Clinical features of ProMisE groups identify different phenotypes of patients with endometrial cancer

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Abstract

Background The Proactive Molecular Risk Classifier for Endometrial Cancer (ProMisE) groups has identified four molecular prognostic groups of endometrial cancer (EC): POLE-mutated (POLE-mt), mismatch repair-deficient (MMR-d), p53-abnormal (p53-abn), p53-wild-type (p53-wt). These groups might have different pathogenesis and risk factors, and might occur in different phenotypes of patients. However, these data are still lacking.

Objective To provide a clinical characterization of the ProMisE groups of EC.

Methods A systematic review and meta-analysis was performed by searching seven electronic databases from their inception to December 2020, for all studies reporting clinical characteristics of EC patients in each ProMisE group. Pooled means of age and BMI and pooled prevalence of FIGO stage I and adjuvant treatment in each ProMisE group were calculated.

Results Six studies with 1, 879 women were included in the systematic review. Pooled means (with standard error) and prevalence values were:

- in the MMR-d group, age = 66.5 ± 0.6; BMI = 30.6 ± 1.2; stage I = 72.6%; adjuvant treatment = 47.3%.
- in the POLE-mt group, age = 58.6 ± 2.7; BMI = 27.2 ± 0.9; stage I = 93.7%; adjuvant treatment = 53.6%.
- in the p53-wt group, age = 64.2 ± 1.9; BMI = 32.3 ± 1.4; stage I = 80.5%; adjuvant treatment = 45.3%.
- in the p53-abn group, age = 71.1 ± 0.5; BMI = 29.1 ± 0.5; stage I = 50.8%; adjuvant treatment = 64.4%.

Conclusion The ProMisE groups identify different phenotypes of patients. The POLE-mt group included the youngest women, with the lower BMI and the highest prevalence of stage I. The p53-wt group included patients with the highest BMI. The p53-abn group included the oldest women, with the highest prevalence of adjuvant treatment and the lowest prevalence of stage I. The MMR-d group showed intermediate values among the ProMisE groups for all clinical features.

Keywords Prognosis · Treatment · Endometrium · Risk assessment · Tumor · Tumour · Carcinoma

Introduction

Endometrial cancer (EC) is the most prevalent gynecologic tumor in the western countries [1]. In the last decades, it also increased in incidence and mortality, due to an inaccurate histopathologic-driven management of patients [1–5]. The current histopathologic risk assessment is indeed poorly reproducible, leading to over- or undertreatment of women, and misinterpretations of findings within clinical trials [5, 6].

In 2013, The Cancer Genome Atlas (TCGA) Research Network has identified four novel prognostic groups of EC based on molecular signatures [7]. Due to technical difficulties and costs of sequencing analysis, a novel molecular classifier, the Proactive Molecular Risk Classifier for
Endometrial Cancer (ProMisE), has been developed based on immunohistochemistry as surrogate of sequencing [2, 6, 8, 9]. Immunohistochemistry is indeed more diffuse in the clinical practice because it is inexpensive and fast [10–16]. ProMisE classifies ECs in the following four prognostic groups: POLE-mutated (POLE-mt), mismatch repair-deficient (MMR-d), p53-abnormal (p53-abn), p53-wild-type (p53-wt). POLE-mt group includes ECs with the best prognosis and the highest mutational load; this group is characterized by mutations in the exonuclease domain of Polymerase-ε (POLE) and is the only one that can be identified exclusively by sequencing. MMR-d group has intermediate prognosis, high mutational load and microsatellite instability; this group can be identified by deficient immunohistochemical expression of mismatch repair protein (MMR). P53-abn group has the worst prognosis, low mutational load, high somatic copy number alteration rate and TP53 mutation; this group can be identified by abnormal immunohistochemical expression of p53. P53-wt group has good-to-intermediate prognosis, low mutational load, low somatic copy number alteration rate, and absence of a molecular signature; this group is identified by excluding molecular signatures of the other groups [2, 3, 6, 8, 9].

