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Title: Practical guidance for mismatch repair-deficiency testing in endometrial cancer

Authors

E. Stelloo^a, A.M.L. Jansen^{a,b}, E.M. Osse^a, R.A. Nout^c, C.L. Creutzberg^c, D. Ruano^a,
D.N. Church^{d,e}, H. Morreau^a, V.T.H.B.M. Smit^a, T. van Wezel^a, T. Bosse^a

Affiliations

^a Department of Pathology, Leiden University Medical Centre, Leiden, The Netherlands

^b Department of Human Genetics, Leiden University Medical Centre, Leiden, The Netherlands

^c Department of Medical and Radiation Oncology, Leiden University Medical Centre, Leiden, The Netherlands

^d Cancer Genomics and Immunology Group, Oxford Centre for Cancer Gene Research, The Wellcome Trust Centre for Human Genetics, University of Oxford. Roosevelt Drive, Oxford, UK

^e Oxford Cancer Centre, Churchill Hospital. Old Road, Oxford, UK

Corresponding author

Dr. Tjalling Bosse

Department of Pathology

Leiden University Medical Centre

PO Box 9600

2300 RC Leiden

The Netherlands

Telephone: 0031(0)715266628

Fax: 0031(0)715266952

Email: T.Bosse@lumc.nl

Abstract

Background: Mismatch repair (MMR)-deficiency analysis is increasingly recommended for all endometrial cancers, as it identifies Lynch syndrome-patients, and is emerging as a prognostic classifier to guide adjuvant treatment. The aim of this study was to define the optimal approach for MMR-deficiency testing and to clarify discrepancies between microsatellite instability (MSI) analysis and immunohistochemical (IHC) analysis of MMR protein expression.

Patients and Methods: 696 endometrial cancers were analyzed for MSI (pentaplex panel) and MMR protein expression (IHC). Agreement between methodologies was calculated using Cohen's Kappa. *MLH1* promoter hypermethylation, dinucleotide microsatellite markers and somatic MMR and *POLE* exonuclease domain (EDM) gene variants (using next-generation/Sanger sequencing) were analyzed in discordant cases.

Results: MSI was found in 180 patients. Complete loss of expression of one or more MMR proteins was observed in 196 cases. A PMS2- and MSH6-antibody panel detected all cases with loss of MMR protein expression. The results of MSI and MMR protein expression were concordant in 655/696 cases ($\kappa=0.854$, $P<0.001$). Ambiguous cases ($n=41$, 6%) included: subclonal loss of MMR protein expression ($n=18$), microsatellite stable or MSI-low cases with loss of MMR protein expression ($n=20$), and MSI-low or MSI-high cases with retained MMR protein expression ($n=3$). Most of these cases could be explained by *MLH1* promoter hypermethylation. Five of seven cases with solitary loss of PMS2 or MSH6 protein expression carried somatic gene variants. Two MSI-high cases with retained MMR protein expression carried a *POLE*-EDM variant.

Conclusion: MSI and IHC analysis are highly concordant in endometrial cancer. This holds true for cases with subclonal loss of MMR protein expression.

Discordant MMR-proficient/MSI-high cases (<1%), may be explained by *POLE*-EDM variants.

Keywords

endometrial cancer, microsatellite instability, mismatch repair

Key message

Based on comprehensive analysis of 696 endometrial cancers, we provided sufficient evidence that a immunohistochemical approach is both practical and feasibly for mismatch repair deficiency testing in routine clinical pathology. Unusual findings, such as suclonal loss of mismatch repair expression mainly due to *MLH1* promoter hypermethylation, were described in detail.

Clinical Trial tissue samples

Translational research was performed on tissue samples from two randomized clinical trials (PORTEC-1 and PORTEC-2). PORTEC-2 trial register number is ISRCTN16228756 and PORTEC-1 was conducted before time of trial registries. Both trials were supported by grants from the Dutch Cancer Society (CKTO 90-01 and CKTO 2001-04).

