Doppler sonography and 3D CT Angiography of Acquired Uterine Arteriovenous Malformations: Report of Two Cases

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ABSTRACT
Uterine Arteriovenous Malformations (AVMs) are rare but life threatening causes of abnormal vaginal bleeding. Accurate clinical and radiological diagnosis is essential because uterine instrumentation that is often used for management of other sources of abnormal bleeding, can lead to massive hemorrhage. Timely diagnosis and early proper treatment can markedly reduce the associated disease mortality. Ultrasound with colour and spectral doppler is the initial imaging modality of choice. Three-dimensional computed tomography (CT) angiography can determine the actual extent of the vascular malformation and helps in pre-interventional planning noninvasively. Uterine AVM can be either congenital or acquired in nature with latter being more common. We hereby report two cases of acquired uterine arteriovenous malformations diagnosed by color doppler sonography and confirmed by three-dimensional CT angiography. Both the cases reported here had previous history of dilation and curettage for abortion. Clinically one patient presented with profuse uterine bleeding and another with meno-metrorrhagia and both cases underwent surgical removal of uterus.

CASE REPORTS

CASE 1
A 38-year-old G3 P2, presented with history of recurrent menometrorrhagia for the past 2 months. She gave previous history of missed abortion at 12 weeks gestational age three years back for which dilation and curettage (D & C) was performed. No other significant past medical or surgical history could be elicited. General examination of the patient was normal. Per-vaginal examination showed slightly enlarged pulsatile uterus with normal vulva and cervix. Gray–scale ultrasound showed presence of tubular anechoic lesions in uterus which showed low impedance, high velocity color flow [Table/Fig-1] on doppler images. Color mosaic pattern was observed representing turbulent flow. Peak systolic velocity was 98cm/sec with resistive index of 0.33. These findings could represent arteriovenous malformation of uterus. Three-dimensional CT angiography was requested to confirm the diagnosis, rule out extra uterine involvement and to localize feeder vessels. CT angiography was performed in 64-slice multi-detector CT which showed bunch of dilated vascular channels within the uterus. The arterial supply was from uterine arteries on both sides which appeared dilated [Table/Fig-2a, b]. Patient refused transcatheter arterial embolization for economic reasons; hence hysterectomy was done after ligation of bilateral internal iliac arteries. Histopathology of the hysterectomy specimen confirmed the diagnosis of uterine arteriovenous malformation.

CASE 2
A 54-year-old G2 P2, presented with profuse vaginal bleeding for the past 3 days. She had undergone curettage for retained products

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of conception 2 years back. No other significant past medical of surgical history could be elicited. General examination of the patient showed pulse rate of 104 beats / min with blood pressure of 90 / 60mm Hg. Emergency transvaginal ultrasound showed presence of bunch of vascular channels replacing uterus [Table/Fig-3a,b]. These vascular channels showed turbulent color flow with peak systolic velocity of 110cm / sec and low resistive index of 0.27. One of these vascular channels showed disproportionate dilatation suggestive of aneurysm formation [Table/Fig-3a,b]. Patient was shifted immediately for CT angiography to confirm the diagnosis of uterine AVM and to localize feeding vessels. CT angiography showed presence of bunch of dilated vascular channels replacing uterine cavity with aneurysm formation in one of vascular channel likely flow related aneurysm [Table/Fig-4a,b]. The arterial feeders were from bilateral uterine arteries which appeared dilated. Although few of the other branches of both internal iliac arteries were dilated, they did not supply the AVM. Patient underwent hysterectomy after ligation of bilateral internal iliac arteries. Histopathology of the hysterectomy specimen confirmed the diagnosis of uterine arteriovenous malformation.

**DISCUSSION**

Uterine arteriovenous malformations (AVMs) are one of the rare causes for abnormal uterine bleeding (AUB) [1,2]. The true incidence of uterine AVMs is not known, however the reported cases are on the rise due to the increasing availability of imaging modalities like ultrasound, CT and magnetic resonance imaging (MRI) [1-3].

