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Abstract. The thermal effect may be a desired outcome or a concerning side effect in laser–tissue interactions. Research in this area is particularly motivated by recent advances in laser applications in diagnosis and treatment of neurological disorders. Temperature as a side effect also limits the maximum power of optical transfer and harvesting of energy in implantable neural prostheses. The main objective was to investigate the thermal effect of a near-infrared laser beam directly aimed at the brain cortex. A small, custom-made thermal probe was inserted into the rat brain to make direct measurements of temperature elevations induced by a free-air circular laser beam. The time dependence and the spatial distribution of the temperature increases were studied and the maximum allowable optical power was determined to be 2.27 W/cm² for a corresponding temperature increase of 0.5°C near the cortical surface. The results can be extrapolated for other temperature elevations, where the margin to reach potentially damaging temperatures is more relaxed, by taking advantage of linearity. It is concluded that the thermal effect depends on several factors such as the thermal properties of the neural tissue and of its surrounding structures, the optical properties of the particular neural tissue, and the laser beam size and shape. Because so many parameters play a role, the thermal effect should be investigated for each specific application separately using realistic in vivo models. © The Authors. Published by SPIE under a Creative Commons Attribution 3.0 Unported License. Distribution or reproduction of this work in whole or in part requires full attribution of the original publication, including its DOI. [DOI: 10.1117/1.JBO.19.1.015009]

Keywords: tissue–light interaction; near-infrared penetration; laser safety.

1 Introduction

Near-infrared (NIR) light finds applications in medicine for diagnostic and therapeutic purposes such as spectroscopic imaging¹ and the treatment of brain tumors.² Infrared light is also tested as a means of wireless power transfer in visual prosthetics.³ Temperature elevation is a complication that has not been addressed sufficiently in these studies. Our laboratory is currently investigating the feasibility of wireless neurostimulation in the central nervous system (CNS) using optically activated microstimulators in the NIR wavelengths.⁴⁵⁶ The project’s motivation is to solve well-known problems caused by the microwires that connect electrode arrays to the outside world.⁴ Microelectrode arrays often fail due to breakage of wire interconnects and the chronic tissue response caused by the tethering forces of the wires.

In humans, a body temperature of greater than 37.5 to 38.3°C is defined as hyperthermia; i.e., elevated body temperature due to failed thermoregulation that occurs when the body produces or absorbs more heat than it dissipates.⁸ Body temperatures exceeding 40.5°C, which usually result from extended exposure to environmental heat, cause a heat stroke. A study of temperature effects on the CNS argues that the brain loses its ability to thermoregulate at the body core temperatures exceeding 40.5°C and the irreversible multiorgan dysfunction begins.⁹ Hyperthermia in the 41 to 44°C range can cause damage in the CNS and neurological symptoms of a heat stroke in some animal models.¹⁰ However, the lasers used in the field of neural prosthetics will most likely involve local and steady temperature elevations in the CNS and the adverse effects should be investigated at the molecular level. For instance, a study performed on the rat spinal cord showed significant morphological and histochemical alterations and signs of diffuse spinal cord lesions of vascular origin with degeneration of nerve cells due to exposure to an ambient temperature of 43°C for 4 h.¹¹ On the other side, the whole or local brain hyperthermia studies indicate that a fluctuation of brain temperature within 4°C may be considered a normal physiological response depending upon the activation of the neurons and blood circulation.¹² The safe level of NIR exposure given by American National Standards for Safe Use of Lasers (ANSI Z136.-2007) does not list the dose for the neural tissue specifically.

Despite these reports that involved extreme temperature applications, others show that even small temperature elevations have observable effects. A study on stimulation induced thermal effects indicates that an increase of 0.5°C causes detectable changes at the cellular level, and 1°C increase can have profound effects.¹³ An implantable neural prosthetic device should be designed with the most conservative criteria considering a potential lifetime of many years. Therefore, we use a conservative value of 0.5°C as a design criterion in this study, although the results can be scaled for other temperature values.

Optical tissue properties (scattering and absorption coefficients) and the number of photons penetrating into the tissue determine the magnitude and the spatial pattern of energy deposition and hence the temperature distribution in neural tissue.¹⁴ The number of penetrating photons at a given location along the tissue surface is a function of the incident light beam parameters,

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e.g., cross-sectional intensity profile, total power, and exposure duration. Scattering is the dominant form of interaction inside the neural tissue at NIR wavelengths. Gray matter, in general, has a lower scattering coefficient than the white matter and as a result, NIR light penetrates deeper into the gray matter.

