RESEARCH ARTICLE

Higher pulse frequency of near-infrared laser irradiation increases penetration depth for novel biomedical applications

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Abstract

Background

The clinical efficiency of laser treatments is limited by the low penetration of visible light used in certain procedures like photodynamic therapy (PDT). Second Harmonic Generation (SHG) PDT is an innovative technique to overcome this limitation that enables the use of Near Infrared (NIR) light instead of visible light. NIR frequency bands present an optical window for deeper penetration into biological tissue. In this research, we compare the penetration depths of 405 and 808 nm continuous wave (CW) lasers and 808 nm pulsed wave (PW) laser in two different modes (high and low frequency).

Methods

Increasing thicknesses of beef and chicken tissue samples were irradiated under CW and PW lasers to determine penetration depths.

Results

The 808 nm CW laser penetrates 2.3 and 2.4 times deeper than the 405 nm CW laser in beef and chicken samples, respectively. 808 nm PW (pulse frequency—500 Hz) penetrates deeper than CW laser at the same wavelength. Further, increasing the pulse frequency achieves higher penetration depths. High frequency 808 nm PW (pulse frequency—71.4 MHz) penetrates 7.4- and 6.0-times deeper than 405 nm CW laser in chicken and beef, respectively.

Conclusions

The results demonstrate the higher penetration depths of high frequency PW laser compared to low frequency PW laser, CW laser of the same wavelength and CW laser with half the wavelength. The results indicate that integrating SHG in the PDT process along with pulsed NIR light may allow the treatment of 6–7 times bigger tumours than conventional PDT using blue light.
Introduction

A significant contribution/application of laser medicine is Photodynamic therapy (PDT) which uses light and a chemical substance called photosensitizer (PS) to generate molecular oxygen and elicit cell death. This phenomenon of phototoxicity has been a promising development with various applications in microbial detection and treatment of infections [1,2] involving even antibiotic resistant bacteria [3], enhanced wound healing [4] and cancer therapy [5]. As a clinically approved minimally invasive therapeutic procedure PDT exerts a selective cytotoxic activity toward malignant cells with demonstrated effectivity in cancers [6,7]—of the skin, head and neck, digestive system [8], urinary system [9], lungs, etc. PDT results in significantly reduced tumour weights compared to untreated samples [3], reducing the number of clonogenic tumour cells. However, its clinical effectivity remains limited to treatment of small tumours [6]. Various technical aspects of the PDT treatment are accountable for the limited effect of PDT in treating large tumours [10,11] with penetration depth of radiation used being a key contributory factor.

Conventional PDT employs the visible electromagnetic (EM) spectrum within the range of 400–750 nm [12,13] to excite the corresponding photosensitisers. However, penetration of visible light through biological tissue is very limited. While red light can penetrate about 6 mm beneath the surface of the skin, blue light penetrates barely 1 mm into tissue [14]. It was shown that the transmittance of visible light through biological tissue increases with increasing wavelength [15,16]. Measurements of light intensity transmittance have demonstrated that the optical penetration depth in biological tissue does not depend on the optical power [17]. Light has its maximum depth of penetration in tissue within the range of 750–950 nm termed as the Near-infrared (NIR) window [18], which has been effectively used for optical imaging in live animals [19]. Thus, a system utilising this band is expected to be more effective in PDT treatment [20,21].

Power transmission in Low Level Laser Therapy (LLLT) decreases with increasing tissue thickness of skin, fat, and muscles [22]. Tissue transmission associated attenuation coefficient was the highest for 660 nm, minimal for 830 nm and the least for 904 nm [22]. Optical transmission of controllable infrared source parameters demonstrates maximum efficiencies of 86% [23] with higher duty cycles transmitting greater power over shorter exposure times while lower duty cycles transmit higher power at lower temperatures. Variations in efficiency of transmission indicate that the power transmission percentage can be appropriately tuned.

