Uterus Didelphys With Cervical Incompetence – A Rare Case Report

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ABSTRACT

Uterus didelphy is one of the congenital uterine anomalies due to defective medial fusion of mullerian ducts. This anomaly is known to have poor reproductive outcome and women with this condition often have to be treated for infertility. An individual physician’s experience with these abnormalities is limited, while for many physicians to report individual cases for collective review in future would be meaningful. In this case report, a 33 year old Gravida 5, had spontaneous abortions 4 with cervical incompetence and having uterus didelphys was managed effectively to deliver a preterm baby by caesarean section.

Keywords: Cervical incompetence, Congenital uterine anomaly, Reproductive performance, Uterus didelphys.

INTRODUCTION

U terine anomalies are associated with both normal and adverse reproductive outcomes. 5–10% of women with recurrent early pregnancy loss and up to 25% of women with late first or second-trimester pregnancy loss or preterm delivery were reported.1-4 Overall, uterine anomalies are associated with difficulty in maintaining a pregnancy and not an impaired ability to conceive.1,2,4 The proper management of infertile women with uterine anomalies is controversial.

Uterus didelphys is a type of mullerian duct anomaly (class III) where there is complete duplication of uterine horns as well as duplication of the cervix, with no communication between them.2 Complete failure of medial fusion of the two mullerian ducts may result in duplication of uterus and cervix with single or double vagina. It occurs between 12th and 16th week of pregnancy and is characterized by two symmetric, widely divergent uterine horns and two cervices. So the uterine volume in each duplicated segment is reduced. Each hemiuteri is associated with one fallopian tube and a longitudinal vaginal septum. True prevalence of this anomaly is unknown as it may be discovered in later part of patient’s life when presenting with infertility.

The incidence of uterus didelphys is 0.03% in women population.2 The chance of seeing a pregnancy continuing to term is significantly reduced, down to only 20%, with 1/3rd of pregnancies ending in abortion and over half in premature deliveries. Only 40% of pregnancies resulted in living children.2

According to literature, Uterus didelphys leads to infertility and recurrent first-trimester pregnancy loss,5 which also causes pregnancy complications like; Preterm birth, preterm premature rupture of membrane (PPROM), breech presentation, placenta previa, placental abruption, IUGR and caesarean section.8-14 In this case report, we present, a 33 years old female of four miscarriages with double uterus and vagina was delivered of a preterm baby by caesarean section from one uterus
showed bulky anteverted uterus while pv in left side showed normal size uterus. Investigations revealed, Haemoglobin 10.6 gm%, TLC 7200/mm3, random plasma glucose 87 mg/dl, chromosomal study of both husband and wife revealed normal chromosomal pattern (Fig 4 and Fig 5). She was advised to reduce weight and was started with folic acid 5 mg/day.

On 7.03.2006 she reported for her 5th pregnancy with the complain of mild bleeding pv and her LMP was on 23.01.2006. USG showed a intrauterine gestational sac of 6 week at right horn of uterus and left horn was empty (Figure 6). Repeat ultrasound on 17.03.2006 showed active intrauterine foetus of size 7 weeks 4 days with no retro placental clot. She was advised complete bed rest and was prescribed with folic acid 5 mg tablet once daily, natural micronized progesterone 200 µg vaginal pessary twice daily and injection human chorionic gonadotrophin 5000 IU intramuscularly twice weekly. Injection human chorionic gonadotrophin was continued till 12 weeks of pregnancy and then discontinued. At 16 weeks and 2 days on 17.05.2006 she came to OPD with the complain of heaviness at lower abdomen and feeling of baby coming out of vagina. On examination uterus was 16 weeks size and irritable. Perspeculum examination showed 1cm long cervix per vaginum with the USG showing cervical length 2.0 cm. For that, Mc-Donald stitch was put on the cervix and sustained release tablet isosuprine hydrochloride 40 mg/day continued throughout the pregnancy. On 22.07.2006 at 25 weeks 5 day she complained of severe lower abdominal pain. Uterus was irritable and urine examination showed no abnormality. She was managed conservatively with oral nifedipine capsule 20 mg tid for 3 days. On 13.08.2006 at 28 weeks 5 days she was given with two doses of injection Betamethasone 12 mg every 12 hourly prophylactically and was repeated on 27.08.2006 at 30 weeks 5 days as the patient was complaining of pain abdomen and had slight bleeding pv. USG revealed placenta at upper segment with small retro placental clot. Hence she was advised absolute bed rest.

