Prenatal Detection of Critical Congenital Heart Disease

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ABSTRACT

Congenital heart disease (CHD) is a leading cause of infant mortality and 30% of fetuses born with CHDs have other associated malformations and chromosomal abnormalities. Prenatal diagnosis also allows parents to opt for termination of the pregnancy.

Keywords: Critical congenital heart disease, Fetal echocardiography, Prenatal diagnosis.

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INTRODUCTION

When discussing congenital heart disease (CHD), consideration of certain points is critical. The first consideration is that CHD is a leading cause of infant mortality, with an estimated incidence of about 4 to 13 per 1000 live births.1,2 Moreover, between 1950 and 1994, 42% of infant deaths reported to the World Health Organization were attributable to cardiac defects.3 The second consideration is that 30% of fetuses born with CHDs have other associated malformations or chromosomal abnormalities. The third consideration is that the vast majority of CHDs affect pregnancies with no previously known risk factors. The fourth consideration is that prenatal diagnosis of CHDs improves the outcome in certain cardiac conditions, which include hypoplastic left heart, transposition of the great arteries, and coarctation of the aorta.4,5 In special situations, prenatal diagnosis of CHDs gives parents the choice of the place of delivery. Prenatal diagnosis may also allow parents the option of termination of the pregnancy.

Antenatal diagnosis of cardiac defects did not witness a great improvement over the last few years.6,7 Furthermore, prenatal detection rates of CHD vary widely.8 Some of this variation can be attributed to differences in examiner experience, maternal obesity, transducer frequency, abdominal scars, gestational age at the time of ultrasound examination, amniotic fluid volume, and fetal position.9,10 Because of the above, efforts are being made to try to improve the detection rates and diagnosis of CHD. These efforts include classroom courses, online courses, training in referred centers, and introduction of guidelines and protocols to diagnose CHD by professional bodies, such as the International Society of Ultrasound in Obstetrics and Gynecology (ISUOG).

HOW TO EXAMINE THE FETAL HEART DURING ROUTINE ANOMALY SCAN

A strict protocol should be followed, which should include the subsequent points. Firstly, examination of all fetuses should be done at gestational age between 18 and 22 weeks. Secondly, the abdominal probe should be done using high-resolution ultrasound. The zoom and cine loop facilities should be used when needed. Thirdly, in all fetuses, a four-chamber view of the heart, with comments on situs, atrial chambers, ventricular chambers, and atrioventricular junction and valves should be viewed. The left and right ventricular outflow tracts should be examined. Fourthly, outflow tracts are imaged by tilting the probe toward the fetal head. The great arteries should be of equal size and should cross at approximately 90° (Figs 3 and 4). Fifthly, all fetuses should have a three-vessel view, which enabled diagnosis of conditions such as coarctation of the aorta, right aortic arch, double aortic arch, and vascular rings (Fig. 5). Sixthly, all fetuses had a three-vessel and trachea view. Using this view, the transverse aortic arch and its relationship to the trachea could be demonstrated. Seventhly, color Doppler mapping should be used to examine the various cardiac structures and to improve the detection of abnormal blood flow. Many congenital heart defects can be seen on the standard four-chamber view (Table 1). However, four-chamber views are not enough to diagnose all cardiac malformations because of many factors, such as the anomaly could be far away from the four-chamber view, the anomaly evolved in utero, routine checklist may be overlooked, and inadequate examination.

Examination of the outflow and outflow trachea tract is essential to diagnose other congenital cardiac defects (Table 2).
Fig. 1: Four-chamber view, showing both atria of similar size, ventricles of similar size, foramen ovale flap in the left atrium.

Fig. 2: Four-chamber view showing both ventricles on equal size, intact septum, moderator band seen in the right ventricle, and different levels of attachment of atrioventricular valve.

Fig. 3: Left outflow tract. The anterior aortic wall is continuous with the intact interventricular septum.

Fig. 4: Right outflow tract. The pulmonary artery bifurcates immediately. The pulmonary valve moves freely.

Fig. 5: The three-vessel view. This view demonstrates the relationship between superior vena cava, aorta, and pulmonary artery. Note the correct position and the size of the three vessels as well as correct alignment.

Table 1: Congenital cardiac defects detected on four-chamber view
- Hypoplastic right and left ventricle
- Atrioventricular canal defect
- Double inlet ventricle
- Ebstein's anomaly
- Single ventricle
- Cardiac tumors
- Large ventricular septal defect

Table 2: Congenital cardiac defects missed on four-chamber view
- Transposition of the great vessels
- Tetralogy of Fallot
- Double outlet right ventricle
- Truncus arteriosus
- Small/moderate ventricular septal defect
- Atrial septal defect
- Coarctation of the aorta
- Total anomalous pulmonary venous return
- Aortic stenosis
- Pulmonary stenosis
DISCUSSION

When we examine the fetal heart during anomaly scan, we aim to screen for any critical congenital heart defect which may affect the life of the newborn. Detection and diagnosis are distinct terms. Detection is the process of discovering a finding, while diagnosis is the process of identifying a disease by examination. In many fetomaternal centers, such as our center in Doha, detection rate is very high, as we were able to discover 10 out of 11 cases of CHD. The accuracy of our diagnosis of CHD is 100%, as all those who were referred to the pediatric cardiologist had their diagnosis confirmed postnatally via echocardiography. This result demonstrates that a high detection rate and accurate diagnosis of CHD is possible.

