Case Report

Spontaneous pregnancy at term with uterus didelphys: a case report

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ABSTRACT

The uterus didelphys results from the absence of fusion of the bilateral mullerian ducts. It is a rare pathology. This malformation concerns 5% of uterine malformations from mullerian ducts and affects one woman in 1,000-30,000. Obstetrical complications of this malformation are numerous. The chance of reaching term for pregnancies with didelphys uterus is reported as 20%–30%. Authors report a case of spontaneous term pregnancy in a 21-year-old primiparous woman with a didelphic uterus. The patient had an unexplained seizure with fetal bradycardia. An emergency cesarean section was performed and allowed the birth of a hypotrophic neonate of 2240g and the discovery of didelphic uterus. Pregnancy developed in the left hemi-uterus. Speculum examination at the end of the procedure showed a longitudinal vaginal septum. There was no associated urinary tract and renal malformation. Scheduled cesarean will be performed from her next pregnancy. The uterus didelphys should be diagnosed early. MRI and 3D echography are necessary for diagnosis. Pregnancy is often complicated, and follow-up needs to be planned. Cesarean section is not systematic.

Keywords: Delivery, Pregnancy, Uterine didelphys, Uterine malformation

INTRODUCTION

The uterus didelphys results from the absence of fusion of the two Mullerian ducts.¹ Both ducts evolve into two separate hemi uteri. Other structures are also separated, ovaries, fallopian tubes, and cervix. The vagina can be single or double.²

This malformation concerns 5% of uterine malformations from mullerian ducts and affects one woman in 1,000-30,000.³,⁴ Congenital uterine malformation is frequently asymptomatic. The vaginal septum may cause dyspareunia, leucorrhea, dysmenorrhea if it is completely obstructed.⁴ Pregnancy anomalies or infertility may lead to the diagnosis. Transvaginal ultrasonography is very useful for the diagnosis. But MRI becomes the first exam to confirm it.² The uterus didelphys may be identified during caesarean section.

This malformation is associated with a renal malformation in 30% of the cases.³ Pregnancy complications with uterus didelphys are numerous such as abortion, malpresentation, premature rupture of membrane, intrauterine growth restriction and preterm birth.² The chance of reaching term for pregnancies with didelphys uterus is reported as 20%–30%.² Through this
presentation, we report a case of pregnancy reaching term on uterus didelphys discovered during an emergency Caesarean section.

CASE REPORT

It is about a 21-year-old patient, primigravida, admitted in obstetric emergency departments for unexplained generalized tonic-clonic seizures, at 39 weeks of Amenorrhea and five days. The pregnancy was spontaneous and normal evolution until then. The patient had no history of high blood pressure or epilepsy. The disease is summarized by the appearance of a generalized tonic-clonic convulsive seizure without prodrome followed by a loss of urine, drooling and post-critical unconsciousness of about one minute, which motivates its consultation. During examination, the blood pressure was 110/60 mmHg, the heart rate was 60 beats / min. She was afebrile. The patient was conscious. She complained of moderate headache. The neurological examination was normal. There was no recurrence of the convolution. Proteinuria was absent in the urine dipstick. There was no uterine contraction. The fetal cardiotocography had shown fetal bradycardia at 100 beats per minute. An emergency cesarean section was performed for fetal bradycardia.

The surgical intervention was performed under general anesthesia. A segmental hysterotomy was performed. The extraction was difficult and allowed the birth of a 2240g male newborn with intrauterine growth restriction. The Apgar score in the fifth minute was 9. The deliverance was normal. At the exteriorization of the uterus outside the peritoneal cavity, we discovered a uterus didelphys (Figure 1).

Pregnancy developed in the left hemi-uterus. The hysterorrhapsy was without particularity. Examination of the vagina at the end of the procedure showed a longitudinal septum with two vaginas and two cervix (Figure 2). The operative sequence was normal. She came out of the hospital on the fifth day. There is no recurrence of the convulsive seizure and the neurological examinations performed were normal. The patient was seen one month after the procedure. There was no neurological abnormality. Echography of the urinary tract and kidney was normal. Scheduled cesarean will be performed from her next pregnancy.

DISCUSSION

The uterus didelphys (type III according to the American Fertility Society's classification: AFS) corresponds to two completely separate hemi-uterus with two cervix. The horns can be asymmetrical. It results from a fusion failure of the mullerian ducts. It develops at the 10th-13th week of embryonic development. Diagnosis is suspected by clinical signs (amenorrhea, dysmenorrhea, infertility, repeated miscarriage, premature delivery). Imaging exams confirm the diagnosis. Hysterography is no longer the first-line examination. 3D echography remains very useful, but MRI remains the gold standard in the exploration of pelvic malformations. In our context, the patient did not receive one of these exams. The diagnosis was made only intraoperatively.

Regarding the outcome of pregnancy in patients with didelphic uterus, Heinonen reported a 30% of miscarriage, 43% of malpresentation, and 82% of cesarean section and 11% of small for gestational age. According to Chan and al, Patients with uterus didelphys had a preterm birth risk of 3.58 times higher (95% CI = 2.00-6.40), with a 3.7-fold increase in malpresentation (95% CI=2.04-6.70). By contrast, the risk of miscarriage is identical. For our case, the pregnancy was reached term but was complicated by a neonatal hypotrophy. Convulsion alone does not explain the low birth weight.

Delivery is most often done by cesarean section. An anomaly of the presentation is observed. Most often, the fetus is in breech presentation. Mechanical obstructed labor may also occur. The hemi-uterus is sometimes a
previa obstacle. However, normal deliveries have been reported even in cases of vaginal septum.

In our case, the delivery route will be a cesarean section scheduled for the next pregnancy. Indeed, in our practice, a history of previous Cesarean section associated with a uterine malformation is an indication of cesarean section. In the literature, the risk of uterine rupture in patients with non-scarring uterus was 1: 15,000. In addition, a study has shown that the risk of uterine rupture in patients with a uterine malformation of Mullerian origin is identical to the general population. But vaginal delivery is rare in these patients. The frequency of cesarean section rises to 69.2%.

There was no renal malformation in our patient. In the literature, a didelphic uterus is often associated with a renal malformation. In 20 to 30% of the cases, the renal malformation is presented. This can be an agenesis, an ectopia, or a malrotation. A systematic study of the kidneys must be performed when a uterine malformation is discovered. This is part of a malformation assessment. The MRI is useful to identify these malformations.

**CONCLUSION**

To conclude, the frequency of the uterus didelphys is rare. The diagnosis must be precocious. MRI and 3D echography are of particular interest for diagnosis. Once diagnosed, follow-up should be closely and regularly as complications are numerous. The delivery route depends on the presentation of the fetus and the occurrence of a previa obstacle by the hemi-uterus during labor. Cesarean section is not systematic.

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