Complete blood vessel occlusion in the chick chorioallantoic membrane using two-photon excitation photodynamic therapy: implications for treatment of wet age-related macular degeneration

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Abstract. Complete blood vessel occlusion is required for the treatment of age-related macular degeneration (AMD). AMD is the leading cause of blindness in developed countries and current treatment regimes have potential to cause collateral damage, or do not remove pre-existing unwanted vasculature. It has been proposed that two-photon excitation (TPE) photodynamic therapy (PDT) can be applied to cause local blood vessel occlusion without damaging surrounding retinal tissues. The in ovo chicken chorioallantoic membrane (CAM) is used as the model for vascularization in the wet form of AMD; novel techniques for the utilization of the CAM are reported. Complete occlusion of CAM vessels ~15 μm in diameter is achieved using the clinically approved photosensitizer Verteporfin (Visudyne®), QLT, Incorporated, Vancouver, British Columbia, Canada) and TPE activation. The average and peak irradiances used for treatment are 3.3 × 10^9 W/cm^2 and 3.7 × 10^11 W/cm^2, respectively. A total fluence of 1.1 × 10^8 J/cm^2 is the dosage required for successful occlusion, and it is expected that for optimal conditions it will be much less. These results are the first proof-of-principle evidence in the literature that indicate TPE-PDT can be used to occlude small blood vessels. Further investigation will help determine the utility of TPE-PDT for treating wet AMD, perhaps through targeting feeder vessels.

Keywords: age-related macular degeneration; two-photon excitation; photodynamic therapy; chorioallantoic membrane.

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1 Introduction

Central vision loss in wet-form age-related macular degeneration (AMD) is the leading cause of blindness in the elderly population in developed countries, and is caused by the growth of choroidal blood vessels into the normally avascular macular region of the retina. This choroidal neovascularization tends to be leaky with fluid exuded into the surrounding tissues, which adds to the vision loss. Currently, there is no cure for wet AMD, but patients do have therapeutic options: laser photocoagulation therapy, antivascular endothelial growth factor (VEGF) therapy, and photodynamic therapy.1,2 In general, treatment is directed toward closing the blood vessels, i.e., occlusion.

Laser photocoagulation therapy cauterizes blood vessels using a high intensity laser beam that causes local tissue heating. This cannot only cause collateral damage, but recurrence rates are high (up to 50%).2,3 Photocoagulation is typically used for extrafoveal neovascularization to minimize damage to the central macula. Anti-VEGF therapy is a relatively new treatment for wet AMD using the drug Macugen® (Eyetech and Pfizer, Incorporated, Cambridge, Massachusetts) but has several drawbacks. It does not stop the growth of blood vessels, but merely slows down growth, and it does not eliminate pre-existing neovascularization.1 The treatment, which is still being tested for optimal drug dose, requires direct intraocular injection, and effects beyond two years are not known.1,4 Currently, the treatment has not shown damage to the healthy tissues surrounding the neovascularization.

The standard treatment regime for wet AMD is photodynamic therapy (PDT), referred to here as one-photon excitation (OPE) PDT, using the porphyrin-based photosensitizer, Visudyne® (QLT Incorporated, Vancouver, British Columbia, Canada). This is activated by light and, in the presence of oxygen (Fig. 1), can produce apoptotic cell death within the treated tissue.3 Visudyne® localizes in areas of new tissue growth and therefore accumulates in the macular neovascularization in wet AMD. In practice, the patient is injected with...
6 mg/m² of Visudyne® over a 10-min period, the drug is allowed to circulate and localize for 15-min, and the neovascular area is then treated with 600 mW/cm² of 680-nm light (Fig. 2) for a period of 83 s (50 J/cm²).² Although successful in closing choroidal neovascularitation, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collagen closing choroidal neovasculature, it has been shown in the cynomolgus monkey model that OPE-PDT can also cause collateral damage to the retina and nearby tissue layers.⁶,⁷

In practice, the success of OPE-PDT is measured by the retention or gain in visual acuity and visualization of vessel leakage. These tests do not allow inspection of surrounding tissues, such as the retinal pigmented epithelium (RPE) and Bruch’s membrane, to quantify collateral damage. Human histological studies of these tissues are rare, as the treated eye must be excised to analyze damage. The first to report histological images of a human eye treated with OPE-PDT was Ghazi et al. in 2001.⁸ One patient was studied and the tissue was received from a submacular membrane resection 27 days after OPE-PDT treatment. Ghazi et al. primarily looked at structural changes in the vasculature; however, they did report structural changes in the RPE that were similar to those in untreated cases. Another important study by Schnurrbusch et al. in 2001⁹ looked at excised eyes from two patients who had undergone surgical extraction of the choroidal neovascularization three months after treatment with OPE-PDT. Schnurrbusch et al. suggest that the RPE degeneration seen in these two cases could be caused by the PDT treatment, but results were inconclusive. Schlötzer-Schrehardt et al. (2002)¹⁰ demonstrated that mild RPE detachment from Bruch’s membrane occurred after traditional OPE-PDT therapy. In this study, one eye from each of three patients with malignant melanoma of the uvea was used. The results from these studies are slightly contradictory and are not statistically relevant; however, damage to retinal tissues has been suggested. Additionally, optic nerve ischemia and retinal detachment have been observed in OPE-PDT treatment of papillary capillary hemangioendothelioma.¹¹ We have hypothesized that two-photon excitation (TPE) PDT could be used to reduce this collateral damage, which may contribute to the need for repeated treatments in OPE-PDT for AMD and other choroidal neovascular diseases.¹²,¹³

TPE is widely used in confocal microscopy.¹⁴ In general, it requires high fluxes of light that can be delivered using ultrashort pulsed and focused laser beams, usually in the near-infrared range (NIR, λ ~ 750 to 950 nm). TPE-PDT has the potential to eliminate the collateral damage in treating AMD, due to the limited focal volume that is excited. The probability of a molecule absorbing two photons simultaneously (each half the energy, i.e., twice the wavelength as in OPE) is extremely small and is proportional to the square of the light intensity. This process occurs only at the focal plane of a tightly focused laser beam, with focal volumes of a few femtoliters (defined by 1/e² of the maximum intensity for a Gaussian beam), using a high numerical aperture lens (NA > 0.9).¹⁵ This highly confined TPE focal volume allows, in principle, selective activation of photosensitizer within the neovascularization, leaving over- and underlying tissues unaffected, even if they contain photosensitizer.

