Case Report

Placental Retention with Accreta in a Uterine Anomaly

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ABSTRACT

A 32 year-old P, G went into spontaneous labour at 37 weeks gestation. Precipitate labour progressed to a normal vaginal delivery (NVD) of a baby boy within 2 hours of onset. She had a retained placenta. The patient was taken to theatre for an examination under anaesthesia and manual removal of placenta. It was discovered that she had uterine anomaly. A calcified accretic placenta was manually removed. The patient was kept on an oxytocin infusion overnight and given intravenous broad spectrum antibiotics. She made a good recovery and was discharged home after 3 days on oral antibiotics.

KEYWORDS: Retained placenta; Uterine anomalies; Life-threatening; Catastrophic haemorrhage; Reproductive outcomes; Maternal mortality.

INTRODUCTION

A retained placenta is a life-threatening condition as it can cause catastrophic haemorrhage and maternal mortality and morbidity. It complicates 2-3% of vaginal deliveries and causes postpartum haemorrhage. Its management must be prompt and effective. The prevalence of uterine anomalies in the general population is 1:201 (0.50%), the commonest types being septate (34%) and bicornuate (39%) of all cases of uterine anomalies. Congenital uterine anomalies are associated with the highest incidence of reproductive failure and obstetric complications. There is scanty data on uterine anomalies and retained placenta in the literature. A retained placenta in a rudimentary horn of a double uterus was described by Wilson in 1955. It is therefore important to document more cases as to alert clinicians and help save lives.

CASE REPORT

A 32 year-old P, G went into spontaneous labour at 37 weeks gestation. She had had a normal antenatal period. In her previous obstetric history, she had delivered NVD 8 years ago a baby boy birth weight 3000 g. The couple had struggled to conceive but had not consulted a clinician to undergone any investigations for subfertility. She had no other medical or surgical history. Precipitate labour progressed to a NVD of a baby boy within 4 hours of onset. The baby’s birth weight was 2710 g. The Apgar scores were 8, 9 and 10 at 1, 5 and 10 minutes respectively. The third stage of labor was managed actively by giving oxytocin 10 international unit (IU) at the delivery of the anterior shoulder. Gentle cord traction was attempted but there were no signs of placental separation after 15 minutes. There was minimal bleeding per vagina. An oxytocin infusion of 40 IU in a litre of normal saline was commenced. After 30 minutes post-delivery, there were still no signs of placental separation. Another attempt at gentle cord traction led to the cord snapping.

The patient was taken to theatre for an examination under anaesthesia and manual removal of placenta. There were second degree tears in the vulva and the cervical cervix or cervical os had contracted significantly. On insertion of the hand, it went right into the fundus...
and no placenta was felt. The empty cavity was roomy. While moving the hand inside a second opening was felt (Figure 1).

This was a case of uterine anomaly only detected in theatre. The newly discovered uterine cavity went right up to the costal margin. There at the fundus an accretic placenta was found. The placenta was sheared off and manually removed. It was a calcified placenta. The second degree tears were repaired. The patient was kept on an oxytocin infusion overnight and given intravenous broad spectrum antibiotics. She made a good recovery and was discharged home after 3 days on oral antibiotics.

DISCUSSION

Retained placenta poses great danger to maternal health as it can lead to catastrophic haemorrhage, the risk is even higher with an adherent placenta. Prenatal diagnosis of abnormal placentation allows anticipation of multidisciplinary team4 management that prevents adverse outcomes. It is important that it is recognized early and prompt steps taken to have it manually removed under anaesthesia. An association of a retained placenta and uterine anomaly has not been described in the literature hence this case is to highlight this association. Uterine morphology can be ascertained outside pregnancy by hysterosalpingography and laparoscopy.5 Magnetic resonance imaging (MRI) can also be a useful tool to diagnosis. Some uterine anomalies may permit normal obstetric outcomes.5

Women with congenital uterine malformations usually have higher incidence of subfertility and complications during pregnancy and delivery.6 The risks include preterm pre-labour rupture of membranes, small for gestational age babies and pre-term delivery.7,8 There are also risks of malpresentation and caesarean section (C-section) delivery,7 and rupture of rudimentary uterine horn.8 The complication of a retained placenta in uterine horn is not described in the literature. Uterine anomalies are associated with both normal and adverse reproductive outcomes.9,10

CONCLUSION

This case reminds clinicians of rare clinical associations that lurk underneath the surface undetected and yet pose significant danger to maternal health. When faced with unusual clinical findings, clinicians must explore other areas that may lead to the discovery of unexpected pathology. This can be life-saving.

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