Optical pacing of the embryonic heart

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Abstract

Light has been used to noninvasively alter the excitability of both neural and cardiac tissue. Recently, pulsed laser light has been shown to be capable of eliciting action potentials in peripheral nerves and in cultured cardiomyocytes. Here, we demonstrate for the first time optical pacing (OP) of an intact heart in vivo. Pulsed 1.875 μm infrared laser light was employed to lock the heart rate to the pulse frequency of the laser. A laser Doppler velocimetry (LDV) signal was used to verify the pacing. At low radiant exposures, embryonic quail hearts were reliably paced in vivo without detectable damage to the tissue, indicating that OP has great potential as a tool to study embryonic cardiac dynamics and development. In particular, OP can be utilized to control the heart rate, and thereby alter stresses and mechanically transduced signaling.

The avian embryo is an important model for studying the mechanisms driving normal development and the abnormalities leading to congenital defects. The embryonic heart is one of the first organs to develop and in avian models begins beating at around 40 hours of incubation. The progression of a single heart tube into a four-chambered heart (cardiac looping) is shaped by mechanical stimuli from the flowing blood and pumping heart which influence molecular and cellular responses that regulate development. Much remains...
unknown about the mechanisms of cardiac looping, in part because of a lack of suitable non-
destructive tools with which to study the minute (< 2 mm) organ. In the adult heart, the
exquisite control achievable using electrical pacing has been critical to advancing our
understanding of cardiac electrophysiology. Unfortunately, electrical pacing of the
embryonic heart is invasive and difficult to achieve consistently and without tissue damage.
A simple noninvasive technique to control heart rate would allow one to manipulate forces
applied to cells in early developing embryos, enabling a new class of experiments exploring
the roles of mechanotransduction and electrical activation.

It has been previously shown that ultraviolet and visible light can noninvasively increase the
excitability of neuronal and myocardial cells. In particular, Gimeno et al demonstrated
that exposure to visible light accelerated the heart rate in the early chick embryo. More
recently, Smith et al employed focused femtosecond pulses (780 nm) from a Ti:Sapphire
laser to induce paced contractions in individual and small groups of cardiomyocytes with
success rates up to 60%. Unfortunately, the high-powered pulses produce reactive oxygen
species that can damage the cell, and optically pacing an entire heart is not currently
possible with this method. In 2005, Wells et al demonstrated that pulsed infrared light could
reliably elicit compound action potentials in mammalian peripheral nerves in a one-to-one
fashion at radiant exposures well below the damage threshold. The stimulation
threshold paralleled the inverse water absorption curve with ~2 μm light producing the
highest damage-to-stimulation threshold ratio. Although the mechanism is not well
understood, it has been suggested that temperature gradients caused by infrared light
absorption open ion channels. To date, comparable pulsed infrared laser stimulation of
cardiomyocytes or cardiac tissue has not been demonstrated. Here, we describe the use of
pulsed infrared light to pace hearts of intact quail embryos, suggesting a promising new tool
for studying embryonic cardiac dynamics and development.

