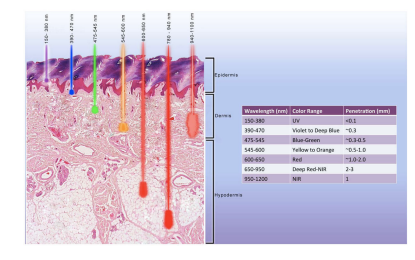


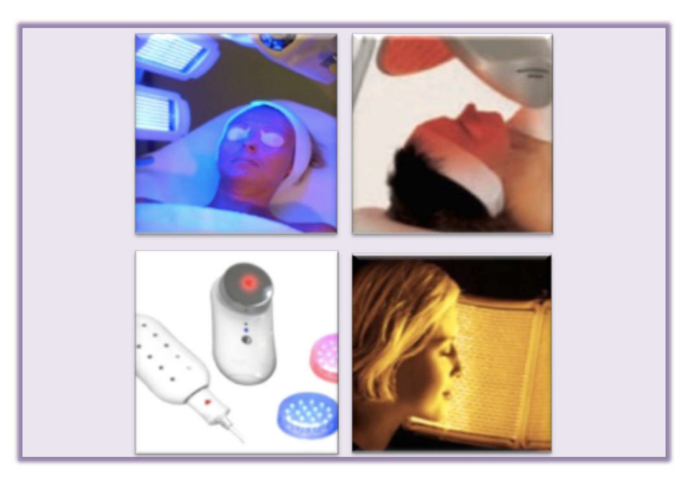
Figure 1.

Mechanism of action of LLLT.

Basic biological mechanism behind the effects of LLLT is thought to be through absorption of red and NIR light by mitochondrial chromophores, in particular cytochrome c oxidase (CCO) which is contained in the respiratory chain located within the mitochondria ^{5–7}. It is hypothesized that this absorption of light energy may cause photodissociation of inhibitory nitric oxide from CCO ⁹ leading to enhancement of enzyme activity ¹⁰, electron transport ¹¹, mitochondrial respiration and ATP production ^{12–14}. In turn, LLLT by altering the cellular redox state can induce the activation of numerous intracellular signaling pathways; alter the affinity of transcription factors concerned with cell proliferation, survival, tissue repair and regeneration^{2,5,6,15,16.}









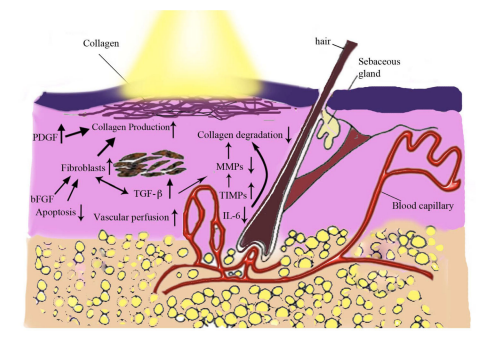


Figure 4.

Possible mechanism of actions for LLLT's effects on skin rejuvenation. LLLT aids skin rejuvenation through increasing collagen production and decreasing collagen degradation. Increase in collagen production occurs by LLLT's increasing effects on PDGF and fibroblast production which happens through decreasing apoptosis, increasing vascular perfusion, bFGF and TGF- β . Decrease in IL-6, and increase in TIMPs which in turn reduce MMPs all aid in reduction of collagen degradation.

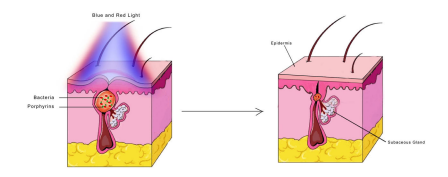


Figure 5.

Illustration of acne treatment with red and blue light.

Red and blue light when used in combination have synergistic effects in treatment of acne. *P. acnes* synthesizes and stores a large amount of porphyrins. Once the porphyrin is exposed to visible light (specifically blue light) it becomes chemically active and transfers to an excited state, resulting in formation of reactive free radicals and singlet oxygen which in turn causes membrane damage in *P. acnes* ^{49,53}. Red light is proposed to exert its effects through reducing the inflammatory process 51,54 .

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Table 1

Examples of LLLT Devices for Dermatological Applications

Supplier	Product name	Wavelength (nm)	Power Density (mW/cm ²)	Standard Dose(J/cm ²)	Application
PhotoMedex (Manchester, UK)	Omnilux	415 (±5) 633 (±6) 830 (±5)	40 105 55	48 126 66	Acne, photodamage, non-melanoma skin cancers, skin rejuvenation, vitiligo and wound healing post elective surgery
Edge Systems (Signal Hill, CA)	Delphia del Sol	420 600–700 700–1000		7.4 J per treatment area	Acne, improving skin texture, firmness and resilience, increasing lymphatic system activity, fine lines, wrinkles and superficial hyperpigmentation
Flip 4 (Sainte-Julie, Quebec, Canada)	Max7	420–700	4		Acne, rejuvenation, injured skin healing including the shortening of the post skin resurfacing erythema duration
Light BioSciences (Virginia Beach, VA)	Gentlewaves	588 (±10)	Variable		Anti-aging
OPUSMED (Montreal, Canada)	LumiPhase-R	660	150		Skin firmness, rhytid depth, wrinkles
Revitalight (Chicago, IL)	Revitalight 747 Hand Spa Food Spa Evolution	420 590 625 940	80 80 80 80	7.2 J per 90 sec. per treatment area	Fine lines, wrinkles, and age spots on the face, neck and hands
Soli-Tone (Woburn, MA)	LumiFacial Lumilift	470 525 590 640	84 27 10 89		Acne, anti-aging, hyperpigmentation, rosacea
DUSA (Wilmington, MA)	BLU-U	417	10		Acne
Curelight (Rehovot, Israel)	iClearXL Clear100XL	405–420 890–900		60	Acne, anti-aging, skin rejuvenation, acceleration of healing of post peel and post surgical suture sites
Lumenis (Santa Clara, CA)	ClearLight Clear100	405-420	200	60	Acne

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Supplier	Product name	Wavelength (nm)	Wavelength (mm) Power Density (mW/cm ²) Standard Dose(J/cm ²)	Standard Dose(J/cm ²)	Application
LIGHTWAVE Technologies	LIGHTWAVE	417			Anti-aging, skin rejuvenation
(Phoneix, AZ)	Protessional Deluxe LED	630			
	System	880			
Dynatronics (Salt Lake City, UT)	Synergie LT2	660	500 mW (total power)	6 J per treatment area	Anti-aging, skin firmness, wrinkles, skin tone and
		880			texture for face and neck