Congenital Heart Diseases associated with Identified Syndromes and Other Extra-Cardiac Congenital Malformations in Children in Lagos

Maladies cardiaques congénitales associées à des syndromes identifiés et extra-Malformations cardiaques congénitales chez les enfants à Lagos

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ABSTRACT
BACKGROUND: Congenital heart diseases are commonly associated with other extra cardiac congenital malformations. OBJECTIVE: To identify congenital heart diseases associated with identified syndromes and other extra cardiac congenital malformations in children in our hospital. METHODS: A prospective descriptive study done on children with congenital malformations referred to the Lagos University Teaching Hospital, Nigeria (LUTH) for echocardiographic evaluation. A thorough 2D assessment of the chambers, septa, heart vessels and concordance of the atrium and ventricle and the great vessels was made. Echo-cardiographic data obtained included M mode direct measure-ments of dimensions of left atrium, aortic root, right ventricular outflow tract, left ventricle in diastole/systole, wall thicknesses – right ventricular wall, interventricular septum, left ventricular posterior wall. Fractional shortening was derived from M mode data. Final diagnosis of the congenital heart disease was recorded.

RESULTS: A total of 101 children with congenital malformations had echocardiography studies done as part of their clinical evaluation, 15(14.9%) were neonates, 53(52.5%) infants 25(24.8%) were aged one to five years and 8(7.9%) were above five years of age. Recognised syndromes were seen in 69(68%) cases. Down syndrome with 54 children contributed 78.3% of those with known syndromes. Other identified syndromes and associations were Marfan’s, Noonan’s, Edwards, Prune Belly, Apert, Ellis-van creveld syndrome and congenital rubella syndrome. Congenital heart diseases were detected in 73(72.3%) patients while 28(27.7%) had no heart defect. The commonest identified congenital heart disease was ventricular septal defect affecting 30 (29.7%) patients.

CONCLUSION: Congenital heart diseases are common in children with congenital malformations. Down syndrome was the most common malformation and the congenital heart disease most associated with the congenital malformation was ventricular septal defect. This study emphasizes the need for cardiac assessment of children with congenital malformations.

Key words: Congenital malformations, congenital heart diseases, children, echocardiography, abnormalities, anomalies.

RÉSUMÉ


RÉSULTATS: Un total de 101 enfants avec des malformations congénitales a echocardiography études réalisées dans le cadre de leur évaluation clinique, 15 (14,9%) étaient des nouveau-nés, 53 (52,5%) des nourrissons 25 (24,8%) étaient âgés de un à cinq ans et 8 (7,9 %) étaient au-dessus de l’âge de cinq ans. Reconnaît syndromes ont été observés dans 69 (68%) des cas. Le syndrome de Down, avec 54 enfants a contribué 78,3% des personnes ayant connu des syndromes. Parmi les autres syndromes de Marfan et les associations ont été, de Noonan, de Edwards, Prune Belly, Apert, Ellis-van Creveld syndrome et le syndrome de rubéole congénitale. Maladies cardiaques congénitales ont été détectées dans 73 (72,3%) patients tandis que 28 patients (27,7%) n’avaient pas d’anomalie cardiaque. La commune a identifié une maladie cardiaque congénitale du septum ventriculaire a été vice affectant 30 (29,7%) patients.

CONCLUSION: Les maladies cardiaques congénitales sont communs chez les enfants atteints de malformations congénitales. Le syndrome de Down est la plus fréquente de malformation cardiaque congénitale et de la maladie associée à la plupart des malformations congénitales a été ventriculaire septal defect. Cette étude souligne la nécessité d’une évaluation cardiaque des enfants ayant des malformations congénitales.


Mots-clés: malformations congénitales, les maladies cardiaques congénitales, des enfants, l’échocardiographie, les anomalies, les anomalies.
INTRODUCTION

Congenital malformations are anatomical defects and chromosomal abnormalities present at birth. Approximately 2% of live births have major congenital abnormality. The incidence is increased in pre-term and small for gestational age infants. Congenital anomalies are among the leading cause of death during the first five years of life in the USA. Congenital malformations accounts for 8.5% neonatal deaths in Lagos University Teaching Hospital, Lagos (LUTH). The externally observed congenital malformations often have other associated internal organ malformations. In the past, due to lack of advanced diagnostic techniques many internal organ malformations were missed. Currently, technological advancement in diagnosis has made it possible to identify the internal organ malformations thus preventing a congenital heart disease to be passed off as pneumonia. This may also have contributed to the seeming increase in congenital malformations in children. A single system can be affected or multiple systems. In a study in Australia, of the 175.9/10,000 cases of congenital malformations which affected all systems, the single most affected system was the heart with 35.9/10,000 cases.