Figure 1 shows the experimental setup. A pulsed infrared diode laser (λ=1.875 μm;
Capella™, Lockheed Martin Aculight) centered at 1875 nm coupled light into a 400 μm core
multimode fiber. A micromanipulator positioned the fiber in close proximity to (500 μm),
but not in contact with the embryo (see Figure 1A), illuminating an area of approximately
0.3 mm² on the inflow region of the heart tube. The stimulation laser employed a red HeNe
laser coupled into the same fiber for aiming, which allowed precise positioning of the
stimulation pulses. The optical power of the HeNe laser light illuminating the sample was
low (0.9 mW) and the sample was illuminated only briefly (~10 s) in order to position the
fiber. The stimulation laser trigger signal was recorded to document the timing and duration
of the stimulation pulses. A laser Doppler velocimeter (LDV, Moor Instruments Ltd.) was
used to monitor the heartbeat of the embryos. The LDV probe was positioned approximately
1 mm from the heart with a micromanipulator (see Figure 1A). The LDV signal was only
intended for heartbeat detection; therefore, the LDV probe was not oriented identically with
every heart, so no significance should be drawn from the shape of the LDV signals. The
trigger from the laser was also directed to a white light emitting diode (LED), which flashed
on and off with the pulsing of the laser and was visible under a video microscope. The
output of the video microscope was recorded by a laptop computer for real-time guidance
and documentation (see movie in supplementary material).
Figure 2 demonstrates optical pacing of an embryonic heart using a pulsed laser. The spikes in the blue traces represent when the laser was emitting light, while the red traces show the heart rate of the embryo. Figure 2a presents a recording from a stage 17 quail embryo in a New culture paced at 2 Hz. The dashed box in 2a is expanded for a close up view in 2b. Clearly the laser pulse and heart rate are synchronized with each laser pulse eliciting a heartbeat. In Fig. 2a, the time interval between successive heartbeats before laser stimulation was 1.58±0.038 s (0.634 Hz) and the interval decreased to 0.4996±0.017 s (2.004 Hz) during pacing. After the stimulation pulses ceased and the heart rate stabilized the time interval between heartbeats increased to 1.44±0.042 s (0.693 Hz). At this point it is not known why the heart rate was slightly higher after pacing than before, but a similar increase occurred in most of the embryos tested. Although we cannot rule out damage, a physiological explanation for the post-pacing rate increase could be enhanced calcium ion influx during pacing. Figure 2c exhibits the heart beating in synchrony with the laser pulses as the frequency of the pulses is changed between 2 and 3 Hz.

The radiant exposure threshold required to pace a day 2 embryonic quail heart in a New culture was determined by subjecting each embryo to a different radiant exposure level and noting whether pacing had occurred. Successful pacing was defined as one to one stimulation, laser pulse to heartbeat, over a twenty second interval. 25 embryos were tested and the results were fit to a normal cumulative distribution function as shown in Figure 3. The threshold (50% probability point) was 0.81+/−0.01 J/cm$^2$ with a t-value of 79.9 and 95% confidence interval between 0.794 and 0.836 J/cm$^2$. The standard deviation of the distribution was 0.036+/−0.016 with a t-value of 2.32. In a shorter study we demonstrated that OP is feasible in embryos cultured on the yolk (see supplemental material). This method of embryo culture is compatible with longer viability of the embryos and would be useful for following long term effects of OP.

Several approaches were used to determine if optical pacing damaged the embryo. Standard transmission electron microscopy (TEM) was employed to examine the ultrastructural detail of cardiomyocytes in the inflow region of the heart tube where the laser was aimed. Figs. 4a-c represent a control embryo (same experimental protocol, but no light exposure), an embryo paced slightly above threshold (0.88 J/cm$^2$) and an embryo paced well above threshold (4.33 J/cm$^2$). In hearts paced well above threshold, the mitochondria are vacuolated and the nuclear envelope and rough endoplasmic reticulum are slightly expanded, whereas the ultrastructure of the cardiomyocytes of embryos paced slightly above threshold looked similar to that of the controls. In all three examples several regions of the heart tube were examined in detail to look for anomalies and no clear indications of cellular damage were identified in the embryo paced slightly above threshold. Some subtle abnormalities in the mitochondrial cristae, small spaces within the cristae, are present in the control and threshold embryo and are likely the result of necessary embryo manipulations during harvest prior to fixation. Other damage assays showed no structural damage to embryo heart cardiomyocytes paced at threshold levels (supplemental material). At this point we did not detect evidence of damage resulting from optical pacing, but further study is warranted.

The mechanism of the observed phenomenon is at this point unclear. However, given the laser parameters used (wavelength, pulse duration and peak power), some inferences can be
drawn. Gimeno et al used continuous visible light to increase the heart rate with maximum effect at 475 nm (photon energy = 2.61 eV). Proposed mechanisms included acetylcholine sensitivity to light and inhibition of ATPase. In a different study, Smith et al showed that a focused near-infrared femtosecond laser caused contraction in cultured neonatal rat cardiomyocytes. A window for this effect was found to occur between 15 and 30 mW average power for an 80 fs, 82 MHz pulse train of 780 nm, using 8 ms exposures applied periodically at 1 to 2 Hz. Mechanistically this effect was attributed to the laser-induced release of intracellular calcium which has been shown in various cell types for these laser parameters. In contrast, our study used near infrared pulses (λ = 1,870 nm) and relatively long (ms) pulses of continuous laser light. Thus the photon energy (0.66 eV) is insufficient to directly drive photochemistry as was the case in Gimeno’s work and the irradiance (W/cm²) is too low for multiphoton effects as in Smith’s work (irradiance from the fs pulses and diffraction limited spot size is roughly 8 orders of magnitude higher than we used). It is more likely that the mechanisms responsible for the effects seen in our study align with those described by Wells et al, who used identical laser parameters in the rat sciatic nerve and concluded that the laser-induced spatio-temporal temperature gradient was responsible for the induction of action potentials (APs) in excitable tissues. It has been shown that non-absorbing wavelengths with otherwise similar parameters do not result in APs in the peripheral nerve and pulse durations that violate the conditions of thermal confinement are similarly unable to induce APs. We have shown here and in previous studies that optical stimulation with these laser parameters is feasible without inducing thermal damage. The question of how the laser-induced thermal gradient ultimately results in the opening of ion channels remains unanswered although several hypotheses are currently under investigation.

