ABNORMAL MENSTRUAL BLEEDING PATTERNS

Includes Menorrhagia, Metrorrhagia, Oligomenorrhea, and Polymenorrhea.
For Amenorrhea, see "Amenorrhea" Guideline. For Dysmenorrhea, see "Dysmenorrhea" Guideline.

INFORMATION REQUIRED  Any history

All Applicants:
- Report of Medical Examination to include the following:
  - Description of bleeding pattern
  - Etiology, if known.
  - Complications, if any.
  - Treatment
  - Recommendations for follow-up over the next 3 years.

Applicants with Menorrhagia or Metrorrhagia:
- Evaluation and work-up required (see comments)
- Copy of related diagnostic tests.

<table>
<thead>
<tr>
<th>CLEARANCE CRITERIA</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Abnormal or irregular menstrual bleeding pattern defined as one of the following:</td>
<td>RN</td>
<td>CLEAR</td>
</tr>
<tr>
<td>2. No significant associated complications, e.g., anemia, dysmenorrhea, or amenorrhea.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Meets clearance criteria 1-2, AND**
- **Polymenorrhea**
  - Resolved, hormonally controlled, or well tolerated, for at least the past 3 months.

**Meets clearance criteria 1-2, AND**
- **Oligomenorrhea (Premenopausal)**
  - Stable menstrual pattern for at least the past 6 months.

**Meets clearance criteria 1-2, AND**
- **Oligomenorrhea (Menopausal)**

**Meets clearance criteria 1-2, AND**
- **Menorrhagia**
  - Evaluation and work-up complete.
  - Resolved, hormonally controlled, or well tolerated, for at least the past 3 months.

**Meets clearance criteria 1-2, AND**
- **Metrorrhagia**
  - Evaluation and work-up complete.
  - Resolved, hormonally controlled, or well tolerated, for at least the past 3 months.

**Does not meet clearance criteria due to one or more of the following:**
- **Undefined abnormal or irregular menstrual bleeding pattern.**

RN

See "Dysfunctional Uterine Bleeding" Guideline.

(continued on next page)
ABNORMAL MENSTRUAL BLEEDING PATTERNS

Does not meet clearance criteria due to one or more of the following:

- **Polymenorrhea**: Not resolved, hormonally controlled, or well tolerated for at least the past 3 months.
- **Oligomenorrhea** (Not menopausal): Menstrual pattern not stable for at least the past 6 months.
- **Menorrhagia or Metrorrhagia**:
  - Evaluation and work-up not complete.
  - Not resolved, hormonally controlled, or well tolerated for at least the past 3 months.
- Significant associated complications.

<table>
<thead>
<tr>
<th>MED ADVISOR</th>
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</thead>
<tbody>
<tr>
<td>Risk varies - assess based on detailed history.</td>
</tr>
</tbody>
</table>

**DIAGNOSTIC CODES**

- 626.3 Menorrhagia
- 626.6 Metrorrhagia
- 626.1 Oligomenorrhea
- 626.2 Polymenorrhea

Cross Reference ICD.9.CM

**NOTES AND INSTRUCTIONS FOR REVIEWERS**

Reviewers to Consider:
- None

**COMMENTS**

**Background:** Any bleeding that is excessive in duration, frequency, or amount is considered abnormal and requires investigation accordingly.

**Evaluation:** Diagnostic tests to consider as appropriate include prolactin, LH, FSH, DHEA-s, free testosterone, LFTs, thyroid function tests, coagulation profile, CBC, and pregnancy test.

**Definitions:**

- **Oligomenorrhea:** Infrequently occurring menses at intervals greater than 35 days.
- **Polymenorrhea:** Frequently occurring menses at intervals of 21 days or less.
- **Metrorrhagia:** Irregularly occurring bleeding.
- **Menorrhagia:** Regularly occurring bleeding excessive in duration or flow, usually defined as 8 days in length or greater.

Literature review available.
AMENORRHEA

Includes Absence of Menses.

Amenorrhea Due to Birth Control Pills or Pregnancy; Guideline not Applicable.

INFORMATION REQUIRED

If history within the past 5 years.

Applicants With Absent Menses, i.e., Absent Bleeding Pattern Less Than 6 Months Duration:
- Report of Medical Examination to include the following:
  - Etiology
  - Specific management recommendations.

Applicants With Amenorrhea, i.e., Absent Bleeding Pattern Greater Than 6 Months Duration:
- Specialist Evaluation (Gynecologist) within the past 6 months to include the following:
  - Etiology
  - Specific management recommendations.
  - Copy of related diagnostic tests.

A. Absence of Menses: Absent Bleeding Pattern Less Than 6 Months Duration.
B. Amenorrhea: Absence of Menses Greater Than 6 Months.

**CLEARANCE CRITERIA**

<table>
<thead>
<tr>
<th>MEETS CLEARANCE CRITERIA</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meets clearance criteria 1-2, AND</td>
<td>RN</td>
<td>CLEAR</td>
</tr>
</tbody>
</table>
| • Amenorrhea resolved (with or without hormonal treatment), i.e., normal menses for at least the past 3 months. | *
| Meets clearance criteria 1-2, AND | MED ADVISOR | *
| • Current amenorrhea | *
| • Etiology determined | *
| • Provider provides specific management recommendations | *

**Diagnosis Considerations**

- Coexisting eating disorder.
- Coexisting polycystic ovarian syndrome (Stein-Leventhal syndrome).
- Coexisting congenital disorders.

**Background:**
Amenorrhea is one of the most common reasons that women seek medical attention. Frequently there is no underlying cause and the abnormal bleeding pattern resolves spontaneously or with hormonal or lifestyle changes.
Secondary Amenorrhea: Most common type. Defined as the absence of menstruation for at least 6 months in a woman with previously normal and regular menses.

Evaluation and Work-Up of Secondary Amenorrhea: A standard work-up commonly includes the following:
- History and physical exam
- Pregnancy test
- TSH, T4
- Glucose
- Prolactin
- Estrogen status assessment, i.e., progesterin challenge or plasma estradiol, FSH, and LH.
- If indicated, testosterone, DHEA-S, 17-hydroxy-progesterone.

Literature review available.
DYSMENORRHEA

All Applicants:
- Report of Medical Examination to include the following:
  - Symptoms
  - Treatment
  - Recommendations for follow-up over the next 3 years.

CLEARANCE CRITERIA

1. No, or mild, pelvic pain.
2. No, or minimal, interference of symptoms with activities of daily living.

Meets clearance criteria 1-2, AND
- If treated, pain well controlled with NSAIDs or oral contraceptive pills.

Does not meet clearance criteria due to one or more of the following:
- Moderate to severe pelvic pain.
- Symptoms significantly interfere with activities of daily living.
- If treated, pain well controlled with narcotics.

DIAGNOSTIC CODES

625.3 Dysmenorrhea

Cross Reference ICD.9.CM

NOTES AND INSTRUCTIONS FOR REVIEWERS:

Reviewers to Consider:
- None

COMMENTS:

Background: Dysmenorrhea is defined as pelvic pain at or around the time of menstruation. There are two types:
- Primary dysmenorrhea - pain without pathological physical findings.
- Secondary dysmenorrhea - pain occurring prior to or during menses, often more severe than primary, having a secondary pathologic (structural) cause.

Symptoms: Symptoms range from mild pelvic discomfort or cramping on the first day of bleeding to severe, intense, cramp-like pain lasting 2-7 days; often associated with gastrointestinal upset, back ache, thigh pain, and headache.

Treatment: Ibuprofen, Naproxen sodium, or Aspirin. [Griffith's 1996]

Literature review available.
PREMENSTRUAL SYNDROME

GYN 3.4

"For Premenstrual Syndrome, "See Mental Health" Guidelines.

INFORMATION REQUIRED

If history within the past 5 years.

All Applicants:

- Report of Medical Examination to include the following:
  - Symptoms
  - Treatment
  - Recommendations for follow-up over the next 3 years.

CLEARANCE CRITERIA

<table>
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<tr>
<th>REVIEWER</th>
<th>GUIDANCE</th>
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<tbody>
<tr>
<td>RN</td>
<td>CLEAR</td>
</tr>
</tbody>
</table>

1. No, resolved, or mild symptoms.
2. If symptoms, well controlled with diuretics, NSAIDs, oral contraceptive pills, diet, or exercise.
3. No, or minimal, interference of symptoms with activities of daily living.

Meets clearance criteria 1-3, AND

- "If treated, treatment does not include antidepressants, narcotics, or counseling.

Does not meet clearance criteria due to one or more of the following:

- Current, moderate to severe, symptoms.
- Symptoms not well controlled with NSAIDs, oral contraceptive pills, diet, exercise, or diuretics.
- Symptoms significantly interfere with activities of daily living.
- Treatment includes antidepressants, narcotics, or counseling.

MED ADVISOR

Risk varies – assess based on detailed history.

DIAGNOSTIC CODES

625.4 Premenstrual Syndrome

Cross Reference ICD.9.CM

NOTES AND INSTRUCTIONS FOR REVIEWERS:

Reviewers to Consider:

- None

COMMENTS:

Background: Premenstrual syndrome is a constellation of symptoms that occur prior to menstruation. Symptoms may be mild or severe enough to interfere significantly with a patient's life.

Symptoms: Depressed mood, mood swings, irritability, difficulty concentrating, fatigue, edema, breast tenderness, headaches, weight gain, depressed mood, and food cravings.

Treatment: No single drug works for all women. Drugs that are used with varying degrees of success include diuretics, ibuprofen or acetaminophen, anti-depressants, alprazolam, Buspirone, magnesium, elemental calcium, Vitamin B6, Vitamin E, and evening primrose oil. Some women obtain relief with diet modification and exercise.

Literature review available.
**INFORMATION REQUIRED: Any history.**

All Applicants:
- Specialist Evaluation (General Surgeon or Oncologist) within the past 1 year to include the following:
  - Documentation of recurrences within the past 5 years.
  - Recommendations for follow-up over the next 3 years.
If Applicable:
- Discharge summary for all related hospitalizations.

