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Svetlana Vrzić-Petronijević¹,²†, Miloš Petronijević¹,², Vojislav Parezanović¹,³, Jelena Stamenković-Dukanac¹,², Zorica Jestrović², Danijela Bratić²

Fetal echocardiography – 25-year experience
Фетална ехокардиографија – 25 година искуства

¹ University of Belgrade, School of Medicine, Belgrade, Serbia; ² Clinical Centre of Serbia, Clinic of Gynecology and Obstetrics, Belgrade, Serbia; ³ University Children's Hospital, Belgrade, Serbia

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† Correspondence to:
Svetlana VRZIĆ-PETRONIJEVIĆ
Clinic of Gynecology and Obstetrics, 11000 Belgrade, Serbia
E-mail: vrzic.dr@gmail.com
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**SUMMARY**

**Introduction/Objective** Congenital heart diseases are the most common congenital anomalies. The objective of the study was to determine reliability, specificity and sensitivity of the fetal echocardiography in detection of congenital heart diseases in referral center for fetal echocardiography.

**Methods** We analyzed 14 500 fetal echocardiography exams (FEC) between 1991 and 2014 which were performed in two tertiary centre.

**Results** The average maternal age at time of diagnosis was 32 years. The mean gestational age at the time diagnosis was 25.9 weeks. The most common indication for FEC were suspicious abnormal cardiac findings in obstetrical screening sonography (50.6%). Among total of 9055 examined fetuses, pathological finding on the fetal heart was found in 638 cases. The most common congenital heart diseases were structural anomalies of the fetal heart in 81%, of which 13.3% were fetuses with hypoplastic left heart syndrome and 11.2% with ventricular septal defect. Of all fetuses with diagnosed congenital heart defect, 46.2% were born alive and had good postnatal prognosis, while 2.7% died in utero, and in early neonatal period died 10.6%. Pregnancy was terminated in 40.4% fetuses with severe congenital heart defect. Sensitivity of diagnostic procedure in our study was 95.9% and specificity was 99.9%.

**Conclusion** Our study proves that FEC is reliable, informative diagnostic tool in detecting congenital heart defects with high specificity and sensitivity. With multidisciplinary aproach it provides optimal time window for improving perinatal outcome.

**Keywords:** prenatal diagnosis; congenital heart defect; perinatal outcome; fetal echocardiography

**INTRODUCTION**

Congenital heart diseases (CHD) are one of the most common congenital anomalies, with incidence 4-13 per 1000 newborns [1]. Early prenatal diagnosis is considered essential because it allows early intervention. Fetal echocardiography as diagnostic procedure was introduced between 1970 and 1980 [2, 3]. With the advancement of technology and interdisciplinary approach prenatal diagnosis of CHD with fetal echocardiography has resulted in a decrease of the morbidity and mortality of these patients [4, 5].

The aim of this study was to analyze recent trends in the indications of fetal echocardiography and to evaluate the reliability of fetal echocardiography as a method for prenatal diagnosis CHD at a tertiary referral center in Belgrade, Serbia.
METHODS

We analyzed the fetal echocardiography databases from Clinic of Gynecology and Obstetrics, Clinical Centre of Serbia and University Children’s Hospital in Belgrade in period from 1991 to 2014, and reviewed the medical records of 9055 fetuses from 8838 pregnancies (217 multiple gestations). We divided all findings on normal and abnormal and then analysed abnormal findings. The clinical data we collected for this retrospective study were gestational age (GA) at the time of diagnosis, maternal age at the time of diagnosis, indications, results of fetal echocardiography and perinatal outcome. Fetal echocardiography was performed by experienced perinatology specialist and pediatric cardiologist.

We classified indications for fetal echocardiography among the diagnosed CHD into 5 categories: abnormal cardiac findings at obstetric ultrasonography, extracardiac findings at obstetric screening, maternal risks, familial risks and teratogenic risk factors.

All results were classified to normal and abnormal. All abnormal results were divided in 3 subclasses: structural anomalies, arrhythmias, and structural anomalies with arrhythmias. Structural anomalies were then stratified in 5 subclasses: valvular/obstructive abnormalities of left/ right heart (HLHS, aortic stenosis, coarctation of the aorta, stenosis and atresia of the mitral valve/ stenosis and atresia of the pulmonary valve, atresia and dysplasia of the tricuspid valve and Mb. Ebstein), shunt lesions (ASD, VSD, AVSD, TAUPV), complex abnormalities (tetralogy of Fallot with or without agenesis of the pulmonary valve, complete transposition of the great arteries, double outlet right ventricle, common pulmonary artery), cardiomyopathy and rare anomalies.

Perinatal outcome and treatment options were analyzed according obtained data for each class of diagnosed CHD.

