

# Impact of the new molecular classification of endometrial cancer: A French cohort study

Jeremie Benichou, Corentin Schwall, Xavier Sastre-Garau, Julie Méreaux, Grégoire Miailhe, Sofiane Bendifallah, Bassam Haddad, Cyril Touboul, Rana Mitri-Frangieh, Yohann Dabi

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# IMPACT OF THE NEW MOLECULAR CLASSIFICATION OF ENDOMETRIAL CANCER: A FRENCH COHORT STUDY 3

- 4 Jeremie Benichou<sup>1</sup>\*, Corentin Schwall<sup>2</sup>\*, Xavier Sastre-Garau<sup>2</sup>, Julie Méreaux<sup>1</sup>, Grégoire
- 5 Miailhe<sup>1</sup>, Sofiane Bendifallah<sup>3</sup>, Bassam Haddad<sup>1</sup>, Cyril Touboul<sup>3</sup>, Rana Mitri Frangieh<sup>2</sup>,
- 6 Yohann Dabi<sup>3</sup>

7 \* These two authors contributed equally to this work and should be considered as joint first8 authors.

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- Unversity of Medicine Paris XII, Department of Obstetrics and Gynecology, Centre Hospitalier Intercommunal, Créteil, France
- University of Medicine Paris XII, Department of Pathology, Centre Hospitalier
   Intercommunal de Créteil, France
- Sorbonne University Department of Obstetrics and Gynecology, Tenon Hospital, AP-HP,
   Paris, France
- 16
- 17 Corresponding author:
- 18 Yohann DABI, M.D
- 19 Sorbonne University, Department of obstetrics and Gynecology, Tenon Hospital AP-HP
- 20 4 rue de la Chine, 75020 Paris, France
- 21 E- mail: yohann.dabi@gmail.com
- 22 Tel: 01.56.01.70.00

- 23 Abstract
- 24

Objective: To evaluate the potential impact of the latest ESGO guidelines for endometrial
 cancer with molecular classification on the management strategy in a French cohort.

27 Methods: All patients treated between January 1st, 2014 and December 31, 2020 for an 28 endometrial cancer at the Centre Hospitalier Intercommunal de Créteil (CHIC, FRANCE) were selected from our prospectively maintained database. All postoperative samples were 29 30 reviewed to confirm histological subtype, myometrial infiltration, cytonuclear grade and 31 presence of lymphovascular emboli. Analysis of p53, MLH1, MSH2, MSH6, PMS2 genes was performed by immunohistochemistry first then a systematic POLE sequencing was 32 33 performed to identify gene mutation. The impact of the latest ESGO 2020 guidelines was assessed regarding adjuvant therapy, surgical strategy, and survival. 34

**Results**: Eighty patients were analyzed, including 70% NSMP (n = 56), 13.75% MSI (n =35 36 11), 10% p53 mutated (n = 8) and 6.25% POLEmut (n = 5). A total of 21 patients (26.3%) were reclassified using the latest ESGO classification. Patients classified at low risk or with 37 advanced / metastatic disease were not reclassified using molecular analysis. Molecular 38 39 analysis and the latest ESGO classification had the most important impact on patients initially classified at intermediate – high risk that were reclassified in intermediate (10/23) and in low 40 41 (4/23) risk. Nine patients (11.3%) were overtreated according to the 2020 ESGO 42 classification: six patients in the low – risk group (4 received vaginal brachytherapy and 2 43 external radiotherapy) and three in the intermediate risk group (3 received external irradiation 44 and 1 received chemotherapy). None of the patients in our cohort would have been undertreated using the 2020 ESGO classification. Patients within the p53 mutated group were 45 46 the most likely to experience recurrence (37.5%, 3/8) and none of the patients POLE mutated 47 recurred.

