ABSTRACT

Time-domain representations of the fetal aortopulmonary circulation were carried out in lamb fetuses to study hemodynamic consequences of congenital diaphragmatic hernia (CDH) and the effects of endothelin-receptor antagonist tezosentan (3 mg/45 min). From the isthmic aortic and left pulmonary artery (PA) flows (Q) and isthmic aortic, PA, and left auricle pressures (P) on day 135 in 10 controls and 7 CDH fetuses (28 ewes), discrete-triggered P and Q waveforms were modelized as Pt and Qt functions to obtain basic hemodynamic profiles, pulsatile waves [P, Q, and entry impedance (Ze)], and P and Q hysteresis loops. In the controls, blood propelling energy was accounted for by biventricular ejection flow waves (kinetic energy) with low Ze and by flow-driven pressure waves (potential energy) with low Ze. Weak fetal pulmonary perfusion was ensured by reflux (reverse flows) from PA branches to the ductus anteriosus and aortic isthmus as reverse flows. Endothelin-receptor antagonist blockade using tezosentan slightly increased the forward flow but largely increased diastolic backward flow with a diminished left auricle pre- and postloading. In CHD fetuses, the static component overrode phasic flows that were detrimental to reverse flows and the direction of the diastolic isthmic flow changed to forward during the diastole period. Decreased cardiac output, flattened pressure waves, and increased forward Ze promoted backward flow to the detriment of forward flow (especially during diastole). Additionally, the intrapulmonary arteriovenous shunting was ineffective. The slowing of car...

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TRANSLATIONAL PHYSIOLOGY

Phasic hemodynamics and reverse blood flows in the aortic isthmus and pulmonary arteries of preterm lambs with pulmonary vascular dysfunction

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Submitted 18 April 2008; accepted in final form 22 September 2008

Rolland PH, de Lagausie P, Stathopoulos E, Leprêtre O, Viudes G, Gorincour G, Hery G, de Magnée C, Paut O, Guys JM. Phasic hemodynamics and reverse blood flows in the aortic isthmus and pulmonary arteries of preterm lambs with pulmonary vascular dysfunction. Am J Physiol Heart Circ Physiol 295: H2231–H2241, 2008. First published September 26, 2008; doi:10.1152/ajpheart.00410.2008.—Time-domain representations of the fetal aortopulmonary circulation were carried out in lamb fetuses to study hemodynamic consequences of congenital diaphragmatic hernia (CDH) and the effects of endothelin-receptor antagonist tezosentan (3 mg/45 min). From the isthmic aortic and left pulmonary artery (PA) flows (Q) and isthmic aortic; PA, and left auricle pressures (P) on day 135 in 10 controls and 7 CDH fetuses (28 ewes), discrete-triggered P and Q waveforms were modeled as P, and Q, functions to obtain basic hemodynamic profiles, pulsatile waves [P, Q, and entrance impedance (Ze)], and P and Q hysteresis loops. In the controls, blood propelling energy was accounted for by biventricular ejection flow waves (kinetic energy) with low Ze and by flow-driven pressure waves (potential energy) with low Ze. Weak fetal pulmonary perfusion was ensured by reflux (reverse flows) from PA branches to the ductus anteriosus and aortic isthmus as reverse flows. Endothelin-receptor antagonist blockade using tezosentan slightly increased the forward flow but largely increased diastolic backward flow with a diminished left auricle pre- and postloading. In CHD fetuses, the static component overrode phasic flows that were detrimental to reverse flows and the direction of the diastolic isthmic flow changed to forward during the diastole period. Decreased cardiac output, flattened pressure waves, and increased forward Ze promoted backward flow to the detriment of forward flow (especially during diastole). Additionally, the intrapulmonary arteriovenous shunting was ineffective. The slowing of cardiac output, the dampening of energetic pressure waves and pulsatility, and the heightening of phasic impedances contributed to the lowering of aortopulmonary blood flows. We speculate that reverse pulmonary flow is a physiological requirement to protect the fetal pulmonary circulation from the prominent right ventricular stream and to enhance blood flow to the fetal heart and brain.

fetal pulmonary circulation; congenital diaphragmatic hernia; pulmonary reverse flows; pulsatile impedance; endothelin receptor inhibitor

THE PULMONARY CIRCULATION in the preterm fetal lamb (see Fig. 1) is hemodynamically unique because the distribution of the combined ventricular output is 40% to the placenta versus only about 7% to the lungs (1, 22, 33, 51, 52). Central vascular shunting leads to a communication of the pulmonary and aortic circulation with driving pressures that would be considered as hypertensive if it persisted after birth (1, 14, 47). These characteristics are functionally optimal for the fetal lung that does not participate in gas exchange and requires only a small amount of blood for development (14, 23, 25, 27, 47). In previous studies we used the preterm fetal lamb model to study lung hypoplasia caused by a congenital diaphragmatic hernia (11, 61) and the porcine thoracic and abdominal aorta model to study anatomical bypass- and stenting-induced hemodynamic changes (38, 39, 48). The purpose of the present study was to investigate two hypotheses that we have formulated based on previous studies. As elaborated below, hypothesis 1 is that phasic pulmonary vascular resistances (PVRs) have been overestimated, and hypothesis 2 is that reverse pulmonary flows contribute to reverse isthmic flows. We also studied the effect of administering an endothelin-receptor antagonist (ERA) tezosentan (TZ) in the left pulmonary artery (LPA).