Introduction

A defect in DNA mismatch repair (MMR) leads to the accumulation of mismatches, insertions and deletions in repeated sequences - a phenomenon named microsatellite instability (MSI). Approximately 20-30% of sporadic endometrial cancers (ECs) display MSI as a consequence of somatic promoter hypermethylation and silencing of *MLH1*. [1] Defective MMR due to pathogenic germline variants in MMR genes causes Lynch syndrome (LS), a tumor predisposition syndrome that accounts for 2% of ECs. [2]

Determination of MMR-deficiency in EC may be important for several reasons. First, recent studies have suggested that tumor molecular features, including MMR-deficiency, may improve prognostication and help guide adjuvant therapy for EC patients. [3, 4] Second, accurate assessment of MMR-deficiency is essential to identify patients with EC caused by LS. However in contrast to colorectal cancer, where consensus guidelines for MMR-deficiency testing have been published, [5] there is no general agreement on screening EC patients for LS. [6, 7] Finally, recent studies have shown that MMR-deficiency in colorectal and urothelial cancer is predictive of response to immunotherapy, [8, 9] suggesting that MMR-deficient ECs may also benefit from these therapeutics.

MMR-deficiency can be detected by either MSI analysis and/or immunohistochemical (IHC) staining, typically for four MMR proteins. The National Cancer Institute microsatellite panel was optimized and correlated with IHC analysis (~95%) to detect MMR-deficiency in colorectal cancer. [10, 11] IHC alone has become standard practice in multiple institutions. Experience in this setting is that while some tumors show uniform and widespread loss of MMR protein expression, cases with subclonal loss of MMR protein expression are also observed. [11, 12] Such cases present with two populations of tumor cells; one with retained expression, and

another with abrupt and complete regional loss of MMR protein expression.[12] Small studies have shown high agreement between MSI and loss of MMR protein expression in EC,[13-15] while others have described subclonal loss of MMR protein expression.[16-20] However, studies identifying the frequency of such staining patterns in large patient series are sparse.

In this study, we sought to establish the optimum method for MMR-deficiency testing by comparison of MSI with IHC analysis in a large series of ECs. We also investigated the frequency of subclonal loss of MMR protein expression and the number of potential LS cases. Cases showing disagreement between methodologies and those with subclonal loss of MMR protein expression were further characterized.

Methods

Study population

The population comprised, 854 ECs from the PORTEC-1 and -2 clinical trials based on availability of formalin-fixed paraffin-embedded slides and sufficient tumor material for DNA isolation.[21, 22] Further details are summarized in Supplementary Table S1 and Supplementary methods.

MSI assay

DNA was isolated as previously described.[23] In cases with subclonal loss of MMR protein expression, tissue sections were used to microdissect the differentially expressed tumor areas. Tumor MSI status was determined as previously reported (Supplementary methods).[4] Tumors initially classified as MSS or MSI-L with concomitant loss of MMR protein expression underwent evaluation of three dinucleotide repeat markers,[24] and reclassified as MSI-H if instability was detected at two dinucleotide markers.

IHC analysis

IHC staining for MLH1, PMS2, MSH2, and MSH6 was performed on all tumors in which MSI status was successfully determined (Supplementary methods). The slides were evaluated in three categories as retained, loss and subclonal loss of protein expression with stromal- and/or lymphocytic cells as internal controls.[16] The cases with subclonal loss of protein expression were re-evaluated to determine the percentage of tumor cells with loss of MMR expression.