There are a number of reports of TPE of photosensitizing drugs, including Photofrin, protoporphyrin IX (PPIX), psoralens, Rose Bengal, hematoporphyrin derivative (HPD), hypocrellins, and Verteporfin (the active ingredient in Visudyne).¹⁶–²⁵ We have confirmed using Verteporfin that, as would be expected, the photophysical processes remain the same after the molecule has reached its first excited state (Fig. 1), regardless of whether excitation occurs through one- or two-photon absorption.¹³,²⁵ Verteporfin has a larger two-photon cross section (31 GM units at 865 nm, where GM is units of Goppert-Meyer and 1 GM = 1 × 10⁻⁴⁹ cm²/s/molecule) compared to other clinical photosensitizers such as Photofrin® (7.5 GM units at 865 nm).¹²,²⁴,²⁶ At 780 nm, the wavelength used here, the value is 61 GM units. Its one- and two-photon excitation spectra can be seen in Fig. 2.

Recently, two articles have been published by the Toronto group demonstrating TPE-PDT in vitro. Karotki et al. (2006)²⁴ and Khurana et al. (2006)²⁶ showed that TPE-PDT of rat pros-
trate endothelial cells using Photofrin and Visudyne® resulted in cell death, with Visudyne® being an order of magnitude more potent than Photofrin. To our knowledge, there is no published evidence for the use of TPE-PDT to completely occlude blood vessels.

Blood vessel occlusion by OPE-PDT has been shown in a number of models, including chicken chorioallantoic membrane (CAM), and the retinas of monkeys, rabbits, and rats. To prove the principle of TPE-PDT for blood vessel occlusion, we have chosen the CAM model, with a number of modifications as described later. The CAM grows between days 3 and 12 of gestation and is used as an external lung and waste exchange system. It is a good model for wet AMD, since it has a wide range of blood vessel sizes (a few microns to several hundred microns in diameter); the vessels are undergoing rapid angiogenesis from day 5 to 9 of gestation and are embedded in transparent tissue similar to the retina; it has been well characterized in studies of tumor angiogenesis, biosensor and detector responses, and PDT itself; moreover, the fertilized chicken eggs are widely available and inexpensive. To further enhance its relevance to wet AMD, only the CAM arteries were used, simulating the feeder vessels that are seen in this disease. In these feeder vessels, blood flows from the choriocapillaries into the subretinal space. In the CAM model, the arteries are the vessels with blood flowing away from the body of the embryo and are the vessels of interest. The veins, with blood flowing toward the embryo, were observed but were not treated.

The ex ovo CAM was used previously in our laboratory to demonstrate its feasibility as a model for AMD, and to study photosensitizer distribution and localization. We demonstrated that when Verteporfin was injected into the blood vessels in microliter amounts, the photosensitizer flowed through the vessels as a compact drug bolus with the leading edge remaining compact, while the trailing edge distributed monoeuropolarly with time. Verteporfin localized most strongly in the upper surface of the CAM (closest to the shell when in ovo), indicating a potential target for TPE-PDT treatment. Using a highly focused laser beam, we were able to demonstrate partial or complete vessel ablation (visible physical tissue damage leading to necrosis), which allowed us to create “leaky” vessels in the absence of exogenous photosensitizer. However, as the focus of this study was to demonstrate the effectiveness of the CAM, we did not study the process of TPE-PDT; nor did we see any evidence of photosensitizer-induced blood vessel occlusion, although similar laser powers were used (40 to 200 mW measured as the beam entered the microscope) to those reported later. The ablation that was observed in this ex ovo model occurred when the CAM was immobilized and slightly dehydrated. Moreover, the uppermost portions of the CAM were photothermally removed as part of the procedure used to close blood vessels photothermally. In the current study, we modified the model so that TPE-PDT treatment and response assessments could be performed in ovo. In this setting, the embryo remains in the shell, reducing potential damage to the CAM, yolk, or embryo, while the egg also retains heat more easily than in the ex ovo system and the CAM stays moist longer (up to 30 min) without external application of Ringer’s solution. The embryo also has increased viability when it remains in the shell. We present novel, proof-of-principle results for TPE-PDT using the CAM, as well as briefly describe novel handling techniques for this membrane in ovo.

2 Materials and Methods

2.1 Chorioallantoic Membrane Preparation

Fertilized chicken eggs were obtained from a local farm (Airrie, Alberta, Canada) and stored at 14°C for up to two weeks. They were removed from the refrigerator at required intervals, cleaned with 70% v/v ethanol, and placed in a 38°C humidified incubator for the duration of their development. The eggs were not rotated so that the position of the embryo would be known, since it always rises to the uppermost portion of the egg. The terminology for the position of the embryo, as it will always move to the uppermost portion of the egg. The bottom side of the egg refers to the portion of the shell directly across from the embryo. Incubation and experimentation were performed with the horizontal axis parallel to the ground. The preparation of the egg is shown in (a), (c), and (d) and described fully in Sec. 2.1.

