Understanding Fetal Hemodynamics Using Cardiovascular Magnetic Resonance Imaging

Liqun Sun\textsuperscript{a} Davide Marini\textsuperscript{a} Brahmdeep Saini\textsuperscript{a} Eric Schrauben\textsuperscript{c} Christopher K. Macgowan\textsuperscript{c,d} Mike Seed\textsuperscript{a,b}

\textsuperscript{a}Division of Cardiology, Hospital for Sick Children, Toronto, ON, Canada; \textsuperscript{b}Department of Paediatrics and Medical Imaging, University of Toronto, Toronto, ON, Canada; \textsuperscript{c}Division of Translational Medicine, Hospital for Sick Children, Toronto, ON, Canada; \textsuperscript{d}Department of Medical Biophysics, University of Toronto, Toronto, ON, Canada

Keywords
Fetal hemodynamics · Cardiovascular MRI · MRI oximetry

Abstract
Human fetal circulatory physiology has been investigated extensively using grey-scale ultrasound, which provides excellent visualization of cardiac anatomy and function, while velocity profiles in the heart and vessels can be interrogated using Doppler. Measures of cerebral and placental vascular resistance, as well as indirect measures of intracardiac pressure obtained from the velocity waveform in the ductus venosus are routinely used to guide the management of fetal cardiovascular and placental disease. However, the characterization of some key elements of cardiovascular physiology such as vessel blood flow and the oxygen content of blood in the arteries and veins, as well as fetal oxygen delivery and consumption are not readily measured using ultrasound. To study these parameters, we have historically relied on data obtained using invasive measurements made in animal models, which are not equivalent to the human in every respect. Over recent years, a number of technical advances have been made that have allowed us to examine the human fetal circulatory system using cardiovascular magnetic resonance (CMR). The combination of vessel blood flow measurements made using cine phase contrast magnetic resonance imaging and vessel blood oxygen saturation and hematocrit measurements made using T1 and T2 mapping have enabled us to emulate those classic fetal sheep experiments defining the distribution of blood flow and oxygen transport across the fetal circulation in the human fetus. In addition, we have applied these techniques to study the relationship between abnormal fetal cardiovascular physiology and fetal development in the setting of congenital heart disease and placental insufficiency. CMR has become an important diagnostic tool in the assessment of cardiovascular physiology in the setting of postnatal cardiovascular disease, and is now being applied to the fetus to enhance our understanding of normal and abnormal fetal circulatory physiology and its impact on fetal well-being.

Background

Early investigations into the circulatory physiology of fetal sheep include those undertaken in the 1930s by Sir Joseph Barcroft at Cambridge University [1]. Sir Geoffrey Dawes elaborated on this work and conducted the first detailed studies of the fetal circulation [2]. Rudolph and
Heymann adopted the techniques pioneered by Barcroft and Dawes for exteriorizing and catheterizing fetal sheep and developed a method for making accurate measurements of the distribution of blood flow in fetal lambs using radioactively labelled microspheres [3]. Their measurements of fetal blood pressure, arterial and venous blood gases, absolute blood flow, and proportions of the combined ventricular output (CVO) delivered to the pulmonary, cerebral, and placental circulations, as well as measurements of the magnitude of the shunts across the foramen ovale, ductus arteriosus, and ductus venosus have provided us with our modern understanding of the normal fetal circulation [4]. The mammalian fetal circulation operates in parallel, not series, with shunts at ductus venosus, foramen ovale, and ductus arteriosus which allow blood returning from the umbilical vein to bypass the liver and lungs. The presence of these unique shunts results in the possibility for the right and left ventricles to have different cardiac outputs. In sheep, the right ventricle provides the more dominant contribution towards the combined ventricular output (CVO). By contrast with the postnatal circulation, the presence of the ductus arteriosus also results in equalization of blood pressure on both sides of the heart. Arterial oxygen saturations are considerably lower in the fetus than postnataally, with the highest oxygen saturations of approximately 85% found in the umbilical vein. The preferential flow, or streaming, of oxygenated blood from the umbilical vein and ductus venosus towards the foramen ovale results in higher saturations in the left heart compared with the right. Hypoxicemic blood delivered to the right heart is directed back to the placenta via the ductus arteriosus, descending aorta, and umbilical arteries, while the relatively low oxygen content of the blood supplied to the fetal lungs is largely responsible for the vasoconstricted state of the pulmonary circulation.

