Monitoring photodynamic therapy with photoacoustic microscopy

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Abstract. We present our work on examining the feasibility of monitoring photodynamic therapy (PDT)-induced vasculature change with acoustic-resolution photoacoustic microscopy (PAM). Verteporfin, an FDA-approved photosensitizer for clinical PDT, was utilized. With a 60-μm-resolution PAM system, we demonstrated the capability of PAM to monitor PDT-induced vasculature variations in a chick chorioallantoic membrane model with topical application and in a rat ear with intravenous injection of the photosensitizer. We also showed oxygen saturation change in target blood vessels due to PDT. Success of the present approach may potentially lead to the application of PAM imaging in evaluating PDT efficacy, guiding treatment, and predicting responders from nonresponders. © 2015 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.20.10.106012]

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1 Introduction

Photodynamic therapy (PDT) is a treatment that takes advantage of toxicity of light-activated photosensitizers to kill targeted diseased tissues.1 It has been successfully applied to a wide range of medical conditions, such as skin conditions and cancer treatment. During a PDT treatment, photosensitizer is activated from ground state to excited state by photons when exposed to a treatment light source; then it decays to a triplet excited state, which interacts with molecular oxygen to create reactive oxygen species including singlet oxygen (¹O₂), which is highly cytotoxic.2 Therefore, there are three interacting components: photosensitizer (PS), light administration, and oxygen involved in PDT. Oxygen level plays an important role, as PDT relies on oxygen in the surrounding environment to generate cytotoxic reactive species. To optimize outcomes in clinical practices, real-time evaluation of PDT treatment is of significant importance and urgently needed.3 Although light dosimetry measurement is important, other factors such as PS distribution, ¹O₂ concentration and light fluence (rate), and biological response monitoring are effective means to evaluate the efficacy of PDT as it is a direct reflection of the treatment outcomes.4

By using PSs with high affinity for vascular endothelial cells, PDT can target vasculature in soft tissue.3 Imaging vasculature is important for these applications. A number of techniques can be used to accomplish this task, for example, magnetic resonance angiography, which is a group of techniques (including dynamic contrast enhancement) to image blood vessel effects based on magnetic resonance imaging.5,6 However, it requires imaging contrast agent, long scanning time, comparatively high cost, and can provide only limited spatial resolution. Positron emission tomography also requires contrast agents to visualize vasculature.5,6 Ultrasound Doppler imaging has been widely applied to vasculature (blood flow) imaging. However, only modest spatial resolution can be provided.10 Recently, optical techniques such as laser Doppler imaging11,12 and laser speckle imaging13 have been proposed to study vasculature. However, these techniques suffer from either limited penetration depth due to highly scattered photons in soft tissue, limited resolution, or difficulty to target specific regions of interest in practice. Optical coherence tomography (OCT) has been applied to PDT.14,15 Jung et al. used time-lapse OCT to image microscale response of in vitro models to PDT.15 Hamdoo et al. used OCT to image tumor cancer to guide PDT in human skin cancer treatment.16 Doppler optical coherence tomography, an extension of OCT, which measures frequency shifts due to moving particles, has been employed to study microvascular tissue response in PDT in animal models.14,15,17,18 Although different methods have been proposed, estimation of oxygen saturation with OCT is still nontrivial.19,20

Photoacoustic (PA) imaging is an imaging modality being intensively studied recently because of its promise for combined high resolution and intrinsic optical contrast, which can reveal important physiological information without the assistance of contrast agents.21 Imaging contrast in PA imaging is primarily provided by optical absorption. As hemoglobin in blood is the dominant absorber in soft tissue to provide optical contrast in the visible wavelength window, PA imaging techniques have been applied to a wide spectrum of topics for both morphological and functional studies related to blood vasculature.22–24 Xiang et al.25 reported using a computed photoacoustic tomography (PAT) imaging system to monitor vascular damage in PDT on a chick chorioallantoic membrane (CAM) tumor model. In their study, a single pulsed laser with a wavelength of 532 nm served as the light source for both PA imaging and PDT. Protoporphyrin IX was used as the PS. They visualized neovascularization in tumor angiogenesis and then verified the capability of PAT for imaging vascular injury induced by PDT. However, only blood vessel size change was studied in their report. Their study was also limited to topical application of PS on...
the CAM model. Mallidi et al.26 used ultrasound-guided PA imaging to map three-dimensional (3-D) atlas of tumor oxygen saturation (SO2) change due in PDT. However, this study is limited to large-scale SO2 maps rather than at single blood vessel level.

