Protocol for the Examination of Specimens From Patients With Carcinoma of the Endometrium

Based on AJCC/UICC TNM, 7th edition, and FIGO 2014 Annual Report
Protocol web posting date: January 2016

Procedure
• Hysterectomy

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CAP Endometrium Protocol Revision History

Version Code
The definition of version control and an explanation of version codes can be found at www.cap.org (search: cancer protocol terms).

Version: Endometrium 3.3.0.0

Summary of Changes
The following changes have been made since the October 2013 release.

The following data elements were modified:
  Specimen Integrity
  Tumor Site
  Tumor Size (changed from required to optional)
  Histologic Type
  Myometrial Invasion
  Tumor Involvement of Cervix
  Peritoneal Ascitic Fluid
  Lymph-Vascular Invasion
  Regional Lymph Nodes
  Distant Metastasis (changed to required only if confirmed pathologically)
  Additional Pathologic Findings

The following data element was added:
  FIGO Stage (not required)
## Surgical Pathology Cancer Case Summary

Protocol web posting date: January 2016

### ENDOMETRIUM: Hysterectomy, With or Without Other Organs or Tissues

Select a single response unless otherwise indicated.

#### Specimen (select all that apply)
- [ ] Uterine corpus
- [ ] Cervix
- [ ] Right ovary
- [ ] Left ovary
- [ ] Right fallopian tube
- [ ] Left fallopian tube
- [ ] Right parametrium
- [ ] Left parametrium
- [ ] Vaginal cuff
- [ ] Omentum
- [ ] Other (specify): ___________________________
- [ ] Not specified

#### Procedure (select all that apply) (Note A)
- [ ] Supracervical hysterectomy
- [ ] Simple hysterectomy
- [ ] Radical hysterectomy
- [ ] Right oophorectomy
- [ ] Left oophorectomy
- [ ] Right salpingectomy
- [ ] Left salpingectomy
- [ ] Right salpingo-oophorectomy
- [ ] Left salpingo-oophorectomy
- [ ] Bilateral salpingo-oophorectomy
- [ ] Omentectomy
- [ ] Peritoneal biopsies
- [ ] Peritoneal washing
- [ ] Other (specify): ___________________________
- [ ] Not specified

#### Lymph Node Sampling (select all that apply)
- [ ] Performed
  - [ ] Pelvic lymph nodes
  - [ ] Para-aortic lymph nodes
  - [ ] Other (specify): ___________________________
- [ ] Not performed
- [ ] Not known

#### Specimen Integrity (Note A)
- [ ] Morcellated hysterectomy specimen
- [ ] Intact hysterectomy specimen
- [ ] Other (specify): ___________________________

+ Data elements preceded by this symbol are not required. However, these elements may be clinically important but are not yet validated or regularly used in patient management.
+ Tumor Site (select all that apply)
+ ___ Anterior endometrium
+ ___ Posterior endometrium
+ ___ Fundus
+ ___ Lower uterine segment
+ ___ Other (specify): ________________________________

+ Tumor Size
+ Greatest dimension: ___ cm
+ Additional dimensions: ___ x ___ cm
+ ___ Cannot be determined (explain): ________________________________

Histologic Type (select all that apply) (Note B)
___ Endometrioid adenocarcinoma
___ Endometrioid adenocarcinoma with squamous differentiation
___ Endometrioid adenocarcinoma, villoglandular
___ Endometrioid adenocarcinoma, secretory
___ Mucinous adenocarcinoma
___ Serous endometrial intraepithelial carcinoma
___ Serous carcinoma
___ Clear cell carcinoma
___ Carcinoïd
___ Small cell carcinoma
___ Large cell neuroendocrine carcinoma
___ Mixed cell carcinoma (specify types and percentages): ________________________________
___ Undifferentiated carcinoma
___ Dedifferentiated carcinoma
___ Carcinosarcoma (malignant mixed Mullerian tumor)
   ___ Homologous type
   ___ Heterologous type (specify heterologous elements): ________________________________
   ___ Epithelial component(s) (specify epithelial cell types and percentages): ________________
___ Other (specify): ________________________________