However, the optical parameters do not determine the temperature distribution alone. Once the optical energy is converted into heat through absorption, the heat spreads into medium by conduction as a function of time. The transient and steady-state temperature distribution depend on both optical and thermal parameters of the neural tissue in question. Although the main objective of this study was to specifically measure the temperature profile inside the rat brain under the exposure of an NIR laser beam with typical parameters needed to activate our microstimulators, the results can also provide practical insights for those who are investigating NIR light–neural tissue interactions in general. In this study, a micro-temperature probe was inserted into the brain cortex in anesthetized rats to directly measure the temperature rise inside in a two-dimensional (2-D) vertical plane while varying the relative position of the NIR laser beam. The maximum allowable light exposure was estimated that caused a threshold level temperature elevation in neural tissue.

2 Methods

2.1 Temperature Probe Preparation

A T type thermocouple (copper-constantan, T301, Unisense, Denmark) made of microwires (50 μm each wire) was wrapped around the tapering part of the tip of a tungsten microelectrode (shank diameter 125 μm) as a backbone for easy penetration into the brain (Fig. 1). The thermocouple wires were covered with a thin layer of epoxy for electrical insulation while maintaining a fast time response. The time constant of the final probe design was measured by dropping hot water droplets on the tip and found to be 60 ms.

2.2 Thermocouple Exposure to Light

Photons absorption directly on the thermocouple wires and the tungsten electrode carrier may result in an overestimation of temperature in tissue. The error introduced by exposure of the probe to the incident light was estimated by inserting the probe in a Petri dish filled with saline and positioning the laser source above the thermocouple junction at 13.5 cm [Fig. 2(b)], the same distance used in the animal experiments. Scattering and absorption in this medium were assumed to be negligibly small at the laser wavelength used (830 nm) and the saline layer above the probe was only 1 mm. On the other hand, the thermal conductivity and the specific heat capacity of saline closely mimic that of neural tissue (0.45 to 0.6 Wm$^{-1}$C$^{-1}$ and 3650 Jkg$^{-1}$C$^{-1}$, respectively). Thus, the saline bath imitated the thermal properties of the brain while eliminating the effects of optical scattering and absorption of light in the medium, which allowed us to assess the thermal effect of light on the probe without perturbing the temperature of the medium.

The steady-state temperature rise was measured as 1.4°C in the thermoprobe with its tip at a depth of 1000 μm in saline [Fig. 2(b)]. The light intensity was also measured at the same position in saline by replacing the probe with a 100 μm glass fiber [Fig. 2(a)], facing up toward the laser, which was connected to a photodiode at the other end. The light beam was attenuated by a small amount in saline (by 2.3%) as predicted. We normalized the in-saline temperature measurements with the experimentally measured values of light in the brain reported earlier in another study from our laboratory. More specifically, the light intensity at the same depth in the brain was about 1.8% of the intensity measured without the brain. This percentage multiplied by 1.4°C gave us the estimated temperature rise in the probe as a result of light exposure when it was in the rat brain. This error term (0.025°C) was <3% of the temperature elevation measured (~1°C) at this point in the tissue during in vivo experiments. Following the same method, the maximum thermocouple heating effect was estimated to occur at the surface and found to be 10%. The error was smaller in deeper regions since the light intensity decreases exponentially by depth (hence the exposure of the probe) while the temperature decline is more gradual due to heat conductance through tissue. It was decided that the temperature error due to light fallen on the probe was small enough to be neglected in this study for the sake of simplicity.

2.3 In Vivo Experiments

Six Sprague–Dawley rats (350 to 500 g) were used in this study. The anesthesia was induced with ketamine (80 mg/kg) and xylazine (12 mg/kg) mixture diluted with saline and further doses of ketamine were given as needed. Marcaine (0.2 mL) was injected at the site of incision as local analgesics. Dexamethasone (3 mg/kg, IM) was injected prior to surgery to reduce edema of the brain. The rectal temperature was continuously monitored and maintained at 36°C using a temperature regulated heating pad. The end-tidal CO$_2$ was observed to stay within the normal range throughout the experiment, which usually lasted about 6 to 7 h including the surgery. All the experimental procedures were approved by the Animal Care Committee at Rutgers University, New Jersey.

