Radical Total Pelvic Exenteration with Concomitant Right Nephrectomy in the Management of Recurrent Endometrioid Ovarian Adenocarcinoma: A Case Report and Literature Review

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Abstract

Ovarian endometrioid carcinomas account for the second largest group of epithelial ovarian cancers and have lower recurrence rates than the more common serous carcinoma. We present the rare case of a woman in her late 40's with a pelvic recurrence of grade 1 endometrioid carcinoma. The patient had previously undergone a total abdominal hysterectomy, bilateral oophorectomy, and omentectomy, as well as 5 cycles of chemotherapy. The right kidney was atrophic and had minimal function secondary to chronic ureteric obstruction by the mass. This recurrence was treated surgically. Total pelvic exenteration was required as the tumour was matted and inseparable from the urinary bladder and rectosigmoid colon. A right nephrectomy was performed as well to remove the atrophic right kidney. The patient made a full recovery following surgery. This case highlights the potential extent of optimal cytoreductive surgery in an uncommon presentation of grade 1 endometrioid carcinoma recurrence.

Keywords
Surgery, Ovarian Epithelial Cancer, Neoplasm Recurrence

Synopsis

We depict a rare presentation of pelvic recurrence of grade 1 endometrioid carcinoma requiring a radical total pelvic exenteration with a concomitant right nephrectomy.

Background

Ovarian cancer is a heterogeneous malignancy with multiple histological subtypes. The most common form of ovarian cancer is epithelial ovarian cancer, of which the serous subtype is most common. The second most common, accounting for approximately 10% of epithelial ovarian cancers, is endometrioid carcinoma [1]. As a result of the rare nature of ovarian endometrioid carcinoma, research and understanding of this subtype, its presentation, management, and prognosis, are limited. Cytoreductive surgery remains the mainstay treatment for this subtype of ovarian cancer, and the extent of tumour resection has been suggested to be a major prognostic factor [2-4].

Here we present the case of a woman in her late 40's with pelvic recurrence of endometrioid ovarian cancer. The patient has consented for this case report to be written and published. The initial histopathology report graded the tumour as grade 1, without any evidence of clear cells, an indicator of aggressiveness. The cancer was International Federation of Gynecology and Obstetrics (FIGO) stage IC2, restricted to the ovaries and on a background of atypical endometriosis. This case highlights the surgical complexity of optimal cytoreductive surgery in recurrent endometrioid ovarian cancer, best achieved by a multidisciplinary approach.

Case Presentation

This woman in her late 40's, with a parity of 1, had a medical history of asthma and a BMI of 37. She was referred from the cancer unit to our cancer centre with a suspected recurrence of ovarian cancer. The patient had a history of grade 1 endometrioid ovarian cancer, FIGO Stage 1C2, treated 4 years prior to the referral in a cancer unit with surgery (total abdominal hysterectomy, bilateral salpingo-oopherectomy, omentectomy, and peritoneal washings) followed by five cycles of chemotherapy (Carboplatin / Paclitaxel). A sixth cycle of chemotherapy was not administered due to toxicity. The patient was followed up with 4 monthly appointments in the first two years post surgery, then 6 monthly appointments. At a follow-up appointment in the third year post surgery, the patient reported mild abdominal discomfort and intermittent lower back pain and had a CA 125 of 42 ku/L. A CT AP was requested and this identified a complex pelvic mass, of approx. 7cm in max. diameter. The imaging was discussed at the Oxford multidisciplinary meeting (MDT) and a decision was made for a diagnostic laparoscopy. A tissue biopsy was obtained and was in keeping with an atypical endometrioma. The patient had GnRH analogues for 6 months followed and serial imaging.

Investigations
A CT abdomen and pelvis identified a 12cm pelvic mass with thickening of the wall in keeping with malignant change. The suspicion of malignancy was confirmed by magnetic resonance imaging (MRI) and positron emission tomography (PET CT) scans which showed a part-cystic part-solid mass with restricted diffusion and high FDG avidity, respectively. The pre-operative MRI pelvis also described adhesions between the pelvic mass and the bowel. Imaging did not suggest any lymph node involvement or distant metastatic disease.

