Cardiac Emergencies in Neonates

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Abstract

Introduction: The diagnosis of cardiac disease in the neonates is not always straight forward because physical examination, ECG, and CXR are often difficult to interpret in the newborn period compared to older infant or child. Although echocardiography is required to precisely define the anatomical abnormality, it is usually possible to define. The most important factors in narrowing down the diagnostic possibilities are:

The clinical presentation
- Shock (ductal dependent systemic circ.) (Grey baby)
- Cyanosis (ductal dependent pulmonary circ.) (Blue Baby) including severe Ebstein’s anomaly
- CHF congestive heart failure (shunt lesions) (Pink Baby)
- Arrhythmia

The timing of the presentation (age)

Associated non cardiac or genetic anomalies

Objective: To Concentrates on few points which make the diagnosis and managements of cardiac emergencies in neonates easier.

Keywords: Neonates; Cardiac; Emergencies

Introduction

The diagnosis of cardiac disease in the neonates is not always straight forward because physical examination, ECG, and CXR are often difficult to interpret in the newborn period compared to older infant or child.

Although echocardiography is required to precisely define the anatomical abnormality, it is usually possible to define the functional abnormality on the basis of the clinical and radiographic findings.

The timing of presentation and severity depends on
- Nature and severity of defect
- The alteration in cardiovascular physiology secondary to the effect of the transitional circulation as
- Closure of ductus/restriction of patent foramen ovale (PFO)
- Fall in pulmonary vascular resistance (PVR)

The most important factors in narrowing down the diagnostic possibilities are
- The clinical presentation
  - Shock (ductal dependent systemic circ.) (Grey baby)
  - Cyanosis (ductal dependent pulmonary circ.) (Blue Baby) including severe Ebstein’s anomaly
  - CHF congestive heart failure (shunt lesions) (Pink Baby)
- The timing of the presentation (age)
- Associated non cardiac or genetic anomalies

So the vital signs and clinical examination in newborns are very important And lead to the right diagnosis: (Right sided obstructive lesions(cyanosis), Left sided obstructive lesions (Collapse), Mixing lesions (cyanosis with CHF), ASD dependent lesions (cyanosis with CHF) And give a chance to start PGE1 when there is any suspicion of PDA dependent lesion (systemic or pulmonary) before the Echo.
Neonatal Presentations


Septicemia, respiratory disorders, persistent pulmonary hypertension of newborn (PPHN), inborn errors of metabolism and so on.

Neonatal Presentations


Septicemia, respiratory disorders, persistent pulmonary hypertension of newborn (PPHN), inborn errors of metabolism and so on.

Cyanosis

Cardiac causes

- PDA dependent pulmonary circulation
- Admixture lesions
- Critical Right ventricular outflow tract obstruction with intracardiac shunt

Case 1

Asymptomatic neonate was discharged on breast feeds
- Suddenly presents at 3 days of life with bluish discoloration
- Sats 70, mild tachypnea and minimal distress, soft ESM, No response to oxygen
- Ventilation -mild increase in saturations (75%)

CXR: Oligemic lung fields and no cardiomegaly Figure 1

Plan: Start Prostaglandin E1 and get Echo

Why PGE1 started, can we do it without echo, what is the probable DD.

Pulmonary atresia with VSD

PDA dependent pulmonary circulation

1. Pulmonary Atresia, Intact Ventricular Septum
2. Pulmonary Atresia, VSD and PDA
3. Pulmonary Atresia with Single Ventricle
4. Severe forms of Epstein's anomaly

What is common to all these lesions

Oligemic lung fields on CXR

Admixture lesions

- Transposition of Great Arteries, intact Interventricular septum
- Total Anomalous Pulmonary venous connection
- Truncus Arteriosus
- Double Outlet Right Ventricle with VSD
- Single ventricle anomalies with or without Pulmonary stenosis

Critical Right ventricular outflow tract obstruction with intracardiac shunt

- Critical Pulmonary Stenosis with interatrial communication
- Tetralogy of Fallot with Critical Pulmonary Stenosis

Pulmonary causes

Primary lung disease

- Respiratory distress syndrome
- Meconium aspiration

PPHN

- Pneumonia
- Tracheo- esophageal fistula

Airway obstruction

- Choanal atresia, laryngotraechomalacia, laryngeal web, vocal cord paralysis
- Extrinsic compression of the lungs
- Pneumothorax
- Chylothorax, Hemothorax
- Diaphragmatic Hernia, Space occupying lesions