Second Harmonic Generation (SHG) is an optoelectronic process mediated by nonlinear material which produces photon up conversion by frequency doubling (or wavelength halving). SHG can be used to allow NIR light to generate visible light of appropriate wavelength to excite PSs and produce the photodynamic effect [24]. This technique overcomes the low penetration depth of visible light while allowing the same photosensitisers to be used effectively in conjunction with NIR light.

NIR light in the range 770–850 nm, projected transcutaneous in Sprague–Dawley rats demonstrated the highest transmission (penetration) through all overlying tissue layers of the spinal cord and blood. 810 nm laser with 150 mW incident power translated into 9 mW reaching the spinal cord through overlying thickness of 18 mm (6% power transmission) [25]. Light at 808 nm penetrates as much as 54% deeper than 980 nm light [26]. In bovine tissue, light at 808 nm with an Initial Power Density (IPD) of 1 W/cm² attenuates over a depth of 3.4 cm delivering 1 mW/cm² in situ [26]. Hence, NIR and especially 808 nm is expected to increase the efficacy of PDT.
Pulsed wave (PW) laser irradiation provides added advantages over continuous wave (CW) lasers. The high instantaneous power fluxes in PW provide efficient energy transfer [27] and the interval between pulses allows for heat dissipation to avoid thermal ablation [28].

In this study we investigated the penetration depths of 808 nm PW in two different modes; high frequency (HF) and low frequency (LF), 808 nm CW laser and 405 nm (the corresponding wavelength for direct excitation of the PS) CW laser using different meat samples representing different kinds of tissue.

Materials and methods

Sample preparation

The Beef (rib) and Chicken (breast) samples were purchase from "Campus Market" (Building 102, 27 Ramat Hagolan Street, Ariel). The samples were kept frozen and the amount needed was thawed to room temperature before each experiment. Beef meat is darker than chicken meat (The appearance of the samples can be seen in Fig 1) due to higher protein content [29]. This contrast was used to represent different types of human internal tissues like muscles and non-muscles respectively for the purpose of measuring penetration depths. Sliced sections of either Beef or Chicken (measuring 2 mm, 4 mm, 6 mm, 8 mm, 10 mm, 12 mm, 14 mm and 18 mm. Thickness error: ±0.2 mm) were placed within a screw-in sample holder made of acrylic glass (providing optical transparency in the bandwidth of the spectrum investigated) with appropriate spacers (Fig 1).

Sample irradiation

Samples placed within the acrylic glass meat holder were irradiated with an 808 nm and 405 nm continuous diode lasers and 808 nm PW (Astrella by Coherent) which is normally used to drive a photo-cathode RF e-gun [30]. We used the PW in two different modes: LF (repetition rate—500 Hz, pulse duration—50 fs, spot diameter—11 mm) and HF (repetition rate—71.4 MHz, pulse duration—5 fs, spot diameter—3 mm). The average power, peak power, energy per pulse, and power density of the LF and HF pulsed laser were 2.55 W, 102 GW, 5.1 mJ, 2.68 W/cm² and 216 mW, 605 W, 3.0 nJ, 3.06 W/cm² respectively. In the case of the 404 nm and 808 nm continuous lasers average powers were 135 mW and 165 mW respectively. The parameters of the lasers used for the irradiation of the samples are summarized in Table 1. Transmitted powers were measured before (transmittance through blank acrylic glass holder—\(I_0\)) and after (transmittance through acrylic glass holder with meat/chicken slices—\(I\)) sample insertion and normalised to those of blank acrylic glass meat holder for stringency of analysis. Power measurements were done using a Newport optical power meter (Model 840) and Newport 883SL photodiode sensor (OD 3) for the CW lasers and Coherent power detector/sensor (OD-COHERENT PowerMax—USB PM10) for the PW.

