Timing of cardiac surgery and other intervention among children with congenital heart disease: A review article

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Abstract

Background: Early diagnosis and improved facilities are necessary for determining the optimal timing of surgery and other interventions in children with congenital heart diseases in Nigeria. This is because late presentation, late diagnosis and delayed surgery can lead to mortality and affect the quality of life among these children.

Objectives: This review article is aimed at enumerating the timing of cardiac surgeries and other interventions and to seek if there is any factor associated with the timing of cardiac surgery.

Methods: A search on PubMed database, World Health Organization libraries, Google scholar, TRIP database, and reference lists of selected articles on timing of cardiac surgery in children was done. The Cochrane Database of Systematic Reviews was also searched. We noted few data from African setting. Key words such as timing of cardiac surgery; children, congenital heart defect were used.

Conclusion: Appropriate timing for cardiac surgery in children with congenital heart disease is very important as late surgical intervention could result in several morbidities and mortality.

Introduction

Congenital heart disease is becoming a common cause of congenital abnormalities in our locale [1]. There is also a dramatic increase of the prevalence of congenital heart disease (CHD) which has risen from 4 to 5 per 1,000 live births among older studies to 12 to 14 per 1,000 live births, reported in the recent literature [2]. In the foregoing therefore, the diagnosis and treatment for congenital cardiac disease has also undergone remarkable progress over the last 6 decades [3]. Accurate timing and intervention significantly improves quality of life, prevents complications and reduces the morbidity and mortality associated with these defects. Worldwide, CHD are the main cardiac anomaly found in children and constitute one of the major causes of infant mortality, particularly in developing countries [4].

The indications and timing of intervention among children with congenital heart disease is decided by the severity and type of the lesion. For instance, balloon pulmonary valvuloplasty is the treatment of choice for valvar pulmonary stenosis and surgery must be done when the peak-to-peak systolic pressure gradient is more than 50 mmHg across the pulmonary valve [5]. For aortic valve stenosis, balloon aortic valvuloplasty appears to be the first therapeutic procedure of choice with intervention feasible when peak-to-peak systolic pressure gradient across the aortic valve in excess of 70 mmHg irrespective of the symptoms or a gradient more than and equal to 50 mmHg with either symptoms or electrocardiographic ST-T wave changes indicative of myocardial perfusion abnormality [6]. The indications for intervention in coarctation of the aorta are significant hypertension and/or congestive heart failure along with a pressure gradient in excess of 20 mmHg across the coarctation [7]. In addition, it is important to remember that the defect once detected should be corrected whether by surgery or by nonsurgical methods as appropriate before ventricular dysfunction, pulmonary vascular change and other complications ensue.
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It is important to note that once ventricular dysfunction, pulmonary vascular change set in, the surgery becomes high risk with an increased chance of postoperative complications and poorer long-term results. Many of these cases may even become inoperable, if timely intervention is not done. Every cardiologist or cardiac surgeon must strive to get a complete diagnosis on a child suspected of having heart disease as soon as possible even if that requires referral to a higher centre. This review article is aimed at determining the timing of cardiac surgeries and other intervention in children with congenital heart disease and to seek if there is any factor associated with timing of cardiac surgery.

Methods

A search on PubMed database, World Health Organization libraries, Google scholar, TRIP database, and reference lists of selected articles on timing of cardiac surgery in children were done. The Cochrane Database of Systematic Reviews was also searched. Some studies were selected that included 2,027 children with congenital heart disease.

Results

In a study done in University of Nigeria Teaching hospital Enugu, Eighty-eight patients, consisting of 59.09% females and 40.91% males with cardiac anomalies presented to the facility over a period of 4 years. The mean age for cardiac surgical closure was 4.4 (4.1) years with minimum and maximum age of 2 months and 18 years respectively. Majority of the patients had surgery after their first birthday and substantial number after their 5th birthday. The gender of patients did not influence the time of surgery. The mean age at cardiac surgery in females was 4.0 (0.5) years while that for males was 4.8 (0.7) years (Table 1).

The time for the closure of cardiac defects was very late in developing countries as seen above, compared with that seen in Pakistan; another developing country, where the mean age at the time of surgery was 6.08 ± 2.80 months (Table 2).

In California, studies done between 1995-1996, 666 children were noted to have undergone ASD closure at a mean age of 5.1 years; median: 4.0 years, and 582 children with VSD closure had theirs at a mean age of 2.8 years; median: 1.1 years, while 394 children with TOF repair had repair at a mean age of 1.7 years; median: .9 years with 177 children who presented with AVC repair had their intervention at a mean age 1.1; median: .6 years (Table 3).