Methylation-specific PCR for *MLH1*

Tumors with loss of MLH1 protein expression underwent testing for hypermethylation status of the *MLH1* 5' regulatory region by methylation-specific PCR, as previously described.[25]

Somatic variant screening

Subject to DNA availability and quality, tumors in which the results of MSI analysis and MMR protein expression were discordant underwent targeted next generation sequencing (NGS) of MMR and *POLE* genes using the Ion Proton™ System (ThermoFisher, MA, USA) as previously described (Supplementary methods).[26] Three additional cases were similarly analyzed using the Ion AmpliSeq Comprehensive Cancer Panel (ThermoFisher) at The Wellcome Trust Center for Human Genetics, UK. Frameshift variants in the polycytosine tract in exon 5 of *MSH6* were analyzed using Sanger sequencing.[27]

Results

Combined analysis of MMR protein expression and MSI was possible in 696 (81%) ECs (Supplementary Table S1). The frequencies of MSS, MSI-H and MSI-L were 74%, 24% and 2%, respectively. Among the 516 tumors assessed as MSS, 496 (96%) showed retained expression of all four MMR proteins (Table 1). The remaining twenty MSS cases showed loss of MMR protein expression as follows: combined MLH1 and PMS2 loss ($n=14$), combined MSH2 and MSH6 loss ($n=3$), solitary MSH6 loss ($n=3$, Figure 1A-D). Of the 11 cases assessed as MSI-L, six displayed combined loss of MLH1 and PMS2 expression (complete in four cases, subclonal in two cases), two cases showed solitary loss of PMS2, a further two cases had solitary MSH6 loss and one case retained expression of all four MMR proteins.

The majority of MSI-H cases (130 of 169, 77%) showed complete loss of MLH1 and PMS2 expression (Table 1). Sporadic MSI due to *MLH1* hypermethylation was observed in 97% of these 130 MSI-H cases. Eight MSI-H cases showed areas of subclonal loss of MLH1 and PMS2 (Figure 1E-H), and six cases displayed subclonal loss of MSH6 in addition to complete loss of MLH1 and PMS2 protein expression (Figure 1I-L). In ten cases (6%), combined loss of MSH2 and MSH6 protein expression was observed. The remaining MSI-H tumors showed solitary loss of PMS2 ($n=8$) or MSH6 ($n=5$) protein expression, or retained expression of all MMR proteins ($n=2$).

Overall, concordance between MSI and IHC analysis was observed in 655 of 696 cases (94%, kappa=0.854; 95%CI 0.811-0.897, $P<0.001$). A PMS2- and MSH6-antibody panel was as effective as the four-antibody panel in detecting MMR protein abnormalities. Twenty-seven concordant cases without *MLH1* promoter hypermethylation were identified as potential LS, but the underlying defect was not further tested. Discordant cases ($n=41$, 6%) included: subclonal loss of MMR

expression ($n=18$), MSS or MSI-low cases with loss of MMR expression ($n=20$), and MSI-low or MSI-high cases with retained MMR protein expression ($n=3$). Details on the sample analysis of discordant cases are shown in Supplementary Figure S1.

All cases with subclonal loss of MMR protein expression ($n=18$) were evaluated in more detail by analyzing MSI in mono- and dinucleotide markers, *MLH1* promoter hypermethylation and somatic MMR- and *POLE*-exonuclease domain (EDM) variants in microdissected tumor areas (Table 2). Among 16 tumors with subclonal *MLH1* and *PMS2* loss, 14 had areas of differential expression that were sufficiently large to permit microdissection. Among these, MSI testing of microdissected areas was concordant with IHC analysis in 11 cases; tumor areas with retained MMR expression were MSS, whereas areas with loss of MMR expression showed MSI-H. A further three tumors showed microsatellite stability of markers in microdissected areas regardless of MMR protein expression (case 13-15, Table 2). All fourteen cases were found to have somatic promoter hypermethylation of *MLH1*. One case with subclonal loss of *MSH2* and *MSH6* protein expression showed microsatellite stability in the differently expressed areas (case 17, Table 2). Unfortunately, both cases with subclonal loss of *MSH2* and *MSH6* protein expression had limited DNA available, and could not be analyzed in more detail.

Analysis of microdissected material from the six MSI-H cases with subclonal loss of *MSH6* in addition to complete *MLH1* and *PMS2* protein loss demonstrated frameshift variants in the polycytosine tract of *MSH6* in areas with *MSH6* loss and stable polycytosine tracts in areas with retained *MSH6*. Five of these cases displayed *MLH1* promoter hypermethylation.

