Ovary

Protocol applies to all primary borderline and malignant surface epithelial tumors, germ cell tumors, and sex cord-stromal tumors.

Protocol revision date: January 2004
Based on AJCC/UICC TNM, 6th edition and FIGO 2001 Annual Report

Procedures
• Cytology (No Accompanying Checklist)
• Incisional Biopsy (No Accompanying Checklist)
• Unilateral Oophorectomy
• Salpingo-oophorectomy
• Subtotal Resection or Removal of Tumor in Fragments
• Hysterectomy with Salpingo-oophorectomy
• Second-look Staging (No Accompanying Checklist)

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Surgical Pathology Cancer Case Summary (Checklist)

Protocol revision date: January 2004
Applies to primary borderline tumors, carcinomas, germ cell tumors, and sex-cord stromal tumors only
Based on AJCC/UICC TNM, 6th edition and FIGO 2001 Annual Report

OVARY: Oophorectomy, Salpingo-oophorectomy, Subtotal Oophorectomy or Removal of Tumor in Fragments, Hysterectomy with Salpingo-oophorectomy

Note: Applies to ovarian primary tumor. If bilateral tumors of 2 different histologic types are present, separate checklists should be used for each tumor.

Patient name:
Surgical pathology number:

Note: Check 1 response unless otherwise indicated.

MACROSCOPIC

Specimen Type (check all that apply)
___ Right oophorectomy
___ Left oophorectomy
___ Right salpingo-oophorectomy
___ Left salpingo-oophorectomy
___ Subtotal right oophorectomy
___ Subtotal left oophorectomy
___ Removal of tumor in fragments
___ Hysterectomy with salpingo-oophorectomy
___ Omentectomy
___ Other (specify): ____________________________
___ Not specified

Primary Tumor Site (check all that apply)
___ Diffuse bilateral ovarian involvement, primary site cannot be determined

Right Ovary
___ Not applicable
___ Parenchymal growth
___ Growth on surface
___ Uninvolved

* Data elements with asterisks are not required for accreditation purposes for the Commission on Cancer. These elements may be clinically important, but are not yet validated or regularly used in patient management. Alternatively, the necessary data may not be available to the pathologist at the time of pathologic assessment of this specimen.

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**Left Ovary**
- ___ Not applicable
- ___ Parenchymal growth
- ___ Growth on surface
- ___ Uninvolved
- ___ Not specified

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**Specimen Integrity (check all that apply)**

**Right Ovary**
- ___ Not applicable
- ___ Intact
- ___ Ruptured
- ___ Fragmented
- ___ Other (specify): ____________________________

**Left Ovary**
- ___ Not applicable
- ___ Intact
- ___ Ruptured
- ___ Fragmented
- ___ Other (specify): ____________________________

**Tumor Size**

**Right Ovary (if applicable)**
Greatest dimension: ___ cm
*Additional dimensions: ___ x ___ cm
___ Cannot be determined (see Comment)

**Left Ovary (if applicable)**
Greatest dimension: ___ cm
*Additional dimensions: ___ x ___ cm
___ Cannot be determined (see Comment)
**MICROSCOPIC**

**Histologic Type (check all that apply)**

- ___ Serous, borderline
- ___ Serous, carcinoma
- ___ Mucinous, borderline
- ___ Mucinous, carcinoma
- ___ Endometrioid, borderline
- ___ Endometrioid, carcinoma
- ___ Clear cell, borderline
- ___ Clear cell, carcinoma
- ___ Transitional cell, borderline
- ___ Transitional cell, carcinoma
- ___ Mixed epithelial, borderline
  
  Specify types: ____________________________
- ___ Mixed epithelial, carcinoma
  
  Specify types: ____________________________
- ___ Undifferentiated
- ___ Granulosa cell
- ___ Germ cell
  
  Specify type(s): ____________________________
- ___ Other(s) (specify): ____________________________

**Histologic Grade**

- ___ Not applicable
- ___ GX: Cannot be assessed
- ___ G1: Well differentiated
- ___ G2: Moderately differentiated
- ___ G3: Poorly differentiated
- ___ G4: Reserved solely for tumors in the undifferentiated category
  
  (WHO classification)

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Pathologic Staging (pTNM [FIGO])

Primary Tumor (pT)
___ pTX [-]: Cannot be assessed
___ pT0 [-]: No evidence of primary tumor
___ pT1 [I]: Tumor limited to ovaries (1 or both)
*___ pT1a [IA]: Tumor limited to 1 ovary; capsule intact, no tumor on ovarian surface. No malignant cells in ascites or peritoneal washings
*___ pT1b [IB]: Tumor limited to both ovaries; capsule intact, no tumor on ovarian surface. No malignant cells in ascites or peritoneal washings
*___ pT1c [IC]: Tumor limited to 1 or both ovaries with any of the following: capsule ruptured, tumor on ovarian surface, malignant cells in ascites or peritoneal washings
 ___ pT2 [II]: Tumor involves 1 or both ovaries with pelvic extension and/or implants
*___ pT2a [IIA]: Extension and/or implants on uterus and/or tube(s). No malignant cells in ascites or peritoneal washings
*___ pT2b [IIB]: Extension to other pelvic tissues. No malignant cells in ascites or peritoneal washings
*___ pT2c [IIC]: Pelvic extension and/or implants (T2a or T2b / IIA or IIB) with malignant cells in ascites or peritoneal washings
pT3 and/or N1 [III]: Tumor involves 1 or both ovaries with microscopically confirmed peritoneal metastasis outside the pelvis (including liver capsule metastasis) and/or regional lymph node metastasis
___ pT3a [IIIA]: Microscopic peritoneal metastasis beyond pelvis
___ pT3b [IIIB]: Macroscopic peritoneal metastasis beyond pelvis 2 cm or less in greatest dimension
___ pT3c and/or N1[IIIC]: Peritoneal metastasis beyond pelvis more than 2 cm in greatest dimension and/or regional lymph node metastasis
____ Any T/Any N and M1 [IV]: Growth involving 1 or both ovaries with distant metastasis.

Note: If pleural effusion is present, there must be positive cytology to assign a case to stage IV. Parenchymal liver metastasis is classified as stage IV.