![Fig. 3 Illustration of the terminology and procedure for windowing the side of an egg to use in in ovo experiments. The terminology for the in ovo embryo is described in (a). The blunt end refers to the end of the egg with the lower radius of curvature and can be visualized as the end with the air sac. The pointed end of the egg has a higher radius of curvature and lacks an air sac. The top side of the egg refers to the position of the embryo, as it will always move to the uppermost portion of the egg. The bottom side of the egg refers to the portion of the shell directly across from the embryo. Incubation and experimentation were performed with the horizontal axis parallel to the ground. The preparation of the egg is shown in (a), (c), and (d) and described fully in Sec. 2.1.](image-url)
the bottom side [Fig. 3(c)]. Cellulose tape was used to cover the top side of the shell and a hole was cut using small dissecting scissors [Fig. 3(c)]. The window was then covered with cellulose tape and the egg returned to the incubator until experimentation on day 8 or 9 [Fig. 3(d)]. The embryo naturally rises to the top, windowed side of the egg, within minutes to hours.

On the day of experimentation, the egg was removed from the incubator, the cellulose tape was discarded, and the window widened by cutting away more shell, if necessary. The CAM remained sufficiently moist if the treatment procedure was kept under 30 min. After treatment was completed, the window could be covered again with fresh cellulose tape and the egg returned to the incubator for observation.

2.2 Photosensitizer Preparation and Administration

A stock solution of Verteporfin (QLT Incorporated, Vancouver, British Columbia, Canada) was prepared by dissolving the lyophilized powder in dimethyl sulfoxide (Sigma, USA), as suggested by QLT Incorporated for optimum solubility and storage. The concentration (1.438×10^{-4} M) was determined spectrophotometrically (λ = 688 nm, ε = 32740 L·mol⁻¹·cm⁻¹). Although dimethyl sulfoxide is used as a stock solution solvent, it is inappropriate to use large amounts of it when injecting into the CAM. Therefore, a liposomal injectate, 0.01-mg/mL Verteporfin to 1.0-mg/mL dioleoyl-phosphatidylcholine (DOPC), was designed to optimize the solubility of Verteporfin and vesicle size and function, while maintaining endogenous components. The injectate was prepared as follows. DOPC in chloroform (25 mg/mL; Avanti Lipids, USA) was added to a glass vial and the chloroform was evaporated off under a gentle stream of nitrogen gas. Verteporfin and phosphate buffer (0.02 M, pH 7.0) were then added and stirred for 30 min to resuspend the dried lipid. Lipid vesicles were formed by subjecting the Verteporfin-lipid solution to 10-min intervals of stirring and sonication in the dark until the solution cleared. We have previously shown that this protocol produces small unilamellar vesicles (SUVs) and some small multilamellar vesicles (SMVs, ~100 nm). These solutions could be kept in the refrigerator for up to one week, with sonication on reuse to disperse any large aggregates. The liposomal solutions were filtered before use with a 0.02-μm hydrophilic cellulose acetate syringe filter (Albet, USA) to remove dust, debris, or remaining aggregates that could potentially clog the capillary needles used for injection.

2.3 Two-Photon Excitation Photodynamic Therapy Equipment

The TPE-PDT setup has been described previously. Briefly, the laser beam was guided through a Zeiss 20× objective lens with a 1.0-cm working distance and 0.4 numerical aperture. Laser intensities were measured at the back plane of the microscope. A 75% loss in intensity has been recorded through the objective lens, so the laser intensities reported have been adjusted to reflect this attenuation. Studies utilizing a Spectra Physics (Palo Alto, California) Tsunami laser (82 MHz or CW) were carried out at 780 nm (Fig. 2). TPE experiments were performed with 100-fs pulse length at a repetition rate of 82 MHz at an average laser intensity of 38 mW, for which the focal volume has previously been determined to be 35±2 μm³ using fluorescence correlation spectroscopy (FCS) with 0.04-μm streptavidin conjugated polystyrene FluoSpheres (Invitrogen, Missasauga, Canada). The radius r of the TPE focal volume is 1.8 μm and the height along the axis of light propagation z₀ is 5.5 μm. The TPE focal volume V_f was calculated using:

$$ V_f = \left( \frac{\pi}{2} \right)^{\frac{3}{2}} r^2 z_0, $$

where V_f is the volume of a symmetrical ovoid and is 35 μm³. This volume is only weakly dependent on the two-photon cross section, so that the focal volume with Verteporfin volume was taken as equivalent to that measured with the FluoSpheres calibration standard.

The average irradiance I(average) of the TPE was calculated as:

$$ I(\text{average}) = \frac{\pi(P)}{2} \left( \frac{n}{\lambda} \right)^2 (1 - \cos \alpha)(3 + \cos \alpha), $$

where $P$ is the average laser power at the sample (0.038 W), $n$ is the refractive index of the immersion media ($n_\text{imm}=1.0008$), $\lambda$ is the wavelength (780 nm), and $\alpha$ is the semiaerture angle of the objective lens [$\alpha=\sin^{-1}(\text{NA}/n) = 2.36 \times 10^3$]. From the average irradiance, the peak irradiance $I(\text{peak})$ can be determined by assuming a Gaussian pulse shape in:

$$ I(\text{peak}) = \left( \frac{\ln 2}{\pi} \right)^{1/2} \frac{2}{\tau f} I(\text{average}), $$

where $\tau$ is the pulse duration [full width at half maximum (FWHM), 100 fs], and $f$ is the pulse repetition rate (82 MHz).

Control experiments were undertaken with several different light sources. White light experiments illustrating OPE-PDT were executed with a dual-gooseneck overhead lamp (150 W, Mille Luce M1000, Novato, California). This light was unfocused and covered the entire surface of the CAM. Other OPE-PDT controls were performed using the previous laser system operating out of pulsed mode, such that the sample received a CW irradiation with a beam waist (ω₀) of 2.4 μm, calculated from:

$$ \omega_0 = \frac{1.22 \lambda}{\text{NA}}. $$

The final control was undertaken with the pulsed light source without photosensitizer injection to rule out thermal effects.