Penetration depth analysis

Beer-Lambert law explaining the exponential attenuation of electromagnetic spectra transmitted through tissue was used to compute the optical penetration depth. Using this law, transmittance (\(T\)) as a ratio of the transmitted light (\(I\)) to the incident light (\(I_0\)) is derived by the equation:

\[
T = \frac{I}{I_0} = e^{-\mu d}
\]

where \(\mu\) is the attenuation coefficient and \(d\) is the thickness of the tissue sample. Penetration
depth ($\delta$) is defined as the depth at which the transmittance inside the tissue equals “$1/e$". Therefore, transmittance can be expressed in terms of the penetration depth as:

$$T = e^{-d/\delta}$$

For each laser, transmittances have been plotted as a function of tissue thickness. Each point in the graph is an average of 2–3 measurements. Each graph has been fitted to the function of $e^{-x/\delta}$ to get the penetration depth ($\delta$) and error ($\Delta \delta$). The least squares analysis was done using MATLAB R2019a software.

**Results**

*808 nm laser penetrates deeper than 405 nm under continuous wave irradiation*

Beef and chicken samples were irradiated with 405 nm and 808 nm CW lasers at 135 mW and 165 mW incident powers respectively and the associated transmittance with varying levels of tissue thickness analysed. 405 nm CW laser reaches a penetration depth of 0.28 ± 0.03 mm in beef samples (Fig 2) while 808 nm CW laser reaches a penetration depth of 0.63 ± 0.16 mm. In chicken samples 405 nm CW laser reaches a penetration depth of 0.48 ± 0.07 mm (Fig 3) while 808 nm CW laser reaches a penetration depth of 1.13 ± 0.21 mm.

| Pulse frequency | Power density | Peak power | Energy per pulse | Average power |
|-----------------|---------------|------------|-----------------|---------------|
| CW 1            | N/A           | N/A        | N/A             | 135 mW        |
| CW 2            | N/A           | N/A        | N/A             | 165 mW        |
| PW LF           | 500 Hz        | 2.68 W/cm² | 102 GW          | 2.55 W        |
| PW HF           | 71.4 MHz      | 3.06 mW/cm²| 605 W           | 216 mW        |

Parameters of the lasers used for irradiation of the samples. **CW 1—405 nm; CW 2—808 nm; PW LF—808 nm, 50 fs; PW HF—808 nm, 5 fs.**

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808 nm pulsed laser irradiation penetrates deeper than continuous wave irradiation

Beef and chicken samples were irradiated with pulsed 808 nm laser (500 Hz repetition rate, 50 fs pulse width) and the associated transmittance with varying levels of tissue thickness were
compared with that of the CW irradiation. Pulsed 808 nm reached a penetration depth of 1.41 ± 0.41 mm in beef samples (Fig 2) while CW 808 nm reached a lower penetration depth of 0.63 ± 0.16 mm. In chicken samples pulsed 808 nm reached a penetration depth of 3.24 ± 0.48 mm (Fig 3) while CW 808 nm reached a lower penetration depth of 1.13 ± 0.21 mm.

### High frequency PW irradiation penetrates deeper than low frequency pulsed laser irradiation

Beef and chicken samples were irradiated with HF 808 nm PW (71.4 MHz, 5 fs) and the associated transmittance with varying levels of tissue thickness were compared with that of the LF 808 nm PW (500 Hz, 50 fs) irradiation. HF pulsed 808 nm reached a penetration depth of 1.70 ± 0.43 mm in beef samples (Fig 2) while LF pulsed 808 nm reached a lower penetration depth of 1.41 ± 0.41 mm. In chicken samples HF pulsed 808 nm reached a penetration depth of 3.56 ± 0.34 mm (Fig 3) while LF pulsed 808 nm reached a lower penetration depth of 3.24 ± 0.48 mm. Table 2 summarizes all the measured penetration depths.

