Reduced Heart Rate and Cardiac Output Differentially Affect Angiogenesis, Growth, and Development in Early Chicken Embryos (Gallus domesticus)

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ABSTRACT

An increase in both vascular circumferential tension and shear stress in the developing vasculature of the chicken embryo has been hypothesized to stimulate angiogenesis in the developing peripheral circulation chorioallantoic membrane (CAM). To test this hypothesis, angiogenesis in the CAM, development, and growth were measured in the early chicken embryo, following acute and chronic topical application of the purely bradycardic drug ZD7288. At hour 56, ZD7288 reduced heart rate ($f_H$) by $\sim$30% but had no significant effect on stroke volume ($-0.19 \pm 0.2 \text{ mL}$), collectively resulting in a significant fall in cardiac output (CO) from $27 \pm 3$ to $18 \pm 2 \text{ mL min}^{-1}$. Mean $f_H$ at 72 h of development was similarly significantly lowered by acute ZD7288 treatment (250 $\mu\text{M}$) to $128 \pm 0.3 \text{ beats min}^{-1}$, compared with $174.5 \pm 0.3$ and $174.7 \pm 0.8 \text{ beats min}^{-1}$ in control and Pannett-Compton (P-C) saline–treated embryos, respectively. Chronic dosing with ZD7288—and the attendant decreases in $f_H$ and CO—did not change eye diameter or cervical flexion (key indicators of development rate) at 120 h but significantly reduced overall growth (wet and dry body mass decreased by 20%). CAM vessel density index (reflecting an increase in both vascular circumferential tension and shear stress) measured 200–400 $\mu\text{m}$ from the umbilical stalk was not altered, but ZD7288 reduced vessel numbers—and therefore vessel density—by 13%–16% more distally (500–600 $\mu\text{m}$ from umbilical stalk) in the CAM. In the ZD7288-treated embryos, a decrease in vessel length was found within the second branch order ($\sim$300–400 $\mu\text{m}$ from the umbilical stock), while a decrease in vessel diameter was found closer to the umbilical stock, beginning in the first branch order ($\sim$200–300 $\mu\text{m}$). Paradoxically, chronic application of P-C saline also reduced peripheral CAM vessel density index at 500 and 600 $\mu\text{m}$ by 13% and 7%, respectively, likely from washout of local angiogenic factors. In summary, decreased $f_H$ with reduced CO did not slow development rate but reduced embryonic growth rate and angiogenesis in the CAM periphery. This study demonstrates for the first time that different processes in the ontogeny of the early vertebrate embryo (i.e., hypertrophic growth vs. development) have differential sensitivities to altered convective blood flow.

Introduction

Angiogenesis results from a complex combination of genetically directed morphogenesis and the influence of local environmental factors, both biochemical and mechanical (e.g., Jones et al. 2006; Lee et al. 2009; Adams and Eichmann 2010; Buschmann et al. 2010; Kaunas et al. 2011; Knudsen and Kleinstreuer 2011; Heinke et al. 2012; Burggren 2013). Chief among the mechanical influences are both blood pressure and flow (Isogai et al. 2003; le Noble et al. 2008). Blood pressure creates both an absolute transmural pressure as well as circumferential stretch (tension) in compliant vessels through the fluctuating blood pressure associated with the cardiac cycle. Blood flow stimulates angiogenesis by generating a shear stress on the inner surface of the vesel that parallels the direction of the blood flow. Endothelial cells comprising the inner layer of the vasculature—and those cells lining the sprouting blood vessel tips—respond to such stressors by proliferating under the influence of vascular endothelial growth factor (VEGF) and other paracrine secretions. New vessels and the eventual throughflow of blood occur as a result (see reviews in Groenendijk et al. 2007; le Noble et al. 2008; Egginton 2011). Arterial (as opposed to venous) identity then arises from the greater pulsatility and shear stresses that eventually emerge in the arterial vasculature (Buschmann et al. 2010). Angiogenesis is crucial to the completion of the capillary network and the connecting of developing arteries and veins. Indeed, failure of angiogenesis in early embryonic development has long been known to lead to rapid embryonic death, as evident from observations of VEGF knockout mice that fail to develop a complete microvasculature (Car- mellet et al. 1996; Ferrara et al. 1996).

While well documented in mature vertebrates, the dependence of early embryonic angiogenesis on the onset of heart beat, blood pressure generation, and blood flow has not been
well established in the embryo. A complicating factor is that blood pressure and flow are not an absolute requirement for organogenesis and growth in the early vertebrate embryo, because development in early embryos will continue for some time when convective blood flow and mass transport are experimentally eliminated (Mellish et al. 1994; Pelster and Burggren 1996; Burggren et al. 2000, 2004). This independence of development and growth from convective blood flow occurs because diffusion provides adequate mass transport during early embryo development. Eventually, a combination of escalating gas exchange and nutrient demands, coupled with increasing diffusion distances of the growing embryo, necessitates the development of an active internal convective blood flow (for review, see Burggren 2004).

