The Transitional Heart: From Early Embryonic and Fetal Development to Neonatal Life

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Abstract
Formation of the human heart involves complex biological signals, interactions, specification of myocardial progenitor cells, and heart tube looping. To facilitate survival in the hypoxemic intrauterine environment, the fetus possesses structural, physiological, and functional cardiovascular adaptations that are fundamentally different from the neonate. At birth, upon separation from the placental circulation, the neonatal cardiovascular system takes over responsibility of vital processes for survival. The transition from the fetal to neonatal circulation is considered to be a period of intricate physiological, anatomical, and biochemical changes in the cardiovascular system. With a successful cardiopulmonary transition to the extrauterine environment, the fetal shunts are functionally modified or eliminated, enabling independent life. Investigations using medical imaging tools such as ultrasound and magnetic resonance imaging have helped to define normal and abnormal patterns of cardiac remodeling both in utero and ex utero. This has not only allowed for a better understanding of how congenital cardiac malformations alter the hemodynamic transition to the extrauterine environment but also how other more common complications during pregnancy including intrauterine growth restriction, preeclampsia, and preterm delivery adversely affect offspring cardiac remodeling during this early transitional period. This review article describes key cardiac progenitors involved in embryonic heart development; the cellular, physiological, and anatomical changes during the transition from fetal to neonatal circulation; as well as the unique impact that different pregnancy complications have on cardiac remodeling. © 2019 The Author(s)
eral types of progenitor cells that are derived from the mesoderm, proepicardium, and neural crest. This eventually leads to the formation of the 4-chambered heart by gestational week 7 via heart looping and complex cellular interactions in utero [2]. Intrauterine fetal life is sustained in a secure, isolated, and hypoxemic environment that is dependent on the mother’s placenta for nutrition, respiration, waste elimination, and metabolism [6]. To facilitate survival in the hypoxemic intrauterine environment, the fetus possesses structural, physiological, and functional cardiovascular adaptations that are fundamentally different from the neonate [3, 4]. At birth, upon separation from the placental circulation, the neonatal cardiovascular system takes over responsibility of vital processes [6].

The transition from intrauterine to extraterine life requires well-orchestrated and complex biochemical, physiological, and anatomical changes in a timely manner to ensure neonatal survival [4]. However, any disruption in early development and transition can have long lasting, adverse, and sometimes fatal consequences (Fig. 1). While it is well known that congenital cardiac malformations alter the hemodynamic transition to the extraterine environ-
soderm posterior 1 and 2 (MESP1/2) are the earliest markers of cardiovascular specification and are expressed at the start of gastrulation [12, 13]. MESP1 represses pluripotent-inducing genes and drives expression of cardiac transcription factors [14]. As cardiac precursors migrate away from the primitive streak to form the cardiac crescent, MESP1/2 is downregulated while activating other transcription factor networks that drive cardiac specification. Early heart development involves myocardial progenitor cells, including first heart field (FHF) and second heart field (SHF) progenitor cells, as well as proepicardial progenitor cells derived from the lateral plate mesoderm and ectoderm-derived cranial neural crest cells (Fig. 2a). These cells emerge via the interaction of inductive and inhibitory signals from the germ layers (endoderm, mesoderm, and ectoderm) during gastrulation [2, 15, 16]. Molecular signals include cardiac differentiation inhibitors (i.e., wntes integrated, Wnt) and cardiac differentiation inducers (i.e., fibroblast growth factor, FGF and bone morphogenetic proteins, BMPs) [16–20]. Disruption in Wnt/β-catenin signaling in endoderm via deletion of β-catenin results in the formation of ectopic cardiac tissue in the overlying mesoderm [21].