### Discussion

As discussed in the introduction, the power of the laser does not affect the penetration depth [14]. Therefore, differences in penetration depth must be related to the wavelength, laser type (CW or PW) and the frequency of the PW. The experiments demonstrate that transmittance of the CW laser passing through chicken samples is 2-times greater on average than through beef samples. Chicken samples are of a lighter color compared to beef accountable to a higher myoglobin content in the latter [29]. Further, transmittance of both high and LF type pulsed lasers passing through chicken samples are also about 2-times greater than through beef samples. This observation corroborates with the trend of increased transmission as one moves from mainly muscular tissue to mainly fatty tissue [31].

The results from CW laser irradiation consistently show that the penetration of NIR 808 nm wavelength (0.63 ± 0.16 mm in beef and 1.13 ± 0.21 mm in chicken) is more than twice higher than the penetration of visible range 405 nm wavelength (0.28 ± 0.03 mm in beef and 0.48 ± 0.07 mm in chicken). Transmittance of light through any specific thickness of tissue improves when a PW is used in contrast to a CW laser, operating at the same wavelength of 808nm. For beef rib samples, 808nm PW–HF and LF irradiation resulted in 2.7- and 2.2-times improved transmittance than that of the CW respectively. For chicken breast, 808nm PW–HF and LF irradiation resulted in 3.2- and 2.9-times improved transmittance than that of the CW respectively.
HF 808 nm PW irradiation resulted in a 20% increase in transmittance through beef rib samples with respect to LF 808 nm PW exposure. In chicken samples, transmittance increased by 10% when HF pulsed irradiation replaced the LF pulsed exposure at 808 nm. HF 808 nm PW penetrates 7.4- and 6.0-times deeper than 405 nm CW laser in chicken and beef, respectively. Increasing diameter of laser irradiation beams are known to account for greater penetration. However, in this case our experiments involved a 3 mm spot diameter for the HF laser and a 11 mm spot diameter for the LF laser (ref. Methods).

A possible explanation for the results is that the high photon density of the PW laser (compare to the CW laser) excites a large percentage of electrons in the sample (close to saturation), rendering the irradiated zone transiently transparent for the rest of the incoming photons. This phenomenon is also indicated by the observed increase in transmission associated with increasing the pulse frequency in the PW regimes. For both HF and LF the pulse duration (5 fs and 50 fs, respectively) is shorter than the lifetime of the excited state, estimated as $10^{-7}$ to $10^{-5}$ s. However, the difference in the repetition rates between the two modes might affect the percentage of the excited electrons in the irradiation area. The low repetition rate (time between pulses = 2 ms) of the LF laser guarantees that the next pulse will reach the sample only after all the excited electrons returned to the ground state. However, due to the high repetition rate (time between pulses = 14 ns) of the HF laser the next pulse will reach the sample while the electrons, excited by the previous pulse, are still in the excited state. This will accumulate electrons in the excited state leading to higher transient transparency as compared to the LF irradiation.

The 4 main NIR optical windows useful for biomedical imaging and therapeutics are classified into bands from—750 to 950 nm, 1100 to 1350 nm, 1600 to 1870 nm and 2100 to 2200 nm [32]. The first window presents the lowest levels of absorption by tissue water as compared to the remaining three bands. Despite this property, often image quality is reduced due to stronger absorption peaks from lipids, hemoglobin and deoxyhemoglobin resulting in blurring due to the molecular process of Rayleigh/Mie scattering [33]. This has necessitated development of special longer NIR wavelength CCD detectors and equipment to overcome the described problems [32] in order to utilize the second, third and fourth optical windows. In contrast, our results demonstrate that increased tissue transparency can be achieved by simply using higher pulse frequencies within the first NIR optical window itself allowing deeper penetration and ease of application.

