Chorioangioma of Placenta

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Abstract

We report a case of Chorioangioma of placenta resulting to hydropsfetalis. A 24 yrs old primigravida with polyhydramnios and large placental chorioangioma at 24 wks of gestation expelled a nonimmune hydropic male baby weighing 900gms. The baby expired after 5 minutes of expulsion.

Keywords: Placental chorioangioma, Hydropsfetalis, Polyhydramnios

Introduction

Placental chorioangioma is the most common benign tumor of placenta. The incidence being 1% of pregnancies. The relationship of vascularised chorioangiomas to adverse pregnancy outcome is well recognized. 50% of all cases will lead to maternal and fetal complications. Ultrasonography and Doppler ultrasonography are useful in establishing the prenatal diagnosis and the prognosis.

Case Report

A 24 yrs old primi- gravida, resident of Kathmandu was admitted in the female surgical ward on 11 chaitra 062 for observation with the diagnosis of 27 +3 weeks of gestation with chorioangioma by USG. She had 2 ante-natal visit, 1st at 18 wks, 2nd at 26+5 weeks of gestation at TUTH.

She attained menarche at 14 yrs. and had regular cycle with average flow without dysmenorrhoea. Her LMP was on18.06.062 and EDD calculated to be on 25.03.063.

No family history suggestive of chorioangioma.

She was non smoker, non alcoholic. In her last ante-natal visit, at 26 weeks of gestation on examination perabdominally she was found to have uterus of 30 weeks size, it was tense in consistency with foetal parts not palpable and FHS was present. Since uterine height was more than the period of gestation urgent USG was advised keeping in mind she might have polyhydramnios, abruptio placentae, multiple pregnancy. USG revealed single live foetus of 24 week gestation with adequate liquor. And a separate placental mass measuring 8.8 x 9 cm size was also noted, giving the impression of chorioangioma of placenta. So she was advised admission for observation. Next day she went into premature labour and expelled an alive male baby weighing 900 gms with Apgar score 1/10, 0/10, 0/10.

Baby was cyanosed, had generalized oedema, and ascitis. There was no gross congenital anomaly.

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On microscopic examination section from umbilical cord showed 2 arteries & 1 vein. Placental section showed multiple chorionic villi lined by trophoblasts with increased vascularity in the mesenchymal cone. The separate tumour mass composed mainly of small blood capillaries along with foetal myxoid area & fibrous tissue.

Her intranatal and post natal period was uneventful and she was discharged on 3rd post natal day.

Discussion

Chorioangioma is the most frequent non trophoblastic tumour of the placenta with an incidence ranging from 0.01 - 1.3% of pregnancies. This term is used to describe abnormal proliferation of vessels arising from chorionic tissue (1). It is the most common benign tumour of the placenta (2). Small placental chorioangiomas are found in 1% of pregnancies and are often of no clinical significance. Large chorioangiomas larger than 5 cm are much less common with an estimated incidence of 1 in 2000-3500 births, but are associated with high perinatal mortality of 40% (3). It is usually single but occasionally multiple. This case also had two placental chorioangiomas with size measuring more than 5 cms. Hpoxia and genetic factors are predisposing factors (4). The majority are small not visible on external surface are asymptomatic. It is variable in shapes and divided by fibrous septa. It most commonly protrudes from the foetal surface of the placenta near the insertion of cord. The size more than 5 cm is associated with complications (5).

There are multiple risk factors for chorioangioma. It is divided into moderate and high risks. Moderate risk factors include age of the patient >35 years, increased BP during second trimester, new paternity and dietetic factor: famine, anorexia. Environmental factors like high altitude, smoking, controlled diabetes mellitus, genetic factor.

BMI more than 25, primiparous, past family history in first degree relatives, pre-eclampsia due to premature placental release, cocaine, tobacco or caffeine use, molar pregnancy, uncontrolled diabetes mellitus, scleroderma, chronic hypertension, anti-phospholipid syndrome, past history of deep vein thrombosis, thrombophilia, past history of placental vascular pathology, IUD, abruptio placentae, IUGR, repeated foetal loss, hyperhomocystinaemia are the high risk factors for choroangioma of placenta (6). In this case clinically and by history she was not suggestive of any of the risk factors mentioned above except she was primigravida.

Maternal complications includes

Polyhydramnios, Precipitate labour, Pre-eclampsia, maternal thrombocytopenia, maternal coagulopathy, Hypertonic uterus, PROM, Haemorrhage- both APH & PPH (6). This case had polyhydramnios, pre term premature rupture of membrane and precipitate labour.

Foetal Complication includes: Abortion due to hypoxia, IUGR, and IUD because the considerable proportion foetal blood passes through the tumour rather than the functional placental tissue. So foetus is supplied by deoxygenated blood and nutrient poor state.

There can be foetal cardiomegaly due to peripheral AV shunts leading to increased foetal cardiac output. The cause of neonatal oedema can be cardia failure or hypoalbuminaemia – because of trasudation of protein from the surface vessels of the tumour or from chorionic foetomaternal bleeding from haemangioma.

Neonatal thrombocytopenia is due to injury within the tumour vessels.

DIC may occur due to release of thromboplastic substance from the haemangioma. Neonatal anaemia is due to sequestration of foetal erythrocytes within the tumour, or massive foetomaternal bleed from the haemangioma or microangiopathic haemolytic anaemia induced by injury inflicted on foetal RBC (5).

Ultrasonogram, maternal serum AFP - increased prenatally , Colour doppler imaging- to differentiate placental chorioangioma from other placental lesions. colour doppler shows no blood flow within the mass, but clear foetal waveforms in its periphery can be demonstrated (7). 3D power doppler angiography-clearly shows highly vascularized placental mass including its feeding vessels& drainage at the foetal surface of the tumour (5).

This case was diagnosed ultrasonographically which was confirmed by histopathological examination. The most effective way of diagnosing choroangioma
is doppler ultrasonography but we could not apply it in this case, it was planned but same night she went into preterm labour.

Various modalities of treatment of chorioangioma of placenta have been reported in different case reports.

1. Treating the cause is by occluding the vascular supply like
   a. Foetoscopic devascularization by ablating the feeding vessels (8).
   b. Microcoil embolization by ultrasound guided laser therapy which leads to tissue coagulation causing complete cessation of tumor blood flow (9).
   c. Intravascular injection of absolute alcohol in the feeding vessels of large tumour - this may be one of the best choices due to its high efficacy, simplicity, safety and very low cost (10).
   d. Ligation of tumor vessels.

2. Treating the foetal complications produced by chorioangioma prenatally by intrauterine transfusion. It is necessary if foetal anaemia is diagnosed by cordocentesis in early pregnancy. There is risk of repeated transfusion, pre-term labour and iron overload, so some form of embolization or tumour devascularization is necessary(8).

3. Symptomatic treatment like amniocentesis: when amniotic fluid index is 40 cm, deepest pocket 12 cm and excessive maternal symptoms.

Prognosis is poor for large tumor. It largely depends on foetal haemodynamic tolerance with large placental chorioangioma. 50% lead to maternal & foetal complication (11).

References