Congenital heart diseases are commonly associated with other extra cardiac congenital malformations and some are particularly seen in certain malformations. This knowledge is very useful in areas where sophisticated diagnostic facilities such as echocardiography machines are absent to make specific diagnosis, guide treatment and hereby reduce mortality. To our knowledge, no such study has been done in our environment ante mortem using echocardiography.

The aim of this study was to document the congenital heart diseases associated with identified syndromes and other extra cardiac congenital malformations in children in Lagos using echocardiography.

SUBJECTS AND METHODS

Over a period from March 2005 to March 2007 a prospective study was carried out on children with congenital malformations referred to the Lagos University Teaching Hospital, Nigeria (LUTH) for echocardiographic evaluation. Lagos University Teaching Hospital is a 760 bed tertiary institution in the former capital and economic nerve centre of Nigeria. It is a major referral centre in Nigeria with patients coming from both private and public health facilities all over Nigeria especially the western part. The heart of each patient was evaluated using a HP Sonos 2000 echocardiography machine and both M mode and 2D findings recorded. Transducers with 7.5/4.5 or 3.5MHz were used as determined by age of the patient. The echocardiographic studies were performed by ENE and AA using standard techniques, views, and measurements. A thorough 2D assessment of the chambers, septa, heart vessels and concordance of the atrium and ventricle and the great vessels was made. Echocardiographic data obtained included M mode direct measurements of dimensions of left atrium, aortic root, right ventricular outflow tract, left ventricle in diastole/systole, wall thicknesses – right ventricular wall, interventricular septum, left ventricular posterior wall. Fractional shortening was derived from M mode data.

All children without obvious external congenital malformations were excluded. Congenital heart disease was defined as a structural abnormality of the heart such as defects in cardiac sepaation, abnormalities of ventriculo-arterial connections, rudimentary or absent chambers, abnormalities of ventricular inflow and outflow, and abnormal vascular connections and structures. Final diagnosis of the congenital heart disease was recorded. Biographic data of patient and parents, clinical diagnosis of the child were then obtained using a structured questionnaire. Consent was obtained prior to recruitment into study.

Data was analysed using SPSS 11.0 software. Frequency distribution tables were generated for the congenital malformations and associated congenital heart diseases.

RESULTS

Demographic Data

A total of 101 children with congenital malformations had echocardiography studies done as part of their clinical evaluation. Their ages ranged from birth to 14 years. Among the 101 children, 15(14.9%) were neonates, 53(52.5%) infants, 25(24.8%) were aged 1to 5 years and 8(7.9%) were above 5 years. (Table 1). Fifty-four (53.5%) of the 101 children with congenital malformations were males while 47(46.5%) were females giving a male: female ratio of 1.1:1. In the different age group categories, the sexes were almost equally shared except for the neonates where the males made up 66.7% of the cases.

The age of the fathers of the children ranged from 26 years to 65 years with a mean of 39.6(7.1) years. The age of the mothers of the children ranged from 23years to 50years with a mean of 34.1(5.9) years. There was no case of consanguinity in all cases studied.

Table 1: Distribution of Patients by Sex and Age

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Number(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age Group</td>
<td></td>
</tr>
<tr>
<td>Neonates</td>
<td>15(14.9)</td>
</tr>
<tr>
<td>Infants</td>
<td>53(52.5)</td>
</tr>
<tr>
<td>1–5years</td>
<td>25(24.8)</td>
</tr>
<tr>
<td>&gt;5years</td>
<td>8(7.9)</td>
</tr>
<tr>
<td>Sex Distribution</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>54(53.5)</td>
</tr>
<tr>
<td>Female</td>
<td>47(46.5)</td>
</tr>
</tbody>
</table>

Congenital Malformations

Recognised syndromes were seen in 69(68%) cases. Down’s syndrome with 54 children contributed 77.1% of those with known syndromes. Other identified syndromes and associations were Marfan’s, Noonan’s, Edwards, Prune Belly, Apert, Ellis-van creveld syndrome and congenital rubella syndrome. (Table 2).

There were 22 musculoskeletal defects, 20 of which were defects of the palate and or lip, choanal atresia and one case of syndactylly and right leg below knee amputation. There were also central nervous system defects seven(6%) and GIT defects one(1%).

Associated Congenital Heart Diseases

Congenital heart diseases were detected in 73(72.3%) patients while
28(27.7%) had no heart defect. The commonest identified congenital heart disease was ventricular septal defect (VSD) affecting 30 patients (Figure 2). This was followed by atrioventricular canal defect (AVCD) affecting 23(22.7%) children; twenty of these children with atrioventricular canal defect had Down syndrome. Other congenital heart diseases were atrial septal defect (ASD) 20(19.8%), patent ductus arteriosus (PDA) 17(16.8%), tetralogy of fallot (TOF) 9(8.9%), aortic root dilatation 3(2.9%) and mitral valve prolapse 3(2.9%). There were two cases of dextrocardia and one case of each of pulmonary stenosis, coarctation of the aorta and double outlet right ventricle. (Table 2) This showed that VSD, AVCD, ASD, PDA and TOF were the top 5 commonest congenital heart diseases.