Scattering constitutes a dominant factor determining tissue penetration within the NIR optical windows [34] with absorbed photons getting trapped in the tissue while other scattered photons either get absorbed or scattered and are detectable at appropriate distance(s) from the incident light source. However, scattering exhibiting a weak dependence on wavelength; the effective use of the NIR optical windows are primarily limited by absorption of blood at shorter wavelengths and water at longer wavelengths [33]. The common minimum absorption trough of the major tissue components (water, deoxygenated and oxygenated blood, melanin, etc.) lies within the 650–900 nm bandwidth [34] which influence the in-vivo optical properties of tissues pertaining to PDT treatment(s). However, due to the large variations of the absorption and scattering parameters biological tissue are not uniformly absorbing and scattering objects [35]. As described in the discussion the higher repetition rate (time between pulses) of the HF laser allows each subsequent pulse to reach the sample while the electrons excited by the previous pulse are still in the higher energy state, enabling an accumulation of excited state electrons leading to transient transparency as compared to the LF irradiation. This phenomenon along with the higher number of photons being delivered by the HF laser overcomes the limits of scattering dependent interactions [34] providing the improved penetration depths as demonstrated by our experiments.
Our experiments demonstrate that 808 nm lasers penetrate deeper than 405 nm CW laser for any given tissue. Irradiation of biological tissue(s) by lasers result in four main types of interactions: reflection, absorption, scattering and transmission [36]. Pogue et al reported some greater effective penetration for pulsed light, but they attributed this to transient alteration of absorption by the PSs [37]. In contrast, our experiments involved no PSs. The results demonstrate a novel discovery that NIR PW irradiation improves the associated penetration depth significantly in contrast to CW exposures. In addition, higher pulse frequency (repetition rate) further enhances the penetration depth compare to lower frequency of the same wavelength. This phenomenon encourages further investigations for novel processes of laser tissue interactions and related applications [36].

Conclusions

Laser power transmission associated attenuation for cancerous tissues being lower than that of normal tissues over all the 4 NIR optical windows [32], our results suggest that adoption of higher pulse frequency irradiation regimes can effectively overcome the penetration depth limitation of using PDT for treating large tumors [10,11].

In the emerging field of nanomedical imaging and laser mediated therapies PEGylated nanoparticles [38] and organic/inorganic nanohybrids [39] have improved drug delivery and circulation to targeted sites [40]. This has allowed for improvements in laser-based PDT. However, PDT is still limited to small tumors due to the penetration depth of the excited light. Besides overcoming the limited penetration depths of visible light, PDT mediated by NIR light also enables development of optoelectronic applications to improve clinical efficiency. SHG allows for conventional photosensitizers (usually operating under visible light) to be used with greater clinical efficiency [24,41] enabling the use of PDT for bigger tumors. Our results indicate that integrating SHG in the PDT process [24] will allow the excitation of PSs with NIR light pulsed appropriately as demonstrated by our experiments reaching 6–7 times higher penetration depths than conventional blue light. In addition, recent developments of using NIR absorbing chromophores [18] like BODIPY, cyanine, porphyrin, phthalocyanine, nano assemblies and etc., applied in conjunction to experimental indications of this study will enhance clinical performance of the photodynamic mechanism for selective ablation of tumors and enabling targeting of sub-cellular organelles. Clinical adaptations of these results which achieve greater penetration than conventional standards are expected to also improve applications of PDT for reducing microbial loads in dermatology [42] and in treatment of emerging diseases like COVID-19 [43].