The right ureter was noted to be obstructed by the mass, leading to hydronephrosis and marked right kidney atrophy. A dimercaptosuccinic acid (DMSA) scan revealed minimal right kidney function.

The case was discussed at the Oxford network multidisciplinary team meeting (MDT) in which a two-stage surgical procedure was supported involving firstly an exploratory laparoscopy to assess resectability of disease, followed by ovarian debulking surgery if suitable.

**Differential Diagnosis**

Endometrioid ovarian cancer may arise from an atypical endometrioma. Distinguishing these two pathologies on imaging may sometimes be challenging due to a number of factors: the ovarian parenchyma adjacent to an endometriotic cyst can be mistaken for an enhancing solid malignant nodule, deep invasive endometriosis of the recto-sigmoid colon can mimic the “mushroom cap” sign associated with invasive malignancy (30)

**Treatment**

The patient was counselled extensively pre-operatively, over multiple consultations with the Gynaecology Oncology, Urology and Colorectal teams. The Cancer Specialist Nurse (CNS) played a central role in supporting her through this process. The Stoma Nurses were also involved as part of stoma planning. The risks and benefits of this surgery, the extent of the surgery, and the option of doing nothing were explored. Early and late surgical complications were discussed, and realistic expectations were set regarding the recovery period, including the need for permanent stomas. The patient was motivated to pursue surgical management. She had a full Anaesthetic work-up prior to surgery. Her diet, level of activity and psychological status were assessed as part of the prehabilitation programme. She was encouraged to maintain a healthy and active lifestyle before the operation.

The final surgical plan was agreed at a Surgical MDT with the Gynaecology Oncology, Urology and Colorectal teams.

**Surgical Procedure:**

**Preparation and Initial Steps**

The surgery was conducted as a joint procedure between Gynaecology Oncology, Urology and Colorectal teams. The patient was first cleaned, draped, catheterised, and positioned with her legs in a modified Llyod-Davies position with flow-trons on. A midline xiphio-pubic laparotomy was performed and the abdomen was opened in layers with a handheld diathermy and Lahey dissecting forceps. The ascending colon and the liver were then mobilised to gain access to the right kidney for the nephrectomy. (Figure 1, 2, 3).

**Right kidney mobilisation**

The right nephrectomy was conducted first as the right kidney had been chronically obstructed by the tumour and appeared atrophic. The ureter was slung, and lower pole, upper pole and lateral attachments were dissected. The renal artery, vein and gonadal vein were then all identified and transected, enabling mobilisation of the right kidney. The ureter was followed to the level of the mass.

**Assessment of mass**

A 15x15cm pelvic mass was identified in the Pouch of Douglas. The mass was seen to be inseparable from the urinary bladder and recto-sigmoid colon and was in close proximity to the left vesico-ureteric junction. The right ureter was encased in the mass. As the tumour was friable, a fragment detached when mobilisation
was attempted. This tissue was sent for frozen section. The frozen section report suggested the tumour was a grade 1 adenocarcinoma, either endometrioid or endometrioid with mucinous component. Pelvic side walls were opened bilaterally, and para-rectal and Latzko spaces developed. Subsequent discussion among the four consultant surgeons present finally led to the conclusion that a total exenteration was required to achieve complete tumour clearance.

Pelvic exenteration

The pelvic exenteration began with the division of the sigmoid colon above the tumour and dissection of the total mesorectal excision (TME) plane to the pelvic floor. A radical cystectomy was performed, and the bladder was mobilised. Subsequent vaginal division and low rectal division then allowed for en-bloc removal of the entire specimen consisting of right kidney, right ureter, urinary bladder, pelvic mass, and recto-sigmoid colon and vaginal cuff. (Figure 4).

