Research Article

The Impact of Near-infrared in Plastic Surgery

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Abstract

Many studies regarding near-infrared, have used near-infrared resources without a water filter or a cooling system, and have proven its thermal effects. With these methods near-infrared energy is mainly absorbed in the superficial tissues and cannot be delivered sufficiently to deeper tissues. As solar near-infrared is filtered by atmospheric water, a water filter is essential in order to simulate solar near-infrared. Thus, these approaches could not sufficiently evaluate the effects of incident solar near-infrared that reaches the human tissue. We have clarified that the near-infrared that simulates solar near-infrared non-thermally affects subcutaneous tissues, including muscle. Importantly, the biological effects of near-infrared have both beneficial applications and deleterious effects. Near-infrared induces dermal heating thermally and non-thermally induces collagen and elastin stimulation, which results in skin tightening, and induces long-lasting vasodilation that may prevent vasospasm and may be beneficial for ischemic disorders and flap surgeries. Near-infrared also non-thermally relaxes and weakens dystonic or hypertrophic muscles to reduce wrinkles and myalgia. Its long-lasting induction of subcutaneous adipocytes may have an application in volume augmentation. However, continuous near-infrared exposure may induce photoaging and thinning of superficial muscles, which results in skin ptosis. Protection against near-infrared should be strongly considered, as over half of the solar energy is near-infrared. Although plastic surgeons are not familiar with the effects of near-infrared, its potential appears to be high and significant. This paper reviews the effects of near-infrared and introduces the new findings and applications of the biological effects of near-infrared in the field of plastic surgery.

Keywords: Near-infrared; non-thermal; biological effects; photoaging.

Introduction

Many studies have demonstrated the effects of sun and ultraviolet (UV) exposure on human tissue. However, the long-term effects of near-infrared (NIR) exposure on human skin and subcutaneous tissues have not been well

investigated. Although various types of UV blocking materials, such as sunblock, sunglasses, films, and fibers are often used to prevent skin damage from UV exposure, and despite the world-wide use of these blocking materials, unwanted physiological effects such as rosacea, erythema ab igne, long-lasting vasodilation (Tanaka, 2009a; Tanaka, 2011a), long-lasting muscle thinning (Tanaka, 2010a, 2011b), and sagging and skin ptosis still occur (Tanaka, 2011c; Tanaka, 2012a, b). Both superficial muscle thinning and skin ptosis develop with age, and though face lift surgeries and skin tightening procedures are proven effective treatments, the mechanisms underlying their development remain unclear.

We previously reported that NIR from 1100 to 1800 nm together with a water-filter that excludes wavelengths between 1400 and 1500 nm penetrates deep into human tissue and it is absorbed by water in the skin (Tanaka, 2009a,b; Tanaka, 2010b; Tanaka, 2011d), hemoglobin in dilated vessels (Tanaka, 2009a; Tanaka, 2011a), myoglobin in the superficial muscle (Tanaka, 2010a; Tanaka, 2011b), and bone cortical mass, and it is scattered by adipose cells (Tanaka, 2011e; Tanaka, 2012b).

Notably, in order to experimentally simulate solar NIR that reaches the skin (Tanaka, 2012a, b); a water filter is required, as solar NIR is filtered by atmospheric water (Anderson, 1981; Gates, 1966). NIR increases the surface temperature and induces thermal effects, so a contact cooling is needed to pursue the properties of NIR (Tanaka, 2011c; Tanaka, 2012a, b).

NIR is used to treat wound healing disorders (Danno, 2001; Horwitz, 1999; Schramm, 2003) and malignant tumors (Bäumler, 1999; Dees, 2002; Kelleher, 1999; Orenstein, 1999). NIR induces dermal heating thermally and non-thermally induces collagen and elastin stimulation, which results in skin tightening. NIR also induces long-lasting vasodilation that may prevent vasospasm and may be beneficial for ischemic disorders and flap surgeries in the field of plastic surgery. Further, NIR non-thermally relaxes and weakens dystonic or hypertrophic muscles to reduce wrinkles and myalgia.

In addition to its therapeutic effects, however, NIR also induces non-thermal DNA damage (Tanaka, 2010c; Tanaka, 2012c) and cell death by apoptosis (Tirlapur, 2001). NIR induces cell death of cancer cells and bone marrow cells (Tanaka, 2011e), which may have a potential application in the treatment of cancer (Tanaka, 2010c; Tanaka, 2012c).