Hypothesis 1. The high fetal PVRs that are consistently used to explain the low pulmonary perfusion have been estimated based on the ratio of the mean transpulmonary pressure gradient to the mean transpulmonary flow (1, 4, 14, 22, 28, 29, 33, 45, 51, 52). Similarly, the input impedance spectrum of pulsed fetal pulmonary artery (PA) hemodynamics, estimated in the frequency-domain analysis, has been used to characterize right ventricular (RV) afterload, proximal arterial compliance, distal vascular resistance, wave-transmission properties, and pulse wave interactions causing phenomena such as reverberations and wave amplification (4, 8, 28, 45, 46, 63). However, frequency domain analysis does not take into account the perfusion of the fetal pulmonary parenchyma by a direct venous admixture from intrapulmonary arteriovenous shunting (iPAVS) (8, 13, 21, 34, 37, 56, 58, 63). The iPAVS (see Fig. 2) is dependent on the supernumerary vessel group that branches from the small PAs at a right angle and acts as a parallel vascular network limiting distal pulmonary perfusion. In lamb fetuses, iPAVS normally occurs during late gestation and the early neonatal period and disappears thereafter. Finally, the frequency domain analysis does not take into account reverse flows from the pulmonary circulation to the ductus arteriosus (DA) and aortic isthmus. These shortcomings of frequency
domain analysis led us to consider that PVRs have been consistently overestimated and that this can be demonstrated by reassessing the preterm lamb fetal hemodynamic based on phasic flow-pressure relationships in impedance profiles and hysteresis loops in the time domain during the cardiac cycle with mathematical modeling of discrete, averaged and timesynchronized pressure (P) and LPA flow (Q) signal waves to obtain their respective $\Omega_P(t)$ and $\Omega_Q(t)$ functions and instantaneous entry impedance ($Z_e$) as the ratio of $\Omega_P(t)$ and $\Omega_Q(t)$ functions (5, 38, 39, 48, 50).

**Hypothesis 2.** The fetal DA is the main vascular outlet of the prominent fetal RV. It forms the pulmonary arch with the main PA and descending thoracic aorta (1, 16, 20, 33, 51, 52). Blood from the fetal RV perfuses subdiaphragmatic organs with 10–15% going to the pulmonary circulation while blood from the left ventricle (LV) perfuses cephalic regions (16, 22, 30, 51). These two arterial flows merge at the junction of the aortic isthmus and DA with the descending aorta. This arrangement implies that output from the RV and LV has opposing effects on flow through the isthmus with LV output causing forward flow and RV output causing backward flow. When reverse flow develops in the aortic isthmus, the preplacental blood with low oxygen saturation coming from the descending aorta can mix with the ascending aortic blood destined to the brain, which can potentially create a risk for cerebral hypoxia (1, 7, 19, 22, 33, 51, 52). In ovine fetuses, a stepwise increase in resistance to placental blood flow lowers oxygen delivery to the brain in association with a major reversal of diastolic blood flow through the aortic isthmus (7, 19). Oxygen delivery to the brain decreases as the net flow through the aortic isthmus becomes retrograde (18, 19, 35, 54).

Understanding the physiology of the aortic isthmus flow as well as its pathological alterations and potential pharmacological controls is important to ensure optimal fetal growth and maturation until delivery and particularly to prevent nonoptimal neurodevelopment (18, 19, 54).

An ultrasonographic study of dynamic phasic events during the last months of gestation in normal fetuses has shown that
forward flow in the aortic isthmus during systole and diastole progressively decreases (54). This finding has been attributed to the increasing preponderance of the RV output and the leveling of the placental vascular resistance as cerebral vascular resistance progressively declines (17, 20, 29, 30, 40, 54). During the cardiac cycle, a brief end-systolic reversal of the isthmic flow velocity profile occurs and steadily increases as gestation progresses (20, 32, 33, 54). Late systolic flow reversal and admixture come from both the isthmus and DA as a result of reflux caused by the delayed onset and prolonged higher acceleration of the ductus flow velocity (54). The ejection and peak flows occur earlier in the pulmonary trunk than in the ascending aorta, whereas the PA peak flow at the aortic end of the DA occurs later than the aortic peak flow. It has been noted that signals in the left PA branch show a brief systolic forward-flow spike followed by late-systolic and diastolic flow reversal (16, 54). The capacitance of the main and branch PA has been implicated in the altered timing of the circulatory events, and the differences might be related to changing vessel diameters (54).

These investigations of the flow velocity led us to the hypothesis that a possible explanation for the slowdown in PA flow waveforms through the main PA and ductus is that the flow enters the pulmonary circulation before returning to the DA and isthmus through reverse pulmonary flows. In this regard, it is interesting to point out that the observation of the onset of flow in LPA before the pulmonary arterial pressure has up to now been considered as a technical artifact (33, 54).

Pulmonary artery hemodynamics, which depend on optimal reciprocal endothelial-epithelial cross talk, contribute to optimal alveolarization and angiogenesis during lung development. However, another critical factor for normal development is the degree to which the fetal lungs are expanded by liquid secreted into the future airways and exiting the trachea (10, 15, 25, 45, 46, 58, 59). Deflation of the fetal lungs during development results in hypoplastic lungs that are deficient in alveoli and exhibit increased proportions of type II alveolar epithelial cells (12, 45, 46, 59). A number of studies on lung hypoplasia (2, 12, 34, 42, 46, 53, 58, 62) have assessed the correlation between fetal lung expansion and prolonged impairment of the signaling necessary for terminal vascularization and alveolarization. In the sheep model of surgically created congenital diaphragmatic hernia (CDH) (34, 58, 62), growth impairment retards development and angiogenesis, especially during the alveolar phase. Reverse flows across the aortic isthmus in combination with reduced blood flow velocities are detectable in the PA of fetuses presenting pulmonary hypoplasia related to CDH (31, 32, 59).