**CLEARANCE CRITERIA**

| 1. Treatment complete. |
| 2. No history of recurrences for at least the past 5 years. |

| Meets clearance 1-2 criteria, AND |
| No use of adjuvant therapy. |
| MED ADVISOR | CLEAR |

| Does not meet clearance criteria due to one or more of the following: |
| Treatment not complete. |
| History of recurrences within the past 5 years. |
| MED ADVISOR | DEFER |

Risk varies - assess based on detailed history.

| Does not meet clearance criteria due to one or more of the following: |
| Current use of adjuvant therapy. |
| MED ADVISOR | | |

**DIAGNOSTIC CODES**

| 183.0  Ovarian Cancer |
| Cross Reference | ICD.9.CM |

**NOTES AND INSTRUCTIONS FOR REVIEWERS:**

Reviewers to Consider:
- None

**COMMENTS:**

Prognostic Factors: At the conclusion of a comprehensive laparotomy, the clinical findings and the histology are used to select postoperative therapy. In addition, new prognostic factors are being evaluated that may also be used in identifying groups of patients in whom more specific histologic treatments or more aggressive therapy is indicated. Clinically pathologic findings determined to be clinically useful include the following:

- FIGO stage
- Histologic subtype
- Histologic grade
- Factors associated with tumor dissemination
  - Malignant ascites
  - Malignant peritoneal washings
  - Tumor excrencences on ovarian surface

Effective 1/15/99
OVARIAN CANCER

Ruptured capsule
Dense ovarian adhesions
Volume of residual disease following cytoreductive surgery

The tumor stage remains the most important prognostic variable. Few trials provide an accurate assessment regarding the long-term survival of patients with early-stage ovarian cancer because earlier studies often included inadequately staged patients. Stage I patients with well- or moderately well-differentiated tumors have a greater than 90% 5-year survival rate. Patients with stage I disease with poor prognostic features are often included in treatment protocols for patients with stage II disease. This group of patients has been termed early-stage disease with unfavorable characteristics. However, there remains limited information regarding the actual survival impact of some of the factors used to characterize patients having an unfavorable prognosis. Rupture of the capsule increases the stage to IC. However, in a Swedish series, no adverse effect on survival could be established for early-stage patients in whom the capsule was ruptured during surgery. Furthermore, in contrast to the established adverse effect of malignant ascites, there is limited information regarding the prognostic significance of positive peritoneal cytology. Tumor adherence in the presence of dense adhesions has also been considered an adverse prognostic factor and such patients should be considered as having stage II disease even in the absence of pathologic confirmation. Tumor size, bilaterality, and cytologically negative ascites have no prognostic significance. The most reliable long-term survival data on accurately staged early-stage ovarian cancer patients are derived from studies of the Gynecologic Oncology Group (GOG). In these studies, unfavorable prognosis early-stage ovarian cancer patients have a 5-year survival rate of approximately 80%.

Patients with stage III disease have a 5-year survival rate of approximately 15% to 20%, which is dependent in large part on the volume of disease present in the upper abdomen. Patients with stage IV disease have less than a 5% 5-year survival rate.

Volume of residual following cytoreductive surgery for patients with advanced ovarian cancer has a significant impact on survival. Following the administration of postoperative cisplatin-based combination chemotherapy, 5-year survival rates for patients with optimal stage III disease (defined as no residual nodule greater than 1 cm in diameter) are approximately 35%.

The true prognostic impact of histologic subtype and grade in patients with epithelial ovarian cancer remains to be determined. In patients with early-stage ovarian cancer, grade is an accepted determinant of risk and is used to assign postoperative therapy as previously discussed. Studies have also identified an adverse prognostic effect of clear cell histology in early-stage ovarian cancer. In advanced-stage patients, mucinous histology and clear cell histology have also been shown to have an adverse prognostic significance. In a GOG analysis, there were no negative second-look laparotomies in patients with mucinous or clear cell tumors. Some but not all studies have also demonstrated that histologic grade has an impact on survival in patients with advanced-stage disease.

Serum CA125 levels frequently reflect the volume of disease and as such, in multivariate analysis, preoperative levels have failed to exert an independent prognostic effect on survival. However, postoperative CA125 levels were shown to be an independent prognostic variable. Most studies have also demonstrated that serum CA125 levels after three cycles of chemotherapy are accurate predictors for the probability of a patient achieving a complete remission. However, the CA125 level after three cycles of chemotherapy cannot be used as a guide for treatment decisions because of the lack of predictive power.

The prognostic significance of age on survival of patients with ovarian cancer has been recognized. Median survival is at least 2 years longer in women under the age of 65 compared with those over 65.

The prognostic significance of DNA ploidy and S-phase fraction has been examined in ovarian cancer. Investigators in Europe have now included aneuploidy in their selection of high-risk, early-stage ovarian cancer patients for adjuvant therapy. Controversy remains, however, as to the nature of the relationship between histologic grade and degree of aneuploidy. In the GOG, aneuploidy has not been included as a criteria for risk in early-stage disease. [MD Consult, 1998]

Literature review available.
OVARIAN CYSTS

Includes Functional Cysts and Complex Cysts.
For Polycystic Ovarian Syndrome, see Polycystic Ovarian Syndrome Guideline.
For Laparoscopy and Laparotomy, see "Gynecology Surgical Procedures" Guideline.

INFORMATION REQUIRED

All Applicants:
- Report of Medical Examination to include the following:
  - Type of cyst.
  - Size, location, and number of cysts.
  - Treatment, e.g., spontaneous resolution, aspiration, hormone therapy, excision.
  - Current status, to include an assessment of cyst(s) progression and/or regression.
  - Recommendations for follow-up over the next 3 years.
- Copy of ultrasound report with interpretation.

If Applicable:
- Copy of laparoscopy report with interpretation.
- Copy of pathology report with interpretation.
- Discharge summary of all related hospitalizations and surgeries.

CLEARANCE CRITERIA

<table>
<thead>
<tr>
<th>REVIEWER</th>
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</table>

1. Functional cyst(s), i.e., simple cysts (follicular), or ovulatory cysts (corpus luteum cyst), noted on examination, current or history of.

2. No, or resolved, symptoms.

3. If current cyst, no evidence of cyst progression or enlargement; determined by examining health care provider.

4. No history of complex cyst(s).

Meets clearance criteria 1 - 4, AND
- Cyst resolved; spontaneously, aspiration, hormone therapy, excision.

Meets clearance criteria 1 - 4, AND
- Current functional cyst, AND
- Cyst size < 2 cm.

Does not meet clearance criteria due to one or more of the following:
- Cyst size > 2 cm.
- Current symptoms.
- Evidence of cyst progression or enlargement; determined by examining health care provider.
- History of complex cyst(s).

DIAGNOSTIC CODES

<table>
<thead>
<tr>
<th>ICD-9-CM</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>620.2</td>
<td>Ovarian Cyst</td>
</tr>
</tbody>
</table>

Cross Reference ICD-9-CM

NOTES AND INSTRUCTIONS FOR REVIEWERS

Reviewers to Consider:
- None
COMMENTS

**Functional Cysts:** Most common ovarian cysts. Usually causes no symptoms and resolve in 1-3 menstrual cycles. Birth control pills suppress ovulation and may decrease size and frequency of functional cysts. The type of cyst can easily be distinguished on ultrasound. A small number of women have recurrent cysts that also respond well to birth control pills.

**Complex Cysts:** Include dermoid cysts, cystadenoma, and endometrioma. These cysts frequently require surgery. Small cysts occasionally require laparoscopy removal. Larger cysts require laparotomy and partial or total oophorectomy. Cysts are considered more serious post menopausal and should be immediately evaluated to rule-out cancer.

**Surgery:** Surgery is indicated to rule-out ovarian cancer or to relieve pain if:

- The cyst is large (>5 cm) or does not resolve after three menstrual cycles.
- Complex cysts (dermoid cysts, cystadenoma, endometrioma).
- Painful or bleeding cysts.
- Torsion of the cyst has occurred.

Literature review available.
**INFORMATION REQUIRED**

**Any history**

<table>
<thead>
<tr>
<th>All Applicants:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specialist Evaluation (Gynecologist) within the past 1 year to include the following:</td>
</tr>
<tr>
<td>- Date of diagnosis</td>
</tr>
<tr>
<td>- Signs and symptoms</td>
</tr>
<tr>
<td>- Treatment</td>
</tr>
<tr>
<td>- Current status</td>
</tr>
<tr>
<td>- Limitations or restrictions of ADLs</td>
</tr>
<tr>
<td>- Recommendations for follow-up over the next 3 years.</td>
</tr>
</tbody>
</table>

**If Applicable:**

| - Copy of most recent ultrasound report with interpretation. |
| - Copy of laparoscopy report with interpretation. |

**CLEARANCE CRITERIA**

1. No, resolved, or mild, symptoms.
2. Signs and symptoms do not interfere with activities of daily living.
3. No, or resolved, irregular or dysfunctional uterine bleeding.