**Formation of Cardiac Chambers: Heart Tube Formation and Looping**

The specification and differentiation of progenitor cells via specific heart fields are critical for embryonic heart development [2, 22]. During the second week of human gestation, the cardiac mesodermal cells migrate toward the anterior direction of the embryo to form the 2 major cardiac progenitor pools: the earliest cells to express MESP1 form the FHF or cardiac crescent and the later wave of MESP1 produces cells that form the SHF (located posterior to the crescent; Fig. 2b) [1, 16, 22, 23]. Both FHF and SHF express specific transcription factors and are the main reservoirs for cardiomyocytes, where mononucleated cardiomyocytes undergo hyperplasia in order to increase cell number for cardiac growth [1, 2, 16, 22, 24, 25].

The specification of cardiac chamber morphogenesis is modulated by several transcription factors, including T-box transcription factors TBX2, TBX3, TBX5, and TBX20; as well as NKX2–5, GATA4, and HAND1 [13]. GATA4 and NKX2–5 in combination with TBX5 and TBX20 promote myocardial specification via the expression of chamber myocardium-specific genes including induction of atrial natriuretic factor, gap junction protein connexins 40/43, and transcriptional repressor ID2, which improves contractibility and patterning of the right ventricular (RV) and left ventricular (LV) bundle branches [25–28]. BMP-induced BX2/3 expression is involved in non-chamber myocardial specification including the atrioventricular canal, outflow and inflow tract [29]. TBX20, an inhibitor of BMP-induced TBX2/3 activation, is broadly expressed throughout the linear heart tube to ensure chamber myocardium specification. Deletion of TBX20 results in defects in chamber development [30]. Furthermore, disturbance of morphogenetic transcription and growth factors including TBX1, TBX5, GATA4, and BMP4 within the SHF-derived cell populations may result in a spectrum of ventricular septal defects at the outflow tract [31]. Ventricular septal defects are characterized by membranous ventricular septum via the fusion of the endocardial outflow tract and atrioventricular canal, which allows blood to pass form the left-to-right side of the heart [32]. If significant in size and unrepaired postnatally, they can result in elevation in pulmonary vascular resistance (PVR), pulmonary arterial pressure, hypoxia, and cyanosis [33].

The SHF of the human embryo shows a dynamic spatio-temporal distribution pattern [34]. During the third week, the FHF fuses at the midline to form the primitive heart tube, which starts to beat at around embryonic day 22 and eventually gives rise to the LV and parts of the right and left atria [2, 16, 22, 24]. Blood flow and contraction causes peristaltic pumping motion in the linear heart tube, which may contribute to the ballooning process since endocardial and myocardial cells alter their shape, size, and proliferation in response to mechanical stress [35–37]. Subsequently, during week 4, the heart tube undergoes rightward looping, with its posterior region moving anteriorly [2, 15, 16, 23, 38]. SHF progenitors, located behind the primary heart tube within the pharyngeal mesoderm, migrate toward the primitive and looping heart tube, contributing to the RV, parts of the atria, septum and outflow tract, and later to the base of the aorta and pulmonary trunk [16, 34]. The cells from the venous poles contribute to the base of the superior and inferior vena cava (SVC and IVC, respectively) [16].

