ABSTRACT

Fetal liver tumors represent approximately 5% of all congenital tumors and hemangiomas account for approximately 15% of all fetal liver tumors. Hemangioma and hemangioendothelioma are benign vascular tumors but they are associated with a significant morbidity and mortality due to high output congestive heart failure and arteriovenous shunting. Ultrasound features of hepatic hemangioma can overlap with arteriovenous malformation. This is a case report of fetal hepatic hemangioma with arteriovenous shunting detected on routine anomaly scan at 20 weeks of gestation in 22 years old Nepalese woman who was asymptomatic. Prenatally, a 3.2 cm x 2.8 cm multicystic mass was seen in the left hypochondrium of the fetal abdomen displacing the stomach. On color Doppler examination, the mass showed internal color flow and multiple tortuous dilated vessels noted around the lesion. After counseling regarding the lesion to the mother and her family; decision to abort the fetus was made. Autopsy was done for the final diagnosis of the lesion, which showed vascular mass in the left lobe of liver insinuating between stomach and spleen.

INTRODUCTION

Hepatic tumors are rare in a perinatal period. They represent about 5% of all neoplasms. Various types of hepatic tumors include hemangioma, hemangioendothelioma, mesenchymal hamartoma and hepatoblastoma. Nearly, 15% of all fetal tumors are hemangiomas, which usually appear on the face, neck and limbs. Hemangioma and hemangioendothelioma are benign vascular tumors but they are associated with a significant morbidity and mortality due to high output congestive heart failure and arteriovenous shunting. They are characterized as either rapidly involuting or non-involving based on their clinical progression. Fetal hepatic hemangiomas are rarely reported; hence, their exact incidence rate remains undetermined. Congenital hepatic hemangiomas, may reach maximum size at birth and, in some cases, gradually disappear. Small hepatic hemangiomas are often asymptomatic and seldom require treatment whereas, huge hepatic hemangiomas (Diameter >
cm) are less frequently reported and can cause serious complications.⁴ Neonatal mortality rates can reach 30–100% if untreated.¹ Hence, studies on the prenatal diagnosis of hepatic hemangiomas are necessary for thoroughly monitoring fetuses for tumor progression and related complications and help obstetricians decide on timely delivery and appropriate neonatal treatments. This is case report of a 22-year-old Primigravida who presented with fetal congenital hepatic hemangioma on routine anomaly scan with arteriovenous shunting.

CASE REPORT

A 22 years-old, healthy, Primigravida Nepalese woman presented for an obstetric anomaly ultrasound scan at 20 weeks of gestational age (GA) to our radiology department. Until then she had only one scan at 8 weeks of pregnancy. A 3.2cm x 2.8cm multicystic mass was seen in the left hypochondrium of the fetal abdomen displacing the stomach (Figure 1).

![Figure 1. Lobulated multicystic mass of approx. 3.2 x 2.8 cm size (arrow) in left hypochondrium of fetal abdomen displacing the stomach.](image)

On color Doppler examination, the mass showed internal color flow and multiple tortuous dilated vessels noted around the lesion (Figure 2). Most of the vessels show venous spectral Doppler waveform and few peripheral vessels shows arterial pattern. The initial diagnosis was made of vascular malformation of abdomen. Except this lesion, there were no other structural abnormalities noted in the fetus. Fetal cardiac size and chambers were normal. There was no history of any medical illness noted in the mother. After counseling regarding the lesion to the mother and her family; decision to abort the fetus was made. Taking the medicolegal written consent from patient, the fetus was aborted at 21 weeks of GA. Autopsy was done for the final diagnosis of the lesion, which showed vascular mass in the left lobe of liver insinuating between stomach and spleen (Figure 3).

![Figure 3. Bluish vascular mass approx 3 x 2.5 cm in left lobe of liver (arrow).](image)

No other morphological abnormalities noted in the fetus. The histopathology report showed the cavernous hemangioma of the liver (Figure 4).

![Figure 4. HE staining showing dilated vascular channels lined by a single layer of endothelial cells.](image)
DISCUSSION

The imaging findings of fetal hepatic tumors may widely overlap, the early detection and prenatal follow up of these tumors are very important for fetal, maternal, and postnatal care. In our case, such tumor presented as multicystic highly vascular mass in the left hypochondrium, diagnosed at routine congenital anomaly scan. The two categories of congenital vascular anomalies of the liver are proliferating tumors, such as hemangiomas, and developmental vascular disorders, such as arteriovenous malformations (AVMs). As proposed by Mulliken and Glowacki and according to the International Society for the Study of Vascular Anomalies (ISSVA), hemangiomas that may affect the liver in children can be classified as: infantile hemangiomas and congenital hemangiomas. Congenital hemangiomas may be Rapidly Involuting congenital hemangioma (RICH) or Non involuting congenital hemangioma (NICH). While small hemangiomas can be asymptomatic lesions incidentally discovered during imaging of the abdomen, larger tumors may act as arteriovenous shunts resulting in high-output congestive heart failure and hepatic dysfunction that may lead to fetal non-immune hydrops. The reported neonatal mortality rate of fetuses diagnosed with liver lesions is 30–100%. This prognosis lead to termination of pregnancy as a possible treatment option prenatally, which was seen in our case too. Since biopsy may result in massive haemorrhage, diagnosis is most often based on characteristic radiological findings of usually well-defined mixed, solid lesions with hypervascularization and fine granular calcifications. Although hemangiomas have been reported hyperechoic or isoechoic, our case showed a multi-septated complex cystic structure with low resistance flow. Associated, peripheral dilated vessels were also noted with low resistance flow. Sonographically, AVMs characteristically appear as echopenic dilated vascular channels replacing liver parenchyma with high-flow Doppler characteristics that lack arterial pulsation. Similar appearance was detected in our case showing anechoic area with peripheral tortuous vessels which is an indication of arteriovenous shunting. Hepatic hemangiomas can occur in either or both lobes of the liver but incidence is more in right lobe than left lobe. The lesions can be solitary or multifocal and vary in size. In case report, the fetus had a left-lobe lesion. When the lesion is in left lobe, the lesion can displace the adjacent organ i.e stomach and spleen such as in our case. Cases of fetal multiple hepatic hemangiomas have been reported rarely.

Fatal complications of huge fetal hepatic hemangiomas include congestive heart failure, consumptive coagulation dysfunction (Kasabach-Merritt syndrome), and liver rupture. Hence, evaluation of intrauterine conditions via serial ultrasound examinations is important in making pregnancy-related decisions, including active management with in-utero treatment, expectant management, or expediting delivery. In resource poor areas like ours, termination of pregnancy is one of the options for the patient, and so it was in our case.

CONCLUSION

Congenital hepatic hemangioma in fetus is not uncommon, but detection of the lesion is crucial in management, as there is overlap in sonographic findings particularly with AVM. And associated, life threatening conditions of hemangioma resulting in increased mortality should always to be considered. Detailed anomaly scan plays an important role to detect any vascular anomaly in fetus. Serial imaging should be performed to evaluate the characteristics of lesion and to avoid any related complications.

REFERENCES