<table>
<thead>
<tr>
<th>Meets clearance criteria 1 - 3, AND:</th>
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<tbody>
<tr>
<td>If symptom management requires treatment, symptoms well-controlled with oral contraceptives only.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Meets clearance criteria 1 - 3, AND:</th>
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<tbody>
<tr>
<td>If on hormone therapy, e.g., aldactone, symptoms well-controlled for at least the past 3 months.</td>
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</table>

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<thead>
<tr>
<th>Meets clearance criteria 1 - 3, AND:</th>
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</thead>
<tbody>
<tr>
<td>If on anti-diabetic agent, e.g., glucophage or Thiazolidinedione (actos), symptoms well-controlled for at least the past 3 months.</td>
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</table>

<table>
<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
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</thead>
<tbody>
<tr>
<td>- If on hormone therapy, e.g., aldactone, symptoms not well controlled for at least the past 3 months.</td>
</tr>
<tr>
<td>- If on anti-diabetic agent, e.g., glucophage or Thiazolidinedione (actos), symptoms not well controlled for at least the past 3 months.</td>
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</table>

<table>
<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
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<tbody>
<tr>
<td>- Current, moderate to severe, symptoms.</td>
</tr>
<tr>
<td>- Signs and symptoms (see comments) interfere with activities of daily living.</td>
</tr>
<tr>
<td>- Current irregular or dysfunctional uterine bleeding.</td>
</tr>
</tbody>
</table>

**DIAGNOSTIC CODES**

- 256.4 Polycystic Ovarian Syndrome
- Cross Reference ICD.9.CM

Effective 2/12/2004
NOTES AND INSTRUCTIONS FOR REVIEWERS

Reviewers to Consider:
- None

COMMENTS

Background: Polycystic ovarian disease (PCOD) is characterized by a state of chronic oligo-ovulation and/or anovulation culminating in oligomenorrhea and/or amenorrhea. The ovaries are usually enlarged and lined with follicles in all stages of development but most are atretic. No ideal treatment exists although hormones and birth control pills are often used to regulate the menstrual cycle. [Griffith's 1998]

Signs and Symptoms: Amenorrhea, oligomenorrhea, obesity, hirsutism, acne, dysfunctional uterine bleeding, infertility, hypertension, virilism, enlarged ovaries, enlarged clitoris, deep voice.

*Literature review available.*
PELVIC INFLAMMATORY DISEASE (PID)

Encompasses a Spectrum of Inflammatory Disorders of the Upper Genital Tract Including Any Combination of:
1) Endometritis; 2) Salpingitis; 3) Tubo-Ovarian Abscess; and 4) Pelvic Peritonitis.

### INFORMATION REQUIRED

| Any history. |

All Applicants:
- Specialist Evaluation (Gynecologist) within the past 1 year to include the following:
  - Number of episodes.
  - Dates of episodes.
  - Treatment
  - Treatment complications, if any.
  - Current status
  - Recommendations for follow-up over the next 3 years.

If Applicable:
- Discharge summary for all related hospitalizations.

### CLEARANCE CRITERIA

<table>
<thead>
<tr>
<th>REVIEWER</th>
<th>GUIDANCE</th>
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</thead>
<tbody>
<tr>
<td><strong>1.</strong> One or two episodes of PID.</td>
<td>RN</td>
</tr>
<tr>
<td><strong>2.</strong> No, or resolved, symptoms.</td>
<td></td>
</tr>
<tr>
<td><strong>3.</strong> Treatment complete and infection resolved for at least the past 3 months.</td>
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</tr>
<tr>
<td><strong>4.</strong> No post treatment complications to include chronic pelvic pain.</td>
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<tr>
<td><strong>5.</strong> No history of tubo-ovarian abscess.</td>
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</table>

**Meets clearance criteria 1-5, AND**

- If hysterectomy for PID, post surgery greater than 3 months.

<table>
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<tr>
<th>REVIEWER</th>
<th>GUIDANCE</th>
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</thead>
<tbody>
<tr>
<td>RN</td>
<td>DEFER</td>
</tr>
<tr>
<td></td>
<td>Deferral letter requires review by the screening manager.</td>
</tr>
</tbody>
</table>

**Does not meets clearance criteria due to one or more of the following:**

- Current symptoms.
- Treatment *not* complete or infection *not* resolved for at least the past 3 months.
- If hysterectomy for PID, post surgery *less than* 3 months.

<table>
<thead>
<tr>
<th>REVIEWER</th>
<th>GUIDANCE</th>
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<tbody>
<tr>
<td>MED ADVISOR</td>
<td>MED ADVISOR</td>
</tr>
<tr>
<td></td>
<td>Risk varies - assess based on detailed history. Consider GYN Accommodation.</td>
</tr>
</tbody>
</table>

**Does not meets clearance criteria due to one or more of the following:**

- More than two episodes of PID, i.e., recurrent infection.
- Post treatment complications to include chronic pelvic pain.
- History of tubo-ovarian abscess.

### DIAGNOSTIC CODES

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>614.0</td>
<td>Pelvic Inflammatory Disease, Acute</td>
</tr>
<tr>
<td>614.1</td>
<td>Pelvic Inflammatory Disease, Chronic</td>
</tr>
<tr>
<td>614.6</td>
<td>Pelvic Inflammatory Disease, With Adhesions</td>
</tr>
<tr>
<td>614.2</td>
<td>Salpingitis</td>
</tr>
<tr>
<td>615.9</td>
<td>Endometritis</td>
</tr>
<tr>
<td>614.2</td>
<td>Tubo-Ovarian Abscess</td>
</tr>
</tbody>
</table>

Cross Reference: ICD.9.CM
PELVIC INFLAMMATORY DISEASE (PID)

NOTES AND INSTRUCTIONS FOR REVIEWERS:

Reviewers to Consider:
- None

COMMENTS:

Background: PID comprises a spectrum of inflammatory disorders of the upper genital tract among women and may include any combination of: 1) Endometritis; 2) Salpingitis; 3) Tubo-ovarian abscess; and 4) Pelvic peritonitis. PID may recur in 23-42% of women with a previous diagnosis. Twenty percent will have a recurrence within 1 year. Whether or not recurrence is caused by a new infection is not always clear. Frequently, pelvic pain persists after resolution of the infection. In a study of more than 100 women admitted to the hospital for treatment of PID, 24% reported persistent pelvic pain 6 months after discharge. Once damaged, fallopian tubes are believed to be more susceptible to bacterial colonization as a result of depressed host defenses, persistent inflammation, or other mechanisms (Kottman, 1995). Chronic PID needs further treatment to prevent infertility and more adhesions. Patients with a history of PID generally require no extra follow-up but are at risk for ectopic pregnancy and complications from the adhesions.

Literature review available.
All Applicants:
- Specialist Evaluation (Gynecologist) within the past 1 year to include the following:
  - Etiology
  - Symptoms
  - Treatment
  - Current status
  - Recommendations for follow-up over the next 3 years.

If Applicable:
- Copy of endometrial biopsy with interpretation.
- Discharge summary for all related hospitalizations.

### CLEARANCE CRITERIA

<table>
<thead>
<tr>
<th>Meets clearance criteria, AND</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Evaluation and diagnostic work-up complete.</td>
<td>RN</td>
<td>CLEAR</td>
</tr>
<tr>
<td>• Etiology determined: benign, or unknown, cause with no underlying pathology.</td>
<td></td>
<td>PCMO FOLLOW-UP</td>
</tr>
<tr>
<td>• No, resolved, or well tolerated, symptoms.</td>
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<td>As recommended by provider.</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Evaluation and diagnostic work-up not complete.</td>
<td>RN</td>
<td>DEFER</td>
</tr>
<tr>
<td>• Etiology not determined.</td>
<td></td>
<td>Requires evaluation and specific diagnosis for clearance. Correlate with diagnostic guideline.</td>
</tr>
<tr>
<td>• Symptoms not resolved or well tolerated.</td>
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<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Etiology determined: Known cause or underlying pathology.</td>
<td>RN</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Correlate with specific diagnostic guideline.</td>
</tr>
</tbody>
</table>

### DIAGNOSTIC CODES

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>626.8</td>
<td>Dysfunctional Uterine Bleeding</td>
</tr>
</tbody>
</table>

Cross Reference ICD.9.CM

### NOTES AND INSTRUCTIONS FOR REVIEWERS:

Reviewers to Consider:
- None
DYSFUNCTIONAL UTERINE BLEEDING

Background: Dysfunctional Uterine Bleeding (DUB) as defined by the American College of Obstetricians and Gynecologists is 'bleeding from the uterine endometrium unrelated to anatomic lesions of the uterus'. Disruption of normal ovarian function, or anovulation, was considered part the definition, and midcycle staining associated with ovulation, menstrual irregularity associated with corpus luteum defect, and uterine bleeding secondary to 'blood dyscrasias, submucous myomas, endometrial polyps, uterine carcinoma, and accidents of pregnancy' was not to be considered dysfunction. By this definition, the diagnosis of DUB is a diagnosis of exclusion. Implicit is that its anovulatory nature has already been demonstrated. Any bleeding that is excessive in duration, frequency, or amount for a particular patient should be considered abnormal and investigated accordingly. [Novak's Textbook of Gynecology, 11th ed.]

Literature review available.
ENDOMETRIAL CANCER

INFORMATION REQUIRED

All Applicants:
- Specialist Evaluation (General Surgeon or Oncologist) within the past 1 year to include the following:
  - Documentation of recurrences within the past 5 years.
  - Recommendations for follow-up over the next 3 years.
If Applicable:
- Discharge summary for all related hospitalizations.

CLEARANCE CRITERIA

1. Treatment: complete.
2. No history of recurrences for at least the past 5 years.

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-2, AND</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>No use of adjuvant therapy.</td>
<td>MED ADVISOR</td>
<td>CLEAR</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment not complete.</td>
</tr>
<tr>
<td>History of recurrences within the past 5 years.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current use of adjuvant therapy.</td>
</tr>
</tbody>
</table>

MED ADVISOR

Clearance Criteria:
- Deferral letter requires review by screening manager.

Risk varies - assess based on detailed history.

DIAGNOSTIC CODES

182.0 Endometrial Cancer
Cross Reference ICD.9.CM

NOTES AND INSTRUCTIONS FOR REVIEWERS:

Reviewers to Consider:
- None

COMMENTS:

Background: Tumors of the uterine fundus comprise the most common group of gynecologic malignancies. Annual incidence figures for the United States have remained stable at 34,000 to 36,000 cases over the past decade. Deaths from disease occur in 4,000 to 5,000 women per year. Women with high risk or advanced disease have a poor prognosis and account for the most uterine cancer deaths.