Here, we have demonstrated a noninvasive method to pace embryonic hearts with pulsed infrared light. Compared to electrical pacing, OP does not require contact, has high spatial precision and avoids stimulation artifacts in electrode recordings. OP was consistently achieved with quail embryos ranging from stage 12 to stage 18 (looped heart stages prior to septation) in both New cultures and cultured on the yolk at radiant exposures well below levels at which damage was apparent. In the future we will make use of OP together with advanced imaging to study mechanotransduction (i.e. shear stress on the endocardium) by controlling the heart rate of the embryo, thereby manipulating mechanical stress. OP will not only enable a new class of experiments in developmental cardiology, but also may become a useful tool for investigating cardiac electrophysiology, single-cell (cardiomyocyte) dynamics, and cardiac tissue engineering. Furthermore, OP may potentially be capable of pacing the adult heart, which could lead to clinical applications.

Methods

Egg preparation

Fertilized quail (Coturnix conturnix) eggs were removed between 48–60 hours post-fertilization from a humidified, forced draft incubator (38 °F). Embryos were removed from the egg and either cultured on the yolk in a sterilized 3.5 cm diameter Petri dish or cultured using the New culture in which the embryo is inverted on an egg-agar substrate. The New culture fully exposes the heart, which facilitates interventions, but decreases lifespan to

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a few days, whereas embryos cultured on the yolk can survive through gestation. Both techniques are routinely employed in the field. Figure 1A shows a healthy 53-hour embryo in the New culture. For pacing experiments, embryos were placed upon a temperature-controlled heating plate (ATC 1000, WPI), which maintained the temperature at 37 °C.

**Radiant exposure calculations**

Radiant exposures (J/cm$^2$) were calculated by dividing the pulse energies by the laser spot size. Pulse energies were measured using a pyroelectric energy meter (PE50BB, Ophir) and spot size was computed using the numerical aperture, fiber diameter and fiber distance from the tissue. Laser light was delivered to the tissue via a 400+/−8 μm - diameter flat-polished multi-mode optical fiber (Ocean Optics) having a numerical aperture of 0.22 +/−0.02. The fiber-to-heart distance was held constant at approximately 500 μm for each trial. The tissue was assumed to be perpendicular to the fiber. While the fiber was oriented at an angle to the membrane of approximately 37 degrees from perpendicular, the curvature of the membrane covering the heart was assumed to offset the angle of the fiber sufficiently to make these effects negligible.

**Threshold measurements**

25 embryos in New cultures were tested to determine the radiant exposure level needed to induce optical pacing. To quickly find to the precise location for optimal stimulation, the laser was turned to a level slightly above threshold and the position of the fiber was optimized, which normally took less than ten seconds. The optical fiber was positioned 500 μm from the embryo. Because the LDV fiber took a significant time to accurately place, for this experiment we relied on the LED flashes and heart motion in the microscope video to determine successful pacing. After a brief waiting period (~ 1 min) with the laser off, the radiant exposure was adjusted and the laser was turned on for 20–30 seconds. Embryos were then either fixed for further experiments or discarded. For each embryo we determined whether there was successful pacing (1) or failed pacing (0) and plotted the results in figure 3. Some embryos took two or three beats to synchronize to the laser pulses, but were counted as successful pacing if the heart rate was consistent after the first couple beats. Conversely, if the heart followed the laser pulses, but skipped more than one beat it was scored unsuccessful. The difference between successful pacing and unsuccessful pacing was easy to distinguish by closely observing the videos.