In this paper, we report our work on monitoring both morphological and functional information variations due to PDT with acoustic-resolution photoacoustic microscopy (PAM). With the wavelength tunability of our laser system, we are able to image SO2 change of areas of interest during PDT, besides structural variations. We studied effects of PDT in a rat ear model with intravenous (IV) injection of PS from a tail vein, and topical application of PS on a CAM model.

2 Methodology

2.1 Experiment Setup

The experiment setup is shown in Fig. 1. We used a tunable nanosecond pulsed dye laser (ND6000, Continuum, Santa Clara, California) as the light source for PA imaging. The laser has a tunable wavelength range between 420 and 900 nm with proper selected dyes. Pulse-repetition rate is 10 Hz and the output energy can reach up to 260 mJ/pulse (with Rhodamine 6G). Pulse duration is ∼10 ns. For the present study, we used Rhodamine 590 (Exciton, Dayton, Ohio), whose emission range with our pumping source is 552 to 584 nm. We utilized 560 nm for structural imaging. For oxygen saturation measurement, 560 and 576 nm were employed. The laser pulse was coupled into a 600-μm multimode optical fiber. The pulses coming out of the fiber were first collimated. After being focused by a microscope objective lens, it was directed to the subject/target by a custom-made imaging probe, which consists of two right-angle prisms. An ultrasound transducer (V214-BB-RM, Olympus) with a 50 MHz center frequency was coupled with the probe acoustically. An optically transparent acoustic lens was positioned under the probe for acoustic focusing. Ultrasonic gel was applied between the acoustic lens and the sample to couple sound waves. PA signals (A-lines) detected by the transducer were digitized by a high-speed digitizer and then transferred to a PC for further processing. To realize raster scanning during the imaging task, the object was positioned on a three-axis motion stage, which was controlled by a PC through a motion control card.27 In the present paper, 15 min were needed for each rastered image acquisition with a field-of-view size of 3.6 mm × 3.6 mm. Lateral and axial resolutions of our system were quantified to be ∼60 and 30 μm, respectively.

Energy used for imaging was ∼3 mJ/pulse, and the laser fluence on animal skin surface was limited to be below 20 mJ/cm2, set by the ANSI safety standard for the wavelength range used here.

A representative whole rat ear 3-D dataset and the corresponding two-dimensional (2-D) image generated with maximum amplitude projection (MAP) method demonstrated the imaging capability of this system.(Fig. 2) In the remaining sections of the present paper, we use 2-D MAP images for quantitative analysis.

A diode laser with a 455 nm wavelength (Ultralasers, Newmarket, Ontario, Canada) was utilized for PDT. The laser beam was directed to an aperture to adjust the beam size for treatment and, thus, to control the optical fluence on the surface of the subjects.

2.2 Photosensitizer

Verteporfin (129497-78-5, Sigma-Aldrich, St. Louis, Missouri) was utilized as the PS. Verteporfin 3.6 mg was first dissolved in 500 μL dimethyl sulfoxide and then diluted with phosphate buffered saline to various concentrations for the experiments. During preparation, handling, and storage, the solution was protected from light.

2.3 Imaging and Data Analysis

To validate the capability of our system to image vasculature change due to PDT, a preliminary study was conducted with the CAM model. We imaged three eight-week-old chick embryos,
which all received 20 min light illuminations. The laser illumination was conducted on embryo #1 without use of PS as a comparison control. Verteporfin 50 μM was applied topically to the regions of treatment 15 min before treatment on the membrane of chick embryos #2 and #3. PAM imaging was performed before and after treatments for the embryos #1 and #2. Then we conducted a longitudinal study with the CAM of embryo #3, for which sequential images were obtained at every 10-min interval.

