Case Report

Mixed Mullerian tumor: a case report

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ABSTRACT

Malignant mixed Müllerian tumor, also known as malignant mixed mesodermal tumor, MMMT and carcinosarcoma, is a malignant neoplasm found in the uterus, the ovaries, the fallopian tubes and other parts of the body that contains both carcinomatous (epithelial tissue) and sarcomatous (connective tissue) components. A 50-year female P1L1A1 with complaints of pervaginum bleeding and pain in abdomen was diagnosed to have carcinosarcoma endometrium for which hysterectomy with bilateral salpingo-opherectomy was done.

Keywords: Carcinosarcoma, Hysterectomy, Malignant mixed Mullerian tumor

INTRODUCTION

Malignant mixed Müllerian tumor, also known as malignant mixed mesodermal tumor, MMMT and carcinosarcoma, is a malignant neoplasm found in the uterus, the ovaries, the fallopian tubes and other parts of the body that contains both carcinomatous (epithelial tissue) and sarcomatous (connective tissue) components. It is divided into two types, homologous (in which the sarcomatous component is made of tissues found in the uterus such as endometrial, fibrous and/or smooth muscle tissues) and a heterologous type (made up of tissues not found in the uterus, such as cartilage, skeletal muscle and/or bone). MMMT account for between two and five percent of all tumours derived from the body of the uterus, and are found predominantly in postmenopausal women with an average age of 66 years. Risk factors are similar to those of adenocarcinoma uterus and include obesity, exogenous oestrogen therapies, and nulliparity. Less well-understood but potential risk factors include tamoxifen therapy and pelvic irradiation.1 A 50-year female P1L1A1 with complaints of pervaginum bleeding and pain in abdomen was diagnosed to have carcinosarcoma endometrium for which hysterectomy with bilateral salpingo-opherectomy was done.

CASE REPORT

A 50-year female P1L1A1 with previous LSCS, tubectomized 11 years back, resident of Namdev Nagar, Wadgaosheri came to emergency ward with chief complaints of pain in abdomen and per vaginum bleeding on and off for 1 month, no history of per vagum discharge. On detailed evaluation patient gives the history of 6 months amenorrhoea followed by bleeding per vaginum for 1 month soaking 4 pads per day associated with passage of clots. Investigations were done with Transabdominal sonography showing bulky uterus thickened and irregular endometrium with multiple cystic changes causing thinning of myometrium with significant peripheral vascularity suggestive of malignant neoplastic etiology - carcinoma endometrium. Dilatation and curettage was done for tissue diagnosis with histopathology report stating no typing possible, after which tissue block was sent for review examination from senior pathologist which was suggestive of carcinosarcoma (malignant mixed Mullerian tumor). Patient was followed with CT (abdomen + pelvis) which showed bulky uterus 16.6 × 9.4 × 8.8 cm with heterogeneously enhancing polypoidal mass lesion involving uterine cavity approx. 8.7 × 5.4 × 5.2 cm.
Intramural fibroid 3.4 × 4.1 cm involving left lower wall. Subserosal fibroid 1.4 × 3 cm involving right lower lateral wall. Both ovaries are visualized and normal, bowel and bladder appears normal. Oncoreference was done and hysterectomy was advised with follow-up USG. Preanesthetic checkup for the patient was done with all routine blood investigations and patient was posted for abdominal hysterectomy with bilateral salpingo oophorectomy, lymph node dissection was done and specimen sent for histopathological examination. Post hysterectomy histopathological report was suggestive of malignant mixed Mullerian tumor without local spread. Tumor not infiltrating myometrium muscle fibers, section studied from parametrium shows adipose tissue free of tumor, no evidence of pelvic and paraaortic lymph node involvement. Ultrasonography was done with no significant abnormality. Review oncoreference was done and pelvic radiation was advised with close follow up, patient was discharged on request after stitch removal and advised follow up after 7 days, patient however lost follow up.

**Figure 1: Specimen showing uterus with polypoid tumor.**

**DISCUSSION**

Malignant uterine neoplasm’s containing both carcinomatous and sarcomatous elements are designated in the World Health Organisation (WHO) classification of uterine neoplasm’s as carcinosarcomas. Gebhardt in 1899 appears to have reported the first case. Meyer, after a personal examination of the slides, accepted it as authentic. Carcinosarcomas representing less than 5% of all uterine tumors, account for 16.4% of all deaths caused by a uterine malignancy. MMMT has been identified in decreasing order of frequency in the vagina, cervix, and ovary and most rarely in the fallopian tubes. There are three main theories regarding the histogenesis of MMMT namely:

- The collision theory suggests that the carcinoma and sarcoma are two independent neoplasms.
- The combination theory suggest that both components and derived from a single stem cell that undergoes divergent differentiation early in the evolution of the tumor.
- The conversion theory suggests that the sarcomatous elements derive from the carcinoma during the evolution of the tumor.

On exploring the literature, we found that W G McCluggage named one more theory: The composition theory suggest that the spindle cell component is a pseudosarcomatous stromal reaction to the presence of the carcinoma.

Outcome of MMMTs is determined primarily by depth of invasion and stage. As with endometrial carcinomas, the prognosis is influenced by the grade and type of the adenocarcinoma, being poorest with serous differentiation. MMMTs are highly malignant; a stage I tumor has an expected five-year survival rate of 50%, while the overall five-year survival rate is less than 20%.1

According to FIGO staging, uterine carcinosarcoma is staged as carcinoma of uterus.

- IA Tumor confined to the uterus, no or <½ myometrial invasion
- IB Tumor confined to the uterus, >½ myometrial invasion
- II Cervical stromal invasion, but not beyond uterus
- IIIA Tumor invades serosa or adnexa
- IIIB Vaginal and/or parametral involvement
- IIIC1 Pelvic node involvement
- IIIC2 Parametrial involvement
- IVA Tumor invasion bladder and/or bowel mucosa
- IVB Distant metastases including abdominal metastases and/or inguinal lymph nodes

Diagnosis of MMMT is most often made postoperatively by histopathological examination and Immunohistochemical (IHC) studies. Microscopically MMMT is composed of both epithelial and mesenchymal elements may be intermittently mixed or be seen as two distinct components.2 The mesenchymal elements may be (a) homologous, containing cells native to the uterus including stromal sarcoma, fibrosarcoma or leiomyosarcoma (2%) or (b) heterologous with mixed components including rhabdomyosarcoma (18%), chondrosarcoma (10%), osteosarcoma (5%) or liposarcoma (1%). Several studies have found concordance of p53 staining between the carcinomatous and sarcomatous components in MMMT.13,14

IHC studies have shown that both the sarcomatous and carcinomatous components often coexpress cytokeratin and vimentin. In the present case, microscopic sections studied showed haphazardly admixed of both epithelial and mesenchymal components. The mesenchymal component showed homologous spindle cell sarcomatous component displaying marked pleomorphism and atypical mitotic activity, suggestive of malignant mixed
Mullerian tumor without local spread, tumor not infiltrating myometrium muscle fibres, section studied from parametrium shows adipose tissue free of tumor, no evidence of pelvic and paraaortic lymph node involvement so stage IA.

CONCLUSION

To conclude MMMT which arise from female genital tract can have both epithelial and mesenchymal component. If epithelial component is benign and mesenchymal component is malignant then it is called as Adenosarcoma. If both epithelial component and mesenchymal component is malignant then it is called as Carcinosarcoma. Malignant mixed Mullerian tumour is a rare, highly aggressive, rapidly progressive neoplasm associated with a poor prognosis.

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