LASER-tissue interactions

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Abstract As new laser devices continue to emerge, it becomes increasingly important for the clinical dermatologist to understand the basic principles behind their operation. A fundamental understanding of how lasers interact with tissue will enable the physician to choose the most appropriate laser for a given clinical situation. Although the physical laws guiding laser design are vastly complex, the fundamental principles of laser-tissue interaction can be summarized as they are applicable to the clinician.

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The role of lasers in dermatology has increased dramatically over the past two decades. A fundamental understanding of laser-tissue interactions is vital for the proper and appropriate use in clinical practice. Several excellent review articles have been published addressing principles of laser physics.1-7 This review will focus on concepts with particular relevance to the clinician.

Properties of laser light

The word laser is an acronym for light amplification by stimulated emission of radiation. In the process of spontaneous emission, a photon will be emitted spontaneously from an excited atom after a given length of time. A laser uses a power source, a lasing medium, and a chamber to stimulate the emission of photons. In stimulated emission, an external power source creates excitation of the atoms in the lasing medium. As unstable atoms release their photons, these photons collide with other excited atoms in the lasing medium and trigger a cascade of reaction resulting in numerous photons released at the same moment of identical wavelength, energy, and phase. As long as the population inversion to an excited state continues, the laser light continues to be amplified.

Laser light has several unique characteristics that differentiate it from other light sources. These key properties (monochromaticity, coherence, collimation, and high power) are the basis for the therapeutic applications of laser energy.

Non-laser light sources, such as intense pulsed light devices, emit light of many wavelengths modulated by a cutoff filter. Laser light, on the other hand, is monochromatic, meaning that all of the light emitted by a laser is of a single, discrete wavelength determined by the lasing medium (see Table 1). The monochromatic nature of laser light is a critical property for the application of laser technology in clinical practice because cutaneous chromophores selectively absorb different wavelengths. The specific wavelength of laser light also affects the distance it can penetrate into tissue. In general, the depth of penetration of laser light increases with increasing wavelength within the
collimated beam by reflecting the light in a chamber between two mirrors that allows the exit of parallel waves only. Because the waves of light run parallel to each other, the tendency toward divergence is low. Owing to this organized pattern of light, a laser beam can be propagated across long distances via optical fibers without loss of light by spreading. In the clinical setting, the beam of light is directed at the clinical target using optical fibers or an articulated arm with a handpiece. The parallel light beam is then focused down to a diffraction-limited spot, or smallest possible spot, through a lens.  

The amplification process within the laser cavity produces a high power density. Power density is a function of energy, power, fluence, and irradiance (see Table 2). Energy and power both quantify the amount of light emitted from a laser. Energy, which is measured in joules, represents work. Power is the rate at which energy is expended (joules per second). Fluence refers to the energy density of a laser beam measured in joules per square centimeter. Irradiance, the power density of a laser beam, is the power of the laser beam divided by the area of the laser beam (spot size) and is expressed as watts per square centimeter. Fluence and irradiance are directly proportional to each other (fluence = irradiance × exposure time). By manipulating the fluence, irradiance, and exposure time, one can tailor the use of laser for specific clinical uses.

### Laser-tissue interactions

Light can interact with tissue in four key ways: transmission, reflection, scattering, and absorption. Transmission refers to the passage of light through a tissue without having any effect on that tissue or on the properties of the light. Reflection refers to the repelling of light off the surface of the tissue without an entry into the tissue. Approximately 4% to 7% of light is reflected off skin. The amount of light reflected increases with increasing angle of incidence with the least reflection occurring when the laser beam directed perpendicular to the tissue. If enough reflection occurs to create a high-intensity beam, damage to unintended targets could occur. For example, an errant 10,600-nm beam can cause damage to the cornea because of its water content whereas a reflected 595-nm pulsed dye laser beam can damage the retina because of its pigment content. Protective eye wear guards against reflected and direct exposure to laser light.

Scattering of light occurs after light has entered the tissue. Scattering is due to the heterogeneous structure of tissue, with variations in particle size and the index of refraction between different parts of the tissue determining the amount of scatter. Scattering spreads out the beam of light within the tissue, resulting in radiation of larger area than anticipated. Scattering also limits the depth of penetration because it can occur backward as well as forward. In skin, most scattered light is due to interaction with dermal collagen. In general, the amount of scattering of laser light is inversely proportional to the wavelength of the laser. Longer wavelengths thus penetrate tissue more deeply. An exception to this rule is laser light beyond the mid-infrared region of the electromagnetic spectrum. Laser light with wavelengths above 1300 nm only penetrate superficially owing to the high absorption coefficient of tissue water.