Cardiac trabeculation occurs after the cardiac looping, it promotes the formation of luminal projections (trabeculae), which consist of myocardial cells enclosed by the endocardial layer [39]. Trabeculation and subsequent compaction of the ventricular myocardium facilitates septation, increases cardiac output, contractility and conductivity, and helps establish the coronary circulation system in the developing heart [39–44]. NOTCH1 regulates trabecular growth via promoting endothelial growth factor neuregulin-1 activity through endocardial Ephrin B2 [41].
Fig. 2. Early cardiac development. a Cardiac cell lineage and specification during development demonstrating the commitment of pluripotent cells toward mature cardiac cell types within the heart development. b Schematic of cardiac morphogenesis in humans. At the second week of gestation, the cardiogenic mesodermal cells migrate toward the anterior side of the embryo to form the FHF or cardiac crescent and SHF that are specified to form specific segments of the PHT, which is patterned along the anteroposterior axis to form the various regions and chambers of the looped and mature heart during weeks 3 and 4. The FHF gives rise to the beating PHT and will eventually give rise to the LV and parts of the right and left atria (RA and LA, respectively). The SHF, located behind the PHT and within the pharyngeal mesoderm by gestational week 3, will contribute to the formation of the RV, parts of the atria and outflow tract, and later to the base of the aorta and pulmonary artery. At gestational week 3, the cells at the venous pole contribute to the formation of the superior and inferior vena cava (IVC, respectively). By gestational week 4, the cardiac neural crest cells migrate in from the dorsal neural tube, forming smooth muscle cells within the aortic and pulmonary arteries. In addition, the proepicardial organ formed by the proepicardial progenitor cell clusters later contributes to the formation of the epicardium. The 4 chambers form by the end of week 7. Wnt, Wingless integrated; FGF, fibroblast growth factor; BMP, bone morphogenetic proteins; FHF, first heart field; SHF, second heart field; OFT, outflow tract; PHT, primary heart tube; PM, pharyngeal mesoderm; VP, venous pole; CNCCs, cardiac neural crest cells; LV, left ventricle; RV, right ventricle; PEO, proepicardial organ; SVC, superior vena cava; IVC, inferior vena cava; EPC, epicardium; and PA, pulmonary artery.
The neuregulin-1 protein binds to the myocardial tyrosine kinase receptor erbB-4 (ERBB4) and dimerizes with ERBB2, thereby activating signaling cascades that modulate cell migration and proliferation. Mutations in ERBB2 result in compromised ventricular contraction [45].

Formation of the Epicardium, Heart Valves, and Parasympathetic Innervation

The embryonic pro-epicardial progenitor cells located at the celomic mesenchyme of the septum transversum will differentiate into cardiac fibroblasts, coronary vasculature, and a small number of cardiomyocytes, which eventually form the outer lining of the (epicardium) [1, 16, 46]. The fourth progenitor population involved in human embryonic cardiogenesis consists of the cranial neural crest cells, which migrate in from the dorsal neural tube through the pharyngeal arches via stromal cell-derived factor 1 as a chemotactic agent [16, 22, 47]. The neural crest cells are involved in septation of the outflow tract and the formation of heart valves, with NOTCH1 contributing to heart valve formation. Several gene mutations in NOTCH1 have now been reported in the human population to be associated with bicuspid aortic valve [32]. Neural crest cells also give rise to smooth muscle cells within aortic and pulmonary arteries, along with the full autonomic and sensory innervation of the heart [16, 22]. Anomalies in the cardiac neural crest are responsible for a multitude of human cardio-cranio-facial defects, such as DiGeorge syndrome, which is associated with TBX1 deletion [48, 49]. Through the looping process, complex interactions of FHF and SHF progenitors, as well as pro-epicardial and cranial neural crest cells, the fetal heart is septated into 4 defined cardiac chambers, which connect to the aorta and pulmonary trunk around gestational week 7 [2, 16, 22].

The Fetal Cardiovascular System

The fetal cardiovascular system is adapted biochemically and structurally to ensure that the highly oxygenated blood is delivered preferentially from the placenta to the brain and the heart while being diverted away from the lungs. The combination of the following contributes to the maintenance of the fetal cardiovascular system: (i) biochemical factors, including vasoregulatory agents such as prostaglandins and endothelin-1 and (ii) anatomical adaptations, such as the presence of 4 shunts: the placenta, ductus venosus (DV), ductus arteriosus (DA), and foramen ovale (FO; Fig.3a) [6, 50–52].

Fetal Biochemical Adaptations: Upregulation of Vasoconstrictors

Endothelin-1 plays a key role in utero-placental circulation by aiding the maintenance of high PVR in the fetal circulation [53, 54]. Expression of vasoconstrictive prostaglandin F2α during early gestation has also been observed to maintain the high PVR, while there is an increase in vasodilating prostaglandin I2 and E2 (PGI2 and PGE2, respectively) nearer to term to aid in the fetal to neonatal transition [5].