Given the differences in terms of molecular background, histologic characteristics and prognosis [17–22], these groups may be considered as different diseases within endometrial cancer landscape. These different entities might also have different pathogenesis and risk factors, and might occur in different phenotypes of patients. Specific clinical features in each ProMisE group may allow hypothesizing tailored prevention strategies and additional treatments (e.g. bariatric surgery and/or diet in the groups associated with obesity) for the single patient in the era of precision medicine [23]. Moreover, specific clinical characteristics may contribute to the prognosis of the ProMisE groups (e.g. younger age, early FIGO stage and/or more common adjuvant treatment may be associated with better prognosis). Therefore, while prognostic and histopathological features of the ProMisE groups were previously summarized [3, 17], this study aimed to provide a clinical characterization of the ProMisE groups of EC, with regards to age, body mass index (BMI), FIGO stage, and adjuvant treatment.

Materials and methods

Study protocol

Methods for each study step (i.e. search strategy, study selection, assessment of risk of bias within studies, data extraction and analysis) were a priori within the study protocol. Each study step was independently completed by two authors (AR, AT). All authors were asked for solution of disagreements. The study was reported according to the Preferred Reporting Item for Systematic Reviews and Meta-analyses (PRISMA) statement [24].

Search strategy

Search strategy was planned using seven electronic databases (i.e. Web of Sciences, Google Scholar, Scopus, ClinicalTrial.gov, MEDLINE, Cochrane Library and EMBASE) from their inception to December 2020. The following text words were alternatively combined: “ProMisE”; “Proactive Molecular Risk Classifier”; PORTEC”; “TransPORTEC”; “TCGA”; “Atlas”; “genome”; “survival”; “prognosis”; “endometrium”; “tumor”; “tumour”; “neoplas*”; “cancer”; “carcinoma”; “endometrioid”; “adenocarcinoma”; “serous”; “clear cell”; “undifferentiated”; “ultramutated”; “hypermutated”; “copy number”; “POLE”; “mismatch repair”; “MMR”; “MMR-d”; “MSI”; “microsatellite instability”; “MLH1”; “MSH2”; “MSH6”; “PMS2”; “EPCAM”; “TP53”; “p53”; “tumor protein 53”; “surrogate”; “immunohistochemistry”; “immunohistochemical”; “marker”; “sequencing”. We considered also all references from each full-text screened study.

Study selection

All peer-reviewed studies reporting clinical characteristics (age, BMI, FIGO stage, adjuvant treatment) of EC patients by each ProMisE group were included in our review. A priori defined exclusion criteria were: case reports, reviews, data not extractable, and studies with patients’ selection based on pathological characteristics of ECs (they were not representative of a real EC population). For studies with overlapping data (i.e. same period of enrollment, study population, institution, and/or findings), if the original data in each study could not be extracted separately, the study with smaller sample size was excluded from the quantitative analysis.

Data extraction

Data were extracted from the included studies without modification and according to the PICO (Population, Intervention, Comparator, Outcomes) items [24]. “Population” of our study was patients with EC. “Intervention” (or risk factor) was the MMR-d, POLE-mt or p53-abn group of EC according to the ProMisE. “Comparator” was not considered because it was not applicable (meta-analysis of prevalence). “Outcomes” were the means ± standard error of age and BMI, and the prevalence of the FIGO stage I and adjuvant treatment in the ProMisE groups of EC.
In the studies with overlapping patient data, duplicate data were excluded and only original data were considered.

**Assessment of risk of bias within studies**

The assessment of risk of bias within studies followed the Methodological Index for Non-Randomized Studies (MINORS) statement [25]. The following six domains related to risk of bias were applicable: (1) aim (if the aim was clearly stated); (2) inclusion of consecutive patients (if all eligible patients during the study period were included); (3) prospective collection of data (if an a priori defined protocol was adopted for data collection); (4) endpoints appropriate to the aim (if two or more authors performed a blind evaluation, re-evaluation or evaluation of study endpoints); (5) unbiased assessment of the study endpoint (if two or more authors performed a blind evaluation, re-evaluation or evaluation of study endpoints); (6) follow-up period appropriate to the aim (if the follow-up time was more than 2 years, which is the minimal follow-up period for patients with endometrial cancer).

