High precision subsurface photodisruption in human sclera

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1 Introduction

Photodisruption occurs when laser power intensity is large enough to induce optical breakdown in a material: the laser pulse transforms the material into plasma. The strong optical intensities required for photodisruption can be achieved at the focus of a high peak power laser beam. Breakdown occurs only where the intensity is larger than the threshold level. Below this threshold level, the laser pulse passes through the material without causing damage. Thus, the location of optical breakdown can be controlled to occur only at the focus of the beam where the intensity exceeds the threshold level. If the laser is focused beneath the surface of a material, subsurface breakdown occurs only at the focus. No damage occurs in the material that the beam was focused through. Complete subsurface incisions of any shape are possible by scanning the focal spot beneath the surface.

Photodisruption is an intensity dependent process. Therefore, pulse energy, which is associated with collateral damage processes, can be decreased by decreasing the pulse duration.

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Abstract. Background and objectives: Femtosecond pulses can generate high precision subsurface photodisruption in transparent tissues, such as the cornea. We used femtosecond laser technology to demonstrate early proof of concept for high precision subsurface photodisruption in the translucent sclera. This technique may ultimately enable novel surgical procedures for the treatment of glaucoma and/or presbyopia. Study design/materials and methods: Microjoule femtosecond pulses from two different sources, 1060 and 775 nm, were used to make subsurface incisions in human sclera in vitro. Scleral tissue was dehydrated to improve translucency at these wavelengths. The beam was focused to a 1.5 (775 nm) or 5 μm spot size (1060 nm) and scanned below the tissue surface at various depths to produce four incision patterns. Results: Photodisruption on the back-surface of the sclera was achieved without damage to overlaying tissue. Several types of intrascleral incisions were made, including transcleral channels and grooves for scleral implants. Conclusions: High precision, subsurface scleral photodisruption can be achieved in vitro for a variety of intrascleral incisions. Further studies are required to determine if this technique is applicable in vivo for actual surgical applications. © 2002 Society of Photo-Optical Instrumentation Engineers. [DOI: 10.1117/1.1482381]

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If femtosecond pulses are used, the pulse energy can be decreased (microjoule level or lower) so that most of the pulse passes through the material with only a small fraction being absorbed. When photodisruption occurs near the threshold energy level, most of the absorbed energy is used to create the plasma with only a small amount of the absorbed energy dissipated by other mechanisms, such as shock wave formation, bubble oscillation, and thermal diffusion. Femtosecond pulses, therefore, allow precise subsurface disruption with almost no collateral damage. The diameter of the disrupted volume can be on the order of microns, allowing computer controlled, scanning delivery systems to create almost any subsurface incisional patterns.

Photodisruption using femtosecond pulses has been used to create high precision subsurface incisions within the transparent cornea, without damaging overlying or surrounding tissues. In contrast to corneal tissue, the sclera scatters visible light, spreading the pulse in both space and time, making it difficult to maintain the tight focus and short pulse duration needed for well-confined photodisruption. The ability to focus
light through sclera is limited, especially at wavelengths less than 800 nm. At longer wavelengths, where scattering is reduced, water in the tissue absorbs laser energy, thereby making the intensity levels necessary for photodisruption harder to achieve. Dehydrating agents, such as Hypaque 76, have been shown to increase scleral transparency by reducing scattering, without damaging the tissue if used for short periods of time. The mechanism of action of this agent can be understood based on the fiber structure of the sclera.

Human sclera is composed of collagen fibrils embedded in a host matrix material. Sclera appears white because of the size (~30–300 nm diameter), packing (285 nm average period), and the index mismatch between the fibers (n = 1.474) and the host matrix (n = 1.345). Bragg scattering for sclera occurs at wavelengths shorter than 800 nm, assuming an index of 1.4 for sclera, thus making sclera an effective diffuser for visible light. Sclera can be made clear by decreasing the index mismatch between the fibers and the host material matrix through dehydration. The index of the background material is raised as the water (n = 1.33) is removed. Several dehydrating agents have been examined previously, including Trazograph and glucose.

Figure 1 shows an example of transparency induced in in vitro human sclera. A tissue that is normally white can be made transparent enough to read through upon dehydration. We hypothesized that if such dehydrating agents cause significant reduction in scattering, then femtosecond pulses of normally scattered wavelengths could be focused strongly beneath the surface to permit subsurface photodisruption, enabling development of a number of potential scleral procedures.

In the first section of this paper, qualitative measurements of laser beam spot sizes through sclera are described to show the need for inducing transparency if subsurface photodisruption is to be achieved at 775 and 1060 nm. In the second part, some examples of surface incisions are shown in human sclera. These incisions are completely subsurface and a variety of shapes with almost no collateral damage.

