CLINIC AND THERAPEUTIC ASPECTS IN DUCTUS-DEPENDENT CONGENITAL HEART DEFECTS – PART I

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ABSTRACT

Congenital heart defects with ductus-dependent circulation have a complex cardiovascular physiology. After birth, the condition of these children is critical, they are asymptomatic and their pulmonary vascular resistance decreases leading to progressive hemodynamic deterioration. Early identification of these defects and the start of appropriate therapy allow to maintain the cardiovascular balance.

Keywords: congenital heart defect, ductus-dependent systemic circulation, persistence of ductus arteriosus, E1 prostaglandin

Congenital heart defects with ductus-dependent circulation are defined as abnormalities, in which the permeability of the ductus arteriosus is mandatory in order to maintain systemic perfusion. Ductus arteriosus permeability is given by the presence of an abnormal communication between the proximal descending aorta and the vicinity of left branch of the pulmonary artery (1).

In the womb, gas exchanges and metabolic reactions occur in placenta, shortcutting the lungs. First breath makes them fill with air and leads to increased pulmonary flow and later cutting the umbilical cord leading to increased systemic vascular resistance and lowering the pulmonary one resulting in the closure of the three shunts. If these events do not occur, the fetal circulation will continue after birth (15-17).

Transition circulation and fetal shunts closure occur as in healthy new-borns even if the new-born has a ductus-dependent heart defect, initially it is asymptomatic (5).

Later, through progressive narrowing of the ductus arteriosus appear the signs reflecting the systemic hypoperfusion (5,7):
• pallor;
• coldness;
• weak felt pulse;
• tachycardia;
• tachypnea;
• hypotension;
• anuria;
• progressive acidosis;
• cardiovascular Collapse.

Ductus-dependent congenital heart defects are divided into ductus-dependent systemic and pulmonary.

Ductus-dependent systemic circulation diseases are:
• critical aortic valve stenosis in the neonate;
• hypoplastic left heart syndrome;
• aortic coarctation;
• interrupted aortic arch.

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CRITICAL AORTIC STENOSIS

When the aortic orifice surface is reduced to less than a quarter of the normal value, overcoming the obstacle and provision of ventricular ejection is achieved by increasing highest systolic pressure in left ventricle to overcome the pressure in the aorta.

Pressure overload causes hypertrophy of left ventricle and gradually left heart failure installs which in time becomes global (8). Hypertrophied left ventricle becomes noncompliant and the diastolic pressure grows causing pulmonary edema.

In critical aortic stenosis, the left ventricle is unable to provide adequate antegrade cardiac output, systemic circulation being ensured by the right ventricle through ductus arteriosus directly into the aorta.

The aortic arch, cerebral and coronary circulation are infused backwards with blood with relatively low oxygen saturation.

As the ductus arteriosus closes, systemic perfusion is compromised realizing a low cardiac output. In this moment, clinical signs of systemic hypoperfusion appear, namely, pallor, tachypnea, tachycardia and hypotension (8.9).

HYPOPLASTIC LEFT HEART SYNDROME

It is a set of heart abnormalities that cause the obstruction of the outflow tract of the left ventricular outflow tract. These abnormalities include: stenosis/ataresia or aortic valve orifice, ascending aorta/aortic arch hypoplasia, hypoplasia or mitral valve stenosis, hypoplasia of left ventricle (2).

Adequate systemic perfusion is ensured by the permeable ductus arteriosus. The flow of blood ejected from the right ventricle gets into the pulmonary arteries through the ductus arteriosus directly into the aorta. From the aorta blood advances backwards through the descending aorta into the lower half of the body and with a relatively low blood oxygen saturation and in the upper half of the body also through the descending aorta to the coronary arteries.

Progressive narrowing of the channel lowers blood pressure, multiorgan hypoperfusion, including acute coronary myocardial hypoperfusion and acidosis, and if ductus arteriosus closes, the prognosis is unfavorable, and the baby dies (5).

After birth, functional signs are minor. With the narrowing of the ductus arteriosus during their first weeks of life, the child’s condition worsens, with the emergence of cyanosis, tachypnea, tachycardia, hypotension, weakly perceptible peripheral pulse or absent, rapidly developing pulmonary edema (3.6). Metabolic acidosis and hypoglycemia appear, indicating systemic hypoperfusion.

CRITICAL AORTIC COARCTATION

Heart congenital defects characterized by narrowing and obstruction of the aorta in the descending part and often in the isthmus.

Aorta obstruction has the following consequences:

- In proximal areas of the obstacle, the hemodynamics is accompanied by increased pressure in the arteriolar bed of the cephalic end, upper limbs and pulmonary bed with left ventricle overload (5,10).
- In post-ductal regions (the arteries supplying the lower limbs and abdominal organs) occurs vascular hypotension, vascularization being ensured by a network of collaterals between subclavian arteries and mammary arteries (5,10).