On 26.09.2006 at 35 weeks 1 day she was admitted in hospital at 6 am with the complain of pain abdomen and decreased foetal movement. Her Mc-Donald stitch was removed and labour was assessed. 12 hours later due to irregular heart rate pattern of the foetus she was taken to the operating room where a lower segment caesarean section was done and a preterm female baby of Apgar score 9 and 10 at 1” and 5” respectively, weighing 2400 gm was delivered. During operation the pregnancy was in right horn of the uterus and left horn of the uterus was ill developed (Figure 7). Her postoperative course was uneventful and she was discharged to home on 8th postoperative day with the infant. No complication was encountered with the baby and the mother during the last 7 years of follow up.

In the mean time, she delivered her second live baby at 33 week by caesarean section on 12. 08.2010. The post partum period after the second issue was uneventful.
Figure 3: Pervaginal exam showing double vagina with a septum in between

Figure 4: Karyotype report of husband of the patient

Figure 5: Karyotype report of the patient

Figure 6: Ultrasound image of pregnant uterus at 6 weeks showing gestational sacs at right horn

Figure 7: Caesarean section of the patient showing the pregnancy at right horn and left horn ill developed

DISCUSSION

The failure of fusion of the two mullerian ducts results in duplication of mullerian structures; a didelphic uterus has two uteri, two endometrial cavities and two cervices. A longitudinal vaginal septum is present in 75% cases. True prevalence of congenital uterine anomalies in population is not known, which varies from 0.1 to 10%. According to a report, the incidence of diagnosed uterine anomalies was close to 0.45%, however the incidence of uterus didelphys was 24.2%, while others found a mean incidence of 11%. Congenital uterine anomalies are associated with higher incidence of reproductive failure and obstetric complications. Women with this form of congenital anomaly required infertility treatment more frequently than women with other uterine anomalies and the overall reproductive performance of uterus didelphys is poor. Others feel that didelphic uterus offers the best chance for a successful pregnancy with a fetal survival rate as high as 64%. Multifetal gestation is rare in women with uterus didelphys. There has also been a triplet pregnancy with uterus didelphys with 72 days lapse between the delivery of the first two fetuses and the third. The presence of a
malformed uterus in a woman, impairs reproductive performance by increasing the incidence of early and late abortions, preterm deliveries as well as the rate of obstetric complications.13,17,18

It was shown a mean 32.9 % abortion rate, a mean 28.9 % preterm delivery rate, a mean 36.2 % term delivery rate with a mean 56.6 % live birth rate in uterus didelphys.7

Uterine anomalies are associated with diminished uterine cavity size, insufficient musculature, impaired ability to distend, abnormal myometrial and cervical function, inadequate vascularity and abnormal endometrial development.7,19,20,21 These abnormalities of space, vascular supply and associated local defects contribute to increased rate of recurrent pregnancy loss and preterm delivery.22 Our patient also had four recurrent abortions where no cause was found. It was found that the spontaneous abortion rate ranges from 32 to 52% and preterm birth rates from 20 to 45%. Recurrent pregnancy loss is due to decreased uterine volume and associated cervical incompetence.22

Our patient had a cervical length of 2.0 cm at USG and in vaginal exam cervical length was 1 cm. According to a report, the cervical incompetence was 30% of mullerian anomaly.22 With cervical encirclage or surgical correction, incidence of term pregnancies in the group with documented cervical incompetence increases from 26 to 63%.28 Abnormal uterine blood flow and decreased muscle mass in uterus didelphys leads to IUGR of foetus.27

It is suggested that abnormal irregular contraction may lead to microablation of placenta leading to placental insufficiency and IUGR.

Women with uterine didelphys accounted for higher proportion of preterm birth < 34 weeks and < 37 weeks than the control.7,28 According to the report 21% of pregnant women with didelphys uterus deliver prematurely.29 Our patient delivered at 35 weeks 1 day. Increased caesarean delivery rate is associated with higher rate of malpresentation and vaginal anomalies such as a longitudinal vaginal septum. According to the report, the highest caesarean section rate (82%) in patients with uterus didelphys.29 Our patient had a cesarean section at 35 weeks 1 day due to irregular fetal heart rate pattern. Spontaneous vaginal delivery as well as caesarean section at term has been reported.

CONCLUSION

Didelphic uterus is a very rare anomaly and it can lead to recurrent pregnancy loss due to decreased uterine volume and associated cervical incompetence. Preterm birth rate is increased due to irregular uterine contraction. By adequate rest, tocolytics and cervical encirclage, a live baby can be awarded.

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