Histologic Grade (Note C)
International Federation of Gynecology and Obstetrics (FIGO) Grading System
(applies to endometrioid and mucinous adenocarcinomas only):
___ FIGO grade 1
___ FIGO grade 2
___ FIGO grade 3

For other carcinomas:
___ G1: Well differentiated
___ G2: Moderately differentiated
___ G3: Poorly differentiated
___ Other (specify): ________________________________
___ Not applicable
Myometrial Invasion (select all that apply) (Note D)

___ Not identified
___ Present
   Depth of invasion: ___ mm

OR, if exact depth of invasion cannot be determined, state:
___ Extent of myometrial invasion cannot be determined (explain): _______________________
___ <50% myometrial invasion
___ ≥50% myometrial invasion
Myometrial thickness: ___ mm
___ Myometrial thickness cannot be determined (explain): _______________________

Tumor Involvement of Cervix (Note E)

___ Not involved
___ Involves cervix without stromal invasion
___ Invasion of cervical stromal connective tissue
___ Cannot be determined (explain): _______________________

Extent of Involvement of Other Organs (select all that apply)

___ Right ovary
   ___ Involved
   ___ Not involved
   ___ Other (explain): _______________________
___ Left ovary
   ___ Involved
   ___ Not involved
   ___ Other (explain): _______________________
___ Right fallopian tube
   ___ Involved
   ___ Not involved
   ___ Other (explain): _______________________
___ Left fallopian tube
   ___ Involved
   ___ Not involved
   ___ Other (explain): _______________________
+ ___ Vagina
   + ___ Involved
   + ___ Not involved
+ ___ Right parametrium
   + ___ Involved
   + ___ Not involved
+ ___ Left parametrium
   + ___ Involved
   + ___ Not involved
+ ___ Omentum
   + ___ Involved
   + ___ Not involved
+ ___ Rectal wall
   + ___ Involved
   + ___ Not involved
+ ___ Bladder wall
   + ___ Involved
   + ___ Not involved
+ ___ Pelvic wall
   + ___ Involved
   + ___ Not involved

+ Data elements preceded by this symbol are not required. However, these elements may be clinically important but are not yet validated or regularly used in patient management.
+ ___ Bladder mucosa and/or bowel mucosa
  + ___ Involved
  + ___ Not involved
+ ___ Other (specify): _________________________________

+ Peritoneal Ascitic Fluid (Note F)
+ ___ Not performed/unknown
+ ___ Negative for malignancy (normal/benign)
+ ___ Atypical and/or suspicious (explain): _____________________________
+ ___ Malignant (positive for malignancy)
+ ___ Unsatisfactory/nondiagnostic (explain): _____________________________

+ Margins (Note G)
+ ___ Cannot be assessed
+ ___ Uninvolved by invasive carcinoma
  + Distance of invasive carcinoma from closest margin: ___ mm
  + Specify margin: _____________________________
+ ___ Involved by invasive carcinoma
  + Specify margin(s): _____________________________

Lymph-Vascular Invasion (Note H)
___ Not identified
___ Present
___ Cannot be determined

Pathologic Staging (pTNM) (Note I)

TNM Descriptors (required only if applicable) (select all that apply)
___ m (multiple primary tumors)
___ r (recurrent)
___ y (posttreatment)

Primary Tumor (pT)
___ pTX: Primary tumor cannot be assessed
___ pT0: No evidence of primary tumor
___ pT1a: Tumor limited to endometrium or invades less than one-half of the myometrium
___ pT1b: Tumor invades greater than or equal to one-half of the myometrium
___ pT2: Tumor invades stromal connective tissue of the cervix, but does not extend beyond uterus
___ pT3a: Tumor involves serosa and/or adnexa (direct extension or metastasis)
___ pT3b: Vaginal involvement (direct extension or metastasis) or parametrial involvement
___ pT4: Tumor invades bladder mucosa and/or bowel mucosa (bullous edema is not sufficient to classify a tumor as T4)

Regional Lymph Nodes (pN) (select all that apply) (Note I)
+ Modifier
+ ___ (sn)