Prognosis: Histopathologic risk factors have been extensively evaluated during the past two decades. Major prognostic factors associated with the uterine component of the tumor are grade or cell type, depth of myometrial invasion, and extension to the cervix. Less important are extent of uterine cavity involvement, lymph-vascular space invasion, and tumor vascularity. Obviously, women whose tumors have spread beyond the uterus have a poor prognosis. The major extrauterine risk factors are adnexal metastases, pelvic or paraaortic lymph node spread, positive peritoneal cytology, peritoneal implant metastases, and distant organ metastases.
A detailed analysis of nearly 1000 patients has been presented by the Gynecologic Oncology Group. The risk for developing recurrent disease was greatest in women whose tumors had metastasized to pelvic or paraaortic lymph nodes, demonstrated gross intraperitoneal spread, or contained unequivocal lymph-vascular space invasion. Not surprisingly, an exceptionally high incidence of recurrence was noted in cases with two or more risk factors. Based on the findings of this and other surgical staging trails, the International Federation of Gynecology and Obstetrics (FIGO) adopted a surgical staging system for uterine fundal cancers in 1988. [DeVita: Cancer: Principles and Practice of Oncology, 5th ed., Copyright © 1997 Lippincott-Raven Publishers]

Surgical Staging of Uterine Fundal Tumors:

<table>
<thead>
<tr>
<th>Stage</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage I</td>
<td>The tumor is confined to the uterine fundus.</td>
</tr>
<tr>
<td>Stage IA</td>
<td>Tumor is limited to the endometrium.</td>
</tr>
<tr>
<td>Stage IB</td>
<td>Tumor invades less than one half of the myometrial thickness.</td>
</tr>
<tr>
<td>Stage IC</td>
<td>Tumor invades more than one half of the myometrial thickness.</td>
</tr>
<tr>
<td>Stage II</td>
<td>The tumor extends to the cervix.</td>
</tr>
<tr>
<td>Stage IIA</td>
<td>Cervical extension is limited to the endocervical glands.</td>
</tr>
<tr>
<td>Stage IIB</td>
<td>Tumor invades the cervical stroma.</td>
</tr>
<tr>
<td>Stage III</td>
<td>There is regional tumor spread.</td>
</tr>
<tr>
<td>Stage IIIA</td>
<td>Tumor invades the uterine serosa, adnexa, or positive peritoneal cytology.</td>
</tr>
<tr>
<td>Stage IIIB</td>
<td>Vaginal metastases are present.</td>
</tr>
<tr>
<td>Stage IIIC</td>
<td>Tumor has spread to pelvic or paraaortic lymph nodes.</td>
</tr>
<tr>
<td>Stage IV</td>
<td>There is bulky pelvic disease or distant spread.</td>
</tr>
<tr>
<td>Stage IVA</td>
<td>Tumor invades the mucosa of the bladder or rectosigmoid.</td>
</tr>
<tr>
<td>Stage IVB</td>
<td>Distant metastases are present [FIGO 1988].</td>
</tr>
</tbody>
</table>


Follow-Up:
- Pap smear every 3 months for two years, then every 6 months for 3 years.
- Chest x-ray once a year [Griffith's, 1999]

Literature review available.
ENDOMETRIAL HYPERPLASIA

All Applicants:
- Specialist Evaluation (Gynecologist) within the past 1 year to include the following:
  - Symptoms
  - Treatment
  - Current status
  - Recommendations for follow-up over the next 3 years.

If Applicable:
- Copy of ultrasound report with interpretation.
- Copy of biopsy report with interpretation.

CLEARANCE CRITERIA

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-4, AND</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>If treated with progestational agents, post treatment greater than 3 months.</td>
<td>RN</td>
<td>CLEAR</td>
</tr>
<tr>
<td>If treated with D&amp;C, post procedure greater than 3 months.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>If treated with hysterectomy, post surgery greater than 3 months.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Does not meet clearance criteria due to one or more of the following:**
- Hyperplasia not resolved.
- Current progesterone use, or post progesterone treatment less than 3 months.
- If treated with D&C, post procedure less than 3 months.
  - If treated with hysterectomy, post surgery less than 3 months.

Does not meet clearance criteria due to one or more of the following:
- Symptoms, i.e., irregular or dysfunctional uterine bleeding.

Does not meet clearance criteria due to one or more of the following:
- Adenomatous cells on endometrial biopsy.

Does not meet clearance criteria due to one or more of the following:
- History of uterine or endometrial cancer.

DIAGNOSTIC CODES

621.3  Endometrial Hyperplasia

Cross Reference  ICD.9.CM
Reviewers to Consider:
- None

COMMENTS:

Background: Endometrial hyperplasia is a pathologic condition that is usually associated with abnormal uterine bleeding. Hyperplasia of the endometrium results from estrogenic stimulation of the endometrium without the usual cyclic modification of progesterone and, therefore, is almost invariably found in anovulatory women. Women with endometrial hyperplasia have 3 treatment options: 1) hormone therapy with progesteronal agents; 2) dilation and curettage, or 3) hysterectomy. Patients treated with hormone therapy may be treated for up to one year and require frequent follow-up during treatment.

Literature review available.
ENDOMETRIOSIS
Includes Uterine Adenomyosis.

1 INFORMATION REQUIRED

All Applicants:
- Specialist Evaluation (Gynecologist) within the past 1 year to include the following:
  - Symptoms to include severity
  - Treatment
  - Current status
  - Recommendations for follow-up over the next 3 years.
- If laparoscopy, copy of most recent laparoscopy report with interpretation.

If Applicable:
- Discharge summary for all related hospitalizations.

2 CLEARANCE CRITERIA

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-6, AND</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment: None.</td>
<td>RN</td>
<td>CLEAR</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-6, AND</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment: Androgenic medication; post treatment greater than 3 months.</td>
<td>RN</td>
<td>CLEAR WITH RESTRICTION. PCMO FOLLOW-UP Periodic evaluation by a board certified gynecologist or an experienced provider.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does not meets clearance criteria due to one or more of the following:</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Treatment: Androgenic medication; current use, or post treatment less than 3 months.</td>
<td>RN</td>
<td>DEFER Delerral letter requires review by screening manager.</td>
</tr>
<tr>
<td>If laparotomy, post surgery less than 3 months.</td>
<td>MED ADVISOR</td>
<td>Risk varies - assess based on detailed history.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does not meets clearance criteria due to one or more of the following:</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>If laparoscopy, diagnosis of moderate to severe disease.</td>
<td>MED ADVISOR</td>
<td>Risk varies - assess based on detailed history.</td>
</tr>
<tr>
<td>Current, moderate to severe, symptoms.</td>
<td>MED ADVISOR</td>
<td>Risk varies - assess based on detailed history.</td>
</tr>
<tr>
<td>If symptoms, controlled with narcotics or pain medication other than over the counter medication, NSAIDs, or oral contraceptive pills.</td>
<td>MED ADVISOR</td>
<td>Risk varies - assess based on detailed history.</td>
</tr>
<tr>
<td>Current irregular or dysfunctional uterine bleeding.</td>
<td>MED ADVISOR</td>
<td>Risk varies - assess based on detailed history.</td>
</tr>
<tr>
<td>History of adenomyosis.</td>
<td>MED ADVISOR</td>
<td>Risk varies - assess based on detailed history.</td>
</tr>
</tbody>
</table>
ENDOMETRIOSIS

Reviewers to Consider:
- None

Background: Endometriosis is endometrial growth at distant sites outside of the uterus such as peritoneal surfaces of the bladder, cul-de-sac, pelvic side walls, broad ligaments, uterosacral ligaments, fallopian tubes, lymph nodes, ovaries, and bowel. More distant sites include the vagina, cervix, abdominal wall, arm, leg, pleura, lung, diaphragm, kidneys, spleen, and gallbladder.

Symptoms: Dyspareunia, dysmenorrhea, dyschezia, chronic pelvic pain, premenstrual spotting, spontaneous abortion, infertility, and luteinized unruptured follicle syndrome.

Treatment: May require 6 months of treatment with androgenic medications, with the usual side effect of hot flashes. Patients often need to wait 3 months after treatment to see if symptoms recur before being released by the GYN. Eighty percent of women using hormonal medications have side effects, i.e., weight gain, fluid retention, fatigue, hot flashes, and amenorrhea. Surgical vaporization of implants is the treatment of choice for severe pelvic pain or to preserve fertility.

Adenomyosis: Adenomyosis is characterized by the ectropic presence of endometrial tissue within the myometrium. It is sometimes referred to as endometriosis interna. The uterus is usually diffusely enlarged, but only rarely to a size greater than 12 weeks gestational size. When it becomes symptomatic, it is characterized by either menorrhagia or dysmenorrhea in the late reproductive years. Treatment is based on symptoms and age and includes antiinflammatory drugs, GnRH agonists, and hysterectomy.

Literature review available.
All Applicants:
- Specialist Evaluation (General Surgeon or Oncologist) within the past 1 year to include the following:
  - Documentation of recurrence within the past 5 years.
  - Recommendations for follow-up over the next 3 years.

If Applicable:
- Discharge summary for all related hospitalizations.

Background: Sarcomas may arise from the endometrium, myometrium, cervix, uterine blood vessels, or a leiomyoma. These diseases are most frequently seen in the fifth decade. The incidence of corpus sarcoma is much higher than that of sarcoma of the cervix. Because all elements of the uterus are mesodermal in origin and ectodermal rests may be present, mixed tumors may occur. A wide spectrum of histopathologic types can be found. Rapid uterine enlargement is a prominent sign of uterine sarcoma, and abnormal bleeding may or may not be present. Pain, anemia, and weight loss are late symptoms. Pulmonary metastases frequently occur early. Surgical excision of the uterus, tubes, and ovaries is the recommended treatment for sarcoma of the uterus.

Sarcomas Types:
- Mixed mullerian sarcoma - heterologous element not native to the mullerian systems, such as cartilage or bone; homologous elements native to the mullerian system.
UTERINE CANCER

- Endometrial stromal sarcoma develops from the stromal component of the endometrium.
- Leiomyosarcoma develops in the myometrium or in a myoma (fibroid).