Obtained data was presented descriptively, by percentage and indicators of diagnostic accuracy: sensitivity and specificity.

RESULTS

Indications

The most common indication for fetal echocardiography was abnormal cardiac finding in obstetric screening in 323 pregnancies (50.6%). The second most common indication was abnormal extracardiac finding in obstetric screening (245 pregnancies - 38.4%). Positive family history was indication for 41 pregnant women (6.4%). Maternal risk factors were indications in 26 (4%) and teratogenic factors in 3 (0.4%) cases.

Fetal echocardiography results in fetuses with congenital heart disease

Among the 9055 pregnancies, CHD was diagnosed in 638 (7.2%) of cases.

Of 638 fetuses with CHD, structural anomaly was found in 515 (81%) of cases. Valvular/obstructive abnormalities of left heart were found in 106 (20.1%) fetuses: HLHS in 70 (13.3%),...
coarctation of the aorta in 22 (4.1%), atresia of the mitral valve in 8 (1.5%) and aortic stenosis in 6 (1.1%) of cases.

Valvular/obstructive abnormalities of right heart were found in 95 (18.0%) fetuses: pulmonary artery atresia in 32 (6.0%), pulmonary artery stenosis in 15 (2.8%), tricuspid valve atresia in 15 (2.8%), tricuspid valve dysplasia in 16 (3.0%), and M. Ebstein in 17 (3.2%) of cases.

Shunt abnormalities were found in 123 (23.3%) fetuses: ASD in 16 (3.0%), VSD in 59 (11.2), AVSD in 46 (8.7%), and TAUPV 2 (0.4%) of cases.

Complex CHD were found in 158 (29.9%) fetuses: T. Fallot in 30 (5.6%), T. Fallot and agenesis of pulmonary artery in 9 (1.7%), univentricular heart in 29 (5.5%), common truncus arteriosus in 10 (1.9%), DORV in 46 (8.7%), TGA in 29 (5.5%), and L-TGA in 5 (0.9%) of cases.

Cardiomyopathy was found in 13 (2.5%) and rare anomalies in 33 (6.2%) fetuses.

Isolated arrhythmia had 110 (17.0%) fetuses. Structural anomaly with arrhythmia had 13 (2.0%) fetuses.

**Perinatal outcome in fetuses with congenital heart disease**

Among 638 fetuses with CHD, 295 (46.2%) were born alive and survived neonatal period. The highest survival rate was noticed in fetuses with arrhythmias (33.5%) and shunt anomalies (24%). Of the total number of diagnosed CHD, the most common anomalies among live-born children were isolated VSD (6.5%), TGA (3.3%) and T. Fallot (2.9%).

In utero, lethal outcome occurred in 17 fetuses (2.7%). The most common structural anomaly in fetuses from this group was HLHS (11.8%).

In the early neonatal period died 68 (10.6%). The highest mortality rates were in cases of cardiomyopathy (30.8%) and obstructive lesions of the left heart (21.7%).

Pregnancy was terminated in 258 (40.4%) cases with CHD.

**Sensitivity and specificity of fetal echocardiography**

Prenatally, 26 neonates with CHD were not diagnosed (12 VSD, 4 stenosis of pulmonary artery, 1 T. Fallot with agenesis of pulmonary artery, 1 atresia of pulmonary artery with VSD, 3 critical coarctation of the aorta, 1 critical stenosis of the aorta in twin pregnancy, 1 TGA, 2 TAUPV and 1 aortic-pulmonary septum).

In 7 fetuses prenatally diagnosed CHD was not confirmed postnatally.

Sensitivity of procedure in our group was 95.2%, and specificity 99.91%.

**DISCUSSION**

Identification of congenital heart defects is important because they are the most common congenital anomalies, and in 50% they are the leading cause of neonatal deaths [6,7]. Pregnant women with increased risk for giving birth to a child with CHD require more detailed evaluation of the fetal heart [8] as well as those with high risk for aneuploidy based on the combined screening [9].
Theoretically, most congenital heart defects can be diagnosed prenatally and a suspicion can be set already at the recommended routine evaluation of fetal anatomy in the second trimester.

In a study from 1997 conducted in Belgrade, basic indications were family history in 42.6%, maternal infection in the first trimester (11.5%), maternal diabetes (9.6%) and indirectly suspected cardiac anomaly because of the clinical manifestations of polyhydramnios or fetal arrhythmia (28.4%) [10]. In 2004, Friedberg reported that the most common indication were family history of CHD (23%), and maternal diabetes (18%). Obstetrical ultrasound indicating CHD accounted for only 13% [11]. Obstetric screening has only recently found an even greater prevalence of cardiac abnormalities.