A microscope stage, altered to hold the eggs, was fabricated, starting from a standard Zeiss upright stage [Fig. 4(a)]. A metal ring, shown in Fig. 4(b), secures the egg horizontally with the window facing upward. This ring was then placed into the microscope stage. The main body of the stage allows for microinjection while observing the blood vessels through the microscope eyepiece using low intensity white light. Injection was performed while visualizing the CAM using a 5× magnification objective lens (Zeiss). The micromanipulator and microinjection system were secured to the stage by a...
rigid arm that allowed the needle to move with the stage when focusing or recentering the needle or blood vessel within the field of view.

The injection system included an $x, y, z$ micromanipulator (MM-33; Sutter Instruments Company, Novato, California) that attaches to the stage arm [Fig. 4(a)]. The manual micro-injector (Sutter) sits to the side of the microscope and Teflon tubing connects it to the needle, which is held in the micromanipulator by a plastic sleeve. The needles used for injection were made by pulling borosilicate glass capillaries with a quick-fill filament (1B100F-4, World Precision Instruments Incorporated, Sarasota, Florida) using a micropipette puller (SP-30, Sutter). The tapered end of the needles were then beveled with a micropipette beveller (BV-10, Sutter) at 10 deg on an optically flat abrasive plate (104D) at 60 rpm to achieve tips that would penetrate thicker blood vessel walls. The average outer diameter of the needle tips was 0.02 mm.

White-light images (still and video) were recorded with a digital camera (Canon PowerShot A70) through the ocular lens of the microscope before and immediately after treatment. The videos found at http://www.chem.ucalgary.ca/research/groups/dcramb appear green due to the dichroic mirror required for laser beam steering. This dichroic also enhances the image contrast between the blood vessels and the surrounding tissues.

### 2.4 Injection and Treatment Parameters

20 eggs in total were used for the experimentations described within this article. Four studies were undertaken (TPE-PDT, and controls 1, 2, and 3), and each of these studies required five trials, with one egg/CAM used for each trial. Injections were always performed in an artery, specifically the largest feeder vessel accessible, downstream from a branch [Fig. 5], since, as mentioned before, the arteries are likely to best represent the feeder vessels seen in wet AMD. The diameter of these vessels was ~200 to 500 $\mu$m. TPE-PDT treatment was always performed on at least the third branch (B3), as illustrated in Fig. 5. These blood vessels were approximately 15 $\mu$m in diameter. For these small vessels, the laser was focused into the main blood stream for treatment. The horizontal and vertical cross sections of the focal volume relative to the treated blood vessels are illustrated in Figs. 5(b) and 5(c).

For the experiments involving application of Verteporfin, 50 $\mu$L of the injectate was administered over a period of several minutes, 2 to 3 $\mu$L at a time, similar to the slow injection...
of Visudyne® clinically.2 Fast injection caused blood flow to cease in the vasculature directly surrounding the injection location, marking the needle, relative to the treatment spot (a). (a) The needle was positioned on an artery feeder vessel, such that it was downstream from a branch and with the flow of blood. The injected drug flows through the feeder vessel and is carried along the branches (B1, B2, and B3). Treatment was performed on the smallest visible branch, normally B3. The dimensions of the excitation volume (faded circle (b) and faded oval (c)) relative to B3 are illustrated. The diameter of B3 (d_B) is ~15 μm. The diameter of the TPE focal volume (d_f) is (b) 3.6 μm and its height (z_f) is (c) 5.5 μm.

3 Results and Discussion
3.1 Chorioallantoic Membrane Preparation
In the past, in ovo windowing of the CAM has been performed through the blunt end of the egg, as reviewed by Richardson and Singh.35 Previous OPE-PDT studies using the CAM have utilized this technique.39–45 In our work, the top side of the egg was windowed, such that there is a larger area of the CAM exposed [Fig. 4(b)]. Removing the shell from the side of the egg also removes the inner-shell membrane, reducing the risk of it adhering to the CAM and interfering with treatment.

Initially, the eggs were windowed while the embryo remained at the top side. However, the CAM is quite sticky and often remains secured to the shell and, if cut or nicked, embryo viability decreases markedly. Yolk leakage also greatly reduces embryo survival.35 Therefore, the shells were always opened with the embryo in the bottom half [Fig. 3(c)]. Richardson and Singh35 recommend draining the albumin at day 3 or 4 and windowing between day 4 and 10. However, we have found that it is best to window the egg at day 4 to 5.5 to maximize embryo viability and surface tension of the CAM for injection.

3.2 Two-Photon Excitation Photodynamic Therapy Responses
TPE-PDT experiments were performed with a pulsed beam (equivalent to 390-nm laser light, Fig. 2) at an intensity of 38 mW and a peak intensity of 4.6 × 10^12 W/pulse for 5 min. The treatment parameters are found in Sec. 2, Materials and Methods, and summarized in Table 1. It was determined using FCS that the TPE volume broadening was not occurring as the laser beam passed through the CAM. The TPE volume was first determined with amino PEG quantum dots (Invitrogen, Mississauga, Ontario, Canada) in water, and then measured in a blood vessel filled with the same quantum dot solution. The effective TPE volume was found to be 37 μm³ (see Appendix).

The TPE-PDT treatments were successful in occluding small blood vessels ~15 μm in diameter in all five trials. Figure 6 shows white-light images before treatment (and injection) and immediately after treatment; preliminary results observing the blood vessels 6 h after treatment are available in the Appendix. In all cases, blood stasis (as noted in the small capillaries branching off of the treated vessel) occurred within the time span of the 5-min treatment, indicating that the irradiance required for occlusion may be less than what was used. This would have to be confirmed in separate experiments, since lower PDT doses may not induce permanent occlusion. White-light imaging was not used during the PDT irradiation to avoid causing additional light exposure, given that the light dose required for one-photon activation is orders of magnitude less than for two-photon.74 With our present experimental setup, it is virtually impossible to observe the CAM several hours posttreatment to determine whether vascular collapse has occurred. Moreover, because imaging the response after treatment with white light may cause additional OPE-PDT damage, we imaged immediately after treatment. This also allowed us to verify that the laser spot remained within the vessel of interest by visual inspection of the laser spot position on the CAM before and after treatment. In principle, movement of the embryo could cause the laser spot to shift off the blood vessel; this did not occur in any of the five trials. It is also very difficult to relocate the treatment spot at later times, given that the vessel virtually disappears in the

