Prenatal Diagnosis of a Rare Case of Double-Outlet Right Ventricle with Tricuspid Atresia

Dear Editor,

Double-outlet right ventricle (DORV) represents a rare group of cardiac anomalies characterized by both great arteries arising primarily from the RV. This disease is a conotruncal anomaly as a complex heart disease and comprises less than 1% of all congenital heart anomalies.\(^1, 2\) The first diagnostic criterion for DORV is the rise of both the aorta and the pulmonary artery from the morphologic RV. Other cardiac malformations are always present with the frequent coexistence of multiple abnormalities. These may include 1 or more ventricular septal defects, left ventricular outflow obstruction, and atrioventricular valve abnormalities.\(^3, 4\) DORV may occur with any atrial situs and many other variations; nonetheless, situs solitus and atrioventricular concordance are the most common findings.\(^4\)

We herein present a very rare case of DORV, in tandem with tricuspid atresia, transposition of the great arteries, pulmonary stenosis, and ventricular septal defect.

A 35-year-old pregnant woman (gravida 2, para 1, and 1 living child with a history of congenital heart disease) was referred for fetal cardiac assessment at 18th weeks of gestational age.

A transabdominal 2D echocardiogram in the axial view showed viscera atrial situs solitus, levocardia, and levoposition. The 4-chamber view demonstrated prominent echo density in tricuspid valve location, expressive of tricuspid valve atresia. Also in this view, a small and hypoplastic RV was detected (Figure 1). The short-axis view of the great arteries determined dextro-malposition of the aorta (right and anterior aorta or D-malposed aorta) (Figure 2). The sagittal outlet view illustrated a small pulmonary diameter, representing pulmonary stenosis (Figure 3). The subcostal or septal view detected a large subpulmonic ventricular septal defect (Figure 4).

Karyotyping was performed on an amniotic fluid sample, but no chromosomal abnormality was detected. We informed the woman that the prognosis of this disease was poor, and she decided to terminate her pregnancy. Pregnancy termination was done at 19 weeks of gestational age.

VanPraagh et al\(^5\) proposed a segmental classification according to the involvement of the 3 major segments of the heart: Group I, DORV with conotruncal anomalies only, involving 1 segment of the heart (great arteries); Group II, DORV associated with malformations of the atrioventricular canal and the ventricles, involving 2 segments (great arteries and ventricles); and Group III, DORV in the setting of heterotaxy syndromes and anomalies of the situs, involving the 3 segments (great arteries, ventricles, and atriia). Our case was classified as Group II DORV. Research has shown a rise in the incidence of chromosomal abnormalities, especially chromosome 22q12 deletion, trisomy 18, and trisomy 13.\(^2\) In our case, the chromosomal karyotype was not abnormal.
The prognosis in patients with DORV depends on the severity of associated lesions. Clinical presentation is influenced by the presence or absence of pulmonary stenosis, the position of the ventricular septal defect in relation to the great arteries, and other associated major defects. Brown et al. concluded that risk factors for DORV included aortic arch obstruction, multiple ventricular septal defects, a large ventricular septal defect, and ventricular outflow obstruction or hypoplasia. The rate of pregnancy termination due to DORV varies between 15.8% and 43.4%. These data suggest that the decision to terminate a pregnancy must be individualized for every patient. Further, more women carrying fetuses with DORV and extracardiac defects might choose to terminate if the diagnosis is made earlier.

Bhatla et al. advocated prototyping in correctly selected cases of multifaceted congenital heart disease by positing that a meticulous anatomical understanding could help surgeons choose potential surgical and cardiac catheterization interventions. They utilized a 3D-printed model to better delineate the anatomy of their patient with DORV.

In conclusion, we presented a rare case of DORV and suggested that this kind of congenital heart disease could be diagnosed by experienced fetal echocardiographers with conventional 2D echocardiography. As the degree and nature of the hemodynamic disturbance will affect the prognosis of any conotruncal anomaly, an accurate prenatal description is important for the counseling of parents, particularly if termination is considered. The detection of serious fetal cardiac abnormalities allows the option of pregnancy termination if they are diagnosed early enough in gestation. The parents in our case opted for termination having been made aware of the poor prognosis. Ideally, fetuses at risk for structural cardiac defects should be scanned between 17 and 24 weeks of gestation. During this period of pregnancy, the heart is large enough for adequate visualization, and there is enough amniotic fluid surrounding the fetus to provide a good ultrasonic window.

References

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