Chorangiomia of Placenta with High Risk Pregnancy: A Case Series

Uma S Andola, Shabnam Karangadan¹, Sainath K Andola¹, Rajashekhar Jewargikar¹
Department of Obstetrics and Gynecology, ¹Resident, Professor and Head of Department, Professor, Department of Pathology, Mahadevappa Rampure Medical College, Gulbarga, Karnataka, India

ABSTRACT

Chorangiomia is the most common benign tumor of placenta, with an incidence of approximately 1%. Tumors larger than 5 cm are associated with maternal and fetal complications. We report four cases of chorangioma that were identified in mostly primigravida (75%) with a mean gestational age of 33 wks (Range 28-33 wks). All except one case, complete placenta was received with mean weight of 550 g (Range 400-800 g) and tumor size of 9.4 cm (Range 0.5-23 cm). Microscopy showed capillary proliferation resembling fetal vessels in enlarged villi covered by trophoblastic epithelium. Three cases had preterm delivery. One also presented with abruptio placenta and intrauterine fetal death. Because of limited documentation of cases and dismal prognosis of these pregnancies, especially with large chorangiomas, it’s of utmost importance to correctly diagnose them with the aid of color Doppler flow. In this series, 75% of cases (3/4), babies survived probably due to small size of tumor and attainment of fetal maturity.

KEY WORDS: Chorangiomia, placenta, placental tumor, vascular tumor

KEY MESSAGES: Even though chorangioma is the most common tumor of placenta, tumors arising from placenta are rare. Because of dismal prognosis of these pregnancies with large lesions, it’s important to correctly diagnose them with the aid of color Doppler flow. This case series demonstrates the need for awareness of such lesions.

INTRODUCTION

Chorangiomia of placenta is the most common benign tumor of placenta, with an incidence of approximately 1% of all pregnancies.[1-3] It is a vascular tumor which is usually small in size and only found after careful examination of the placenta. Tumors larger than 5 cm are associated with maternal and fetal complications.[1,2,4]

Limited documentation of cases and dismal prognosis of these pregnancies, especially with large chorangiomas, has prompted us to present this case series.

CASE REPORT

Four cases of placental chorangioma were diagnosed in the Department of Pathology, M. R. Medical College, Gulbarga, between 2011 and 2013. The clinical and pathological findings of the cases are summarized in Table 1.

The patient’s age ranged from 23 to 28 yrs (mean 24 yrs). Most of them were primigravida (3/4). Gestational age at diagnosis ranged from 28 to 39 wks (Mean 33 wks). Clinical presentation was in active stage of labor, bleeding PV and with retained bits of placenta post delivery. Size of the tumor ranged from 0.5 to 23 cm (mean 9.4 cm). All except Case 3, complete placenta was received [Figure 1a]. Placental weight of the three cases ranged from 400 to 800 g (mean 550 g). Ultrasound scan report was available in Case 1 which was reported as 10 × 10 cm placental hematoma [Figure 1b].

All cases showed capillary proliferation resembling fetal vessels in enlarged villi covered by trophoblastic epithelium [Figures 1c and 1d]. Lymphoplasmacytic infiltrate along with scattered Hofbauer cells were noted. Areas of myxoid change and calcification were also seen. In between these hemangiomatous areas are seen normal and degenerated villi.

All four cases delivered by spontaneous normal vaginal delivery, of which one was still born. Cases 1 and 4’s babies were having low birth weight as well (0.8 and 1.8 kg, respectively).
placemomgaly, growth restriction, abruptio placenta, perinatal death and maternal pre-eclampsia. Large tumors probably act as arteriovenous shunts and cause complications. In this series, two cases that were more than 5 cm, had preterm delivery and Case 1 also presented with pre-eclampsia, abruptio placenta and intrauterine fetal death. Case 4 was associated with maternal rheumatic heart disease and polyhydramnios. IUGR and low birth weight was seen in Cases 1 and 4.

Grossly, these are usually nodular, fleshy lesions connected to the chorionic plate and often bulge from the fetal surface of placenta. Cut surface is variable, from dark red, soft appearance similar to a blood clot to a firm white lesion similar to an infarct.[9] In the present series, size ranged from 0.5 to 23 cm and weight ranged from 400 to 800 grams. Several giant chorangiomas have been mentioned in the literature. Largest has been reported by Arodi et al. weighing 1500 g and 30 × 20 × 5 centimeters.[6] In India, the largest chorangioma weighed 2240 grams.[7] In the present series, the largest (Case 1) weighed 800 g and measured 23 × 10 × 6 centimeters. It was associated with pre-eclampsia, abruptio placenta, preterm delivery, low birth weight and intrauterine fetal death.