Neurologic causes

- Drug-induced depression of respiratory drive,
- Intracranial hemorrhage,
- Post asphyxia cerebral dysfunction,
- Central apnea
- Respiratory neuromuscular dysfunction
Cardiac Emergencies in Neonates

- Spinal muscular atrophy,
- Infant botulism,
- Neonatal myasthenia gravis

**Hematologic causes**
Methemoglobinemia or polycythemia cyanotic with normal Pao2

How to identify cardiac vs respiratory causes of cyanosis (Table 1)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Respiratory pattern</th>
<th>Saturation difference between right upper limb (Rt) and lower limb (LL)</th>
<th>PCO₂</th>
<th>Response to 100% oxygen</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiac disease</td>
<td>Increased respiratory rate with minimal distress</td>
<td>Usually no difference unless specific lesions</td>
<td>Normal or low</td>
<td>No significant change</td>
</tr>
<tr>
<td>Primary pulmonary disease</td>
<td>Increased respiratory rate with distress</td>
<td>No difference</td>
<td>High</td>
<td>Increased saturations</td>
</tr>
<tr>
<td>PPHN</td>
<td>Increased respiratory rate with distress</td>
<td>&gt;10% Rt &gt; LL</td>
<td>Normal or high</td>
<td>May or may not change</td>
</tr>
</tbody>
</table>

**Hyperoxia Test**

Differentiate mainly cardiac from respiratory cyanosis (Table 2)

If resting saturations are less than 95%.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>At Fio2 = 0.21 Pao2 (Saturation %)</th>
<th>At Fio2 = 1.00 Pao2 (Saturation %)</th>
<th>PaCo2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>&gt;70 (&gt;95)</td>
<td>&gt;300 (100)</td>
<td>Normal</td>
</tr>
<tr>
<td>Pulmonary disease</td>
<td>50 (85)</td>
<td>&gt;150 (100)</td>
<td>111211</td>
</tr>
<tr>
<td>Neurological disease</td>
<td>50 (85)</td>
<td>&gt;150 (100)</td>
<td>High</td>
</tr>
<tr>
<td>Methemoglobinemia</td>
<td>&gt;70 (&gt;85)</td>
<td>&gt;200 (&gt;85)</td>
<td>Normal</td>
</tr>
<tr>
<td>Cardiac disease</td>
<td>&gt;40-60 (75-93)</td>
<td>&lt;150 (&lt;93)</td>
<td>Normal</td>
</tr>
<tr>
<td>PPHN</td>
<td>Preductal 40-70 (75-95)</td>
<td>Variable</td>
<td>Normal/high</td>
</tr>
<tr>
<td></td>
<td>Post ductal &lt;40 (75)</td>
<td>Variable</td>
<td></td>
</tr>
</tbody>
</table>

**Limitations of hyperoxia test**
- False positive
- False negative

Pulmonary disease with a massive intrapulmonary shunt may not respond to oxygenation.

**Differential cyanosis**
Preductal saturation > Post-ductal saturation
- Persistent Pulmonary Hypertension of the newborn (PPHN)
- Interrupted aortic arch (Figure 2), critical coarctation of the aorta and critical aortic stenosis.

**Lower limb blue, Rt upper limb pink**

**Table 2: Hyperoxia test.**

**Interpretation of hyperoxia test**
- PaO₂ more than 250 mmHg excludes cyanotic congenital heart disease
- PaO₂ is more than 150, cyanotic CHD unlikely
- PaO₂ less than 150, Cyanotic CHD

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**Reverse differential cyanosis**

Pink Lower limb, blue upper limb

The post ductal saturation higher than the preductal saturation.

We notice it in many cases like:
- TGA with (critical coarctation of the aorta, interrupted aortic arch, critical aortic stenosis)
- TGA with PPHN.

