Factors associated with Synchronous Endometrial and Ovarian Cancer, Review of a Case

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Abstract

Background: Synchronous endometrial and ovarian cancer (SEOC) account from 50% to 70% of all synchronous female genital tract malignancies. The incidence of SEOC is among 5% of women with endometrial cancer (EC) and 10% with ovarian cancer (OC).

SEOC are more frequent in younger, obese, premenopausal and nulliparous women. It has been proved that SEOC has a better prognosis than a single-organ disease with metastasis.

Aim: Our aim is to describe clinicopathologic characteristics and factors associated with SEOC, by reviewing the recent literature published and describing a very representative case.

Keywords: Ovarian cancer, Malignancies, Endometrial cancer, Nulliparous women, Metastasis

Case Report

A 54-year-old perimenopausal woman complained of abnormal uterine bleedings and lower abdominal pain for 6 months. Personal antecedents were smoking, hypertension, chronic obstruction pulmonary disease, obstructive sleep apnea, obesity (BMI 42.7 kg/m²) and nulliparity [1].

The first exploration showed high tumor markers (Ca125 890 UI/ml, Ca19.9 960 UI/ml) and an enlarged uterus with thickened endometrium, but the gynecological examination was unreliable.

A diagnostic hysteroscopy showed an abnormal growth on the entire endometrial cavity with atypical vascularization. Biopsy was consistent with low-grade endometrial adenocarcinoma.

The pelvic magnetic resonance (MR) showed a large uterus, occupied by an endometrial carcinoma 4 mm close to the uterine serosa. There was an invasion to the cervix stromal, a left 20 cm heterogeneous and a hypovascular multiloculated adnexal mass adhered to mesosigma. No ascites or carcinomatosis were found.

A computerized tomography (CT) was performed finding a 6 cm solid left renal tumor suspicious of malignancy, and unspecific pelvic and paraaortic lymph nodes (Figure 1). Histology and radiology raised suspicions of an advanced endometrial cancer and synchronous renal tumor. After reviewing the case, we decided to perform a complete endometrial cancer staging procedure by laparoscopy, with intraoperative analysis of the adnexal mass and left nephrectomy.

In December 2015, the patient underwent surgery. The intraoperative analysis showed an endometrioid endometrial carcinoma invading less than 50% of the myometrium, and a carcinoma whose primary origin could not be confirmed as ovarian.
A laparoscopy was performed, finding a 20 cm left adnexal mass with a large uterus, but no carcinomatosis or ascites. Breathing difficulties forced us to change to laparotomy. The complete staging surgery consisted in: total abdominal hysterectomy, bilateral pelvic, pelviperitoneectomy, bilateral salpingo-oophorectomy and paraaortic lymphadenectomy to the level of the left renal vein, total sub-colonic omentectomy, appendectomy, left nephrectomy and splenectomy. Postoperative recovery was complex, with multifactorial shock, pneumonia, and sepsis.

Final histopathology results revealed a synchronous endometrioid endometrial and an ovarian cancer associated to endometriosis. A chromophobe renal carcinoma was also found. The 4 cm ovarian endometrioid adenocarcinoma presented lymphovascular space invasion, and a stage IIIA1 in the FIGO scale, because of paraaortic lymph node metastasis (a total of 2 in 40 paraaortic and none of 38 pelvic nodes were metastatic from ovarian carcinoma).

The 85 mm endometrioid endometrial adenocarcinoma was a remarkably differentiated FIGO stage IA, classified by the European Society of Medical Oncology (ESMO) as a low risk endometrial cancer.

We decided to complete the treatment with adjuvant standard chemotherapy (Carboplatin-paclitaxel six courses) followed by vaginal brachytherapy (21 Gy) that was well tolerated. The patient is currently free of disease.

Discussion

Synchronous cancers account for 0.7-1.8% of all gynecologic malignancies. Among them, synchronous ovarian and endometrial cancers are the most frequent (40-53%) [2]. Abnormal uterine bleeding is the most often reported symptom associated to EC (36%), which can explain why synchronous OC is also diagnosed at an early stage [3].