Fetal Structural Adaptations: The Four Shunts in Fetal Circulation

By the 10th week of gestation, the fetal circulation has transitioned from passive gas exchange by both the yolk sac and the placenta to being placenta dominant (Fig. 3a) [4]. The placenta is not only the source of nutrition but also an organ of waste elimination for the fetus; the supply of nutrients, as well as the exchange of oxygen and waste products, takes place in the intervillous space [6]. It is a low resistance vascular bed, which promotes the fetal-to-maternal exchange of deoxygenated blood under low pressure [3, 50]. Despite the low partial pressure of oxygen in the placenta, oxygen delivery to fetal tissues remains adequate due to the combined high ventricular output and the presence of fetal hemoglobin, which has a higher oxygen affinity than adult hemoglobin [50, 55, 56].

Following oxygenation in the intervillous spaces, relatively oxygenated fetal blood is carried by the umbilical vein to the liver [5, 57]. The DV allows 50–60% of umbilical vein blood to bypass the hepatic circulation and enter the IVC, mixing with desaturated IVC blood [6, 58]. The remainder of blood perfuses the liver, ultimately merging with desaturated blood from the lower part of the body via the IVC to enter the right atrium [6]. In the right atrium, blood is diverted into 2 streams, with more than half traversing the FO in the inter-atrial septum to enter the left atrium [23, 50, 51, 59]. This oxygen-saturated blood then passes through the LV and mixes with the pulmonary venous return to be pumped through the ascending aorta toward the carotid and coronary arteries [5, 23, 50, 60]. The remainder of IVC blood flow mixes with desaturated blood from the superior vena cava, which first enters the RV through the tricuspid valve and, subsequently, the pulmonary artery [5, 57]. The high PVR results in the majority of blood that leaves the RV being preferentially shunted through the DA, bypassing the pulmonary circulation and going directly into the descending aorta. As a result, only 8–10% of total cardiac output passes through the high-resistance pulmonary cir-
Fetal circulatory [5, 50, 57, 59, 61, 62]. Approximately 40–60% of blood that enters the descending aorta either supplies the umbilical artery to the lower limbs or is reoxygenated at the placenta [57].

**Fetal Combined Ventricular Output and Heart Rate**

Normal fetal heart rate ranges from 110 to 160 beats per minute (Table 1) [63], with the low resistance placental bed providing a low systemic vascular resistance (SVR) and absorbing over 40% of the combined cardiac output from both ventricles [6]. Increased PVR combined with low SVR results in the right-to-left shunting of blood through the FO and the patent ductus arteriosus (PDA), with the RV accounting for two-thirds of cardiac output perfusing the lower half of the body and placenta. LV output is directed toward the coronary and carotid arteries, supplying the heart muscle wall and brain [51, 57, 64–66]. The amount of blood entering the pulmonary circulation...
varies during pregnancy, increasing from just over 10% of the combined cardiac output at mid-gestation to around 25% by 30 weeks’ gestation [5]. Perfusion rate of fetal tissues is higher than in the adult, with normal ranges of 470–500 mL/kg/min measured by Doppler ultrasound (Table 1) [64].

**The Transition to Neonatal Circulation**

The neonatal period involves dramatic and rapid physiological changes [57]. The transition to the postnatal environment triggers cardiomyocytes to switch from a fetal hyperplastic (cell proliferation) to hypertrophic (increase in cell size) growth pattern [67, 68]. Hypertrophy of cardiomyocytes is achieved via karyokinesis without cytokinesis (mitotic cell division without cytoplasmic separation), resulting in the formation of multinucleated myocardial cells to promote myocardial growth [69–71]. As such, in the first few months of postnatal life, myocardial cell volume increases approximately 30-fold in humans. Cardiopulmonary adaptation at birth involves an intricate series of timely biochemical and structural modifications that are required for a successful cardiopulmonary transition from the fetal to neonatal circulation, which includes a switch in the site of gas exchange from the placenta to the neonatal lungs [6].