Fetal sheep studies have also been used to investigate fetal circulatory adaptations to acute hypoxia, whereby the presence of fetal shunts and differential changes in the cerebral, coronary, pulmonary, and systemic circulations result in redistribution of blood flow to favor conservation of myocardial and cerebral oxygen delivery at the expense of the musculoskeletal system, abdominal viscera, and lungs [5]. This response, which is detectable in humans using ultrasound through changes in cerebral arterial pulsatility, is frequently referred to as “brain-sparing physiology” [6]. However, in the setting of chronic hypoxemia, there is some return to more normal flow distribution (and cerebral arterial Doppler velocity profiles) with secondary adaptations to reduced oxygen delivery including the slowing of growth and reductions in fetal activity [7]. These secondary adaptations are associated with reductions in fetal oxygen consumption, which presumably serve to protect the fetus against hypoxic injury.

Over recent decades, magnetic resonance imaging (MRI) techniques for assessing cardiovascular function and hemodynamics in postnatal patients have played an increasingly important role in clinical practice [8]. Anatomical cine cardiovascular magnetic resonance (CMR) provides the most accurate means to measure ventricular volumes and ejection fraction, while cine phase contrast flow assessment represents the noninvasive gold standard for the measurement of vessel flow, providing unique quantification of valvular regurgitation and shunt volume [9]. In conjunction with invasive pressure measurements, this approach allows for the precise measurement of systemic and pulmonary vascular resistance [10].

CMR provides approaches to myocardial tissue characterization, including the quantification of regional and diffuse myocardial fibrosis as well as techniques for measuring myocardial perfusion. The opposing magnetic properties of oxy- and deoxy-hemoglobin can be exploited for direct visualization and quantification of tissue and blood oxygenation [11]. One approach to magnetic resonance oximetry based on blood relaxometry involves the application of a T2 preparation sequence for the quantification of T2 combined with a modified Look-Locker inversion recovery (MOLLI) sequence for measuring the T1 of blood [12]. The combination of any pair of vessel T1 and T2 measurements provides an accurate measurement of oxygen saturation and hematocrit in vitro, although in vivo validation of this approach to fetal oximetry has not yet been provided. The measurement of the T2 of flowing blood is challenging, and requires careful attention to the refocusing interval, choice of T2 preparation times, and consideration of artifacts arising from turbulent flow, local variations in the magnetic field, and partial voluming. However, assuming these challenges can be overcome and ignoring the negligible contribution of dissolved oxygen to the oxygen content of fetal blood, oxygen content can be quantified as the product of oxygen saturation and hematocrit by 1.36 (the amount of oxygen in milliliters bound to 1 g of hemoglobin). Fetal oxygen delivery can then be calculated as the product of umbilical vein oxygen content and umbilical blood flow, while fetal oxygen consumption is the product of umbilical venous flow by the umbilical vеноarterial difference in oxygen content.

The adaptation of CMR techniques for use in fetal subjects has required technical development to counter arti-
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Human fetal MRI oximetry data therefore support the presence of the same streaming of oxygenated blood from the umbilical vein to the left heart via the ductus venosus and foramen ovale that we have demonstrated in fetal sheep using 4D flow MRI (a volumetric cine phase contrast MRI technique) [27]. The hemodynamics of the ductus venosus result in a remarkable mechanism whereby two columns of blood moving at different speeds and with different oxygen saturations coexist in the suprapericardial inferior vena cava. The well-oxygenated blood returning from the placenta via the ductus venosus is directed toward the foramen ovale, presumably to ensure an adequate supply of glucose and oxygen to the brain and heart, the most metabolically active organs in utero. Streaming also results in transport of more deoxygenated blood back to the placenta via the ductus arteriosus, descending aorta, and umbilical arteries. The deoxygenated blood that passes into the right heart and pulmonary trunk is also presented to the pulmonary circulation, where its relatively low oxygen content maintains the high pulmonary vascular resistance typical of the fetal circulation [4] (Fig. 2).