We also examined the feasibility of our technique with a rat ear model. Experimental procedures followed the laboratory animal protocol approved by the University of Alberta Animal Use and Care Committee. An animal anesthesia system was utilized while image data were collected. Verteporfin was injected via a tail vein of 60-g-weight Sprague Dawley (Charles River Laboratories, Wilmington, Massachusetts) rats (n = 3 animals) with a dose of 2 mg/kg, as suggested in the literature. Before an experiment, the rats’ ears were processed with hair removal cream for 10 min and then cleaned with warm water. PDT was started 20 min after each of the injection. Light irradiance on the skin surface for the treatment was ~80 mW/cm². During the experiment, the treatment was temporally stopped every 5 min for PAM imaging. To quantify blood vessel structural change due to PDT, we compared the diameter of one selected major blood vessel in the PAM images induced by the treatment for each animal. Only blood vessels in the treatment areas with diameter around two to three times larger than the imaging lateral resolution were selected for analysis. Full width at half maximum of the identical feature was used to measure the diameter in longitudinally acquired images for comparison. For a single blood vessel, measurements were taken at multiple locations along the axial direction of the vessel, and data were presented as mean ± standard error.

Oxygen saturation of target blood vessels in the treatment area was measured at multiple time spots longitudinally during PDT in rats’ ears (n = 3) to demonstrate SO₂ variations. We utilized the dual-wavelength SO₂ assessment method described in Ref. 29. At each target blood vessel location, 100 PA measurements with both wavelengths were recorded and averaged to suppress noise effects.

3 Results

Results of the CAM imaging are shown in Fig. 3. The two treatment regions in the PAM images [Figs. 3(a) and 3(b) (chick embryo #1) and Figs. 3(c) and 3(d) (embryo #2)] received the same illumination irradiance (20 mW/cm²) before and after the same PDT duration (20 min). Figures 3(a) and 3(b) are the PAM images of the comparison control area taken before and after this treatment. Although vasculature remained the same in the image of the control area [Figs. 3(a) and 3(b)], a dramatic change was visualized in the treatment region [Figs. 3(c) and 3(d)]. Major blood vessel structures (with diameters <100 μm) were destroyed. Capillary networks too small to be resolved were seen as cloudy signal areas. Such capillary beds were also partially damaged, leaving bleeding/leaking spots (bright spots) with extravasation. We measured the size change of blood vessels with an original diameter of 242 ± 20 μm [boxed in Figs. 3(e) and 3(f)] along with the treatment, as is shown in Fig. 4. On average, the diameter of the blood vessel experienced a steady decrease of up to ~60% of its original value. After the therapy, discontinuities appeared in the circled vessel, which may be thrombi after the treatment.

Resultant images of the rat ear imaging are shown in Fig. 5 with the same color map. Although no obvious morphological changes were observed in the control group animals as is shown in Fig. 5(a), both PAM images and white-light pictures of the treatment area [darkened area in the left image of Fig. 5(b)] showed destruction of blood vessels. Absence of blood vessel structure was observed in the white-light pictures, with enlargement of the bleached area during the treatment. This suggested variations of vasculature and was confirmed by destruction of vessel structures in PAM images [top row of Fig. 5(b)]. We measured diameter change of one ~200-μm target vessel in the treatment areas induced by PDT in each of the three rats (labeled with dashed rectangular). It should be noted that more serious destruction of blood vessel structures in the 15-min PAM image.

![Fig. 3](image-url)  
Structural variations of blood vessel vasculature on a chorioallantoic membrane model. (a) and (b) control group; (c) and (d) vasculature change before and after a 15-min treatment with 20 mW/cm² light power; (e) and (f) vasculature before and after a 30-min photodynamic therapy (PDT) with 10 mW/cm² optical power.