Category (pN)
___ pNX: Cannot be assessed
___ pN0: No regional lymph node metastasis
___ pN1: Regional lymph node metastasis to pelvic lymph nodes
___ pN2: Regional lymph node metastasis to para-aortic lymph nodes, with or without positive pelvic lymph nodes
___ No nodes submitted or found

+ Data elements preceded by this symbol are not required. However, these elements may be clinically important but are not yet validated or regularly used in patient management.
Pelvic lymph nodes:
___ No pelvic nodes submitted or found

Number of Pelvic Lymph Nodes Examined
Specify: ___
___ Number cannot be determined (explain): ______________________

Number of Pelvic Lymph Nodes Involved
Specify: ___
___ Number cannot be determined (explain): ______________________

Para-aortic lymph nodes:
___ No para-aortic nodes submitted or found

Number of Para-aortic Lymph Nodes Examined
Specify: ___
___ Number cannot be determined (explain): ______________________

Number of Para-aortic Lymph Nodes Involved
Specify: ___
___ Number cannot be determined (explain): ______________________

Other lymph nodes:
Specify site: ______________________

Number of Other Lymph Nodes Examined
Specify: ___
___ Number cannot be determined (explain): ______________________

Number of Other Lymph Nodes Involved
Specify: ___
___ Number cannot be determined (explain): ______________________

Distant Metastasis (pM) (required only if confirmed pathologically in this case)
___ pM1: Distant metastasis (includes metastasis to inguinal lymph nodes, intraperitoneal disease, or lung, liver, or bone metastasis. It excludes metastasis to para-aortic lymph nodes, vagina, pelvic serosa, or adnexa)
Specify site(s), if known: __________________________

+ FIGO Stage
+ I: Tumor confined to the corpus uteri
  + ___ IA: No or less than half myometrial invasion
  + ___ IB: Invasion equal to or more than half of the myometrium
+ ___ II: Tumor invades cervical stroma, but does not extend beyond the uterus
  + ___ IIIA: Tumor invades the serosa of the corpus uteri and/or adnexae
  + ___ IIIB: Vaginal involvement and/or parametrical involvement
+ ___ IIIC: Metastases to pelvic and/or para-aortic lymph nodes
  + ___ IIIC1: Positive pelvic nodes
  + ___ IIIC2: Positive para-aortic nodes with or without positive pelvic lymph nodes
+ ___ IV: Tumor invades bladder and/or bowel mucosa, and/or distant metastases
  + ___ IVA: Tumor invasion of bladder and/or bowel mucosa
  + ___ IVB: Distant metastasis, including intraabdominal metastases and/or inguinal nodes

+ Data elements preceded by this symbol are not required. However, these elements may be clinically important but are not yet validated or regularly used in patient management.
+ Additional Pathologic Findings (select all that apply) (Note J)
+ ___ None identified
+ ___ Hyperplasia without atypia
  + ___ Simple without cytologic atypia
  + ___ Complex without cytologic atypia
+ ___ Atypical hyperplasia
  + ___ Simple
  + ___ Complex
+ ___ Endometrial intraepithelial neoplasia (EIN)
+ ___ Other (specify): ___________________________

+ Ancillary Studies
  Note: For reporting molecular testing, immunohistochemistry, and other cancer biomarker testing results, the CAP endometrium biomarker template should be used. Pending biomarker studies should be listed in the Comments section of this report.

+ Clinical History (select all that apply) (Note K)
+ ___ Lynch syndrome
+ ___ Other (specify): ___________________________

+ Comment(s)
Explanatory Notes

A. Specimen Type
In rare occasions when an endometrial carcinoma is not suspected, the pathologist may receive a supracervical hysterectomy specimen removed by laparoscopy. The type of procedure should be recorded. It has been reported that hysterectomies performed using certain laparoscopic techniques result in the finding of venous tumor emboli that are likely to be iatrogenic.¹ The significance of morcellation techniques in unsuspected endometrial cancer cases is not known, but there is theoretical risk of spreading tumor cells to the pelvis and peritoneal cavity. Therefore, reporting of such a procedure is important (and listed under Specimen Integrity in the case summary).