**ERA blockade.** Alterations in pulmonary vascular reactivity and smooth muscle cell proliferation, in which nitric oxide (NO) and endothelin (ET-1) appear to be key factors, are prominent causes of abnormalities in the pulmonary circulation (for review, see Refs. 6, 41, and 53). In the lamb model, an increased pulmonary blood flow at 1 wk of life has been associated with alterations in ET-1 signaling followed by changes in NO signaling (6, 43). Various pharmaceutical agents have been used to successfully decrease PA pressures and resistances in patients (for review, see Refs. 6, 41, and 58). However, even though CDH can be detected in a fetus by ultrasound, an early intervention is poorly effective without tocolysis (24, 26).

Preterm vascular intervention in the fetal lung therefore is a rational approach to enhance the likelihood of survival during the life-threatening period of extrauterine life (26). TZ, a dual ET-1 antagonist that blocks both endothelin types A and B (ET_{A} and ET_{B}) receptors, currently is the only ET-1 receptor antagonist that can be delivered systemically in the last week of gestation when vascular and alveolar pulmonary development enters the terminal phase (9). Therefore, we also investigated the contribution of preterm vasoconstricting processes in both situations by evaluating the effect of a dual ERA, pharmacologically designed for intravenous delivery (9). The protocol was applied to control fetuses with normal lung expansion and to experimental lamb fetuses with restricted lung expansion induced by surgically created CDH (62). Concomitantly, the three-dimensional organization of fetal lungs was investigated under scanning electron microscopy (SEM) (44), with a special focus on the supernumerary vessels considered as pulmonary arteriovenous shunts.

**MATERIALS AND METHODS**

**Animals and surgical procedures.** The Animal Use and Care Committee of the Marseille School of Medicine reviewed and approved all procedures and protocols used in this study that was in compliance with the requirements of the Animal Welfare Act and the Guide for the Care and Use of Laboratory Animals (National Research Council, 1996). Twenty-eight Merinos d’Arles ewes (National Institute of Agricultural Research, Ewe breeding in dry countries, INRA 868, Montpellier, France) were synchronized using intravaginal fluorogestone (30 mg) and pregnant mare serum gonadotrophin (40 IU, 14 days later) 3 days before mating. Forty-five days after mating, pregnancy testing was carried out using pregnancy-specific glycoprotein assays. Ewes were enrolled at the rate of two per week throughout the study.

Before surgery, the animals were fasted for 24 h and anesthetized by a slow infusion of a bolus of pentobarbital sodium (250 mg) in an external jugular vein and a lumbar intrathecal administration of 1% maraine (2 ml) as previously described (61). Pentobarbital sodium was titrated to achieve a continuous narcosis throughout the surgical procedure. Six ECG leads and one transcutaneous arterial oxygen saturation/plethysmographic probe (HP78354C) were placed. Animals were ventilated via endotracheal tubing using a controlled oxygen-air mixture. Intravenous (ear and vena cava) and intra-arterial (superficial femoral artery, 6 Fr-introducer sheath, BSC, Natick, MD) lines were placed for perfusion with Ringer lactate solution (15 ml·kg⁻¹·h⁻¹) and endovascular hemodynamic monitoring of heart rate, blood pressure, and oxygen saturation (Mistral monitoring devices, Agilent Technologies) (38, 49, 50). Aseptic techniques were used and prophylactic antibiotics (penicillin G streptomycin, 1 g) were administered before and after surgery (48). In ewes carrying twins, only one fetus underwent instrumentation.

The diaphragmatic hernia was created under sterile conditions at day 75 as follows (61). A midline abdominal incision was made, and the uterus was externalized. Following a hysterotomy and a local pregnancy testing was carried out using pregnancy-specific glycoprotein assays. Ewes were enrolled at the rate of two per week throughout the study.

On the 135th day of gestation, the ewes were reoperated upon using the same anesthetic and monitoring procedures (61). Three pediatric ECG leads were placed on the upper body of the fetus. After a local
infiltration of lidocaine (2 ml, 1% solution), the left axillary artery and vein of the delivered left forelimb of the fetus were exposed and ultrathin-thick-walled 22-gauge polyvinyl catheters were advanced through the vessels to the ascending aorta and superior vena cava, respectively. A left thoracotomy (5th intercostal space) and pericardial incision (and further suspension) were performed to expose the heart and great vessels (see Fig. 1). Using an appropriate intravenous placement system (Angiocath, Travenol, Deerfield, IL) through purse-string sutures (6/0), 22-gauge polyvinyl catheters were introduced into the LPA and main PA. The catheters were connected to Stöckert III CDX III pressure transducers (COBE Cardiovascular, Arvada, CO). A testing of the catheter-transducer system using the flush method showed no secondary resonance peak up to a frequency of 250 Hz and a low mean damping coefficient (<0.15%) (54, 55). A 2-Fr Mikro-Tip catheter-pressure-transducer probe (SPR-249/Millar Instruments, Houston, TX) was likewise implanted into the left auricle. For the specific requirements of this study, indwelling catheters connected to fluid-filled pressure probes were inserted for the delivery of dips in the PA. Identical catheters, i.e., the same model and length (40 cm), were placed in the PA and aorta to ensure uniform measurements. To rule out the possibility of artificial decay of the pressure waveform suspected with fluid-filled catheters (54), we checked that the waveforms obtained after generating a R-wave trigger of the ECG were superposable using the CDX III pressure transducers and 1.8-Fr Mikro-Tip transducer (model SPR-612, Millar Instruments). Probes were precisely positioned by fluoroscopic guidance. To measure blood flow, 4- and 6-mm RS ultrasonic flow transducers (Transonics, Ithaca, NY) were placed around the LPA and ascending aorta on the aortic isthmus, i.e., between the DA and brachiocephalic trunk, respectively (see Fig. 1). Catheters were flushed with 2 ml of heparinized saline (2.5 IU/ml) to avoid a thrombotic obstruction. Control animals underwent the same procedure on the 135th day of gestation.