48	<b>Conclusion</b> : Around one in 4 patients were reclassified in a more accurate prognostic group
49	using molecular diagnosis and the latest ESGO guidelines which could decrease the use of
50	adjuvant therapies to spare morbidity.
51	Keywords: endometrial cancer; molecular classification; ESGO guidelines; survival; risk

- assessment; prognostic
- 54

#### 55 Introduction

Endometrial carcinoma (EC) is currently the most common gynecological pelvic 56 malignancy in developed countries, accounting for 57.8% of new cases of gynecological 57 58 cancers in the US in 2020 (1). Preoperative assessment of the risk of lymph node invasion is 59 currently based on histotype and grade in patients that do not exhibit lymph node invasion on preoperative MRI (2). These parameters have been shown to have poor reproducibility (3). 60 61 The generalization of the sentinel lymph node procedure even in patients classified 62 preoperatively at low risk has significantly reduced the complication risk and the morbidity 63 rates and reshuffled the cards (4,5). However, preoperative accurate assessment of lymph node 64 invasion risk still maters to both anticipate adjuvant therapies and inform patients accordingly (6). Besides, lymphovascular space invasion, which could be very relevant to refine risk 65 group, is hardly assessed on preoperative biopsy (7.8). All of these factors result in partial 66 67 preoperative assessment potentially leading to inadequate surgical gestures. Moreover, the postoperative risk of recurrence assessment has been shown to have a limited predictive value 68 as some patients at "low - risk" experience recurrences sometimes with a short delay 69 70 following treatment (9).

71 In 2013, the Cancer Genome Atlas (TCGA) research network group performed an 72 integrated genomic characterization of 373 endometrial carcinomas (EC) using sequencing 73 and array-based technologies (10). Based on these findings, the ProMisE classification has 74 identified four molecular groups of EC with different prognoses(11): the POLE-mut group 75 (POLEmut), the mismatch repair-deficient group (MMRd), the p53-abn group is classified as 76 "high copy number" and the p53-wild-type group (p53-wt) or "non-specific molecular profile" 77 (NSMP). More than individually, the ProMise classification appears to be a beneficial and 78 complementary contribution to the 2013 ESMO classification. Talhouk et al in 2017 reported 79 that regarding the main oncological outcomes (OS, DFS and PFS), ProMisE use alone seems to perform as well as ESMO, or even better when postoperative parameters are 80

considered(12). The new ESTRO ESGO ESP 2020 guidelines have integrated the molecular classification into the management algorithms, with a modification of the risk groups and therefore of the medical and surgical management of endometrial cancers (13). The ultimate goal of applying accurate prognostic classification using molecular subtypes is to eventually reduce iatrogenic morbidity by decreasing indications of unindicated adjuvant therapies according to ESGO 2020 guidelines while efficiently reserving these treatments for patients truly at high risk.

88 To date, potential impact of these new guidelines on prognostic assessment and 89 management of patients with endometrial cancers has not been evaluated in a French cohort to 90 assess its external validity.

91

#### 93 Materials and methods

94 The protocol was validated by the Research Organization Committee of the Centre
95 Hospitalier Intercommunal de Créteil on September 26, 2019. Written consent was obtained
96 for all patients as part of the PELVIMASS protocol (CPP No. 2016-A01381-42)

97

#### 98 <u>Population</u>

All patients treated between January 1<sup>st</sup>, 2014 and December 31, 2020 for an endometrial cancer at the Centre Hospitalier Intercommunal de Créteil (CHIC, FRANCE) were selected from our prospectively maintained database. Patients for whom the tissue was not usable due to alterations during preservation or due to poor quality of DNA's extractions were excluded. Young patients < 18 years, those with rare histological forms, and those with numerous missing data were not included.

Data of interest were abstracted from patients' chart, including socio demographic characteristics, preoperative imaging and pathological analysis, prospective management including surgery and adjuvant therapies as well as survival data.

108

#### 109 Prospective management

Patients were treated in accordance with European recommendations at the time of prospective management(14,15). Preoperative management included clinical examination, pelvic ultrasonography and abdomino-pelvic MRI to determine loco-regional extension, lymph node involvement and distant metastases. Tumors' markers such as cancer antigen 125 (CA125) were measured in patients with type II tumors. The 2009 - FIGO classification was used to classify tumors (16).

Follow-up consisted of a clinical examination every 4 months for 3 years, then every 6 months for 2 years and then annually. Depending on the clinical findings, the histological type of the tumor and the initial extension of the tumor, a thoraco-abdomino-pelvic CT scan 119 could be requested as well as a biological evaluation including tumor markers CA125 for120 non-endometrioid tumors.