Fetal Congenital Heart Disease

Using a combination of MRI blood flow and oximetry measurements, concepts regarding the impact of cardiac malformations and placental dysfunction on fetal circulatory physiology have been examined. Mean vessel flows and oxygen saturations obtained using the MRI techniques described above in a preliminary group of fetuses with a range of more severe forms of congenital heart disease (CHD) reveal many of the expected changes in fetal circulatory physiology [26]. For example, when one side of the heart is obstructed, there is a compensatory increase in blood flow through the unobstructed side. For example, in hypoplastic left heart syndrome, ascending aortic flow is significantly reduced, while flow across the main pulmonary artery and ductus arteriosus is markedly increased. Similarly, in lesions characterized by right heart obstruction like tetralogy of Fallot and tricuspid atresia, we have observed significant increases in ascending aortic flow with diminished flow in the main pulmonary artery and arterial duct. In transposition of the great arteries we have noted a reversal of the normal relationship between main pulmonary artery and ascending aortic flows. In the normal fetal circulation, main pulmonary artery flow exceeds ascending aortic flow. This is possible in the fetal circulation because of the presence of shunts at the foramen ovale and ductus arteriosus, and presumably reflects the greater venous return to the right ventricle. In transposition, ascending aortic flow is higher.

Human Fetal Circulatory Physiology by MRI

Normal Fetal Circulation

Using a combination of blood flow measurements made with cine phase contrast MRI with metric optimized gating and oxygen saturation measurements made using T2 mapping, we have collected preliminary reference ranges for the distribution of blood flow and oxygen transport in the late-gestation human fetus [26]. The results are in keeping with previous estimates regarding the human fetal circulation made by Rudolph which were based on his invasive measurements made in fetal sheep [4]. Pulmonary and cerebral blood flow are higher in the human fetus compared with the sheep fetus, the latter being readily explained by the larger size of the human brain compared with the sheep. Umbilical flow is lower in the human, although fetal oxygen delivery is similar due to the higher fetal hematocrit levels in the human [4] (Fig. 1).

In common with the sheep fetus, the oxygen saturations of the blood in the late-gestation human left heart and ascending aorta (65%) are approximately 15% higher than in the right ventricle and pulmonary artery (50%). Human fetal MRI oximetry data therefore support the
than main pulmonary artery flow, likely due to the discordant ventriculoarterial connections that result in the aorta being connected to the dominant right ventricle. Despite these variations in the outputs of the right and left heart, fetal organ perfusion appears to be reasonably well maintained. For example, pulmonary blood flow is rather stable across all of these forms of CHD, and similar to pulmonary blood flow in normal controls. If superior vena caval flow is taken as a surrogate for cerebral blood flow, then cerebral blood flow is also well maintained in most forms of CHD. The exception to this may be in those fetuses with more severe forms of Ebstein’s anomaly. Fetal Ebstein’s anomaly is associated with the lowest CVOs we have observed in fetuses with CHD, and appears to be prone to lower cerebral blood flow. Other types of CHD with single ventricle physiology are also associated with a reduction in CVO in the range of 10–20%. While cerebral and pulmonary perfusion is relatively well maintained, this reduction in cardiac output is associated with a drop in umbilical blood flow.

