Prenatal Diagnosis of Aorta-Porto-Umbilical Vein Fistulas with Left-Sided Inferior Vena Cava

Jiawei Zhou, PhD1 Qichang Zhou, PhD1 Qianghai Peng, PhD1 Shi Zeng, PhD1

1 Department of Ultrasonography, The Second Xiangya Hospital, Central South University, Changsha, Hunan, People’s Republic of China

Address for correspondence Shi Zeng, PhD, Department of Ultrasound, The Second Xiangya Hospital, Central South University, 139 Renmin Road (M), Changsha 410011, Hunan, People’s Republic of China (e-mail: doctorzshi@163.com).

Case Study

A 32-year-old, gravida 2 para 1 woman was referred to our center for fetal echocardiography at 30\(\frac{1}{4}\) weeks of gestation following routine ultrasound screening because of cardiomegaly. Family history was negative for congenital heart malformations, and there was no history of medication use during pregnancy. The cardiothoracic ratio was 0.49, and there was no sign of hydrops. Echocardiography showed normal segmental anatomy and a perimembranous ventricular septal defect. On detailed examination, it was noted that the dilated intra-abdominal part of umbilical vein coursed into the liver like a helix. Doppler imaging demonstrated a 28 × 24 mm vascular structure in the middle part of the liver and revealed a “saw-toothed” arterial flow pattern with an extremely high-peak velocity (240 cm/s) in the lesion (►Fig. 1). Subtle examination of the lesion on sagittal and transverse views using color and power Doppler revealed that the following vessels were in continuity with the lesion: an arterial vessel originating directly from the abdominal aorta, portal veins, and a dilated umbilical vein (►Fig. 2). Moreover, the subrenal inferior vena cava (IVC) ran upward along the left side of the aorta and then anteriorly crossed the aorta at the level of the kidney to join the normal IVC on the right (►Fig. 3). All three hepatic arteries and ductus venosus were visible and coursed normally. Cordocentesis yielded a normal karyotype (46, XY). The pregnancy was uneventful and showed no sign of hydrops, and spontaneous vaginal delivery occurred at 37 weeks' of gestation. An extended right hepatectomy was successfully performed at the age of 3 months. There were no complications, and sonography showed no obvious residual lesions upon follow-up after 3 months.

Discussion

Congenital arteriovenous shunts in the liver of fetuses are very rare, especially when involved with the abdominal aorta.1 Portal hypertension and/or congestive heart failure are the main presentations in postnatal life. Although the progress of the arteriovenous malformation in the fetus depends on the size of the lesion, the related arteries and the associated anomalies, most published isolated cases showed good outcomes including spontaneous closure2 and successful surgery.3

This anomaly has mostly presented alone, with a few cases been associated with other congenital defects, such as duodenal atresia and trisomy 21.1 To the best of our knowledge, the combination of defects described in this case, namely, aorta-porto-umbilical vein fistulas with a left-sided IVC, has not been previously reported. Left-sided IVC is a rare form of IVC malformation. A search of the

Abstract

Keywords

► fetal

► fistulas

► left-sided inferior vena cava

Congenital arteriovenous fistulas involved with the abdominal aorta are very rare. Left-sided subrenal inferior vena cava (IVC) with normal connection to the heart is also rare and has not been reported prenatally. In this article, we described a fetus with aorta-porto-umbilical vein fistulas combined with a left-sided IVC.
Fig. 1  Doppler revealed “saw-toothed” arterial flow pattern with extremely high-peak velocity (240 cm/s) in the lesion.

Fig. 2  Doppler imaging demonstrated a vascular structure in the middle part of the liver. The following vessels were in continuity with the lesion: an arterial vessel originating directly from the abdominal aorta, portal veins, and dilated umbilical vein. Ao, abdominal aorta; Sp, spine; St, stomach; UV, umbilical vein.
Prenatal ultrasonography can facilitate the prenatal detection, provide close observation, and evaluate the complications and hemodynamic status of such lesions.

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References

Fig. 3 The arrows demonstrate the course of IVC: the subrenal IVC ran upward along the left side of aorta, and then crossed the aorta anteriorly at the level of the kidney to join the normal IVC on the right. Ao, abdominal aorta; IVC, inferior vena cava; St, stomach.