**Case Report**

**Breus’ mole/chorangiosis/chorangioma of the placenta: a dilemma with a rare fetal outcome report**

Nitika Sobti¹*, Ankita Chandna², Bhawna Narula³

¹Department of Obstetrics and Gynecology, Max Health Care, Gurugram, Haryana, India
²Department of Obstetrics and Gynecology, Max Super Specialty Hospital, Shalimar Bagh, New Delhi, India
³Department of Pathology, Max Super Specialty Hospital, Shalimar Bagh, New Delhi, India

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*Correspondence:
Dr. Nitika Sobti,
E-mail: drnitikasobti@gmail.com

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**ABSTRACT**

Massive Subchorionic Thrombohematoma (MST) is a rare condition in which there is a massive collection of blood between the placental membranes and uterine wall separating the villous chorionic plate from villous chorion. It is relatively rare and is poorly understood. Many theories have been proposed to explain the etiology of Breus mole; some suggest it is a fetal haemorrhage, while others claim it has a maternal-origin thrombosis of placental vessels. A 30-year-old healthy Indian pregnant woman was presented at Max Hospital, Shalimar Bagh Delhi, India, during her second pregnancy with a complaint of fever. On routine level-2 ultrasonography (USG) done at 18.6 weeks of gestation showed thick placenta. No fetal tumours or any other anomalies were noted on that scan which was followed by a detailed scan which confirmed a solitary mass arising from fetal side 103x64x82 mm S/O chorioangioma. Serial growth and doppler USG were conducted to monitor placental function, tumor characteristics and future anatomy. The subject received steroids to enhance fetal lungs maturation at Week 30, iron/calcium supplements, Ecosprin tablets, and progesterone support. At 32.5 weeks, the subject developed deranged sugars followed by gestational hypertension at 34.1 weeks. Ultrasonography also showed fetal growth restriction with large chorioangioma. The subject underwent a successful elective caesarean section at 34.4 weeks. On placental examination, 10 cm large mass encasing ¾ of the placenta was identified as a large subchorionic hematoma/chorioangioma (800 g). This study concludes that early identification of a large chorioangioma aids in consequent fetal surveillance, management of maternal symptoms, and delivery planning discussions even if the pathological diagnosis turns out to be Breus’ mole with underlying chorangiosis postnatally.

**Keywords:** Breus’ mole, Chorioangioma, Chorangiosis, Perinatal outcomes, Placenta

**INTRODUCTION**

The placenta is the largest organ which is the mirror of maternal and fetal status. It is important for appropriate maternal and fetal development and functions. Therefore, histological examination of the placenta provides sensitive, specific, and scientific information for evaluating the nature and extent of the placental injury. Massive subchorionic thrombohematoma (MST) is a rare condition in which there is a massive collection of blood between the placental membranes and uterine wall separating the villous chorionic plate from villous chorion.¹² It was first termed Breus’ mole and described in 1892. The frequency of MST is reported between 0.03%–0.08%; its etiology and pathogenesis are yet unknown. The outcomes of MST are dependent on its site
and size. If MST is near the cord insertion, it can lead to umbilical venous obstruction, cord compression, and decreased fetal perfusion. Therefore, MST is related to complications including fetal growth restriction (FGR) and intrauterine fetal death resulted due to placental insufficiency.3,4

In some cases, there is a placental change that results in excessive formation of blood vessels within the terminal chorionic villi called chorangiosis.1,2 Presence of chorangiosis is associated with a high mortality rate (39%) and congenital malformations (42%).5 Differential diagnosis of chorangioma includes chorangiosis and chorangiomatosis, though in our case, it was diagnosed as Breus’ mole with underlying chorangiosis. Therefore, it needs very close follow-up and management during the antenatal period for a successful fetal outcome.5

A case of an Indian female is reported with large placental chorioangioma on antenatal diagnosis, the antenatal management, and maternal and fetal outcome and the postnatal diagnosis.

CASE REPORT

A 30-year old healthy Indian pregnant woman with fever was presented at Max hospital, Gurugram, India, during her second pregnancy. She was evaluated by a level 2 ultrasonography (USG) at 18.6 weeks of gestation that showed placentomegaly. There were no fetal tumours or any other anomalies noted on that scan followed by a USG for monitoring growth which showed bulky placenta with a discrete hypoechoic mass S/O large chorioangioma. The ultrasound done at 31.4 weeks confirmed a solitary mass arising from fetal side 103 × 64 × 82 mm S/O chorioangioma (Figure 1).