We proceeded to perform detailed analysis of the 23 cases with discordant MSI status and MMR protein expression by examination of dinucleotide markers, *MLH1* promoter hypermethylation and/or NGS of the MMR- and *POLE* genes (Table

3). The two MSI-H cases (case 19-20) with retained MMR protein expression had a *POLE*-EDM variant, p.(V411L), and p.(A428T). The *POLE*-EDM p.(V411L) mutant case also harbored a truncating (p.(R563*)) and missense p.(R107W) variant in *PMS2*. The solitary MSI-L case with retained MMR protein expression (case 21) showed stability in dinucleotide markers and no somatic MMR or *POLE*-EDM gene variants.

One of 12 cases classified as MSS or MSI-L despite combined loss of MLH1 and PMS2 protein expression, showed mobility shifts in the dinucleotide markers (case 24, Table 3). Analysis of the *MLH1* promoter was successful in 11 of these cases, and revealed promoter hypermethylation in ten cases, while the single case lacking *MLH1* promoter hypermethylation was found to harbor a pathogenic *POLE*-EDM variant, p.(P286R).

Of the two MSI-L tumors with solitary PMS2 loss (case 34-35, Table 3), only one had sufficient DNA quality for further analysis. This confirmed MSI in the dinucleotide markers and revealed two likely pathogenic somatic *PMS2* variants, a start loss (p.(Met1?)) and a frameshift variant (p.(Val302Thrfs*4)).

Four of five cases classified as MSS/MSI-L with solitary MSH6 loss were informative for further analysis. All four showed microsatellite stable dinucleotide markers. Three tumors carried two ($n=1$) or one ($n=2$) pathogenic *MSH6* variants, while one tumor carried one somatic VUS predicted to affect function by two out of three protein prediction software used (Table 3). Case 41 with loss of MSH2 and MSH6 protein expression and a MSS phenotype had limited DNA, and was therefore excluded for further analysis.

Discussion

Accurate identification of MMR-deficiency in EC may be important to identify patients with a higher risk of recurrence,[3, 4, 28] and those whose tumors may be a consequence of LS. Similarly to two small studies, we demonstrated high agreement (94%) between MSI and IHC analysis in 696 ECs.[13, 14] Most discordant cases involved loss of MMR protein expression and a MSS/MSI-L phenotype and could be explained by *MLH1* promoter hypermethylation or MMR variants. In addition, subclonal loss of MMR protein expression generally corresponded to *MLH1* promoter hypermethylation and subclonal MSI within microdissected area of the tumor.

Importantly, the present study demonstrated that <3% of cases displayed subclonal loss of MMR protein expression. The fact that MSI and *MLH1* promoter hypermethylation were commonly found in areas with subclonal loss of *MLH1* and *PMS2* protein expression indicates sporadic intratumor heterogeneity.[16, 18, 19] However, *MLH1* germline epimutations cannot be totally excluded.[2] Subclonal loss of *MSH6* expression, either in conjunction with or without *MSH2* protein expression was also previously observed in EC but the underlying molecular mechanisms remain unclear.[12, 20] In accordance with our findings, subclonal loss of *MSH6* in cases with complete loss of *MLH1* and *PMS2* protein expression has been related to secondary MSI events in *MSH6*. [20, 29] Although numbers are limited, subclonal loss of MMR protein expression is not associated with LS.

Our data suggest that cases with subclonal loss of MMR protein expression are best classified as MMR-deficient, even though the areas with retained expression are MSS. With regard to MMR-deficiency as a prognostic or predictive marker, it remains to be determined whether subclonal loss of MMR protein expression has the same biological behavior as tumors with MMR-proficiency. In view of the limited numbers of cases with subclonal loss of MMR protein expression (~3%),

(inter)national collaborations are essential to obtain sufficient cases for such an analysis. Pending these future studies, we suggest for uniformity to classify tumors with 10% subclonal loss of MMR protein expression, as being MMR-deficient.