They noted that Asians tended to be older at surgery for all types of cardiac defects yet gender or race were noted not to have any effect on age at surgical closure of congenital heart defects. They however noted an association between the place of domicile and age at operation especially for ASD and TOF with no attendant link between distance to surgical centre and age at operation. It was seen that among 467 live births to Massachusetts residents between 2004 and 2009, were among 916 children with congenital heart disease, 126 (13.8%) had delayed diagnosis. Rates of prenatal CHD diagnosis increased from 44.9% in 2004 to 63.8% in 2009, whereas rates of delayed diagnosis decreased from 17.1% to 10.6% over the same time period. Delayed diagnosis was associated with delivery outside a tertiary hospital and isolated congenital cardiac defect (Figure 1).

Brief discussion of results on some studies as regards timing of cardiac surgery

Accurate timing and intervention in children with congenital heart disease (CHD) significantly improves quality of life, prevents complications and reduces the morbidity and mortality associated with these defects. Worldwide, CHD are the main heart diseases found in children and constitute one of the major causes of infant mortality, particularly in developing countries. In fact, nine out of 10 new-borns with CHD will live in a developing country where cardiac care is suboptimal or unavailable entirely. Seventy percent of those infants in a developing nation will require medical or surgical intervention within their first year of life, and 30% of those children will die without treatment in that first year.

Table 1: Age at surgical closure of cardiac anomaly. (UNTH ENUGU) n=88.

| Age at Surgery | Frequency | % |
|----------------|-----------|---|
| ≤ 1            | 14        | 15.9 |
| >1-5           | 47        | 52.4 |
| > 5            | 27        | 30.7 |
| Total          | 88        | 100 |

Table 2: Age at surgery of cardiac in Pakistan n=120 [8].

| Age at Surgery | Frequency | % |
|----------------|-----------|---|
| 6.08 ± 2.80 Months | 51        | 42.5 |
| 37.10 ± 19.94 Months | 69        | 57.5 |
| > 5            | 0         | 0   |
| Total          | 120       | 100 |

Table 3: Age of closure in Massachusetts [9].

| Defect Age at Closure (Mean) | Frequency |
|------------------------------|-----------|
| ASD 5.1 years; median: 4.0 years | 666       |
| VSD 2.8 years; median: 1.1 years | 582       |
| TOF 1.7; median: 9 years       | 394       |
| AVC 1.1; median: .6 years     | 177       |

Figure 1: Trends in timing of diagnosis by birth year are shown in Massachusetts [10].
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In developing countries like Nigeria, most surgical intervention for all cardiac lesions takes place between 1 and 5 year of age. In a study done in University of Nigeria Teaching hospital Enugu, it is noted that the mean age for cardiac surgical closure was 4.4 (4.1) years with minimum and maximum age of 2 months and 18 years respectively. It was also noted that gender of patients did not influence the time of surgery. The time for the closure of cardiac defects was very late in developing countries as seen above, compared with that seen in Pakistan another developing country, where the mean age at the time of surgery was 6.08 ± 2.80 months [8].

In California, studies done between 1995-1996, 666 children were noted to have undergone closure at earlier age. They noted that Asians tended to be older at surgery for all types of cardiac defects yet gender or race were noted not to have any effect on age at surgical closure of congenital heart defects. They however noted an association between the place of domicile and age at surgical closure [9]. In a study in Massachusetts, it was noted that among 916 children with congenital heart disease, 126 (13.8%) had delayed diagnosis. Delayed diagnosis was associated with delivery outside a tertiary hospital and isolated congenital cardiac defect [10]. The study above is similar to what is obtainable in the western world. For instance, Roheena, et al. [11] noted in their study that the earliest time of surgical intervention for their subjects is less than 1 year [11]. The reason for this late surgical intervention in this part of the world could be explained by the fact that surgical closure or any intervention for cardiac lesions is very costly and most people can’t afford it due to the harsh economic situations in the country. Moreover, surgery in this region is not done routinely due to lack of facilities and enabling environment.

**Review on time of repair of various congenital heart disease**

**Time for surgical closure of Atrio septal defect (ASD):** Spontaneous closure is rare if defect is more than 8 mm at birth and rare after age 2 years. Very rarely an ASD can enlarge on follow up. It is therefore, important to note that ideal time of closure should be about three months when there is a fall of pulmonary vascular resistance. For ASDs, Indication for closure includes ASD associated with right ventricular volume overload and the ideal age of closure in asymptomatic child is usually 2-4 years and 4-5 years for sinus venosus [12].