Regional Lymph Nodes (pN)
___ pNX: Cannot be assessed
___ pN0: No regional lymph node metastasis
___ pN1 [IIIC]: Regional lymph node metastasis
Specify: Number examined: ____
Number involved: ____

Distant Metastasis (pM)
___ pMX: Cannot be assessed
___ pM1 [IV]: Distant metastasis
  *Specify site(s), if known: ____________________________

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Implants (only applies to borderline tumors) (check all that apply)
___ Not applicable/none sampled
Noninvasive (epithelial) implants
___ Not present
___ Present
   Specify site(s): ____________________________
Noninvasive (desmoplastic) implants
___ Not present
___ Present
   Specify site(s): ____________________________
Invasive implants
___ Not present
___ Present
   Specify site(s): ____________________________

Summary of Organs/Tissues Microscopically Involved by Tumor
(check all that apply)
___ One ovary
___ Both ovaries
___ Omentum
___ Uterus
___ Peritoneum
___ Other organs/tissues
   Specify all: ____________________________

*Venous/Lymphatic (Large/Small Vessel) Invasion (V/L)
*___ Absent
*___ Present
*___ Indeterminate

*Additional Pathologic Findings (check all that apply)
*___ None identified
*___ Endometriosis
   *___ Ovarian
   *___ Extraovarian
*___ Endosalpingiosis
*___ Other(s):
   *Specify site(s) and type(s): ________________________________

*Comment(s)
I. Cytologic Material
A. Clinical Information
   1. Patient identification
      a. Name
      b. Identification number
      c. Age (birth date)
   2. Responsible physician(s)
   3. Date of procedure
   4. Other clinical information
      a. Relevant history
         (1) currently pregnant
         (2) abnormal uterine bleeding pattern
         (3) previous therapy (hormonal, radiation, chemotherapy)
         (4) previous tumors or operations of possible relevance
         (5) family history of ovarian or breast cancer
      b. Relevant findings (eg, radiologic studies, aspiration of cyst, laboratory data, ascites)
      c. Clinical diagnosis
      d. Procedure (eg, brushing, washing, other)
      e. Operative findings (eg, rupture-preoperative, intraoperative, or postoperative)
      f. Type(s) or site(s) of specimen(s)
         (1) ascitic fluid
         (2) peritoneal washings (specify site)
         (3) brushings (specify site)
         (4) cyst fluid (specify site)
         (5) fine-needle aspirate (specify site)
         (6) cytology preparation of tissue (touch preparation) (specify site)
         (7) pleural fluid
         (8) other

B. Macroscopic Examination
   1. Specimen
      a. Unfixed/fixed (specify fixative)
      b. Number of slides received, if appropriate
      c. Quantity and appearance of fluid specimen, if appropriate
      d. Other (eg, tissue received for cytologic preparation)
      e. Results of intraprocedural consultation
   2. Material submitted for microscopic evaluation (eg, smear, cytocentrifuge, touch or filter preparation, cell block)
   3. Tissues submitted for special studies (specify) (eg, frozen tissue for immunohistochemistry, flow cytometry)

C. Microscopic Evaluation
   1. Adequacy of specimen (if unsatisfactory for evaluation, specify reason)
   2. Tumor, if present
      a. Histologic type, if possible (Note A)
      b. Other characteristics, as pertinent
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3. Additional cytologic findings, if present
4. Results/status of special studies (specify)
5. Pathologic stage
6. Comments
   a. Correlation with intraprocedural consultation, as appropriate
   b. Correlation with other specimens, as appropriate
   c. Correlation with clinical information, as appropriate

II. Incisional Biopsy
A. Clinical Information
   1. Patient identification
      a. Name
      b. Identification number
      c. Age (birth date)
   2. Responsible physician(s)
   3. Date of procedure
   4. Other clinical information
      a. Relevant history
         (1) currently pregnant
         (2) abnormal uterine bleeding pattern
         (3) previous therapy (hormonal, radiation, chemotherapy)
         (4) previous tumors or operations of possible relevance
         (5) family history of ovarian or breast cancer
      b. Relevant findings (eg, radiologic studies, aspiration of cyst, laboratory data, ascites)
      c. Clinical diagnosis
      d. Procedure
      e. Operative findings (eg, rupture-preoperative, intraoperative, or postoperative)
      f. Type(s) or site(s) of specimen(s)
B. Macroscopic Examination
   1. Specimen
      a. Fixed/unfixed (specify fixative)
      b. Number of pieces, size or size range
      c. Descriptive features
      d. Orientation, if designated
      e. Results of intraoperative consultation
   2. Tissues submitted for microscopic evaluation
      a. Submit entire specimen
      b. Frozen section tissue fragment(s) (unless saved for special studies)
   3. Tissues submitted for special studies (specify) (eg, frozen tissue for immunohistochemistry, flow cytometry)
C. Microscopic Evaluation
   1. Tumor, if present
      a. Histologic type (Note A)
      b. Histologic grade (Note B)
      c. Invasion
      d. Other features of possible prognostic or therapeutic significance
2. Additional pathologic findings, if present (specify); and relation to tumor, if pertinent
3. Results/status of special studies (Note C)
4. Pathologic stage
5. Comments
   a. Correlation with intraoperative consultation, as appropriate
   b. Correlation with other specimens, as appropriate
   c. Correlation with clinical information, as appropriate

III. Unilateral Oophorectomy or Salpingo-oophorectomy

A. Clinical Information
1. Patient identification
   a. Name
   b. Identification number
   c. Age (birth date)
2. Responsible physician(s)
3. Date of procedure
4. Other clinical information
   a. Relevant history
      (1) currently pregnant
      (2) abnormal uterine bleeding pattern
      (3) previous therapy (hormonal, radiation, chemotherapy)
      (4) previous tumors or operations of possible relevance
      (5) family history of ovarian or breast cancer
   b. Relevant findings (eg, radiologic studies, laboratory data, ascites)
   c. Clinical diagnosis
   d. Procedure
   e. Operative findings (eg, rupture-preoperative, intraoperative, or postoperative)
   f. Type(s) or site(s) of specimen(s)
   g. Identification of areas for special study
      (1) rupture site(s)
      (2) adhesions suspicious for tumor
      (3) resection margin(s), if pertinent