In ductus arteriosus closure, the entire cardiac output must pass through the bottleneck to the descending aorta. When the coarctation is severe (critical), the left ventricle cannot provide adequate antegrade flow into the descending aorta, therefore, the systemic flow to the lower end of the body is ductus-dependent. (11,12)

In severe forms of the disease, signs of life appear in 7-10 days, through signs of congestive heart failure brutally installed and clinically presented through: tachypnea, cyanosis (especially in the legs), cold lower extremities weakly perceptible pulse or absent in femoral and Pedi arteries, acute renal failure (oliguria, anuria), shock, oxygen saturation difference between the upper and lower limbs (5,11). Also, there is a difference of more than 10-15 mmHg in blood pressure between upper and lower limbs. Presence of pulmonary rales stasis and hepatomegaly appear.

INTERRUPTED AORTIC ARCH

Congenital heart defect is the discontinuity between the ascending and the descending aortas and a critical obstruction in the systemic circulation.

The flow in the descending aorta depends on the presence of permeable ductus arteriosus. With the decrease of the resistance in the pulmonary vascular bed (after the first week of life), increases the pulmonary flow with the occurrence of heart failure. When the ductus arteriosus has a tendency to closure, appear tissue hypoxia, metabolic acidosis and prerenal failure (5,12).
As through ductus arteriosus, the lower half of the body receives blood, here cyanosis often occurs. If it is associated with large DSV, it will improve pulmonary blood oxygen saturation.

PHARMACOLOGICAL TREATMENT OF CONGENITAL HEART DEFECTS WITH DUCTUS-DEPENDENT SYSTEMIC CIRCULATION

Recommendations:
- Oxygenation improvement with caution to avoid closure of the ductus arteriosus, opting to maintain oxygen saturation of 75-85% (5).
- Maintaining the permeability of ductus arteriosus by administering a dose of prostaglandin E1 \(0.05-0.1\) mcg/kg/min (4).
- Patients diagnosed late will require high doses of prostaglandin E1 \(0.15-0.20\) mcg/kg/min with careful monitoring of adverse effects: respiratory depression and even apnea, hypocalcemia, hypoglycemia, hypotension, tachycardia or bradycardia, convulsions (4,13).
- After ductus arteriosus permeability, overloading the pulmonary circulation using inhibitors of angiotensin converting enzyme (Captopril 1 mg/kg) shall be avoided when blood pressure is normal or high, or phosphodiesterase inhibitors – Sildenafil \(0.5-1\) mg/kg/dose shall be used when the blood pressure is low (13,14).
- Diuretics are used, the preparation of choice Spironolactone \(1\) mg/kg in 1 to 3 doses may increase the retention of salt at 1-2 mg/kg/day in 1-3 divided doses.
- In severe forms of congestive heart failure, Furosemide \(1\) mg/kg/dose is associated until diuresis of more than 3 ml/kg/hour is achieved, the dose can be repeated over 6-12 hrs; in the absence of the effect, the dose can be repeated after 2 hours.
- The increase circulation to normal volume using the volume-expander (0.9% saline or Ringer Lactate) in the amount of 10 ml/kg administered intravenously.
- Severe metabolic acidosis must be treated with 4.2% sodium bicarbonate (in an amount \(2\) mEq/kg/dose) administered intravenously very slowly, the equivalent of \(2-4\) ml/kg/dose with adequate ventilation.
- Correction of rhythm disorders
- To maintain heart compliance, shall be administered (13,14):
  - Inotropic – Dopamine, in case of reduced cardiac contractility; administered \(5-20\) mcg/kg/min intravenously with infusion on the pump.
  - Digoxin \(0.04\) mg/kg saturation dose every 8 hours in the first day, followed by a maintenance dose, the \(1/4\) dose of the saturation dose in two divided doses.

The child will be monitored, the saturation dose being achieved through intravenous administration and the maintenance dose may be administered orally.

If clinical signs appear, such as food refusal, vomiting, impaired general condition, or electrocardiographic signs (PQ prolongation, underlevied ST, ventricular arrhythmia), it is considered that there was an overdose or poisoning with Digoxin, and the following measures shall be taken:
- Digoxic administration termination;
- Oxygen therapy;
- Correction of electrolyte disturbances;
- Vigorous anti-arrhythmic treatment in bradyarrhythmias – Atropine \(0.02-0.03\) mg/kg and in tachyarrhythmias – Lidocaine in bolus of \(0.5-1\) mg/kg the initial dose, the subsequent doses \(0.02-.0.3\) mg/kgc/min.

CONCLUSIONS

In the neonatal period, multiple congenital heart defects may be overlooked due to patient’s ductus arteriosus contributing to maintenance of a poor clinical picture but with extremely severe evolution at the closure of ductus arteriosus.

REFERENCES