The mean age of SEOC patients is 53 years old [3]. Nearly 1 in 10 women under 50 years old that are diagnosed with EC, will have a synchronous OC [4]. Comparing the histologic subtypes, the endometrioid tends to appear in younger patients. In Sozen et al. article, 48% of woman with SEOC were premenopausal [3]. In Oranratnaphan’s et al. study, patients with synchronous cancers were younger, diagnosed at earlier stages, more frequent nulliparous and with better prognosis than the patients in the metastatic group [5].

Risk factors associated with synchronous ovarian and endometrial cancer are hyperestrogenous conditions such as: obesity, perimenopause, chronic anovulation, polycystic ovarian syndrome, estrogen-producing ovarian tumors and unopposed estrogen replacement therapy [1,4]. This suggests that a hormonal “estrogenic effect” may account for the development of these simultaneous cancers [1]. However, most studies have involved a small number of patients, therefore these predictors for tumor development and recurrence have not yet been clearly established [1].

SEOC in patients appears with different clinical characteristics compared to patients with isolated EC/OC, and it is usually classified as a single-organ advanced disease. In patients with abnormal uterine bleeding, the ovaries should be assessed by imaging test; and in patients presenting with an adnexal mass, assessment of the endometrium is essential [6].

The detection of an endometrial and ovarian synchronous tumor is very important, to perform an accurate complete staging surgery, that could avoid reintervention and adjuvant treatment.

In this particular case, we had the suspicion of an ovarian cancer, therefore a pelvic and paraaortic lymphadenectomy was performed, in order to give the patient a proper adjuvant therapy. In addition to this, our patient had multiple risk factors, so she was not eligible for a new surgery after histologic confirmation. In case of suspicion or doubt, an intraoperative study of the tumor specimen is recommended. On the other hand, in young patients those want to preserve fertility, an appropriate treatment plan before surgery will be beneficial [6]. Lymphadenectomy should be performed in every patient with SEOC [7].

As determined in our case, in SEOC, ovarian tumor is usually unilateral, lymphovascular space invasion is related to the ovarian carcinoma and lower Ca125 level at diagnosis [7]. Consistent pathologic criteria are crucial to distinguishing between synchronous and metastatic disease [4] Scully et al. described a list of clinicopathologic postoperatively features used to differentiate endometrial disease with metastasis to the ovary, from ovarian disease with metastasis to the endometrium; updated by Ulbright and Roth (Table 1) [3,8]. Histopathological diagnosis of SEOC still lacks a clear definition; immunohistochemistry and molecular analysis are very useful to a more accurate diagnosis [6,9].

**Table 1** The clinicopathological differential diagnostic criteria of SEOC described by Scully, et al.

| No. | Primary Endometrial Cancer with ovarian metastases (ECOM) | Primary Ovarian Cancer with endometrial metastases (OCEM) | Independent Endometrial and Ovarian tumors (SEOC) |
|-----|-----------------------------------------------------------|----------------------------------------------------------|--------------------------------------------------|
| 1   | Histologic similarity of the tumors                       | Histologic similarity of the tumors                      | Histologic dis-similarity of the tumors           |
| 2   | Large endometrial and small ovarian tumors                | Large ovarian and small endometrial tumors               |                                                  |
| 3   | Atypical endometrial hyperlasia additionally present     | No atypical endometrial hyperlasia                       | Atypical endometrial hyperlasia                  |
| 4   | Deep myometrial invasion.                                 | Location in ovarian parenchyma.                         | No or only superficial myometrial invasion of endometrial tumor. |
Endometrioid cancer is an uncommon histological type of ovarian cancer and it is thought to be developed under the influence of endometriosis and a hyperestrogenic environment. However, further studies should be developed to confirm the hypothesis.

In Oranratanaphan et al., the possibility of tumor heterogeneity must be taken into account in the evaluation of ploidy findings. Aneuploidy with similar DNA indices or diploidy of both tumors.*

The most common histological type of both endometrial and ovarian cancers is endometrioid carcinoma [2]. Endometrioid cancer is an uncommon histological type of ovarian cancer and it is thought to be developed under the same conditions as endometrial cancer. The exact etiology of synchronous primary cancers is unknown, [5] but endometriosis and a hyperestrogenic environment may be involved [3,5]. However, further studies should be developed in order to confirm that hypothesis.