**TEM preparation**

The torso of the embryo was excised and immediately fixed by immersion in the triple aldehyde-DMSO mixture of Kalt and Tandler$^{22}$. After rinsing, the tissues were postfixed in ferrocyanide-reduced osmium tetroxide$^{23}$. After again rinsing, they were soaked overnight in acidified uranyl acetate$^{24}$. Thin sections were sequentially stained with acidified uranyl acetate$^{24}$ followed by Sato’s triple lead stain as modified by Hanaichi et al.$^{25}$ and examined in a JEOL 1200 electron microscope.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.
Acknowledgments

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Figure 1. OP setup

**a**, Photograph of a 53 hour quail embryo in the New culture. **b**, Block diagram of the setup of the optical pacing experiment. Embryos in the Petri dish under the microscope were stimulated by a pulsed infrared laser, while a laser Doppler velocimeter probe measured blood flow. The LDV output and trigger pulse from the laser were recorded to verify pacing. Also, the trigger pulse from the laser activated a white light LED which was observable in the video. The video output from the microscope camera was recorded at video rate (29.97 fps) by a laptop computer.
Figure 2. Pacing of the embryonic quail heart

(a,b) The trigger pulse (blue) from the pulsed laser is superimposed on the LDV recordings (red) of heart rate. (a) presents a recording from a stage 17 (59 hour) quail embryo in a New culture paced at 2 Hz. The laser pulse duration was 1 ms and the radiant exposure was 0.92 J/cm² per pulse. The laser pulses were directed toward the inflow region of the heart tube. The laser pulses were turned on and off several times to demonstrate the robustness of optical pacing. The heart rate increased from 0.634 to 2 Hz when the stimulation laser pulses were started and decreased to 0.693 Hz by the end of the trace in 2a. (b) The dashed box in 2a is expanded for a close up view in 2b. Clearly the trigger pulse and LDV signal were synchronized with each laser pulse eliciting a heartbeat. (c) presents a recording from a stage 14 (53 hour) quail embryo in a New culture. The laser pulse duration was 2 ms and the radiant exposure was 0.84 mJ/cm² per pulse. The frequency of the laser pulses were varied from 2 Hz to 3 Hz and back to 2 Hz to demonstrate the ability of the embryo heart to follow the pulse frequency. Laser stimulation (blue) and heart rate (red) traces were calculated from the video by plotting the pixel intensity of LED flashes triggered by the laser and intensity at the edge of the heart wall.
Figure 3. Threshold measurement
25 embryos in New cultures were exposed to varying radiant exposures and the results were plotted in terms of pacing probability (successful pacing = 1, unsuccessful pacing = 0). The data were fit to a normal cumulative distribution function (blue line). The threshold (50% probability point) was 0.81+/−0.01 J/cm$^2$ with a t-value of 79.9 and 95% confidence interval between 0.794 and 0.836 J/cm$^2$. The standard deviation of the distribution was 0.036+/−0.016 with a t-value of 2.32.
Figure 4. Transmission electron microscopy (TEM) after the OP procedure

a-c, The TEM images show a typical cardiomyocyte in the inflow (sinoatrial) region of the heart tube of three different quail embryos, where laser light was directed. Each embryo was between 52 to 58 hours of development. The torso of each embryo was excised and fixed immediately after OP. a, Control embryo. The embryo was exposed to the OP experimental procedure, except that the pacing laser was not turned on during the experiment. b, Embryo paced slightly above threshold. The embryo heart was paced for approximately 20 seconds with 2 ms pulses (2.64 ml/pulse) at 2 Hz. c, Embryo paced well above threshold. The embryo heart was paced for approximately 20 seconds with 4 ms pulses (13 ml/pulse) at 2 Hz. The micrograph image was taken from a border region separating severely damaged tissue (partially ablated) from apparently healthy tissue. The mitochondria are vacuolated and the nuclear envelope is swollen, as are elements of the rough ER (large arrow). The cells and mitochondria in 4a and b have a similar appearance and show no signs of damage from the pulsed laser light. Some subtle abnormalities in the cristae present in cells from both threshold paced and unpaced control hearts are due to the unavoidable delay in excising the embryonic torso. N -nucleus, M - mitochondria.