Before any measurements, a 20-min period with continuous pressure and flow monitoring was allowed for hemodynamic stabilization. After the oxygenation status of the fetus was checked, the vascular reactivity in the LPA was investigated in response to ACh (15 μg/ml, 1 ml/min, 2 min infusion) (33), and another 30-min period was allowed for stabilization. Basal hemodynamics parameters were recorded three times for 30 s before the start of the infusion of TZ (Actelion, Basel, Switzerland) into the LPA (120 μg/ml, 555 μg/min, overall dose: 3 mg/45 min). At the end of the experiment, the fetal lamb was euthanized with potassium chloride (15% wt/vol, 10 ml) and midazolam (20-mg bolus). The fetal heart and lungs were removed and immersed in ph 7.3 buffer solution pending a histological and SEM examination.

Aortic and pulmonary hemodynamics. Analog dimensional and ECG signals were digitized using an ATMIO-16X analog-to-digital converter (16 bits ADC, 1,000 Hz sampling frequency, National Instruments, Austin, TX) and recorded for at least 50 consecutive cardiac cycles (standard recording time, 30 s) using a customized data capturing window in the Lab View 7.0 software (National Instruments) (5, 38, 39, 48, 50). Throughout the experiment, a parallel acquisition of processed and raw data from each input channel was performed using an iox2 data acquisition and analysis system (EMKA Technologies, Falls Church, VA). The cycle-to-cycle wave variability was overcome by generating a steady-state cycle (256 points) from the time-averaged P and Q cardiac cycles calculated using linear interpolation according to the following formula:

\[
\text{Mean period } T = \frac{1}{N_c} \sum_{j=1}^{N_c} N_j,
\]

where \(N_j\) is the number of points within each cardiac cycle, \(N_c\) is the number of cycles, and \(f\) is the sampling frequency. A mathematical modeling of the discrete P and Q signal waves was then performed to obtain the respective \(\Omega_P(t)\) and \(\Omega_Q(t)\) analytical functions after the fractionation of curves into successive segments separated by two opposite extreme points scaled to measured values offset in the two-dimensional coordinate axis and settling into an increasing or decreasing nature of the waveform segment. In the basic mathematical model, the curve is considered as the second derivative of a sigmoidal function and extracted the elementary segment between the minimum and maximum. The initial waveform was synthesized by summing individual segments and fitting the curve to the original pulse wave so as to obtain the error that was treated in the same manner. The final \(\Omega_P(t)\) and \(\Omega_Q(t)\) analytical functions corresponded to the sum of the serial functions. The model was applied to both the P and Q waves. The quadratic error normalized to the original waveforms was low regardless of the experimental conditions.

The mean hemodynamics of the P and Q waves were calculated as the systolic (maximum), diastolic (minimum), pulsatile (\(A = |\max - \min|\)), and mean values from their respective \(\Omega_P(t)\) and \(\Omega_Q(t)\) functions. The results were expressed as millimeters of mercury and milliliters per minute. The algebraic mean pressure (\(P\)) and flow (\(Q\)) values were calculated according to the following formulas:

\[
\overline{P} = \frac{1}{T} \int_0^T \Omega_P(t) \, dt \quad \text{and} \quad \overline{Q} = \frac{1}{T} \int_0^T \Omega_Q(t) \, dt
\]

respectively, where \(T\) is the mean cardiac cycle period value.

The parametric equations of hysteresis loop for P-Q relationships were defined as follows:

\[
\begin{align*}
X &= \Omega_P(t), \\
Y &= \Omega_Q(t).
\end{align*}
\]

Peripheral resistance (\(R_p\), in mmHg·min⁻¹·ml⁻¹) and mean input impedance (\(Z_e\), mmHg·ml⁻¹·min) were calculated according to the following formulas, respectively:

\[
R_p = \frac{\overline{P}}{\overline{Q}} \quad \text{and} \quad Z_e = \frac{\Delta P}{\Delta Q} (\Delta P = P_{\max} - P_{\min}, \Delta Q = Q_{\max} - Q_{\min})
\]

The instantaneous \(Z_e\) (\(Z_e\), in kN·s·m⁻²) during the cardiac cycle was calculated as the ratio of the function of the pressure and flow according to the following formula:

\[
Z_e = \frac{\Omega_P(t)}{\Omega_Q(t)} \sum_{k=0}^{2} \sum_{i=0}^{2} \xi_P \xi_Q
\]

Scanning electron microscopy. Electron microscopy using a Quanta 200 SEM (FEI, Eindhoven, The Netherlands) was performed on the fetal heart and lungs that were removed at the end of the experiment and stored in 10% formalin (44). Specimens were prepared as follows. After being rinsed with the fixative solution, the surgical specimens were fixed for another 48 h. Block samples of the diaphragmatic and cephalic pulmonary lobes were made and further fixed for 24 h before being cut into 3-mm serial sections. The sections were submitted to sputter coating with a 30-nm gold layer with or without prior dehydration in progressive alcohols (low vacuum mode, 1 Torr with water vapor).