Surgical Staging of Uterine Fundal Tumors:

<table>
<thead>
<tr>
<th>Stage</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage I</td>
<td>The tumor is confined to the uterine fundus.</td>
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<tr>
<td>Stage IA</td>
<td>Tumor is limited to the endometrium.</td>
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<td>Stage IB</td>
<td>Tumor invades less than one half of the myometrial thickness.</td>
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<tr>
<td>Stage IC</td>
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<td>Stage III</td>
<td>There is regional tumor spread.</td>
</tr>
<tr>
<td>Stage IIIA</td>
<td>Tumor invades the uterine serosa, adnexa, or positive peritoneal cytology.</td>
</tr>
<tr>
<td>Stage IIIB</td>
<td>Vaginal metastases are present.</td>
</tr>
<tr>
<td>Stage IIC</td>
<td>Tumor has spread to pelvic or paraaortic lymph nodes.</td>
</tr>
</tbody>
</table>

| Stage IV | There is bulky pelvic disease or distant spread.                             |
| Stage IVA| Tumor invades the mucosa of the bladder or rectosigmoid.                    |
| Stage IVB| Distant metastases are present [FIGO 1988].                                 |

Prognosis for Uterine Sarcomas: Stage is the most significant predictor of outcome for women with uterine sarcomas. Patients whose tumors are confined to the uterus have a survival rate of 60% to 70% following surgical resection. Major sites of failure include the pelvis, upper abdomen, and lung. Few well-conducted prospective adjuvant therapy trials have been accomplished, so a precise role for either adjuvant irradiation or chemotherapy remains undefined.

As has been noted for endometrial carcinoma, adjuvant pelvic irradiation may reduce the rate of pelvic failure without improving survival if more patients succumb to distant failure. Pelvic irradiation and local tumor control may be an important issue in tumors with extension to the cervix. However, so few patients are placed in this category that meaningful treatment data are not available. Very few patients with tumor spread outside of the uterus can be curatively treated. Some women with small-volume regional disease have obtained long-term survival following external-beam irradiation. However, most patients with advanced or recurrent disease ultimately experience disease progression and die. [DeVita: Cancer: Principles and Practice of Oncology, 5th ed., Copyright © 1997 Lippincott-Raven Publishers]

Expected Course and Prognosis [Griffith's, 1998]:

<table>
<thead>
<tr>
<th>Grade</th>
<th>5 Year Survival (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IAG1</td>
<td>98</td>
</tr>
<tr>
<td>IAG2</td>
<td>85</td>
</tr>
<tr>
<td>ICG3</td>
<td>60</td>
</tr>
<tr>
<td>IIA/B</td>
<td>60</td>
</tr>
<tr>
<td>IIIB</td>
<td>40</td>
</tr>
<tr>
<td>IV</td>
<td>15</td>
</tr>
</tbody>
</table>

Follow-Up:
- Pap smear every 3 months for two years, then every 6 months for 3 years
- Chest x-ray once a year [Griffith's, 1999]
All Applicants:
- Report of Medical Examination within the past 1 year to include the following:
  - Symptoms
  - Clinical evidence of GI, GU, or GYN obstruction.
  - Current status, to include comment about stability of fibroids over the past 6 months.
  - Treatment history, to include surgery.
  - Assessment of need for medical intervention over the next 3 years.
  - Recommendations for follow-up over the next 3 years.
- Copy of most recent ultrasound report with interpretation to include size, location, and number of fibroids.

If Applicable:
- Discharge summary for all related hospitalizations.

**CLEARANCE CRITERIA**

<table>
<thead>
<tr>
<th>Meets clearance criteria 1 - 5, AND</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Size stable for at least the past 6 months; determined by examining health care provider; may be determined by pelvic exam or ultrasound.</td>
<td>RN</td>
<td>CLEAR</td>
</tr>
<tr>
<td>- No irregular or dysfunctional uterine bleeding.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>- No, or mild, symptoms.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>- If mild symptoms, well controlled non-narcotic medication, e.g., with NSAIDs or birth control pills.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>- No clinical evidence of GI, GU, or GYN obstruction.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Does not meet clearance criteria due to one or more of the following:**

- Size not stable for at least the past 6 months.
- If leiomyomectomy or hysterectomy, post surgery less than 3 months.

**Does not meet clearance criteria due to one or more of the following:**

- Irregular or dysfunctional uterine bleeding.
- Moderate to severe symptoms.
- Symptoms controlled with narcotics or pain medication other than NSAIDs or birth control pills.
- Clinical evidence of GI, GU, or GYN obstruction.

**DIAGNOSTIC CODES**

218.0 Uterine Fibroids (Leiomyomas)
Cross Reference ICD.9.CM

**NOTES AND INSTRUCTIONS FOR REVIEWERS**

Reviewers to Consider:
- None
UTERINE LEIOMYOMAS (FIBROIDS)  

**Background:** Uterine leiomyomas are well circumscribed, pseudo-encapsulated benign tumors of the uterus. The majority are asymptomatic and are only suspected from pelvic exam. Approximately 20-40% of all women have uterine fibroids. Fibroids frequently regress after menopause. Approximately 10% recur following myomectomy. (Griffith's, 1998)

**Symptoms:** The majority are asymptomatic and are only suspected from pelvic examination. The most common symptom is abnormal uterine bleeding. Hypermenorrhea most common. Secondary anemia with associated symptomatology may result. Pressure on the bladder may result in suprapubic discomfort, urinary frequency. Pressure on the rectosigmoid may result in low back pain. Edema and varicosities of the lower extremities may result from large tumors. Pain may result from twisted, pedunculated myomas or degenerating, hemorrhagic or infected myomas. Rapid growth particularly in perimenopausal or postmenopausal may indicate sarcoma (Griffiths, 2003).

**Treatment:** Treatment must be individualized. Patients with minimal symptoms may be managed with iron preparations and analgesics.

**Conservative management:** Asymptomatic myomas of less than 14 week’s size gestation should be closely observed with pelvic examinations and ultrasonography at 3-6 month intervals, as long as size is stable. Usually regress after menopause.

**Nonsurgical therapies:**
- Lutenizing hormone releasing hormone (LHRH) agonists induce an abrupt artificial menopause with cessation of bleeding and shrinkage of myomas. Not recommended for more than 6 months. May be useful in perimenopausal patients or as an adjunct to surgery.
- Myolysis by needle cautery or cryotherapy. Long term outcome is unknown.
- Uterine artery embolization average 50% shrinkage; painful.

**Surgical Measures:**
- Surgical management is indicated in the following situations:
  - Excessive uterine size (> 14 weeks gestation) or excessive rate of growth.
  - Submucous location if associated with hypermenorrhea.
  - Pedunculated myomas may undergo torsion, pain, necrosis and hemorrhage.
  - Symptomatic from pressure on bladder or rectum.
  - If differentiation from ovarian mass is not possible.
  - If there is associated pelvic disease, i.e., endometriosis, pelvic inflammatory disease, etc.

(Griffith's, 2003)

**Follow-Up:**
- Newly diagnosed uterine myoma, if symptomatic or excessive size, 2-3 months with pelvic exam and ultrasound.
- Consider CA-125 antigen.
- Monitor hemoglobin and hematocrit, if bleeding excessive.
- If uterine size and symptoms are stable, monitor every 6 months. (Griffith's, 2003)

Literature review available.
BARTHOLIN GLAND INFECTIONS
Includes Infection, Abscesses, and Cysts

INFORMATION REQUIRED
If history within the past 5 years.

All Applicants:
- Report of Medical Examination to include the following:
  - Number of episodes
  - Symptoms
  - Treatment
  - Recommendations for follow-up over the next 3 years.

CLEARANCE CRITERIA

1. Infection, abscess, or cyst resolved.
2. No, or resolved, symptoms.
3. If treated, treatment complete.
4. If history of incision and drainage or marsupialization, post procedure greater than 6 weeks.

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-4, AND</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Single episode.</td>
<td>RN</td>
<td>CLEAR</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-4, AND</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Multiple episodes, i.e., recurrent.</td>
</tr>
<tr>
<td>• Definitely treated with marsupialization.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-4, AND</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Multiple episodes.</td>
</tr>
<tr>
<td>• No history of marsupialization or failed marsupialization.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Infection, abscess, or cyst not resolved.</td>
</tr>
<tr>
<td>• If treated, treatment not complete.</td>
</tr>
<tr>
<td>• If history of incision and drainage or marsupialization, post procedure less than 6 weeks.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>RN</td>
<td>DEFER</td>
</tr>
<tr>
<td></td>
<td>Deferal letter requires review by screening manager.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Current symptoms.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>MED ADVISOR</td>
<td></td>
</tr>
<tr>
<td>Risk varies - assess based on detailed history.</td>
<td></td>
</tr>
</tbody>
</table>

DIAGNOSTIC CODES

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>616.3</td>
<td>Bartholin Gland Abscess</td>
</tr>
<tr>
<td>616.2</td>
<td>Bartholin Gland Cyst</td>
</tr>
</tbody>
</table>

Cross Reference: ICD 9 CM

NOTES AND INSTRUCTIONS FOR REVIEWERS:

Reviewers to Consider:
- None
Background: Many women have a single infection that is easily treated and requires no special follow-up. Recurrent abscesses, however, may require treatment by marsupialization.

Marsupialization: Process of raising the borders of an evacuated tumor or abscess sac to the edges of the wound and stitching them there to form a pouch. The interior sac suppurates and gradually closes by granulation.

Literature review available.
All Applicants:
- Specialist Evaluation (General Surgeon or Oncologist) within the past 1 year to include the following:
  - Documentation of recurrences within the past 5 years.
  - Recommendations for follow-up over the next 3 years.

If Applicable:
- Discharge summary for all related hospitalizations.