Each domain was judged by authors as “low risk”, “unclear risk”, or “high risk” of bias if data were “reported and adequate”, “not reported”, or “reported but inadequate”, respectively.

**Data synthesis**

Means of age and BMI, and prevalence of FIGO stage I and adjuvant treatment in each ProMisE group of EC were calculated for each included study and as pooled estimate. They were graphically reported on forest plots, with 95% confidence interval (CI).

The inconsistency index \( I^2 \) was used to assess statistical heterogeneity among included studies, as previously described [26, 27]. Heterogeneity was considered null for \( I^2 = 0 \), minimal for \( I^2 < 0.25 \), low for \( I^2 < 0.50 \), moderate for \( I^2 < 0.75 \) and high for \( I^2 \geq 0.75 \).

All analyses were performed by adopting the random effect model of DerSimonian and Laird.

Data analysis was performed by Review Manager 5.3 (Copenhagen: The Nordic Cochrane Centre, Cochrane Collaboration, 2014) and Comprehensive Meta-Analysis (Bios tat,14 North Dean Street, Englewood, NJ 07631, USA).

**Results**

**Study selection**

Electronic searches identified 5,784 studies. 895 studies remained after duplicates removal. 52 studies remained after title screening. 21 studies were evaluated for eligibility after abstracts screening. Lastly, six studies were included in the systematic review and five studies in the meta-analysis [2, 6, 8, 9, 28, 29].

Figure S1 graphically shows the study selection process.

**Study and patients’ characteristics**

Our qualitative analysis included a total of 1,879 women with EC from retrospective cohorts. Mean age of patients ranged from 42.9 ± 5.6 to 66.9 ± 0.7, and mean BMI ranged from 29 ± 7.7 to 33 ± 1. Overall, prevalence of Stage I and adjuvant treatment was 75.1% and 48.8%, respectively. Of total EC, 23.9% were classified in MMR-d, 10% in POLE-mt, 51% in p53-wt, and 15% in p53-abn groups.

Characteristics of the included studies and patients are shown Tables S1 and S2, respectively.

**Risk of bias within studies assessment**

All included studies were judged at low risk of bias in all domains, except for “Inclusion of consecutive patients” domain. In particular, four studies were judged at “unclear risk” of bias in this domain because they did not report if all eligible patients during the study period were included [2, 6, 9, 28]. Furthermore, Britton et al. performed a patient’s selection based on age, only including women aged 49 or younger [9]. The remaining study was judged at low risk of bias.

Assessment of risk of bias within studies is graphically shown in Figure S2.

**Meta-analysis**

All duplicate patient data were excluded from the meta-analysis: the study by Britton et al. was excluded because of an overlap of patients with the other included studies [9]; for the Talhouk 2017 study, only the new patients (“confirmation cohort”) were included in the meta-analysis, while patients overlapping with Talhouk 2015 (“discovery cohort”) were excluded [2, 6]; the study by Kolehmainen et al. was included only in the analysis of FIGO stage, since it reported neither standard errors for age and stage nor data regarding adjuvant treatment [28]. Finally, five studies with 1,622 were included in the meta-analysis of FIGO stage, while four studies with 1,018 women were included in the meta-analyses of age, BMI and adjuvant treatment.

Pooled means with standard error of age was 66.5 ± 0.6 (95% CI 65.3–67.7%) in the MMR-d group, 58.6 ± 2.7 (95% CI 53.4–63.9) in the POLE-mt group, 64.2 ± 1.9 (95% CI 60.6–67.9) in the p53-wt group, 71.1 ± 0.5 (95% CI 70.2–72) in the p53-abn group (Fig. 1). Statistical heterogeneity among studies was high for each group, with the exception for the p53-abn group where it was moderate (\( I^2 = 86.4; F^2 = 96; F^2 = 99.7; F^2 = 66 \), respectively).
Pooled means with standard error of BMI was $30.6 \pm 1.2$ (95% CI 28.3–33) in the MMR-d group, $27.2 \pm 0.9$ (95% CI 25.6–29) in the POLE-mt group, $32.3 \pm 1.4$ (95% CI 29.6–34.9) in the p53-wt group, $29.1 \pm 0.5$ (95% CI 28.2–30) in the p53-abn group (Fig. 2). Statistical heterogeneity among studies was high for each group ($I^2 = 93.6$; $I^2 = 82.4$; $I^2 = 98.7$; $I^2 = 72.4$, respectively).

