Summary
Disease (CHD) in clinically sub-optimal settings. It is presented to provide simple differences between certain conditions commonly seen in children with CHD for training doctors and nurses and other staff working within the speciality. In the majority of clinical presentations, a comprehensive, well structured training of a multidisciplinary team and carefully chosen hospital equipment and resources, can permit CHD to be safely and effectively treated by palliative or curative procedures. In our experience using this strategy, outcomes are almost comparable to those in advanced centers across the world. Among nations with severely restricted general and specialist healthcare resources, several issues must be overcome to diagnose and treat children with congenital heart diseases (CHD). The principal challenges to address are:

Provider issues
1. Lack of primary and tertiary specialist facilities to support, diagnose, treat, implement follow-up care and preventive measures within the community so that avoidable complications of these diseases can be identified and minimized.
2. Lack of trained personnel in most of the specialties needed to support pediatric cardiac services

Patient related issues
1. Maternal health issues affecting prematurity, dysmaturity, nutrition inadequacy during pregnancy threatening fetal development;
2. Child development issues: from insufficient nutritional support during post-natal growth; regionally limited, poor or absent primary healthcare; lack of preventative measures to reduce complications in such presentations ((for example, rheumatic fever, complicating CHD); With careful preparation using the algorithms designed by CardioStart International, safely conducted complex operations can lead to good clinical outcomes throughout the peri-operative period. A vitally important component of these, is the “Dry Run Checklist” which allows the local team to confirm adequacy of equipment and disposables throughout the perioperative period. *[These are available on request]

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Introduction
Congenital heart diseases (CHD) are made up of structural or functional heart defects which are present from birth and give rise to disease as growth takes place. The worldwide prevalence of CHD is 8-10/1000 live births; among these, 33% to 50% of such defects may be considered critical: they normally require intervention within the same hospital admission, and ideally, are dealt with early on during the first year of life. In locations across the globe in which healthcare resources are poor, several related and unrelated issues may prevent diagnosis and treatment being available to those children in an adequate time period:
1. Poor ante-natal performance during pregnancy;
2. Malnutrition;
3. Medications, alcohol, recreational drugs and smoking history during pregnancy leading to co-existing morbidities;
4. (Frequent) late presentation of CHD beyond infancy;
5. Lack of early referral;
6. Illiteracy of the parents;
7. Large numbers of births taking place within homes lacking adequate sanitation;
8. Large numbers of births that are un supervised by a doctor familiar with neonatal care;
9. Inadequate number of cardiologists and cardiac surgeons;
10. Lack of regional specialized cardiac centers;
11. Limited understanding and knowledge of CHD by local primary health care providers (physician, pediatrician, internist, etc.)
To overcome these obstacles the optimal strategy should involve the training and education of the local teams. Thus, a review of the essentials for the treatment of the most prevalent CHD is presented in following sections.

A: Understanding congenital heart diseases
Measurement of oxygen saturation by clinical observation and bedside pulse oxymetry testing, assists separation of congenital heart diseases into two main categories:
1- Acyanotic CHD: included in this category are children with near normal or normal arterial oxygen saturation shortly after birth. Examples: Some ventricular septal defects (VSD), atrial septal defect (ASD), patent ductus arteriosus (PDA), coarctation of aorta, moderate pulmonary stenosis, and some mild cases that belong to the spectrum of Tetralogy of Fallot.
2- Cyanotic CHD: In this category are those with clearly reduced arterial oxygen saturation that does not improve.
Examples: Tetralogy of Fallot (TOF), transposition of great arteries, tricuspid atresia, truncus arteriosus, total anomalous pulmonary venous drainage, hypoplastic left or right heart, and certain peri-natal cardiac tumors.

B: Basics of Surgery for Congenital Heart Diseases
According to the clinical presentation, the surgical treatment of CHD can be grouped into two main approaches:
1- Palliative Procedures (staged)
2- Complete Repair (primary, definitive operation)
As a surgical program grows and becomes more established, some of the techniques described below may be modified as experience develops, and from personal choice and preferences of individual surgeons.