<table>
<thead>
<tr>
<th>CLEARANCE CRITERIA</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Treatment complete.</td>
<td></td>
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<tr>
<td>2. No history of recurrences for at least the past 5 years.</td>
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</table>

Meets clearance criteria 1.2, AND"
International Federation of Gynecology and Obstetrics—Classification of Vaginal Cancer:

Stage 0  Carcinoma in situ
Stage I  Carcinoma limited to the vaginal wall
Stage II Carcinoma involving the subvaginal tissues but not extending to the pelvic wall
Stage III Carcinoma extending to the pelvic wall
Stage IV Carcinoma extending beyond the true pelvis or involving the mucosa of the bladder or rectum.
Stage IVA Involvement of adjacent organs (bladder, rectum)
Stage IVB Involvement of distant organs

Prognosis: "The prognosis for patients with vaginal squamous cell carcinoma depends primarily on the extent of disease at the time of diagnosis. Stage I disease treated with radiation therapy results in 5-year survival rates of 80% to 90%. Five-year survival rates for higher stages are 45% to 58% for stage II, 25% to 40% for stage III, and up to 10% for stage IV. The overall 5-year survival rate is approximately 45." [Ryan: Kistner's Gynecology: Principles & Practice, 6th ed., Copyright © 1995 Mosby-Year Book, Inc.]

"The rates of local control, distant metastasis, and survival are all correlated strongly with tumor stage. Tumor size also appears to be an important predictor of outcome. Chyle and colleagues reported a higher rate of local and distant failure for tumors larger than 5 cm in diameter; Kirkbride and colleagues reported a significantly better survival rate for patients with tumors smaller than 4 cm in diameter; and Stock and colleagues reported better survival when disease was limited to one third of the vaginal canal. Most investigators have been unable to find a correlation between tumor site and outcome. However, Chyle and colleagues reported higher rates of local recurrence and overall relapse in patients with posterior wall lesions, and Kucera and Vavra reported a significantly better survival rate for patients whose tumors involved the upper one third of the vagina. Tumors that involve the entire vagina tend to have a poorer prognosis, probably reflecting the larger size of these lesions.

Investigators disagree about the influence of histologic grade and type on outcome. Several investigators have reported a correlation between increasing grade of squamous carcinomas and recurrence, whereas others have found no correlation. Chyle and colleagues reported significantly poorer survival and local control rates for patients with adenocarcinoma, but other investigators found no difference in outcome for patients with squamous carcinomas or adenocarcinomas." [DeVita: Cancer: Principles and Practice of Oncology, 5th ed., Copyright © 1997 Lippincott-Raven Publishers]

Recurrence: Recurrences often occur locally and are more common in higher stages. Distant metastases occur later and usually involve the lung and bone. Radical surgery may be attempted in selected cases of isolated, local recurrence.

"Chemotherapy is being evaluated for a possible role in cases with systemic recurrence. [Ryan: Kistner's Gynecology: Principles & Practice, 6th ed., Copyright © 1995 Mosby-Year Book, Inc.]

Literature review available.
**INFORMATION REQUIRED**

All Applicants:
- Report of Medical Examination to include the following:
  - Number of episodes within the past 2 years.
  - Etiology
  - Symptoms
  - Treatment
  - Recommendations for follow-up over the next 3 years.
- Copy of related diagnostic tests, i.e., tests to rule out sexually transmitted diseases.

**CLEARANCE CRITERIA**

<table>
<thead>
<tr>
<th>Meets clearance criteria 1-2, AND</th>
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<tr>
<td>- Resolved for at least the past 3 months.</td>
<td>RN</td>
<td>DEFER</td>
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</table>

Does not meet clearance criteria due to one or more of the following:
- Not resolved for at least the past 3 months.
CONTRACEPTION

Includes Oral Contraception, Barrier Contraception, Intrauterine Devices (IUD), and Long-Acting Progestins.

### INFORMATION REQUIRED

**All Applicants:**
- Report of Medical Examination to include the following:
  - Current menstrual pattern
  - Complications

**Applicants Using Norplant:**
- Insertion date
- If Norplant inserted greater than 2 years ago and applicant wishes to keep Norplant in place during PC service: Applicant Personal Statement verifying plan to use barrier method of contraception at 5 years post the insertion date.

**Applicants Using an IUD:**
- Insertion date
- Type of IUD
- Need for removal or replacement over the next 3 years.

**Applicants Using Depo Provera:**
- Date injections started
- Date of last injection

### CLEARANCE CRITERIA

<table>
<thead>
<tr>
<th>Meets clearance criteria 1 - 3, AND</th>
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<tbody>
<tr>
<td>• Contraceptive method - <strong>Oral Contraceptives</strong>, includes 91-day regimen (Seasonale).</td>
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<th>Meets clearance criteria 1 - 3, AND</th>
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<tr>
<td>• Contraceptive method – <strong>Contraceptive Patch</strong></td>
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<tr>
<th>Meets clearance criteria 1 - 3, AND</th>
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<tr>
<td>• Contraceptive method - <strong>Barrier, e.g., Diaphragm, Cervical Cap with Spermicide</strong> (see Comments).</td>
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<th>Meets clearance criteria 1 - 3, AND</th>
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<tr>
<td>• Contraceptive method - <strong>NuvaRing</strong></td>
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<th>Meets clearance criteria 1 - 3, AND</th>
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<tr>
<td>• Contraceptive method - <strong>IUD</strong></td>
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<th>REVIEWER</th>
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<td>RN</td>
<td>CLEAR WITH RESTRICTION</td>
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PCMO FOLLOW-UP
Periodic evaluation, to include pelvic exam, by a qualified gynecologist or an experienced provider.

(continued on next page)
CONTRACEPTION GYN 8.1

### Male Condom

**FDA Approval Date:** Latex: use started before premarket approval was required. Polyurethane: cleared in 1989; available starting 1995.

**Description:** A sheath placed over the erect penis blocking the passage of sperm.

**Failure Rate:** 11 (a,b)

**Some Risks:** Irritation and allergic reactions (less likely with polyurethane).

**Protection from Sexually Transmitted Diseases (STDs):** Except for abstinence, latex condoms are the best protection against STDs, including gonorrhea and AIDs.

**Convenience:** Applied immediately before intercourse; used only once and discarded. Polyurethane condoms are available for those with latex sensitivity.

**Availability:** Nonprescription

*OMS Policy: OMS recommends the use of condoms that do not contain the spermicide, nonoxynol-9 (N-9). N-9 may increase the transmission of HIV (see OMS policy memo dated 8/25/2000).*

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<table>
<thead>
<tr>
<th>DIAGNOSTIC CODES</th>
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<tbody>
<tr>
<td>V25 Contraception</td>
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<tr>
<td>Cross Reference ICD.9.CM</td>
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<tr>
<th>NOTES AND INSTRUCTIONS FOR REVIEWERS</th>
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<tr>
<td>Reviewers to Consider:</td>
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<tr>
<td>None</td>
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<th>COMMENTS</th>
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<tbody>
<tr>
<td><strong>CONTRACEPTIVE METHODS</strong> (Source: Adapted from the Food and Drug Administration 12/03)</td>
</tr>
<tr>
<td>Male Condom</td>
</tr>
</tbody>
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### Meets clearance criteria 1 - 3, AND

- Contraceptive method - Depo Provera
- Status post first Depo Provera injection greater than 3 months.
- Documentation of most recent Depo Provera injection.

<table>
<thead>
<tr>
<th>PCMO FOLLOW-UP</th>
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<tbody>
<tr>
<td>Norplant should be removed at 5 years post the insertion date.</td>
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<tr>
<td>If removal is not possible in country, a barrier method of contraception must be used until Norplant can be safely removed.</td>
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<tr>
<th>Does not meet clearance criteria due to one or more of the following:</th>
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<tr>
<td>- Norplant inserted greater than 2 years ago and applicant wishes to keep implants in place during PC service; No Applicant Personal Statement verifying plans to use barrier method of contraception at 5 years post the insertion date (see Comments).</td>
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<th>Does not meet clearance criteria due to one or more of the following:</th>
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<tr>
<td>- Significant complications.</td>
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<td>- Unstable amenorrhea, i.e., moderate or severe break through bleeding.</td>
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<tr>
<th>MED ADVISOR</th>
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<tr>
<td>Risk varies - assess based on detailed history.</td>
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Effective 2/12/2004
Female Condom
FDA Approval Date: 1993
Description: A lubricated polyurethane sheath shaped similarly to the male condom. The closed end has a flexible ring that is inserted into the vagina.
Failure Rate: (number of pregnancies expected per 100 women per year): 21
Some Risks: Irritation and allergic reactions
Protection from Sexually Transmitted Diseases (STDs): May give some STD protection; not as effective as latex condom
Convenience: Applied immediately before intercourse; used only once and discarded.
Availability: Nonprescription

Diaphragm with Spermicide
FDA Approval Date: Use started before premarket approval was required.
Description: A dome-shaped rubber disk with a flexible rim that covers the cervix so that sperm cannot reach the uterus. A spermicide is applied to the diaphragm before insertion.
Failure Rate: (number of pregnancies expected per 100 women per year): 17
Some Risks: Irritation and allergic reactions, urinary tract infection. (c) Risk of toxic shock syndrome, a rare but serious infection, when kept in place longer than recommended.
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: Inserted before intercourse and left in place at least six hours after; can be left in place for 24 hours, with additional spermicide for repeated intercourse.
Availability: Prescription

Lea’s Shield
FDA Approval Date: 2002
Description: A dome-shaped rubber disk with a valve and a loop that is held in place by the vaginal wall. Covers the upper vagina and cervix so that sperm cannot reach the uterus. Spermicide is applied before insertion.
Failure Rate: (number of pregnancies expected per 100 women per year): 15
Some Risks: Skin irritation, spotting, discomfort (female and male partners), urinary tract infection. Theoretical risk of toxic shock syndrome.
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: Inserted before intercourse and left in place at least 8 hours after; can be left in place for up to 48 hours, with additional spermicide for repeated intercourse.
Availability: Prescription

Cervical Cap with Spermicide
FDA Approval Date: Prentiff Cap—1988; FemCap—2003
Description: A soft rubber cup with a round rim, which fits snugly around the cervix.
Failure Rate: (number of pregnancies expected per 100 women per year): Prentiff Cap—17; FemCap—23
Some Risks: Irritation and allergic reactions, abnormal Pap test. (c) Risk of toxic shock syndrome, a rare but serious infection, when kept in place longer than recommended.
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: May be difficult to insert; can remain in place for 48 hours without reapplying spermicide for repeated intercourse.
Availability: Prescription

*SOM Policy: OMS does not supply the cervical cap due to difficulties with proper fitting and use.