The necessity to protect against NIR has not been well investigated to date. Fair skin with lower concentrations of melanin and a thin dermis might allow NIR to penetrate deeper into human tissue, and damage superficial muscles compared to darker skin with dense melanin and a thick dermis (Tanaka, 2011c; Tanaka, 2012a, b). In addition to natural NIR, human skin is increasingly exposed to artificial NIR from medical devices and electrical appliances (Schieke, 2003; Schroeder, 2008). Furthermore most sunscreens only block UV, but not visible light and NIR (Tanaka, 2011c; Tanaka, 2012a, b), and sunglasses and glasses are unable to block NIR exposure. Thus, individuals using sunscreens and glasses should further equip themselves for protection against NIR (Darvin, 2010; Meinke, 2011; Pujol, 1993; Schieke, 2003; Schroeder, 2008; Tanaka, 2011c; Tanaka, 2012a, b), as we are exposed to tremendous amounts of NIR (Tanaka, 2011c).

Previous studies regarding NIR have reported its application in the industrial and the agricultural fields, but have not well investigated the effects of NIR exposure in the medical fields, particularly in the field of plastic surgery.

Findings from our previous studies suggest that we should consider the biological effects of NIR, which have both potentially useful applications and detrimental physiological effects.
Methods to Evaluate Non-Thermal Biological Effects of NIR

In previous studies of NIR, a light source emitting wide wavelengths of NIR were used as an NIR source (Frank, 2004; Tanaka, 2012a). The temperature of the superficial layer of a culture fluid in a laboratory dish under NIR irradiation rises immediately, as NIR is mainly absorbed by water. The energy of NIR then decreases as it penetrates deeper, and will not reach enough target cells in the base. Therefore, these previous studies were only able to describe the optical and thermal effects of NIR, and are not useful to examine non-thermal biological effects of NIR (Tanaka, 2012a, b).

NIR induces dermal heating, which results in the tightening of skin laxity (Bitter, 2000; Chan, 2008; Goldberg, 2000; Ross, 2000; Tanaka, 2009a, b; Zelickson, 2006). In previous studies (Danno, 2001; Kim, 2005; Kligman, 1982) NIR devices without a water filter or contact cooling were used to investigate thermal effects on the human body. NIR increases the skin surface temperature and induces perspiration and vasodilation, as NIR is primarily absorbed by water and hemoglobin. Subsequently, a substantial amount of energy is absorbed in the superficial layers of skin, and only limited NIR energy can be delivered to deeper tissues. Therefore, these previous studies only describe superficial and thermal effects of NIR, and are not useful to examine non-thermal biological effects (Tanaka, 2012a, b).

Sunlight that reaches the human skin contains solar energy composed of 6.8% UV light, 38.9% visible light and 54.3% infrared (IR) (Kochevar, 1999). The IR spectral region is arbitrarily divided according to wavelength into sub-regions of NIR (760–3000 nm), middle IR (3000–30 000nm), and far IR (30 000 nm–1 mm). NIR from the sun is selectively filtered by atmospheric water (Anderson, 1981; Gates, 1966), and NIR that reaches the Earth’s surface readily penetrates the superficial layers of the skin (Tanaka, 2010a, 2011c, 2012a, 2011e) (Fig. 1, left).

Wavelengths below 1100 nm will be absorbed by melanin in the skin. Wavelengths between 1400 and 1500 nm and those above 1850 nm will be absorbed by water in the skin, which results in heating and may induce painful sensations and burns (Kelleher, 1999). Filtering out the wavelengths below 1100 nm, around 1450 nm, and above 1850 nm enabled the delivery of NIR to deeper tissues (Davenport, 2006) and also simulated the solar NIR that contracts human skin under natural conditions.

An NIR device that emits a spectrum of NIR wavelength from 1100 to 1800 nm together with a water-filter that excludes wavelengths between 1400 and 1500 nm as well as those above 1850 nm simulates the natural exposure of skin to NIR and allows for the evaluation of solar NIR that reaches the skin under natural conditions. NIR increases the surface temperature and induces thermal effects, thus, in order to reduce the skin surface temperature, perspiration, and blood vessel dilation, contact cooling is required. These specific wavelengths and the cooling system enables NIR to be delivered to the deeper tissues without pain or epidermal burns (Davenport, 2006; Goldberg, 2007), and allows for the investigations of properties of NIR (Tanaka, 2012a, b) (Fig. 1).
Discussion

Properties and Chromophore of NIR

NIR exhibits both wave and particle properties and it is strongly absorbed by water, hemoglobin, and myoglobin (Tanaka, 2011c). As a consequence, NIR can affect the subcutaneous tissues, including muscles, with its high permeability (Tanaka, 2011c) (Fig. 2).