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white-light images when it becomes occluded [Fig. 6(a)], and also because the CAM is undergoing rapid angiogenesis. Marking the treatment area, for example using Teflon locator rings, dying the CAM, referencing shell fragments on the CAM surface, or relying on the pretreatment photographs, was attempted. Teflon rings and dying caused damage to the surrounding vasculature, so that changes solely due to TPE-PDT could not be isolated. Since the blood vessel pattern changes significantly over a few hours, identifying the treated vessel just by visual inspection was not feasible. Alternate lighting and imaging approaches to overcome this limitation and allow longer term response assessment, while still avoiding OPE-PDT artifacts, are being explored.

It is evident in Fig. 6(b) that the treated vessel shuts down completely: not only does the blood flow stop, but the red blood cells are completely evacuated from the treated region. The movement of blood in surrounding vessels can be viewed with the supplementary videos provided at http://www.chem.ucalgary.ca/research/groups/dcramb. The observed emptying of blood from the vessels is similar to the findings of Lange et al.\(^{39}\) using OPE-PDT, and suggests that some level of vascular collapse has occurred; whether this is permanent occlusion has yet to be determined. In OPE-PDT, it has been shown that, in general, treated blood vessels undergo an initial period of stasis before apoptosis causes vascular collapse.\(^{5}\)

In clinical OPE-PDT therapy for AMD, it is observed that although vascular collapse is immediate, reperfusion can occur over a period of days and weeks after treatment. The human studies conducted suggested that reperfusion occurs in vessels that were occluded, owing to a local inflammatory response, rather than directly occluded by PDT. It was also observed in clinical trials that almost all treated patients experienced a reoccurrence of some fluorescein leakage one to three months after OPE-PDT.\(^{30}\) However, this leakage is widely believed to be due to a recurrence of angiogenesis. Because we are observing the blood vessel closure immediately after treatment, there is some concern that the vessels have not closed permanently.

| Table 1 Summary of TPE-PDT and control experimental parameters. * indicates energy per pulse calculated from parameters provided. |
|---------------------------------|-----------------|----------------|----------------|-----------------|-----------------|
| **Photosensitizer concentration** | CAM TPE-PDT (Verteporfin) | Control 1 | Control 2 | Control 3 | Endothelial TPE-PDT (Photofrin) (Ref.24) | Endothelial TPE-PDT (Verteporfin) (Ref.26) |
| mg/mL | 1.0×10^{-2} | 1.0×10^{-2} | 1.0×10^{-2} | 0 | 2.5×10^{-2} | 7.2×10^{-3} |
| Wavelength (nm) | 780 | Broad | 780 | 780 | 850 | 850 |
| NA of objective lens | 0.4 | N/A | 0.4 | 0.4 | 1.2 | 1.2 |
| Immersion substance | Air | Air | Air | Air | Water | Water |
| Pulse width (fs) | 100 | - | - | 100 | 300 | 300 |
| Repetition rate (MHz) | 82 | - | - | 82 | 90 | 90 |
| Power (mW) | 38 | 50 | 38 | 38 | 10 | 7.4 |
| Energy per pulse (nJ/pulse) | 0.46 | 50 | 38 | 0.46 | 0.11* | 0.11* |
| Average irradiance (W/cm²) | 3.3×10^6 | 6 | 2.1×10^{-3} | 3.3×10^6 | 7.5×10^6 | - |
| Peak irradiance (W/cm²) | 3.7×10^{11} | 6 | 2.1×10^{-3} | 3.7×10^{11} | 2.6×10^{11} | - |
| Fluence (J/cm²) | 1.1×10^8 | 1800 | 0.63 | 1.1×10^8 | 6300 (50% killing) | 750 (90% killing) |
Although we have demonstrated immediate occlusion by TPE-PDT, there is some evidence of partial reperfusion in the longer term (i.e., 6 h posttreatment, see Appendix). It is therefore important to address the term “immediate” when discussing the observation of vascular collapse and blood stasis caused by PDT treatment. We describe “immediate” within this work as seconds after the TPE-PDT treatment stops. Immediate stasis in a clinical sense (characterized by lack of vessel leakage) can be anywhere from several hours (i.e., 5 h\(^5\)) to one week.\(^6\) Clinically, the observation of changes in blood flow has not been documented in the time frame of a few seconds after terminating treatment. Therefore it is difficult to compare our TPE-PDT results with the clinical OPE-PDT, as the observations are on vastly different time scales. However, there is evidence in the literature that suggests OPE-PDT vessel closure, or at least stasis, can occur during blood vessel treatment and in the same time scale as described here.

For instance, Li et al. (2006)\(^52\) investigated real-time imaging of vasculature in the rat xenograft model of Dunning prostate cancer during PDT using Photofrin as the photosensitizer. Li et al. found that although vessels had somewhat varying responses, significant decreases in blood flow (measured in vascular index) were observed with an average decrease of 76%. In some cases, complete stasis was observed without recovery up to 10 min after treatment. The vessels observed by Li et al.\(^52\) are four to five times larger than the ones observed in the TPE-PDT treatment described in our current study. Additionally, Lim et al. (2006)\(^53\) observed vascular occlusion, implying stasis (from fluorescein injection), as well as complete vessel shutdown using the photosensitizer DH-I-180-3 in an EMT6 mammary tumor model. These observed vessel changes occurred on a time scale of a few minutes after treatment. In both of these cases, successful OPE-PDT treatment was reported, indicating that the results we see within seconds of terminating treatment are indicative of effective TPE-PDT.