- 121
- 122 Pathological et molecular analysis

123 All postoperative samples were reviewed to confirm histological subtype, myometrial 124 infiltration, cytonuclear grade and presence of lymphovascular emboli. A systematic analysis 125 of p53, MLH1, MSH2, MSH6, PMS2 genes was performed first by immunohistochemistry. 126 Immunohistochemical staining was performed on a Ventana BenchMark Ultra© machine, 127 according to the protocols of the various antibody suppliers. The Thermo Fisher© monoclonal 128 antibody (DO-7 clone) was used for p53 testing. Results were characterized in 2 categories: a 129 heterogeneous positivity classified the sample as wild type. A strong and diffuse positivity ( 130 over-expression) or a complete absence of marking (negative) classified the sample as 131 abnormal. A systematic POLE sequencing was performed to identify gene mutation. This was 132 first screened by HRM (High Resolution Matching) PCR to select samples with suspected 133 POLE gene mutation. In order to precisely characterize the type of mutation, a gene 134 sequencing technique (Next-Generation Sequencing or NGS) was performed on the samples 135 previously selected by HRM.

In cases of loss of expression of immunophenotypic markers or ambiguity of the
immunostaining, a molecular technique was used using Idylla© (Biocartis, Mechelen,
Belgium). Eight cases had microsatellite instability searched using PCR prior the initiation of
this study.

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#### 141 Assessment of the new ESGO 2020 classification impact

All patients were reclassified according to the new ESGO 2020 classification, using
molecular analysis. The new risk group was then compared with the initial risk assessed

144 during prospective management. The impact of the new ESGO 2020 guidelines was assessed

145 by comparing adjuvant therapy and surgical strategy.

Survival of patients according to histological characteristics, prognostic risk group, and bymolecular group were analyzed.

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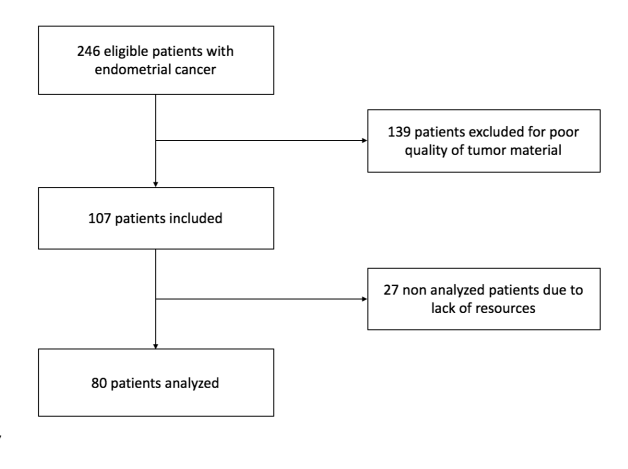
149 <u>Statistical analysis</u>

150 The data used were collected on a secure Excel sheet (Microsoft Corporation, Redmond, WA, 151 USA) and all statistical analyses were performed using the freely available online R software 152 (version 1.3.1093). For all analyses performed, a p-value < 0.05 was considered to indicate a 153 statistically significant difference. Categorical variables were compared using a Chi2 or 154 Fisher test according to the number of participants, and quantitative variables were compared 155 using a Student's t test. Kaplan-Meier survival curves were generated to assess recurrence-free 156 survival and overall survival according to the groups determined. The log-rank test was used 157 compare survivals. to

#### 1 Results

Among the 246 eligible patients, 107 patients were included and 139 patients were excluded due to poor quality of tumor material. Eventually, 27 unselected patients did not undergo molecular analysis due to lack of resources during the COVID 19 pandemic and thus were excluded leading to a total of 80 patients analyzed (Figure 1).

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Figure 1: Flow chart of the study

9

10 <u>Characteristics of the population</u>

The main characteristics of the patients included are displayed in Table 1. The mean age was 66 years old (range 34-87 years old) with an average BMI of 31kg/m<sup>2</sup> (range 20-51kg/m<sup>2</sup>). Diagnosis was obtained through endometrial biopsies in 76% (61/80) cases and operative hysteroscopy in 24% (19/80) cases. Patients that could not undergo molecular analysis due to COVID 19 pandemic were similar to those that did (Supplementary Table 1 and 2).