The NIR spectrum is a result of the overtones and combination of bond stretching vibrations from O-H, C-H, and N-H groups (Weyer, 1985). Water is a polar molecule possessing hydrogen bonds with an electrical dipole moment, and will be resonated by NIR. NIR will be resonated and absorbed in O-H intramolecular hydrogen bonds and the electrical dipole moment (Tsai, 2001).

NIR induces an increase in water retention in the dermis and induces perspiration, vasodilation, and the expression of collagen and elastin (Tanaka, 2009a). Both collagen and elastin possess helical structures rich in hydrogen bonds. Humans have biological defense mechanisms in which induced hydrogen bonds and helical structures are resonated by NIR and absorb NIR to protect the subcutaneous tissues against this radiation.

NIR non-thermally induces degeneration of myoglobin, resulting in apoptosis of vascular smooth muscle cells and long-lasting vasodilation (Tanaka, 2011a). Both hemoglobin and myoglobin have heme binding sites and alpha helices. Our results showing long-lasting muscle thinning and vasodilation induced by NIR suggest that NIR might resonate and damage heme. However, our collagen, elastin, and cancer studies suggest that NIR may mainly resonate hydrogen bonds, helical structures, alpha helices, and DNA (Tanaka, 2011c). Alpha helices are thought to be resonated by NIR (Nevskaya, 1976). Both hemoglobin and myoglobin are oxygen-carrying proteins with many alpha helices. It is possible that NIR induces resonance of helical structures in the oxygen-carrying proteins and degenerates proteins containing hydrogen bonds and helical structures, which result in damage to the storage and transport of oxygen. This could be one of the mechanisms of apoptosis (Tanaka, 2011c).
NIR induces photochemical changes and affects a large volume and depth of tissue (Anderson, 1981). Actively proliferating cells show increased sensitivity to NIR (Karu, 1994; Tafur, 2008), and NIR induces strand breaks and apoptosis (Tirlapur, 2001) as well as death of cancer and bone marrow cells (Tanaka, 2010c, 2011e, 2012c). NIR is used to treat wound healing disorders (Danno, 2001; Horwitz, 1999; Schramm, 2003) and malignant tumors (Bäumler, 1999; Dees, 2002; Kelleher, 1999; Orenstein, 1999). While NIR appears to damage tumor tissue, it has also been shown to reduce cellular protein damage produced by biological oxidants in normal cells (Kujawa, 2004).

Subcutaneous adipocytes located above superficial muscles are effective and reasonable for the temperature retention and protection from NIR, because fatty tissue can scatter NIR optically (Tanaka, 2010a; Tanaka, 2011b) and fatty acids are the major NIR absorbing materials in soft tissues (Tsai, 2001). Oil in the liquid phase is transparent, whereas oil in the solid phase is highly scattering to NIR (Van Veen, 2005). The long-lasting induction of subcutaneous adipocytes may protect the underlying tissues, including the panniculus carnosus in animal and superficial muscles in human, against NIR damage. Because muscles contain hemoglobin and myoglobin which strongly absorb NIR, muscles are easily damaged by NIR (Srinivasan, 2003).

We previously reported that NIR thickens the dermis and induce the expression of collagen (Tanaka, 2009b), elastin, and water-binding proteins (Tanaka, 2009a, b) without scar formation (Tanaka, 2010b) to increase the amount of fluid protecting subcutaneous tissues from NIR. Further, NIR non-thermally induces vasodilation (Tanaka, 2011a), muscle thinning (Tanaka, 2010a), muscle relaxation (Tanaka, 2011b), bone marrow damage (Tanaka, 2011e), increased superficial granular adipocytes (Tanaka, 2011e), a cytocidal effect on cancer cells (Tanaka, 2010c, 2012c), stimulation of stem cells (Tanaka, 2011c,e), and DNA damage of mitotic cells (Tanaka, 2010c, 2012c).

Other than the skin, the optic nerve which is the only exposed component of the central nervous system also has a defence mechanism against NIR on multiple levels; rich blood flow in the retina, hyaluronic acid in the lens, tears in sclera, and fat in the eyelid. These are effective protective mechanisms, as water, hemoglobin, and fat are all ideal materials for blocking exposure to and consequences of NIR.

Further studies are needed to investigate the range and mechanisms of the biological effects of NIR in humans.