**Timing of surgical repair of atrio-ventricular-septal defect (AVSD):** Timing of intervention varies, depending on the type of congenital heart defect. For instance, in complete atrio-ventricular-septal defect (AVSD) with uncontrolled congestive heart failure, surgery is usually done as soon as possible after optimization [9-13]. Furthermore, incomplete Atrioventricular septal (AVSD) with controlled heart failure, complete surgical repair by 3-6 months of age is advocated and pulmonary artery banding may be necessary if risk of cardiopulmonary bypass is considered high. In Partial AVSD, surgery at about 2-3 years of age is advocated, however, associated significant AV regurgitation may necessitate early surgical intervention. Pulmonary artery banding should be done if risk of cardiopulmonary bypass is considered high [13-17]. Regrettably, in the developing country and sub-Saharan Africa, children with atrio-septal defects were maintained on anti-failure regimen with regular follow up until surgical intervention is possible. This makes some of them inoperable due to high pulmonary vascular resistance.

**Time of closure of ventricular septal defect (VSD):** In large (non-restrictive) VSD, where diameter of the defect is approximately equal to diameter of the aortic orifice, surgery should be done immediately. However, in children where right ventricular systolic pressure is systemic with high pulmonary vascular resistance, surgery may be impossible [18-22]. There have been general indications for surgical closure of VSD, which include patients at any age with large defects in whom clinical symptoms and failure to thrive cannot be controlled medically, infants between 6 and 12 months of age with large defects associated with pulmonary hypertension, children older than 24 months with a Qp: Qs ratio greater than 2 and children with a supra-cristal (doubly committed) VSD of any size (because of the high risk for aortic valve regurgitation) [20]. Contraindications to surgical closure of VSD include severe pulmonary vascular disease nonresponsive to pulmonary vasodilators.

Generally, in symptomatic infants with VSD, pulmonary vascular disease is usually prevented when surgery is performed within the 1st year of life except if there is evidence of shunt becoming restrictive [14-24]. For large VSD with uncontrolled congestive heart failure, closure should be done as soon as possible while in large VSD with severe pulmonary artery hypertension, closure should be done at 3-6 months of age. For moderate VSD where pulmonary artery systolic pressure is 50%-66% of systemic pressure, closure should be at 1-2 years of age or earlier if one episode of life threatening lower respiratory tract infection or failure to thrive ensues [18-24]. In the case of small sized VSD, with normal pulmonary artery pressure and left to right shunt >1.5:1, it is important to institute closure by 2-4 years [14-19]. Small outlet VSD (<3mm) without aortic valve prolapse should be on 1-2 yearly follow up to look for development of aortic valve prolapse. In addition, small outlet VSD with aortic valve prolapse without aortic regurgitation, should be closed by 2-3 years of age irrespective of the size and magnitude of left to right shunt [25,26].

Small outlet VSD with any degree of aortic regurgitation (especially in supra cristal type) should be closed whenever aortic regurgitation is detected. While small peri-membranous VSD with aortic valve prolapse with no or mild aortic regurgitation should be reviewed with 1-2 yearly follow up to look for any increase in aortic regurgitation [25,26].
For situations like Small Peri-membranous VSD with aortic cusp prolapse with more than mild aortic regurgitation, surgery is done whenever aortic regurgitation is detected. However, for small VSD with one or more episodes of infective endocarditis, early VSD closure is recommended [25,26]. The mode of closure for VSDs is usually surgical closure, device closure should however be done in muscular VSD especially for children weighing more than 15 Kg. In addition, pulmonary artery banding is usually indicated for multiple (Swiss cheese) VSD in very large VSD with almost a single ventricle, in infants with low weight (less than 2Kg) and associated co-morbidity like chest infection [18-25]. In countries where facilities for closure are not available, late closure is the norm and at times Eisenmenger’s complex could result from late surgical closure.

Timing for pulmonary stenosis: Timing for pulmonary stenosis also depend on certain factors. For instance, in right ventricular dysfunction, immediate intervention (valvuloplasty) irrespective of gradient should be done. In case with normal right ventricular function, balloon dilatation if Doppler gradient (peak) of more than 60mmHg. In neonates balloon dilatation is indicated if right ventricle dysfunction/ mild hypoplasia or hypoxia is present [35,36].