![Fig. 4](image-url)  
Size change of an ~242 ± 20-μm target blood vessel along with a 30-min PDT.
may be due to dilation of soft tissue caused by PDT. Neverthe-
less, the quantitative analysis of blood vessel change did show
size decrease. We repeated the experiments (n = 3 animals), and
the results are presented in Fig. 6 as mean ± standard error
of the measurement. Target vessels shrank to ~40% of their original
size after 30-min treatments. This change agrees with previous
reports in the literature describing blood vessel diameter decrease
due to blood vessel constriction, which is a normal biological
response to PDT. This was followed by destruction of
blood vessel architecture, which was observed after a 45-min
treatment.

We longitudinally examined the oxygen saturation levels of
target vessels during the treatment to demonstrate the capabil-
ities of PAM in real-time monitoring of oxygen levels in PDT,
and the data presented in Fig. 7(a) are the average SO₂ change
on one target vessel in each of the three animals, and in Figs. 7
(b)–7(d), we plot the data points collected on all the three ani-
imals. Measurements of SO₂ in the target vessels were obtained
at 15-min intervals during treatments. A quick decrease of oxy-
genation was observed after the first treatment interval, which
then gradually recovered even with the presence of the treatment
light. The recovery reached a level slightly lower than its original
status, along with constriction of blood vessels after 30 min
of treatment. As destruction of blood vessels begins to appear
after 30 min of treatment, as is shown in Fig. 5, oxygen levels
measured 45 min after the initiation of PDT showed significant
inconsistency. Nevertheless, an immediate drop-recover cycle
was observed in the three treated animals (n = 3).

Fig. 5 Vasculature change of a rat ear model with a 45-min PDT: (a) results from the control group with only
treatment light but no photosensitizer and (b) white-light pictures and PAM images of the treatment area.

Fig. 6 Target blood vessels (≈200 µm) size change of rat ear models.

Fig. 7 (a) Average oxygen saturation (SO₂) level change of target blood vessels with 60 min PDT and
(b) to (d) the change curve of each of the three samples.
4 Conclusion and Discussion

We report our initial feasibility studies of parametric imaging of vasculature response to PDT with PAM with both structural and functional parameters. We measured size and oxygen level variations in single blood vessels during PDT.

We observed consistent major blood vessel size decrease along the PDT. We believe this is due to blood vessel constriction described in previous reports. This follows destruction of the blood vessel and thrombus, which can be observed as blood vessel discontinuities in PA images. Microcapillaries whose dimensions are below the resolving capabilities of our imaging system were observed as cloudy structures. During PDT, destruction of these structures lost PA signals, leaving black areas in the PA images. However, it is interesting to note that in some areas, bright spots can be observed after the treatment. We believe this is micro-hemorrhaging due to damage of small blood vessels, but this still needs careful studies to verify.

PA imaging has the capability of imaging oxygen saturation in blood vessels via assessing the relative concentration of oxygen and deoxyhemoglobin. To demonstrate that this can be utilized in monitoring PDT, we longitudinally measured SO2 level change in target vessels during the treatment. We observed a quick SO2 level decrease after the treatment was initiated, which we believe was a result of immediate consumption of oxygen due to generation of reactive species. This followed a gradual oxygenation recovery in the vessels afterward, which ended at a level that was lower than its original status. However, we would like to note that relatively big variations of SO2 levels were observed at the end of each treatment experiment. The inconsistence might be due to the complex status of blood vessels including blood vessel structural damages.

This study, however, has several limitations. Structural information monitoring was not in real time. The treatment was stopped temporarily to perform the measurements as mechanical scanning of the target was required by our current imaging system. This limitation could be solved by development of a faster imaging system. Though only superficial vasculature parameters were imaged in the present report, PAM is scalable to deeper depths at the cost of sacrificing spatial resolution, so imaging to depths of 2 to 3 mm is anticipated. Whereas our system is immediately useful for medical conditions such as port wine stain, one can turn to micro-endoscopic PA imaging systems for imaging internal body structures.

A number of directions can be enlisted as our future work. Examples include studying responses to PDT in terms of these parameters, such as blood flow velocity, PA imaging of PS distribution before PDT, PA-guided light dosimetry estimation, etc. These efforts will significantly benefit the PDT practices such as personal treatment planning, therapy monitoring, and prognosis.

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References


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