Microcystic, elongated and fragmented (MELF) pattern in endometrial carcinoma: clinicopathologic analysis and prognostic implications

Jinghua Song, PhDabcde, Huajun Li, PhDabcde*, Hongyan Guo, PhDabcde, Yuhan Cai, PhDabcde

Abstract

To assess the clinical value of microcystic, elongated, and fragmented (MELF) pattern in Chinese patients with endometrial endometrioid carcinoma. A total of 189 patients with endometrial endometrioid carcinoma were retrospectively analyzed in Peking University Third Hospital from January 2017 to December 2019. We analyzed the association of MELF pattern with the histopathologic data and prognosis of the patients, while immunohistochemistry was performed. The frequency of MELF pattern was 17.99% (34/189). MELF pattern was associated significantly with tumor size, myometrial invasion, histological grade, International Federation of Gynecology and Obstetrics stages, lymphovascular space invasion, and lymph node metastasis. According to multivariate logistic regression analysis, lymphovascular space invasion [95% confidence interval 1.021–48.485, P = .048] was a significant predictor of lymph node involvement. However, MELF pattern was not a significant predictor (95% confidence interval 0.054–2.279, P = .400). Loss of expression for mismatch repair proteins was observed in 10 MELF + cases (29.41%) and 54 MELF− cases (34.84%), respectively. All patients were followed up for 36.8 ± 8.9 months (18–54 months). Only 1 patient with MELF pattern was diagnosed with vaginal recurrence 28 months after the surgery. MELF pattern was associated with adverse histologic findings in endometrial endometrioid carcinomas. However, MELF pattern was statistically not a valuable predictor of lymph node metastasis and it needs more studies to show whether MELF pattern has an impact on the prognosis of patients with endometrial endometrioid carcinoma. MELF pattern may be important for identifying those patients who need comprehensive staging surgery.

Abbreviation: FIGO = International Federation of Gynecology and Obstetrics, IHC = immunohistochemistry, LNM = lymph node metastasis, LVSI = lymphovascular space invasion, MELF = microcystic, elongated, and fragmented, MMR = mismatch repair, MSI = microsatellite instability.

Keywords: lymph node metastasis, elongated, endometrial endometrioid carcinoma, "microcystic, and fragmented (MELF)” pattern, mismatch repair

1. Introduction

Endometrial carcinoma is the most frequent gynecological neoplasia in women, with an increase in incidence and mortality over the past few decades. Endometrial endometrioid carcinoma is the most common histologic subtype. Although patients with low-grade (grades 1 and 2) endometrioid carcinoma have better outcomes than those with high-grade (grade 3) endometrioid carcinoma, a subset of patients with low-grade endometrioid carcinoma have a recurrence and adverse prognosis. Therefore, there is a constant need for novel prognostic factors which may improve patient risk stratification. Among these, microcystic, elongated, and fragmented (MELF) pattern of myometrial invasion has recently been related to increased risk of lymphovascular space invasion (LVSI), lymph node metastasis (LNM) and extra-uterine disease. However, the biological and prognostic significance of MELF pattern in endometrial endometrioid carcinomas remains uncertain.

This study was conducted to elucidate clinicopathologic features and the prognostic value of MELF pattern in Chinese patients with endometrial endometrioid carcinoma. For that purpose, we retrospectively reviewed data of 189 consecutive patients with endometrial endometrioid carcinoma and analyzed the clinicopathologic and prognostic features. Furthermore, some immunohistochemical analyses were applied to elucidate the nature of tumor cells in MELF pattern.

*Correspondence: Huajun Li, Center for Reproductive Medicine, Department of Obstetrics and Gynecology, Peking University Third Hospital, Beijing 100191, China (e-mail: chnafly@1967@126.com).

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2. Materials and methods

2.1. Study population and clinical information

This study examined 189 consecutive patients with endometrial endometrioid carcinoma that were resected with total hysterectomy and bilateral salpingo-oophorectomy with or without lymphadenectomy at the Department of Obstetrics and Gynecology, Peking University Third Hospital (Beijing, China), between January 2017 and December 2019. All patients provided written informed consent according to institutional guidelines. Patients were informed that the resected specimens were going to be stored by the Pathology Unit of the Peking University Third Hospital and might potentially be used for scientific research, and that their privacy would be maintained. Clinical and demographic information was collected from patient charts, including age at surgery, adjuvant chemotherapy and/or radiation therapy, recurrence and survival status.