**Neonatal Biochemical Adaptations**

Changes in the activity of cyclo-oxygenase appear to be involved in adaptation to the extraterine environment [5]. After birth, there is an inhibition of cyclo-oxygenase-2 enzyme pathways, which results in a reduction of the vasoconstrictor thromboxane. After the third trimester, there is an upregulation of endothelial nitric oxide (NO) synthase, which results in greater synthesis of the vasodilator NO. As a direct consequence of the increased oxygenation associated with lung inflation, there is also a surge in NO at birth [5, 59]. NO causes smooth muscle relaxation and arteriolar vasodilation, leading to lower PVR [59]. Vasodilation is further encouraged by the increased secretion of vasodilating PGI2 and PGE2 [6].

Acute adaptations to postnatal life are also managed by catecholamine release [52]. During delivery, there is a surge in circulating catecholamines, which act to strengthen myocardial contractility and increase heart rate. This leads to sufficient elevation of cardiac output in order to meet the increased metabolic demands associated with spontaneous thermogenesis, feeding, and breathing [52, 57]. With increasing gestation, fetal cortisol and thyroxine levels also rise gradually until term gestation, further increasing during labor and peaking just after birth [52]. Cortisol is important for maturation of pulmonary surfactant, increasing catecholamine secretion, as well as increasing the density of vasodilatory β-adrenoreceptors in the heart and lungs [52]. Furthermore, both cortisol and thyroxine activate sodium pumps within the alveoli to facilitate lung fluid removal.

**Neonatal Structural Adaptations: Neonatal Circulation and Modified Shunts**

The transition from fetal to postnatal circulation requires the removal or modification of the 4 fetal shunts (placenta, DV, DA, FO) to eliminate the umbilico-placental circulation, resulting in a large increase in SVR (Fig. 3b, Table 1) [51, 52, 61]. As SVR rises and stabilizes, PVR declines and the right-to-left shunt through the DA reverses to become a left-to-right shunt. RV output is also directed fully to the pulmonary circulation, leading to an increase in biventricular stroke volume [51, 61, 72–74]. Any perturbations to the transitional process may result in neonatal cardiopulmonary dysfunction. For instance,
higher left atrial pressures \([3, 52, 59, 77, 78]\). This causes atrium resulting in a left-to-right shunt direction and nary blood flow and pulmonary venous return to the left the pulmonary artery into the lungs, increasing pulmo-
lar partial pressure of oxygen and oxygen saturation \([50]\).

Cutting the umbilical cord eliminates the low-resis-
tance placental flow and stimulates peripheral and central chemoreceptors, which combined with other factors (thermal, neuroendocrine, and mechanical stimuli), con-
tributes to the onset and maintenance of neonatal respira-
tory efforts. A subsequent increase in SVR and reduction in right heart preload results \([6]\), with the SVR continuing to increase due to cold stress and catecholamine surges \([76]\). Initiation of breathing stimulates a series of events that are responsible for the shift from fetal to postnatal circulatory patterns. Hypoxemia can occur in the neonata-
tal period via ventilation/perfusion mismatch, hypoven-
tilation (impaired ventilatory function and reduced respira-
tory compliance), inadequate pulmonary surface area (i.e., pulmonary hypoplasia, diaphragmatic hernia), and increased physiological dead space, including respiratory distress syndrome \([75]\).

Fetal production of lung fluid decelerates shortly be-
fore full-term birth, with labor providing a strong driving force for lung fluid removal \([3, 52, 59]\). With the resorp-
tion and removal of lung fluid from the alveoli, surfactant secretions by type II pneumocytes into the acinus support the retention of the first breaths of air into the newly in-
flated lung \([52]\). Lung aeration allows the establishment of alveolar surface tension, as well as negative interstitial and intrapleural pressures, thereby increasing the alveo-
lar partial pressure of oxygen and oxygen saturation \([50]\).