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18	endometrial cancer 72.5% (58/80).
17	6.3% (5/80) had para-aortic lymph node involvement (Table 1). Most patients had stage I
16	On preoperative MRI, 13.8% of patients (11/80) had pelvic lymph node involvement and

Characteristics	Final population
	N = 80 (%)
Age in years (mean $\pm$ sd)	66 ± 11.6
Body mass index (kg/m <sup>2</sup> ) mean ( $\pm$ sd)	31 (± 7.1)
Nulliparity	18 (26.5)
High blood pressure	43 (54)
Diabetes	14(18)
Menopausal	70 (87.5)
Bleeding	70 (87.5)
FIGO MRI stage	
IA	25 (31.25)
IB	33 (41.25)
II	5 (6.25)
III	7 (8.75)
IV	6 (7.5)

Surgery	
HBSO	78 (97.5)
Total Hysterectomy and ovarian sparing	2 (2.5)
Omentectomy	11 (13.8)
Appendectomy	6 (7.5)
Pelvic sentinel node	3 (3.8)
Pelvic lymphadenectomy	30 (37.5)
Para-aortic lymphadenectomy	26 (32.5)
Inguinal lymphadenectomy	3 (3.8)
External beam radiotherapy	32 (40)
Neoadjuvant chemotherapy	1 (1.3)
Adjuvant chemotherapy	21 (26.3)
Brachytherapy	56 (70)
Preoperative ESMO	
Low	22 (27)
Intermediate	29 (39)
High	26 (32,5)
NĂ	3

29 Table 1: Characteristics of the study population, treatments received by patients and

30 ESMO/ESGO 2013 preoperative classification. HBSO:

31 Total hysterectomy with bilateral salpingo-oophorectomy

32 NA: Not assessed

33 Patients' management

34 Lymph node staging was performed by sentinel node procedure in 3.8% cases, by pelvic

35 lymphadenectomy in 37.5% and para-aortic lymphadenectomy in 32.5% of the cases. No

36 lymph node staging was performed in 56% (45/80) and 17.5% of the patients (14/80)

37 underwent secondary surgery for lymph node staging.

38 Discrepancy between pre and postoperative histology occurred in 11.6% (7/60) and 10.5%

- 39 (2/19) of patients diagnosed by endometrial biopsy and operative hysteroscopy, respectively.
- 40 Regarding adjuvant therapies, brachytherapy, external radiotherapy and chemotherapy were

41 used in 70% (56/80), 40% (32/80), and 26.3% (21/80), respectively (Table 1).

42

### 43 Comparison of ESGO 2020 and ESMO 2016

- 44 The NSMP group was the most represented (70%, 56/80), followed by the MSI (13.75%;
- 45 11/80), the mutated P53 (10%, 8/80) and the POLEmut (6.25%, 5/80) groups (Table 2).
- 46 Morphological characteristics of tumors according to molecular group are described in
- 47 Supplementary Table 3.
- 48

	N = 80 (%)
Histological type	
Endometrioid	70 (87.5)
Serous	7 (8.75)
Serous + Endometrioid	1 (1.25)
Clear cell	2 (2.5)
Grade	
Low grade	62 (77.5)
High grade	18(22.5)
LVSI	
0	37 (46)
<5	13 (16)
>5	30 (38)
Molecular group	
POLE	5(6.25)
MSI	11(13.75)
NSMP	56 (70)
P53	8 (10)

50 Table 2: Histological and molecular characteristics of the study population

A total of 21 patients (26.3%) were reclassified following application of the new ESGO 2020 classification (table 3). Concordance between the two classifications regarding postoperative risk was observed in 73.7% (59/80). Patients classified at low risk or with advanced / metastatic disease were not reclassified using molecular analysis. Molecular analysis and the latest ESGO classification had the most important impact on patients initially classified at intermediate – high risk that were reclassified in intermediate (10/23) and in low (4/23) risk.

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ESMO 2016			Intermediate		
ESGO 2020	Low	Intermediate	high	High	Advanced/metastatic
Low	18 (75%)	2 (8%)	4 (17%)	0	0
Intermediate	0	10 (50%)	10 (50%)	0	0
Intermediate high	0	0	9 (64%)	5 (36%)	0
High	0	0	0	15 (100%)	0
Advanced metastatic	0	0	0	0	7 (100%)

61

62 Table 3: Number of patients classified into risk groups according to ESMO 2016 and ESGO

63 2020 recommendations. Proportions are calculated based on the ESGO 2020 group size.