Onset of blood pressure and flow generation—a metabolically expensive process—occurs well in advance of the embryo’s absolute need for material transport, a phenomenon that has been termed “prosynchronotropy” (Territo and Burggren 1998). Unanswered is the question of why cardiac contraction and blood convection occur earlier than actually required for bulk transport. An as yet untested hypothesis is that the initial beating of the heart is not for material transport but rather to provide the pulsatile pressure and flow—and the attendant circumferential stretch and shear stresses—in the peripheral circulation as a necessary stimulant for angiogenesis in the early embryo (Burggren 2004). The hypothesis of prosynchronotropy has recently been refined to that of synangiotropism, namely, that blood pressure and flow appear just in time for the need for angiogenesis rather than bulk transport (Burggren 2013). To test this hypothesis, this study used the purely bradycardic drug ZD7288 to induce chronic bradycardia in the chicken embryo from 72 to 120 h of development. Bradycardia increases pressure in the arterial vascular beds as a result of the lengthened period of diastolic pressure runoff, thus potentially increasing vascular circumferential stress. If bradycardia induces a decreased cardiac output (CO), it will additionally reduce shear stress on the vascular endothelial lining.

In early stages of avian development, bradycardia is not accompanied by a compensatory increase in stroke volume, and so a reduction in CO accompanies heart rate reduction (Bowers et al. 1996; W. Burggren and M. Yamada-Fisher, unpublished manuscript). Using experimentally induced bradycardia as a tool for lower CO, we have also assessed rates of embryonic development, growth, and angiogenesis in both control populations and those chronically exposed to either Pannett–Compton (P-C) saline or ZD7288.

Material and Methods

Source of Experimental Animals

Fertilized eggs of the white leghorn chicken (Gallus domesticus L.) were obtained from Texas A&M University (College Station, TX) and shipped to the University of North Texas (Denton, TX). Eggs were incubated in a Hova-Bator Styrofoam incubator at a constant temperature of 37.5° ± 0.5°C and 55%–60% humidity. After 48 h of incubation, the eggs were removed from the incubator and transferred into a shell-less culture system.

Shell-Less Culture

To facilitate drug delivery and measurements, shell-less embryo cultures were prepared after the method of Hamamichi and Nishigori (2001). This method enables growth of the embryo for up to 5 d, while allowing full access to the embryo and its vasculature for manipulation and observation (fig. 1). To prepare a shell-less culture, each incubated egg was removed from the incubator and positioned horizontally at room temperature on an egg crate for 10 min to allow the embryo to be positioned correctly for harvesting. The egg surface was then lightly sprayed with 70% ethanol to reduce microbial contamination from the egg surface and then allowed to air-dry. Each egg was cracked, and the whole egg contents were transferred into an autoclaved Kimax crystallizing dish (60 mm × 35 mm) under aseptic conditions. Any air bubbles surrounding the yolk were removed with a sterile Kimwipe. The dish was covered with clear polyethylene film secured with an elastic rubber band. Cultured embryos were then numbered and reincubated at 37.5° ± 0.5°C for further development and use. Care was taken to include only cultures with intact yolks that had the blastodisc positioned on the uppermost side of the yolk.

Experimental Design and Embryo Populations

Three populations of shell-less culture embryos were created. The control embryos were untreated in any way and were simply grown in the culture conditions described above. P-C saline–treated embryos grown in culture received topical application of P-C saline (2.2 mol L⁻¹ NaCl, 0.21 mol L⁻¹ KCl, pH = 7; after Stern and Holland 1993) to determine any mechanical or physiological effects of fluid placement on the embryo. The Po₂ of the culture medium, which was air saturated, was ~150 mmHg. Finally, a group of embryos, grown in culture as described above, received topical application of the bradycardic agent ZD7288. This drug blocks the Iₒ cardiac pacemaker channels in the heart pacemaker, reducing heart rate without creating an inotropic cardiac effect (Yusuf and Camm 2003; Luo et al. 2006).

Drug Solutions and Delivery Methods

ZD7288 was obtained from Tocris Bioscience (Ellisville, MO). A stock solution of 10 mg ZD7288 in 1 mL of autoclaved distilled water was made and stored at −20°C. Subsequent solutions of specific concentrations were made using P-C saline (Pannett and Compton 1924), which was originally developed as a superior medium to standard chick Ringer solutions for growing blastodiscs in vitro.