The physiological transition to the extraterine envi-
ronment increases the systemic arterial oxygen tension, and the decrease in circulating PGE\(_2\) (including placental prostaglandins) promotes the constrictive effect of oxygen on the ductal tissue \([50, 52, 81, 82]\). Furthermore, endothelin-1 constricts the DA in response to oxygen \([50, 61]\). In term-born babies, functional closure of the DA commences immediately after the umbilical cord is cut and may take between 1 and 4 days \([23]\). Over the subse-
quent 4–8 weeks, permanent anatomical closure occurs via endothelial proliferation and fibrosis, giving rise to the ligamentum arteriosum \([80]\). In some term infants and many preterm infants, the process of physiologic ductal closure can be maladaptive, delayed, or even ar-
rested \([80]\). If the situation is considered hemodynam-
ically significant, medical therapy will be required to aug-
ment closure of the DA, with possible surgical ligation depending on severity \([80]\). Despite closure (either spontaneous or with medical therapy), the hemodynamic environment may induce DA opening and the persist-
tence of the DA can have deleterious hemodynamic con-
sequences \([83]\). Immediately after birth, the DV ceases to function, resulting in complete perfusion of the liver and reduction in blood returning to the IVC. However, the structural closure may take between 3 and 7 days in a term-born baby \([52, 84]\). The structure disappears within 2 weeks after birth and ultimately becomes the ligamentum venosum \([23]\).

Echocardiography studies of healthy term neonates have expanded our understanding of the transitional physiology and hemodynamics \([72, 74]\). These studies have revealed that on day 2 postpartum, there is a small reduction in LV basal diameter, mitral valve inflow velocity time integral, and systolic velocity of the lateral wall and septum \([72]\). On the other hand, there is a small in-
crease on day 2 of life in RV dimensions including mid-
cavity diameter, RV antero-inferior basal diameter, and end-diastolic area. The differential adaptive physiological responses of the RV and LV, pulmonary hemodynamics, and shunt characteristics may relate to loading conditions and patent DA closure [72, 73]. Any perturbations to the transition process may result in dysfunction in neonatal cardiopulmonary function.

**Neonatal Cardiac Output, Heart Rate, and Blood Pressure**

With the functional closure of FO and DA, output from the ventricles equalizes and overall cardiac output increases by nearly double [51, 61]. The increase gradually returns to lower levels within the first 24 h [52]. At term, the neonatal cardiac output is approximately 200 mL/kg/min [85], which is more than twice that of adults relative to body weight [57]. However, the neonatal myocardium is stiffer with fewer myofibrils in a more disordered arrangement, with a low arterial systolic and diastolic blood pressure of 60 and 30 mm Hg, respectively. Based on the Frank-Starling relationship, this leads to a limited increase in stroke volume for a given ventricular filling volume [57, 86]. Thus, the neonatal myocardium is initially more dependent on heart rate to increase cardiac output, with heart rates around 160–180 beats per minute in the first 30 min following delivery and then stabilizing in the range of 100–160 beats per minute (Table 1) [6, 57]. Shortly thereafter, the LV hypertrophies as it takes over the systemic circulation and the RV remodels to being the thin-walled, crescent shaped chamber that supplies the lower pressure pulmonary circulation [3]. By 3 weeks’ postnatal age, pulmonary pressure has normally fallen below systemic pressure and by 3–6 months after birth, the classical LV dominant pattern of adulthood is established with increased SVR-stimulating ventricular hypertrophy [57].

**Impact of Pregnancy Complications on Early Cardiac Remodeling**

A number of clinical conditions and diseases exist that can affect transitional physiology and lead to long-term cardiovascular health complications, such as congenital heart diseases. While the full discussion of these diseases and conditions is beyond the scope of this review, herein we have focused on more common complications that affect transitional physiology and have been identified as novel long-term cardiovascular risk factors.