Laser light absorption by specific tissue targets is the fundamental goal of clinical lasers. According to the Grothus-Draper law, light must be absorbed by tissue to produce an effect in that tissue. The absorption of the photons of laser light is responsible for its effects on the tissue. The components of the tissue that absorb the photons

### Table 1: Commonly used dermatologic lasers

<table>
<thead>
<tr>
<th>Wavelength (nm)</th>
<th>Laser</th>
<th>Chromophore</th>
</tr>
</thead>
<tbody>
<tr>
<td>308 (UV-B)</td>
<td>Excimer</td>
<td>DNA/RNA</td>
</tr>
<tr>
<td>323 (green)</td>
<td>KTP</td>
<td>Hemoglobin</td>
</tr>
<tr>
<td>585-600 (yellow)</td>
<td>Q-switched Nd:YAG</td>
<td>Tattoo ink (red)</td>
</tr>
<tr>
<td>694 (red)</td>
<td>Pulsed dye</td>
<td>Hematoporphyrins</td>
</tr>
<tr>
<td>755 (infrared)</td>
<td>Q-switched ruby</td>
<td>Tattoo ink (black, blue)</td>
</tr>
<tr>
<td>810 (infrared)</td>
<td>Long-pulsed ruby</td>
<td>Melanin</td>
</tr>
<tr>
<td>1064 (infrared)</td>
<td>Q-switched Nd:YAG</td>
<td>Tattoo ink (blue, black, green)</td>
</tr>
<tr>
<td>1320 (infrared)</td>
<td>Long pulsed Nd:YAG</td>
<td>Water</td>
</tr>
<tr>
<td>1450 (infrared)</td>
<td>Diode</td>
<td>Water</td>
</tr>
<tr>
<td>1540 (infrared)</td>
<td>Er:glass</td>
<td>Water</td>
</tr>
<tr>
<td>2940 (infrared)</td>
<td>Er:YAG</td>
<td>Water</td>
</tr>
<tr>
<td>10,600 (infrared)</td>
<td>carbon dioxide</td>
<td>Water</td>
</tr>
</tbody>
</table>

### Table 2: Measurements of laser output

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spot size</td>
<td>= laser beam cross-sectional area</td>
</tr>
<tr>
<td>Fluence</td>
<td>= watts × seconds/cm² = joules/cm²</td>
</tr>
<tr>
<td>Irradiance</td>
<td>= laser output × pulse duration / spot size</td>
</tr>
<tr>
<td></td>
<td>= Watts/cm²</td>
</tr>
</tbody>
</table>
preferentially depend on wavelength. These light-absorbing tissue components are known as chromophores. Frequently targeted chromophores in the skin include melanin, hemoglobin, and water, as well as exogenous tattoo inks. Absorption of energy by a chromophore results in conversion of that energy into thermal energy.

Selective photothermolysis

The theory of selective photothermolysis refers to laser energy absorption by a target chromophore without significant thermal damage to surrounding tissue. To achieve selective photothermolysis, the laser must produce a beam of light with a wavelength preferentially absorbed by the chromophore in the lesion. Equally important, the pulse duration of the laser beam must be shorter than the thermal relaxation time of the chromophore to prevent the spread of thermal energy beyond the targeted chromophore. The thermal relaxation time is defined as the time needed for the chromophore to cool to half of its peak temperature after laser irradiation, which is proportional to the square of the size of the chromophore.

In general, smaller objects cool faster than larger ones. For example, 0.5- to 1.0-μm melanosomes have a thermal relaxation time of approximately 1 microsecond whereas 10- to 100-μm capillaries have a thermal relaxation time of approximately 1 millisecond. If the pulse width is greater than the thermal relaxation time, nonspecific thermal damage occurs because of heat diffusion. Finally, the energy delivered to the site (fluence) must be high enough to destroy the chromophore within the pulse duration. Based on the theory of selective photothermolysis, the wavelength, pulse duration, and fluence of a laser can be tailored to result in selective damage to the lesions without nonspecific thermal damage to the surrounding tissues.

Lasers emitting visible light were designed on the basis of selective photothermolysis. These lasers include the pulsed dye, Q-switched ruby, Nd:YAG, and alexandrite lasers. The wavelength of each laser is preferentially absorbed by certain chromophores (see Table 1). Primary cutaneous chromophores targeted by visible light include melanin and oxyhemoglobin. Oxyhemoglobin has three main absorption peaks (418, 542, and 577 nm), whereas melanin has a broad overlapping absorption band ranging across ultraviolet, visible, and near infrared spectrum with decreasing absorption as the wavelength increases. Water absorbs mainly infrared laser energy. Depth of penetration affects the ability to treat certain chromophores at certain depths with certain wavelengths. For example, the dominant absorption peak for hemoglobin is at 420 nm, but this wavelength only penetrates to the dermal-epidermal junction (100 μm), which would limit use of a laser of this wavelength for cutaneous vascular lesions. The smaller absorption peak of oxyhemoglobin at 577 nm is more useful clinically because of the deeper penetration achieved at this wavelength.

