Case Report

Secretory Endometrial Adenocarcinoma: A Rare Sequelae in a Postmenopausal Woman Following Tamoxifen Therapy for Breast Cancer

Sonam Sharma

Department of Pathology, Kalpana Chawla Government Medical College, Karnal, Haryana, India

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INTRODUCTION

Endometrial adenocarcinoma of secretory type is an uncommon variant of well-differentiated endometrioid adenocarcinoma and accounts for only 1%–2% of all endometrial carcinomas.[1] It is histologically a reminiscent of normal early secretory endometrium (17–22 days) and is frequently seen as a focus within the other endometrioid carcinomas rather than in pure form. It occurs predominantly in postmenopausal women and often presents with abnormal vaginal bleeding. Many known risk factors similar to those of other endometrial cancers and polyps such as obesity, hypertension, diabetes mellitus, exogenous estrogen as well as progestin administration, and nulliparity are associated with it.[2] However, tamoxifen therapy has been rarely implicated in its pathogenesis. Nevertheless, this tumor is generally low grade and has a good prognosis.[3] Herein is described an extremely uncommon case of secretory adenocarcinoma of the endometrium which developed in a 60-year-old woman due to the long-term intake of tamoxifen as a treatment for her breast cancer.

CASE REPORT

A 60-year-old woman, G4 L4, presented to the gynecological outpatient department with the chief complaint of bleeding per vaginum for the past 2 months. She had four full-term normal vaginal deliveries and had attained menopause 12 years back. Her medical history revealed that she had undergone right-sided mastectomy for breast cancer 8 years back, for which she had further received six cycles of chemotherapy and had been taking oral 10 mg tamoxifen twice daily since then. She was also diagnosed with hypertension 6 months back and was on oral telmisartan 40 mg/day. However, she denied any history of estrogen replacement therapy after menopause. Her family history of any cancer (breast/uterine/colon/ovarian) or of any other major illnesses was non-contributory. On general physical examination, she had an average built and was anemic. Per abdomen examination revealed mild tenderness in the lower abdomen with no evidence of ascites or any organomegaly. The external pelvis and per speculum examination was unremarkable. Bimanual pelvic examination revealed an open external os and a firm, mobile cervix. The uterus was anteverted, irregular, soft,
and enlarged with free bilateral fornices. No adnexal masses could be palpated. On per rectal examination, the rectal mucosa was free. All other systemic examinations were within the normal limits. Her routine hematological investigations revealed microcytic hypochromic blood picture. Urine and blood cultures were negative. Kidney and liver function tests were normal. Serum antibodies to the human immunodeficiency virus, hepatitis B surface antigen, and syphilis were negative. X-ray chest was normal. Transvaginal sonography (TVS) showed heterogeneous irregular endometrium of 22.2 mm thickness. Papanicolaou smear was performed which was negative for dysplasia or any malignancy. Fractional curettage specimen consisting of multiple fragments of soft, gray–brown material together measuring 2 cm × 1 cm × 1 cm in size was histopathologically examined. Microscopic sections showed a well-differentiated adenocarcinoma with dominant papillary and glandular patterns, consisting of tall columnar epithelial cells showing mild-to-moderate pleomorphism and having small, vesicular, and uniform nuclei with fine nuclear chromatin. The cytoplasm of these cells was clear and exhibited a severe degree of supranuclear and subnuclear vacuolization. The mitotic rate was 2–3/high-power field, and most of the mitosis was atypical. Areas of the stromal invasion were also evident [Figure 1]. A diastase sensitive Periodic Acid-Schiff (PAS) stain showed positivity with the presence of eosinophilic fine granular material in the vacuolated cells, and the stains for mucin were negative [Figure 1b-Inset]. On immunohistochemistry (IHC), tumor cells were positive for estrogen receptor (ER), progesterone receptor (PR), p53 and exhibited a high Ki-67 proliferation index of 60% [Figure 2]. Based on these histomorphological and immunohistochemical findings, a diagnosis of secretory adenocarcinoma of the endometrium was made. The patient underwent total hysterectomy with bilateral salpingo-oophorectomy.

Grossly, the resected specimen consisted of a globoid uterus with the cervix and bilateral adnexae. The uterus along with the cervix measured 7 cm × 6 cm × 5 cm in size. On cut section, the uterus showed no polyoidal lesions of the endometrium or any apparent invasion of the myometrium. There was a generalized thickening of the endometrium and the myometrium measured 3 cm in thickness. The cervix, bilateral tubes, and ovaries were unremarkable. Microscopic sections from the uterus revealed a tumor of 0.4 cm in size at its greatest dimensions which had approximately 3 mm downward invasion into the myometrium. The morphology of the tumor as well as IHC and PAS stain positivity was identical to that of the curettage specimen examined earlier. A final diagnosis of secretory adenocarcinoma of the endometrium (Grade 1 and Stage 1A) was rendered. The postoperative period of the patient was uneventful. She is under regular monthly follow-up and has no fresh complaints.