1.1. Palliative Procedures for Congenital Cardiac Lesions
A palliative operation is chosen to produce an improvement on abnormal circulation and heart function, minimize some of the anatomical and functional features of the disorder and worsening of abnormal growth; it is usually performed in children who are considered to be too young for corrective surgery at the regional center dealing with complex pediatric heart disease. The goal is to lessen cyanosis, control heart failure and prepare the circulation for later correction, anticipating natural growth. Each procedure is chosen to bring relief of symptoms, improvement in regional blood flow, and better oxygenation. The success of such operation is influenced by the child’s growth, and absence of co-morbidities (and new onset diseases that commonly occur in most children).
It is anticipated that as the infant grows, age and body weight development indices will then permit a fully corrective surgical technique to be employed. The objectives in performing a palliative procedure are:
a) To provide symptomatic relief;
b) Allow the impact of the cardiac malformation to be better tolerated; c) Produce an improvement in the patient’s general clinical condition.
A reduction in degree of cyanosis and improved hemodynamics usually produce a lowering in resting respiratory rate and effort of breathing, which leads to better food intake. A growth spurt may therefore be achieved during the interval before the child undergoes complete repair. In the twentieth century, two classic palliative procedures emerged during development of neonatal and pediatric cardiac surgery procedures which are performed in clinical practice. Since their inception, modifications to the surgical technique have evolved:  

**Aortopulmonary shunt:**  
This procedure aims to directly connect blood from the ascending aorta or its branches to the pulmonary artery at the point distal to any obstruction: the operation is designed to increase pulmonary blood flow and promote an enlarge-

**Pulmonary artery banding**  
In this scenario, pulmonary blood flow is excessive due to the intra-cardiac lesion, and if untreated, flooding of the pulmonary circulation will allow pulmonary hypertensive changes to occur. This operation is designed to carefully reduce pulmonary blood flow and pressure thereby protecting the pulmonary vasculature from progression towards irreversible pulmonary vascular disease. Multiple or large VSDs in infants weighing less than 3kg may present so, and the improvement in pulmonary dynamics and drop in respiratory rate may assist with better feeding, particularly at the breast.  
More complex palliative procedures including the staging for the Norwood operation for hypoplastic left heart syndrome and the Glenn operation are outside the scope of this review.  

**Indications and timing of palliative procedures:**  
**Aortopulmonary shunt**  
Most patients with pulmonary obstruction, presenting with cyanosis, dyspnea, and/or failure to thrive, usually benefit from a shunt procedure. This is a frequent operative choice in centers with limited experience, facilities or support staff, and/or when the patient may be judged as too small for adequate total repair. Examples are Tetralogy of Fallot less than 3 months of age; and Pulmonary Atresia with or without VSD.  

**Surgical techniques**  
An aortopulmonary shunt can be performed through a mid-line sternotomy or through a right

Figure 1- Aortopulmonary shunts: classic and modified Blalock-Taussig shunt  

or left thoracotomy using one of the techniques described below:  
1- **Classic Blalock-Taussig-Thomas shunt:**  
This technique is now less commonly used. It consists of anastomosing the subclavian artery to the pulmonary artery on the side opposite the aortic arch.  
With certain surgical technical modifications, the subclavian artery can be anastomosed to the pulmonary artery on the same side of aortic arch. The surgical approach is usually carried out through an upper thoracotomy centered on the upper border of the 3rd or 4th intercostal space. Among the essential surgical instruments are a straight and curved (Cooley) vascular clamps to permit uncluttered surgical access and retain hemostatic control throughout. A surgical headlight is also required. If performed on the left, careful attention must be given to avoid strenuous retraction of the left lung against the heart, which may produce poor venous return (Figure 2).  

Figure 2- Classic Blalock-Taussig-Thomas shunt
II- Modified Blalock-Taussig-Thomas shunt (MBT Shunt)
In this procedure, a polytetrafluoroethylene (PTFE) tube graft (3-5mm, depending on age at presentation) is interposed between the subclavian or innominate artery and the right or left pulmonary artery using the same left or right thoracotomy access as described above. The surgical procedure requires the surgeon to apply careful judgment of the length of the graft chosen to provide sufficient flow, avoid kinking of the graft (too long) or puckering of the native vessels (too short) as the child grows during the “surgical interval” between operations. It is the most commonly performed shunt procedure (Figure 3).

2.1. Complete Repair of Congenital Cardiac Lesions
A Complete repair is feasible in many extra-cardiac as well as intra-cardiac congenital anomalies.

A- Repair of extra-cardiac lesions:
In these presentations, there is usually no need for cardiopulmonary bypass. The commonest examples are PDA, and coarctation of aorta.

PDA:
A spontaneous closure can be reasonably expected in small PDAs in full term babies at the age of 3 months; however, large PDAs are unlikely to close.