Although this is the first demonstration of complete blood vessel occlusion, there have been several other reports of TPE experiments.\(^13,16,18,23,25,54,55\) In particular, Karotki et al. (2006)\(^54\) and Khurana et al. (2006)\(^6\) measured the two-photon spectra of Verteporfin and Photofoir, and showed the quadratic dependence on TPE light dose of apoptotic cell death in a monolayer of cultured vascular endothelial cells. The corresponding treatment parameters are summarized in Table 1. Although the irradiation setup for TPE-PDT irradiation (which was done using a confocal laser-scanning microscope) was very different from that used in these CAM studies, the average and peak irradiances were of the same order of magnitude and also similar to those of King and Oh (2004),\(^55\) who reported singlet oxygen production using 5-(and 6)-chloromethyl-2',7'-dichlorodihydrofluorescein diacetate, acetyl ester with a peak power density (or peak irradiance) of 3 \(\times 10^{11}\) W/cm\(^2\).

The fluence for TPE-PDT blood vessel occlusion (1.1 \(\times 10^{6}\) J/cm\(^2\)) represents the total light dose delivered, assuming that the focal spot remains focused at the same spot on the blood vessel during treatment. However, any movement of the embryo can cause slight movements of the CAM. During these five trials, we assume that the laser spot remained somewhere within the treated vessel, because after the 5-min treatment, the laser spot was still within the blood vessel. We feel that this is a valid assumption, as any strong movement will cause the laser to wander off of the vessel, and in our experience it does not return to exactly the same location. Estimating the maximum movement to be 15 \(\mu\)m (the diameter of the vessel) reduces the fluence to 1.6 \(\times 10^{6}\) J/cm\(^2\). This is also much greater than the value reported by Khurana et al.\(^6\) The difference is attributed to the intermittent application of light during the laser beam scanning process reported. In fact, one might expect that lower TPE-PDT dose levels are needed to kill a 0.23 \(\times 0.23\)-mm area of monolayer cultured cells than to close a vessel of 15 \(\mu\)m diameter. Also, blood vessel closure occurred within the 5-min treatment, indicating the fluence may be too large (i.e., treatment time could be shortened).

The pulse energy used here (0.46 nJ/pulse) is lower than in some other TPE-PDT studies, e.g., 2.2 nJ/pulse to 200 \(\mu\)J/pulse have been reported with beam waists varying from 0.4 \(\mu\)m to several millimeters.\(^16,17,54,55\) However, the pulse energy does not take into account the size of the treatment (focal) volume or the laser repetition rate, and therefore is not a relevant PDT dose parameter to use in comparison to different TPE-PDT studies.

Additionally, direct comparison between light dose for OPE-PDT and TPE-PDT can be misleading. Because of the miniscule cross sections for TPE, significant peak intensities
are needed to drive the process. Therefore, one should consider the excitation rate of the sensitizer, instead of fluence, when comparing OPE versus TPE. As is demonstrated in the control experiments, even at high fluence, no collateral damage was observed.

The measured focal volume \((z_0=5.5 \, \mu m, r=1.8 \, \mu m; 35 \, \mu m^3)\) for our studies fits easily within the 15-\(\mu m\) blood vessels (Fig. 5), and the laser focal spot was focused within the upper half of the blood vessel for all experiments. The blood vessel wall is transparent and very thin (\(\sim 1.5 \, \mu m\)) compared to the height of the laser spot, and it is difficult to determine whether or not the laser spot was held directly on the blood vessel wall. However, in larger vessels the focal volume could be focused on the vessel wall to target directly the endothelial cells, as it is believed that the destruction of these cells during apoptosis is the cause for vascular collapse.5

The thickness of a blood vessel wall is typically \(\sim 10\%\) of the blood vessel diameter,36 so that in vessels \(>55 \, \mu m\) diameter, the beam could be localized within the vascular wall. In fact, a study of feeder vessel photocoagulation55 indicated that the closure of vessels of 50 \(\mu m\) diameter or smaller, lead to an increase in visual acuity and a reduction in choridal neovascularization (CNV) extent. However, in this study extensive collateral damage to the retina in the region of the feeder vessels was also observed.

### 3.3 Controls

Each of the following control exposures, summarized in Table 1, was carried out using five eggs, each CAM receiving treatment to one blood vessel.

#### 3.3.1 Control 1: can blood vessel closure be achieved with white light?

Control 1 used the same 50-\(\mu L\) injection to determine if this was an appropriate drug dose and to provide a visual comparison for blood vessel closure in the TPE-PDT treatment. Several minutes after photosensitizer injection, the entire CAM surface was irradiated at 6 W/cm\(^2\) with broad-spectrum white light from an overhead goose-neck lamp for 5 min (1800 J/cm\(^2\)). Images were recorded immediately after treatment ended.

Control 1 (white light with Verteporfin injection, Table 1) was performed to illustrate that enough photosensitizing drug was present to induce a (single-photon) photodynamic response, and that wide-beam PDT treatment causes extensive vasculature damage, as illustrated in Fig. 7. This was not an attempt to replicate the actual OPE-PDT treatment that occurs clinically, as the appropriate equipment (i.e., 680-nm laser) and Visudyne\textsuperscript{®} were not available. It is evident that the OPE-PDT effect occurred strongly in the specific artery into which Verteporfin was injected, yet there was little damage to the surrounding veins, probably due simply to the dilution of the photosensitizer. Stasis can be observed throughout this entire vessel, confirming the sensitivity of these CAM vessels to Verteporfin OPE-PDT. The supplementary video shows that the blood flow essentially stopped throughout the entire artery and stasis also occurred in some of the smaller (\(\sim 15\) to 30 \(\mu m\) diameter) directly adjacent neighboring veins, whereas the larger portions of the adjacent vein (\(>30 \, \mu m\) diameter) still maintained a constant blood flow. These results