![Fig. 1 Temperature microprobe design. Only the tapering part of the tungsten electrode tip is shown.](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics/015009-2-January-2014-Vol.19(1)/Ersen-Abdo-Sahin-Temperature-elevation-profile-inside-the-rat-brain-induced-by-a-laser-beam/)

![Fig. 2 Experimental setup for optical (a) and temperature (b) measurements in saline.](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics/015009-2-January-2014-Vol.19(1)/Ersen-Abdo-Sahin-Temperature-elevation-profile-inside-the-rat-brain-induced-by-a-laser-beam/)
A 4 × 6 mm cranial opening was made immediately rostral to the bregma on the right side of the central fissure. The connective tissue over the dura was carefully removed, but the dura was left intact. Dehydration of brain tissue was prevented using a pool of mineral oil or saline (three experiments with each) over the cortex filling the cavity to the rim of the skull opening.

After positioning the head perfectly horizontally in the stereotaxic frame, the probe was inserted into the brain (through a small hole in the dura that is off the laser path) at an angle of 45 deg using a three-axis micromanipulator (Fig. 3). A free-air, NIR laser beam (74 mW, 830 nm, DLS-500-830FS-100, StockerYale, Canada) with a Gaussian intensity profile and a circular footprint of 0.4 mm in diameter was aimed at the probe from 13.5 cm above the cortex. The laser activation pattern and data acquisition were controlled through MATLAB™.

The laser intensity profile was verified (at a distance of 13.5 cm) using a commercial photodiode with a very small active area (A0.08 mm², G9842, Hamamatsu, GaAs PIN photodiode) as shown in Fig. 7 on the top. The laser profile was Gaussian with a circular cross section. The radius was taken as the $\sqrt{2\sigma}$ value of the Gaussian intensity curve, which is the horizontal distance where the intensity profile drops down to $1/e$ of its peak in the center. This diameter was equal to that of a flat profile beam with the same peak power intensity and the same total power as the Gaussian profile laser that was used.

Figure 4 shows a sample temperature signal with the sensor in the brain gray matter at a depth of 500 μm. Because the baseline brain temperature was different in each animal due to variations in the homeostasis of the animal and the room temperature, the differential temperature elevation from the baseline ($\Delta T$) was taken as the temperature signal caused by the laser. The trend in the temperature signal did not follow a simple exponential curve but rather had multiple regions [Fig. 4(a)]. There was an initial fast rising region [expanded in Fig. 4(b)] which was
followed by a ramp-like segment for the rest of the laser ON phase. An analytical function that consists of the products of multiple exponential terms would probably give the best fit to the experimental data. Considering the thermal noise level of the signals, we chose a simpler but a more robust model. The initial segment (first 1.5 s) of temperature signals was modeled with a single exponent $\Delta T = T_{inf}*[1 - \exp(-t/\tau)]$ where $t$ is the time) and the last 12 s of the curve with a linear ramp ($\Delta T = a + bt$, where $a$ and $b$ are the constants) as shown by the dash and solid lines in Fig. 4(a). The linear line was extrapolated back to time zero to find the $T_{inf}$ parameter of the exponential formula, leaving the time constant ($\tau$) as the only unknown. The baseline was calculated by averaging the data points within 1.5 s window immediately before the onset of the laser. The temperature elevation at each measurement point was taken at time equals to 5 s from the line fit, subtracted from the baseline, and used for a 2-D temperature plot.

3 Results

The temperature elevation as a function of depth at the center of the laser beam is shown in Fig. 5. The temperature decline by depth was fitted by a single exponent, as shown over the plot. The space constant, which was defined as the depth where the intensity was reduced to 1/e of its maximum, was found as 0.96 mm. If extrapolated, the plot intersects the $y$-axis at 2.6°C, i.e., when the depth equals to zero.

The map of temperature elevations within the 2-D vertical plane is shown in Fig. 6. The maximum temperature values were measured near the surface at the beam center. The temperature profile quickly drops in the horizontal direction nearly following the same trend of the laser intensity profile. In the vertical direction, however, the intensity profile extends a few times the laser diameter. The directional asymmetry is a result of the beam direction entering the brain as well as the large directionality factor of neural tissue ($\theta \approx 0.9$).

3.1 Statistical Analysis

The experiments were divided into two groups; those with saline (three experiments) and those with mineral oil (three experiments) that was used to fill the cranial hole above the dura. The Student $t$ test indicated a significant difference between the two groups in terms of the time constant of the initial rising phase ($p < 0.005$). The group means were 0.45 ± 0.10 and 0.28 ± 0.08 s, respectively (Fig. 7). The temperature measurements taken at $t = 5$ s of the temperature curve were also significantly different between saline and mineral oil when the corresponding points in the 2-D plane were paired ($p < 0.002$, n-way analysis of variance).