B. Histologic Type
For consistency in reporting, the histologic classification of endometrial carcinoma and hyperplasia proposed by the World Health Organization (WHO), shown below, is recommended.²

Endometrioid adenocarcinoma
Endometrioid adenocarcinoma with squamous differentiation
Endometrioid adenoacarcinoma, villoglandular
Endometrioid adenocarcinoma, secretory
Mucinous adenocarcinoma
Serous endometrial intraepithelial carcinoma
Serous carcinoma
Clear cell carcinoma
Low-grade neuroendocrine tumor (carcinoid)
High-grade neuroendocrine carcinoma (small cell neuroendocrine carcinoma)
High-grade neuroendocrine carcinoma (large cell neuroendocrine carcinoma)
Mixed cell carcinoma
Undifferentiated carcinoma
Dedifferentiated carcinoma
Carcinosarcoma (malignant mixed mullerian tumor)
  Homologous type
  Heterologous type
  Epithelial components

¹ The term mixed carcinoma should only be used when 2 or more distinct subtypes of endometrial carcinoma are identified, each representing more than 10% of the tumor. Optimally, the diagnosis is made on examination of a hysterectomy specimen, but if only a smaller specimen is available, any amount of a second tumor category suffices for the diagnosis. When a carcinoma is classified as “mixed,” the major and minor types and their relative proportions should be specified. High-grade tumors with ambiguous features should be classified as “carcinoma, subtype cannot be determined”; however, this is a very infrequent situation and every effort should be made to subclassify such tumors. It should be noted that for mixed endometrioid and serous carcinomas, studies have found variable results regarding tumor behavior based on percentage of the serous component. Some studies have found that tumors with >25% serous component behave like pure serous carcinomas, whereas other studies have shown that tumors with <10% serous component also behave like pure serous carcinomas.³,⁴ It is important to be aware that some serous carcinomas may display a glandular architecture.⁵ Thus, when a gland-forming endometrial carcinoma shows high-grade nuclear features, the diagnosis of serous carcinoma should be considered. Finally, the term endometrial intraepithelial carcinoma is discouraged because it is not uncommon for these lesions to be associated with extraterine spread.⁶,⁷ Instead, the term serous endometrial intraepithelial carcinoma should be used.

In addition, carcinosarcoma (also referred to as malignant müllerian mixed tumor [MMMT]) has been added to the above list of tumors in the case summary. Carcinosarcoma is a high-grade endometrial neoplasm that is staged like endometrial carcinomas because it is thought to represent a high-grade metaplastic carcinoma. The diagnosis of carcinosarcoma requires presence of both a malignant epithelial component and a malignant mesenchymal (sarcomatous) component in the neoplasm, which should not merge.
Proposed Criteria Defining Endometrial Carcinoma versus Endometrial Hyperplasia

1. Irregular infiltration of glands associated with an altered fibroblastic stroma (desmoplastic response), or
2. Confluent glandular pattern (cribriform growth), or
3. Extensive papillary growth pattern, or
4. Severe cytologic atypia (G3 nuclear atypia)

Some investigators have offered specific measurements to assess confluent glandular growth more objectively. Kurman and Norris proposed 1.9 mm as a cutoff, whereas Longacre and colleagues proposed 30% as a cutoff. Extensive papillary growth has also been quantitatively measured in 1 study and defined to be at least 4.2 mm in diameter to warrant the diagnosis of carcinoma. However, it is important to note that different investigators did not find these parameters to have the same predictive value.

C. Histologic Grading

The International Federation of Gynecology and Obstetrics (FIGO) grading system for carcinomas of the uterine corpus is only officially designated for endometrioid carcinomas and is based on architectural features as follows:

- Grade 1: 5% or less nonsquamous solid growth pattern
- Grade 2: 6% to 50% nonsquamous solid growth pattern
- Grade 3: >50% nonsquamous solid growth pattern

Notable nuclear atypia, which exceeds that which is routinely expected for the architectural grade, increases the tumor grade by 1.

In addition, the following guidelines should be used in grading:

1. The squamous component of endometrioid adenocarcinoma should not be graded because the degree of differentiation typically parallels that of the glandular component.
2. Because mucinous carcinomas are closely related to endometrioid carcinomas, they can be graded by the same criteria.
3. Serous, clear cell, transitional, small cell, undifferentiated carcinomas, and carcinosarcomas are generally considered to be high grade and it is not recommended to assign a FIGO grade to these tumor types. When the case summary is being completed, these should be designated as “not applicable” for histologic grade.
4. In mixed carcinomas, the highest grade should be assigned.