Laser parameters and tissue interactions

Beam characteristics

An important feature of the light produced by a laser is how the intensity is distributed across the beam diameter. Most cutaneous lasers produce a beam with a Gaussian profile in which the intensity peaks at the center of the beam and attenuates at the periphery. Clinically, this results in the necessity of treating tissue with some overlap of the laser beam to deliver energy to tissue in a more uniform manner. Other modes of operation for lasers result in a doughnut-shaped or target-shaped distribution of energy delivery, with the intensity of the laser light at the edge of the beam being greater than in the center or fluctuating across the diameter of the beam. These modes tend to approach a more constant intensity profile, thus overlapping needs to be kept to a minimum to prevent over heating of the tissue at the periphery of the laser beam. These modes are somewhat limited in clinical use because of having a much larger minimum spot size than the fundamental mode.

Spot size

The spot size of a laser is equivalent to the laser beam cross section. The spot size directly affects the fluence and the irradiance of a laser beam, as mentioned above. Fluence and irradiance are inversely proportional to the square of the radius of the spot size. As such, halving the spot size increases the energy density or power density by a factor of 4. Spot size is also important clinically owing to laser scattering in the skin. A small spot size allows more scattering both backwards and sideways than a larger spot size. This results in a more rapid reduction of energy fluence in tissue than with a larger spot size. A large spot size of 7 to 10 mm is needed for maximal penetration of laser light to mid-dermal or deeper targets. Increasing depth of penetration levels off with spot sizes of 10 to 12 mm.

Pulse duration

Laser light can be delivered in a continuous wave or a pulsed wave. Continuous wave lasers emit a constant beam of light that may result in nonselective tissue injury. Pulsed delivery of laser light allows for more selective tissue damage. The duration of time of exposure to a laser beam determines the rate at which the laser energy is delivered. Pulse duration ranges from very short (nanoseconds) as in the case of Q-switched lasers to long (milliseconds) as in the case of hair removal lasers. The pulse duration of any given laser will be determined by the thermal relaxation time of the intended target (Table 3). The thermal relaxation time is generally proportional to the size of the target structure. For example, Q-switched lasers target very small structures with short thermal relaxation times such as melanosomes and tattoo ink in the case of the Q-switched Nd:YAG laser. Increasing the pulse duration of
Surface cooling

When the chromophore contained in the target for ablation, such as melanin in a hair follicle, is located deeper in the skin than unintended targets such as epidermal melanin, the selectivity of the laser for its intended target can also be improved by surface cooling. There are three principal delivery methods for surface cooling: precooling, parallel cooling, and postcooling. Precooling consists of cooling the temperature of the epidermis immediately before the laser pulse, usually by cryogen spray. In contrast, parallel cooling occurs at the same time as the laser pulse usually by use of a water-cooled sapphire tip. Parallel cooling is preferable for laser devices with longer pulse duration. Postcooling with ice helps reduce pain and edema but likely has little effect on the distribution of laser-induced thermal injury.

Photothermal ablation and nonablative dermal remodeling

Infrared lasers such as the carbon dioxide (10,600 nm) and erbium:YAG (2940 nm) lasers target water as a chromophore. Both ablative and nonablative infrared lasers heat tissue by using water as a target chromophore. The change in tissue temperature and the rate of heating determines the tissue response. Cellular injury with subsequent inflammation and repair occur after tissue temperature increases by only 5°C to 10°C. Deactivation of cellular enzymes occurs at temperatures of 40°C to 45°C, with reversible damage that becomes irreversible with sustained exposure. Temperatures above 60°C lead to denaturation of most proteins, whereas further increases above 70°C lead to denaturation of DNA. Vaporization of tissue water with cell shrinkage, hyperchromasia, membrane rupture, protein denaturation, and collagen hyalination occurs at temperatures of 60°C to 140°C. This results in coagulation, with blanching macroscopically. With sustained exposure to these temperatures, explosive vaporization occurs. At temperatures of 300°C to 1000°C, tissue ablation occurs with smoke generation.