B. Macroscopic Examination
1. Specimen
   a. Organs/tissues received (specify)
   b. Unfixed/fixed (specify fixative)
   c. Number of pieces
   d. Dimensions (measure attached tissues individually)
   e. Orientation of specimen
   f. Results of intraoperative consultation
2. Ovary or ovary-tube, if fused into single mass*
   * If fused ovary and tube are identifiable separately on sectioning, describe tumor in each, including relation to one another.
   a. Size, weight, or volume, as appropriate
   b. Outer surface (describe)
      (1) adhesions, roughening, granularity (largest dimension or proportion of total area involved)
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(2) rupture (Note D)

c. Sectioned surface of specimen or opened cyst(s)
d. Tumor (Note E)
   (1) location (Note F)
      i. cortex
      ii. medulla
      iii. hilus
      iv. combination
      v. replaces specimen
   (2) dimensions (and proportion of entire specimen, if appropriate)
   (3) solid and cystic components
      i. proportion of each
      ii. number of cysts, if easily countable
      iii. size range of each component
      iv. location(s), if pertinent
      v. contents of cyst(s)
      vi. lining of cyst(s)
      vii. papillary or polypoid excrescences
      viii. roughening, etc (largest dimension or proportion of total area involved)
e. Identification of areas for special study (eg, rupture site[s], adhesions suspicious for tumor)
f. Resection margin(s), if pertinent
g. Additional pathologic findings, if present

3. Fallopian tube, if not fused with ovary into single mass
a. Tumor, if present
   (1) size
   (2) location and relation to ovarian tumor
   (3) descriptive features
b. Additional pathologic findings, if present

4. Tissues submitted for microscopic evaluation (Note G)

5. Tissues submitted for special studies (specify) (eg, frozen tissue for immunohistochemistry, flow cytometry)

C. Microscopic Evaluation

1. Ovary or ovary-tube, if fused into single mass
a. Tumor
   (1) site(s) of origin (Note H)
   (2) location (Note F)
   (3) surface (if possible, distinguish origin on surface from invasion onto surface by subjacent tumor)
   (4) extent of invasion in ovary
      i. cortex
      ii. medulla
      iii. hilus
      iv. combination
      v. replaces specimen
   (5) extent and distribution of invasion of tube, if involved
   (6) low-power pattern (single mass, discrete nodules, etc) (Note I)
   (7) histologic type (note mixtures) (Note J)
(8) histologic grade (Note B)
(9) venous/lymphatic vessel invasion (Note K)
(10) other features of possible prognostic or therapeutic significance
b. Status of any specially designated resection margins
c. Additional pathologic findings, if present (specify); and relation to tumor, if pertinent (Note L)
2. Fallopian tube, if not fused with ovary into single mass
a. Tumor
   (1) site(s) of origin (Note H)
   (2) location
   (3) extent
   (4) histologic type or grade, if different from that of ovarian tumor
   (5) venous/lymphatic vessel invasion (Note K)
   (6) other features of possible prognostic or therapeutic significance
b. Status of any specially designated resection margins
c. Appearance of epithelium adjacent to tumor (Note H)
d. Additional pathologic findings, if present (specify); and relation to tumor, if pertinent
3. Results/status of special studies (Note C)
4. Pathologic stage
5. Comments
   a. Correlation with intraoperative consultation, as appropriate
   b. Correlation with other specimens, as appropriate
   c. Correlation with clinical information, as appropriate

IV. Subtotal Resection or Removal of Tumor in Fragments
A. Clinical Information
1. Patient identification
   a. Name
   b. Identification number
   c. Age (birth date)
2. Responsible physician(s)
3. Date of procedure
4. Other clinical information
   a. Relevant history
      (1) currently pregnant
      (2) abnormal uterine bleeding pattern
      (3) previous therapy (hormonal, radiation, chemotherapy)
      (4) previous tumors or operations of possible relevance
      (5) family history of ovarian or breast cancer
   b. Relevant findings (eg, radiologic studies, laboratory data, ascites)
   c. Clinical diagnosis
   d. Procedure
   e. Operative findings (eg, rupture-preoperative, intraoperative, or postoperative)
   f. Type(s) or site(s) of specimen(s)
   g. Identification of areas for special study
      (1) rupture site(s)
      (2) adhesions suspicious for tumor
B. Macroscopic Examination

1. Specimen
   a. Organs/tissues received
   b. Unfixed/fixed (specify fixative)
   c. Number of pieces
   d. Dimensions (measure organs and tissues individually)
   e. Orientation of specimen
   f. Results of intraoperative consultation
   g. Outer surface (describe)
      (1) adhesions, roughening, granularity (largest dimension or proportion of total area involved)
      (2) rupture (Note D)
   h. Sectioned surface of specimen or opened cyst(s) #
      # If fused ovary and tube are identifiable separately on sectioning, describe tumor in each, including relation to one another.
   i. Tumor (Note E)
      (1) location (Note F)
         i. cortex
         ii. medulla
         iii. hilus
         iv. combination
         v. replaces specimen
   j. Dimensions (and proportion of entire specimen, if appropriate) of solid and cystic components
      (1) proportion of each
      (2) number of cysts, if easily countable
      (3) size range of each component
      (4) location(s), if pertinent
      (5) contents of cyst(s)
      (6) lining of cyst(s)
      (7) papillary or polypoid excrescences
      (8) roughening, etc (largest dimension or proportion of total area involved)
   k. Outer surface (describe)
   l. Identification of areas for special study
      (1) rupture site(s)
      (2) adhesions suspicious for tumor
      (3) resection margin(s), if pertinent

2. Additional pathologic findings, if present

3. Tissues submitted for microscopic evaluation (Note G)

4. Tissues submitted for special studies (specify) (eg, frozen tissue for immunohistochemistry, flow cytometry)

C. Microscopic Evaluation

1. Tumor
   a. Site(s) of origin (Note H)
   b. Location (Note F)
   c. Surface (if possible, distinguish origin on surface from invasion onto surface by subjacent tumor)
   d. Extent of invasion
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(1) cortex
(2) medulla
(3) hilus
(4) combination
(5) replaces specimen
e. Extent and distribution in tube, if involved
f. Low-power pattern (single mass, discrete nodules, etc) (Note I)
g. Histologic type (note mixtures) (Note J)
h. Histologic grade (Note B)
i. Venous/lymphatic vessel invasion (Note K)
j. Other features of possible prognostic or therapeutic significance