D. Myometrial Invasion

Assessing myometrial invasion may be difficult. Depth of invasion should be measured from the endomyometrial junction to the deepest point of invasion, which may not be easy because the endomyometrial junction in normal conditions is often irregular. In these cases, it is always helpful to look for compressed, nonneoplastic endometrial glands at the nearby endomyometrial junction or even at the base of the tumor. Carcinoma involving adenomyotic foci should not be interpreted as invasive carcinoma. However, the distinction between invasive carcinoma and carcinoma involving adenomyosis may be difficult, because in some cases invasive carcinoma may not elicit stromal response. In the absence of adenomyosis uninvolved by tumor in other sections of the specimen, a diagnosis of adenomyosis involved by adenocarcinoma should be made with caution. CD10 staining is not helpful in this differential diagnosis because stromal cells surrounding foci of invasive carcinoma are also frequently CD10 positive. There are no rules for determining how to measure the depth of invasion in the rare cases where myoinvasive carcinoma is only encountered in foci of adenomyosis involved by carcinoma. In such cases, it is advised that the distance from the adenomyotic focus to the deepest area of invasion be measured. Therefore, if there is a tumor with a 2-mm focus of myoinvasion from a focus of adenomyosis in the deep myometrium, it is still considered as having <50% myometrial invasion (FIGO stage IA).
E. Cervical Involvement
Cervical involvement by endometrial carcinoma has been traditionally divided into 2A when there was only secondary involvement of the endocervical epithelium and 2B when the endometrial carcinoma invaded the cervical stroma. Recently, it has been shown that involvement of the surface endocervical epithelium and/or endocervical glands (either by direct extension or drop metastases) does not have any prognostic significance. Therefore, the American Joint Committee on Cancer (AJCC)/FIGO staging system considers stage II disease only when cervical stromal involvement is seen.

F. Peritoneal Washings or Ascites Fluid
The prognostic significance of presence of tumor cells in peritoneal washings or ascites fluid is controversial. There are studies that indicate either a worse prognosis or no alteration of prognosis on the basis of positive cytology. Consequently, the newly adopted staging system no longer utilizes positive cytology to alter stage. When collected, however, cytology results should be recorded.

G. Margins
The paracervical soft tissue is the only true margin in total hysterectomy specimens, and reporting the status of this margin is usually not performed; conversely, reporting the status of the vaginal and parametrial margins in a radical hysterectomy specimen is optional.

H. Lymph-Vascular Invasion
Lymph-vascular invasion (LVI) indicates whether microscopic lymph-vascular invasion is identified. LVI includes lymphatic invasion, vascular invasion, or lymph-vascular invasion. According to AJCC/International Union Against Cancer (UICC) convention, LVI does not affect the T category indicating local extent of tumor unless specifically included in the definition of a T category.

I. TNM and FIGO Staging of Endometrial Carcinoma
The TNM staging system for endometrial cancer endorsed by the AJCC and the UICC, and the parallel system formulated by FIGO are recommended, as shown below.

According to AJCC/UICC convention, the designation “T” refers to a primary tumor that has not been previously treated. The symbol “p” refers to the pathologic classification of the TNM, as opposed to the clinical classification, and is based on gross and microscopic examination. pT entails a resection of the primary tumor or biopsy.
adequate to evaluate the highest pT category, pN entails removal of nodes adequate to validate lymph node metastasis, and pM implies microscopic examination of distant lesions. Clinical classification (cTNM) is usually carried out by the referring physician before treatment during initial evaluation of the patient or when pathologic classification is not possible.

Pathologic staging is usually performed after surgical resection of the primary tumor. Pathologic staging depends on pathologic documentation of the anatomic extent of disease, whether or not the primary tumor has been completely removed. If a biopsied tumor is not resected for any reason (eg, when technically infeasible) and if the highest T and N categories or the M1 category of the tumor can be confirmed microscopically, the criteria for pathologic classification and staging have been satisfied without total removal of the primary cancer.