Such result is very encouraging as only a few years ago the detection rate of CHD was suboptimal. A European study between 2000 and 2005 involving 16 countries and including $3.3 \times 10^6$ live births showed the prenatal detection rate of CHD is only 20%. Many other studies done in European countries showed that the detection rate varies dramatically, between 20 and 80%, emphasizing the need for additional studies and an improvement in detection rate.11-13

Another study by Galindo et al14 involving 33 centers in Spain, and including 1,060 major CHDs between 2004 and 2006, reported that the prenatal detection rate of CHD in routine second-trimester scans was as low as 52.6% (95% confidence interval (CI) 45.6–60.8). The study identified two important predictors of an increased detection rate by center: firstly, a systematic approach to the examination of the heart, showing at least the four-chamber view and the outflow tracts (prevalence ratio 1.3, 95% CI 1.0–1.8) and secondly, the availability of specialists in fetal echocardiography to perform the examination (prevalence ratio 1.4, 95% CI 1.1–1.9).15 The study advised that prenatal detection of CHD should be globally strengthened by way of a uniform approach to the cardiac examination along with improving examiner training and skills. Moreover, authors of the study recommended close collaboration with fetal heart specialists to help accomplish this aim.15 A prospective observational study at a London teaching hospital reported similar conclusions. Authors encouraged continuous feedback-based training of health care professionals, a low threshold for echocardiography referrals, and convenient access to fetal heart specialists as techniques to improve the effectiveness of a screening for CHDs.11

A study by Marek et al,16 including all pediatric and fetal patients with CHD over a 21-year period in the Czech Republic, reported a discrepancy in the detection rate of various cardiac defects. For example, the prenatal detection rate for hypoplastic left heart was 95.8% (Fig. 6), whereas detection of transposition of the great arteries was only 25.6%.

Prenatal detection of CHD is essential because it has a large impact on the appropriate management of affected infants during labor and delivery. For example, some babies may need to be delivered in a cardiac operation room for quick intervention. When the fetus is delivered, there are critical changes in the fetal circulation which include initiation of breathing, conversion of the fetal circulation from a parallel circuit to a series circuit, increase in the systemic vascular resistance, decrease in pulmonary vascular resistance, and closure of fetal shunt pathways. Detection of CHD during the antenatal period allows parents and physicians to choose the place of delivery and prepare a multidisciplinary team for the baby’s arrival, which includes a fetal medicine specialist, pediatric cardiologist, neonatologist, anesthesiologist, and pediatric cardiac surgeon. These preparations improve the prognosis of the infant.

The accepted gestational age for performing fetal echocardiography is between 18 and 22 weeks. However, a small proportion of CHDs become evident or more obvious as pregnancy progresses. Thus, the cardiac evaluation can be normal at 18 weeks, although a significant malformation is diagnosed later on in the pregnancy or after birth. This is true of some cases of aortic or pulmonary stenosis, cardiac tumors, and cardiomyopathies (Fig. 7). Fortunately, it is rare for a life-threatening malformation to arise after 20 weeks’ gestation. In addition, minor lesions, such as small ventricular septal defects can be overlooked because of the limitation of ultrasound resolution (Fig. 8). Other defects, such as persistent arterial duct and atrial septal defect cannot be predicted prenatally as these communications are always present.
Prenatally. Thus, there are confidence limits with even detailed fetal heart scanning. These confidence limits may be much wider with poor image quality. For this reason, it is crucial to counsel the parents on these limitations of antenatal scanning and to ensure that they are aware that while prenatal screening presents a huge benefit to their fetus, the detection is not 100%.

**WHAT IS NEW?**

**The New 3D/4D-based Spatiotemporal Imaging Correlation in Fetal Echocardiography**

Spatiotemporal imaging correlation is an automated device incorporated into the ultrasound probe and has the capacity to perform slow sweep to acquire a single three-dimensional (3D) volume. This acquired volume is composed of a great number of two-dimension (2D) frames. This volume can be analyzed and reanalyzed as required to demonstrate all the required cardiac views. It also provides the examiner with the ability to review all images in a looped cine sequence. Spatiotemporal imaging correlation has the potential to improve our ability to diagnose CHD. This potential becomes possible because of unlimited number of images available for viewing, additional information not available in the standard 2D ultrasound images, information can be shared with other colleagues remote from the site of examination for second opinion, and the patient can be counseled in great detail in light of new information.

In conclusion, the majority of prenatally detectable cases of CHD occur in patients without any risk factors or extracardiac anomalies. Several studies have reported the detection rate of CHD; however, these figures vary widely. Regardless, these studies all came down to the general agreement that an improvement in detection rate is necessary, as it can greatly improve the prognosis of the infant. It also allows hospital systems to prepare for the arrival of the infant by making sure a multidisciplinary team is available. Detection rate can be improved by implementing a uniform, systematic approach to the fetal cardiac examination and by ensuring that fetal echocardiography is performed by specialists who are familiar with the prenatal diagnosis of CHD.

**REFERENCES**


**Fig. 7:** A case of rhabdomyoma which is a benign cardiac tumor; however, it could be associated with tuberous sclerosis which is autosomal dominant condition

**Fig. 8:** Ventricular septal defect can occur in any part of the interventricular septum and is often associated with other cardiac defects