2. Status of any specially designated resection margins

3. Additional pathologic findings, if present; and relation to tumor, if pertinent (Note L)
   a. Endometriosis
   b. Abnormalities of surface epithelium or surface epithelial inclusion glands or cysts
   c. Others

4. Results/status of special studies (Note C)

5. Pathologic stage

6. Comments
   a. Correlation with intraoperative consultation, as appropriate
   b. Correlation with other specimens, as appropriate
   c. Correlation with clinical, information, as appropriate

V. Hysterectomy with Salpingo-oophorectomy and Removal of Attached and/or Unattached Organs or Tissues (Staging Procedure)

A. Clinical Information
   1. Patient identification
      a. Name
      b. Identification number
      c. Age (birth date)
   2. Responsible physician(s)
   3. Date of procedure
   4. Other clinical information
      a. Relevant history
         (1) currently pregnant
         (2) abnormal uterine bleeding pattern
         (3) previous therapy (hormonal, radiation, chemotherapy)
         (4) previous tumors or operations of possible relevance
         (5) family history of ovarian or breast cancer
      b. Relevant findings (eg, radiologic studies, laboratory data, ascites)
      c. Clinical diagnosis
   d. Procedure
   e. Operative findings (eg, rupture-preoperative, intraoperative, or postoperative)
   f. Type(s) or site(s) of specimen(s)
   g. Identification of areas for special study
(1) rupture site(s)
(2) adhesions suspicious for tumor
(3) resection margins, if pertinent

B. Macroscopic Examination

1. Specimen
   a. Organs/tissues included
   b. Unfixed/fixed (specify fixative)
   c. Number of pieces
   d. Dimensions (measure attached tissues individually)
   e. Orientation of specimen
   f. Results of intraoperative consultation

2. Ovary or ovary-tube if fused into single mass
   a. Size, weight, or volume, as appropriate
   b. Outer surface (describe)
      (1) adhesions, roughening, granularity (largest dimension or proportion of total area involved)
      (2) rupture (Note D)

3. Sectioned surface of specimen or opened cyst(s)
   a. Tumor (Note E)
      (1) location (Note F)
         i. cortex
         ii. medulla
         iii. hilus
         iv. combination
         v. replaces specimen
      (2) dimensions (and proportion of entire specimen, if appropriate) of solid and cystic components
         i. proportion of each
         ii. number of cysts, if easily countable
         iii. size range of each component
         iv. location(s), if pertinent
         v. contents of cyst(s)
         vi. lining of cyst(s)
         vii. papillary or polypoid excrescences
         viii. roughening, etc (largest dimension or proportion of total area involved)
   b. Identification of areas for special study (eg, rupture site[s], adhesions suspicious for tumor, resection margin[s], if pertinent)
   c. Additional pathologic findings, if present

4. Fallopian tube, if not fused with ovary into single mass
   a. Tumor, if present
      (1) size
      (2) location and relation to ovarian tumor
      (3) descriptive features
   b. Additional pathologic findings, if present

5. Contralateral ovary (Note M)
   a. Size, weight, or volume, if appropriate
b. Outer surface (describe)
c. Sectioned surface
d. Tumor, if present (handle as for predominant ovarian mass)
e. Other lesions (specify)

6. Uterus
   a. Descriptive features of endometrium, myometrium, and serosa
   b. Tumor, if present
      (1) descriptive features
      (2) location (depth of myometrial invasion, if appropriate)
      (3) relation to ovarian tumor (separate or adherent)
   c. Other lesions, if present (specify)

7. Omentum (Note G)

8. Regional lymph nodes
   a. Number and size range at each designated location
   b. Tumor, if identifiable
   c. Other lesions, if present (specify)

9. Other staging biopsy specimens (specify)
   a. Tumor, if present
      (1) descriptive features
      (2) location
   b. Other lesions, if present (specify)

10. Other organ(s) or tissue(s) removed
    a. Type, dimensions, and other descriptive features
    b. Tumor
       (1) location and relation to ovarian tumor (separate or adherent)
       (2) size and distribution within organ or tissue
    c. Resection margins, if applicable
    d. Other lesions, if present (specify)

11. Tissues submitted for microscopic evaluation (Note G)

12. Tissues submitted for special studies (specify) (eg, frozen tissue for immunohistochemistry, flow cytometry)

**B. Microscopic Evaluation**

1. Ovary or ovary-tube if fused into single mass
   a. Tumor
      (1) site(s) of origin (Note H)
      (2) location (Note F)
      (3) surface (if possible, distinguish origin on surface from invasion onto surface from subjacent tumor)
      (4) extent of invasion in ovary
         i. cortex
         ii. medulla
         iii. hilus
         iv. combination
         v. replaces specimen
      (5) extent and distribution of invasion of tube, if involved
      (6) low-power pattern (single mass, discrete nodules, etc) (Note I)
      (7) histologic type (note mixtures) (Note J)
      (8) histologic grade (Note B)
      (9) venous/lymphatic vessel invasion (Note K)
(10) other features of possible prognostic or therapeutic significance
b. Status of any specially designated resection margins
c. Additional pathologic findings, if present (specify); and relation to tumors, if pertinent (Note L)
2. Fallopian tube if not fused with ovary into single mass
   a. Tumor
      (1) site(s) of origin (Note H)
      (2) location
      (3) extent
      (4) histologic type or grade, if different from that of ovarian tumor
      (5) venous/lymphatic vessel invasion (Note K)
      (6) other features of possible prognostic or therapeutic significance
   b. Status of any specially designated resection margin(s)
   c. Appearance of epithelium adjacent to tumor (Note H)
   d. Additional pathologic findings, if present (specify); and relation to tumor, if pertinent
3. Contralateral ovary (Note M)
   a. Tumor, if present (handle as for predominant ovarian mass)
   b. Additional pathologic findings, if present
      (1) endometriosis
      (2) abnormalities of surface epithelium
      (3) surface epithelial glands or cysts (Note L)
4. Uterus
   a. Tumor, if present (Note N)
      (1) histologic type
      (2) histologic grade
      (3) location
      (4) extent including depth of invasion of wall, if suspected to be primary in endometrium
      (5) venous/lymphatic vessel invasion
   b. Endometrium uninvolved by tumor
   c. Additional pathologic findings, if present (Note O)
5. Omentum (Note P)
   a. Tumor, if present
      (1) histologic type, if different from ovarian tumor
      (2) histologic grade, if different from ovarian tumor (Note P)
      (3) invasive or noninvasive (Note P)
   b. Additional pathologic findings, if present (Note O)
6. Regional lymph nodes (at each location, if separately designated)
   a. Number
   b. Number involved by tumor
      (1) histologic type, if different from ovarian tumor
      (2) histologic grade, if different from ovarian tumor (Note P)
   c. Additional pathologic findings, if present (Note O)
7. Other staging biopsy specimens at each location, if so designated
   a. Tumor, if present
      (1) histologic type, if different from ovarian tumor
      (2) histologic grade, if different from ovarian tumor (Note P)
   b. Additional pathologic findings, if present (Note O)
8. Other organs or tissue removed
   a. Tumor, if present
      (1) location, distribution, and extent
      (2) histologic type, if different from ovarian tumor
      (3) histologic grade, if different from ovarian tumor (Note P)
   b. Resection margins, if applicable
   c. Additional pathologic findings, if present (Note O)
9. Results/status of special studies (Note C)
10. Pathologic stage
11. Comments
   a. Correlation with intraoperative consultation, as appropriate
   b. Correlation with other specimens, as appropriate
   c. Correlation with clinical information, as appropriate