64

#### 65 **Impact of the molecular classification**

66 Twelve patients (15%) had lymph node involvement on final analysis. Of these patients, 50%

67 were p53 mutated and 33% had no specific molecular profile. All of these patients were

classified as high risk (58%) or advanced/metastatic (42%). Patients in the p53mutated group

69 had lymph node involvement in 75% cases (6/8). The distribution of patients with lymph node

70 involvement by molecular group and prognostic classification is presented in Table 4.

	Pelvic lymph node	Para-aortic lymph	Lymph node
	involvement	node involvement	involvement
	(N=7)	(N=9)	(N=12)
Molecular group			
• POLE mutated (N=5; 6.25%)	0	0	0
• MSI (N=11; 13.75%)	0	2 (18%)	2 (18%)
• NSMP (N=56; 70%)	3 (5.4%)	2 (3.6%)	4 (7.1%)
• P53 mutated (N=8; 10%)	4 (50%)	5 (62.5%)	6 (75%)
ESGO Risk Group 2020			
• Low (N=24; 30%)	0	0	0
• Intermediate (N=20; 25%)	0	0	0
• Intermediate-high (N=14; 17%)	0	0	0
• High (N=15; 19%)	4 (27%)	4 (27%)	7 (47%)
• Advanced/metastatic (N=7; 9%)	3 (43%)	5 (71%)	5 (71%)

78 of lymph node involvement were calculated according to the size of the molecular groups or

the 2020 risk groups

82 Impact of the ESGO 2020 classification

<sup>77</sup> Table 4: Node involvement by molecular group and ESGO 2020 risk group. The proportions

The 2020 ESGO classification could have spared secondary surgery for staging in 21.4% (3/14) of patients classified at low or intermediate risk. According to the latest ESGO 2020 guidelines, 86% (69/80) of our patients could have benefited from the sentinel lymph node procedure (only intermediate-high risk or high - risk patients with FIGO stage >2 are not eligible).

A total of 9 patients (11.3%) were overtreated according the 2020 ESGO classification: Six patients in the low – risk group (4 received vaginal brachytherapy and 2 external radiotherapy) and three in the intermediate risk group (3 received external irradiation and 1 received chemotherapy).

92 None of the patients in our cohort would have been undertreated.

- 93
- 94

#### 95 <u>Survival analysis</u>

96 The median follow-up time was 25 months (0-64). During follow-up, 12 patients relapsed 97 (15%) and 7 patients died (9%). The 2020 ESGO postoperative risk groups but not molecular 98 subtypes were significantly associated with disease free survival (p < 0.001) and overall 99 survival (p = 0.005) (Figure 2 and 3). Survival curves according to the histological type, the 100 FIGO stage, the cytonuclear grades, the presence of lymphovascular emboli and the ESMO 101 2016 classification are available in the Supplementary Figure 1

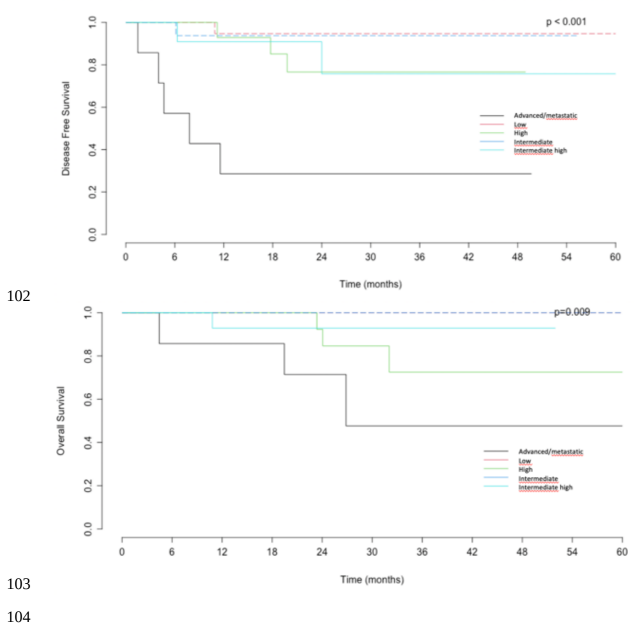


Figure 2: Recurrence-free survival (left) and overall survival (right) stratified by risk groups
 according to ESGO/ESTRO/ESP 2020 (in months). There were a significative difference
 between groups (p-value 0.001 and 0.009 respectively).