Microscopically, typical chorangioma is composed of proliferation of fetal blood vessels, usually supported by scant connective tissue and covered by trophoblast. Villus expansion is caused by proliferation of blood vessels, which is diagnostic. Capillary, cavernous, endotheliomatous, fibrosing, and fibromatous tumors have been differentiated, but such precision is unwarranted as the clinical outcome depends more on the size of the tumor than on its composition.[6] All the four cases in the present series were of capillary type. Occasionally, chorangiomas are associated with infarction and degenerative changes, such as hyalinization, necrosis, myxoid stromal changes or calcifications. Myxoid change and calcifications were noted in Case 1 which was also observed by Bagga et al.[7]

Clinical manifestations are rare and usually associated with tumors greater than 5 cm. Large chorangiomas are rare but clinically significant because of their well-known association with maternal and fetal complications such as polyhydramnios, hydrops fetalis, fetal heart failure, preterm delivery, fetal anemia and thrombocytopenia.

Prenatal diagnosis is ideally by ultrasound with color Doppler by demonstration of a well-circumscribed placental mass with similar or higher echogenicity to the surrounding tissue, and a clearly delineated vascular supply. Done by Asokan et al. in 1978.[8] Bromley and Benacerraf

Table 1: Clinical and pathological findings of placental chorangiomas

<table>
<thead>
<tr>
<th>Age (yrs)</th>
<th>Gravida</th>
<th>Gestational Age (wks)</th>
<th>Clinical presentation</th>
<th>Birth weight (kg)</th>
<th>Tumor size (cm)</th>
<th>Placental weight (grams)</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>28</td>
<td>1</td>
<td>28</td>
<td>Bleeding PV due to abruptio H/o pre-eclampsia</td>
<td>0.8</td>
<td>23</td>
<td>800</td>
<td>Intrauterine foetal death</td>
</tr>
<tr>
<td>20</td>
<td>1</td>
<td>32</td>
<td>Active labor</td>
<td>2.8</td>
<td>10</td>
<td>450</td>
<td>Alive and healthy</td>
</tr>
<tr>
<td>23</td>
<td>2</td>
<td>39</td>
<td>Bleeding PV due to retained placenta</td>
<td>3.1</td>
<td>0.5</td>
<td>—</td>
<td>Alive and healthy</td>
</tr>
<tr>
<td>26</td>
<td>1</td>
<td>34</td>
<td>Active labor H/o Maternal RHD and Polyhydramnios</td>
<td>1.8</td>
<td>4</td>
<td>400</td>
<td>Alive and healthy</td>
</tr>
</tbody>
</table>

DISCUSSION

Chorangiomas are hamartomas of placenta, first described by Clarke in 1798.[3] They are also designated as placental angioma, chorionangioma, angiomyxomas and vascular hamartoma of placenta. The incidence of these tumors is 1 in 9000 to 1 in 50,000 placentas. When careful study and sectioning of placentas is undertaken, prevalence may be as high as 1 in 100. Chorangiomas occur more frequently in Caucasian than in African-American mothers, more often with multiple gestations.[3] The present series describe four cases of placental chorangioma.

Various hypotheses have been proposed for its pathogenesis. Some authors regard them as hamartoma, rather than a true neoplasm. Other factors include environmental factors like high altitude, hypoxic stimulus suggested by elevated nucleated red blood cells in the fetal circulation and vascular growth factors.[3]

Clinical manifestations are rare and usually associated with tumors greater than 5 cm. Large chorangiomas are rare but clinically significant because of their well-known association with maternal and fetal complications such as polyhydramnios, hydrops fetalis, fetal heart failure, preterm delivery, fetal anemia and thrombocytopenia.
stated that the sonographic appearance of a chorioangioma was indistinguishable from that of a placental hemorrhage. They speculated that the use of color Doppler sonography could have helped them in differentiating these lesions.[8] Similar error was made in one of the cases in this series (Case 1), mistaken as placental hematoma.

Chorioangioma with complications before fetal viability requires interventions. Because of the dismal prognosis of these pregnancies, especially with lesions larger than 5 cm, it’s important to correctly diagnose these lesions and to closely follow up these pregnancies, preferably with the aid of color Doppler.[2] Except Case 1, all babies are alive and healthy. The 28 wks gestation fetus was delivered by normal vaginal delivery following severe ante partum hemorrhage due to abruptio placenta. Baby expired due to birth asphyxia. In 3 out of 6 chorangiomas Zalel et al.[2] described pregnancies that ended before completion of 32 wks of gestation.

Chorangiomas can be confused with subamniotic hematoma, placental teratoma, degenerated myoma and placental infarction on gross morphology but microscopy is characteristic.

CONCLUSION

Even though chorangioma is the most common tumor of placenta, tumors arising from placenta are rare. Because of dismal prognosis of these pregnancies, especially with lesions larger than 5 cm, it’s of utmost importance to correctly diagnose them and to closely follow up these pregnancies, preferably with the aid of color Doppler flow. In the present series, 75% of cases (3/4), babies survived probably due to small tumor size and attainment of fetal maturity. The present case series demonstrates the need for awareness of such lesions, and emphasizes the importance of timely antenatal diagnosis and intervention.

REFERENCES


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