Statistical analysis. Data treatment and statistical analyses were performed using Systat 10.0 software (SPSS, Chicago, IL). Results were expressed as means ± SD unless otherwise specified. An analysis of variance was performed with either the Kruskall-Wallis nonparametric test (Scheffé’s post hoc test) or the Mann-Whitney test when there are only two groups (49).
RESULTS

Technical outcome. A total of 21 fetuses, including 14 control and 7 CDH fetuses, completed the entire protocol by fulfilling the required inclusion criteria. Eleven fetuses were excluded because of ischemic status (7 cases), weak vascular reactivity to ACh (2 cases), and surgically provoked bleeding (2 cases). Weight gain was lower in CDH fetuses than in control fetuses (2.42 ± 0.24 vs. 3.08 ± 0.32 kg; P < 0.01). Preprocedural blood gas tension, postprocedural blood gas tension, and pH in the superior caval vein in CDH fetus versus controls were as follows: PO2, 20.32 ± 4.67 versus 19.10 ± 4.47 mmHg; PCO2, 48.98 ± 7.61 versus 51.82 ± 7.61 mmHg; and pH 7.25 ± 0.06 versus 7.22 ± 0.11 (P > 0.1). There was no significant difference in heart rate between CDH fetuses and controls (overall mean baseline heart rate, 161 ± 14 beats/min).

SEM findings in the left lung. In controls, photomicrographs of the left lung (Fig. 2) demonstrated normal long PA branches alongside the airways, and alveoli were nonexpanded and uniformly distributed throughout the lung. Supernumerary vessels were identified arising orthogonally from the PA between the normal long branches. These vessels supplied the capillary arterial bed in the immediate vicinity of the PA branch from which they arose. A surface tridimensional examination also showed that some supernumerary vessels allowed the direct venous admixture by creating a network of connections between 100- and 500-μm PA branches and similarly sized veins. The supernumerary vessels anatomically can result in iPAVS that may potentially exclude alveoli from the PA blood flow.

In CDH fetuses (Fig. 3), SEM demonstrated clear evidence of an impaired development of alveolar and vascular structures in the left lung. Some PA branches were severely thickened and opened, and adjacent bronchi were embedded within nonvascularized regions with proliferating parenchymal cells showing no alveolar organization. The degree of impairment was variable with regions showing persistent vascuarization and alveolarization but with substantial disorganization compared with controls. Intimal proliferation, splitting of elastic laminae, medial and adventitial hyperplasia, and thickening were characteristic features of PA remodeling. The areas of neoangiogenesis and bronchiogenesis with the underdeveloped and interlaced dead-end tubes were consistently observed in the less impaired regions of the hypoplastic lung. No iPAVS was detectable in the hypoplastic lung.

Phasic hemodynamics in the controls. The pulsatile pressure range in both the fetal PA and AO was not incidental. Indeed, although mean pressure values were not significantly higher in the PA than in the AO, the minimal and peak pressures in the PA and AO were identical. This finding strongly indicates that vascular compliances were elevated in both vascular beds (Table 1). The isthmic pulsatile pressure profile consistently showed a symmetric sine contour centering on the systolic peak that accounted for 77 ± 6% of the cardiac cycle (Fig. 4). In contrast, the systolic flow peak preceded the systolic pressure peak and the ensuing post-systolic flow consisted of a marked backward flow. The isthmic $Z_e$ profile showed that most of the aortic flow encountered low forward phasic impedance. The forward impedance associated with the aortic pressure-ascending phase was framed by two impedental events having asymptotic boundaries, whereas the diastolic phasic impedance was negative. The boundaries were mathematically attributable to changes in flow direction: the end

Fig. 3. Photomicrographs (scanning electron microscopy) of hypoplastic left lung (transverse section of the diaphragmatic lobe) in a lamb fetus with CDH and bronchopulmonary dysplasia. Left: proliferating parenchymal cells, nonvascularized and nonalveolarized regions (arrow), mixed with areas of milder injury. The bronchi and arteries are wide open. Arteries show medial and adventitial hyperplasia (details in inset). Right: intense foci of immature, anarchic neoaangiogenesis (arrow), neobronchiogenesis, and alveolarization.
systolic Z change coincided with the decrease in systolic flow, and reversal occurred when the zero-flow value was reached. The values showed the opposite pattern when the diastolic forward flow was observed in the early systolic forward flow. The flow-pressure hysteresis loop depicted how the increasing aortic pressure was associated with the end of the forward-flow segment. The reversal of isthmic flow provided the energy necessary to supply blood to the cephalic and coronary arteries.

In contrast with the biphasic profile observed in the aortic circulation, the PA hemodynamics were more complex. The PA systolic pressure peak (lasting 31 ± 6% of the cardiac cycle) displayed a transient shoulder that was synchronous with PA systolic flow and occurred before the isthmic pressure peak. The diastolic PA pressure peak that was synchronous with the left atrial pressure peak was no longer detectable in the pulmonary wedge pressure profile, being a reaction to left atrial contraction and emptying. The pulsatilie LPA flow consisted of three consecutive forward peak waves interspersed with a substantial reverse flow. The sharp PA ejection peak generated the early shoulder on the systolic pressure peak. Reverse flows were synchronized with the systolic and diastolic peaks. The diastolic peak was induced by left atrial pressure rise. The PA Z profile showed a series of temporal events delineated by asymptotic boundaries. The PA Z profile showed that Z against forward PA flow was 10 times lower than Z against reverse PA flow. The flow-pressure hysteresis loop showed that LPA flow reached its maximum when PA pressure was at only 10% of maximum and then abruptly dropped to zero-flow status. The decreasing PA pressure period was associated with the backward flow except during the mid-diastolic forward-flow segment corresponding to left auricular loading before a contraction.

