Long-term survival of patients with mismatch repair-deficient, high-stage ovarian clear cell carcinoma

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ABSTRACT

Aims: Gynaecological cancer patients with germline mutations appear to have a better prognosis than those with sporadic malignancies. Following the observation of long-term survival in a patient with stage III ovarian clear cell carcinoma (CCC) and possible Lynch syndrome (LS), DNA mismatch repair (MMR) protein immunohistochemistry was performed in a series of high-stage CCC and correlated with patient outcomes.

Methods and Results: Thirty-two consecutive cases of stage III/IV ovarian CCCs accessioned between 1992 and 2015 were examined. The tumours from two patients (6%), including the index case, showed loss of MSH2/MSH6 expression while MLH1/PMS2 staining was retained. The index patient subsequently developed colonic and rectal carcinomas that were also MSH2/MSH6 deficient while the second patient had a genetically confirmed germline MSH2 mutation. All other tumours showed retained expression of the four MMR proteins. The two patients with MMR protein-deficient tumours were alive 160 months and 124 months following surgery whereas the median survival of patients with MMR protein-intact CCCs was 11.8 months (75th and 25th percentiles of 8.1 months and 39.3 months, respectively), with 21 patients deceased due to tumour.

Conclusions: Larger studies are required but high-stage, MMR protein-deficient CCCs may have a relatively favourable prognosis.

INTRODUCTION

Five major subtypes of tubo-ovarian adenocarcinoma are recognised comprising high-grade serous carcinoma, low-grade serous carcinoma, mucinous carcinoma, endometrioid carcinoma (EC) and clear cell carcinoma (CCC). The latter two tumours each account for approximately 10% of ovarian cancers in Western countries but CCC is relatively common in the Japanese population accounting for up to one quarter of cases (1). Ovarian CCC is often associated with endometriosis and demonstrates distinctive molecular alterations including frequent ARID1A and PIK3CA mutations, findings that may become important as specific targeted therapies are developed (1,2).
In recent years it has been appreciated that patients with Lynch syndrome (LS), due to germline mutations in genes encoding for the DNA mismatch repair (MMR) proteins MLH1, PMS2, MSH2 or MSH6, have an increased risk (approximately 10%) of developing ovarian neoplasia and the majority of these tumours are ECs or CCCs (3). Therefore reflex immunohistochemistry for MMR proteins has been proposed in these cases (4-6).

Traditionally, ovarian CCC has been considered a clinically aggressive malignancy and therefore regarded as high-grade for management purposes. However, the prognosis of surgically staged CCC confined to the ovary is relatively favourable with 5-year survival rates of 80-90%, although the prognosis of stage IC tumours with ovarian capsular involvement and/or positive peritoneal fluid cytology is more guarded (7-9). High-stage CCCs with lymph node, omental, peritoneal and/or distant metastases have been reported to have a very poor prognosis with inferior survival rates compared to equivalent stage high-grade serous carcinomas (1,8,10). This may partly reflect the relative insensitivity of CCC to standard chemotherapy.

In routine practice we identified a possible LS patient with stage III ovarian CCC who had surprisingly long disease-free survival leading us to speculate whether LS-associated CCC might have a better prognosis than sporadic CCC, analogous to findings in colorectal carcinoma where tumours demonstrating MMR protein deficiency and/or microsatellite instability (MSI) have relatively favourable outcomes (11,12). Therefore in the present study we reviewed all high-stage (≥stage III) CCCs presenting in our department over a 24-year period (1992-2015), correlating the findings with MMR protein expression and clinical outcomes. The study was performed according to the Declaration of Helsinki and received institutional ethics approval (KEMH reference 11014, 1/2/2016).

CASES AND RESULTS
A total of 32 patients with high-stage ovarian CCC were identified in the study period. Twenty-nine patients had stage III carcinomas based upon the presence of omental, pelvic and/or para-aortic lymph node metastases while 3 patients had stage IV disease based upon cytologically confirmed malignant pleural effusions (2 cases) or cervical lymph node metastases (1 case). All stage IV cases also had omental and/or abdominal lymph node involvement. Immunohistochemistry was performed on representative full face sections of each tumour as previously described with

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appropriate internal positive control staining in all cases (13). Two CCCs (6.3%), summarised below, demonstrated loss of MSH2/MSH6 staining with preserved MLH1/PMS2 expression (Figure 1). All other tumours showed retained expression of all four MMR proteins.

Mismatch repair protein-deficient CCCs (n=2)

Case 1 (index case)
A 41-year-old underwent simple hysterectomy for menorrhagia. The uterus showed benign appearances but a nodule was noted in the pouch of Douglas intraoperatively and biopsy revealed CCC. She then underwent bilateral salpingo-oophorectomy, omentectomy and bilateral pelvic lymph node dissection. Both ovaries revealed clear cell carcinoma, the larger right ovarian tumour being up to 25mm diameter. The tumour had solid and glandular growth patterns and there was a prominent peritumoural lymphoplasmacytic infiltrate (Figure 1). Endometriosis was not identified. Metastatic CCC was present in 1/5 left external iliac lymph nodes but there was no evidence of metastasis in the omentum or in 87 additional pelvic nodes, and peritoneal fluid cytology was also negative. The patient received adjuvant pelvic radiotherapy. Thirteen years later she underwent surgery for concurrent mucinous carcinoma of the sigmoid colon and signet ring cell carcinoma of the rectum. Both tumours were confined to the bowel wall (pT2) but metastatic carcinoma (of probable sigmoid origin) was present in 1/16 pericolic nodes. Immunohistochemistry showed loss of MSH2/MSH6 expression in both tumours, similar to the ovarian CCC. The patient had no evidence of tumour recurrence at last known follow-up 160 months after her initial gynaecological surgery.