**Intrauterine Growth Restriction**

IUGR fetuses do not grow according to their genetic growth potential, which can be caused by impaired placenta including utero-placental vascular insufficiency [7, 87]. When over a third of the fetal villous vasculature is abnormal, the Doppler resistance indices indicate an increase in impedance to flow and resistance in the umbilical artery [88]. In severe cases, where >60% of the villous tree is damaged, it may couple with absent end-diastolic flow or reversed end-diastolic flow, promoting fetal distress when the gaseous exchange in the intrauterine environment has deteriorated [88–91]. In monochorionic twins, selective IUGR may occur, which is when unequal sharing of the placental mass results in poor fetal growth in one of the twins [92]. The use of fetal ultrasound allows for identification of the velocity, direction, and patterns of blood flow to diagnose selective IUGR, as well as to differentiate it from twin-twin transfusion syndrome. Unlike selective IUGR, in twin-twin transfusion syndrome, there is a large inequality in blood sharing between the monochorionic twins via placental blood vessel connections, such that one twin acts as the donor and the other as the recipient. This results in changes in urine output and amniotic fluid volume, which can lead to fetal hydrops in the recipient due to volume overload [92].

IUGR offspring demonstrate early cardiac changes that emerge during fetal life and persist into neonatal life, including a relatively hypertrophied ventricular septum, LV dilatation, decreased myocardial reserve, a more globular cardiac morphology, and abnormal early postnatal hemodynamic adaptations [8, 93–95]. Furthermore, IUGR neonates exhibit an increase in left myocardial performance index (a marker of ventricular dysfunction), an inability to increase LV stroke volume, as well as reduced heart rate in the first week of postnatal life when compared with appropriate-for-gestational-age controls [8, 96]. Changes in cardiomyocyte functionality including altered contractile machinery in the form of shorter sarcomere length and delayed cardiomyocyte maturation may help to explain the cardiac dysfunction associated with IUGR [97, 98]. It is likely that these early changes are of long-term clinical significance, given that epidemiologic studies have consistently shown an association between IUGR and an increased rate of cardiovascular mortality in adulthood [99]. In line with this, it has been demonstrated that many of the cardiovascular changes that first emerge during fetal life, including cardiac morphological changes, subclinical myocardial dysfunction, arterial remodeling, and impaired endothelial function, persist long-term [94].
Preeclampsia

Preeclampsia is a common hypertension-related pregnancy complication that is characterized by proteinuria and maternal organ dysfunction (such as renal insufficiency, liver, and placental dysfunction) post 20 weeks' gestation, affecting over 8% of pregnancies worldwide [9, 100]. Preeclampsia results in utero-placental hypoxia, where the maternal blood oxygenation is normal, but the utero-placental circulation is impaired [101–103]. This can be induced via shallow placenta implantation due to restricted extravillous trophoblast invasion [104, 105]. Hypoperfusion of the placenta may result in growth restriction of the fetus and oligohydramnios, which often lead to preterm delivery of the fetus [106–109].

Exposure to a hypoxic intrauterine environment is associated with offspring vascular, metabolic, and cardiac modifications including aortic wall thickening, loss of NO vessel modulation, altered LV function, and increased susceptibility to ischemia and metabolic syndrome [110–112]. Recently, it has become apparent that offspring of hypertensive pregnancies may also have microvascular genotypic and phenotypic abnormalities that may persist beyond the perinatal period and into adulthood [113–116], which may account for their significantly lower threshold for the development of cardiovascular-related diseases [110]. While studies assessing the specific impact of preeclampsia on cardiac remodeling during the transition from fetal to neonatal life remain limited, further investigations are needed given that adolescents and young adults who were exposed to maternal preeclampsia display unique LV remodeling patterns including greater relative wall thickness, smaller internal cavity diameter, and lower longitudinal peak systolic strain [112, 117].