Uterine AVMs are classified broadly into congenital and acquired, of which acquired AVMs are more common, as in our cases [1,2]. Abnormal differentiation of primitive vascular structures during embryogenesis leads to development of congenital uterine AVM. The common causes for acquired AVMs are iatrogenic resulting from prior dilation and curettage, uterine surgery or therapeutic abortion, direct uterine trauma, uterine malignancy and gestational trophoblastic disease [3,4]. Both the cases reported here gave previous history of dilation and curettage. The difference between congenital and acquired AVMs are that congenital AVMs have multiple feeding arteries with a central nidus and many draining veins, contrary to acquired AVMs which have multiple small arteriovenous fistulae between intramural arterial branches and myometrial venous plexus [1,2].

Uterine AVM have been reported in ages ranging from 18 years to 72 years. The incidence of uterine vascular malformations are relatively higher in patients after abortion than in postpartum or outpatient groups as reported in a study [5]. Affected patients usually present with menorrhagia or menometrorrhagia, like in our cases [3]. Few cases may present with heavy uterine bleeding requiring transfusion.

The diagnosis of uterine AVM is clinically important as therapeutic uterine instrumentation like D & C which is done in cases of retained products of conception, may aggravate the bleeding in cases of uterine AVM [3].

The initial imaging modality of choice for uterine AVM is sonography with color and spectral doppler analysis [6]. It also helps in ruling out other conditions causing abnormal uterine bleeding like retained products of conception, trophoblastic disease, sarcoma of uterus, uterine artery pseudoaneurysm and pelvic varicose veins. In both the cases reported here, sonography with color doppler accurately suggested the diagnosis of uterine AVM. Gray scale ultrasound of uterine AVM commonly shows heterogeneous myometrium with multiple hypo to anechoic tubular structures. In few cases, grey scale ultrasound may show focal inhomogeneity or a mass resembling leiomyoma or a polyp. Application of color doppler sonography will reveal abnormal hypervascularity within the lesion with color mosaic of aliasing and flow reversal. Spectral doppler ultrasound shows low resistive index values with high velocity flow [6].

The role of three-dimensional CT angiography is to define the extent of involvement, to rule out extraterine involvement, to differentiate between congenital and acquired uterine AVM non-invasively and to delineate the feeder vessel / uterine arteries [7,8]. In both the cases reported here, CT angiography helped in answering all the questions posted above. In patients considered for embolization, CT angiography can reduce the study time, radiation dose and contrast required for the procedure and it is also useful for anatomical conceptualization to the gynaecologist in patients planned for surgery [9]. Magnetic resonance imaging (MRI) shows presence of flow voids in the region of vascular malformation and MR angiography provides the details about the vascular supply and drainage. However MRI is a costly radiological investigation and acquisition time is relatively longer for MRI and MR angiographic images compared to CT angiography. Hence CT angiography is preferred in unstable patients with heavy bleeding, those who cannot afford MRI and when MRI is contraindicated. Digital subtraction angiography (DSA) though gold standard, is usually reserved for patient planned for transcatheter embolization, because of its invasive nature.

Past medical history is also important in patients presenting with AUB and can suggest the possibility of uterine AVM in proper clinical setting. Both the cases reported here gave similar past history of dilation and curettage and in both cases ultrasound with doppler and three-dimensional CT angiography helped in confirming the diagnosis and planning the treatment.

Hence a complete clinical history combined with radiological investigations like color doppler ultrasonography and CT angiography can help in making accurate diagnosis at the
earliest and reducing patient mortality. The treatment options for uterine AVMs include surgery and transcatheter uterine artery embolization [2,6]. Transcatheter uterine artery embolization has become an effective alternative to surgical intervention for uterine AVMs with major advantage of maintaining child bearing capacity [6]. Hysterectomy is preferred if patient does not want future pregnancies or when transcatheter embolization fails [1]. Since both our patients had completed their family and were economically poor to afford transcatheter uterine artery embolization, they underwent hysterectomy.

Therefore, in any patient with unexplained uterine bleeding, color doppler ultrasonography should be performed at the earliest to facilitate accurate diagnosis. Three-dimensional CT angiography helps in confirming the diagnosis of uterine AVM and provides valuable cross sectional anatomical details in patients planned for surgery or embolization.

REFERENCES