The two-dimensional nature of shell-less culture was effective in allowing an even distribution of solution and drug to the embryo and its extraembryonic tissues. Preliminary observations using P-C saline solution tinted with Evans blue dye showed that
flushed each embryo’s surface with P-C saline or ZD7288 so-
embryo’s heart. This chronic drug delivery system slowly
then was placed through the cellophane wrap directly over each
glued into the tip of a 50-mm-long PE 240 polyethylene tube
simultaneously. The tip of a 10-mm-long PE 100 tubing was
containing ZD7288 or P-C saline were secured on the syringe

topoical application of a 5-μL drop of solution directly above the
embryo’s body wall resulted in broad, even distribution of the
solution across both the embryo and the surrounding chorio-
allantoic membrane (CAM). At the same time, excess solution
ran off the surface of the embryo and collected remotely at the
far edges of the culture dish, thus preventing accumulation of
fluid over the embryo itself (fig. 1).

In acute experiments, the dish containing a cultured embryo
was briefly removed from the incubator. A single predetermined
dose of ZD7288 solution was topically applied from a sterilized
transfer pipette inserted through the polyethylene wrap cover.
Acute experiments were carried out on embryos of the follow-
ing ages: 55–56, 72, 96, and 120 h.

For chronic dosing, three dishes with 48-h embryos were
placed in a Lyon Reptille incubator (model RL182) set at
37° ± 0.5°C and 55%-60% humidity (fig. 1). A syringe pump
(model 352; Sage Instruments, Boston, MA) was also placed
within the incubator to ensure precise temperature control of
drug or saline delivery. Three Bristoline 10-cm³ glass syringes
containing ZD7288 or P-C saline were secured on the syringe
pump and used for continuous drug delivery to three embryos
simultaneously. The tip of a 10-mm-long PE 100 tubing was
 glued into the tip of a 50-mm-long PE 240 polyethylene tube
and attached by a plastic on/off connector to each syringe and
then was placed through the cellophane wrap directly over each
embryo’s heart. This chronic drug delivery system slowly
flushed each embryo’s surface with P-C saline or ZD7288 solu-
tion at a rate of 200 μL h⁻¹ for a 24-h exposure period, for
a total of 4.80 mL of delivered solution by age 72 h. Control
embryos were cultured within the incubator at the same tem-
perature and for the same time period but did not receive P-
C saline or ZD7288.

Treatment and Heart Measurement Protocols

Each of the lids of the incubators, in which the acute pop-
ulations were housed, had a transparent section through which
the beating heart of the cultured embryo could be visually
observed. Heart rate (fH) in embryos from P-C saline–treated
and ZD7288 groups was measured by counting the number of
heart beats in two successive 15-s intervals separated by a 10-
s period and transformed to beats per minute. After this initial
heart rate measurement, embryos in their culture dish were
then briefly removed from the incubator. In one acute popu-
ation, a single 5-μL aliquot of P-C saline was applied topically
to the embryo. In the experimental population, bradycardia
was induced by a single 5-μL aliquot of one of the range of
concentrations (30, 250, and 1,000 μM) of topically applied
ZD7288. The control population was left untreated. Embryos
were then immediately placed back into the incubator. Heart
rate continued to be taken every 10 min after administration
of the drug for ~10 h. This protocol was carried out at 56, 72,
96, and 120 h of development.

For chronic treatments, embryos were dosed with 7.5, 10,
15, 20, and 30 μM ZD7288 for 24 h, starting at 48 h and
continuing through 72 h of development. In preliminary ex-
periments used to refine the protocol, chronic treatment of 30
μM ZD7288 at a rate of 200 μL h⁻¹ (about 40 5-μL drops h⁻¹)
achieved the same degree of bradycardia as the maximum acute
bradycardic response produced by a single drop of 5 μL of 500
μM ZD7288. The subsequent maximum chronic concentration
used was thus adapted from these results.

Blood Velocity and Flow Measurements

Blood flow measurements were made in shell-less culture em-
bryos from 56 to 60 h of development. This stage of de-
velopment is ~10 h younger than the stage at which some of the
other measurements were made but was also the stage yielding
the most precise signals with the lowest variation. Dorsal aortic
blood velocity was measured with a 20-MHz pulsed Doppler
flowmeter and a 0.5-mm-diameter piezoelectric crystal (Iowa
Doppler Products, Iowa City, IA) positioned over the dorsal
aorta at a 45° angle. Analog signals were logged onto an eight-
channel PowerLab III data acquisition system (ADInstruments,
Colorado Springs, CO). Mean aortic velocity and fH were cal-
culated by the data acquisition software. The diameter of the
dorsal aorta was measured with a filar micrometer eyepiece.
From these measures, stroke volume (μL) and CO (μL/min)
were calculated. (Note that dorsal aortic blood flow measured
at this point does not include CO going to the developing
embryonic head [Hu and Clark 1989].)