2.2. Histopathologic evaluation and immunohistochemistry (IHC)

In all cases, 2 experienced gynaecological pathologists established the histological diagnosis of endometrial endometrioid carcinoma after an extensive and careful evaluation of tumor specimens, according to the WHO Classification of Tumours of Female Reproductive Organs.[6] Histopathologic findings, including the extent of myometrial invasion, invasion to the uterine cervix, LVSI and LNM, were evaluated to confirm the initial diagnosis. Finally, the International Federation of Gynecology and Obstetrics (FIGO) staging system published in 2009 was applied to all patients. Patients with mixed carcinoma (endometrioid carcinoma and serous/clear cell carcinoma) were excluded because high-grade components can influence patient survival.[7]

IHC was performed with the labeled streptavidin–biotin peroxidase detection system.Mismatch repair (MMR) proteins status was determined with the antibodies (MLH1, MSH2, MSH6 and PMS2) in the setting of intact control stromal/lymphocyte staining. Cases were considered as showing stable immunophenotype (MMR+) if any tumor cell nuclei showed positive staining, and unstable immunophenotype (MMR−) if all tumor cell nuclei were negative in the presence of internal positive controls. Stromal/lymphocyte staining as well as nonneoplastic endometrial glands were used as positive internal controls. The expression profile of p53 was evaluated by estimating the proportion of nuclear staining of tumor cells. Cases in which nuclear staining was observed in at least 10% of cancer cells were classified as a p53-stained group. Cases were classified as “p53 wild type” (p53-wt: focal and/or heterogeneous staining pattern) and “p53 immunohistochemically mutated” (diffuse expression in at least 75% of tumor cell nuclei); cases showing complete absence of staining in tumoral nuclei were considered as “null phenotype”.

2.3. Assessment of MELF pattern

MELF pattern was initially recognized by Lee, Vacek and Belinson,[8] but the term MELF was first defined by Murray et al.[9] The histological appearance of invasive glands, as cystic-dilated or slit-like, lined by flattened, endothelial-like epithelium or squamoid tumor cells, with eosinophilic cytoplasm, often with intraluminal tufts or fragmented, alongside with small groups or isolated tumor cells, led to their denomination as “microcystic, elongated and fragmented glands”.

2.4. Statistical analysis

All statistical analyses were performed using SPSS 22.0 for Windows (IBM SPSS Statistics, IBM software, Armonk, NY).

Values were given as mean ± standard deviation (SD) or median (interquartile range). Continuous variables were tested for normality by the Kolmogorov–Smirnov test. The analysis of the differences between groups was assessed by the Welch t test or the Mann-Whitney-Wilcoxon test for parametric or nonparametric data respectively. The Chi-square test was applied to compare proportions of categorical variables. The multivariate logistic regression analysis was used to study the possible correlation between MELF pattern and risk of lymph node metastasis. Significance was defined as P < .05.

3. Results

3.1. Clinical characteristics and clinicopathologic parameters of endometrial endometrioid carcinoma

Representative photographs of MELF pattern in endometrial endometrioid carcinoma were presented in Fig. 1. Table 1 presents the clinical characteristics and clinicopathologic parameters with MELF pattern. The frequency of MELF pattern was 17.99% (34/189). The presence of MELF pattern was associated with tumor size (P = .003), deep myometrial invasion (P < .001), histological grade (P < .001), FIGO stage (P < .001), LVSI (P < .001), and LNM (P = .001). There was no significant difference in patient age and cervical stroma involvement between MELF+ patients and MELF− patients. The result

Figure 1. Representative images of MELF pattern in endometrial endometrioid carcinoma (A); elongated gland lined by simple squamous epithelium and columnar epithelium, with lumens containing eosinophilic tumor cells (B); microcystic gland with neutrophilic infiltration (C). HE staining: (A) × 50; (B) × 200; (C) × 400. MELF = microcystic, elongated, and fragmented.
of tumor marker CA 125 was also related to MELF pattern (P < .001). The proportion of adjuvant therapy was significantly higher in MELF pattern (P < .001). MELF pattern was present only in low-grade but not in high-grade endometrioid carcinomas.

Table 2 shows the relationship between prognostic factors and lymph node involvement. According to multivariate logistic regression analysis, L V S I [95% confidence interval 1.021–48.483, P = .048] was a significant predictor of lymph node involvement. However, MELF pattern was not a significant predictor (95% confidence interval 0.054–2.279, P = .400).