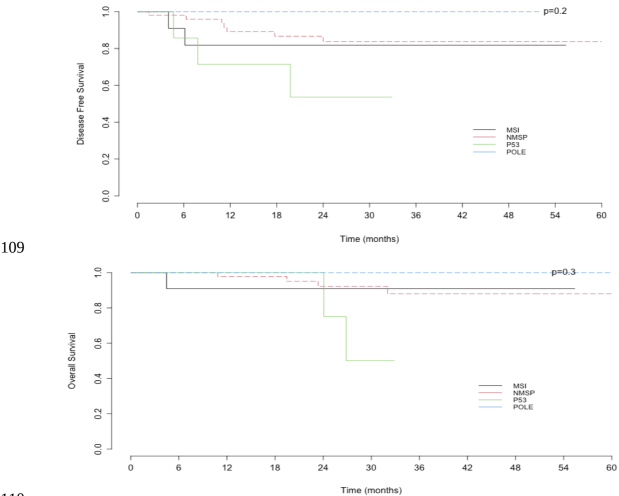


Figure 3: Recurrence-free survival (left) and overall survival (right) stratified by molecular
groups according to ESGO/ESTRO/ESP 2020 (in months). There was not a significant
difference between groups neither for recurrence – free nor overall survival (p-value =
0.2 and 0.3, respectively)

116 Relapses occurred on average at 9 months (1 - 23) and were localized as follow: locoregional 117 (vagina = 1, rectum = 3, parametrium = 2, pelvic non - specified = 6), lymph node (para-118 aortic, n = 6) and distant (peritoneum = 5, lung = 5, liver = 5 and bone = 1). Patients within 119 the p53 mutated group were the most likely to experience recurrence (37.5%, 3/8), followed by those MSI (18%, 2/11) and NSMP (12.5%, 7/56). None of the patients with a POLE 120 121 mutation recurred. Location of recurrence varied with the molecular subtype. In patients with 122 NSMP, 86% (6/7) had distant recurrence. All p53 patients had a distant relapse and one 123 patient also had locoregional recurrence. Patients in the MSI group had pelvic recurrences 124 without distant lesions.

- 125 Patients were most likely to die during follow up when p53 mutated (25%; 2/8) than when
- 126 part of the NSMP (7%, 4/56) or MSI (9%, 1/11) groups (p=0,3)
- 127 The distribution and characteristics of patients who recurred or died according to their risk
- 128 group or molecular status are presented in Tables 5 and Supplementary Table 4.
- 129

	Patients that	Patients that died	
	had recurrence	during follow - up	
	during follow -	(N=7)	
	up		
	(N=12)		
Molecular group			
POLE mutated (N=5)	0	0	
• MSI (N=11)	2 (18%)	1 (9%)	
• NSMP (N=56)	7 (12.5%)	4 (7%)	
• P53 mutated (N=8)	3 (37.5%)	2 (25%)	
ESGO Risk Group 2020			
• Low (N=24; 30%)	1 (8.3%)	0	
• Intermediate (N=20; 25%)	1 (8.3%)	0	
• Intermediate-high (N=14; 17%)	2 (16.7%)	2 (28.6%)	
• High (N=15; 19%)	3 (25%)	2 (28.6%)	
• Advanced/metastatic (N=7; 9%)	5 (41.7%)	3 (42.8%)	
ESGO Risk Group 2016			
• Low (N=18; 23%)	1 (8.3%)	0	
• Intermediate (N=12; 15%)	0	0	
• Intermediate-high (N=23; 28%)	2 (16.7%)	1 (14.3%)	
• High (N=20; 25%)	4 (33.3%)	3(42.8%)	

• Advanced/metastatic (N=7; 9%)	5 (41.7%)	3 (42.8%)		
Table 5: Distribution in molecular groups and prognosis of recurrence and death				

