Absent pulmonary valve syndrome (APVS) is a rare conotruncal anomaly consisting of a severely hypoplastic pulmonary valve with annular stenosis, aneurysmal dilatation of main pulmonary artery with dilatation of one or both pulmonary artery branches, and a ventricular septal defect. Here, we report a prenatal echo diagnosis of APVS in a 27-year-old primi gravida at 20 weeks of gestation confirmed on fetal autopsy. A ‘bow tie’-like hypoechoic shadow in fetal cardiac ultrasound observed by us in a modified four-chamber view was suggestive of aneurysmal dilatation of branch pulmonary arteries. The consequences of continuation of pregnancy including immediate neonatal complications and possible medical and multistaged surgical interventions were well explained. Parents opted for medical termination of pregnancy. Autopsy findings of the fetus were consistent with the prenatal echo diagnosis of APVS. The presence of patent ductus arteriosus seen in the autopsy may be the cause of severe heart failure evidenced by the abnormally large congested liver, dilated right heart chambers, and tricuspid valve annulus. We infer that the prenatal diagnosis of APVS may be possible with a high degree of accuracy with characteristic fetal echocardiographic findings such as ‘bow tie’-like or ‘ballooning’-like shadows observed in this case. The presence of ductus confirms definite fetal loss and the parents can be counselled accordingly. However, when the ductus is absent, decision-making is difficult as the fetus is going to survive.

Keywords Absent of pulmonary valve syndrome • Pulmonary artery dilatation • Prenatal cardiac diagnosis • Tetralogy of Fallot

Introduction

Cheevers1 reported a case of absent pulmonary valve syndrome (APVS) in 1847. APVS is a rare conotruncal anomaly in which the pulmonary valve is absent or rudimentary, usually associated with a malalignment type of ventricular septal defect and stenotic pulmonary annulus.2 It is considered as a variant of tetralogy of Fallot (TOF) and ~30% of these children have absent ductus arteriosus.3 The presence of ductus arteriosus in this anomaly will cause severe pulmonary regurgitation leading to congestive heart failure and fetal loss.4 Here, we report a case of the prenatal diagnosis of APVS at 20-week gestation, with autopsy correlation.

Case report

A 27-year-old primi was referred for fetal echocardiogram at 20 weeks of gestation in view of atrial enlargement noted elsewhere on routine ultrasound examination. Fetal echocardiogram obtained by various conventional and modified views showed normal situs, levocardia, normal systemic and pulmonary venous drainage, and large subaortic ventricular septal defect with 50% overriding of aorta (Figure 1A). M-mode measurement values of fetal heart were compared with normal reference ranges for gestational age with corresponding percentile showed mitral valve 4 mm (50th), tricuspid valve 5.2 mm (90th), mitral valve orifice/tricuspid valve orifice ratio 1.3 (>95th), right pulmonary artery 4.8 mm, left pulmonary artery 8.4 mm (>95th), and width of right and left ventricles 8.8 mm (>95th) and 7.9 mm (>80th), respectively. The cardio-thoracic ratio (0.54) was more than 95th percentile, indirectly gives high central venous pressure. Great artery relationship was normal, significant pulmonary artery stenosis and regurgitation was noted by continuous-wave Doppler interrogation across the pulmonary annulus (Figure 1B). A ‘bow tie’-like hypoechoic shadow observed by us at the left side of the heart in a
modified four-chamber view obtained by tilting the probe posteriorly was suggestive of dilated branch pulmonary arteries (Figure 1C). These findings on fetal echocardiogram were consistent with APVS.

Counselling was done, regarding the consequences of continuation of pregnancy including the possibility of intrauterine complications endangering the pregnancy such as congestive heart failure and fetal hydrops. Immediate neonatal complications, such as respiratory distress due to the bronchial compression of the dilated pulmonary artery, and need of multistage surgical procedures were also explained. The parents opted for the medical termination of pregnancy.