VI. Second-look Staging
A. Clinical Information
1. Patient identification
   a. Name
   b. Identification number
   c. Age (birth date)
2. Responsible physician(s)
3. Date of procedure
4. Other clinical information
   a. Relevant history
      (1) currently pregnant
      (2) abnormal uterine bleeding pattern
      (3) previous therapy (hormonal, radiation, chemotherapy)
      (4) previous tumors or operations of possible relevance
      (5) family history of ovarian or breast cancer
   b. Relevant findings (eg, radiologic studies, laboratory data)
   c. Clinical diagnosis
   d. Procedure
   e. Operative findings
   f. Type(s) or site(s) of specimen(s)
B. Macroscopic Examination
1. Specimen
   a. Organs/tissues received
   b. Unfixed/fixed (specify fixative)
   c. Number of pieces
   d. Dimensions (measure attached tissues individually)
   e. Orientation of specimens
   f. Results of intraoperative consultation
2. Tumor
   a. Size
   b. Descriptive characteristics
3. Additional pathologic findings, if present
4. Tissues submitted for microscopic evaluation
   a. Submit entire specimen
   b. Frozen section tissue fragment(s) (unless saved for special studies)
5. Tissues submitted for special studies (specify) (eg, frozen tissue for immunohistochemistry, flow cytometry)

C. Microscopic Evaluation

1. Tumor
   a. Site(s) of origin of tumor
   b. Histologic type, if different from ovarian tumor
   c. Histologic grade, if different from ovarian tumor (Note P)
   d. Invasive vs noninvasive
   e. Changes due to therapy, if present

2. Additional pathologic findings, if present (Note O)

3. Results/status of special studies (specify) (Note C)

4. Comments
   a. Correlation with intraoperative consultation, as appropriate
   b. Correlation with other specimens, as appropriate
   c. Correlation with clinical information, as appropriate

Explanatory Notes

A. Histologic Type

It is recommended that the World Health Organization (WHO) Classification and Nomenclature of Ovarian Tumors be used because of its wide acceptance. An abbreviated form of this classification is shown below.

WHO Classification of Malignant Ovarian Tumors

Surface Epithelial-Stromal Tumors

Histologic Type (Epithelial Component)

- Serous
- Mucinous
- Endometrioid
- Clear cell
- Transitional (including Brenner)
- Squamous
- Mixed
- Undifferentiated

(Degree of Malignancy of Epithelial and/or Stromal Component)

- Borderline (of low malignant potential)†
- Malignant
  - Carcinoma
  - Sarcoma
  - Both (malignant mesodermal mixed tumor)

Germ Cell Tumors

- Dysgerminoma
- Yolk sac tumor (endodermal sinus tumor)
- Immature teratoma
- Mixed malignant germ cell tumors (specify types)
- Cancer with dermoid cyst (specify type)
- Other (specify)
Sex Cord-Stromal Tumors
   Granulosa cell tumor
   Other (specify)

These tumors should be further subclassified according to the location of the neoplastic cells (e.g., “cystadeno-,” “surface,” or both) and the quantity of their stromal component. When the stromal component predominates, “adenofibro-” appears in the diagnostic term. This addition may be important since malignant ovarian tumors in which the neoplastic cells are surrounded by abundant benign fibromatous tissue appear to have a better prognosis than those without such a component. Surface involvement by neoplastic cells elevates the substage in stage 1 cases and indicates a higher likelihood of extraovarian peritoneal involvement.

Kurman and his group\(^2\) have recently challenged the concept of borderline neoplasia, providing evidence that most so-called borderline tumors in the serous category should be designated “atypical proliferating” because they are rarely fatal, but that a small subset of serous borderline tumors with a micropapillary or cribriform pattern are fatal in a substantial number of cases, and should be designated “micropapillary carcinoma” even in the absence of invasion of the stromal component of the tumor. This proposal has not been widely accepted by gynecological pathologists,\(^3\)\(^-\)\(^4\) however, satisfactory reproducibility of the histological distinction between the 2 proposed forms of serous borderline tumor has not yet been established. Evaluation of additional series of cases by other groups of investigators is warranted before change in the WHO terminology is considered.

Another proposal by several groups of investigators\(^5\)\(^-\)\(^7\) is the reinterpretation of so-called mucinous borderline tumors of intestinal type associated with pseudomyxoma peritonei as metastatic tumors of intestinal (almost always appendiceal) type in almost all cases. This proposal should be considered by staging groups in the near future.

B. Histologic Grade for Surface Epithelial Stromal Tumors

Numerous grading systems, including architectural, nuclear, and combined architectural and nuclear systems, as well as schemas that incorporate additional features (e.g., appearance of tumor margin, inflammatory cell reaction, and vascular space invasion) have been used for ovarian cancers. This protocol does not recommend any specific grading system since several types that have been evaluated have proved to have prognostic significance.\(^8\)\(^-\)\(^10\) For the sake of uniformity, however, it is recommended that 3 grades be used, as shown below.