The timing for PDA closure is as follows:
- Large/ moderate PDA, with congestive heart failure (CHF) and/or pulmonary hypertension (PAH): Early closure (by 3-6 months) is recommended.
- Moderate PDA, no Congestive Heart Failure (CHF): closure at 6 months-1 year of age. If the infant is exhibiting failure to thrive, closure can be accomplished earlier.
- Small PDA: closure can be carried out at 12-18 months of life.
- Silent PDA: Closure is not recommended. This entity is defined as a small ductus arteriosus with normal pulmonary artery pressure which cannot be identified clinically, except by Doppler echocardiography.

Coarctation of aorta
Surgical treatment currently represents the preferred option for the correction of coarctation of aorta (CoAo).

The defect is best approached through a limited left posterolateral thoracotomy based on the upper border of the 4th interspace, and is usually divided between clamps or ligated (Figures 4a and 4b) taking care to avoid the phrenic nerve and Abott’s artery. If ligation only is chosen, the separate ligating sutures should be 2/0 or 3/0, carefully tied without subtraction to avoid “cheese-cutting” the friable duct, pulling on the vessel or worse, causing tearing off from the pulmonary artery or aorta. The surgeon should be prepared to immediately open the adjacent pericardium in that scenario to retrieve the retracted PA segment.

The timing of intervention should be based on the clinical status at presentation, rather than waiting for developing hypertension and left ventricle dysfunction:
• Low gradient: Intervention is not indicated if Doppler gradient across coart segment is <20 mmHg with normal LV function. It should be reassessed at regular intervals as the child grows and blood pressure measurements should be taken from right, left arms and one leg. Pulse delay should be checked and recorded if found.
• CoAo with LV dysfunction / CHF or severe upper limb hypertension (for age): Immediate intervention. In this scenario, considerable care should be taken during the operation to avoid retraction of the lung against the pericardium (which may produce a reduction in left atrial filling and subsequent bradycardia).
• CoAo with normal LV function, no CHF and mild upper limb hypertension: Intervention may be undertaken beyond 3-6 months of age
• CoAo with no hypertension, no CHF, normal LV function: Intervention can be made at 1-2 years of age.

The defect is best approached through a left posterolateral thoracotomy centered on the upper border of the 3rd left interspace using one of the following techniques (if made lower, it is very difficult to apply the appropriate clamps and gain surgical access).
• Resection and end–to–end anastomosis (Figure 5a and 5b),
• Augmentation of the coarcted segment

B- Repair of intra-cardiac lesions:
These usually require cardiopulmonary bypass. The commonest defects are VSD, ASD, and Tetralogy of Fallot.

Atrial Septal Defect (ASD)
A spontaneous closure is rare if the defect is >8 mm at birth and if still evident after age 2 years.
The timing of repair is based on the symptomatology secondary to pulmonary blood overload, as follows. (If a child is well and the lesion is picked up by auscultation on routine review, many surgeons will choose to do this operation during school holidays or at parents’ convenience. It is usually non-urgent):
• Asymptomatic ASD: These can be repaired at 2-4 years.
• Symptomatic ASD in infancy - CHF, severe Pulmonary Artery Hypertension (PAH): (8%-10% of cases). For these, early closure is recommended, and it should be fully confirmed that this is the only cardiac lesion present.
• Late presenting ASD: Elective closure should be carried out at presentation irrespective of age and, especially, if there is right heart volume overload and pulmonary vascular resistance with subclavian flap, or synthetic patch -polytetrafluoroethylene (PTFE).

Note: Occasionally, it may be necessary to introduce one of the clamps through a lower intercostal space to gain a suitable clamp position. The use of silastic slings after careful, discrete (localized) dissection of the proximal and distal aortic segments is an important adjunct to the surgical technique.

Figure 5a – Juxta-ductal coarctation of aorta. Figure 5b - End–to–end anastomosis of the coarcted segment.

(PVR) is within operable range (PVRIO<10WU. m2 – PVRIO. Pulmonary vascular resistance index can be assessed using a pure oxygen inhalation challenge).

ASD closure is most commonly approached through a median sternotomy under cardiopulmonary bypass. In centers with a large experience it is occasionally managed by upper right thoracotomy, centered on 4/5th interspace. This can be a helpful operative approach in girls who may seek to avoid a central chest scar. If the later procedure is done, very careful cannulation techniques must be used to avoid accidental line loss; the incision must not be placed anteriorly so to avoid risking a dimpling deformation of the right breast later on, when the patient reaches puberty.

An autologous pericardial patch is usually used to close the defect. (Figure 6a and 6b). If future surgery is anticipated for an additional cardiac lesion, it is best to also use commerical pericardial patches to then draw the pericardium closed, (if such materials are available). This is carried out on terminating cardiopulmonary by-pass and de-cannulation.