Timing for intra cardiac repair of Teratology of Fallot (TOF): In teratology of fallot (TOF), total correction should be at 1-2 years of age but if there is significant cyanosis (SaO2< 70%) or history of spells despite medical therapy especially plasma or saline phaeresis, then systemic to pulmonary artery shunt (Blalock Taussig shunt) should be done at less than 3 months, but if the child is more than 3 month, shunt or correction depending on anatomy and surgical centers' experience should be done [34-37].

Timing for repair of VSD with pulmonary atresia: For children with VSD with pulmonary atresia, repair should be at 3-4 years, if right ventricle to pulmonary artery conduit is required. Systemic to pulmonary artery shunt should be done if the child is symptomatic [34-36]. In TOF-like conditions where two ventricular repair is possible (DORV), timing of surgery also varies. For instance, for stable cases who are mildly cyanosed, repair at 1-2 years of age if conduit is not required while repair is at 3-4 years of age if conduit is required. Systemic to pulmonary artery shunt (Blalock-Taussig shunt) should be done if the child presents earlier with significant cyanosis (SaO2<70%) [34-38].

Timing for repair of transposition of great arteries (TGA): For transposition of great arteries (TGA), timing of intervention depends on the integrity of the interventricular septum. In TGA with Intact interventricular septum, if the child is less than 3-4 weeks of age; Arterial switch operation should be done immediately. If the patient is more than 3-4 weeks of age at presentation; it is expedient to assess the left ventricle mass and possibility of regression by echocardiography [37,39]. If the left ventricle is regressed, Senning /Mustard operation should be done at 3-6 month. It is very important to point out here that rapid two stage arterial switch approach would depend on institutional practice [37,38]. For example, if the left ventricle is still functional, very early arterial switch operation should be carried out. In borderline left ventricle; Senning or Mustard; or arterial switch operation is indicated. Adequacy of left ventricle for arterial switch operation can be assessed by echocardiography in most cases. In cases where a child presents with TGA with ventricular septal defect; Arterial switch operation by 3 months of age is necessary [38-40].

Several reasons have been propounded for late presentation and surgical intervention especially in a resource poor country. This ranges from late presentation due to ignorance to low socioeconomic class, unavailability of adequate
facilities, loss to follow-up despite proper diagnosis. Other reasons are refusal of surgery by some parents, unavailability of adequate manpower in the area of corrective heart surgery, some turbulent post-op cases and lack of proper coordination and synergy in all specialties involved in open heart surgery due to non-routine practice.

**Factors associated with timing of surgical closure**

Some authors have noted that gender or race had no effect on age at surgical closure of congenital heart defects. This was corroborated by Chang, et al. [38], who noted that although Asians tended to be older at surgery for all types of cardiac defects yet gender or race were noted not to have any effect on age at surgical closure of congenital heart defects. They however noted an association between the place of domicile and age at operation especially for ASD and TOF with no attendant link between distance to surgical centre and age at operation. In a study in among 460, 467 live births among Massachusetts residents between 2004 and 2009, Liberman, et al. [39], identified 916 CCHD cases, of which 126 (13.8%) had delayed diagnosis. Rates of prenatal CCHD diagnosis increased from 44.9% in 2004 to 63.8% in 2009, whereas rates of delayed diagnosis decreased from 17.1% to 10.6% over the same time period. They opined that delayed diagnosis was indeed associated with delivery outside a tertiary hospital.

**Conclusion**

Appropriate timing for cardiac surgery in children with congenital heart disease is very important as late surgical intervention could result in several morbidities and mortality. Literature search showed late timing of cardiac surgery in developing world especially in Africa. Factors such as late detection or diagnosis, poor counselling techniques used on our patients’ parents especially in the area of follow up visits, poor health education to encourage parents/caregivers to present with their children early are all seen as associated factors been implicated in delayed timing of cardiac surgery especially in developed country.

**Recommendation**

Early detection or diagnosis (all hands must be on deck), proper counselling of our patients’ parents especially in the area of follow up visits, more aggressive health education to encourage parents/caregivers to present with their children early are recommended. Preventive Cardiology through routine mass education and community cardiac outreach with screening ECG and Echocardiography is also very advisable. Setting up well-structured foundation for indigent children with CHD and other cardiovascular diseases, training and re-training of medical staff and government providing an enabling environment for more corrective surgeries to be performed in our hospitals should be advocated.

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