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>Cannot be assessed</td>
</tr>
<tr>
<td>1</td>
<td>Well differentiated</td>
</tr>
<tr>
<td>2</td>
<td>Moderately differentiated</td>
</tr>
<tr>
<td>3</td>
<td>Poorly differentiated (tumors with minimal differentiation seen in very small foci)</td>
</tr>
</tbody>
</table>

Note: Grade 4 is reserved solely for tumors in the undifferentiated category (WHO classification).
Epithelial Cancers

As a rule, both architectural and nuclear features are evaluated. The prognostic significance of the grading varies with the type of tumor.

**Serous.** Usually, architectural features parallel nuclear features (ie, the extent of gland and papillae formation versus the quantity of solid growth correlates with well versus moderate versus poor differentiation). Exceptions exist, however, such as certain tumors with a solid growth pattern in the form of small nests exhibiting a high degree of nuclear maturation and often containing numerous psammoma bodies. Tumors in the latter category are assigned grade 1 despite their solid architecture.

**Mucinous.** Architectural and nuclear features are both evaluated. The most important determination, however, is whether the tumor is borderline or carcinoma (WHO).

Many mucinous tumors that lack obvious stromal invasion contain cysts and glands lined by malignant instead of atypical epithelium. Such tumors have been designated “intraglandular carcinoma” or borderline tumor “with intraepithelial carcinoma.” These tumors appear to have an excellent prognosis but one that appears to be slightly worse than that of borderline tumors lacking this feature.

**Endometrioid.** These tumors can be graded according to the system suggested for similar tumors of the uterine corpus.

- **Grade 1:** 5% or less of a non-squamous, solid growth pattern
- **Grade 2:** 6% to 50% of the neoplasm has a solid non-squamous pattern
- **Grade 3:** More than 50% of the tumor shows a solid (non-squamous) growth pattern

Notable nuclear atypia, inappropriate for the architectural grade, raises the grade of a grade 1 or grade 2 tumor by 1 grade. Adenocarcinomas with squamous differentiation are graded according to the nuclear grade of the glandular component.

**Clear cell.** Although it is recommended that these carcinomas be graded with the use of general guidelines, many investigators have not been able to correlate the grade with the prognosis.

**Transitional cell.** Nuclear grading, with a 3-tiered grading scale, is recommended.

**Squamous cell.** Histologic grading, using a 3-tiered grading-scale, is recommended.

Borderline Tumors

Grading of ovarian borderline tumors may be done but has not been proved prognostically significant, stage for stage. There is some evidence, however, that nuclear grading of associated peritoneal disease is prognostically important in cases of serous borderline tumors.
Germ Cell Tumors
Immature teratomas are graded on the basis of the quantity of embryonal elements, almost always neuroectodermal, that they contain. Other primitive germ cell tumors are not graded.

Granulosa Cell Tumors
Two groups of investigators have found that nuclear grading is effective in determining prognosis.

C. Special Studies
Special studies include histochemical and immunohistochemical staining, which are helpful diagnostically in occasional cases; flow cytometry; DNA image analysis; quantitative microscopy; hormone receptor studies; molecular genetic studies; chromosome analysis; and others. At present, the use of special studies for prognostic and therapeutic purposes remains controversial and is under continuing investigation.

D. Rupture of Tumor
Site refers to the location of rupture within a complex tumor that may be partly benign, partly borderline, and partly invasive or partly mature and partly immature if the tumor is a teratoma. In such cases, it may be helpful to the gynecologist to know what component of the tumor has ruptured to formulate an opinion whether benign, borderline, or malignant cells may have spilled into the abdominal cavity.

E. TNM and Stage Groupings
In view of the role of the pathologist in the staging of cancers, the staging system for ovarian cancer endorsed by the American Joint Committee on Cancer (AJCC) and the International Union Against Cancer (UICC), as well as the parallel system formulated by the International Federation of Gynecology and Obstetrics (FIGO) are recommended.

By 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 unfeasible) 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.
**TNM and FIGO Staging Systems for Ovarian Carcinoma**

<table>
<thead>
<tr>
<th>Primary Tumor (T)</th>
<th>TNM</th>
<th>FIGO</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>—</td>
<td>—</td>
<td>Primary tumor cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>—</td>
<td>—</td>
<td>No evidence of primary tumor</td>
</tr>
<tr>
<td>T1</td>
<td>I</td>
<td>I</td>
<td>Tumor limited to ovaries (1 or both)</td>
</tr>
<tr>
<td>T1a</td>
<td>IA</td>
<td>I</td>
<td>Tumor limited to 1 ovary; capsule intact, no tumor on ovarian surface. No malignant cells in ascites or peritoneal washings*</td>
</tr>
<tr>
<td>T1b</td>
<td>IB</td>
<td>I</td>
<td>Tumor limited to both ovaries; capsule intact, no tumor on ovarian surface. No malignant cells in ascites or peritoneal washings</td>
</tr>
<tr>
<td>T1c</td>
<td>IC</td>
<td>I</td>
<td>Tumor limited to 1 or both ovaries with any of the following: capsule ruptured, tumor on ovarian surface, malignant cells in ascites or peritoneal washings</td>
</tr>
<tr>
<td>T2</td>
<td>II</td>
<td>II</td>
<td>Tumor involves 1 or both ovaries with pelvic extension and/or implants</td>
</tr>
<tr>
<td>T2a</td>
<td>IIA</td>
<td>II</td>
<td>Extension and/or implants on uterus and/or tube(s). No malignant cells in ascites or peritoneal washings</td>
</tr>
<tr>
<td>T2b</td>
<td>IIB</td>
<td>II</td>
<td>Extension and/or implants to other pelvic tissues. No malignant cells in ascites or peritoneal washings</td>
</tr>
<tr>
<td>T2c</td>
<td>IIC</td>
<td>II</td>
<td>Pelvic extension and/or implants (T2a or T2b/IIA or IIB) with malignant cells in ascites or peritoneal washings</td>
</tr>
<tr>
<td>T3 and/or N1</td>
<td>III</td>
<td>III</td>
<td>Tumor involves 1 or both ovaries with confirmed peritoneal metastasis outside the pelvis (including liver capsule metastasis and/or regional lymph node metastasis [N1])</td>
</tr>
<tr>
<td>T3a</td>
<td>IIIA</td>
<td>III</td>
<td>Microscopic peritoneal metastasis beyond pelvis (no macroscopic tumor)</td>
</tr>
<tr>
<td>T3b</td>
<td>IIIB</td>
<td>III</td>
<td>Macroscopic peritoneal metastasis beyond pelvis 2 cm or less in greatest dimension</td>
</tr>
<tr>
<td>T3c and/or N1</td>
<td>IIIC</td>
<td>III</td>
<td>Peritoneal metastasis beyond pelvis more than 2 cm in greatest dimension and/or regional lymph node metastasis</td>
</tr>
<tr>
<td>(M1)</td>
<td>IV</td>
<td>IV</td>
<td>Distant metastasis (excludes peritoneal metastasis), including positive cytology in a pleural effusion or parenchymal liver metastasis</td>
</tr>
</tbody>
</table>

* The presence of nonmalignant ascites is not classified. The presence of ascites does not affect staging unless malignant cells are present.