#### 133 **Discussion**

134 In this first report of a French cohort following the latest issue of ESGO guidelines for 135 endometrial cancer, around <sup>1</sup>/<sub>4</sub> of the patients were reclassified into a more accurate group of 136 prognosis. Molecular analysis and the latest ESGO classification had the most important 137 impact on patients initially classified at intermediate - high risk that were reclassified in intermediate (10/23) and in low (4/23) risk. The 2020 ESGO classification could have spared 138 139 secondary surgery for staging in 21.4% of patients classified at low or intermediate risk. A 140 total of 9 patients (11.3%) were over-treated according to the 2020 ESGO classification: six 141 patients in the low - risk group (4 received vaginal brachytherapy and 2 external 142 radiotherapy) and three in the intermediate risk group (3 received external irradiation and 1 143 received chemotherapy). None of the patients in our cohort were undertreated. The 2020 144 ESGO postoperative risk groups but not molecular subtypes were significantly associated 145 with disease free survival (p < 0.001) and overall survival (p = 0.005).

146 In our cohort, the molecular group distribution included a higher proportion of NSMP tumors 147 than the study of Kommoss et al.(17) and the meta-analysis of Raffone et al. (18) that 148 included 2818 patients, but with an equivalent proportion of POLEmut and p53 mutated. 149 Patients diagnosed either in the POLEmut group or in the p53 mutated group were little 150 represented (16.75%). These two groups are associated with extreme prognoses with very low 151 and high risk of recurrence / death, respectively. Such discrepancy in the repartition of the 152 molecular groups could be explained by the limited number of patients included, leading to an 153 over-representation of patients classified NMSP that have mild benefit of the molecular 154 subtype assessment. Regarding pathological characteristics, our findings were consistent with 155 previous study(12,19). In the studies by Talhouk et al, the POLEmut group was composed of 156 92% of endometrioid tumors including 58% of low-grade tumors and 58% with LVSI. Of 157 note, a significant proportion of POLEmut patients in our cohort had a myometrium 158 infiltration > 50% (80%) with 40% LVSI which are poor prognostic factors for recurrence

159 and survival. Our findings highlight the limited value of these parameters to assess the risk of 160 recurrence and advocate for molecular diagnosis use to decrease adjuvant therapies in patients 161 with excellent prognosis. On the other side, patients with p53 mutation usually have 162 numerous factors associated with bad prognosis with 75% serous tumors with 88% of LVSI 163 and infiltration of myometrium > 50%. In our cohort, significant survival differences existed 164 by ESGO 2020 groups but not by the different molecular groups. These results are conflicting 165 with those reported by Talhouk et al. (12,20,21) that found that compared to the "non -166 specific molecular profile" group, the risk was reduced by 77% in overall survival rates and 167 84% in recurrence-free survival rates for the POLE group, whereas the risk of death or 168 recurrence was multiplied by 3.29 and 2.19 times respectively for the p53 mutated group. The 169 main issue with molecular analyses remain the availability of the technic, limited by both the 170 cost and the time – consuming procedure. In the case of the POLE mutation research, High 171 Resolution Melting (HRM) screening of candidates for gene sequencing by NGS allows to 172 limit the final cost of the analysis, with a unit cost of 10 € for HRM against 120 € for Next 173 Generation sequencing (NGS). The time required for molecular biology analysis of POLE 174 mutations or microsatellites can be long when confirming cases in NGS or for microsatellite 175 analysis. McConechy et al. reported a concordance rate of more than 93% for the diagnostic 176 performance of immunohistochemistry and molecular biology (22). This problematic is 177 relative for p53 analysis as IHC has a high performance (Se: 90-100%, Sp: 94%, PPV: 98%, NPV: 74%)(3). When adjuvant therapy decision relies on molecular analysis, the delay to 178 179 obtain results is crucial. This is all the more important since these patients could exhibit bad 180 prognostic factors that could encourage clinicians to prescribe unindicated adjuvant therapies. 181 In our cohort, 9 patients had unindicated adjuvant treatment according to ESGO 2020 182 guidelines. In the PORTEC 3 study, side effects (neuropathy, alopecia, hematological, 183 gastrointestinal, auditory side-effects, pain etc.) were significantly more important in the 184 group treated by chemotherapy in combination with radiotherapy (23). In the study by De