Preterm Birth

Preterm birth prior to 37 weeks’ gestation is one of the leading causes of morbidities and mortality [118], though survival rates have continued to increase due to advances in prenatal and postnatal care [119–121]. It is a complex condition resulting from multiple etiologic pathways including preeclampsia, multiparity, and IUGR, though the majority of preterm births are spontaneous with unknown causes [122–124]. Given that premature infants begin extrauterine life prior to maturation of the fetal circulation, preterm neonates often possess many immature functional and structural characteristics [3, 67]. The immature cardiovascular circulation increases susceptibility to a suboptimal fetal to neonatal transition [8, 52, 125].

In addition, in preterm neonates, the immature DA muscular wall alongside elevated levels of circulating PGE2 and NO levels often results in an unresponsive ductal constriction and subsequent delayed ductal closure [6, 52, 126]. The DA remains open for >4 days of age in 10% of preterm neonates born between 30 and 37 weeks’ gestation; 80% of those born between 25 and 28 weeks’ gestation; and 90% of those born between at 24 gestational weeks’ gestation [3, 52]. In term-born neonates, functional closure usually occurs within the first 3 days of postnatal life [127–129]. The FO can also remain patent in many premature infants and can have adverse consequences on pulmonary hyperperfusion and systemic hypoperfusion due to the increase in pulmonary flow, left-to-right shunting, and sustained ductal patency [3].

Bronchopulmonary dysplasia (BPD) is the most common morbidity of preterm infants born <30 weeks’ gestation and is the most common lung disease of infancy [130]. The pathophysiology is characterized by interrupted lung development and dysmorphic pulmonary capillaries [131]. Over 30% of infants with BPD will develop pulmonary hypertension, including nearly 50% of infants with severe BPD [132, 133]. Those who survive generally demonstrate resolution of pulmonary hypertension concordant with pulmonary vascular growth. However, longitudinal studies of BPD survivors have shown a life-long compromised pulmonary function, as well as an increased risk of RV dysfunction and possible recurrent pulmonary hypertension in adulthood [130].

Preterm birth, even in the absence of BPD and other related comorbidities, can affect myocardium structural development via altering cardiomyocyte maturation. In animal studies, the early exposure to a high resistance and hypoxic ex utero environment causes the immature hyperplastic fetal cardiomyocytes to undergo significant hypertrophy, increased interstitial collagen deposition, and abnormal functional properties during postnatal development [67, 68]. Similarly, recent studies in humans using echocardiography and MRI have highlighted that preterm infants display disproportionate postnatal hypertrophy, accompanied with a reduction in LV diastolic function and a persistent reduction in RV systolic function [10, 134, 135].

The consequences of exposure of the immature heart to the extrateruterine environment persist beyond the neonatal period. Studies using echocardiography and MRI have highlighted the early impact on long-term cardiovascular development associated with prematurity, with potentially adverse remodeling in infants, children, adolescents, and young adults [10, 112, 135–137]. In addition to the unique geometric and functional cardiac changes at rest, the pre-
term cardiac phenotype is also characterized by a reduced myocardial functional reserve [138], which together may explain their increased risk of heart failure as early as childhood and into young adulthood [34, 139–141].

**Conclusion**

The development of the heart begins as early as the third week of gestation with the 4-chamber fetal heart formed by gestational week 7. It involves complex biochemical signals, interactions, and specification of myocardial progenitor cells and heart tube looping. The transition from the fetal to neonatal circulation is considered to be a period of intricate physiological, anatomical, and biochemical changes in the cardiovascular system. With a successful cardiopulmonary transition to the extraterine environment, the fetal shunts are functionally modified or eliminated, enabling independent life. Early and accurate diagnosis of fetal congenital heart defects and investigations into the impact of pregnancy complications on the offspring are now feasible with the usage of medical imaging tools including ultrasound and MRI. Identifying abnormalities in early cardiogenesis and the cardiovascular transitional physiology from fetal to neonatal life is essential for the immediate and long-term cardiovascular health risk of neonates.

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