3.2. Immunohistochemical findings
Loss of expression for MMR proteins was observed in 10 MELF + cases (29.41%) and 54 MELF− cases (34.84%), respectively. The details regarding type of protein loss were shown in Table 1. However, there was no significant differences

### Table 1
Clinical characteristics and clinicopathologic parameters of all study patients.

| Parameter                              | MELF + (n = 34) | MELF− (n = 155) | χ²/t     | P     |
|----------------------------------------|-----------------|-----------------|----------|-------|
| Age (yrs)                              | 58.03 ± 7.93    | 52.71 ± 10.50   | 2.784    | .061  |
| Tumor size (cm)                        | 2.12 (2.02)     | 1.88 (1.86)     | −2.980   | .003  |
| CA 125 (U/ml)                          | 47.11 (66.60)   | 15.99 (12.94)   | −3.962   | <.001 |
| Myometrial invasion                    |                 |                 |          |       |
| <1/2                                   | 12 (35.29%)     | 136 (87.74%)    | 45.152   | <.001 |
| ≥1/2                                   | 22 (64.71%)     | 19 (12.26%)     |          |       |
| Cervical stroma involvement            |                 |                 |          |       |
| Absent                                 | 27 (79.41%)     | 139 (89.68%)    | 2.749    | .097  |
| Present                                | 7 (20.59%)      | 16 (10.32%)     |          |       |
| Lymph node metastasis*                 |                 |                 |          |       |
| Absent                                 | 24 (70.59%)     | 112 (91.80%)    | 10.707   | .001  |
| Present                                | 10 (29.41%)     | 10 (8.20%)      |          |       |
| Lymphovascular space invasion          |                 |                 |          |       |
| Absent                                 | 10 (29.41%)     | 125 (80.65%)    | 35.863   | <.001 |
| Present                                | 24 (70.59%)     | 30 (19.35%)     |          |       |
| Histological grade                     |                 |                 |          |       |
| 1                                      | 5 (14.71%)      | 69 (44.52%)     | 20.837   | <.001 |
| 2                                      | 29 (85.29%)     | 66 (42.58%)     |          |       |
| 3                                      | 0                            | 20 (12.90%)     |          |       |
| FIGO stage                             |                 |                 |          |       |
| Stage I/II                             | 21 (61.76%)     | 140 (90.32%)    | 18.019   | <.001 |
| Stage III/IV                           | 13 (38.24%)     | 15 (9.68%)      |          |       |
| Adjuvant therapy                       |                 |                 |          |       |
| No                                     | 2 (5.88%)       | 86 (55.48%)     | 27.571   | <.001 |
| Yes                                    | 32 (94.12%)     | 69 (44.52%)     |          |       |
| Immunophenotype MMR                    |                 |                 |          |       |
| Stable                                 | 24 (70.59%)     | 101 (65.16%)    | 0.367    | .545  |
| Instable                               | 10 (29.41%)     | 54 (34.84%)     |          |       |
| MLH1-PM2                              | 28 (82.35%)     | 111 (71.61%)    | 1.653    | .199  |
| Negative                               | 6 (17.65%)      | 44 (28.39%)     |          |       |
| MSH2-MSH6                             | 30 (88.24%)     | 145 (93.55%)    | 1.148    | .284  |
| Positive                               | 4 (11.76%)      | 10 (6.45%)      |          |       |
| Negative                               | 24 (70.59%)     | 141 (90.97%)    | 0.242    | .623  |
| Mutated                                | 4 (11.76%)      | 14 (9.03%)      |          |       |

FIGO = International Federation of Gynecologists and Obstetricians, MELF = microcystic, elongated and fragmented pattern, MMR = mismatch repair.

*Analysis of 156 patients with lymphadenectomy.

Values are expressed as mean ± SD or median (interquartile range) for parametric or non-parametric data respectively.

### Table 2
Results of univariate and multivariate analyses of odds ratios in the logistic regression model with lymph node metastasis as the dependent variable.

| Variables                        | Univariate analysis | Multivariate analysis |
|----------------------------------|----------------------|-----------------------|
|                                  | OR                   | 95% CI                | P        | OR           | 95% CI                | P        |
| Age (yr)                         | 0.984                | 0.923–1.119           | .579     | 1.017        | 0.894–1.183           | .738     |
| Tumor size (cm)                  | 0.746                | 0.804–2.234           | .186     | 1.340        | 0.448–2.243           | .261     |
| MELF                             | 0.351                | 0.087–2.651           | .273     | 0.480        | 0.054–2.279           | .400     |
| Myometrial invasion              | 2.916                | 0.482–17.661          | .244     | 2.904        | 0.458–18.432          | .258     |
| Lymphovascular space invasion    | 6.785                | 1.090–42.251          | .040     | 7.035        | 1.021–48.485          | .048     |
| Cervical stroma involvement      | 2.856                | 0.395–20.626          | .298     | 4.832        | 0.176–40.537          | .447     |
| Histological grade               | 1.326                | 0.060–29.068          | .858     | 0.685        | 0.722–32.317          | .768     |

CI = confidence interval, MELF = microcystic, elongated and fragmented pattern, OR = odds ratio.
between the 2 groups in the distribution of the MMR proteins alterations. Nevertheless, a statistical trend has been observed. Our data showed a higher prevalence of MSH2-MSH6 loss in MELF+ group (11.76% in MELF+ cases vs 6.45% in MELF− cases) but a higher frequency of MLH1-PMS2 loss in MELF− group (28.39% in MELF− cases vs 17.65% in MELF+ cases) (Fig. 2).