3.2 Optical Power Versus Temperature

The laser source had an average power of 14.8 mW (74 mW operated at 20% duty cycle) and a Gaussian intensity profile beam with a circular footprint. The diameter was adjusted to be 0.4 mm at the dura surface using lenses to focus the beam. This diameter was calculated by finding where the light intensity drops to 1/e times the peak in the center, which corresponds to the diameter of a flat intensity profile beam that has the same peak and total power as the Gaussian beam. The average laser power density, i.e., the total power divided by the beam area, was calculated as 14.8 mW/0.1256 mm$^2 = 118$ mW/mm$^2$. At this power level, the maximum temperature measured in the beam center was 2.1°C at 250 µm below the surface. By extrapolating the curve in Fig. 5 back to zero, the temperature at the pia matter surface was estimated to be 2.6°C.

A temperature elevation of 1°C causes profound effects in neural tissue. Thus, a more conservative value of 0.5°C was taken as the maximum allowable temperature in our calculations. The signal-to-noise ratio (SNR) at the thermocouple output was very poor for temperature changes less than a degree.

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Fig. 5 Temperature elevation as a function of depth at the center of the laser beam. The data points represent the mean±standard error of measurements from all six animals. The dashed line is an exponential curve fit (correlation 0.99) with a single exponent.

Fig. 6 Two-dimensional temperature elevation inside the rat brain due to a laser beam of 14.8 mW average power. Measurements were made in steps of 250 µm vertically and 100 µm horizontally and then interpolated. The bell-shaped curve above shows the laser intensity profile aligned with the 2-D temperature plot.
Celsius. In order to improve the SNR, the laser power was intentionally kept high and the results were scaled down to find the optical power level corresponding to 0.5°C temperature elevation (linear interpolation that corresponds to 0.5°C temperature elevation). The optical properties play as much of a role as the thermal parameters of the neural tissue in defining the temperature distribution. Therefore, the temperature increase was a linear function of power within the temperature ranges observed in this study (Fig. 8). The laser power was varied by adjusting the pulse width while keeping the intensity and frequency the same. Vascular mechanisms of temperature regulation did not seem to play a large role within the range studied here, as suggested by the linearity of the plot in Fig. 8. The laser power limit was found as 22.7 mW/mm² or 2.27 W/cm² by linear interpolation that corresponds to 0.5°C temperature elevation (118 mW/mm² × 0.5/2.6 = 22.7 mW/mm²).

4 Discussion

4.1 2-D Temperature Profile

Optical properties play as much of a role as the thermal parameters of the neural tissue in defining the temperature distribution inside the brain. NIR wavelengths can penetrate more easily into neural tissue than the visible light. The dominant form of the interaction with the neural tissue at NIR wavelengths is scattering. Since the gray matter has a lower scattering coefficient than the white matter, NIR light can penetrate deeper into the gray matter. The laser wavelength (830 nm) chosen in this study is one of the several wavelengths in the NIR range at which commercial semiconductor lasers diodes are available. The specific wavelength is not very critical for the general conclusions of this study since the optical properties of tissue do not vary drastically within the NIR region. The temperature elevation of the beam center at a depth of 1150 μm is around 0.9°C in Fig. 6 while the same temperature increase at the surface in the horizontal direction occurs at 750 μm. Thus, the spatial gradient of temperature decrease is steeper in the horizontal direction than the vertical direction. This trend is similar to the optical intensity profile observed in a previous study (their Fig. 4 in Ref. 5), except that the temperature decline is much less steep in all directions because of heat conduction through the medium.

The laser beam profile and diameter are also among the factors that determine the light intensity profile inside the neural tissue as predicted by Monte Carlo simulations. A larger beam size may yield smaller increases in the local temperatures near the surface for the same total power that needs to be delivered. This would make sense only if the same level of optical power can be harvested at the targeted implant with the larger beam size. The Gaussian profile has a central peak of $1/\left(2\pi\sigma^2\right)$ times the total power of the source. A flat profile circular beam can help to reduce this spatial peak effect in the center.

4.2 Sources of Measurement Error

A steady-state was assumed to be reached within the 5 s of laser application. From the time course of the thermocouple signal, however, it was evident that the temperature was still rising slowly at the end of the pulse train. The 5 s time frame was chosen, while recognizing that this is not the steady-state measurement, to be able to finish the 2-D temperature map within a few hours before the animal’s condition began to deteriorate under anesthesia. Other factors influencing the measurements included motion artifacts due to breathing and pulsation of the heart. We also saw the disturbances from the air circulation in the room particularly in the measurements made near the brain surface. The time constant of the probe (60 ms) was significantly shorter than the 5-s measurement period and therefore dismissed as a possible error source. The heat dissipation to the outside through the electrode shank that carried the thermocouple was also assumed to be negligible.