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References
1. Demidova TN, Hamblin MR. Photodynamic Therapy Targeted to Pathogens. International Journal of Immunopathology and Pharmacology. 2004; 17(3):245–254. https://doi.org/10.1177/039463200401700304 PMID: 15461858
2. Xu C, Akakuru OU, Ma X, Zheng J, Zheng J, Wu A. Nanoparticle-Based Wound Dressing: Recent Progress in the Detection and Therapy of Bacterial Infections. Bioconjugate Chemistry. 2020; 31(7):1708–1723. https://doi.org/10.1021/acs.bioconjugchem.0c00297 PMID: 32538089
3. Dharmaratne P, Sapugahawatte DN, Wang B, et al. Contemporary approaches and future perspectives of antibacterial photodynamic therapy (aPDT) against methicillin-resistant Staphylococcus aureus (MRSA): A systematic review. European Journal of Medicinal Chemistry. 2020; 200:112341. https://doi.org/10.1016/j.ejmech.2020.112341 PMID: 32505848
4. Oyama J, Ramos-Milare CFH, Lera-No nose DSSL, et al. Photodynamic therapy in wound healing in vivo: a systematic review. Photodiagnosis and Photodynamic Therapy. 2020; 30:101682. https://doi.org/10.1016/j.pdpdt.2020.101682 PMID: 32032780
5. Hamblin MR. Photodynamic Therapy for Cancer: What’s Past is Prologue. Photochemistry and Photobiology. 2020; 96(3):506–516. https://doi.org/10.1111/php.13190 PMID: 31820824
6. Agostinis P, Berg K, Cengel KA, Foster TH, Girotti AW, Gollnick SO, et al. Photodynamic therapy of cancer: an update. CA: a cancer journal for clinicians. 2011; 61(4), 250–281. https://doi.org/10.3322/caac.20114 PMID: 21617154
7. Yang K, Zhang Z, Du J, Li W, Pei Z. Host–guest interaction based supramolecular photodynamic therapy systems: a promising candidate in the battle against cancer. Chemical Communications. 2020; 56 (44):5865–5876. https://doi.org/10.1039/d0cc02001j PMID: 32432243
8. Yano T, Wang KK. Photodynamic Therapy for Gastrointestinal Cancer. Photochemistry and Photobiology. 2020; 96(3):517–523. https://doi.org/10.1111/php.13206 PMID: 32676439
9. Bedi N, Reddy D, Ahmed HU. Targeting the cancer lesion, not the whole prostate. Translational Andrology and Urology. 2020; 9(3):1518–1525. https://doi.org/10.21037/tau.2019.09.12 PMID: 32676439
10. Minnes R, Weitman H, Ehrenberg B. The effect of lipid composition, bilayer phase and temperature on the uptake of hematoporphyrin by liposomal membranes. Journal of Porphyrins and Phthalocyanines. 2007; 11(08):577 –585. https://doi.org/10.1142/s1088424607000679
11. Allegra A, Pioggia G, Tonacci A, Musolino C, Gangemi S. Oxidative Stress and Photodynamic Therapy of Skin Cancers: Mechanisms, Challenges and Promising Developments. Antioxidants. 2020; 9(5):448. https://doi.org/10.3390/antiox9050448 PMID: 32455998
12. Minnes R, Weitman H, You Y, Detty MR, Ehrenberg B. Dithioporphyrin Derivatives as Photosensitizers in Membranes and Cells. The Journal of Physical Chemistry B. 2006; 112(10):3268–3276. https://doi.org/10.1021/jp0768423 PMID: 18278897
13. Abrahamse H, Hamblin MR. New photosensitizers for photodynamic therapy. Biochem J. 2016; 473 (4):347–364. https://doi.org/10.1042/BJ20150942 PMID: 26862179
14. Clement M, Daniel G, Trelles M. Optimising the design of a broad-band light source for the treatment of skin. Journal of Cosmetic and Laser Therapy. 2005; 7(3–4):177–189. https://doi.org/10.1090/14764170500344575 PMID: 16414906
15. Ash C, Dubec M, Donne K, Bashford T. Effect of wavelength and beam width on penetration in light-tissue interaction using computational methods. Lasers in Medical Science. 2017; 32(8):1909–1918. https://doi.org/10.1007/s10103-017-2317-4 PMID: 28900751
16. Jacques SL (2013). Optical properties of biological tissues: a review. Physics in medicine and biology, 58(11), R37–R61. https://doi.org/10.1088/0031-9155/58/11/R37 PMID: 23686068
17. Arslan H, Doluğan YB, Ay AN. Measurement of the Penetration Depth in Biological Tissue for Different Optical Powers. Sakarya University Journal of Science. 2018;1–1. https://doi.org/10.16984/ saufenbilder.332802
18. Smith AM, Mancini MC, Nie S. Bioimaging: second window for in vivo imaging. Nat Nanotechnol. 2009; 4(11):710–711. https://doi.org/10.1038/nnano.2009.326 PMID: 19898521
19. Frangioni JV. In vivo near-infrared fluorescence imaging. Curr Opin Chem Biol. 2003; 7(5):626–634. https://doi.org/10.1016/j.cpb.2003.08.007 PMID: 14580568