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**Fig. 7** Representative images of control 1 (a) and (b) before and (c) and (d) immediately following treatment. (a) shows the vasculature before injection and irradiation of the entire CAM with white light. The arrows indicate the direction of blood flow in the arteries (white) and the veins (black). The white box indicates the area magnified in (b). Image (c) is after 5-min white light illumination. The artery is visibly occluded, as well as some of the small surrounding veins. Collapsed areas of the vessel are indicated by the white arrows, while the blood flow in the major veins is indicated by the thin black arrows. The white box in (c) illustrates the area magnified in (d).
are similar to those found by Lange et al. (2001),\textsuperscript{39} where Verteporfin was used to induce OPE-PDT in the CAM, although at much higher drug doses (2 to 20 \( \mu \text{g} \)) than we used here (0.5 \( \mu \text{g} \)), but much lower light dose (600 mW/cm\textsuperscript{2}). In that study, complete vascular collapse was evidenced by the disappearance of blood vessels in the treated area. There is confirmation that stasis is being observed in the current control experiments, since in all five trials [illustrated in Figs. 7(c) and 7(d) and supplementary videos] blood flow completely ceased in small vessels (diameter >30 \( \mu \text{m} \)), whereas there was still some movement in vessels greater than 30 \( \mu \text{m} \) diameter immediately after treatment. There is some evidence of the start of vascular constriction, where the blood vessels have become thinner [marked by the white arrows in Figs. 7(c) and 7(d)].

3.3.2 Control 2: does a cw laser beam cause the same blood vessel closure seen in two-photon excitation?

The purpose of control 2 was to check if focused cw laser irradiation at 780 nm causes the same blood vessel closure as seen in TPE with short-pulse irradiation at 780 nm, i.e., is there any significant contribution from one-photon excitation at this wavelength? Treatments were performed with otherwise similar parameters (50-\( \mu \)L injection, 780 nm, 38 mW for 5 min). However, the effective spot sizes were not identical (1.8 and 2.4 \( \mu \text{m} \) for pulsed and cw, respectively), so that the total fluences were slightly different. The peak irradiance with TPE was some 11 orders of magnitude higher than the average (constant) irradiance with cw treatment. Figure 8 shows that there was no blood vessel occlusion (or any other subocclusive responses) in any of the five cw controls, so that the responses in the TPE-experiments can be attributed to two-photon activation. It should be noted that the blood slows nominally throughout treatment due to the cooling of the embryo. The slight decrease in overall vessel diameter seen in Fig. 8(b) is discussed next.

3.3.3 Control 3: does a pulsed laser beam cause thermal damage to the blood vessels?

The objective with control 3 was to check for any thermal effects due to the pulse-laser 780-nm irradiation, in the absence of photosensitizer. As shown in Fig. 9 and the supplementary videos, no structural damage or effect on blood flow was observed. We have shown previously that TPE-PDT ablation of CAM blood vessels could be achieved with much shorter irradiation times, using higher irradiances and burning through the upper portions of the membrane prior to ablating the blood vessel itself,\textsuperscript{27} but in that case the thermal damage was quite visible. It has recently been reported by Karotki et al. (2006)\textsuperscript{25} that photomechanical damage in cultured endothelial cells appears at peak irradiances of about 8 \( \times 10^{11} \text{ W/cm}^2 \), while König, Rieman, and Fischer\textsuperscript{58} reported a threshold of 6.8 \( \times 10^{11} \text{ W/cm}^2 \). Our peak irradiance (3.7 \( \times 10^{11} \text{ W/cm}^2 \)) is below these threshold levels. Although the safety margin is not great, the lack of vessel closure in control 3 also suggests that there was no significant photomechanical damage.

We note that there was an overall decrease in vessel diameters (both treated and untreated) in Figs. 6(b) and 8(b). Initially, we attributed this to slight PDT effects from imaging in white light, or possibly a drug response. However, because we see the same effect in the CAM with no Verteporfin injected [Fig. 9(b)], we performed another control (see Appendix) that illustrates that the decrease in vessel diameter is due to air exposure of the CAM, causing slight drying of the membrane and slight cooling.

4 Conclusions

The proof-of-principle experiments presented here show that focal TPE-PDT with Verteporfin is capable of causing complete blood vessel occlusion in the chick CAM model. The experimental parameters, including wavelength, drug dose, and light dose, have not been completely optimized and this is part of ongoing work. The controls used show that the occlusion is not due to OPE-PDT or thermal damage from the pulsed laser source. Comparison to experimental OPE-PDT results found in the literature indicates that although we could not observe blood vessel closure over long periods of time, the treatment was successful. These results are encouraging for the development of TPE-PDT as a potential precise focal (e.g., feeder vessel targeted) treatment of choroidal neovascularization in wet AMD. Finally, these results suggest that with the development of better TPE-PDT sensitizers (higher TPE cross sections with good \( ^1 \text{O}_2 \) quantum yield), the treatment times and/or light doses can be decreased substantially.
5 Appendix

5.1 TPE Volume Measurement

A comparison between the TPE volume in water to a blood vessel within the CAM was investigated in order to estimate treatment volume broadening due to scatter of the laser light in the CAM tissue. A solution of amino PEG CdSe/ZnS quantum dots (Invitrogen, Mississauga, Canada) in water was prepared and aliquoted into a covered well slide. Using FCS, the average autocorrelation function (ACF) was collected (Fig. 10, average of two 30-s runs). The same quantum dot solution was then injected into the CAM blood vessel. During the injection, an air bubble was introduced into the needle (and subsequently the blood vessel) before and after the quantum dot solution. This prevents movement of blood, allowing the quantum dot solution to be inside the vessel without mixing with the blood. The ACF was then collected in two blood vessels, approximately 200 μm in diameter (Fig. 10). These are not the same diameter as those treated using TPE-PDT: the larger diameter and wall thickness of the vessels examined using FCS should lead to a greater broadening of the TPE volume than in the smaller (15 μm) PDT-treated vessels. Thus, the measurements represent the largest broadening of the TPE volume compared to water.