The nearest point of measurement to the cortical surface was 250 μm below the dura and the temperature plot was extrapolated back to the pial surface. The measurements at the pial or dural surface were not reported here because of large variations seen in those measurements. In our observations, the dura matter thickness was also one of the major factors that influenced how much light penetrates into the brain and hence the temperature elevation profile inside. The inter-animal variations in the dura thickness caused large differences in our results as well. Despite these imperfections and disturbances, an adequate level of reproducibility was achieved in the measurements to make a temperature map by averaging the results from multiple animals.
4.3 Maximum Allowable Optical Power

The laser power corresponding to 0.5°C temperature increase (2.27 W/cm²) is almost two orders of magnitude larger than what was needed (75 mW/cm²) to stimulate the ventral horn neurons in the rat cervical cord at a depth of 2.35 mm from the dorsal surface. This suggests that our microstimulators can be implanted deeper in neural tissue without exceeding the temperature limitation on the surface. In a simulation study, the maximum power for the same temperature increase was found as 325 and 250 mW/cm² for the human gray and white matters, respectively. Almost an order of magnitude difference in simulated and measured power levels (2.27 versus 325 mW/cm² for the gray matter) must result from several factors, including the contribution of heat dissipation from blood flow that was not considered in the simulations. Another source of discrepancy is the medium filling the cavity above the brain cortex, which was assumed to be air in the simulations and was either saline or mineral oil in the experiments of this study. The thermal conductivity and the specific heat capacity for air and saline are substantially different (0.03 versus 0.45 to 0.6 Wm⁻¹ K⁻¹, and 1005 versus 3650 Jkg⁻¹ K⁻¹, respectively) and air would dissipate much less heat. The finding that the time constants measured with saline and mineral oil as the filling material are statistically different suggests that the heat dissipation through the skull opening was an important contributor to the measurements. The thermal conductivity and specific heat capacity of the skull are closer than air to that of saline (1.30 Wm⁻¹ K⁻¹ and 1606 Jkg⁻¹ K⁻¹, respectively) and thus the experimental conditions of this study are expected to better mimic the chronic implant conditions where the skull opening would presumably be a tiny hole for the passage of an optical fiber for delivery of NIR light.

On the optical side, the reported values of the scattering coefficient for rat and human gray and white matters vary in a large range; a parameter that would determine the penetration depth and hence the region of maximal temperature effect. We experimentally found the scattering coefficient to be around 108 cm⁻¹ in the rat brain gray matter. The above discussion exemplifies how many different factors account for the thermal effect of a light beam penetrating into tissue. In general, the experimental results may be considered more reliable, if the conditions can closely mimic the targeted application, while being cognizant of the shortcomings of the experimental setup. For instance, anesthesia may reduce the blood flow and suppress the vascular regulation mechanisms of temperature control. If anesthesia indeed has a significant influence, it would push the estimated maximum allowable optical power toward more conservative values.

4.4 Future Work

A computer bioheat model can provide insights as to which parameters the results are most sensitive to, including the properties that cannot be easily modified experimentally, and thereby increase the level of confidence in these experimental results. A computer simulation can also perform a transient time analysis that would allow one to study the heat propagation through the volume in time.

Finally, the degree to which these results can translate to the other parts of the CNS, such as the spinal cord, and to other species is arguable since optical parameters are significantly different between the white and gray matters and between species. These concerns warrant further experimentation for the extension of these results to other neural tissues.

5 Conclusions

The thermal effects of NIR laser exposure in the rat brain are reported here in anesthetized preparations. The noteworthy outcome is that the significant levels of optical energy (2.27 mW/mm²) can be injected into the brain tissue before the thermal effect reaches a critical level that may influence the homeostasis of the local neurons. As a comparison, this optical power is about two orders of magnitude higher than the energy levels required to wirelessly activate neural tissue via photodiode-based floating devices implanted into the rat cervical spinal cord. Nonetheless, the thermal effects should be tested for each specific application since the temperature map depends on many parameters such as the thermal properties of the tissue and the surrounding structures, the optical properties of the CNS region, and the laser beam size and shape.

Acknowledgments

This study was funded by the National Institute of Health Grant (NIBIB, R01 EB009100). We thank David S. Freedman for commenting on this paper.

References


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