**Beneficial Effects of NIR**

NIR induces dermal heating and non-thermally induces collagen and elastin stimulation, resulting in skin tightening (Tanaka, 2009a, b). NIR denatures collagen and promotes the generation of new collagen, resulting in tighter skin (Goldberg, 2007; Zelicke, 2006), and induces elevated levels of collagen production markers (Orringer, 2005). NIR also increases the amounts of collagen in human dermal fibroblasts, and can result
in clinical improvement of skin texture (Lee, 2006). NIR induces high collagen density in the dermis, resulting in epidermal smoothness without scar formation, which provides safe, consistent, and long-term effects of skin rejuvenation (Tanaka, 2010b).

Pre-exposure of NIR prevents UV-induced toxicity (Danno, 1992; Frank, 2004; Menezes, 1998), and this effect is independent of heat shock protein induction and cell division (Menezes, 1998). These findings support the hypothesis that NIR prepares skin to better resist the subsequent damage from UV or NIR (Tanaka, 2012a, b).

NIR is known to be of therapeutic benefit in the treatment of musculoskeletal disorders and healing of indolent wounds (Ceylan, 2003; Webb, 1998). NIR induces long-lasting vasodilation for an increase in blood circulation by causing apoptosis of vascular smooth muscle cells, which may prevent vasospasm and may be beneficial for ischemic disorders and flap surgeries.

NIR increases mitochondrial metabolism (Karu, 1999; Passarella, 1984; Yu, 1997; Wilden; 1998), facilitates wound healing, and promotes angiogenesis in skin (Conlan; 1996), bone (Yaakobi, 1996), nerve (Assia, 1989), and skeletal muscle (Bibikova, 1994; Oron, 2006).

NIR also non-thermally relaxes and weakens dystonic or hypertrophic muscles to reduce wrinkles and myalgia. Thus, this simple technique of using NIR may offer an alternative method to relax superficial muscles.

We previously reported that NIR increased subcutaneous adipocytes on the panniculus and CD34-positive cells surrounding the subcutaneous adipocytes. Adipogenesis is tightly associated with angiogenesis, and the expression of adipocytes is linked to the development of its vasculature (Christiaens, 2010). Thus, induction of subcutaneous adipocytes may have an application for volume augmentation.

Our previous study revealed that NIR increased subcutaneous and bone marrow adipocytes, CD34-positive hematopoietic stem cells in bone marrow, and cortical bone mass (Tanaka, 2011e). Stimulation of stem cells by NIR irradiation might be beneficial for applications in regenerative medicine.

NIR is also an essential tool in cancer detection and imaging, as actively proliferating cells show increased sensitivity to NIR (Karu, 1994; Tafur, 2008). NIR also induces non-thermal DNA damage of mitotic cells especially in prophase, metaphase, and anaphase due to the absence of nuclear lamin protection (Tanaka, 2010c, 2012c), and suppresses the proliferation of various kinds of malignant cells (Tanaka, 2010c, 2011c, 2012c).

NIR may be beneficial in multiple fields of plastic surgery, as the schedule reduces discomfort and side effects, deep subcutaneous tissues are accessible, and facilitates repeated irradiations.

These findings indicate that NIR has a wide range of biological effects, and future studies are warranted to develop these findings into potential useful and beneficial techniques and applications.
**Deleterious Effects of NIR**

NIR exerts biologic effects on human skin (Schieke, 2003). NIR causes skin changes similar to those observed in solar elastosis, and enhance UV-induced dermal damage (Kligman, 1982). NIR activates mitogen-activated protein kinases and induces gene transcription, and it is likely to increase collagen degradation (Kim, 2006; Schieke, 2003; Schroeder, 2008). Epidemiological data and clinical reports imply that NIR is not innocuous to human skin (Dover, 1989; Kligman, 1984; Schieke, 2003).

Several diseases such as rosacea and erythema ab igne might be induced by chronic NIR exposure. Rosacea is a chronic cutaneous disorder characterized by centrofacial persisting erythema, telangiectasia, papules, pustules, edema, and ocular involvement. Rosacea affects all races, though is more common in Caucasians and fair-skinned populations (Berg, 1989). NIR should be considered as a critical factor in the development and aggravation of rosacea, since the distribution of erythema is most prominent on the facial convexities (Bae, 2009).