**Regional Lymph Nodes (N): TNM Classification**

<table>
<thead>
<tr>
<th>N</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>NX</td>
<td>Regional lymph nodes cannot be assessed</td>
</tr>
<tr>
<td>N0</td>
<td>No regional lymph node metastasis</td>
</tr>
<tr>
<td>N1</td>
<td>Regional lymph node metastasis</td>
</tr>
</tbody>
</table>

**Distant Metastasis (M): TNM Classification**

<table>
<thead>
<tr>
<th>M</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>MX</td>
<td>Distant metastasis cannot be assessed</td>
</tr>
<tr>
<td>M0</td>
<td>No distant metastasis</td>
</tr>
<tr>
<td>M1</td>
<td>Distant metastasis (excludes peritoneal metastasis)</td>
</tr>
</tbody>
</table>
Stage Groupings

<table>
<thead>
<tr>
<th>TNM Stage Groupings</th>
<th>FIGO Stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage IA T1a N0 M0</td>
<td>Stage IA</td>
</tr>
<tr>
<td>Stage IB T1b N0 M0</td>
<td>Stage IB</td>
</tr>
<tr>
<td>Stage IC T1c N0 M0</td>
<td>Stage IC</td>
</tr>
<tr>
<td>Stage IIA T2a N0 M0</td>
<td>Stage IIA</td>
</tr>
<tr>
<td>Stage IIB T2b N0 M0</td>
<td>Stage IIB</td>
</tr>
<tr>
<td>Stage IIC T2c N0 M0</td>
<td>Stage IIC</td>
</tr>
<tr>
<td>Stage IIIA T3a N0 M0</td>
<td>Stage IIIA</td>
</tr>
<tr>
<td>Stage IIIB T3b N0 M0</td>
<td>Stage IIIB</td>
</tr>
<tr>
<td>Stage IIIC T3c N0 M0</td>
<td>Stage IIIC</td>
</tr>
<tr>
<td>Any T N1 M1</td>
<td>Stage IV</td>
</tr>
</tbody>
</table>

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 following 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 prior to 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).

**Vessel Invasion**

By AJCC/UICC convention, vessel invasion (lymphatic or venous) does not affect the T category indicating local extent of tumor unless specifically included in the definition of a T category. In all other cases, lymphatic and venous invasion by tumor are coded separately as follows.

**Lymphatic Vessel Invasion (L)**
- LX: Lymphatic vessel invasion cannot be assessed
- L0: No lymphatic vessel invasion
- L1: Lymphatic vessel invasion

**Venous Invasion (V)**
- VX: Venous invasion cannot be assessed
- V0: No venous invasion
- V1: Microscopic venous invasion
- V2: Macroscopic venous invasion

**F. Location of Tumor**

If an ovarian tumor does not replace the parenchyma, its location within the ovary may provide a clue as to its origin and nature. For example, confinement to or centering of a tumor in the hilus generally favors metastatic carcinoma over primary carcinoma or an origin from some structure in the hilus. Also, location of a malignant component in a tumor that is predominantly benign may be important because a focal carcinoma extending to the external surface of a tumor is probably associated with a poorer prognosis than an otherwise similar focus located centrally. Distinction of origin on the surface of the ovary from invasion of an underlying carcinoma may be important in the decision to classify a tumor as either an ovarian or as a primary peritoneal tumor (see Note O).

**G. Suggestions for Sampling for Microscopic Examination**

**Surface Epithelium**

The external surface of the ovary should be handled as gently as possible; rubbing or scraping it or allowing it to dry should be avoided.

* The surface epithelium and its inclusion glands and cysts and adjacent stroma are generally considered the source of most epithelial cancers of the ovary but have not been studied carefully by most pathologists. One reason is the great fragility of the surface epithelium, resulting in its absence in most microscopic specimens. Occasionally, dysplastic lesions and carcinoma can be identified in the surface epithelium and its inclusions if they are examined carefully. Gentle handling of the specimen and meticulous microscopic examination are especially important in patients who have had a prophylactic oophorectomy because of a family history of ovarian cancer with or without breast cancer or documentation of a BRCA mutation. It may be
possible to detect tiny but potentially fatal carcinomas by careful examination of the ovaries from such patients. 

**Primary Tumor**

One section for each 1 to 2 cm of tumor largest dimension is generally recommended, with modification based on the degree of homogeneity or heterogeneity of the tumor and the difficulty of diagnosis. 

Some sections should include the ovarian surface closest to the tumor on gross examination, with the number depending on the degree of suspicion of surface involvement.

Adhesions of tumor, sites of rupture, and resection margins, if pertinent, should be sampled and labeled specifically if necessary for microscopic identification.

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**Sampling Issues:** The recommendation for the number of sections to be taken of an ovarian tumor that may be at least focally malignant is a general guideline, with the pathologist determining in each case how many sections are necessary. If a tumor is obviously malignant and homogeneous throughout on gross examination, large numbers of sections are not required in most cases. In contrast, if there is great variability in the gross appearance of the sectioned surfaces or opened cysts, and the presence or extent of malignant change cannot be easily determined on gross examination, it may be necessary to take large numbers of sections to sample the tumor adequately. Mucinous tumors, particularly those with solid areas, solid teratomas, and primitive germ cell tumors, often require especially careful gross examination and judicious and extensive sampling. Often, additional sampling of a tumor that poses a problem in differential diagnosis is more informative than special studies such as immunohistochemical staining.

**Fallopian Tube(s)**

- One section of each, if no gross lesion is present.
- Representative sections of tumor, if present, to determine its distribution and the appearance of the adjacent epithelium (Note F).