CAM Vessel Density Index Measurement

CAM vessel density index protocols were created using previ-
sously published conventions (Strick et al. 1991; Höper and Jahn
1995; Corona et al. 1999). Images of each embryo were taken
at × 3.5 using a Javelin camera mounted on a Leica WILD
M3Z microscope. To measure the CAM vessel density index of
the extraembryonic vasculature, Image Pro software was used
to draw five concentric circles (200, 300, 400, 500, and 600
μm) centered directly over the umbilical stock of the embryo.
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Figure 2. Technique for measuring chorioallantoic membrane (CAM) vessel density index using concentric circles at 100-μm intervals, all centering around the root of the umbilical stock. CAM vessel density index was calculated by counting the number of clearly visible vessels that intersected each circle around its entire circumference. A color version of this figure is available in the online edition of *Physiological and Biochemical Zoology*.

Vessel Branch Order, Length, and Diameter Measurements

Images taken to assess the CAM vessel density index were also used to measure number of vessels per branch order, vessel length, and vessel diameter. Branch order was quantified as follows: first order being the main umbilical vessel extending from the embryo on both sides, second order being the vessels branching off the first order vessel, and so forth. Both left and right sides of the embryo were measured for statistical comparison.

The number of vessels per branch order was measured to the fourth branch order, beyond which visual quantification became imprecise. Vessel length was measured from the beginning of the vessel to the end of the vessel, where it branches off into the next order. Measurements were repeated and then averaged. Vessel diameter was measured at 25% and 75% of the length of the vessel to ensure that representative measurements of the vessel were obtained. Both vessel length and vessel diameter were measured out to the third branch order.

Eye Diameter/Cervical Flexure Measurement

Eye diameter (μm) and cervical flexure (°) are conventional measures of development during early developmental stages. In the current experiments, these variables were assessed with Image-Pro software (ver. 4.1). Calibrated images taken with a dissecting microscope and camera at a total magnification of ×16 were used for measurement.

Embryo Dry and Wet Mass Measurements

After the vasculature measurements, embryos were removed from culture to acquire wet and dry mass. The embryo was carefully cut away from the surrounding vessels, and extraembryonic membranes were pulled off of the embryo with forceps, blotted with filter paper to absorb excess liquid, and placed into a preweighed plastic dish for wet mass measurement. The dish containing the embryo was then placed into a drying oven at 80°C for 48 h and then weighed to obtain dry mass.

Statistical Analyses

Effects of development and chronic exposure to ZD7288 on heart rate, eye diameter, cervical flexure, and wet and dry mass were assessed using one-way ANOVA, followed by Tukey post hoc tests to determine pairwise differences. Two-way ANOVA was used to assess statistical differences in CAM vessel density index caused by chronic exposure to ZD7288 and acute heart rate caused by acute exposure to ZD7288 and development, respectively. Two-way ANOVA was also used to assess the difference between the number of vessels per order, vessel length, and vessel diameter. Where significant differences were found, Tukey post hoc tests were used for pairwise comparisons. We used t-tests to determine whether there was a difference between the number of vessels on the right and the left side of embryo. All variables are represented as mean ± 1 SE, with α = 0.05. All statistical analyses were performed using SigmaStat and SigmaPlot software.

Results

Heart Rate and Blood Flow in Control and Saline-Treated Populations

Mean $f_0$ at 56 h of development in control embryos was ∼130–140 beats min⁻¹, rising to ∼170 beats min⁻¹ by 72 h of development (fig. 3). As expected from prior studies, $f_0$ began to plateau at ∼220 beats min⁻¹ at 96 h and increased only slightly to 230 beats min⁻¹ at 120 h (fig. 3).

Blood velocity in the dorsal aorta at 56 h ranged from 18 to 22 mm s⁻¹ at peak systolic ejection, falling back to 0 during diastole (fig. 4), indicating little or no windkessel effect that would otherwise maintain substantial aortic blood flow during diastole.

Stroke volume at 56 h in saline-treated embryos was approximately 0.19 μL, leading to a CO of ∼27 μL min⁻¹ at $f_0$ of 140 beats min⁻¹ in control populations (fig. 5). Acute topical
Acute Cardiovascular Effects of ZD7288

Initial experiments were conducted to determine the effective dose ranges for ZD7288-induced bradycardia in ~56-h chicken embryos. $f_{Ht}$, expressed as a percentage of control values, decreased significantly with as little as 30 μM (fig. 6). The most pronounced effect occurred at 500 μM, with no further significant drop in $f_{Ht}$ at concentrations as high as 1,000 μM. Subsequent experiments were thus limited to doses in the 30–500 μM range.