Colonic conduit and end colostomy

Following the pelvic exenteration, a colonic conduit and end colostomy were required. First, the splenic flexure was mobilised. The proximal descending colon was then divided, and the distal section used to create the colonic conduit with the left ureter (Bricker procedure). The colonic conduit and proximal segment of the descending colon were matured through the left abdomen and two stomas produced in a vertical orientation.

Closure

Before closure, bulky aorto-caval lymph nodes that had been noticed earlier were removed. The pelvis was washed out and a Jackson-Pratt drain inserted. The abdomen was then closed in layers. The total blood loss was 1500mls.

Outcome and Follow-Up

Due to the complex and extensive nature of the surgery, the patient was admitted to the Intensive Care Unit post-operatively and stayed as an inpatient for the next 11 days before discharge. There were no significant complications in this period. The patient was readmitted four weeks post surgery with superficial wound dehiscence and urosepsis, which were managed with intravenous antibiotics (grade 1 Clavier Dindo complications). She recovered well following this admission. The final histopathology report of the removed section revealed the tumour was matted to the bladder and the rectum, but there was no evidence of invasion of the wall of these structures. The en-bloc specimen had clear margins. The cavo-aortic lymph nodes removed were normal with no evidence of metastatic carcinoma. These findings were discussed in MDT, along with immunohistochemistry results, and the ultimate decision was that adjuvant chemotherapy or radiotherapy was not needed. This decision was subsequently conveyed to the patient who had made a full recovery post-surgery. A Benchmark imaging (CT chest, abdomen, pelvis) and MRI pelvis were performed 4 months following the total exenteration (Sep 2023) and were unremarkable.

Discussion

Endometrioid carcinomas are the second most common epithelial ovarian cancer. In contrast to the more common serous carcinomas, endometrioid carcinomas present earlier, at a younger age and have better long-term outcomes [5-7]. Several studies have suggested that endometrioid carcinomas have higher rates of both 5-year overall survival (80.6%) and progression-free survival (68%) compared to other ovarian cancer subtype. Additionally, they have a lower recurrence rate, especially at lower grades [1, 6]. Relapse patterns appear to further differentiate endometrioid carcinomas from serous carcinomas, with endometrioid carcinomas having a much higher proportion, at nearly 50%, recurring solely in the pelvis, while serous carcinoma relapse tends to be much more diffuse [6]. Here we discuss an interesting, rare presentation of a pelvic recurrence of low grade endometrioid carcinoma.

Endometriosis is a risk factor for the development of ovarian cancer, mostly specific subtypes including endometrioid and clear cell carcinomas. Up to 42% of endometrioid adenocarcinomas are associated with endometriosis. Patients with endometrioid ovarian adenocarcinoma arising from endometriosis tend to
present at a younger age, lower stage, and lower grade than those without associated endometriosis (8). The transformation from benign to atypical endometriosis to endometriosis-associated ovarian cancer involves a combination of: oxidative stress, inflammation, molecular genomic alterations, hyperoestrogenism (32). Some of the molecular abnormalities encountered in endometriosis-associated ovarian cancer include: the activation of oncogenic KRAS and PI3 K pathways and the inactivation of tumour suppressor genes PTEN and ARID1A (3). Some of the key mutations involved in the malignant transformation and progression, including AT-Rich Interaction Domain IA (ARID1A) mutations, may have potential to be effective chemotherapy targets [8, 9].

In a patient with endogenic hyperoestrogenism related to obesity, a new pelvic mass diagnosed three years from the initial surgery may represent as, rather than a recurrence, a de novo lesion progressing from benign endometriosis to atypical endometriosis and then endometrioid ovarian adenocarcinoma.