Pooled prevalence of FIGO stage I was 72.6% (95% CI 67–77.6%) in the MMR-d group, 93.7% (95% CI 87.4–97%) in the POLE-mt group, 80.5% (95% CI 75.2–84.9%) in the p53-wt group, 50.8% (95% CI 44.6–56.9%) in the p53-abn group (Fig. 3). Statistical heterogeneity among studies was moderate for the MMR-d group ($I^2 = 35.8$), null for POLE-mt and p53-abn group ($I^2 = 0$), and low for the p53-wt group ($I^2 = 57.2$).

Pooled prevalence of adjuvant treatment was 47.3% (95% CI 43–63.9%) in the MMR-d group, 53.6% (95% CI 43–63.9%) in the POLE-mt group, 45.3% (95% CI 26–66.1%) in the p53-wt group, 64.4% (95% CI 48.4–77.7%) in the p53-abn group (Fig. 4). Statistical heterogeneity among studies was high for each group, with the exception for the POLE-mt group where it was null ($I^2 = 86.8$; $I^2 = 0$; $I^2 = 93.9$; $I^2 = 72.4$, respectively).

Pooled estimates of means or prevalence of the clinical features in the ProMisE groups are concisely reported in Table S3.

Discussion

Main findings and interpretation

This study aimed to provide a clinical characterization of the ProMisE groups of EC to hypothesize possible prevention strategies and additional treatments that may be tailored on the single patient, and to further explain prognosis data across the ProMisE groups. We found that the POLE-mt group included the youngest women, with the lower BMI and the highest prevalence of FIGO stage I. The p53-wt group included patients with the highest BMI and the lowest prevalence of adjuvant treatment. The p53-abn group included the oldest women, with the highest prevalence of adjuvant treatment and the lowest prevalence of FIGO stage I. The MMR-d group showed intermediate values among the ProMisE groups for all clinical features.

MMR-d group

The intermediate values for all clinical characteristics in this group are in accordance with survival data, which showed an intermediate prognosis [2, 3, 6–8]. In fact, in our previous study, we found that the prognostic value of MMR defect signature was affected by prognostic clinicopathological features. In particular, the difference with the p53-wt group in terms of overall survival, disease-specific survival and progression-free survival became not significant when normalized for clinicopathological factors [3]. Moreover, it has to be shown that the prognostic value of the MMR defect signature may also be affected by the study population. In particular, in early stage endometrioid ECs, it shows an
unfavorable prognostic role, while in high-grade ECs it may improve the prognosis [15, 30, 31].

In addition, we found that these patients were obese. Such a finding is in agreement with the hormone-driven pathogenesis of these tumors the first phase of the pathogenesis. In fact, although a MMR defect signature is associated with lower hormone-responsiveness, MMR-d ECs arise from atypical endometrial hyperplasia, which is an hormone-responsive lesion at least in its earliest phases [32, 33]. However, the mean BMI in the MMR-d group was lower compared to p53-wt group; such a difference might be due to the presence of patients with Lynch syndrome, which may develop MMR-d regardless of an initial estrogenic stimulation [32].

Fig. 2 Mean body mass index in ProMisE groups of endometrial cancer for each study and as pooled estimate

Fig. 3 Forest plot of prevalence of FIGO stage I in ProMisE groups of endometrial cancer, including individual study and pooled data
The clinical features of the patients in the POLE-mt group seem to outline a specific phenotype: the women in this group were the youngest ones and showed the lowest BMI. These findings might suggest POLE-mt cancers are less “estrogen-related” if compared to the prototypical Bokhman type I EC \[7, 34\]. In this regard, the high mutational load of POLE-mt ECs might support the onset of estrogen-independent mechanisms \[7\]. Thus, these carcinomas might also be less responsive to hormonal therapy.