**Uterus**

- Tumor grossly present: sections necessary to determine its extent, including depth of invasion of myometrium if tumor possibly originated in endometrium, and to determine its relation to ovarian tumor (metastatic to, metastatic from, independent primary).

**Omentum**

- Representative sampling of grossly identifiable tumor.
- Multiple representative sections when no tumor is detected grossly (see Note P).
- For borderline tumor or immature teratoma with grossly apparent implants, multiple sections of the implants (see Note P).

**Lymph Nodes**

- Representative sections of grossly positive lymph nodes.
- If lymph nodes appear free of tumor, samples of every node in the specimen(s).
Other Staging Biopsy Specimens
- Embed entirely unless grossly positive for tumor.
- If grossly positive, a representative section.

Other Organ or Tissue Removed
- Sections adequate to determine presence or absence and location and extent of tumor, if present.
- Resection margins, if applicable.

H. Site(s) of Origin of Tumor
When a tumor involves both the ovary and the fallopian tube, it may be difficult to determine the primary site of the tumor in some cases. Typically, the tumor predominates and is obviously primary in one or the other organ, almost always the ovary. Occasionally, however, the ovary and tube are fused to form a solid or cystic mass with destruction of most or all landmarks. In such cases the tumor is almost always designated a primary ovarian cancer because the frequency of ovarian cancer is much greater than that of tubal cancer. Finding what appears to be in situ carcinoma in the tube adjacent to the main tumor mass is not always a reliable criterion for the diagnosis of a primary carcinoma of the tube since carcinoma that has extended into the tube can grow along its mucosal surface and closely simulate carcinoma in situ.

I. Growth Pattern of Tumor
Whether an ovarian cancer is growing as a single mass or in discrete nodules is 1 of several clues that it is primary or metastatic, respectively. Many metastatic tumors that are not large enough to replace the ovary form 2 or more separate nodules, in contrast to primary cancers, which usually form single nodules or masses.

J. Mixtures of Histological Types of Tumors
If neoplasia of more than 1 cell type accounts for 10% or greater of a surface epithelial cancer, the tumor is diagnosed as one of mixed cell types. Therefore, it is important to identify and estimate the proportion(s) of various cell types within epithelial cancers. Quantitation of various epithelial cell types within a specimen, as well as quantitation of tumor types within primitive germ cell tumors, may be important prognostically. Although unproven, it is reasonable to assume that the quantity of the more malignant elements in tumors that have varying combinations of benign, borderline, and carcinomatous components also has prognostic significance.

K. Vessel Invasion
Although the prognostic significance of vascular space invasion in ovarian cancer has not been demonstrated by multivariate analysis, studies of cancers in many other organs suggest that this finding might also be important in ovarian cancer. In addition, vascular space invasion is much more common in cancers that are metastatic to the ovary and therefore is worth noting, particularly if there is doubt about the primary or metastatic nature of the tumor.

L. Other Lesions
The presence of other lesions, such as endometriosis and atypicality of surface epithelium or surface epithelial inclusion glands or cysts, is occasionally important in
determining the origin of an ovarian cancer. Also, the presence of endometriosis, particularly if it is in continuity with an endometrioid cancer (either carcinoma or stromal sarcoma), is a very important clue as to the primary nature of the ovarian tumor in cases in which it may be difficult otherwise to exclude metastasis from a synchronous or asynchronous cancer of the uterine corpus.

M. Contralateral Ovary
“Contralateral ovary” refers to the ovary that is non-dominant, either because it is involved by a tumor that is similar to but smaller than the dominant ovarian tumor, or because it appears negative for tumor or contains only what appears to be metastatic tumor on gross examination. If the contralateral ovary contains only focal tumor, the gross and microscopic examination should concentrate on determining whether the tumor is independently primary or metastatic from the dominant ovary. Metastatic involvement is supported by the same criteria that are used to distinguish primary and metastatic cancers in general, such as the presence of multiple nodules, surface implants, and vascular space invasion, which favor metastasis.

N. Uterine Tumor
When carcinoma involves the ovary and uterus, it may be difficult to determine whether one is dealing with a primary ovarian carcinoma with spread to the uterus, vice versa, or independent primary tumors. This problem arises most commonly when the tumors in both organs are of endometrioid type but also exists occasionally when the tumors are of serous or other cell types. There are numerous criteria for determining which of the 3 alternative explanations for the coexistence of ovarian and uterine tumors is correct. The size and distribution of the tumors, the presence of a precancerous lesion in either organ (atypical hyperplasia of the endometrium, endometriosis or adenofibroma of the ovary), microscopic comparisons of the tumors, DNA ploidy findings, and molecular genetic studies of the 2 tumors have all been used to facilitate the differential diagnosis. Over-reliance on a single criterion, however, has resulted in a lack of consensus in the literature. The very good prognosis of endometrioid carcinomas confined to the uterine corpus and 1 or both ovaries, however, suggests that in most cases of this combination, the tumors are independently primary.

O. Other Lesions
The occurrence of benign fallopian tube type epithelium in the peritoneum (endosalpingiosis) and in lymph nodes (mullerian inclusion glands, endosalpingiosis) has raised the possibility of multicentric origin of serous malignant tumors in the ovary and on the peritoneum or in lymph nodes. Therefore, it is important to identify the above benign lesions when they are present. Currently, serous cancers that are characterized by a grossly predominant mass on the peritoneum or in the omentum with no involvement of the ovary or with tumor in the ovary that invades to a depth of less than 5 mm are generally considered to be primary peritoneal cancers. An interesting finding in association with some serous borderline tumors of the ovary is an independent primary serous borderline tumor in a pelvic or para-aortic lymph node arising from a mullerian glandular inclusion.

P. Omentum
In 1 study, 43 microscopic metastases were detected in 22% of grossly negative specimens of omentum, with a mean tumor diameter of 6.7 mm. Because of this type of
experience, and because the finding of a single metastatic lesion on microscopic examination may alter the management of the patient, the omentum should be adequately sampled in those cases. In cases of serous borderline tumors and immature teratomas of the ovary, it is also suggested that multiple sections of grossly recognizable implants be taken, since the implants of these tumors may vary from noninvasive to invasive, and from mature to immature, respectively. Identification of a single invasive or immature implant may alter the prognosis and therapy.

References


**Bibliography**