Sponge with Spermicide
FDA Approval Date: 1983 (Not currently marketed)
Description: A disk-shaped polyurethane device containing a spermicide.
Failure Rate: (number of pregnancies expected per 100 women per year): 14-28
Some Risks: Irritation and allergic reactions, difficulty in removal. (c) Risk of toxic shock syndrome, a rare but serious infection, when kept in place longer than recommended.
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: Inserted before intercourse and protects for repeated acts of intercourse for 24 hours without additional spermicide; must be left in place for at least six hours after intercourse; must be removed within 30 hours of insertion. Is discarded after use.
Availability: Nonprescription; not currently marketed

*SOM Policy: OMS does not supply, nor recommend the use of, the cervical sponge.
Spermicide Alone

**FDA Approval Date:** Use started before premarket approval was required. Since November 2002, only one active ingredient has been allowed.

**Description:** A foam, cream, jelly, film, suppository, or tablet that contains nonoxynol-9, a sperm-killing chemical

**Failure Rate** (number of pregnancies expected per 100 women per year): 20-50 (studies have shown varying effectiveness rates)

**Some Risks:** Irritation and allergic reactions, urinary tract infections (g)

**Protection from Sexually Transmitted Diseases (STDs):** None

**Convenience:** Instructions vary; check labeling. Inserted between 5 and 90 minutes before intercourse and usually left in place at least six to eight hours after.

**Availability:** Nonprescription

---

Oral Contraceptives – Combines Pill

**FDA Approval Date:** First in 1960; most recent in 2003

**Description:** A pill that suppresses ovulation by the combined actions of the hormones estrogen and progestin. A chewable form was approved in 2003.

**Failure Rate** (number of pregnancies expected per 100 women per year): 1-2

**Some Risks:** Dizziness; nausea; changes in menstruation, mood, and weight; rarely, cardiovascular disease, including high blood pressure, blood clots, heart attack, and strokes

**Protection from Sexually Transmitted Diseases (STDs):** None

**Convenience:** Must be taken on daily schedule, regardless of frequency of intercourse. Women using the chewable tablet must drink 8 oz. of liquid immediately after taking.

**Availability:** Prescription

---

Oral Contraceptives – Progestin-Only Minipill

**FDA Approval Date:** 1973

**Description:** A pill containing only the hormone progestin that reduces and thickens cervical mucus to prevent the sperm from reaching the egg.

**Failure Rate** (number of pregnancies expected per 100 women per year): 2

**Some Risks:** Irregular bleeding, weight gain, breast tenderness, less protection against ectopic pregnancy

**Protection from Sexually Transmitted Diseases (STDs):** None

**Convenience:** Must be taken on daily schedule, regardless of frequency of intercourse.

**Availability:** Prescription

---

Oral Contraceptives – 91-Day Regimen (Seasonale)

**FDA Approval Date:** 2003

**Description:** A pill containing estrogen and progestin, taken in 3-month cycles of 12 weeks of active pills followed by one week of inactive pills. Menstrual periods occur during the 13th week of the cycle.

**Failure Rate** (number of pregnancies expected per 100 women per year): 1-2

**Some Risks:** Similar to oral contraceptives—combined pill

**Protection from Sexually Transmitted Diseases (STDs):** None

**Convenience:** Must be taken on daily schedule, regardless of frequency of intercourse. Since users will have fewer periods, they should consider the possibility that they might be pregnant if they miss scheduled periods. May have more unplanned bleeding and spotting between periods than with 28-day oral contraceptives.

**Availability:** Prescription

---

**Patch (Ortho Evra)**

**FDA Approval Date:** 2001

**Description:** Skin patch worn on the lower abdomen, buttocks, or upper body that releases the hormones progestin and estrogen into the bloodstream.

**Failure Rate** (number of pregnancies expected per 100 women per year): 1-2 (Appears to be less effective in women weighing more than 198 pounds.)

**Some Risks:** Similar to oral contraceptives—combined pill

**Protection from Sexually Transmitted Diseases (STDs):** None

**Convenience:** New patch is applied once a week for three weeks. Patch is not worn during the fourth week, and woman has a menstrual period.

**Availability:** Prescription
Vaginal Contraceptive Ring (Nuva Ring®)

FDA Approval Date: 2001

Description: A flexible ring about 2 inches in diameter that is inserted into the vagina and releases the hormones progestin and estrogen.

Failure Rate (number of pregnancies expected per 100 women per year): 1-2

Some Risks: Vaginal discharge, vaginitis, irritation. Similar to oral contraceptives--combined pill

Protection from Sexually Transmitted Diseases (STDs): None

Convenience: Inserted by the woman; remains in the vagina for 3 weeks, then is removed for 1 week. If ring is expelled and remains out for more than 3 hours, another birth control method must be used until ring has been reinserted continuously for 7 days.

Availability: Prescription

Storage: NuvaRing has a shelf life of two years at 2-8 °C in the Pharmacy, followed by a shelf life of 4 months stored below 86°F (30°C) at the users' site. NuvaRing must be stored at room temperature 77°F (25°C). Temperatures can range from 59-86°F (15-30°C). Direct sunlight and storing above 86°F (30°C) should be avoided.

OMS Policy: OMS does not supply Nuva Ring® due to overseas procurement and storage difficulties.

Post-Coital Contraceptives (Preven and Plan B)

FDA Approval Dates: 1998-1999

Description: Pills containing either progestin alone or progestin plus estrogen

Failure Rate (number of pregnancies expected per 100 women per year): Almost 80 percent reduction in risk of pregnancy for a single act of unprotected sex

Some Risks: Nausea, vomiting, abdominal pain, fatigue, headache

Protection from Sexually Transmitted Diseases (STDs): None

Convenience: Must be taken within 72 hours of having unprotected intercourse.

Availability: Prescription

Injection (Depo-Provera)

FDA Approval Date: 1992

Description: An injectable progestin that inhibits ovulation, prevents sperm from reaching the egg, and prevents the fertilized egg from implanting in the uterus.

Failure Rate (number of pregnancies expected per 100 women per year): less than 1

Some Risks (serious medical risks from contraceptives are rare): Irregular bleeding, weight gain, breast tenderness, headaches.

Protection from Sexually Transmitted Diseases (STDs): None

Convenience: One injection every three months.

Availability: Prescription

Additional Information: The most common side effects of Depo Provera include the following: menstrual irregularities (various altered bleeding patterns). As women continue using Depo Provera, fewer experience irregular bleeding and more experience amenorrhea. By one year approximately 55% experience complete amenorrhea. Unusually heavy or continuous bleeding, however, is not a usual side effect of Depo Provera. The recommended dose is 150 mg DMPA every 3 months (13 weeks) by deep, intramuscular injection in the gluteal or deltoid muscle. If Depo Provera is administered under the prescribed schedule, i.e., within 5-7 days of a normal period, no further contraceptive protection is needed.
CONTRACEPTION

Some Risks: Irregular bleeding, weight gain, breast tenderness, headaches, difficulty in removal
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: Implanted and removed by health-care provider in minor outpatient surgical procedure; effective for up to five years.
Availability: Prescription. In July 2002, Norplant's manufacturer announced that it will no longer distribute the Norplant system. Women using the system should contact their doctors about what their contraceptive options will be after the five-year expiration date of their Norplant systems.

Additional Information: Causes complication in approximately 50% of users. These complications, usually amenorrhea or frequent bleeding, require evaluation by a GYN. The findings are almost always negative. A woman on Norplant with amenorrhea needs access to reliable pregnancy testing. A woman on Norplant with frequent bleeding needs a gynecologic evaluation to rule out abnormalities that are causing the bleeding. If the exams are negative, she requires no special follow-up. Norplant must be surgically removed or replaced every 5 years.

Removal of Norplant After 5 Years: The manufacturers "strongly recommend the removal of Norplant System after five years. As the Norplant System is a progestin-only contraceptive method, it is presumed that if a pregnancy occurs it may be more likely to be an ectopic pregnancy. In addition, since the extended use of the Norplant System has not been studied, we cannot recommend use beyond five years in patients who are using another method of contraception." [Beamish, 1997]

IUD (Intrauterine Device)
FDA Approval Date: 1976
Description: A T-shaped device inserted into the uterus by a health professional.
Failure Rate (number of pregnancies expected per 100 women per year): less than 1
Some Risks: Cramps, bleeding, pelvic inflammatory disease, infertility, perforation of uterus
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: After insertion by physician, can remain in place for up to one or 10 years, depending on type.
Availability: Prescription.
Additional Information: Associated with an increased risk of pelvic infection, which may be higher in less hygienic situations abroad.