The specific clinical phenotype of the POLE-mt group might also contribute to the favorable prognosis of this group \[2, 3, 6–8\]. In fact, the younger age of these patients might positively affect overall survival, given the lower likelihood of death, comorbidity, and severe side effect of chemotherapy associated with the young age. The higher prevalence of FIGO stage I (93.3%) may also contribute to the good prognosis. Such prevalence might be due to a low-tumor aggressivity with slow progression, or to more frequent check-ups in younger women with consequently earlier EC diagnosis. The high frequency of adjuvant treatment in POLE-mt EC (53.6%) is likely explained by the high prevalence of high-grade ECs in this group (especially considering that the vast majority is at FIGO stage I), and is a further factor that may improve prognosis \[35, 36\].

Although age and FIGO stage might have a role in the good prognosis of the POLE-mt group, in our previous study we found that POLE mutation appeared as the molecular signature least affected by other prognostic clinicopathological factors \[3\]. For this reason, the fact that more than 50% of these women currently undergo adjuvant treatment may constitute an overtreatment.

**p53-wt group**

Patients in this group were obese and showed the highest BMI among all groups. This would support the estrogen-driven pathogenesis making inroads to possible prevention strategy based on diet and bariatric surgery for patients with severe obesity. The high BMI, the good-to-intermediate prognosis and the high prevalence of low-grade endometroid carcinomas in this group reflect the prototypical type I EC according to the Bokhman’s classification \[34\]. Patient age and stage I prevalence were intermediate. These findings may contribute to the good-to-intermediate prognosis in these patients.

**p53-abn group**

The p53-abn group appears to embody a specific phenotype of patient, including the oldest women among the ProMisE groups. Such feature is consistent with the high prevalence of serous carcinoma, which arises in a background of atrophic endometrium \[7\]. Moreover, these patients were non-obese and with the lowest prevalence of FIGO stage I (48.8%). These characteristics, along with the high prevalence of serous histotype, make these cancers assimilable to the type II endometrial cancer of Bokhman’s classification \[42\]. The highly unfavorable clinical profile of the p53-abn group likely contribute to the poor prognosis of this group. Indeed, in our previous study we found that unfavorable
clinicopathological factors sensibly worsened the prognosis of the p53-abn group [3]. However, we also found that the unfavorable prognostic value of the p53-abn signature remained significant even after adjusting for other clinicopathological factors. In this regard, the prognosis remains the worst one although most EC of this group (64.4%) undergo adjuvant treatment.

**Strengths and limitations**

To the best of our knowledge, this study may be the first systematic review and meta-analysis to provide a clinical characterization of the ProMisE groups of EC. This study hypothesized that the ProMisE groups might benefit from specific prevention strategies and additional treatments (e.g. diet, bariatric surgery, hormonal therapy) to be tailored on the single patient. Moreover, the provided characterization may be useful to further explain prognosis data across the ProMisE groups. The overall quality of the included studies is very high, given that almost all the domains related to risk of bias were judged at low risk of bias and no one was considered at high risk.

The low number of included studies might be a limitation of our meta-analysis. Nonetheless, the presented data were devoid of patient overlap due to the exclusion of all duplicate data.

**Conclusion**

The clinical characterization of ProMisE groups of EC shows that molecular signatures were associated with different phenotypes of patients. The POLE-mt group includes the youngest women, with the lowest BMI and the highest prevalence of FIGO stage I. The p53-wt group includes patients with the highest BMI. The p53-abn group includes the oldest women, with the highest prevalence of adjuvant treatment and the lowest prevalence of FIGO stage I. The MMR-d group showed intermediate values among the ProMisE groups for all clinical features. The clinical characterization of these groups may suggest different pathogenetic mechanisms and may also contribute to explain the prognostic data of the ProMisE groups, with potential impact on the patient management.