The average ACF from water was fitted using the ACF for a single diffusing species:\textsuperscript{59,60}

\[ G(\tau) = G(0) \left( 1 + \frac{8D\tau}{r_0} \right)^{-1/2} \left( 1 + \frac{8D\tau}{z_0} \right)^{-1/2}, \]  

where \( \tau \) is the lag time, \( D \) is the diffusion constant, \( r_0 \) is the laser beam radius at its focus, \( z_0 \) is the \( 1/e^2 \) radius in the \( z \) direction, and \( G(0) \) is

\[ G_{SB}(0) = \frac{1}{N_AV_{\text{eff}}(C)}, \]

where \( N_A \) is Avogadro’s number, \( V_{\text{eff}} \) is the effective TPE volume, and \( \langle C \rangle \) represents the time-averaged concentration of quantum dots. The previously determined radius (\( r = 1.8 \) μm) and height (\( z_0 = 5.7 \) μm) of the focal volume were held constant for a laser power of 100 mW. The concentration (\( C = 6.1 \pm 0.6 \times 10^{-10} \) M) and diffusion coefficient (\( D = 3.7 \pm 0.1 \times 10^{-11} \) m/s) extracted from the average water ACF fit were then used to determine the TPE volume parameters for the CAM blood vessels (\( r = 1.5 \pm 0.6 \) μm and \( z_0 = 7.9 \pm 0.8 \) μm). These results are summarized in Table 2. The change in \( G(0) \) notable in Fig. 10 is due to a slight dilution of the dot solution upon injection. However, the decays of the ACFs are indistinguishable, confirming no change in TPE focal volume.

The calculated TPE volumes for the water and blood vessel trials are the same within measurement error [\( V_{\text{f(well slide)}} = 37 \) fL and \( V_{f(vessel)} = 36 \) fL]. When the fitted values of the radius and height are considered, these parameters are similar for both water and the blood vessel. Identical or similar values of \( r \) and \( z_0 \) are not as pertinent as the focal volume, which has much lower variance. These results indicate that there is little to no TPE focal volume broadening when irradiating through the CAM tissue.

5.2 Control 4: Why Does the Overall Diameter of the CAM Vessels Decrease Over Time?

The majority of the vessels in the CAM showed a decrease in diameter over time during TPE-PDT treatment. The vessel constriction was observed for vessels with and without Verteporfin injection. This suggests that the mode for vessel shrink-
age is independent of the PDT procedure. In order to illustrate this observable trend, three eggs were prepared as described in Sec. 2.1 of the article. Each of the three eggs was left open and untreated (with laser light or photosensitizer) in a darkened room for a period of 15 min. As illustrated in Fig. 11(a), the images at time zero indicate a normal CAM, and the blurred look of the vessels are indicative of fast blood flow.

The second image [Fig. 11(b)], was taken approximately 10 min later, and it is evident that the vessels have slightly decreased in diameter and the blood flow has slowed (vessels are more clearly depicted). The videos that demonstrate this slowing of blood flow can be seen with the other supplementary videos at http://www.chem.ucalgary.ca/research/groups/dcram

5.3 Preliminary TPE-PDT Responses after 6 H
In an attempt to evaluate for long-term blood vessel occlusion, TPE-PDT, as described in Sec. 2.4, was performed on five eggs in a separate trial. Instead of imaging directly after treatment as described in Sec. 3.2, a low-intensity white light was used to verify that the laser spot had remained stationary during treatment, and the blood vessel of interest had indeed been occluded. The eggs were returned to an incubator in a darkened room to reduce exposure to light. Six hours after TPE-PDT, the treated blood vessels were observed. Of the five eggs in this trial only two were useable after 6 h incubation. At this time, it was shown that the treated vessel in one of the embryo’s CAM had partially reperfused, but blood stasis could still be observed downstream of the treatment spot (Fig. 12 and supplementary videos). In another CAM, there was also some evidence of reperfusion. Of the three remaining eggs, the original treated blood vessel could not be observed.

Of the three other eggs, in two the feeder vessel injected into had shut down, indicating either the injection site had bled out or that a wider PDT effect had occurred from stray white light. In the remaining egg, the CAM had dried out significantly, and although blood flow in the main vessel was observed, the smaller, treated vessel could not be visualized. It is noted that these data do not provide an appropriate set of statistics to make sweeping conclusions about the whether or not the blood vessel closure using the treatment parameters described herein is transient or permanent. It has been suggested that the permanent success of PDT (marked by blood flow stasis and permanent vascular damage) is reliant on the amount of circulating photosensitizer and light dose. It is anticipated that permanent vessel closure will be achieved when the drug and TPE light dose has been optimized.

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Table 2 Summary of parameters used to determine the focal volume broadening in the CAM, measured using amino PEG quantum dots.

<table>
<thead>
<tr>
<th></th>
<th>Radius, ( r ) (( \mu m ))</th>
<th>Height, ( z_0 ) (( \mu m ))</th>
<th>Diffusion Coefficient, ( D ) (m/s)</th>
<th>Concentration, ( C ) (M)</th>
<th>Focal Volume, ( V_f ) (fl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>1.8</td>
<td>5.7</td>
<td>( 3.7 (\pm 0.1) \times 10^{-11} )</td>
<td>( 6.1 (\pm 0.6) \times 10^{-12} )</td>
<td>37</td>
</tr>
<tr>
<td>CAM Vessel</td>
<td>1.5 (\pm 0.6)</td>
<td>7.9 (\pm 0.8)</td>
<td>( 3.7 \times 10^{-11} )</td>
<td>( 6.1 \times 10^{-12} )</td>
<td>36</td>
</tr>
</tbody>
</table>

Note: Errors in brackets indicate the errors determined from fit. The errors indicate the parameters allowed to float, while the parameters with no reported error were fixed.

Fig. 11 Exposure to air causes the vessels in the CAM to constrict slightly and for blood flow to slow. Image (A) was taken at time zero, and image (B) was taken approximately 10 min later, after the egg was left open in a darkened room.

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