Periodic Abstinence
FDA Approval Date: N/A
Description: To deliberately refrain from having sexual intercourse during times when pregnancy is more likely.
Failure Rate (number of pregnancies expected per 100 women per year): 20
Some Risks: None
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: Requires frequent monitoring of body functions (for example, body temperature for one method).
Availability: Instructions from health-care provider

Trans-Abdominal Surgical Sterilization - Female (Falope Ring, Hulka Clip, Filshie Clip)
FDA Approval Date: Early 1970s
Description: The woman's fallopian tubes are blocked so the egg and sperm can't meet in the fallopian tube, preventing conception.
Failure Rate (number of pregnancies expected per 100 women per year): less than 1
Some Risks: Pain, bleeding, infection, other post-surgical complications, ectopic (tubal) pregnancy.
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: One-time surgical procedure that requires an abdominal incision.
Availability: Surgery

Sterilization Implant - Female (Essure System)
FDA Approval Date: 2002
Description: Small metallic implant that is placed into the fallopian tubes. The device works by causing scar tissue to form, blocking the fallopian tubes and preventing conception.
Failure Rate (number of pregnancies expected per 100 women per year): less than 1
Some Risks: Mild to moderate pain after insertion, ectopic (tubal) pregnancy.
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: Minor surgical procedure, permanent sterilization. Device is inserted through the vagina using a catheter. Women must rely on another birth control method during the first three months, until placement is confirmed with an X-ray procedure.
Availability: Prescription

Effective 2/12/2004
CONTRACEPTION

Surgical Sterilization - Male
FDA Approval Date: N/A
Description: Sealing, tying, or cutting a man's vas deferens so that the sperm can't travel from the testicles to the penis.

(b) Failure Rate (number of pregnancies expected per 100 women per year): less than 1
Some Risks (serious medical risks from contraceptives are rare): Pain, bleeding, infection, other minor postsurgical complications
Protection from Sexually Transmitted Diseases (STDs): None
Convenience: One-time surgical procedure.
Availability: Surgery

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Effective 2/1/2004

Page 7 of 7
TERMINATION OF PREGNANCY
Includes Therapeutic and Elective Abortion.

INFORMATION REQUIRED
- If history within the past 1 year.

All Applicants:
- Report of Medical Examination to include the following:
  - Report of negative pregnancy test if procedure within the past 2 months.

CLEARANCE CRITERIA

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1. No, or resolved, medical complications.
2. No, or resolved, emotional complications.
3. Post abortion greater than 2 months.

Meets clearance criteria 1-3, AND
- Negative pregnancy test if procedure within the past 2 months.

Does not meet clearance criteria due to one or more of the following:
- Post abortion less than 2 months.

Reviewers to Consider:
- Medevac Policy: Following a Medevac for termination of pregnancy, Volunteer may be cleared to return to service as early as 3 weeks following the procedure. Clearance is dependent upon a thorough medical evaluation and counseling clearance.

Background: Less than 2% of individuals have medical complications following termination of pregnancy.

Literature review available.

Effective 1/15/99
**Mammography**

**Female Applicants Age 50 or Over:**
- Mammography report with interpretation *within* the past year to include the following:
  - Recommendations for follow-up over the next three years.

**If Applicable:**
- Comparison films

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<td>1. Mammography report includes one of the following findings:</td>
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**Meets clearance criteria, AND**
- **Negative**
  - Radiologist or examining health care provider recommend routine follow-up, **OR** no recommendations specified.

- **RN**
  - **CLEAR WITH RESTRICTION**
    - Mamrogram Accommodation
    - PCMO FOLLOW-UP
      - Annual mammogram.

**Meets clearance criteria, AND**
- **Benign Finding (Includes Densities, Irregularities, and Nodularities).**
  - Radiologist or examining health care provider recommend routine follow-up.

- **RN**
  - **CLEAR WITH RESTRICTION**
    - Mamrogram Accommodation
    - PCMO FOLLOW-UP
      - Annual mammogram.

**Meets clearance criteria, AND**
- **Probably Benign Finding (Includes Densities, Irregularities, and Nodularities).**
  - Radiologist or examining health care provider recommend a short interval follow-up to confirm no change in an abnormal finding.

- **RN**
  - **DEFER**
    - Defer until resolved or stable as per radiologist or examining health care provider.

**Meets clearance criteria, AND**
- **Needs Additional Evaluation**
  - Radiologist or examining health care provider recommend additional views, magnification, ultrasound, or advanced mammographic studies.

- **RN**
  - **DEFER**
    - Defer until resolved as per radiologist or examining health care provider.

**Meets clearance criteria, AND**
- **Suspicious Finding**
  - Radiologist or health care provider recommend biopsy.

- **RN**
  - **DEFER**
    - Defer until resolved as per radiologist or examining health care provider.

**Meets clearance criteria, AND**
- **Highly Suspicious Finding**
  - Radiologist or examining health care provider recommend biopsy.

- **RN**
  - **DEFER**
    - Defer until resolved as per radiologist or examining health care provider.

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Effective 1/15/99
MAMMOGRAPHY

DIAGNOSTIC CODES

87.37  Mammography

Cross Reference  ICD.9.CM

NOTES AND INSTRUCTIONS FOR REVIEWERS:

Reviewers to Consider:
- Applicants cleared to a mammography country should bring, to their country of assignment, their most recent mammogram films for comparison.

COMMENTS:

Background: In addition to regular breast examination, the Public Health Service (and other groups) recommend that women over the age of 50 receive mammography every one to two years. Mammography is expected to detect 2-4 cancers per 1000 women who are regularly screened. The detection rate is expected to be higher for women having their first mammogram.

Literature review available.
### INFORMATION REQUIRED

**All Applicants:**

- Report of Medical Examination to include the following:
  - Date of surgery
  - Type of surgical procedure
  - Reason for surgery
  - Etiology, if known.
  - Post surgical complications, if any.
  - Treatment, if any, e.g., hormone replacement.
  - Recommendations for follow-up over the next 3 years.
- Copy of pathology report if underlying malignant etiology.

If Surgery Within the Past 1 Year:

- Discharge summaries for all related hospitalizations.
- Documentation of release from surgical care.

### CLEARANCE CRITERIA

<table>
<thead>
<tr>
<th></th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>History of gynecologic surgery and surgical procedure within the past 1 year.</td>
<td>RN</td>
</tr>
<tr>
<td>2</td>
<td>No post surgical complications.</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Benign etiology, i.e., no history of malignancy.</td>
<td></td>
</tr>
</tbody>
</table>

Meets clearance criteria 1 - 3, AND

- Laparoscopy (Diagnostic)
- Clinical post surgery greater than 6 weeks.
<table>
<thead>
<tr>
<th>Procedures</th>
<th>MED ADVISOR</th>
<th>RN</th>
<th>DEFER</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Meets clearance criteria 1 - 3, AND</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>- Salpingectomy</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>- Status post surgery greater than 3 months.</td>
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<tr>
<td><strong>Meets clearance criteria 1 - 3, AND</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>- Ovarian Cystectomy (via Laparoscopy or Laparotomy)</td>
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<td></td>
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<tr>
<td>- Status post surgery greater than 3 months.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Meets clearance criteria 1 - 3, AND</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>- Oophorectomy (via laparoscopy or laparotomy)</td>
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<td></td>
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<tr>
<td>- Status post surgery greater than 3 months.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td><strong>Does not meet clearance criteria due to one or more of the following:</strong></td>
<td></td>
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<tr>
<td>- Status post surgery less than the time specified for each procedure.</td>
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<td></td>
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</tr>
<tr>
<td><strong>Does not meet clearance criteria due to one or more of the following:</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>- Post surgical complications.</td>
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<td></td>
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<tr>
<td><strong>Does not meet clearance criteria due to one or more of the following:</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>- Malignant etiology.</td>
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</tr>
</tbody>
</table>

**DIAGNOSTIC CODES**

- 54.21 Laparoscopy
- 54.19 Laparotomy
- 66.39 Tubal Ligation
- 68.4 Hysterectomy
- 66.6 Salpingectomy
- 65.5 Oophorectomy, Bilateral
- 65.3 Oophorectomy, Unilateral

Cross Reference ICD.9.CM

**NOTES AND INSTRUCTIONS FOR REVIEWERS**

Reviewers to Consider:
- Shorter deferral period with adequate documentation and review by Medical Advisor.

**COMMENTS**

Literature review and abstract available.

Effective 11/15/2002
OTHER GYNECOLOGY DISORDERS
Includes Ectopic Pregnancy, Pelvic Mass, and Uterine Prolapse.

<table>
<thead>
<tr>
<th>INFORMATION REQUIRED</th>
<th>Any history:</th>
</tr>
</thead>
</table>

All Applicants:
- Report of Medical Examination
If Medical Advisor Requests:
- Specialist Evaluation (Gynecologist)

<table>
<thead>
<tr>
<th>CLEARANCE CRITERIA</th>
<th>REVIEWER</th>
<th>GUIDANCE</th>
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</table>

Applicant presents with a history of one or more of the following disorders:

<table>
<thead>
<tr>
<th>1. Ectopic Pregnancy</th>
<th>MED ADVISOR</th>
<th>Risk varies - assess based on detailed history</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. Solid Pelvic Mass or Tumor (includes Ovary)</td>
<td>MED ADVISOR</td>
<td>Requires evaluation and specific diagnosis for clearance. Consult with diagnostic guideline</td>
</tr>
<tr>
<td>3. Uterine Prolapse</td>
<td>MED ADVISOR</td>
<td>Risk varies - assess based on detailed history</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>633.0 Ectopic Pregnancy</td>
</tr>
<tr>
<td>789.3 Pelvic Mass</td>
</tr>
<tr>
<td>518.1 Uterine Prolapse</td>
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</tbody>
</table>

Cross Reference ICD.9.CM

<table>
<thead>
<tr>
<th>NOTES AND INSTRUCTIONS FOR REVIEWERS:</th>
</tr>
</thead>
</table>

Reviewers to Consider:
- None

<table>
<thead>
<tr>
<th>COMMENTS:</th>
</tr>
</thead>
</table>

Ectopic Pregnancy: Extrainuterine pregnancy - any pregnancy existing outside the confines of the uterine cavity.

Uterine Prolapse: Occurs when the integrity of supporting structures is lost. The allows the uterus to descend into the vagina. In advanced cases, complete protrusion with inversion of the vagina occurs. Approximately 1 in 10 women experience some degree of prolapse. Signs and symptoms include pelvic pressure and low back pain, dyspareunia, and difficulty with urination and defecation. Treatment includes Kegel exercises, estrogen replacement therapy, and surgery.

Literature review available.