**DISCUSSION**

Tamoxifen, a non-steroidal triphenylethyl compound, belongs to a class of drugs that are known as selective ER modulators. It has both the estrogenic and antiestrogenic effects depending on the difference in the ER expression of the target tissue, differential ER conformation on ligand binding as well as varied expression, and binding of coregulator proteins to the ER.[4] Due to its estrogen antagonistic properties, it
has emerged as the first antineoplastic agent that has been extensively used for ER-positive breast cancers. It has not only significantly reduced the recurrences from the early stages of breast cancer but also has decreased the chances of contralateral breast cancer development.[5] Nevertheless, it also has a strong estrogenic agonistic effect on the vaginal epithelium, the uterine myometrium, and the endometrium of the postmenopausal women as well as the liver and the bones, which cannot be underestimated as this action of tamoxifen can potentially be detrimental as it may increase the risk of adenomyosis, adenomyomatous polyps, leiomyoma, cervical polyps, ovarian cysts, benign cystic hyperplasia of the endometrial stroma, endometrial polyps, and endometrial cancers.[2,6,7]

Tamoxifen-associated endometrial carcinogenesis has been observed in approximately 3.05/1000 women per year.[8] The risk of endometrial cancer increases two- to three-fold after a tamoxifen exposure of up to 5 years, and a median tamoxifen cumulative dose of 29 g has been suggested as a threshold for its development. However, the presence of other known risk factors of endometrial cancer such as obesity, previous estrogen use, and hypertension has also been documented to increase the chances of endometrial carcinoma in women taking tamoxifen.[9] The mechanisms involved in such tumorigenesis are still not well understood. Nevertheless, various molecular genetic alterations involving the cellular oncogenes such as mutations in Phosphatase and tensin homolog (PTEN), K-ras, and the presence of microsatellite instability have been implicated to play a crucial role in its pathogenesis.[5,10] However, these mutations are segregated on the basis of histological subtypes rather than tamoxifen exposure, indicating that tamoxifen may only act as a tumor initiator through estrogen agonistic activity on the endometrium.[11]

Secretary adenocarcinoma of the endometrium is a very rare subtype of well-differentiated endometrioid adenocarcinoma, which has been seldom associated with tamoxifen therapy. According to the pertinent world literature, till now, only two previous case reports have highlighted this rare association.[10,12] In the current case, the use of tamoxifen since last 8 years in addition to hypertension might have triggered the initiation of this tumor. Another important finding was that this rare tumor was ER, PR, and p53 positive, which was unlike the case reported by Wu CJ et al.[10] Researchers have documented that tamoxifen can cause an increase in PR as well as sex hormone-binding globulin and a significant decrease in ER which can, in turn, enhance the progestational changes of this tumor.[13] Therefore, the mechanisms by which tamoxifen promotes endometrial carcinogenesis are still controversial and need exploration as it can also act independent of its action as an estrogen agonist. Nevertheless, although the existing data on tamoxifen-associated secretory endometrial carcinoma is limited, this tumor has an excellent prognosis irrespective of the tamoxifen therapy status of the patient so far. However, it needs to be differentiated histologically from other endometrial pathologies such as clear cell carcinoma, atypical hyperplasia with secretory change, endometrioid adenocarcinoma with squamous differentiation, lipid-rich endometrioid carcinoma, low-grade mucinous carcinoma, metastatic renal cell carcinoma, Arias–Stella reaction, and clear cell change of pregnancy.[3]

**Conclusion**

This case is a relevant valuable addition to the spectrum of tamoxifen-associated endometrial carcinomas and also underscores that the prolonged utilization of tamoxifen after breast cancer should be reconsidered in all the premenopausal and postmenopausal users. It is necessary to evaluate the pathological changes on the female genital tract, especially endometrium, both before and during the period of administration of this drug. In such circumstances, vaginal and endometrial cytology, TVS, fractional curettage, and frequent gynecological follow-up can aid in preventing its deleterious effects. However, more insight is required in understanding the genesis behind the endometrial carcinomas associated with tamoxifen therapy, both for the early diagnosis of the disease and also for developing better alternative treatment methods so as to improve the survival rate of such patients in near future.

**Declaration of patient consent**

The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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**Conflicts of interest**

There are no conflicts of interest.

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