Figure 1: Ultrasound examination of the uterus at 31.4 weeks of gestation.

Serial growth and doppler USG were conducted to monitor placental function, tumor characteristics and future anatomy. In past, the subject had regular menstrual cycles, and the earlier antenatal period was normal. Two years back, the subject had uneventful caesarean delivery for breech and had a healthy female child.

Figure 2: Ultrasound examination of the uterus after week 34 of gestational age.

The subject was counselled and kept under strict fetal-maternal surveillance. Fetal medicine and genetic consultation were also sought. Weekly USGs were done to see for any increase in the size of the placental tumor and to monitor fetal growth. The subject received steroids to enhance fetal lungs maturation at week 30. She was advised to take iron/calcium supplements, ecosprin tablets, and progesterone support. At week 32.5, the subject developed deranged sugars, and at week 34.1, she developed gestational hypertension which was managed conservatively. Serial USG showed FGR with large chorioangioma (Figure 2). In view of high BP readings with deranged sugars and FGR, the decision to deliver the baby by caesarean section was taken in consultation with neonatologist. The subject underwent a successful elective caesarean section at week 34.4 and delivered a healthy female child of 1.89 kg with an Apgar score of 8, 9, 8. She was shifted to neonatal intensive care unit for observation. After the operation, was BP and blood sugar of the subject were within the normal ranges. Both mother and baby were discharged on day 5.

Gross examination revealed a large nodular cystic lesion (blood clot) measuring 10 × 5 cm covering a large portion of the placental disc on the fetal surface (Figure 3 and Figure 4). Microscopic examination revealed an intervillous hematoma and fibrinous deposits directly beneath the chorionic plate with chorangiosis (increased vessels in terminal villi; (Figure 5). A large amount of blood was stored in the subchorionic space. No hematoma or infarction was observed on the maternal surface of the placenta. Based on these findings, MST with chorangiosis was diagnosed. It was recommended chromosomal analysis of the baby as well which the
subject was not willing to do. The subject had an uneventful postoperative course.

**Figure 3: Large nodular cystic lesion.**

**Figure 4: Cut section of large nodular cystic lesion.**

**Figure 5: Microscopic examination of umbilical cord A) at low magnification B) at medium magnification C) at high magnification.**

**DISCUSSION**

This was a case report study at Max Hospital, Gurugram, India, on a 30-year-old female subject who reported an MST, a rare condition with underlying chorangiosis. Chorangiosis is a rare placental vascular anomaly which is hypoxia-related angiogenesis. Both conditions can be associated with increased FGR and intrauterine fetal death.\(^7\) The diagnosis is often made with histopathological examination of placenta after delivery.

Massive subchorionic thrombohematoma may be reported as homogenous, heterogeneous, or hypoechogenic mass in the chorion. Its appearance is different from the ultrasonic appearance of normal placental tissue, and can occur as placentomegaly.\(^8\) In a study, elastography showed clearly the differentiation of a hematoma from the placenta in relation to a case of possible placenta previa.\(^9\) However, with USG prenatal diagnosis is generally challenging. In our case, the diagnosis of MST with USG was not possible because of severe oligohydramnios. Though, magnetic resonance imaging (MRI) results of MST vary, MRI is reported useful in differentiating MST from rest of the placental diseases, such as chronic abruption oligohydramnios sequence, placental abruption, and placental mesenchymal dysplasia. A high signal at the rim on T1-weighted images represented thrombohematoma while a low signal at the marginal zone on the T1- and T2-weighted images represented it. In addition, steady-state free precession MRI is used to show the arrangement of the placenta, umbilical cord, hematoma, and foetus clearly. In a previous study, of 14 subchorionic hematomas, 9 were confirmed by MRI.\(^9,10\) Overall, these results suggest that MRI is optimal for the diagnosis of MST when USG diagnosis is difficult. The MST etiology remains obscure. In MST, a large blood amount, especially of maternal origin, collects and separates the chorionic plate from the villous chorion. DNA analysis revealed that a large amount of the blood (85%) in the thrombus is of maternal origin.\(^11\) Fetal villous haemorrhage or obstruction with further accumulation of blood and separation of the chorionic plate massively is proposed to cause MST.\(^12\) Further, maternal blood stasis in the subchorionic space causes thrombosis.\(^3,4\) Recently MST is reported following thrombolytic therapy and in patients with thrombophilic conditions.\(^13\) Chorangiosis is a rare condition and found in up to 5% of pregnancies. A large study reported chorangiosis with the incidence of 3% among 1614 deliveries which was associated with placental lesions including foetal artery thrombosis, infarcts, and chronic villitis.\(^14,15\) Villous hypervascularity in which individual terminal villi comprises vessels in excessive numbers has been classified as chorangiosis. There should be >10 terminal villi containing >10 capillaries per villous in 10 medium power fields occurring in multiple areas (minimum 3) of the placenta.\(^5\)