Resurfacing lasers are high-energy pulsed lasers that generate photothermal ablation that occurs with rapid heating when tissue absorbs enough laser energy to cause the water in the tissue to vaporize. Damage also occurs in the tissue surrounding the ablated area due to thermal diffusion and scatter (zone of thermal damage). The CO₂ laser has a high absorption coefficient for tissue water that allows minimal residual thermal damage if the power density is significant enough to cause tissue vaporization that outpaces the speed of thermal diffusion; the energy delivered is less than the thermal relaxation time; or the fluence is only enough to result in only a thin layer of collagen being denatured. With each pass of the CO₂ laser, a predictable thickness of tissue is vaporized. The depth of ablation of the CO₂ laser is approximately 30 μm per pass; however, the underlying zone of thermal damage from heat diffusion is typically 300 μm or more because the thermal relaxation time of the 30-μm tissue layer heated by the laser is less than 1 μs.

Collagen shrinkage and remodeling require an additional one to two passes after the initial pass results in epidermal ablation. The total depth of dermal injury consists of the thermally damaged base in addition to the thin layer of collagen being denatured. The depth of ablation can be accurately controlled by adjusting the power, spot size, and number of passes used. Residual thermal damage gradually increases with each pass of the CO₂ laser while the ablation depth of each pass below the papillary dermis decreases due to decreasing water content. While increasing depth of thermal injury increases clinical response up to a point, overheating due to an excess number of passes or rapid overlapping of pulses can result in scarring. Overlap of the CO₂ laser spot should be 10% or less. The short-pulsed erbium:YAG laser has a higher absorption coefficient than that of the CO₂ laser, resulting in more efficient absorption of erbium laser energy by water-containing tissue. Given the higher coefficient of absorption of the erbium:YAG laser by tissue water, each pass produces a tissue ablation to a depth of 2 to 5 μm per pass with an underlying zone of thermal necrosis of 10 to 15 μm. Pulse stacking does not seem to increase the depth of thermal damage in contrast to the CO₂ laser. Less depth of thermal injury results in shorter healing time with erbium:YAG laser treatment but less skin-tightening effect due to collagen shrinkage than the CO₂ laser. Modulated erbium:YAG lasers with extended pulse durations up to 500 microseconds result in larger zones of thermal necrosis compared with traditional short-pulsed laser systems. These larger zones of collateral damage result in enhanced tissue tightening effects approaching the CO₂ laser.

Nonablative laser resurfacing results in collagen remodeling by inducing selective heating of tissue water in the dermis without injury to the epidermis. The fundamental difference compared with ablative lasers is the slower rate of delivery of laser energy and the use of surface cooling. With the use of cooling devices, the energy of the

| Table 3 | Thermal relaxation times of common laser targets |
|-----------------------------------------------|
| Target                                    | Size (μm) | Thermal relaxation time |
| Tattoo ink particle                        | 1         | 10 ns                  |
| Melanosome                                | 1         | 1 μs                   |
| Erythrocyte                               | 7         | 20 μs                  |
| Epidermis                                 | 50        | 1 ms                   |
| Blood vessel                              | 50        | 1 ms                   |
| Ectatic blood vessel                      | 100       | 15 ms                  |
| Hair follicle                             | 200       | 20-100 ms              |
laser beam largely bypasses the epidermis and is absorbed deeper within the papillary dermis. The clinical importance of this treatment approach is that it offers a modality with significantly less morbidity and downtime than with ablative laser therapy. Although heating of dermal collagen can be achieved by a variety of lasers with different wavelengths, the most effective nonablative lasers are infrared lasers (1000-1500 nm) that target water in the dermis. Infrared laser therapy is achieved by a variety of lasers with different wavelengths, including monochromicity, coherence, collimation, and high power, are the basis for the therapeutic applications of laser energy. The energy, power, fluence, and irradiance are specific parameters of laser light that can be adjusted for specific clinical uses. The theory of selective photothermolysis guides the clinical application of lasers visible spectrum. High-energy pulsed ablative lasers in the infrared range result in the vaporization of water-containing tissues. Nonablative laser therapy causes heating of dermal collagen while sparing the epidermis. Cooling devices optimize delivery of laser energy to deeper targets. Further refinements of laser-tissue interactions will result in the enhancement of existing technology as well as novel clinical applications in the future.

Conclusions

Given the rapid evolution of laser therapy in dermatology, a clear understanding of the principles of laser light physics and tissue interactions is vital for proper management of patients. The unique properties of laser light, including monochromicity, coherence, collimation, and high power, are the basis for the therapeutic applications of laser energy. The energy, power, fluence, and irradiance are specific parameters of laser light that can be adjusted for specific clinical uses. The theory of selective photothermolysis guides the clinical application of lasers visible spectrum. High-energy pulsed ablative lasers in the infrared range result in the vaporization of water-containing tissues. Nonablative laser therapy causes heating of dermal collagen while sparing the epidermis. Cooling devices optimize delivery of laser energy to deeper targets. Further refinements of laser-tissue interactions will result in the enhancement of existing technology as well as novel clinical applications in the future.

References