Case 2
A 45-year-old presented with a pelvic mass and was found at laparotomy to have bilateral, 6-7cm ovarian tumours and multiple omental tumour deposits measuring up to 15mm. Histological examination demonstrated bilateral ovarian CCC showing a predominantly solid growth pattern with confirmed omental metastases. There was a relatively mild and focal tumour-associated lymphoplasmacytic infiltrate more prominent in the omentum. Endometriosis was not identified. The uterus showed benign findings and peritoneal fluid cytology was negative. Lymphadenectomy was not performed. Post-operatively the patient received standard platinum-based adjuvant

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chemotherapy. She had a family history of colorectal and ovarian cancer and had undergone multiple screening colonoscopy examinations beginning 5 years prior to her presentation with ovarian neoplasia. On most occasions colonoscopy revealed 1 or 2 polyps which were variably of adenomatous or hyperplastic type but there was no invasive colonic malignancy. Subsequently genetic analysis confirmed LS due to a germline MSH2 mutation. There was no evidence of tumour recurrence at last known follow-up 124 months following her gynaecological surgery.

Mismatch repair protein-intact CCCs (n=30)
The median age of the patients was 52 years (range 37-73 years). Nineteen patients (63%) had histologically documented endometriosis either in the ovary affected by tumour or in other pelvic sites. Omental sampling was performed in all patients and was positive for CCC in 22 cases (73%) while peritoneal fluid cytology was positive in 21/25 (84%) patients. Pelvic and/or paraaortic nodal metastases were present in 13/15 patients (87%) who underwent lymphadenectomy. Follow-up data, available in 29 cases, are summarised in Figure 2. The overall median follow-up interval was 9.6 months (range 0.8-119), with a median of 14.1 months (range 2.5-54.9) for those patients who were disease-free clinically at last follow-up. The median survival was 11.8 months with the 75th and 25th percentiles being 8.1 and 39.3 months, respectively.

DISCUSSION
Lynch syndrome-associated and MMR protein-deficient ovarian carcinomas are most commonly of clear cell or endometrioid subtype (5,6,14), although, conversely, only 2-14% of these tumours demonstrate MMR protein deficiency and/or microsatellite instability (MSI) (15-18). The current study, albeit restricted to high-stage CCC specifically, supported these findings since MMR protein-deficiency was identified in only 2/32 tumours (6.3%). Interestingly, both of these tumours were bilateral, a generally uncommon finding in ovarian CCC although there are limited data on high-stage cases (19). Genetic analysis confirmed the diagnosis of LS in one case but this was not performed in the second patient and therefore it is uncertain whether she had LS or a ‘Lynch-like’ syndrome which may also be associated with MMR protein-deficient neoplasia (20,21).
The prognosis of high-stage ovarian CCC is considered very poor and this is illustrated by the present series where 22 of 31 patients (71%) with follow-up information died of metastases with a median survival of less than 1 year. However, surprisingly long-term survival of at least 124 and 160 months was recorded in the two stage III patients with possible or confirmed LS. Similarly, a LS patient with stage III CCC in the series of Niskakoski et al survived at least 156 months (14). Recently Bennett and colleagues reported MMR protein deficiency in 6/109 unselected ovarian CCC and two of these patients, both with likely LS, had stage III disease (17); one patient (who also had endometrial carcinoma) died of tumour 14 months after diagnosis but the other was disease free at 93 months. These findings, albeit based upon a small number of cases, raise the possibility that high-stage LS-associated and/or MMR protein-deficient ovarian CCCs have a better prognosis than similar stage sporadic tumours. In this regard it is noteworthy that in a multicentre European study of LS-associated ovarian tumours the overall 5-year survival for stage III/IV cases was 59% compared with an expected survival of approximately 30-35% in high-stage ovarian carcinoma generally (22). However, this study had limited histological correlation and also included non-epithelial tumours, and therefore correlation with a CCC phenotype is not possible.

Treatment details were not available in most patients in this series and therefore we were unable to determine whether prolonged survival in the two patients with MMR protein-deficient CCCs was associated with a more favourable response to adjuvant therapy. We note that Case 2 was unlikely to have been cured surgically due to the extensive nature of her disease at presentation. Interestingly, a recent study of endometrial carcinomas found that patients with probable LS and high-stage disease who received adjuvant therapy showed better outcomes (23). Improved survival of LS patients may also apply to those with low-stage CCC but this will be more difficult to determine given the generally favourable outcome in this group of patients. It has been suggested that improved survival in LS-associated carcinomas may be due to the diploid nature of these tumours, reduced tumour viability due to genomic instability, and/or the prominent immune reaction that often characterises such cases (24). In regard to the latter, it is noteworthy that the one of the CCCs presented herein was associated with a prominent lymphoplasmytic infiltrate.

In summary, we describe two patients with possible/ confirmed LS and long-term survival following the diagnosis of stage III ovarian CCC. While larger studies
are required, these findings raise the possibility that MMR protein-deficient CCCs have a relatively favourable prognosis, analogous to colorectal carcinoma.

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REFERENCES


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**FIGURE LEGENDS**

Figure 1. Case 1. A. Ovarian clear cell carcinoma. The tumour is associated with a prominent lymphoplasmacytic infiltrate. B. Higher magnification showing solid and glandular architecture. Part of a follicular cyst is present at upper right. Immunohistochemistry on corresponding sections shows retained expression of MLH1 (C) but loss of MSH2 (D) in the tumour cells (insets at high magnification).

Figure 2. Survival curve of mismatch repair protein-intact, high-stage ovarian clear cell carcinomas.
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