Successful Outcome of a Pregnancy Complicated by a Voluminous Placental Chorangioma: Case report with Review of the Literature

Dilani Lokuhetty and Maiya Al Bahari

Abstract

Placental chorangioma is a rare, benign, placental neoplasm associated with a myriad foetomaternal and neonatal complications. Voluminous chorangiomas have frequent complications and adverse outcomes. The successful outcome of a pregnancy complicated by a voluminous chorangioma is reported herein, with the literature review. Chorangiomas mostly occur in primiparous women over 30 years of age, complicated by polyhydramnios, placenta praevia, ante-partum haemorrhage, premature separation of membranes, premature labour, toxemia of pregnancy, hydrops foetalis and midterm intrauterine death. The neonatal period may be complicated by effects of low birth weight, cardiomegaly, cardiac failure, anaemia, thrombocytopenia and unsuspected congenital anomalies. This case had an unusually uneventful antenatal period for 36 weeks, culminating in a medically unassisted, spontaneous, vaginal, breech delivery with a successful pregnancy outcome, in spite of having a voluminous chorangioma.

Keywords: Voluminous chorangioma, Placenta, Pregnancy outcome.

Introduction

Placental chorangioma described by Clarke in 1789 is usually <5mm in size, occurring in 0.5-1% of pregnancies. Voluminous chorangiomas (VC) >5cm are rare, with prevalence ranging from 1:3500 to 1:9000 births. Chorangioma is a benign vascular neoplasm arising from the placental chorionic mesenchyme. Some consider it a hamartoma arising from angioblastic tissue of the primitive chorionic mesenchyme. Most chorangiomas are incidental, microscopic, more frequent in primiparous women over 30 years and in twin pregnancies. Pregnancy induced hypertension and diabetic complications occur seven and three times more frequently in those affected respectively. VC’s are often associated with polyhydramnios, ante-partum hemorrhage and premature labor with severe pregnancy complications. Amniodrainage facilitates continuation of pregnancy in polyhydramnios. VC’s are also more likely to cause breech presentation, placenta praevia with ante-partum hemorrhage and premature membrane separation. Intrauterine midterm fetal demise is a major complication, as well as non-immune hydrops foetalis. Overall mortality rate of VC’s is around 30%.

Successful pregnancy outcomes complicated by VC is relatively under documented. We report herein, a case of VC with an uneventful antenatal period culminating in a successful outcome at 36 weeks of gestation, with review of the literature. To our knowledge this is one of the largest chorangiomas hitherto documented.

Case Report

A 31 year old gravida 6, para 5 woman from a remote area in the Sultanate of Oman with no significant present/past obstetric history, was diagnosed with a large (>12cm) placental chorangioma by ultrasonography, at 33 weeks of gestation. Imaging confirmed a breech presentation, intrauterine growth retardation (IUGR) and polyhydramnios. She was under regular surveillance at the Sultan Qaboos University hospital (SQUH) and was booked for an elective LSCS. However spontaneous labor commenced at home at 36 weeks of gestation, a few days before the elective LSCS. She had a male baby by vaginal, breech delivery with an intact perineum en route to the SQUH. Her delivery was carried out by family members without medical assistance. The baby weighing 2.15kg was grunting, had tachypnoea, and cyanosis, with sub costal recession and reduced air entry on arrival at hospital. Respiration improved after immediate clearance of the airway and administration of oxygen in the hospital. The baby was on antibiotics for five days. Subsequent neonatal examination and investigations were normal with no evidence of congenital lesions, congestive cardiac failure, anaemia, thrombocytopenia or cardiomegaly. The mother and baby were both well by the fifth day of hospital stay and was discharged. The placenta weighed 1297.6g, with placental disc measuring...
19×18×3cm, umbilical cord measuring 55cm in length and fetal membranes. The placenta had a smooth, brownish bosselated mass measuring 13×12×6cm centrally close to the cord insertion on the fetal surface. (Figure 1) The mass comprised solid nodular brownish areas and blood filled spongy areas separated by bands of fibrous tissue. (Figure 2) Histology confirmed a chorangioma with areas of ischaemic necrosis. (Figure 3)

**Figure 1:** Placental disc with the bosselated brownish mass on the fetal surface.

**Figure 2:** Cut surface of the bosselated brownish mass.

**Figure 3:** Histological appearance of the chorangioma (hematoxyline and eosine stain 20×)

### Discussion

The singleton pregnancy of this multiparous woman of 31 years, was complicated by a VC, measuring 13×12×6cm. Her antenatal period was uneventful until 33 weeks of gestation, when the chorangioma was diagnosed with polyhydramnios and IUGR. The following are attributed to the polyhydramnios:

1. fluid transudation from the large vascular area of the VC
2. transudation caused by obstruction to blood flow by VC’s located close to cord insertion.
3. placental functional insufficiency caused by shunting of blood into the chorangial vascular bed.