**Phasic hemodynamics in CDH fetuses.** Although still a symmetric systolic peak centered on the cardiac cycle, the aortic pressure waveform in CDH fetuses was drastically flattened due to elevated PAo min and decreased pulse pressure, whereas PAo max remained constant (Table 1 and Fig. 5). The phasic component of the pressure driving systemic blood flow fell 50% and, consequently, the pulsatilie-to-static hemodynamic ratio changed from 1:1 to 1:4 in CDH fetuses. These hemodynamics changes were highly detrimental to reverse

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**Table 1. Mean aortopulmonary hemodynamics in control and CDH fetal lambs**

<table>
<thead>
<tr>
<th>Basal situation</th>
<th>Control Fetuses</th>
<th>CDH Fetuses</th>
</tr>
</thead>
<tbody>
<tr>
<td>P&lt;sub&gt;AO&lt;/sub&gt;, mmHg</td>
<td>26±7</td>
<td>126±11</td>
</tr>
<tr>
<td>Q&lt;sub&gt;AO&lt;/sub&gt;, ml/min</td>
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<td>-126±11</td>
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<td>38±10</td>
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<td>Q&lt;sub&gt;LPA&lt;/sub&gt;, ml/min</td>
<td>-126±11</td>
<td>-148±34*</td>
</tr>
<tr>
<td>P&lt;sub&gt;LPA&lt;/sub&gt;, mmHg</td>
<td>13±8</td>
<td>8±4*</td>
</tr>
</tbody>
</table>

Values are means ± SD; n = 10 control and 7 congenital diaphragmatic hernia (CDH) fetuses. P, pressure; Q, flow; Z, entry impedance; A0, aorta; PA, pulmonary artery; LPA, left PA; LA, left atria; TZ, tezosantan. *P < 0.01 vs. control under basal situation; †P < 0.01 vs. control after TZ treatment; ‡P < 0.01 vs. CDH bronchopulmonary dysplasia basal situation.
flows: the isthmic flow profile showed a major redistribution of forward-flow throughout the cardiac cycle. The ventricular systolic flow peak lasted 41 ± 5% of the cardiac cycle, and the forward diastolic peak was strong enough to cross over the zero-flow value. Isthmic Ze was dramatically higher during both systole and diastole and for both forward and reverse flows. The aortic P-Q loop shifted to the right as a result of increased diastolic pressure. A rapid decrease in pressure-driven systolic forward flow occurred before crossing the zero flow. Reverse flow was transient with the pressure peak and did not resume until telediastole.

In the PA, the higher minimal pressure and lower pulse pressure with no change in systolic pressure resulted in a coalescence of three PA pressure contours into a single systolic peak centered on the cardiac cycle. There was a parallel drop in LPA. Systolic LPA forward flow was prolonged to 8 ± 2% of the cardiac cycle, and the second forward-flow peak shifted to the right as a result of the pressure peak coalescence. As the pressure increased in the LPA, the flow turned to a long-lasting diastolic forward flow due to the drastic decrease in the left auricular pressure that accompanied the decreased PA flow. Damped flows encountered dramatically elevated Ze, especially the diastolic forward flow that was almost cancelled out by 50 times higher Ze. The P-Q hysteresis loop indicated that the LPA flows were restricted to the spindle-shaped systolic forward flow and the curved reverse flow associated with the maximum pressure value. The remaining interspersed half of the cardiac cycle was characterized by near-zero flows.

**Pulsatile hemodynamics in the controls treated with TZ.** Perfusion of TZ in the LPA for 45 min achieved stabilization after 25 to 35 min (data not shown). Changes in the isthmic hemodynamics included an increase in minimal pressure and reverse flow resulting in a decrease in pulse pressure and an increase in pulse flow (Table 1 and Fig. 6). Asymmetry accompanied the shortening of the systolic aortic pressure peak (lasting 51 ± 5% of the cardiac cycle, P < 0.05 vs. basal value). With no change in forward Ze, systolic isthmic flow peak was reduced by one-fourth and still preceded the systolic pressure peak, contributing to the shoulder on QAo systolic contour. The diastolic reverse flow reverse increased due to a tenfold decrease in Ze (increase, QAo min > PAo min). The aortic P-Q loop depicts the sharpening effect of TZ perfusion on the initial systolic P-Q slope with the induction of a drastic drop in the P-Q slope of the reverse flow plateau that lasted throughout diastole.

After perfusion of TZ, the diastolic pressure in the PA increased, leading to a decrease in the pulse pressure, accompanied by an increase in pulse and mean flow as a result of an increase in both forward and reverse flows. The flow-induced pressure formed an enlarged symmetric peak lasting 68 ± 6% of the cardiac cycle (P < 0.05 vs. basal value). The enhanced LPA ejection was accompanied by a substantial increase in LPA pulse flow. The first diastolic flow peak that was in the forward direction in controls was converted into a transient lowering of the backward flow. The left auricular loading phase and pressure substantially decreased after TZ perfusion due to the increased backward flow in the PA. The forward and backward flows encountered higher Ze due to a coalescence of previously separated impedental events. The hysteresis loop shifted to the right due to the increased diastolic pressure and clearly demonstrated that the forward flow was due solely to...
the enhancement of the systolic peak flow and higher relief in the late diastolic phase. The systolic flow began at a low threshold of diastolic reverse flow and provided more energy for the ensuing systolic forward flow.

Pulsatile hemodynamics in CDH fetuses treated with TZ. After TZ perfusion, the mean isthmic flow reversed due to a marked decrease in the systolic forward flow accompanied by a twofold increase in the reverse flow (Table 1 and Fig. 7). The enhancement of the systolic forward flow was immediately followed by a drastic increase in the reverse flow that progressively returned to a low-grade forward plateau. The diastolic forward $Z_e$ increased markedly. The aortic P-Q hysteresis loop demonstrated that that the early systolic forward flow peak was held at a plateau for a longer time but at a lower level. When the maximum pressure was reached, the plateau suddenly changed into an amplified reverse flow before returning to an almost completely forward flow throughout diastole.