The time course of ZD7288 effects was then established, along with the effects of the carrier, P-C saline. The dose- and time-dependent effects of acute exposure to three different concentrations of ZD7288 and P-C saline in 72-h embryos are indicated in figure 7. $f_{Ht}$ of embryos acutely treated with ZD7288 typically decreased significantly ($P<0.01$) within 30 min of application at doses 30, 250, and 500 μM, with the largest effects occurring at 1–2 h following exposure but lasting at least 4–5 h (fig. 7).

$f_{Ht}$ at 56 and 72 h decreased from ~140 to slightly more than 100 beats min$^{-1}$ and from 170 to ~140 beats min$^{-1}$, respectively, following acute topical application of ZD7288 (figs. 5A, 7). Stroke volume at 56 h did not change significantly ($P>0.05$) following an acute application of ZD7288 and the attendant longer period of diastolic filling accompanying bradycardia (fig. 5B). As a consequence of a fall in $f_{Ht}$ without a compensatory increase in stroke volume in embryos acutely exposed to ZD7288, CO decreased significantly ($P<0.01$) from ~27 to 18 μL min$^{-1}$ (fig. 5C).

Mean $f_{Ht}$ in control and P-C saline–treated (acute) embryos at 72 h of development were statistically identical at 175 ± 0.3 and 175 ± 1 beats min$^{-1}$ ($P>0.05$), respectively (fig. 7). Acute dosing with 5 μL of all three concentrations of ZD7288 produced a significant decrease from control $f_{Ht}$ values within 15 min of drug delivery (fig. 7). Maximum effects were evident within 45–75 min of treatment. Thirty microns of ZD7288 produced an $f_{Ht}$ of ~102 beats min$^{-1}$ that was significantly lower than the control and P-C saline groups ($P<0.001$) but significantly higher than the more profoundly reduced $f_{Ht}$ of the 250 and 500 μM ZD7288 groups ($P<0.001$), which were not significantly different from each other (one-way ANOVA, $P=0.830$; fig. 7).

Acute ZD7288 treatment (500 μM) continued to create a significant ($P<0.001$) bradycardia at both 96 and 120 h of development. This effect was unrelated to delivery vehicle, since treatment with P-C saline caused no significant change in $f_{Ht}$ from control at these stages.

Hemodynamic Effects of Chronic Treatment with ZD7288

Heart rates at 72 h of development in control embryos (169 ± 3 beats min$^{-1}$) and those chronically treated with P-C saline from 48 to 72 h (172 ± 3 beats min$^{-1}$) were not significantly different ($P>0.05$). Chronic treatment with 7.5, 10, 15, 20, and 30 μM ZD7288 all created a bradycardia that was significantly different from both the control and the P-C saline groups ($P<0.001$) yet not significantly different from each other ($P>0.05$; fig. 8). Chronic dosing of ZD7288 decreased $f_{Ht}$ by ~33% from 171 ± 3 beats min$^{-1}$ to an average for all ZD7288 doses of 116 ± 3 beats min$^{-1}$ at 72 h of development. The higher doses were accompanied by increased embryo mor-
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Figure 5. Acute effects of ZD7288 on heart rate (A), stroke volume (B), and cardiac output (C) in 56-h embryos. Note that acute ZD7288 treatment reduces heart rate, but there is no compensatory increase in stroke volume, resulting in a net decrease in cardiac output. Means ± 1 SE are plotted; n = 6 for each group, P-C, Pannett-Compton.

Figure 6. Dose response curve showing percent decrease of heart rate for acute dosing of different concentrations of ZD7288 (ranging from 30 to 1,000 μM) on 56-h embryos. Numbers indicate n values for each measurement.

Vessel Number per Branch Order and ZD7288 Treatment

To quantify the number of blood vessels for each branching order, both sides of the embryos were measured. In the control embryos, there was no significant difference between the number of vessels on the left and the right side of the embryos (P = 0.934). Therefore, the left side of the embryo was used for treatment comparisons and statistical analysis. Vessel num-

CAM Vessel Density Index and ZD7288 Treatment

There was no significant effect of the various ZD7288 concentrations on f/1 in chronic experiments (i.e., they were statistically identical populations). Consequently, the CAM vessel density index data for all embryos chronically dosed with the various doses of ZD7288 were merged together to form a total of three groups for statistical analysis: control, P-C saline–treated, and ZD7288-treated groups. CAM vessel density index of control embryos at a distance of 200, 300, and 400 μm from the umbilical stalk was 3.2 ± 0.1, 6.2 ± 0.33, and 9.5 ± 0.44 intersects μm⁻¹, respectively (fig. 9).