Other important pathologies have also been noted to be associated with endometrioid carcinoma that should be understood and considered, including endometrial cancer. The rare presentation of primary tumours in both the endometrium and ovary in synchronous endometrial and ovarian carcinoma, is recognised to be a separate entity to either pathology with different prognoses and treatment implications [10-12]. These tumours were previously thought to be synchronous independent tumours, however molecular analysis has established that they have a common clonal origin [13]. The 2023 FIGO staging of endometrial cancer, which incorporates molecular findings, classifies these tumours as stage IA3 when certain criteria are met: unilateral disease, no capsular spread, less than 50% myometrial invasion, absence of substantial/extensive lympho-vascular space invasion (LVSI). These tumours have a better prognosis and do not require adjuvant chemotherapy [14].

The Carboplatin/Paclitaxel regimen as adjuvant treatment has not been proven to result in survival benefit for low-grade endometrioid ovarian cancer [15]. Current National Comprehensive Cancer Network (NCCN) guidelines [16] for grade 1 endometrioid ovarian carcinoma are Carboplatin/Paclitaxel or hormonal treatment, such as aromatase inhibitors, leuprolide acetate, tamoxifen. Novel, biomarker-driven therapies, are currently being investigated for this histological subtype: Bouquet (NCT04931342 GOG-3051) is a multicentre clinical trial which is currently recruiting patients with persistent or recurrent low-grade endometrioid ovarian cancer and other rare ovarian tumours that are not amenable to curative surgery.

Until new treatments options are identified, surgery with maximal cytoreductive effort remains the mainstay treatment for this histological subtype of ovarian cancer [17-25]. Achieving resection of all macroscopic disease (R0) is the single independent factor for survival. In their meta-analysis including 6885 patients with advanced ovarian cancer, Bristow et al (2023) have shown there is a 5.5% increase in median survival time with each 10% increase in maximal cytoreduction [2].

In the case we presented, a radical total exenteration was required to achieve R0. A right nephrectomy was performed to remove the right kidney, which was non-functioning. The Bricker technique for urinary diversion involved spatulation of the remaining ureter and anastomosis to a segment of descending colon used as a conduit. Total cystectomy with urinary diversion is a standard procedure in the context of anterior or total exenterations. The use of a segment of descending colon should be considered in patients undergoing end colostomy, to avoid the need for primary small bowel anastomosis.

In conclusion, ovarian endometrioid carcinoma of the ovary is an uncommon histological subtype of ovarian carcinoma for which the mainstay of treatment is surgery. Cytoreductive effort should be maximized to achieve R0, especially in young patients. The complexity of the operation should not be a deterrent factor if surgery is carried out in a multidisciplinary environment with robust clinical governance.

**Learning Points**

- Endometrioid ovarian carcinoma can arise in a background of endometriosis.
- Carboplatin/Paclitaxel adjuvant chemotherapy has limited survival benefit in low-grade endometrioid
ovarian adenocarcinoma, a subtype of ovarian cancer for which novel biomarker therapies are still being developed.

- Maximal cytoreductive effort with the aim to achieve R0 is key to the management of recurrent low-grade endometrioid ovarian cancer.
- Complex operations with multidisciplinary involvement may be required in order to remove all macroscopic disease.

**Author Contributions**

(I) Conception and design: Hooman Soleymani Majd, Raghavskandhan Ramachandran

(II) Administrative support: Hooman Soleymani Majd

(III) Provision of study materials or patients: Hooman Soleymani Majd

(IV) Manuscript writing: All authors

(V) Final approval of manuscript: All authors

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**Conflict of Interest**

The authors have no conflicts of interest

**References**


Figures

Figure 1: En bloc specimen containing right kidney, right ureter, recto-sigmoid colon, pelvic tumour, urinary bladder, and vaginal cuff.
Figure 2: Sling guided below abdominal aorta after bowel mobilisation using Cattell-Braasch and Mattox manoeuvres.

Figure 3: Abdominal aorta (red sling) and IVC.

Figure 4: En bloc specimen after right nephrectomy and mobilisation of right ureter; Mattox manoeuvre
Cattell-Braasch maneuver

Sling guided below abdominal aorta at L2-L3 level, using O'Shaughnessy forceps

Mattax manoeuvre