Our results show that approximately half of all fetal echocardiographic exams were performed because of abnormal cardiac findings at the obstetric screening (50.6%). Next common indication was extracardiac finding at obstetric screening (38.44%). Other indications were: familial (6.41%), maternal (4.06%) and teratogenic risk factors (0.47%).

Prenatal diagnosis of congenital heart disease allows parents to receive information on the diagnosis, the further course of pregnancy and possible treatment before or after birth, making easier for them to make a decision whether to continue the pregnancy or to terminate it earlier.

In our investigation that included 9055 fetuses, CHD was diagnosed in 638 of cases. The incidence of CHD was 7.2%. This percentage is significantly higher than the incidence of CHD in the general population, because it is the population of pregnant women with increased risk for the development of fetal CHD. It is noticed that the prevalence of newborns with CHD is persistent over the last 15 years [12].

The most frequent were valvular obstructive lesions of the left heart, isolated in 201 (38.1%), and associated with other anomalies in total of 310 (58.7%). The most common of obstructive abnormalities were the hypoplasia of the left heart diagnosed in 70 (66.3%) cases and pulmonary artery atresia (5.94%).

Isolated shunt lesions were diagnosed in 123 (23.3%) and associated with other anomalies in total of 221 (41.7%). Prenatal shunt and valvular lesion ratio is 1.6: 1, and it does not correlate to the postnatal relationship, which is 1.3: 1. Thus, our result correspond to those in the literature, where this ratio is 1.2-1.4:1 [13, 14]. Those findings are understandable considering specific characteristics of fetal circulation.

Prenatally diagnosed complex congenital heart disease reduces neonatal morbidity and mortality. Out of the total number of diagnosed CHD, 295 (46.2%) newborns survived the fetal and neonatal period. These children had hemodynamically stable anomalies with a relatively favorable prognosis. The higher survival rate was noted in a group of children with: isolated VSD (6.5%), TGA (3.3%) and T. Fallot (2.9%). There were no survivals in group with hypoplastic left heart (0/70).

Lethal outcome in utero occurred in 17 fetuses (2.7%). The incidence of intrauterine fetal death in a group of fetuses with CHD was 2.7% (17/638). Supraventricular tachycardia with heart failure was the leading cause of intrauterine demise in 35.3%, followed by HLHS in 11.8%.
In the early neonatal period died 68 children (10.6%). The highest mortality rate was among the neonates with cardiomyopathy (30.8%) and obstructive lesions of the left heart (21.7%). In our country the prognosis of these pregnancies is unfavorable due to limited treatment options (heart transplantation e.g.), so the choice of treatment is usually reduced to palliative care.

The outcome of pregnancy after the diagnosis CHD depends on many factors, but certainly the most important is the type of defect, its prognosis and impact on quality of life.

Termination of pregnancy may be the option in cases of complex heart defects known to have poor prognosis (HLHS) which could be diagnosed already at the level of screening, but also in anomalies showing significant deterioration during repeated prenatal checks. In our study termination of pregnancy was done in 258 (40.4%) cases after diagnosis of CHD. All fetuses had complex heart defects, and some also had a chromosomal aberrations or extracardiac anomalies. In Serbia, termination of pregnancy for medical indications is legally defined and is not limited by gestational age.

The sensitivity of fetal echocardiography in our study was 95.9% and specificity 99.9%. The positive predictive value was 95.9% and the negative predictive value was 99.9%. These values were correlated with the results obtained from large global studies. The sensitivity is around 42%, a specificity of about 98%. The positive predictive value of echocardiography is about 90%, and negative predictive value was 93% [15].

The rate of detection in tertiary institutions is around 85-95% [16]. The largest discrepancies are related to the atroventricular morphology and outlets of the great arteries. Our study shows that high accuracy rate could be achieved by obstetrician if sonographer is experienced and preferably trained by pediatric cardiologist specialized in fetal echocardiography.

In our study, 26 neonates with CHD were not diagnosed prenatally (12 VSD, 4 stenosis of pulmonary artery, 1 T. Fallot with agenesis of pulmonary artery, 1 atresia of pulmonary artery with VSD, 3 critical coarctation of the aorta, 1 critical stenosis of the aorta in trigeminal pregnancy (in one of triplets), 1 TGA, 2 TAUPV and 1 aortic-pulmonary septum). In 7 fetuses diagnosis made prenatally wasn’t confirmed postnataally. False positive fetal echocardiography findings are rare making the specificity of this method high, over 90%. Most frequent false positive diagnosis were VSD and coarctation of the aorta.

Our study proves that fetal echocardiography is reliable, informative diagnostic tool in detecting congenital heart defects with high specificity and sensitivity. With multidisciplinary aproach, given the fact that Serbia is still country in transition with limited options for congenital heart defects treatment, it provides optimal time window for improving perinatal outcome.

REFERENCES