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**Declarations**

**Conflict of interest** The authors report no conflict of interest.

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**References**

1. Siegel RL, Miller KD, Jemal A (2015) Cancer statistics, 2015. CA Cancer J Clin 65:5–29
2. Talhouk A, McConkey MK, Leung S, Li-Chang HH, Kwon JS, Melnyk N et al (2015) A clinically applicable molecular-based classification for endometrial cancers. Br J Cancer 113:299–310
3. Raffone A, Travaglino A, Mascolo M, Carbone L, Guida M, Insabato L, et al (2019) TCGA molecular groups of endometrial cancer: pooled data about prognosis. Gynecol Oncol. [Epub ahead of print], https://doi.org/10.1016/j.ygyno.2019.08.019
4. Gilks CB, Oliva E, Soslow RA (2013) Poor interobserver reproducibility in the diagnosis of high-grade endometrial carcinoma. Am J Surg Pathol 37:874–881
5. Hoang LN, McConkey MK, Kobel M, Han G, Rouzbahman M, Davidson B et al (2013) Histotype-genotype correlation in 36 high-grade endometrial carcinomas. Am J Surg Pathol 37:1421–1432
6. Talhouk A, McConkey MK, Leung S, Yang W, Lum A, Senz J et al (2017) Confirmation of ProMisE: a simple, genomics-based clinical classifier for endometrial cancer. Cancer 123:802–813
7. Cancer Genome Atlas Research Network, Kandoth C, Schultz N, Cherniack AD, Akbani R, Liu Y, et al (2013) Integrated genomic characterization of endometrial carcinoma. Nature 497:67–73
8. Kommos S, McConney MK, Kommos F, Leung S, Bunza A, Magrill J et al (2018) Final validation of the ProMisE molecular classifier for endometrial carcinoma in a large population-based case series. Ann Oncol 1(29):1180–1188
9. Britton H, Huang L, Lum A, Leung S, Shum K, Kale M et al (2019) Molecular classification defines outcomes and opportunities in young women with endometrial carcinoma. Gynecol Oncol 153:487–495
10. Kim G, Kurnit KC, Djordjevic B, Singh C, Munsell MF, Wang WL et al (2018) Nuclear β-catenin localization and mutation of the CTNNB1 gene: a context-dependent association. Mod Pathol 31:1553–1559
11. Raffone A, Travaglino A, Cerbone M et al (2020) Diagnostic accuracy of p53 immunohistochemistry as surrogate of TP53 sequencing in endometrial cancer. Pathol Res Pract 216(8):153025
12. Raffone A, Travaglino A, Cerbone M et al (2020) Diagnostic accuracy of immunohistochemistry for mismatch repair proteins as surrogate of microsatellite instability molecular testing in endometrial cancer. Pathol Oncol Res 26(3):1417–1427
13. Travaglino A, Raffone A, Saccone G et al (2019) Nuclear expression of β-catenin in endometrial hyperplasia as marker of pre-malignancy. APMSIS 127(11):699–709
14. Stelloo E, Nout RA, Osse EM, Jürgenliemk-Schulz IJ, Jobsen JJ, Lutgens LC et al (2016) Improved risk assessment by integrating molecular and clinicopathological factors in early-stage endometrial cancer—combined analysis of the PORTEC cohorts. Clin Cancer Res 22:4215–4224
15. Stelloo E, Bosse T, Nout RA, MacKay HJ, Church DN, Nijman HW et al (2015) Refining prognosis and identifying targetable pathways for high-risk endometrial cancer; a TransPORTEC initiative. Mod Pathol 28:836–844
16. Raffone A, Travaglino A, Saccone G, Cieri M, Mascolo M, Mollo A et al (2019) Diagnostic and prognostic value of ARID1A in endometrial hyperplasia: a novel marker of occult cancer. APMSIS 127:597–606
17. Raffone A, Travaglino A, Mascolo M et al (2020) Histopathological characterization of ProMisE molecular groups of endometrial cancer. Gynecol Oncol 157(1):252–259. https://doi.org/10.1016/j.ygyno.2020.01.008
18. Travaglino A, Raffone A, Stradella C et al (2020) Impact of endometrial carcinoma histotype on the prognostic value of the TCGA molecular subgroups. Arch Gynecol Obstet 301(6):1355–1363