Uteroplacental insufficiency is the basic mechanism of FGR. In severe FGR cases, at least half of the surface of
the placenta could be covered with haemorrhage under the chorionic plate. In such instances, the finding of normal uterine artery doppler waveforms at presentation is a favorable sign (prognostic) of perinatal survival. In addition, in MST, the site of severe haemorrhage is important whether entirely resides within the membranes and faraway from the placental disc or detach implantation of placenta. In our case, the subject had FGR with gestational diabetes, gestational hypertension and anaemia. Grossly there was placentomegaly and large nodular cystic lesion (blood clot) measuring 10×5 cm covering a large portion of the placental disc on the fetal surface with microscopic features of chronic villitis. Massive subchorionic thrombophlebitis is related to poor pregnancy outcomes. Fung et al. reported that only six of ten pregnancies with MST resulted in a live birth, only 2 of which reached full term. Another study compared seven survivors and seven non-survivors with MST and concluded that MST can be diagnosed in the second trimester by USG examination of the placenta. In this case, the subject was diagnosed as having a large chorangioma about 10 cm encompassing ¾ of the placenta with 1 cm in diameter. Though, on histological examination diagnosis of Breus’ mole with underlying chorangioma was made. The fetal outcome was normal, and the infant had weight 1.89 kg at birth. The infant was preterm with a gestational age of 34 weeks at birth. The management of gestational hypertension and FGR and preterm delivery was done for the successful fetal outcome. As shown in our case, Breus’ mole placental chorangiomas/chorangiosis are not always associated with bad outcomes. Study also favours that early diagnosis results in improved outcomes with close follow-up of affected pregnancies and use of therapeutic intervention as indicated in case complications arise. Although the incidence is low nearly 1 in 1200 placenta, detailed screening of placenta by USG preferably incorporating modalities such as colour flow imaging and MRI might allow better perinatal outcomes. Thus, we suggest that Breus’ mole with chorangioma, a marker of chronic hypoxia is associated with fetal and maternal complications like gestational diabetes, hypertension, severe anaemia, FGR, intrauterine death and congenital anomalies. Diagnosed it due to antenatal suspicion of a large chorangioma and followed the subject with a successful subject outcome. This case study emphasizes the importance of comprehensive sonographic evaluation in pregnancy, especially in the presence of a suspected placental lesion.

Henceforth, obstetricians and pathologists should be aware of this entity and should send the placenta for pathological testing in suspected cases and should report this in the pathology report of each patient.

CONCLUSION

In general, Breus’ mole occurs randomly without warning. It is important to share the diagnostic criteria of placental pathology, and radiological (antenatal) and histopathological (after delivery) along with close antenatal management for the successful fetal-maternal outcome. Great care should be taken in evaluating the entirety of the placenta. However, there is a potential for significant progression in size, vascularity and form which can result in late-onset fetal and maternal complications. Early identification of a placental (chorioangioma) aids in consequent fetal surveillance, management of maternal symptoms, and delivery planning discussions even if the pathological diagnosis turns out to be Breus’ mole with chorangiosis postnatal. Till date to the best of our knowledge, the exact incidence of Breus’ mole and chorangioma has not been documented, and this uncommon and underreported entity has diagnostic implications. Its etiopathogenesis has remained a dilemma in the 18th and 19th century. The significance of Breus’ mole does not depend upon the size but on their site of appearance. Therefore, early recognition of placental pathology and timely intervention is of utmost importance for a successful fetal outcome.

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