2 Focusing Through Hydrated and Dehydrated Sclera

Tissue from human globes, not suitable for transplantation, was obtained from the Midwest Eye Bank and Transplantation Center one to four weeks after harvest. Full thickness scleral pieces (approximately 1 cm square and approximately 0.5 mm thick) were cut from the limbal region. Samples were preserved for up to several months in 95% ethanol solution. Prior to experimentation, samples were soaked in physiological saline solution for several hours. Some samples were then dehydrated in Hypaque 76 until becoming maximally transparent, usually within 15 min.

The transmission spectrum of the tissue was measured in a spectrophotometer. The spectra obtained were forward scattered spectra—the light scattered in a small cone angle around the incident beam—as no integrating spheres were used. The forward scattered light is believed to be an indicator of the light that is focusable through the sclera. As shown in Figure 2, transmission of hydrated sclera is never greater than 10%, peaking at 1700 nm. Upon dehydration, the transmission greatly increases across the entire spectrum, especially for wavelengths in the near infrared. The transmission is mainly controlled by scattering in the visible spectrum be-
cause of absence of absorbing chromophores in this region in sclera. Attenuation occurs at longer wavelengths due to the absorption spectrum of water, most notably seen by the strong peaks at 1450 and 1900 nm. Transmission at 775 and 1060 nm is above 60%. The difference in transmission is not expected to affect the results of photodisruption since photodisruption depends on the intensity of the pulse and not the linear absorption.

Since scattering is the dominant mechanism controlling transmission, transmission spectra of sclera are not good indicators of the focussability of a laser beam beneath the surface. The next experiment was performed to qualitatively describe the size of the focal spot on the backsurface of sclera. This study was performed only qualitatively since: (1) inhomogeneities in the sclera change the spot size from location to location, (2) the technology of the camera used is not suited to take exact measurements due to dynamic nonuniformity and gain curves, and (3) the exact spot size obtained in photodisruption will be altered by nonlinear processes, such as self-focusing. However, the spot images obtained are enough to explain the observed physical processes.

The spot size measurement setup is shown in Figure 3. Since different lasers were used, the profiles were approximately made to be the same by passing the beam through a single mode step index fiber to obtain a \( LP_{01} \) (near Gaussian) spatial mode. The intensity of the beam was attenuated to be low enough to prevent nonlinearities in the fiber. The beam was then focused using a 0.5 numerical aperture (NA) aspheric lens (Thorlabs C240TM) to the backsurface of a piece of sclera sandwiched between a microscope slide and cover slip. The backsurface of the sclera was imaged using a 20\( \times \) or 40\( \times \) microscope objective.

Before the tissue was examined, the size of the unscattered beam was measured by using an equivalent slab of water (0.5 mm) between the glass slides used to mount the tissue. Figure 4(a) shows the minimum spots obtained for 775 and 1060 nm after focusing through a full thickness (0.5 mm) piece of tissue. The square in the top right hand corner is 10 \( \mu \)m on edge. In the bottom left corner is the beam after passing through an equivalent thickness of saline. The unscattered beam is smaller than 10 \( \mu \)m in diameter. The large size is due to underfilling the focusing lens with the incident beam and ab-

![Fig. 3 Spot size measurement setup. A clean spatial mode created by a single mode optical fiber was focused through the sclera mounted between a microscope slide and a cover slip. The backsurface of the tissue was imaged to a vidicon with a 20\( \times \) or 40\( \times \) microscope objective.](image)

![Fig. 4 Minimum spot sizes of a focused beam through human sclera. The bottom left corner shows the smallest spot obtained by focusing through an equivalent thickness 0.5 mm slab of water. A square 10 \( \mu \)m on edge is shown in the top right corner. The bar beneath each picture is the color map: left is not light (black), right is maximum light intensity (white). (a) In normal hydrated tissue, focusing at 775 and 1060 nm is not possible because of scattering. Image taken with 20\( \times \) magnification. (b) When water is removed from the tissue using a dehydrating agent, the tissue becomes clear as the index is matched between the collagen fibers and host matrix. The smallest focusable spot through sclera is about the same size as the beam focused through water. Image taken with 40\( \times \) magnification.](image)
3 Subsurface Photodisruption in Sclera Using a Dehydrating Agent: Methods

The scleral tissue sample (typically 0.5 mm thick) was placed between two microscope slides (each 1 mm thick) and mounted on a computer controlled translation stage in order to move the tissue with respect to the laser focus as shown in Figure 5. All incisions were made by single-shot laser exposure: the sample was translated fast enough so that each piece of treated tissue was hit only once by the laser. Typical translation speeds were 5 mm/s, which yields 5 μm laser spot spacing for a 1 kHz pulsed laser system. Different surgical patterns were formed by scanning the spot in a raster pattern to achieve the desired incision. The beam’s initial position was determined using the visible light produced by photodisruption of glass.