Ventricular Septal Defects (VSD)
Approximately 30%-40% of moderate or small defects (restrictive) close spontaneously, the majority closing by 3-5 years of age or becoming hemodynamically insignificant without shunt flow on serial echocardiographic assessments. Spontaneous closure is uncommon in large VSDs, some of which (approximately 10%) die in first year of life, primarily due to CHF when the VSD is large and non-restrictive (= diameter of the aortic orifice). The timing of repair is based on the symptomatology characterized by the presence of CHF, respiratory infection and/or PAH. In those children presenting with cardiac or other comorbidities, a careful review of the peri-operative management in Intensive Care equipment, nursing skills and medications that are available should be made. Even after a perfect surgical result, in the OR, a child’s most challenging time may be in the post-operative period.

Large VSD with uncontrolled CHF: Repair should be carried out as soon as possible, after accomplishing maximizing general hemodynamic improvement before the child undergoes anesthesia.

Large VSD with severe PAH: Repair at 3-6 months.

Moderate VSD with PASP 50%-66% of systemic pressure: Repair is recommended between 1-2 years of age or earlier if one episode of life-threatening lower respiratory tract infection or failure to thrive.

Small sized VSD with normal PAP, left to right shunt >1.5:1: Closure by 2-4 years.

Small outlet VSD with any degree of aortic regurgitation (AR): Surgery whenever AR is detected.

Small VSD with one previous episode of infective endocarditis: Early VSD closure is recommended.

Surgical technique
The VSD is approached through a median sternotomy under cardiopulmonary bypass and closure is accomplished by use of a pericardial patch, (autologous pericardium, Dacron patch or PTFE membrane are popular). (Figure 7a and 7b). Great care should be given in regard to suture placement to avoid ensnaring one of the aortic cusp, conduction system, and deforming the tricuspid valve leading to valvular incompetence. Some surgeons do not use pledgets throughout. Some prefer a single, continuous suture technique.
This may suffer from the risk of rupture and unzipping which can cause dynamic sub-aortic obstruction (and require emergent re-operation). A four-quadrant interval continuous technique reduces the probability of this complication occurring. Surgical approaches are now most commonly carried out by opening the right atrium and gentle upward retraction of the septal cusp of the tricuspid valve, but some circumstances may require right ventriculotomy or, by direct access through aortotomy/pulmonary truncal opening to gain access to the immediate sub-aortic or sub-pulmonary margin of the defect.

Tetralogy of Fallot
This accounts for 15% of all cyanotic heart diseases. All patients need surgical repair but in patients coming from rural locations, the primary diagnosis may be sometimes delayed well into the patient’s teen years. The timing of repair is often based on the severity cyanosis if the patients are seen soon after birth. They may require a different peri-operative strategy depending on factors relating to surgical and anesthetic perfusion and nursing experience, and the availability of specialist management, operative and perfusion equipment in such small sized patients.

1. **TOF stable, minimally cyanosed**: Total correction can be usually, safely carried out at 1-2 years of age or earlier according to the institutional policy.

2. **TOF with significant cyanosis (SaO2<70%) or spells despite therapy**
   - <3 months: A BT shunt may be performed, scheduling full correction after a growth period.
   - >3 months: Shunt or correction depending on anatomy, growth, co-morbidities and surgical centers’ experience

TOF is approached through a median sternotomy under cardiopulmonary bypass. The repair consists of closure of VSD with a patch (usually bovine pericardium or Dacron patch) and augmentation of the RV outflow tract and the stenosed pulmonary valve and arteries (Figures 8a and 8b).
The results of surgery usually show substantial improvement in patient’s tolerance to exercise, improved growth and resolution of cyanosis. Patients should be advised that there may be some distortion of the pulmonary artery following growth and that surgical revion may be required.

**Summary**

1. PDA, CoAo, ASD & VSD with their respective anatomic variants and the lesions found in TOF represent > 80% of all CHD for which a center that is located in a resource-poor environment may reasonably develop the capability to repair them.

2. Pediatricians/ cardiologists/ other health care providers should strive to get a complete diagnosis on a child suspected of having heart disease, even if that requires referral to an advanced center.

3. Development of surgical protocols: these are vital tools to help with in pre-operative planning, peri-operative management; they also help provide a smooth passage through the essential fatures of management in each clinical presentation: diagnosis and surgical decision, the optimal timing of intervention. Such a strategy will help achieve good outcomes in the more common congenital heart diseases which present to resource-poor centers.

**References**