There was no significant difference in CAM vessel density index between control and treatment groups (P > 0.05) proximally at 200, 300, and 400 μm from the embryo umbilical stalk (fig. 9). However, chronic bradycardia from the application of ZD7288 induced a significant decrease in CAM vessel density index at a distance of 500 μm from the umbilical stalk, down to 12.2 ± 400 intersects μm⁻¹ from a control value of 14.1 ± 700 intersects μm⁻¹ (P < 0.001). Surprisingly, chronic application of P-C saline also induced a significant decrease in CAM vessel density index at 500 μm, down to 12.5 ± 400 intersects μm⁻¹ compared with the control, 14.3 ± 700 intersects μm⁻¹ (P = 0.020).

At a distance of 600 μm from the umbilical stalk, there was also a significant decrease (P < 0.001) in the CAM vessel density index of both the ZD7288 (16.6 ± 600 intersects μm⁻¹) and the P-C saline (18.5 ± 600 intersects μm⁻¹) embryos as compared with the control group (19.7 ± 700 intersects μm⁻¹; fig. 9).
Figure 7. Heart rate at 72 h of development as a function of control, Pannett-Compton (P-C) saline treatment, and acute ZD7288 dosing at three concentrations. Each point is the mean ± SE of a 15-min period. Letters indicate statistical difference between groups, starting at 20 min after dosing; n values are shown for each condition.

Figure 8. Heart rate at 72 h of development following 24 h of chronic ZD7288 dosing. Horizontal lines indicate statistically identical groups. Means ± 1 SE are plotted; n values are in parentheses. P-C, Pannett-Compton.

Vessel Length and Diameter and ZD7288 Treatment

Vessel length in the first branch order was not significantly different within treatment groups. In the second order, however, ZD7288-treated embryos showed a significant 40% decrease from both the control and the P-C saline embryos (fig. 10B). In the third branch order, the P-C saline- and ZD7288-treated embryos both showed a significant 28% decrease in vessel length from control embryos (two-way ANOVA, Tukey pairwise comparison, P < 0.05; fig. 10B).

Not surprisingly, as branch order increased, vessel diameter decreased (fig. 10C). Within the first and second branch orders, there was a significant 19%–14% decrease, respectively, of vessel diameter in ZD7288-treated embryos from P-C saline and control embryos (two-way ANOVA, Tukey pairwise comparison, P < 0.05). Vessel diameter in the third branch order was not significantly different between any of the groups (fig. 10C).

Developmental Effects of Chronic ZD7288 Treatment

Eye diameter at 72 h of development was not significantly affected (ANOVA, P = 0.061) by treatment. Values for eye diameter were 487 ± 20 μm for control (n = 34), 555 ± 20 μm for P-C saline (n = 15), and 512 ± 10 μm for the values of ZD7288 spanning the dose range from 7.5 to 30 μM (n = 40). Similarly, no significant difference was found in a second indicator of developmental progress—cervical flexure—between the three groups (P = 0.126), with values of 125.5° ± 5.5° (n = 13), 130.5° ± 7.6° (n = 13), and 127.8° ± 4.9° (n = 14) in control, P-C saline-, and ZD7288-dosed embryos, respectively.

Body Mass Effects of Chronic ZD7288 Treatment

Mass was used an indicator of growth, independent of development, which was assessed as described above with eye diameter and cervical flexure. Wet mass of the control (14.1 ± 0.9 mg) and P-C saline–treated (14.8 ± 0.8 mg) embryos was not significantly different (P = 0.131); consequently, these populations were grouped together in subsequent analyses. Wet masses of embryos dosed with 7.5, 10, 15, 20, and 30 μM of ZD7288 were also not significantly different (P = 0.053) from each other (fig. 11A). Collectively, wet mass (11.6 ± 0.8 mg) of the pooled ZD7288 treatment group was 22% smaller than the P-C saline–treated group (14.8 ± 0.5 mg; P < 0.001).

Dry mass of the control (1.2 ± 0.1 mg) and P-C saline–treated (1.0 ± 0.3 mg) embryos was not significantly different (P = 0.335); consequently, these populations were grouped together in subsequent analyses. Dry masses of embryos dosed with 7.5, 10, 15, 20, and 30 μM of ZD7288 were also not significantly different from each other (P = 0.175; fig. 11B). Collectively, dry mass (1.2 ± 0.4 mg) of the pooled ZD7288 treatment group was 33% smaller than the control group (0.8 ± 0.04 mg; P < 0.001), mirroring the changes described for wet mass.
Critique of ZD7288 and Its Application

ZD7288, which blocks the \( I_f \) cardiac pacemaker channels in the heart pacemaker, is widely regarded as a pure bradycardic drug that reduces \( f_H \) without producing inotropic changes in heart performance (e.g., Yusuf and Camm 2003; Luo et al. 2006). Acute dosing of ZD7288 at 30-, 250-, and 500-\( \mu \)M concentrations caused a bradycardia that was dependent on dose and time (fig. 7). The lowest concentration of 30 \( \mu \)M ZD7288 caused a significant decrease in \( f_H \), but recovery occurred much more quickly than with the higher concentrations. Acute dosing at 72 h of development with 250 or 500 \( \mu \)M ZD7288 caused the same degree of bradycardia (fig. 7). Similarly, the bradycardia induced by chronic ZD7288 measured at 72 h of development was both large and independent of dose above 7.5 \( \mu \)M (fig. 8).