Femtosecond lasers were used to demonstrate the possibility of achieving subsurface photodisruption. Two different lasers were used because of the availability issues. (This paper is not meant to be a comparison between the two lasers.) The first laser had typical operating parameters of 5 μJ, 1 kHz, 500 fs, and 1.06 μm, and the second at 5 μJ, 1 kHz, 150 fs, and 775 nm. For both lasers, the beam was focused through a 0.5 NA aspheric lens (ThorLabs C240TM) corrected for wavefront flatness. The spot size in air was measured to be 5 μm full width half maximum (FWHM) at the focus of the lens for the 1060 nm light and 1.5 μm for the 775 nm light. No compensation was made for potential aberrations produced by the microscope slides and dehydrated sclera. Tissue inhomogeneities inherent in the tissue and nonlinear self-focusing of the beam should permit controlled backsurface photodisruption.

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also make it difficult to determine the spot size in the tissue. The goal here was simply to produce a surgical effect in a normally scattering tissue.

Treated scleral samples were fixed in 2% glutaraldehyde and prepared for scanning electron microscopy (SEM) immediately after laser exposure. Since this process includes dehydration, distortion secondary to shrinkage and gross movement of structures is possible.  

4 Subsurface Photodisruption Results
In all of the examples that follow, three pictures are shown in the figures. Part (a) shows the electron micrograph of the

Fig. 7 Femtosecond laser created transcleral channel. (a) A channel with a minimum width was demonstrated. The channel width is estimated at 10 \( \mu \)m or less from this cross section. A plane was scanned in order to find the channel upon dissection. (Laser parameters: 4.2 \( \mu \)J, 150 fs, 775 nm, 1 kHz, 2 \( \mu \)m FWHM spot, 1.5 \( \mu \)m spot separation.) The minimum width channel was made through human sclera. (b) Potential application of creating a drainage passageway. (c) Scan pattern.

Fig. 8 Cross section of channel created in human sclera. A channel the width of the beam was created in the sclera using 1.0 \( \mu \)m spot separation in the vertical direction (perpendicular to the tissue surface). The channel penetrates the full scleral thickness as verified in top and bottom SEM examination. (a) The channels are not seen to penetrate the entire surface here because the sectioning cut intersects only a portion of the channel. (Laser parameters: 5 \( \mu \)J, 500 fs, 1060 nm, 1 kHz, 5 \( \mu \)m FWHM spot, 0.1 \( \mu \)m vertical spot separation.) (b) Such channels may be useful for altering the hydraulic conductivity of sclera for increased fluid outflow. (c) Raster pattern for the laser focus.
treated tissue. Part (b) shows the treatment geometry in a full globe. Part (c) shows the scanning pattern of the laser spot.

4.1 Partial Thickness Channel Creation

Figure 6 shows the backsurface ablation created by focusing the femtosecond laser pulses through the tissue. Note that no damage occurred above (superficial to) the ablated region, even though the laser beam was focused through this portion of the tissue. There is no technical limit to create deeper channels, since the beam is moved from the inside of the tissue outwards. This incision pattern could be used to perform transscleral procedure analogous to deep sclerectomy, where a block of inner surface sclera is removed with minimal disruption to overlying layers.

4.2 Full Thickness Channel Creation

Figure 7 demonstrates the narrowest transscleral channel possible using our experimental setup. The beam was focused through the tissue to the inner surface and then advanced outward (towards the focusing lens). A plane was cut by the laser in order to find the incision after sectioning. The channel has an estimated width of 10 μm, which is probably even narrower but is obscured by the electron micrograph processing. This incision pattern could be used to perform transscleral procedure analogous to deep sclerectomy, where a block of inner surface sclera is removed with minimal disruption to overlying layers.

4.3 Creation of Tissue Pores

A grid of full-thickness pores were created using the pattern shown in Figure 8(b). This treatment may be useful in changing the bulk properties of the tissue, including the tissue’s hydraulic conductivity. Two representative channels were identified after sectioning, each approximately 10 μm wide. Penetration of the laser beam to the superficial and deep surfaces was indicated by a regular array of photodisrupted tissue at the channel openings at both surfaces (not shown).