Boer et al., toxicities and quality of life scores were higher (with more severe symptoms) in the radiochemotherapy group than in the radiotherapy alone group (p<0.001) and seemed to improve over time (non-significant results at 12 months from the end of treatment) (24). While survival has improved over the years, recent research has focused on quality of life after treatment. It seems essential to adapt adjuvant therapies to the molecular profile by limiting indications to selected subtypes

191 Molecular diagnosis also impacts surgical staging strategy. De kerdaniel et al. (25) 192 found surgical under-staging occured in 26% of the cases according to the 2010 guidelines. 193 Older patients (>70 years) were more often under-staged than younger patients (<70 years) 194 (p=0.037). In a recent meta-analysis, He et al. reported a 6% rate of positive lymph nodes (7 / 195 118) in POLE mutated patients and no significant association between the POLE mutated 196 status and the risk of lymph node involvement (OR 0.41; p=0.47) (26). These results are in 197 line with our findings that no POLEmut patients had lymph node involvement that could 198 benefit from less morbid procedure such as sentinel lymph node. Similarly, patients p53 199 mutated are at high risk of lymph node involvement and could benefit from per-operative 200 lymph node analysis to decide immediate complete lymphadenectomy, avoiding secondary 201 surgery. The search for MSI status by immunohistochemistry (more accessible and faster) 202 and the efficacy of antiPD-1 (27) treatments on these tumors in case of treatment failure 203 reinforces the necessity for MSI systematic testing. The RAINBO (Refining Adjuvant 204 treatment IN endometrial cancer Based On molecular profile) program, led by the 205 TransPORTEC study group, will bring interesting insight on the value on molecular subtype-206 based strategy.

Some limitations of our work deserve to be mentioned. This is a retrospective, observational, single-center study with a limited number of patients included. Our follow up could have been too short to diagnose some recurrences or death which might have bias the results. However, it has been demonstrated that the higher rate of recurrence is within the first 211 two years of follow up (28,29). The proportions of patients with POLE, p53 and MSI were 212 insufficient which have limited the full exploration of their prognostic impact. Eventually, a 213 significant number of patients did not undergo lymph node staging at all which clearly limit 214 the extent of our conclusions, especially as many cancer centers now perform sentinel lymph 215 node procedures even in high-risk patients. This also underline the benefit of molecular 216 subtype assessment was more likely underestimated in this cohort. One issue with molecular 217 diagnosis is that it depends of the quality of the DNA used, which is directly impacted by cold 218 ischemia duration, transport duration, delay prior fixation and the quality of the latter (29). 219 The retrospective inclusion of the cases limited the control of the conditions of conservation 220 of the slides. 221

222

223

## 225 Conclusion

226	Around one in 4 patients were reclassified in a more accurate prognostic group using
227	molecular diagnosis and the latest ESGO guidelines. which would significanly impact the use
228	of adjuvant therapies and help plan surgical strategy. Systematic molecular subtype
229	assessment will require easier and faster access to genetic plateforms to enable short circuits
230	useful to impact endometrial cancer strategy. Eventually, it will help plan therapeutic strategy
231	and decrease the use of adjuvant therapies to spare morbidity.
232	
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#### **Figures and table captions**

- 340 Figure 1: Flow chart of the study.
- Figure 2: Recurrence-free survival (left) and overall survival (right) of relapse risk groups
   according to ESGO/ESTRO/ESP 2020 (in months)
- Figure 3: Recurrence-free survival (left) and overall survival (right) of molecular groups by
   ESGO/ESTRO/ESP 2020 (in months).
- Table 1: Characteristics of the study population, treatments received by patients and
   ESMO/ESGO 2013 preoperative classification
- 347 Table 2: Histological and molecular characteristics of the study population
- Table 3: Number of patients classified into risk groups according to ESMO 2016 and ESGO
   2020 recommendations. Proportions are calculated based on the ESGO 2020 group size.
- Table 4: Node involvement by molecular group and ESGO 2020 risk group. Pelvic lymph
  node involvement may be associated with para-aortic lymph node involvement. Paraaortic involvement may be associated with pelvic involvement. A total of 12 patients
  were classified as N+. Here are calculated the proportions of lymph node involvement
  according to the size of the molecular groups or the 2020 risk groups
- 355 Table 5: Distribution in molecular groups and prognosis of recurrence and death