SEOC are usually diagnosed at an earlier stage (48% FIGO stage I for both EC and OC) and lower histological grade [2,6]. Similar molecular genetic karyotypic abnormalities in both tumors. Metastases; OCEM: Ovarian Cancer and Endometrial Metastases; SEOC: Synchronous Endometrial and Ovarian Cancer. * The possibility of tumor heterogeneity must be taken into account in the evaluation of ploidy findings.

The prognosis of patients with SEOC with different tumor FIGO stages, like our case, seems to be mainly dependent on the prognosis of the advanced cancer [9]. General prognostic factors related were: young age, premenopausal status, no obesity, low FIGO stage, low-grade and low-risk histology, low level of cancer antigen Ca125 at diagnosis, complete staging or debulking surgery and good performance status [9]. For Bese et al., other prognostic factors are: performing lymphadenectomy, an absence of omental metastasis, and no residual disease [7].

High level of Ca125 at diagnoses was an independent risk factor for recurrence [3]. In a multicenter study by Song et al., pre-treatment serum Ca125 and tumor stage of the ovary were found to have prognostic significance on PFS and OS [1] Sozen H et al., concluded that synchronous endometrioid type, in primary EC and OC, had different clinical histopathological characteristics and favorable prognosis, compared to the other histologic types of these tumors [3]. Oranratanaphan et al., estimated that 5-year PFS was 64% and OS was 92.8%, compared to 41% and 48.5% in EC with ovarian metastasis [5].

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Treatment recommendations should take into account the risk of relapse of the respective tumor compounds. Therapy usually depends on the stage and grade of ovarian cancer, because this neoplasm is characterized by a worse prognosis and greater risk of recurrence [9]. The main adjuvant treatment is chemotherapy, followed by radiotherapy, but is more controversial if indicated [2]. In our case only chemotherapy was used, because radiotherapy requirements were not meet.

Although we only report about one single case with a short follow-up and in retrospect, it is representative for the subject of our analysis. It would be appropriate to conduct a comprehensive analysis of cases, or a multicenter study with longer follow-up. In order to determine the prognosis of adjuvant treatment by stage, a comparison between SEOC, primary EC with ovarian metastases (ECOM) and primary OC with endometrial metastases (OCEM) should be conducted.

We conclude that in the case of a suspected synchronous tumor, Scully et al. clinicopathologic criteria [10] and intraoperative assessment are recommended. More aggressive therapeutic approaches should be considered, for patients with SEOC, in order to perform a complete surgery and adjuvant therapy. More prospective studies would be required.

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| Direct extension into the adnexa | Direct extension from ovary into outer wall uterus | No vascular space invasion, surface implants or predominant hilar location in parenchyma. |
|---------------------------------|-----------------------------------------------|-----------------------------------------------------------------------------------|
| Vascular space invasion in myometrium, surface implants or combination in ovary. | Ovarian tumor located in parenchyma. | Ovarian tumor located in parenchyma. |

| 5 Spread elsewhere in typical pattern of endometrial carcinoma. | Spread elsewhere in typical pattern of ovarian carcinoma. | Absence of other evidence of spread of ovarian tumor. |
|------------------|------------------|------------------|
| 6 Ovarian tumor unilateral (80-90%) of cases) in a single mass. | Ovarian tumor unilateral (80-90%) of cases). | Ovarian tumor unilateral (80-90%) of cases). |
| 7 Ovarian endometriosis absent. | Ovarian endometriosis absent. | Ovarian endometriosis absent. |
| 8 Aneuploidy with similar DNA indices or diploidy of both tumors*. | Aneuploidy with similar DNA indices or diploidy of both tumors*. | Different ploidy of DNA indices, if aneuploidy, of the tumors*. |
| 9 Similar molecular genetic karyotypic abnormalities in both tumors. | Similar molecular genetic karyotypic abnormalities in both tumors. | Dissimilar molecular genetic or karyotypic abnormalities in both tumors. |
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