**Classification of Cyanotic CHD**

Decreased PBF
- TOF/TGA, VSD, PS/DORV, VSD, PS/TA, VSD, PS/SV, PS/Pulmonary atresia

Increased PBF
- TGA, VSD/Truncus/TAPVC (admixture)/SV without PS/TA, VSD without PS

*Chest X ray (figure 3)*

*Figure 3: High PBF (left) and low PBF (right).*

- Rule out respiratory issues
- Assess pulmonary blood flow and presence of Cardiomegaly

**Increased PBF features on CXR**
- Cardiomegaly
- Increased vascularity

**Shock**

**Case 2**

D3 old baby – suddenly sick
- Poor feeding, Not passing urine
- Color not looking good – pale
- Cold peripheries, Tachypnoea, tachycardia poor pulses, BP 32/20, metabolic acidosis, raised lactates

- A healthy newborn who presents after 48-72 hrs of life with sudden onset of pallor, grey appearance and breathing difficulty.
- Not passing urine and not taking feeds over last 4-6 hrs
- Metabolic acidosis
- Start prostaglandin E1 suspecting duct dependent systemic circulation??
- Other measures to stabilize including ventilation and inotropes (unless echo rules out cardiac lesion).

**Cardiac causes of shock:**
- Duct dependent systemic circulation and Left ventricular outflow Tract obstructions
  - Critical Aortic stenosis and coarctation, Interrupted aortic arch
  - Hypoplastic Left Heart Syndrome (HLHS)
- Obstructive TAPVC
- Rhythm disturbances: Tachyarrhythmias and Brady arrhythmias (e.g. complete heart block)

**Prostaglandin E1**

How do we start Prostaglandin?
- Dose - 0.001-0.4 microgram/kg/min infusion
- Higher doses 0.1 mic/kg/min should be used to reopen the closed PDA (or if there is sudden onset of severe cyanosis or shock).
- Once the duct has opened, dose can be reduced to a minimum

If no Response to prostaglandin Infusion (Table 3).

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Management</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transposition of Great Arteries with Intact Ventricular Septum and a restrictive interatrial communication</td>
<td>Needs emergency Balloon Atrial Septostomy</td>
</tr>
<tr>
<td>Obstructed TAPVC</td>
<td>Emergency surgery</td>
</tr>
<tr>
<td>Non Cardiac Diagnosis</td>
<td>Treatment of etiology</td>
</tr>
</tbody>
</table>

*Table 3*

**Congestive heart failure**

**Cyanotic heart disease with high pulmonary flow**
- Truncus Arteriosus
- Single ventricle physiology without Pulmonary Stenosis
- Transposition of Great Arteries with VSD
- Double Outlet Right Ventricle with VSD
Cardiac Emergencies in Neonates

- Total Anomalous Pulmonary venous connection anotic heart disease with high pulmonary flow
  - Not significantly cyanotic due to high pulmonary blood flow
  - Saturation may vary in the range of 90’s.
  - Naked eyes will not pick up the cyanosis
  - Pulse oximetry - early detection

Acyanotic heart disease with high pulmonary flow
- Preterm with significant post tricuspid shunt lesions (e.g. VSD, PDA, Aorto- Pulmonary window)
- Severe valvular regurgitant lesions (e.g. Mitral regurgitation associated with AV canal defects)
- Anomalous Left Coronary Artery from Pulmonary Artery (ALCAPA)
- Cardiomyopathy

Rhythm disturbances
- Tachyarrhythmias
- Brady arrhythmias (e.g. complete heart block, High degree second heart block)

Non Cardiac causes
- High output states like anemia, thyrotoxicosis, systemic Arteriovenous malformations (g.e. Vein of Galen)

Clinical features
- Difficulty in feeding
- Sub costal indrawing
- Sweating with feeds
- Tachypnoea
- Tachycardia
- Gallop rhythm and hepatomegaly.

Arrhythmias
- SVT
- Brady arrhythmias with Long QTc

Neonatal Interventions
- BAS-Ballon atrial Septostomy
- BPV-Ballon pulmonary valvuloplasty
- BAV-Ballon aortic valvuloplasty
- Ballon Coarct angioplasty
- PDA stenting

Conclusion
- Right sided obstructive lesions(cyanosis) - critical PS, pulmonary atresia
- Left sided obstructive lesions (Collapse) – critical AS, IAA, COA, HLHS
- Mixing lesions (cyanosis with CHF) – TGA
- ASD dependent lesions (cyanosis with CHF) TAPVC, Tricuspid atresia

Important massage
- Vital signs and clinical examination in newborns are very important
- Starting PGE1 when there is any suspicion of PDA dependent lesion (systemic or pulmonary) before the Echo.

Bibliography

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