We hypothesized that the changes in fetal brain metabolism and reduced newborn brain size associated with CHD, as demonstrated by Limperopoulos et al. [28], result from reductions in cerebral oxygenation arising from interruption of the normal streaming of oxygenated blood from the placenta to the fetal brain. Our findings confirmed an average 10% reduction in the oxygen saturation of blood supplied to the developing brain in subjects with complex CHD [29]. The mechanisms leading to this aortic desaturation depend on the cardiac anatomy. For example, in fetuses with transposition of the great arteries, we found that the normal streaming of blood
from the ductus venosus to the left atrium results in well-oxygenated blood being directed to the pulmonary circulation, while the blood supplied to the brain is derived largely from more deoxygenated blood returning from the caval veins (Fig. 3). In fetuses with tetralogy of Fallot, the oxygenated blood crossing the foramen ovale passes into the left ventricle and aorta in the normal way but is diluted by more deoxygenated blood shunting from right to left across the ventricular septal defect. In hypoplastic left heart syndrome, no streaming of oxygenated blood is possible because there is essentially only one outlet from the heart, so the entire fetal circulation is supplied by blood with the same oxygen content. In addition to the disruption of streaming of oxygenated blood toward the brain, in single-ventricle hearts oxygen delivery to the fetus is diminished through a reduction in CVO. For example, in fetuses with more severe forms of Ebstein’s anomaly, fetal CMR reveals a reduction in CVO of up to 50% compared with normal fetuses [30, 31]. While Ebstein’s anomaly subjects appear to have reasonably normal placental function, their lower CVO results in reduced umbilical flow which results in reduced fetal oxy-
gen delivery. Although fetuses with Ebstein’s anomaly demonstrate increases in oxygen extraction, which partially compensates for the reduced oxygen delivery, body and brain growth remains impaired. In addition to the reductions in umbilical flow seen in fetuses with single-ventricle hearts, CHD appears to be associated with modest reductions in the oxygen content of umbilical venous blood. This may reflect abnormal structural differences that have been described on pathologic examination of the placenta in pregnancies affected by fetal CHD [32–34]. It is interesting to consider whether underdevelopment of the fetal placental vasculature resulting from impaired fetal perfusion of the placenta may result in a unique form of placental insufficiency.

Our fetal CMR findings in patients with CHD appear to support the concept that the disordered prenatal brain growth that is typical of CHD could be caused by the disruption of normal fetal cardiovascular physiology resulting in diminished substrate delivery. Recent fetal brain blood oxygen level-dependent (BOLD) MRI data are also in keeping with reductions in cerebral tissue oxygenation in fetuses with CHD [35]. However, recent deep genetic analyses have revealed widespread DNA damage in patients with CHD, which is more severe in the setting of associated noncardiac organ system malformations and neurodevelopmental delay [36]. Interestingly, some of these de novo mutations are also found in children with neurodevelopmental problems having normal hearts. These results have led to speculation that the neurodevelopmental problems commonly affecting nonsyndromic children with CHD are more likely to be genetic in origin rather than the result of abnormal fetal physiology or postnatal hemodynamics and brain injury. One approach to testing this hypothesis is to attempt to augment fetal brain oxygenation using maternal hyperoxygenation in fetuses with CHD. Using T2 mapping, we observed significant increases in the oxygen content of human umbilical venous blood during an episode of acute maternal hyperoxygenation, raising the possibility that cerebral oxygenation could be improved in fetuses with some forms of CHD through inhaled maternal supplemental oxygen [37]. A trial investigating the potential neuroprotective efficacy of chronic maternal hyperoxygenation in fetuses with single-ventricle CHD is being conducted at our center, although one study of chronic intermittent maternal hyperoxygenation in the setting of left ventricular hypoplasia found a reduction in brain size in fetuses treated with oxygen [38].

**Fetal Growth Restriction**

CMR has been used to study placental and fetal cardiovascular physiology in intrauterine growth restriction (IUGR) [39]. In fetuses with late-onset IUGR, umbilical vein oxygen saturation and flow are reduced, resulting in diminished fetal oxygen delivery, which is associated with lower fetal oxygen consumption. IUGR fetuses have lower oxygen saturations in the ascending aorta and higher superior vena caval flow, in keeping with fetal animal models of “brain-sparing physiology” (Fig. 4). By contrast, increased oxygen extraction and a more modest reduction in oxygen saturations in the main pulmonary artery are associated with reductions in pulmonary blood flow, in keeping with hypoxic pulmonary vasoconstriction. Although the correlation we observed between umbilical
vein oxygen saturation and flow is less striking, it is perhaps in keeping with the notion that placental disease is associated with stealing of diastolic blood flow into the cerebral circulation, high placental resistance, reduced umbilical flow, and impaired placental oxygen exchange.