Long-term exposure to sources of heat and NIR, such as fires and stoves, results in erythema ab igne (Findlayson, 1966), which is clinically characterized by a reticular hyperpigmentation and telangiectasia accompanied histologically by epidermal atrophy, vasodilation, dermal melanin and hemosiderin deposits. Long-term exposure of NIR from various heat sources is thought to induce reticulated erythema and results in histopathological changes similar to those seen in solar-damaged skin (Page, 1988). The occurrence of telangiectasia appeared to increase with age, increased sunbathing, and poor pigmentation ability (Berg, 1989).

These lesions may develop thermal keratosis, such as hyperkeratosis, keratinocyte dysplasia, and dermal elastosis, which are similar to the changes that occur in actinically damaged skin (Arrington, 1979). Several reports indicate that carcinomas arise from heat-induced erythema ab igne (Hewitt, 1993; Jones, 1988; Kligman, 1984). Similar to UV, NIR induces photoaging and potentially photocarcinogenesis (Schieke, 2003). In addition, skin tumors in mice appeared faster after irradiation with the full lamp spectrum containing UV, visible, and NIR compared to irradiation with UV alone (Bain, 1943).

NIR is attenuated by thick water-containing dermis. Thus, skin with sparse melanin and a thin dermis allows NIR to penetrate deeper into tissue than skin with dense melanin and a thick dermis (Tanaka, 2011c; Tanaka, 2012a, b). The mean area of the facial surface that is covered with wrinkles is significantly larger in Caucasians than in African Americans, and characteristics of age-related periorbital changes seem to occur at a more accelerated rate in Caucasians (Odunze, 2008). In addition, fair skin is more sensitive to skin aging (Guinot, 2002; Nagashima, 1999). These findings support the observation that fair skin tends to wrinkle and sag earlier in life (Rawlings, 2006; Tsukahara, 2004), as fair skin is thinner and is more susceptible to NIR damage to the underlying superficial muscles than dark skin (Tanaka, 2011c; Tanaka, 2012a).
Continual long-term exposure to solar NIR causes superficial muscle thinning and the muscle extensions to the dermis, which ultimately leads to skin ptosis (Fig. 4). Additional factors thought to contribute to brow ptosis include the gradual loss of forehead skin elasticity and a reduction in the tone of the frontalis muscles (Knize, 1996; Niechajev, 2004). The use of NIR for smoothing forehead wrinkles also caused brow ptosis. Although further studies are required to confirm our results, these results may have major implications in superficial tissue aging and skin ptosis.

Plastic and reconstructive surgeons are usually exposed to NIR from the microscope for an extended period of time during microsurgery, thus, surgeons should consider protection of eyes from NIR. NIR that is absorbed by the anterior segment (the cornea, aqueous, and lens) can produce clouding of the cornea and cataract (Aly, 2011; Lydahl, 1984; Zaret, 1976). The International Commission on Non-Ionizing Radiation Protection recommended protection to avoid the thermal injury of the cornea and possible cataractogenesis.

Fig. 4: Superficial Tissue Aging Induced by Solar NIR Radiation

Conclusion

In order to simulate solar NIR that reaches the skin and to investigate the properties of NIR, the NIR source must possess a water filter and contact cooling.

Appropriate NIR induces dermal heating thermally and non-thermally induces collagen and elastin stimulation, which results in tightening of skin laxity. NIR also non-thermally induces long-lasting vasodilation, which may prevent vasospasm and may be beneficial for ischemic disorders, vasospasm, and flap surgeries. Further, NIR relaxes and weakens dystonic or hypertrophic muscles to reduce wrinkles and myalgia, which might have an application for treating muscle disorders, and causes long-lasting induction of subcutaneous adipocytes, which may have an application for volume augmentation. NIR might be beneficial for regenerative medicine based on its stimulation of stem cells. Cancer detection and imaging may gain from NIR, due to cancer's high sensitivity to NIR, and NIR induction of non-thermal DNA damage of mitotic cells could be useful in cancer treatment. These new techniques using NIR may be beneficial in many fields of plastic surgery.

Importantly, solar NIR may cause unexpected photoaging, muscle thinning and stimulation of stem cells, including cancer stem cells, in areas of the body exposed to the sun. Although various kinds of sunscreen materials are often used to prevent skin damage from UV exposure, these materials block neither visible light nor NIR. Therefore, skin should be protected with clothing or sunscreen that not only blocks UV, but also NIR, in order to prevent photoaging, skin ptosis, and oncogenicity. However, additional non-
thermal studies are needed to investigate the biological effects of NIR in humans.

Disclosure

None of the authors of this study have a conflict of interest.

References