4.4 Scleral Pocket Creation

Intrascleral pocket incisions were successfully created [Figure 9(c)], using the pattern schematized in Figure 9(a). A 5 μm spot separation was used to first create the pocket, which was then opened manually with a forceps. A metal pin was placed into the pocket to demonstrate the complete opening, while the surface quality of the separated tissue plane at both the superficial and deep portions of the pocket are shown in Figure 9(d). This procedure may be useful for creating scleral pockets for implants to treat presbyopia.

4.5 Subsurface Volume Pattern Photodisruption

To evaluate the completely enclosed photodisruption pattern, a rectangular prism geometry was scanned beneath the surface as shown in Figure 10(b) using the 775 nm laser. The spot separation in this pattern was 3 μm in every dimension. The rectangular prism was 100 μm wide, 100 μm deep, and 3 mm long. Subsequently, the sample was cut perpendicular to the tissue surface to expose a 100×100 μm square cross section. The incision as shown in Figure 10(a) shows two significant
features. First, the tissue has not been altered above and below the incision. Second, there is debris in the treated volume that maintains some lamellar structure of the sclera. Irregularities in the shape of the photodisrupted area may be due to a variety of reasons, including scattering of the laser light in the tissue, laser beam stability (energy and position), and processing distortion.

In order to compare the quality of incisions between 775 and 1060 nm lasers, the 1060 nm laser was used to cut a similar geometry. Figure 11 shows a 50 × 50 μm cross section of a rectangular prism cut with a 5 μm spot separation using the 1060 nm laser. In this instance, there is no damage above and below the incision, but the debris in the incision has a different structure of that shown in Figure 10(c) using the 775 nm laser. When viewed at higher magnification in Figure 12, a sharp boundary between the photodisruption debris and the adjacent tissue is seen. The debris from the 1060 nm laser ablations is quite disorganized, as compared with the more regular structure seen with the 775 nm ablations shown in Figure 10(c). Adjacent to this area is a 2–10 μm wide region that contains undisrupted, but debris-coated fibers, which were not present in 775 nm ablations.

5 Discussion

We have demonstrated in an in vitro model that human sclera can be made surgically accessible for high precision intrasceral photodisruption if the sclera’s natural light scattering properties are reduced by dehydration. Similar to observations in the cornea, minimal collateral damage was observed in the surrounding structures adjacent to the photodisrupted volume.21 A variety of incisional geometries can be created as long as the pattern is formed in a deep to superficial manner, so that cavitation bubbles produced during photodisruption do...
not interfere with the focusing of the laser beam. Partial and full-thickness transcleral channels, subsurface pockets, and subsurface volumetric photodisruption are examples of possible geometries. In patterns where surface openings are not created (or are inadequate for the volume of debris) the photodisrupted material remains inside the tissue, appearing bright in the electron micrographs. Although the mechanism accounting for the debris coating adjacent collagen fibers is not known, ejection of a portion of the debris into the surrounding tissue seems likely.

A number of additional studies is required to determine if and how ultrashort laser pulses can be used for actual surgical procedures in the sclera. First, in vivo studies, must determine if enough optical clearing occur after subconjunctival injection of Hypaque to allow low energy photodisruption with femtosecond pulses. Second, physiologic effectiveness must be evaluated. For example, in glaucoma procedures, residual tissue debris may block aqueous flow, although hydraulic conductivity may be altered enough in the treated volume to increase fluid outflow. In this case, the photodisrupted debris could help to keep laser created transscleral channels open. Finally, in vivo testing is required to evaluate effects evaluate effects that may not be apparent using this in vitro techniques, such as toxicity from the dehydrating agent or wound healing after laser ablation.22 In addition, similar techniques for subsurface photodisruption may be possible in other tissues, such as the skin, where similar optical clearing effects are possible.23

The differences seen between the 1060 nm 500 fs laser and the 775 nm 150 fs laser are interesting, although many other parameters also differed in the two systems, including wavelength, spot size, pulse duration, and scanning density. More detailed investigations are necessary to characterize the temporal and spatial profile of the pulses applied, since the standard FWHM measurement of the temporal duration may not be adequate to determine the breakdown characteristics of the pulse.24 Regardless, both laser sources provide a method of subsurface photodisruption in human sclera, which have the potential to be used in a clinical procedure.

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Fig. 12 Higher magnification of lesion zone. (a) A boundary between solid debris and debris-coated fibers. (b) Debris seems to be coating the fibers adjacent to the debris region. (c) The debris is disorganized collagen bundle.
References

7. HYPAQUE 76, diatrizoate meglumine 76%, Nycomed Inc., Princeton, NJ.
11. This procedure was provided by the Kellogg Eye Bank for preserving sclera for transplantation.