There was no dose response found, suggesting that even the lowest concentration of ZD7288 saturates all the \( I_f \) cardiac pacemaker channels in the heart. In isolated mouse hearts, even smaller doses of ZD7288 (1 \( \mu \)M) were sufficient to produce stable and profound bradycardia (67% heart rate decrease) within 10 min of drug exposure, with a similar saturation effect (i.e., no further increase in heart rate) of ZD7288 at the higher concentrations (Stieber et al. 2003).

Heart rate of control embryos and embryos receiving chronic application of P-C saline were not significantly different (fig. 8). This indicates that the bradycardic effects caused by the application of ZD7288 were strictly pharmacological and not the result of the physical application of the vehicle by which the drug was administered, that is, P-C saline.

The highest concentration of 30 \( \mu \)M ZD7288 used in our chronic experiments (which did not produce a significantly greater bradycardia than the far lower concentrations) nonetheless proved to be a lethal dose to \(~50\%\) of the embryos. Additional studies to determine the effects of ZD7288 in older chicken embryos would help characterize this drug for potential use in future cardiovascular studies.

Bradycardia, Pulse Pressure, and Angiogenesis

Bradycardia and the associated increased diastolic duration increases pulse pressure in the heart and central arterial vessels in the chicken embryo (Buschmann et al. 2010). Increased pulse pressure in turn will create greater blood vessel wall distension (circumferential tension), increasing the degree of mechanical distension on the endothelial cells lining the blood vessels and potentially stimulating angiogenesis through the paracrine effects of VEGF (Groenendijk et al. 2007). In \(~3\)-d-old chicken embryos, the central aorta exhibits very modest cyclic stretch: only an approximately 3% diameter excursion between systole to diastole (Buschmann et al. 2010). Vessels more peripheral to the aorta showed no measurable rhythmic distension associated with the cardiac cycle (Buschmann et al. 2010). These observations of low vascular compliance in the early chicken embryo, which would create little to no windkessel effect to maintain a diastolic runoff of blood flow, are compatible with the measurements of the current study, showing central aortic blood velocity falling sharply to 0 in early diastole (fig. 4).

The pulsatile blood flow of the early embryonic arterial system (fig. 4) is likely to create significant and variable shear stress along the walls of the developing vascular system. Moreover, the long-standing predictions that wall shear stress is constant through the vascular tree has been replaced with measurements showing both high regional variability as well as highest shear stress in the proximal vasculature, where total cross-sectional diameters are lowest and blood velocity is highest (see Stroev et al. 2007; Reneman and Hoeks 2008). Yet in
our experiments, induced bradycardia and the attendant changes in blood flow (and associated shear stress) and pulse pressure (and associated mechanical wall distention) did not stimulate angiogenesis (as quantified by the CAM vessel density index) in vessels falling within the range of 200–600 μm from the embryo’s umbilical stock (fig. 9), the very region predicted to have the highest shear stresses. In fact, chronic bradycardia and the attendant decrease in CO actually produced a small but significant decrease in CAM vessel density index in the most peripheral vasculature (500–600 μm from the embryo) when compared with the control and P-C saline populations experiencing higher heart rates. A viable explanation could be that decreased heart rate and CO compromises perfusion and the bulk transport of materials necessary for angiogenesis, especially in the CAM periphery. The effects of ZD7288 were pronounced in the periphery of the CAM most likely as a cause of timing of drug application during development. The ZD7288 infusion began at 48 h of development after the initial vessel branch order had been formed. The effect of ZD7288 was seen in the outer branch orders in the smaller vessels that experienced angiogenesis during the experimental 24-h treatment.

The number of blood vessels increases with each increase in branch order as branch order increases in control embryos, as expected for a normal progression of angiogenesis in the CAM of control embryo. However, the embryos treated both with P-C saline and with ZD7288 exhibited a decrease in the number of blood vessels per branch order, indicating possible reduction in CAM vessel proliferation (fig. 10). ZD7288-treated embryos experiencing significant bradycardia had much shorter vessels in the outer branch order compared with control embryos, and while their vessel diameter was also smaller, the effect was not as pronounced. These data further indicate that bradycardia, the associated changes in blood flow and pulse pressure, and the anticipated changes in shear stress on the growing vessel walls do not stimulate angiogenesis at this stage of development, contrary to our initial hypothesis. Vessel diameter in the highest branch order was unaltered, showing additionally that a decreased \( f_H \) does not alter the size of the actual blood vessel formed but alters the process of normal angiogenesis, as represented by vessel number and length. These findings suggest several new avenues for research, including, for example, investigation of the ontogeny per se of the endothelial response to shear and circumferential stresses.