It is important to note that in endometrial cancer, as in cancer of other organs, the validity of T stage depends upon the adequacy and completeness of the surgical staging.

**Primary Tumor (T):**

<table>
<thead>
<tr>
<th>TNM Category</th>
<th>FIGO Stage</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>(-)</td>
<td>Primary tumor cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>(-)</td>
<td>No evidence of primary tumor</td>
</tr>
<tr>
<td>T1</td>
<td>I</td>
<td>Tumor confined to corpus uteri</td>
</tr>
<tr>
<td>T1a</td>
<td>IA</td>
<td>Tumor limited to endometrium or invades less than one-half of the myometrium</td>
</tr>
<tr>
<td>T1b</td>
<td>IB</td>
<td>Tumor invades one-half or more of the myometrium</td>
</tr>
<tr>
<td>T2</td>
<td>II</td>
<td>Tumor invades stromal connective tissue of the cervix</td>
</tr>
<tr>
<td>T3</td>
<td>III</td>
<td>Local and/or regional spread as specified in T3a and T3b, and in FIGO IIIC1 and IIIC2</td>
</tr>
<tr>
<td>T3a</td>
<td>IIIC1</td>
<td>Regional lymph node metastasis to pelvic lymph nodes</td>
</tr>
<tr>
<td>T3b</td>
<td>IIIC2</td>
<td>Regional lymph node metastasis to para-aortic lymph nodes with or without positive pelvic lymph nodes</td>
</tr>
<tr>
<td>T4*</td>
<td>IVA</td>
<td>Tumor invades bladder mucosa* and/or bowel mucosa*</td>
</tr>
</tbody>
</table>

* Presence of bullous edema is not sufficient evidence to classify a tumor as T4.

**Regional Lymph Nodes (N):**

<table>
<thead>
<tr>
<th>TNM Stage</th>
<th>FIGO Stage</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>NX</td>
<td></td>
<td>Regional lymph nodes cannot be assessed</td>
</tr>
<tr>
<td>N0</td>
<td></td>
<td>No regional lymph node metastasis</td>
</tr>
<tr>
<td>N1</td>
<td>IIIC1</td>
<td>Regional lymph node metastasis to pelvic lymph nodes</td>
</tr>
<tr>
<td>N2</td>
<td>IIIC2</td>
<td>Regional lymph node metastasis to para-aortic lymph nodes with or without positive pelvic lymph nodes</td>
</tr>
</tbody>
</table>

* Regional lymph nodes include the pelvic, obturator, internal iliac (hypogastric), external iliac, common iliac, para-aortic, presacral, and parametral lymph nodes.

**Distant Metastasis (M):**

<table>
<thead>
<tr>
<th>TNM Stage</th>
<th>FIGO Stage</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>M0</td>
<td></td>
<td>No distant metastasis</td>
</tr>
<tr>
<td>M1</td>
<td>IVB</td>
<td>Distant metastasis (includes metastasis to abdominal lymph nodes [other than para-aortic], and/inguinal lymph nodes; excludes metastasis to vagina, pelvic serosa, or adnexa)</td>
</tr>
</tbody>
</table>

**TNM Stage Groupings**

<table>
<thead>
<tr>
<th>Stage 0</th>
<th>Tis</th>
<th>N0</th>
<th>M0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage IA</td>
<td>T1a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IB</td>
<td>T1b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage II</td>
<td>T2</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIIA</td>
<td>T3a</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIIB</td>
<td>T3b</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIIC1</td>
<td>T1-T3</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IIIC2</td>
<td>T1-T3</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IVA</td>
<td>T4</td>
<td>Any N</td>
<td>M0</td>
</tr>
</tbody>
</table>
Stage IVB  Any T  Any N  M1

**TNM Descriptors**

For identification of special cases of TNM or pTNM classifications, the “m” suffix and “y,” “r,” and “a” prefixes are used. Although they do not affect the stage grouping, they indicate cases needing separate analysis.

The “m” suffix indicates the presence of multiple primary tumors in a single site and is recorded in parentheses: pT(m)NM.