Thirty cases (83.33%) showed a wild-type pattern for p53 in MELF+ patients. Among MELF− patients, 141 cases (90.79%) showed a wild-type pattern for p53. There were no significant differences between the 2 groups in the p53 phenotype.

3.3. Survival and recurrence

All the 189 patients were followed up for 36.8 ± 8.9 months (18–54 months). Only 1 patient with MELF pattern was diagnosed with vaginal recurrence 28 months after the surgery. She, diagnosed as FIGO stage IIIA, underwent laparoscopic hysterectomy with bilateral salpingo-oophorectomy and pelvic lymphadenectomy, and underwent chemotherapy. She underwent partial upper vaginectomy and radiotherapy for the recurrence. One patient without MELF pattern died of heart failure 6 months after the diagnosis of endometrial cancer. The other patients were followed up with no local recurrence or systemic metastasis occurred.

4. Discussion

Endometrial endometrioid carcinomas show various patterns of myometrial invasion. There have been described 5 myoinvasive patterns, respectively diffusely infiltrating, broad front, adenomyosis-like, microcystic, elongated, and fragmented (MELF) glands and adenoma malignum, each having morphological and prognostic particularities. The frequency of MELF pattern was 17.99% in our study. MELF pattern is reported with variable frequencies, ranging between 9.4% and 23.1%.[3,4,10–12] The frequencies fluctuated markedly. It may be that some studies included different pathologic types of endometrial cancer. On the other hand, it may also be due to insufficient understanding of the MELF pattern, which had led to an underestimation. The histologic pattern of myometrial invasion in endometrial carcinomas may be a possible predictor for tumor evolution.[13] The biological characters and prognostic significance of MELF pattern remained unclear, although several studies have investigated its clinicopathologic features. Our study showed that MELF pattern was associated with adverse histologic findings such as larger tumor size, deeper myometrial invasion, LVSI and LNM in patients with endometrial endometrioid carcinoma. These findings in our study have been demonstrated in some previous studies. One study showed MELF pattern was associated significantly with larger tumor size, myometrial invasion of more than 50%, advanced FIGO stages, LNM and LVSI, papillary architecture, and mucinous differentiation among the patients with low-grade endometrioid carcinoma.[12] MELF pattern was more common in low-grade endometrioid carcinoma.[14] However, Tresserra F et al observed MELF pattern can be seen in high-grade endometrioid adenocarcinoma of the endometrium.[14] In our study, MELF pattern was found exclusively in low-grade endometrioid carcinomas. Han et al found there was a significant correlation between MELF pattern and cervical stroma involvement.[15] In our study, MELF pattern was not associated with cervical stroma involvement.

The association between MELF pattern and LNM remained uncertain. Several studies proved that MELF pattern was associated significantly with LNM.[3,10,11,16,17] The high probability of LNM in MELF pattern can lead to better therapeutic management, where the role of lymphadenectomy in the surgical management of endometrial cancer remains controversial. Given the favorable evolution in most cases of low-grade endometrial carcinoma, lymphadenectomy is generally avoided because of the serious potential short-term and long-term sequelae, such as lower limb lymphedema, vascular or nerve injury, symptomatic lymphocyst and chylous ascites.[18] In this context, identification of MELF pattern could represent an indication for subsequent lymphadenectomy. Furthermore, sentinel node mapping is increasingly being utilized for endometrial cancer staging.[19] However, there were previous studies demonstrated that MELF was a univariate but not multivariate predictor of LNM.[5] Amy S et al also reported that MELF pattern is not an independent risk factor for LNM or extrauterine metastasis.[11] Our study, in agreement with those studies, showed that patients with MELF pattern were more likely to have LNM than those without MELF pattern (29.41% vs 8.20%), while MELF pattern was

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Figure 2. Immunohistochemistry of MELF pattern in endometrial endometrioid carcinoma. MELF-pattern glands are moderate positive for MSH2 (A), strong positive for MSH6 (B), weak positive for MLH1 (C) and weak positive for PMS2 (D). MELF = microcystic, elongated, and fragmented.
MELF pattern was statistically not a valuable predictor of lymph node metastasis. MELF pattern was associated with adverse histologic findings in endometrial endometrioid carcinomas. The MELF pattern may be important for identifying those patients who need comprehensive staging surgery. Nonetheless, its implication in affecting survival and recurrences is unclear and further larger studies are needed to clarify the exact role of MELF in prognosis and adjuvant therapy.

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Author contributions

Data curation: Yuhan Cai.
Methodology: Hongyan Guo.
Writing – original draft: Jinghua Song.
Writing – review & editing: Huajun Li.

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