The Ductus Arteriosus, a Vascular Outsider, in Relation to the Pulmonary Circulation

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Abstract

The muscular ductus arteriosus (DA) has many unique characteristics setting it apart from the adjoining elastic arteries. Preparation for neonatal closure takes place in utero with the development of intimal thickening. Ductus-specific gene and protein expression patterns were demonstrated during this process. We postulated that the closing process, with increased expression of progerin, might reflect aspects of premature ageing. Persistent patency of the DA, can be congenitally or immaturity based. During embryonic development the sixth pharyngeal arch arteries are the last to develop. In avian embryos this arch is divided into a proximal (part of the future pulmonary artery) and a distal part (on the left side the origin of the DA). The consequence is that the pulmonary arteries have a double vascular contribution being a proximal sixth arch artery component and
a distal true pulmonary artery segment. We have conclusive evidence that this is not encountered in the human embryo. Both sixth arch arteries and the right and left pulmonary arteries connect separately and at distinct locations to the pulmonary trunk side of the aortic sac. The insertion of the wall of the DA between the proximal and distal part of the left pulmonary artery, referred to as pulmonary coarctation, constitutes a congenital malformation and not a remnant of what is encountered during normal development. This configuration is found almost exclusively in combination with pulmonary atresia leading to neonatal interruption of the proximal left pulmonary artery with consequences for surgical or intervention repair.

**Keywords**

Vascular development · Patent ductus arteriosus · Persistent ductus arteriosus · Congenital heart disease · Pulmonary atresia · Pulmonary ductal coarctation · Juxtaductal pulmonary coarctation

### 33.1 The Normal Closing Ductus Arteriosus

The DA is a functional shunt before birth, directing oxygen-rich blood to the systemic circulation, bypassing the pulmonary circulation. Perinatally, the DA shows a physiological contraction followed by anatomical closure after birth. Eventually the closed DA is remodeled into a fibrous strand, the ligamentum arteriosum [1]. The characteristics of the muscular DA are reflected in its histological make-up, setting it apart from the elastic adjoining arteries [2]. Around 17 weeks of human gestation there is the start of formation of typical intimal thickenings (cushions). The first indication is the subendothelial appearance of hyaluronic acid and detachment of the endothelium [3]. The process of hyaluronic acid production continues during ductal maturation and even after birth [4]. In the second half of gestation the maturation process progresses and marked intimal thickenings develop containing media-derived synthetic smooth muscle cells and fine elastin fibers [5]. Accompanying the first physiological contractions the DA develops degeneration of the inner media, showing apoptosis and extracellular matrix filled mucoid lakes [5]. A recent study shows that this process is accompanied by a different lamin A/C to progerin balance. Progerin, an indicator of vascular ageing, is increased in the mature DA [6]. This observation has lead us to postulate that a process akin to premature ageing might accompany postnatal ductal closure [6]. In Fig. 33.1, schematic and correlating histological features of the DA are presented.

More advanced methodology has allowed to shift focus to the unique gene and protein profiles of the DA. The data derive from DNA microarray profiling [7], transcriptional profiles [8] and gene profiling after single-cell microcapture isolation [9]. These studies unravel pieces of the puzzle underlying DA closure and maturation. Results show that the important relaxing factor, PGE2, inhibits elastogenesis, clarifying the muscular nature of the DA [10]. Furthermore the endothelin receptor EP4 promotes hyaluronic acid–induced intimal thickening [4].
Fig. 33.1 Schematic presentation of normal closure, delayed closure and persistent patency of the ductus arteriosus with corresponding histology and histopathology (Modified after [1]).
With increased information, showing the distinct characteristics of smooth muscle and endothelial cells (Fig. 33.2) we gain insight into the pathological and delayed DA closure.

### 33.2 Persistent Patency Versus Delayed Closure of the DA

Several causes lead to a non-adequate to non-closure of the DA. The most common cause is delayed closure after premature birth because of immaturity of the ductal wall [5]. A different cause for ductal patency is a structural wall abnormality inhibiting anatomical closing. We have described that a DA with an additional and marked subendothelial elastic lamina (Fig. 33.1) is not capable of permanent anatomical closure [2]. As this anomaly can only be detected by histology, it poses a problem to differentiate it clinically from delayed closure. Very little is known about the genetic or environmental background of persistent DA. A few genes have been discovered like the TFAP2B gene mutation [11] that lead to persistent patenty.

### 33.3 Normal Sixth Pharyngeal Arch Formation and Connections to the Surrounding Arteries

During development, pharyngeal arch arteries connect the aortic sac bilaterally to the dorsal aortae. The sixth arch, important for DA and pulmonary artery formation, has a separate status [12]. In various animal species it has been described that the putative left and right pulmonary arteries connect end to side to the sixth pharyngeal arch artery [13]. We have shown in quail embryos that this connection results from ingrowth of the pulmonary sprout into the sixth arch artery opposed to an outgrowth towards the lung mesenchyme [12]. This implies that the proximal part of the sixth arch artery forms the proximal part of the left pulmonary artery while the distal end develops the
characteristic DA structure. This necessitates a major remodeling process to achieve the mature situation in the human fetus and neonate in which the DA and left and right pulmonary arteries have a distinct connection site with the pulmonary trunk (Fig. 33.3). Investigation of human embryos, before 6 weeks of age, revealed that the ingrowth of the pulmonary arteries together with the proximal part of the sixth arch is incorporated into the pulmonary side of the aortic sac. Thus the distal part of the left-sided sixth pharyngeal artery develops into the left-sided DA. The distal right-sided sixth arch disappears together with the distal part of the right-sided dorsal aorta early in development by selective apoptosis [14]. These observations in the human embryo are relevant for understanding some forms of congenital heart disease and related complications.

33.4 Pulmonary Coarctation (PC) and Proximal Left Pulmonary Artery Interruption

DA problems can exist in right heart congenital heart disease with narrowing to atresia of the right ventricular outflow tract and hypoplasia of the pulmonary trunk. The most extreme is complete absence of the DA with the question whether a sixth arch artery ever developed. The VEGF120/120 mouse revealed that with increase of
mild pulmonary stenosis towards complete atresia the DA became hypoplastic to atretic. In the most severe cases major aorto-pulmonary collateral arteries (MAPCAs) developed [15]. We published data on a set of human neonatal specimens with various degrees of pulmonary stenosis and atresia that had developed a PC [16]. We assumed that this anomaly mimicked the aortic coarctation in which DA tissue extends within the inner circumference of the aortic arch in which the roof of the coarctation always consisted of elastic aorta tissue (Fig. 33.2). More recent observations showed that PDC could lead to a complete obstruction of the proximal left pulmonary artery. DA tissue was, however, not only observed on the inside lining of the pulmonary artery but could completely encircle the lumen [17], being thus interposed between the proximal and distal part of the left pulmonary artery. Progressive narrowing by DA tissue could result in left pulmonary artery interruption. We concluded that a more left lateral insertion of the DA into the left pulmonary artery is not a normal position but a congenital anomaly. Reversed hemodynamics through the DA might play a secondary role as cause for PC.

### 33.5 Future Directions and Clinical Implications

- Is PC exclusively linked to anomalies with pulmonary atresia or can it also be associated with less severe obstructions of the right ventricular outflow tract and pulmonary valve?
- Is there a link to the 22q11 deletion syndrome and thus to mutations within the second heart field and its interaction with neural crest cells?
- A clinical implication of the complete encircling of the DA tissue of the left pulmonary artery is that by surgical restoration of the left pulmonary artery it should be aimed to completely excise the DA tissue or completely stent the DA-infested region.

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