The “y” prefix indicates those cases in which classification is performed during or after initial multimodality therapy (ie, neoadjuvant chemotherapy, radiation therapy, or both chemotherapy and radiation therapy). The cTNM or pTNM category is identified by a “y” prefix. The ycTNM or ypTNM categorizes the extent of tumor actually present at the time of that examination. The “y” categorization is not an estimate of tumor before multimodality therapy (ie, before initiation of neoadjuvant therapy).

The “r” prefix indicates a recurrent tumor when staged after a documented disease-free interval and is identified by the “r” prefix: rTNM.

The “a” prefix designates the stage determined at autopsy: aTNM.

**Additional Descriptors**

**Residual Tumor (R)**

Tumor remaining in a patient after therapy with curative intent (eg, surgical resection for cure) is categorized by a system known as R classification, shown below.

RX  Presence of residual tumor cannot be assessed
R0  No residual tumor
R1  Microscopic residual tumor
R2  Macroscopic residual tumor

For the surgeon, the R classification may be useful to indicate the known or assumed status of the completeness of a surgical excision. For the pathologist, the R classification is relevant to the status of the margins of a surgical resection specimen. That is, tumor involving the resection margin on pathologic examination may be assumed to correspond to residual tumor in the patient and may be classified as macroscopic or microscopic according to the findings at the specimen margin(s).

**Regional Lymph Nodes and Sentinel Lymph Nodes (sn): Isolated Tumor Cells**

Isolated tumor cells (ITCs) are single cells or small clusters of cells not more than 0.2 mm in greatest dimension. Lymph nodes or distant sites with ITCs found by either histologic examination (eg, immunohistochemical evaluation for cytokeratin) or nonmorphological techniques (eg, flow cytometry, DNA analysis, polymerase chain reaction [PCR] amplification of a specific tumor marker) should be so identified. There is currently no guidance in the literature as to how these patients should be coded; until more data are available, they should be coded as “N0(i+)” with a comment noting how the cells were identified.

There is little data to assign risk for nonsentinel lymph node metastasis based on the size of the metastasis in the sentinel lymph node. However, the size criteria for micrometastasis and macrometastasis is adopted from the experience in breast carcinoma. Micrometastasis is defined as a metastasis measuring greater than 0.2 mm but equal to or less than 2 mm.

**J. Additional Findings**

**Endometrial Intraepithelial Neoplasia**

In part because of poor agreement in the diagnosis of atypical hyperplasia of the endometrium under the WHO criteria, a new diagnostic terminology, endometrial intraepithelial neoplasia (EIN), has been proposed. EIN describes a clonal expansion of premalignant endometrial glands with endometrioid features, but without invasion. Microscopic criteria proposed to diagnose this lesion include:
K. Clinical History

Colon carcinoma is the most common malignancy in hereditary nonpolyposis colon cancer (HNPCC; Lynch syndrome). However, endometrial carcinoma develops before colon carcinoma in >50% of women with HNPCC.\(^{20-23}\) Still, the reported series of HNPCC-related endometrial carcinomas are much smaller in number than those reported for HNPCC colonic carcinoma. Histopathologic features suggestive of HNPCC-related carcinoma are well characterized in the colon, but not in the uterus. However, when examining an endometrial carcinoma in a patient under 50 years of age or with a personal or family history of colon carcinoma, it is important to consider the possibility of an HNPCC-related endometrial carcinoma. In these cases, testing for defective DNA mismatch repair may be performed by immunohistochemistry (MLH1, MSH2, MSH6, and PMS2 antibodies are commercially available). Loss of MSH2 expression essentially always indicates Lynch syndrome and MSH6 is related to MSH2. HNPCC-related endometrial carcinoma is predominantly associated with MSH2 mutations, and MSH6 mutations in particular.\(^{20-23}\) PMS2 loss is often associated with loss of MLH1 and is only independently meaningful if MLH1 is intact. In addition, PCR assays can be used to detect high levels of microsatellite alterations (MSI), a condition that is definitional for defective DNA mismatch repair. This testing is performed on paraffin-embedded tissue and compares the results of tumor DNA to those of nonneoplastic tissues from the same patient. Please refer to the CAP endometrial cancer biomarker reporting template on www.cap.org for further details.

References


**Bibliography**