Some cases had more than one cardiac defect. ASD, Atrial septal defect; AVCD, Atrioventricular canal defect; COA, Coarctation of the aorta; DORV, Double outlet right ventricle; MVP, Mitral valve prolapse; PDA, Patent ductus arteriosus; PS, Pulmonary stenosis; TOF, Tetralogy of fallot; VSD, Ventricular septal defect.

9(8.9%), aortic root dilatation 3(2.9%) and mitral valve prolapse 3(2.9%). There were two cases of dextrocardia and one case of each of pulmonary stenosis, coarctation of the aorta and double outlet right ventricle. (Table 2) This showed that VSD, AVCD, ASD, PDA and TOF were the top 5 commonest congenital heart diseases. Figure 1.

Down Syndrome with 54(53.5%) children was the single most common congenital malformation with 77% of them having a congenital heart disease. Only seven (6.9%) cases did not have a heart defect. Among the 47 who had congenital heart defects, 41(87.2%) had ventricular septal defect 21 (20.8%) and atrioventricular canal defect 20(19.8%). The other syndromes and their associated heart defects are also shown in Table 2.

Musculoskeletal abnormalities followed Down’s Syndrome with 22(21.8%) cases. Twenty of these were cleft lip and/or palate. Fourteen (70%) of the 20 cases with cleft lip and/or palate had no heart defect. There were two cases of cleft lip and none had a heart defect. Of the eight cases with cleft palate six had no cardiac defect while the other two had atrial septal defect and Tetralogy of Fallot with patent ductus arteriosus respectively. Ten (9.9%) patients had cleft lip and palate and six of them had no heart defect. The four with heart defects comprised of two cases of ventricular septal defect, a case of atrial septal defect and a case of dextrocardia with complete atrioventricular canal. (Table 2).

DISCUSSION
The congenital heart diseases can occur either on their own or in association with other malformations in an individual. In this study, congenital heart diseases were detected in 73(72.3%) of the 101 patients with congenital malformations. This is a very high rate. The incidence of congenital heart diseases in children varies quite considerably, depending on the study population and methodology.
There were two cases of Edward’s syndrome especially ventricular septal defect. The two cases had double outlet right ventricle and a combination of atrial and ventricular septal defects respectively.

Ellis-van Creveld syndrome (chondroectodermal dysplasia) is reported to be associated with single atrium. The only case in this report had tetralogy of Fallot. Turner’s syndrome was associated with coarctation of the aorta as expected. Apert’s syndrome is reported to be occasionally associated with congenital heart defects. The case with Apert’s syndrome in the study had no congenital heart disease. Prune Belly Syndrome is a urinary tract anomaly usually has no association with heart defects and the only case in the report had no congenital heart disease.

Lesions of the musculoskeletal system followed Down’s syndrome with 22 cases. Twenty of them were defects of the palate and or lip. Whereas 100% of the cases with cleft lip had no cardiac defect, 75% of those with cleft palate had cardiac defects and 60% of those with both cleft lip and palate had no cardiac defect. It therefore appears that a patient with cleft palate is more likely to have an associated heart defect than a patient with cleft lip. More subjects with cleft lesions need to be studied to confirm this observation.

Congenital anomalies are a major cause of morbidity and mortality with congenital heart diseases contributing to this. Identification and correction of any identified lesion will reduce the mortality burden from this cause. There is need for an early and proper cardiovascular evaluation of all children with congenital anomalies. Assessment of all newborns with Down’s syndrome with an echocardiogram is the standard recommendation. This requires facilities and trained personnel for cardiac evaluation and surgery to cater for this group of patients in our sub region.

There has been a dramatic improvement in the prognosis of children with congenital heart diseases in regions where the technology and personnel to carry out cardiac interventions are available. It is reported that in 1986 60% of deaths from congenital heart disease occurred in the first year of life, whereas in the 1990s the majority of deaths occurred in adults over the age of 20 years. It is predicted that 78% of the babies born with congenital heart disease today will survive into adulthood. A national policy for Paediatric heart care and International collaboration to build effective and sustainable cardiac surgery programmes are required.

This result confirms the well known association of some congenital abnormalities such as Down’s, Edwards, Turner’s, Marfan’s and rubella syndromes in our sub Saharan African environment. It demonstrates a high association of congenital heart diseases with extra cardiac congenital malformations and Down Syndrome to be the most commonly seen congenital anomaly with an 87% rate of associated congenital heart disease. This knowledge serves as a guide for the region in evaluating children with congenital malformations. It shows that the reported associations of some congenital malformations with congenital heart defects in the developed countries are similar to what pertains in the tropics. Hence management of the patient can be guided by these associations when sophisticated equipment for thorough diagnosis is absent.

REFERENCES

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