Despite showing evidence of brain-sparing, newborns with IUGR have smaller brains than controls, in keeping with the brains of newborns with CHD. The relative importance of these delays in brain growth and maturation compared with postnatal physiologic and environmental factors has yet to be determined. However, the interpretation of studies investigating the relationship between placental insufficiency and neurodevelopmental outcomes has often been complicated by difficulties in distinguishing between milder forms of placental insufficiency and constitutional small size for gestational age. Of interest, in our study of late-onset IUGR, which used a composite scoring system incorporating fetal growth and Doppler parameters, placental histology, and neonatal nutrition to diagnose the presence of fetal growth restriction, 1/3 of the fetuses with late-onset IUGR had normal blood flow distribution by MRI and normal cerebroplacental ratio by Doppler. We believe that this group of fetuses, which likely have a very stable and milder form of placental insufficiency, may frequently go undetected in clinical practice.

The improved sensitivity of fetal MRI to chronic fetal hypoxia and its versatility for looking at placental, circulatory, and cerebral physiology and brain development may be helpful as new approaches to improving the outcomes of fetuses exposed to a hostile in utero environment are sought. A number of MRI methods for investigating placental physiology in human and animal models are currently being explored, with the broad aim of addressing the relatively poor in vivo characterization of placental pathology and limited options for treating placental disease that are currently available. Dynamic contrast-enhanced MRI has been used to quantify placental perfusion in animal models [40]. Other techniques including placental BOLD and magnetic resonance spectroscopy have also been used to explore placental function [41, 42].

**Future Developments**

The development of fetal CMR has provided a new research tool for studying fetal cardiovascular physiology and the impact of abnormal circulatory hemodynamics on fetal development [30, 32, 43–46]. Some centers have suggested there may be a clinical role for fetal CMR in the diagnosis of congenital malformations affecting the arterial and venous anatomy [22, 23]. Potential future clinical indications for fetal CMR also include improved diagnosis of fetal cardiac malformations, the monitoring of fetal therapy, and the assessment of placental function. Lloyd et al. [47] have explored the potential role of fetal CMR for the evaluation of complex cardiac anatomy, particularly in late gestation, when ultrasound imaging of the fetal heart may be hampered by adverse fetal lie, oligohydramnios, and reflection of the ultrasound beam by the
bony thorax. As the range of fetal treatments available broadens, the need for more advanced fetal cardiovascular imaging may emerge as a means of therapeutic monitoring. For example, cine phase contrast flow quantification has been used to demonstrate an increase in pulmonary blood flow following decompression of pulmonary venous obstruction in fetuses with hypoplastic left heart syndrome with restrictive atrial septum [45]. Similarly, improvements in fetal cardiac output have been demonstrated following transcervical nonsteroidal anti-inflammatory drug-induced ductal constriction in fetuses with Ebstein’s anomaly complicated by a circular shunt [31]. MRI has also been used to interrogate the potential to enhance the growth of underdeveloped left heart structures through the pulmonary vasodilatory action of maternal hyperoxygenation [46]. Finally, preliminary work on the development of MRI measurement of fetal hema-tocrit indicates that in future, decision-making about the need for repeat blood transfusions in the setting of fetal anemia could be guided by MRI [48, 49].

Fetal CMR techniques developed for human imaging may also be useful for studying animal models of human disease and for evaluating experimental fetal therapies in animal models. Very high quality measurements of blood flow and blood oxygen content are possible using a blood pressure waveform to trigger the MRI acquisition. The pressure trace is obtained using an indwelling fetal arterial catheter and imaging quality is optimized using maternal anesthesia to restrict maternal and fetal motion.

Examples of applications of fetal CMR that have been used to study animal models include the visualization of myocardial infaracts induced by coronary artery ligation and the quantification of the distribution of blood flow across the fetal sheep circulation [50, 51]. Thus, the fetal sheep physiology work that inspired the development of human fetal CMR may now be enhanced by this versatile imaging medium for examining cardiovascular physiology, which now provides an alternative to conventional invasive techniques.

**Statement of Ethics**

The research ethics board of the Hospital for Sick Children approved the study, and written consent was obtained from every mother prior to study enrolment.

**Disclosure Statement**

The authors declare that they have no conflicts of interest.

**Funding Sources**

None.

**Author Contributions**


**References**