**Saline Application and Angiogenesis**

Unexpectedly, the mere topical application of P-C saline produced a small but significant decrease in CAM vessel density index in the most peripheral vessels examined (fig. 9). The most parsimonious explanation is that chronic P-C saline application to the embryo’s surface washed out molecules important to normal CAM vessel development. A very wide range of proteins (e.g., transcription factors) and other molecules are important to angiogenesis (e.g., Chang and Hla 2011; Gianni-Barrera et al. 2011) and could be disturbed by extracellular pools of P-C saline and the resulting concentration imbalances produced by saline moving across the surface of the growing CAM. Since the application of both P-C saline and ZD7288 decreased vessel density in the more peripheral regions of the CAM, this suggests that there may be a specific microenvirom-
mental balance that maintains normal CAM blood vessel development, at least in the peripheral if not more central regions of the CAM. The washout effect of saline-treated embryos was mainly seen at the periphery of the CAM, most likely because the initial branch orders were formed before treatment and the smaller vessels of the outer branch orders showed the effect that the washout of angiogenic factors had on decreasing vessel development. P-C saline was the vehicle for ZD7288. It is possible that this washing away of necessary angiogenic factors—rather than a bradycardic effect of ZD7288 by itself—could have affected the development of vessels in the ZD7288-treated embryos as well.

Influence of Cardiac Output on Development and Growth Rates

Eye diameter is a standard method of tracking the rate of embryo development in the chicken embryo, not only because eye diameter is easy to visualize but also because the eyes show large, linear increase in size during 72–120 h (Romanoff 1960). Similarly, decreasing cervical flexure is taken as an indication of increasing embryonic development (Männer et al. 1995). There was no significant difference in either eye diameter or cervical flexure between any of the chronically treated groups, indicating that development rate (as distinct from growth) of the embryos was not altered by chronic bradycardia. Reduced CO induced by partial ligation of the ventricular outflow tract had no effect on eye diameter or cervical flexure in day 3–4 chicken embryos (Burggren et al. 2004). In this study, pharmacologically induced bradycardia was not compensated for by an increase in stroke volume (fig. 5), and so CO declined as a result of ZD7288 treatment at 56 h. Other studies on early chicken embryos have similarly shown that f_H is a strong indicator of CO. For example, Bowers et al. (1996), examining ~84-h embryos, demonstrated that CO is linearly related to f_H (and preload) while end-stroke volume is linearly related to diastolic volume. Consequently, given the strong evidence for the fact that bradycardia decreased CO in embryos both slightly younger (56 h) as well as slightly older (84 h), we assume here that embryos acutely and chronically treated with ZD7288 and measured at 72 h (i.e., bracketed by the published data cited above) similarly experienced decreased CO directly as a result of bradycardia. We interpret from our data, then, that normal rates of development (evident from no disturbance to normal rates of eye development and cervical flexure) can continue, despite the abnormally low CO produced by a ZD7288-induced bradycardia. Expressed differently, normal progression of the embryo through successive early developmental stages is not dependent on normal levels of CO.

Does experimentally reduced CO directly influence growth (evident from embryo mass through cell division or hypertrophy, for example) as distinct from development rate measured by eye diameter and cervical flexure? Indeed, chronic bradycardia and the attendant reduction in CO significantly decreased both wet and dry body mass (fig. 11). Thus, experimentally reduced CO slowed absolute growth (measured by reduced body mass) but not development per se, as measured by eye diameter and cervical flexure, aligning with earlier experimental findings of Burggren et al. (2004).

Conclusion

Collectively, our data suggest that bradycardia at 48–72 h of development does not produce the hypothesized stimulation of angiogenesis, actually having the opposite effect in the most
peripheral regions of the CAM vasculature. However, the accompanying reduced CO induced in this study has produced two interesting findings. First, the rate of development, expressed by the key developmental landmarks of eye diameter and cervical flexure, is not affected by reduced CO. Yet embryonic growth, assessed by wet and dry weight, is significantly reduced. These findings are important in light of the fact that the convective flow of blood is not required for continued oxygen consumption or development at these same developmental stages (Burggren et al. 2000; Burggren 2004). This study indicates that convective blood flow is indeed required for absolute growth of the embryo. Thus, it would appear that in early embryonic states, while diffusion of oxygen across the embryo body wall and of nutrients and waste products between various body compartments can maintain normal oxygen consumption, absolute growth is indeed dependent on adequate convective blood flow. This study presents the novel finding that different processes in the ontogeny of the early vertebrate embryo have differential sensitivities to perturbations in convective blood flow.

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