After TZ perfusion, the CDH-induced flattening of the systolic pressure peak in the PA was shorter in duration (lasting $55 \pm 3\%$ of the cardiac cycle). The mean flow was in a forward direction and increased due to a marked increase in the diastolic forward flow. The enhancement of systolic forward impedance was noticeable due to a drastic decrease in the impedental events throughout the remaining cardiac cycle. When compared with baseline, the P-Q hysteresis loop clearly showed an increase in the magnitude of the forward systolic peak and a marked enhancement in both the duration and magnitude of diastolic reverse flow.

**DISCUSSION**

This study assessed the pressure and flow wave relationships in near-term fetal lambs. These waves were modeled into parametric phasic hemodynamics synchronized with an R-peak trigger of the ECG with anterograde flow being zeroed and positive, the reverse flow being negative, and the “reverse” impedance (a term devoid of physical meaning) being defined as the impedance encountered by reverse flow. The specific sites of investigation (see Fig. 1) were the aortic isthmus and LPA in the aorta-PA-DA fetal vascular network. This anatomically complicated network is interconnected by central cardiac shunts and presents large outlets, i.e., the supraaortic/brain/coronary arteries and thoracic aorta/placenta as well as middle-sized tributary effluents, i.e., the PA branches (3, 4, 8, 10, 34, 63).

The most outstanding finding in control fetuses was that the energy required for blood circulation was supplied in a sequential manner by two components. The first component involved the synchronous aortic and pulmonary systolic flow waves (kinetic energy) produced by biventricular cardiac ejection. The second component involved a recoil of compliant arterial walls following respective flow-related distension pressure waves (potential energy) (5, 38, 39, 48, 50). A unique feature of fetal pulmonary arterial circulation is that phasic, pulsatile hemodynamics is equivalent to static hemodynamics. The systolic pressure peak generated by prominent right ventricle ejection encounters low phasic arterial impedance and is synchronized with reverse LPA flow that is highly sensitive to left auricular events due mainly to efficient PAVS (28). The same results were previously obtained in the preterm fetal lamb by...
Invasive measurements of pressure and flow with "manual" postsynchronization to the R wave of the ECG (33, 54). The first hypothesis of this study was that phasic PVR has been overestimated. Our data show that weak fetal pulmonary perfusion is not accounted for by elevated PVRs. Indeed, an analysis of phasic hemodynamics (i.e., 50% of the pulsed flow vs. static component) revealed that the Zr in PA branches was low. The contributing components are a substantial reflux from the LPA (and other PA branches) to the DA and aortic isthmus and functional iPAVS. The potential impact of iPAVS on pulmonary hemodynamics is supported by high sensitivity to atrial events elicited by flow in the PA and its branches (21, 23, 28, 29, 63).

Our second hypothesis was that a reverse pulmonary flow ultimately directed into the isthmus promoted forward cerebral and coronary blood flow in the later stages of the cardiac cycle. This would potentially be more energy efficient since the pressure wave generated by the outgoing blood was used to propel the isthmic flow. In addition, this flow distribution would prevent oxygen-depleted visceral blood in the descending aorta from reaching the cerebral and coronary arteries.

The data in CDH fetuses revealed that the static component of the pulmonary circulation became dominant over the phasic component. The pressure-driven phasic component of biventricular ejection was reduced by 50% (pulsatile-to-static ratio, 1:4). The RV and LV ejection volume markedly decreased, but cardiac unloading prevented the blood pressure from rising (1, 3, 4, 22, 23), and the well-documented increase in arterial impedance was observed in the hypoplastic lung. These hemodynamic changes were detrimental for the reverse flows, and the isthmic flow changed to the forward direction during diastole. This response was physiologically appropriate to preserve brain oxygenation since maintaining the reverse flow such as the reverse pulmonary flow decreased would have required the reverse isthmic flow from the descending aorta and visceral circulation. Our results indicated that the blood flowing to the coronary and brain arteries was slowed but not contaminated by poorly oxygenated blood. It was estimated that the drop in the amount of oxygenated blood reaching the brain was <50%.

The contributing factors to diminished pulmonary flows included morphometric (and functional) PA alterations and the presence of nonvascularized regions with proliferating parenchymal cells showing no alveolar organization. These factors were enhanced by the absence of iPAVS and increased sensitivity to left auricular hemodynamics (14, 34, 61, 63). The observation of an asynchronous decrease in kinetic and subsequent potential energies in CDH fetuses supported the prominence of flow changes over pressure changes and suggests a cardiac origin. A poor propulsion of the aortopulmonary blood flow was attributed to a combination of concomitant factors including decreased cardiac output, damping of pressure waves and pulsatility, and heightened phasic impedance. The shortening and damping of the systolic flow waveform are in agreement with previous studies in the developing heart, showing that chronic pulmonary insufficiency causes global cardiac dysfunction, e.g., an impairment of functional reserve and myocardial contractility, but that diastolic function was preserved (23, 28). In clinical settings, neonates with CDH have a much higher prevalence of cardiac defects than normal infants and a lowered LV-to-RV ratio correlates a poor outcome and pulmonary hypertension (24, 32, 60). These results strongly advocate a therapeutic strengthening of the failing cardiac function that may recover with the restoration of the appropriate PA flow. Specifically, it may be of therapeutic importance to determine whether or not the CDH-induced changes in the reverse flows are related to the changes in ventriculo-arterial coupling and thereby to LV-to-RV-ratio lowering.