Compromised placental function with altered feto-maternal circulation possibly causes growth retardation and even foetal demise. VC’s possibly act as space occupying lesions within the uterine cavity causing breech presentation and placenta praevia. Any one single case however may not have all these complications.

Pregnancy progressed to 36 weeks of gestation in this woman with a VC, polyhydramnios, IUGR and breech presentation. She went into premature labor and had an uneventful vaginal delivery. The location is likely to be critical in determining the outcome of chorangiomas, as those as small as 2cm located at the outer placental edge has been associated with complications. Though this chorangioma was massive, its central location would have allowed the pregnancy to continue up to the 36 weeks of gestation, with a relatively uneventful antenatal period. She also had a medically unassisted, spontaneous, vaginal delivery with a successful outcome, possibly facilitated by her multiparous status.

While several chorangiomas have measured up to 12cm, only a few have measured 13cm or more. Table 1 compares the clinical/pathological features of these VC’s. VC that measured >13cm in the older literature have progressed undetected till delivery, with fatal outcomes. More recent VC’s measuring 13cm or more have resulted in live births by cesarean sections, with significant foetal/neonatal complications. This case is unique in comparison, as only low birth weight was encountered as a complication of prematurity. Fetal/neonatal complications of VC’s such as hydrops fetalis, cardiomegaly, cardiac failure, hemolytic anemia and thrombocytopenia were absent. Unsuspected/suspected congenital anomalies have also been described in infants with pregnancies complicated by chorangiomas. This neonate lacked any congenital abnormality.

Chorangiomas comprise benign proliferating networks of congested capillary blood vessels. Unusual histological features documented include myxoid change and a haemangiopericytomatous pattern. Adjacent villi have shown arrested maturation. Rare proliferating malignant trophoblast have been detected among proliferating capillaries with high mitotic activity and large areas of necrosis in chorangiocarcinomas. Hitherto only few chorangiocarcinomas are documented, with its’ very existence challenged in the absence of metastasis. This chorangioma did not have unusual/malignant features.
Three infants with VC’s and polyhydramnios also have had neonatal haemangiomatosis. All three had undergone amniodrainage with possible embolization of precursor endothelial cells from placental vessels causing haemangiomatosis.

Though the focus has been on chorangioma size, vascularity of the chorangioma is suggested to be more important in predicting its outcome, in view of its blood flow from the foetal circulation.

Conclusion

VC’s may not adversely impact pregnancy outcome as demonstrated by this case. However considering the high incidence of complications, those identified with chorangiomas by ultrasound scanning during early pregnancy; need close monitoring and management of possible foeto-maternal complications. Delivery in a hospital setting and postnatal investigation of the neonate for congenital anomalies and, or the complications described above are recommended.

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Table 1: Features of chorangiomas measuring ≥ 13 cm

<table>
<thead>
<tr>
<th>Studies</th>
<th>Size</th>
<th>Placental location</th>
<th>Diagnostic method</th>
<th>Delivery and outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manitoba, Canada 12</td>
<td>14 × 11 × 8 cm</td>
<td>Separate mass attached by vascular pedicle to the placenta.</td>
<td>Undetected till delivery.</td>
<td>Fatal outcome. Macerated stillbirth at 39 weeks of gestation.</td>
</tr>
<tr>
<td>London, United Kingdom 13</td>
<td>16.6 × 10 × 6 cm</td>
<td>On maternal aspect of placenta.</td>
<td>Undetected till delivery.</td>
<td>Fatal outcome. Delivered live at 32 weeks of gestation with death after 69 hours.</td>
</tr>
<tr>
<td>Bornova, Turkey 14</td>
<td>13 × 10 cm</td>
<td>Not mentioned.</td>
<td>Ultrasound scan and colour Doppler flow imaging at 32 weeks of gestation.</td>
<td>Fatal outcome. Delivered live at 33 weeks of gestation with death on the 3rd day.</td>
</tr>
<tr>
<td>London, United Kingdom 15</td>
<td>14.3 cm in diameter</td>
<td>Not mentioned.</td>
<td>Ultrasound scan and colour Doppler flow imaging.</td>
<td>Live birth. Delivered by elective caesarean section at 31 weeks and 6 days of gestation, following cordocentesis.</td>
</tr>
<tr>
<td>Kaohsiung, Taiwan 16</td>
<td>15.5 cm in diameter</td>
<td>Medio lateral to cord insertion, embedded within placenta.</td>
<td>Ultrasound scan at 29 weeks of gestation.</td>
<td>Live birth. Delivered by emergency cesarean section at 35 weeks of gestation.</td>
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References


