RARE CASE REPORTS OF COMPLETE UTERINE SEPTUM WITH LONGITUDINAL VAGINAL SEPTUM PRESENTING AS DIDELPHIC/ UNICORNUATE UTERUS

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HOW TO CITE THIS ARTICLE: Kalpana B, Gupta R. Rare case reports of complete uterine septum with longitudinal vaginal septum presenting as didelphic/ unicornuate uterus. J. Evolution Med. Dent. Sci. 2017;6(94):6920-6922, DOI: 10.14260/jemds/2017/1501

PRESENTATION OF CASE

Müllerian duct anomalies (MDA) comprise of a heterogeneous group of genital malformations accounting for an incidence of 3%-4% in the general female population. Use of improved imaging modalities such as 3D transvaginal ultrasound (TVUS) and magnetic resonance imaging (MRI) has helped clinicians to better identify and characterise such anomalies.1 Septate uterus with cervical duplication and longitudinal vaginal septum is a rare Müllerian anomaly, hence its true incidence is not known. We report 2 cases of this rare anomaly.

CASE REPORT 1

A 23 years old female, married for 1.5 years presented to us with complaints of primary infertility. Her menstrual cycles were regular (3/30) with dysmenorrhea and dyspareunia. Per speculum examination revealed a longitudinal vaginal septum and 2 cervixes. Hysterosalpingogram (HSG) (Fig. 1) and TVUS suggested 2 uterine cavities with 2 cervixes. A right complex ovarian cyst was also seen on TVUS. MRI confirmed the diagnosis as uterine didelphys with adenomyosis with right ovarian chocolate cyst with left PCOD. There was no evidence of renal anomaly. We proceeded with hysteroscopy.

Laparoscopy revealed a broad single smooth uterine fundus with bilateral normal tubes. Right ovary had a chocolate cyst for which cystectomy was done and left ovary was normal. Hysteroscopy revealed a complete uterine septum extending till external os and a longitudinal vaginal septum as shown in Fig. 2.

CASE REPORT 2

A 20 years old female, married for 3 years presented to our clinic as case of primary infertility. She had regular menstrual cycles with no dysmenorrhea or dyspareunia. She had undergone laparotomy with right salpingo-oophorectomy for right chocolate cyst at the age of 16 years. Her HSG report was suggestive of left unicornuate uterus as shown in Fig. 3. She had a USG report suggestive of unicornuate uterus with absent right kidney and a chocolate cyst in left ovary. She also had an MRI report, which suggested a possibility of Uterus Didelphys with left ovarian chocolate cyst. As there was a variation between HSG and MRI impression, we proceeded with hysterectomy.

Hysteroscopy revealed a small uterine cavity deviated to the left side, left ostium was seen but right ostium was not visualised suggestive of left unicornuate uterus. On laparoscopy, the uterine fundus was smooth and broad. Right tube and ovary were absent. Left tube was normal and left ovary had Polycystic Ovarian Disease (PCOD).

In view of these inconclusive findings, we probed to look for another opening in the perineum and found a small hole of 0.2 cm on right side of vulval outlet with mucus and pus coming out of it. We dilated the opening and let the pus out. We introduced the hysteroscope through the same opening and visualised the right half of vagina and cervix and gently proceeded with hysteroscopy. The uterine cavity was small and the right ostium was normal. There was a complete uterine septum extending from uterine fundus to external os with a longitudinal vaginal septum. Only after Hysteroscopy, this diagnosis was confirmed.

For both patients, vaginal septum was excised followed by unification of 2 cervixes. Transcervical resection of uterine septum was done with resectoscope. The post-operative period was uneventful. Patients were given estradiol valerate in post-operative period to promote endometrial healing and to prevent intrauterine adhesion formation.

A follow-up hysteroscopy showed a normal uterine cavity.

DIFFERENTIAL DIAGNOSIS

Complete septate uterus with longitudinal vaginal septum with uterus didelphys.

DISCUSSION

This unusual Müllerian anomaly is inconsistent with the theory of linear caudal to cephalad Mullerian fusion. ESHRE classification has divided the female genital tract anomalies into uterine, cervical and vaginal anomalies. Uterine anomalies are subdivided into 7 classes:

- U0 Normal uterus.
- U1 Dysmorphic uterus.
- U2 Septate uterus.
- U3 Bicornoral uterus.
- U4 Hemi-uterus.
- U5 Aplastic.
- U6 Unclassified Malformations.

Among them, the septate uterus is the commonest. These uterine anomalies may be associated with cervical and/or vaginal anomalies. However, complete septate uterus with a cervical septum and longitudinal vaginal septum is a rare uterine anomaly.

Patients described in literature have presented with a variety of obstetric and gynaecologic symptoms, some have been asymptomatic. Diagnosis is made by careful examination and imaging. MRI provides good cervical imaging and is considered the best non-invasive method for
differentiating septate, bicornuate and didelphys uterus. However, in our cases MRI findings were inconsistent with hysterosalpingography findings. 3D TVUS is emerging as a better non-invasive diagnostic tool for Müllerian anomalies.[1] Khaled Abd et al compared the different imaging modalities to differentiate septate and bicornuate uterus.[1] They found sensitivity of 77.4% and specificity of 60% for HSG. MRI showed sensitivity of 93.5% and specificity of 80%. The 3D ultrasound showed the highest diagnostic parameters with sensitivity of 96.7% and specificity of 100%. However, if the diagnosis is inconclusive with the various imaging modalities, hysteroscopy is the best way to confirm the diagnosis.

Our patients came with infertility and variable findings on USG, HSG and MRI without any conclusive diagnosis. Direct visualisation with hysteroscopy revealed the final diagnosis. It also facilitated to perform the corrective surgery to improve the fertility and future obstetric outcome. Resection of the vaginal septum is easy and commonly performed. Hysteroscopic resection of a uterine septum to improve the obstetric outcome is the gold standard and is indicated in females with unexplained infertility and previous poor obstetric outcomes. Seddigheh et al reported a pregnancy rate of 67% and a live birth rate of 57.5% in infertile women who underwent hysteroscopic septoplasty.[2] Kazem Nouri et al reported overall pregnancy rate of 69% and live birth rate (LBR) was 49% after hysteroscopic septoplasty.[3]

The union of the two cervices is another area of controversy. Vijay A et al avoided the resection of cervical septum to avoid cervical incompetence in future.[4] However, Meen Jiun Seet et al, SC Ribeiro et al and ME Parsanezahed et al favoured resection of cervical septum.[5,6,7] ME Parsanezahed et al found that resection of cervical septum during Hysteroscopic metroplasty of uterine septum makes the procedure safer, easier and less complicated than the procedure with preservation of the cervical septum.[7] The reproductive outcome is the same in the two procedures. Hence, we also proceeded with unification of cervices. Most of the recent studies favour non-invasive imaging modalities as the preferred diagnostic tests for Müllerian anomalies. However, in our cases non-invasive imaging modalities gave a misleading diagnosis. Confirmatory diagnosis was made only with hysteroscopy. Hysteroscopy has got a therapeutic value in addition to its diagnostic abilities. Hysteroscopy should be recommended when imaging modalities do not give a confirmative diagnosis, especially in symptomatic patients.

Figure 1. HSG Image showing 2 Uterine Cavities with 2 Cervices

Figure 2. Hysteroscopy Image showing Cervical and Longitudinal Vaginal Septum

Figure 3. HSG Image showing Left Unicornuate Uterus with Left Tube

FINAL DIAGNOSIS
Complete Uterine Septum with Longitudinal Vaginal Septum.

REFERENCES