Our findings in preterm fetuses indicate that experimental CDH had a greater impact on pulmonary flow than on pulmonary pressure. Previous investigations of the effects of prolonged tracheal obstruction (TO) and subsequent increased lung expansion on pulmonary hemodynamics in fetal sheep showed that an equalization of intraluminal pressure markedly increased pulmonary flow in TO fetuses (45, 46, 59). Most changes in the pulmonary flow waveform occurred during diastole with few changes during systole. The diastolic pulmonary flow was mainly responsible for generating the backward compression and expansion waves that interact with the pulmonary vascular bed to cause retrograde flow (23, 29, 45, 46). Knowing the extent of the impact on backward flow caused by the enhancing effect of TO on the forward flow and by the increased luminal pressure related to liquid filling might be of critical importance for therapeutic decision making. In regard to therapy to restore impaired pulmonary circulation in the hypoplastic lung, a potentially interesting finding of our study was the presence of an intense foci of neoangiogenesis and bronchiogenesis with interlaced immature and dead-end tubes in less disorganized regions of the lung. The epithelial cells in distal airspaces of the fetal lung expressed VEGF, suggesting that the epithelium might regulate a temporal and spatial development of alveolar capillaries (25, 53, 58, 61). Since VEGF is considered as an endothelial cell survival factor that can improve oxygen and cytokine injury (25, 53, 62), it is tempting to speculate that the aberrant angiogenesis represents an attempt by the still-organized alveolar cells to restore the appropriate vascularization through persistent VEGF secretion.

Signaling pathways mediating the pulmonary vascular response that was either impaired in the hypoplastic lung or heightened in the case of hypertension have still not been clearly defined (for review, see Refs. 53 and 58). In control fetuses, we observed that endothelin receptor (ER) blockade had a slight favorable effect on the forward flow in both the aorta and LPA, whereas it drastically enhanced the backward flow during the prolonged diastolic period. The net effect of these two changes was a decrease in left auricular pre- and postloading. Forward flow was favored during systole in the aorta and during diastole in the PA where decreasing vascular impedance enhanced back pumping from the pulmonary circulation. The consequences of TZ perfusion on both forward and backward flow were correlated with the changes in cardiac cycle (e.g., shortening of systolic duration) and arterial effects (e.g., vascular wall stiffness and elasticity), respectively. These hemodynamic effects were likely mediated by NO: recent findings in juvenile lambs demonstrated that the increase in mean blood flow observed in TZ-treated lambs was due to an increase in NO synthase activity in lung tissue that was not observed in control lambs (42).

Our observation of a drastic 10-fold decrease in aortic impedance after TZ perfusion was consistent with the presence of ET receptors in the aorta (11, 41). Similarly, the TZ effects in the PA were consistent with a different spatial distribution of the ER along the pathway as well as with the differences in the
responses of ET\textsubscript{A} and ET\textsubscript{B} receptors to ET-1 (9). The vasoconstricting properties of ET-1 were inhibited in the proximal segments of the PA where ET\textsubscript{A} receptors are predominant (11, 41, 62). In distal segments of the PA where ET\textsubscript{B} receptors that mediate vasodilatation through activation of the NO/prostanoid signaling system are predominant, TZ strongly enhanced the forward flow during diastole. Since forward diastolic flow is dependent on vasodilatation in smaller PAs, this finding strongly supported the view that acute TZ perfusion did not elicit the vasoconstricting effects associated with ET\textsubscript{B} receptor blockade (9, 41).

In CDH fetuses the hemodynamic effects of ER blockade were limited to a slight increase in forward flow during systole in the aorta and during diastole in the PA. The mechanisms regulating these processes mainly involved a decrease in the corresponding impedances. The pharmacological effects of ERAs in CDH fetuses were unclear. Previous studies have provided contrasting evidence. One study showed that the ET\textsubscript{B} receptor agonist sarafotoxin had no effect, whereas another showed that the ET\textsubscript{A} receptor antagonist BQ-123 caused a greater pulmonary vasodilatation in the CDH than in the control animals (42, 61). The findings that TZ was inefficient for acute blockade of ET\textsubscript{A} receptors in lamb fetuses with CDH argued against the intravenous administration of soluble mixed ET\textsubscript{A} receptor antagonists to improve the hypoplastic lung perfusion. The present results have a number of therapeutic implications. Together with functionally correct iPAVS, the backward flow in the PA is a physiological necessity to protect the pulmonary circulation from the right heart stream and, conversely, to supply blood to the fetal brain and placenta during the diastole. Since these backward flows must change to forward flows by the time of DA closure at birth, they may be deleterious in the case of impaired pulmonary vasculature, e.g., in CDH fetuses. Pulmonary hypoplasia occurs early in development when the lung is still in the terminal air sac phase and when growth retardation continues into the final 2 wk of gestation when the lung enters the alveolar phase. From a therapeutic point of view, it is unrealistic to hope for the recovery of the pulmonary proliferation in nonvascularized, nonalveolarized regions. On the other hand, less affected regions showing signs of neangiogenesis and alveolarization, even anarchic in nature, can be therapeutic targets, especially for attempting to initiate backward-to-forward conversion of pulmonary flows. As mentioned above, the hemodynamic effect of TO should be investigated with regard to forward-backward flow conversion. The poor effectiveness of ER blockade is disappointing. A possible alternative therapeutic approach to achieve hemodynamic objectives in addition to restoring vascularization and alveolarization in moderately affected pulmonary regions might consist of the implanting of a flow wave-filtering device (possibly resorbable) in the PA (5, 50). The purpose of such a device would be to restore the forward blood flow through the PA by blocking the backward flow. Further studies to design such a device are needed.

REFERENCES