20. Xu X, Lu H, Lee R. Near Infrared Light Triggered Photo/Immuno-Therapy Toward Cancers. Frontiers in Bioengineering and Biotechnology. 2020; 8. https://doi.org/10.3389/fbioe.2020.00488 PMID: 32528941

21. Swamy PCA, Sivaraman G, Priyanka RN, et al. Near Infrared (NIR) absorbing dyes as promising photosensitizer for photo dynamic therapy. Coordination Chemistry Reviews. 2020; 411:213233. https://doi.org/10.1016/j.ccr.2020.213233

22. Barbosa RI, Guirro ECO, Bachmann L, Brandino HE, Guirro RRJ. Analysis of low-level laser transmission at wavelengths 660, 830 and 904 nm in biological tissue samples. Journal of Photochemistry and photobiology. B, Biology. 2020 Aug; 209:111914. https://doi.org/10.1016/j.jphotobiol.2020.111914 PMID: 32516626

23. Abbas HA, Triplett GE. Infrared Transmission Characteristics and Laser Tissue Interaction of Albino Pigskin Using Pulsed NIR Laser Light. International Scholarly Research Notices, vol. 2013, Article ID 451208, 8 pages, 2013. https://doi.org/10.1155/2013/451208.

24. Kachynski A, Pliss A, Kuzmin A, Ohulchansky TY, Baev A, Qu J, et al. Photodynamic therapy by in situ nonlinear photon conversion. Nature Photonics. 2014; 8(6):455–461. https://doi.org/10.1038/nphoton.2014.90

25. Byrnes KR, Waynant RW, Iliev IK, et al. Light promotes regeneration and functional recovery and alters the immune response after spinal cord injury. Lasers in Surgery and Medicine. 2005; 36(3):171–185. https://doi.org/10.1002/lsm.20143 PMID: 15704098

26. Hudson DE, Hudson DO, Wininger JM, Richardson BD. Penetration of Laser Light at 808 and 980 nm in Bovine Tissue Samples. Photomedicine and Laser Surgery. 2013; 31(4):163–168. https://doi.org/10.1089/pho.2012.3284 PMID: 23441909

27. Ando T, Xuan W, Xu T, et al. Comparison of Therapeutic Effects between Pulsed and Continuous Wave 810-nm Wavelength Laser Irradiation for Traumatic Brain Injury in Mice. PLoS ONE. 2011; 6(10). https://doi.org/10.1371/journal.pone.0026212 PMID: 22028833

28. Joensen J, Øvsthus K, Reed RK, et al. Skin Penetration Time-Profile for Continuous 810 nm and Superpulsed 904 nm Lasers in a Rat Model. Photomedicine and Laser Surgery. 2012; 30(12):688–694. https://doi.org/10.1089/pho.2012.3306 PMID: 23025702

29. Listrat A, Lebret B, Louveau I, et al. How Muscle Structure and Composition Influence Meat and Fleshy Quality. The Scientific World Journal. 2016; 2016:1–14. https://doi.org/10.1155/2016/3182746 PMID: 27022618

30. Nause A, Fridman A, Rosenzweig J, Fukasawa A, Roussel R, Spataro B. First Operation Of A Hybrid E-Gun At The Schlesinger Center For Compact Accelerators In Ariel University. In: Proceedings of IPAC2019. Geneva, Switzerland: JACOW Publishing; 2019:1171–1174.