Autopsy of the fetus was performed with the consent of the parent. The situs was normal and the size of the fetus and fetal heart measured 240 and 27 mm, respectively. The liver was enlarged, congested, and extending up to the left side of the abdomen (Figure 2A). Intra-cardiac autopsy findings included aneurismal dilatation of pulmonary arteries and its branches with preferential aneurismal dilatation of left pulmonary artery measuring 7 mm in diameter (Figure 2B), left oriented infundibulum, stenotic pulmonary annulus (2 mm) (Figure 2C), large conoventricular septal defect (9 mm) (Figure 2D), and ductus arteriosus (Figure 2E). Enlarged liver in the presence of dilated right heart chambers and tricuspid valve (Figure 2F) indicated severe intrauterine congestive heart failure and impending hydrops.

**Discussion**

Conotruncal anomalies can be diagnosed in prenatal life by fetal echocardiogram with a high degree of accuracy. Difficulties in defining the spatial relationship of great arteries limit the accuracy of diagnosis in some instances. APVS can be suspected when we see any associated echocardiographic findings like aneurysmal dilatation of pulmonary arteries, and its branches appearing as ‘bow tie’- or ‘balloon’-like hypoechoic shadow as we have observed (Figure 1A and C). The presence of both stenotic and regurgitant Doppler gradient at the pulmonary annulus differentiate from uncomplicated TOF (Figure 1B). Studies conducted by Andrew
et al. showed that conotruncal anomalies pose poor prognosis for the fetus and 64% of the foetuses with a prenatal diagnosis of conotruncal anomalies died except in cases of uncomplicated TOF. The mortality is comparable with the Allan et al. study which showed mortality ranged from 50 to 65%. In our autopsy study, we found that the liver was abnormally large and congested with tricuspid valve annular dilatation, signifying intrauterine congestive heart failure, and impending hydrops endangering the fetal life. The absence of ducal flow between aorta and pulmonary artery is essential for the in utero survival of a fetus with APVS.

In fetuses with TOF, APVS, patent ductus arteriosus, and an unrestrictive ventricular septal defect, the regurgitant flow from the aorta through ductus and absent pulmonary valve fills not only the right ventricle but also the left ventricle. The diastolic overload of both ventricles is probably incompatible with further fetal life and results in severe heart failure, fetal hydrops, and fetal loss. A possible additional mechanism causing fetal demise could be a significant diastolic run-off not only from the systemic vascular bed of fetus but also from the critical placental vascular bed, leading ultimately to fetal hypoxaemia. In the present case, the presence of ductus made changes in fetal physiology leading to fetal heart failure (Figure 2A, E, and F).

In a study by Pinsky et al., 40% of children with APVS had developed respiratory distress during early infancy secondary to aneurismal dilatation of pulmonary arteries compressing bronchi resulting in massive lobar emphysema. The most critical period for patients with APVS was in infancy. Autopsy done at neonatal death showed that ductus was almost always absent.

Normally infundibulum is fairly short and vertical, whereas in APVS, the orientation of infundibulum can be towards right and horizontal with corresponding preferential aneurismal dilatation of right pulmonary artery, compressing the origin of the middle lobe bronchus, or left oriented with corresponding preferential aneurismal dilatation of left pulmonary artery, compressing left main bronchus and left upper lobe bronchus. In our case, infundibulum was left oriented.

The severity of the heart disease, pulmonary complications, long-term medical management, complications of staged surgical interventions, and outcomes need to be explained to the parents. The possibility of developing intrauterine heart failure and hydrops in APVS should also be explained to the parents. Deletions in chromosome 22q1.2 in most of the patients with APVS were studied by Johnson et al. This supports a specific genetic and embryologic mechanism involving the interaction of the neural crest and primitive aortic arches as a possible cause of APVS.

The prenatal diagnosis of APVS may be possible with a high degree of accuracy with characteristic fetal echocardiographic findings such as ‘bow tie-like’ or ‘ballooning-like’ shadows observed in this case. The presence of ductus confirms definite fetal loss and the parents can be counselled accordingly. However when the ductus is absent, decision making is difficult as the fetus is going to survive. Further studies on APVS may be required with respect to natural courses, survival of fetus with or without PDA, and outcomes of medical and surgical management in APVS for a reasonable prenatal counselling influences the decision of medical termination.

**Conflict of interest:** None declared

**References**