The interpretation of MSI-L cases remains controversial in EC and it is uncertain whether such cases are best considered as MSS or MSI. Similar numbers of DNA slippage events were observed in MSS and MSI-L ECs.[30] To date, no extensive research on the clinical implications of MSI-L in ECs has been performed, and the number of MSI-L cases in our study ($n=11$) was too low to permit such an analysis as well. However, most of these showed loss of MMR protein expression and would generally be regarded as abnormal by strategies that rely on IHC alone. Noteworthy, several studies have also shown MSI-L and MSS in association with loss of MMR expression and/or pathogenic germline MMR variants.[13, 14, 31]

Our study shows high agreement between IHC and MSI analysis, but not 100%. Of note, other assessments of DNA defects by IHC analysis e.g. *HER2* gene amplification only reaches 69-98% agreement.[32] Assessment of MMR protein expression is preferred over MSI analysis for the following reasons: lower costs, widely available, and determination of affected MMR gene. Our findings confirm the utility of testing MMR-deficiency using a PMS2- and MSH6-IHC approach,[33] which can be followed by MLH1- and MSH2-IHC in case loss of PMS2 or MSH6 was observed. To overcome suboptimal fixation, drawback of IHC analysis, pathologists can rely on IHC analysis in pre-operative EC specimen.[23] IHC with standard well accepted techniques would appear adequate to identify EC patients with LS and to serve as a biomarker for trials of EC patients harboring MMR-deficiency.

It is debatable whether not screening for germline MMR variants is a limitation of this study. Of note, 5% of all cases in this study can be classified as potential LS (no *MLH1* promoter hypermethylation). Somatic screening of the discordant cases

did show somatic variants but not in all cases. However, we cannot exclude the possibility of missed large genomic rearrangements within the tested genes, which is a limitation of NGS. Further analysis would improve understanding the molecular basis of the discordant cases, however, this study did not aim to determine the sensitivity and specificity of the two methodologies to identify LS. MSI and IHC analysis are highly concordant therefore germline testing is not needed to conclude which approach is best suitable for identifying patients with LS.

In conclusion, MSI and IHC analysis are highly concordant, also in cases with subclonal loss of MMR expression, therefore, an IHC approach is sufficient for determining MMR-deficiency in EC. Pathologists should be aware of the MMR protein expression patterns, including subclonal loss, to ensure correct classification in daily diagnostic pathology.

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Disclosure/Conflict of interest

The authors have declared no conflicts of interest.

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Legends to tables and figures

Figure 1. Representative images of MMR protein expression in EC. MMR protein expression of a MSS case with subclonal loss of MLH1 and PMS2 protein expression (A-D, case 13), a MSI-H case with complete loss of MLH1 and PMS2 and subclonal loss of MSH6 protein expression (E-H), a MSS case with loss of MSH6 protein expression (I-L, case 38). A-E-I) MLH1 protein expression, B-F-J) PMS2 protein expression, C-G-K) MSH2 protein expression and D-H-L) MSH6 protein expression. Scale bar represents 50 μ M.

Table 1. Details on the MSI status and MMR protein expression in early-stage EC ($n=696$).

Table 2. Details on ECs with subclonal loss of MMR protein expression.

Table 3. Details on ECs with discordant MSI status and MMR protein expression.

Supplementary Figure S1. Flowchart of sample analyses of cases with discordant MSI status and MMR protein expression. Explained cases are further described in the white colored lines and unexplained/failed cases in the grey colored lines in Table 2 and Table 3.

MSI – microsatellite instability, IHC – immunohistochemical, MMR – mismatch repair, MSS – microsatellite stable, MSI-L/H – microsatellite unstable with low- or high-frequency.

Supplementary Table S1. Clinicopathological characteristics according to the MSI status in early-stage EC ($n=696$).