19. Travaglino A, Raffone A, Gencarelli A et al (2020) TCGA classification of endometrial cancer: the place of carcinosarcoma. Pathol Oncol Res 26(4):2067–2073
20. Travaglino A, Raffone A, Mollo A et al (2020) TCGA molecular subgroups and FIGO grade in endometrial endometrioid carcinoma. Arch Gynecol Obstet 301(5):1117–1125
21. Travaglino A, Raffone A, Mascolo M et al (2020) TCGA molecular subgroups in endometrial undifferentiated/dedifferentiated carcinoma. Pathol Oncol Res 26(3):1411–1416
22. Travaglino A, Raffone A, Mascolo M et al (2020) Clear cell endometrial carcinoma and the TCGA classification. Histopathology 76(2):336–338
23. McAlpine JN, Gilks CB (2019) Precision medicine in endometrial carcinoma. Pathol Oncol Res 153(11):932–937
24. Zakhour M, Cohen JG, Gibson A et al (2017) Abnormal mismatch repair and other clinicopathologic predictors of poor response to preoperative progesterin treatment in young women with endometrial complex atypical hyperplasia and well-differentiated endometrial adenocarcinoma: a consecutive case series. BJOG 124(2):194–200
25. Slim K, Nini E, Forestier D, Kwiatkowski F, Panis Y, Chipponi J (2003) Methodological index for non-randomized studies (minors): development and validation of a new instrument. ANZ J Surg 73:712–716
26. Raffone A, Travaglino A, Saccone G et al (2020) Diabetes mellitus is associated with occult cancer in endometrial hyperplasia. Pathol Oncol Res 26(3):1377–1384
27. Raffone A, Travaglino A, Saccone G et al (2019) Diabetes mellitus and responsiveness of endometrial hyperplasia and early endometrial cancer to conservative treatment. Gynecol Endocrinol 35(11):932–937
28. Kolehmainen AM, Pasanen AM, Koivisto-Korander RL, Büttzow RC, Loukovaara MJ (2020) Molecular characterization in the prediction of disease extent in endometrial carcinoma. Eur J Obstet Gynecol Reprod Biol. (Epub ahead of print) https://doi.org/10.1016/j.ejogrb.2020.10.031
29. Timmerman S, Van Rompuy AS, Van Gorp T, Vanden Bempt I, Brems H, Van Nieuwenhuyzen E, Han SN, Neven P, Victoor J, Laenen A, Vergote I (2020) Analysis of 108 patients with endometrial carcinoma using the PROMISE classification and additional genetic analyses for MMR-D. Gynecol Oncol 157(1):245–251
30. Moroney MR, Davies KD, Wilberger AC, Sheeder J, Post MD, Berning AA et al (2019) Molecular markers in recurrent stage I, grade 1 endometrioid endometrial cancers. Gynecol Oncol 153:517–520
31. Bosse T, Nout RA, McAlpine JN, McConeney MK, Britton H, Hussein YR et al (2018) Molecular classification of grade 3 endometrioid endometrial cancers identifies distinct prognostic subgroups. Am J Surg Pathol 42:561–568
32. Ferreira AM, Westers H, Albergaria A, Seruca R, Hofstra RM (2009) Estrogens, MSI and Lynch syndrome-associated tumors. Biochim Biophys Acta 1796(2):194–200
33. Kahlau M, Cohen JG, Gibson A et al (2017) Abnormal mismatch repair and other clinicopathologic predictors of poor response to progestin treatment in young women with endometrial complex atypical hyperplasia and well-differentiated endometrial adenocarcinoma: a consecutive case series. BJOG 124(10):1576–1583
34. Bokhman JV (1983) Two pathogenetic types of endometrial carcinoma. Pathol Oncol Res 151(1):10–17
35. Colombo N, Creutzberg C, Amant F, Bosse T, González-Martín A, Ledermann J, et al (2016) ESMO-ESGO-ESTRO consensus conference on endometrial cancer: diagnosis, treatment and follow-up. Ann Oncol 27:16–41
36. Abu-Rustum NR, Yashar CM, Bean S, et al (2019) NCCN clinical practice guidelines in oncology (NCCN Guidelines®)—Uterine neoplasms. Version 3.2019 – February 11, 2019

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