31. Han D, Xu J, Wang Z, et al. Penetrating effect of high-intensity infrared laser pulses through body tissue. RSC Advances. 2018; 8(56):32344–32357. https://doi.org/10.1039/c8ra05285a

32. Sordillo LA, Pu Y, Pratavieira S, Budansky Y, Alfano RR. Deep optical imaging of tissue using the second and third near-infrared spectral windows. J Biomed Opt. 2014 May; 19(5):056004. https://doi.org/10.1117/1.JBO.19.5.056004 PMID: 24805808.

33. Puet al Y., “Two-photon excitation microscopy using the second singlet state of fluorescent agents within the tissue optical window,” J. Appl. Phys., 114 (15), 153102 (2013). http://dx.doi.org/10.1063/1.4825319.

34. Sandell JL, Zhu TC. A review of in-vivo optical properties of human tissues and its impact on PDT. J Biophotonics. 2011; 4(11–12):773–787. https://doi.org/10.1002/jbio.201100062 PMID: 22167862

35. Perekatova V, Subochev P, Kleshnin M, Turchin I. Optimal wavelengths for optoacoustic measurements of blood oxygen saturation in biological tissues. Biomed Opt Express. 2016; 7(10):3979–3995. Published 2016 Sep 13. https://doi.org/10.1364/BOE.7.003979 PMID: 27867709

36. Lin JT (2016) Progress of medical lasers: Fundamentals and applications Med Devices Diagn Eng 2: https://doi.org/10.10371/MDDE.1000111

37. Pogue BW, Lilge L, Patterson MS, Wilson BC, Hasan T (1997). Absorbed photodynamic dose from pulsed versus continuous wave light examined with tissue-simulating dosimeters. Appl. Opt. 36, 7257–7269. https://www.osapublishing.org/ao/fulltext.cfm?uri=ao-36-28-7257&id=61107. https://doi.org/10.1364/ao.36.007257 PMID: 18262435

38. Jokerst JV, Lobovkina T, Zare RN, Gambhir SS. Nanoparticle PEGylation for imaging and therapy. Nanomedicine. 2011; 6(4):715–728. https://doi.org/10.2217/nmn.11.19 PMID: 21718180

39. Liang C, Zhang X, Wang Z, Wang W, Yang M, Dong X. Organic/inorganic nanohybrids rejuvenate photodynamic cancer therapy. Journal of Materials Chemistry B. 2020; 8(22):4748–4763. https://doi.org/10.1039/d0tb00098a PMID: 32129418
40. Suk JS, Xu Q, Kim N, Hanes J, Ensign LM. PEGylation as a strategy for improving nanoparticle-based drug and gene delivery. Advanced Drug Delivery Reviews. 2016; 99:28–51. https://doi.org/10.1016/j.addr.2015.09.012 PMID: 26456916

41. Zhou X, Chen Y, Su J, Tian X, Luo Y, Luo L. In situ second-harmonic generation mediated photodynamic therapy by micelles co-encapsulating coordination nanoparticle and photosensitizer. RSC Adv., 2017, 7, 52125–52132. https://doi.org/10.1039/C7RA07334H

42. Queiros C, Garrido PM, Silva JM, Filipe P. Photodynamic therapy in dermatology: Beyond current indications. Dermatologic Therapy. 2020. https://doi.org/10.1111/dth.13997 PMID: 32654315

43. Dias LD, Bagnato VS. An update on clinical photodynamic therapy for fighting respiratory tract infections: a promising tool against COVID-19 and its co-infections. Laser Physics Letters. 2020; 17 (8):083001. https://doi.org/10.1088/1612-202x/ab95a9