

# Clinical Policy: Endometrial Ablation

Reference Number: CP.MP.106

Date of Last Revision: 03/22

<u>Revision Log Coding Implications</u>

See <u>Important Reminder</u> at the end of this policy for important regulatory and legal information.

#### **Description**

This policy describes the medical necessity guidelines for an endometrial ablation. Endometrial ablation is a minimally invasive surgical procedure used to treat premenopausal abnormal uterine bleeding. Although this procedure preserve the uterus, endometrial ablation is indicated for those who have no desire for future fertility. The two major classifications of endometrial ablation procedures are first generation resectoscopic techniques and second generation non-resectoscopic methods. Quality of life may improve following endometrial ablation procedures.

## Policy/Criteria

- I. It is the policy of health plans affiliated with Centene Corporation<sup>®</sup> that endometrial ablation using an FDA approved device is **medically necessary** when all the following criteria are met:
  - **A.** One of the following indications:
    - 1. Menorrhagia unresponsive to at least 3 months of hormonal or medical therapy (unless contraindicated to such therapy);
    - 2. Abnormal uterine bleeding, including residual menstrual bleeding after at least 6 months of androgen therapy in a female to male transgender person;
  - **B.** Cervical cytology or HPV testing and gynecological exam excludes significant cervical disease;
  - C. Endometrial sampling prior to the procedure has excluded malignancy or hyperplasia;
  - **D.** No structural anomalies, such as fibroids or polyps that require surgery or represent a contraindication to an ablation procedure, or previous transmyometrial uterine surgery (including classical cesarean);
  - **E.** If anatomic or pathologic conditions exist that may result in a weakened myometrium, only a resectoscopic endometrial ablation is appropriate;
  - **F.** Does not have any of the following contraindications:
    - 1. Premenopausal with future desire for fertility;
    - 2. Untreated disorders of hemostasis;
    - 3. Pregnancy at time of procedure;
    - 4. Intrauterine device at time of procedure;
    - 5. Active pelvic infection.
- II. It is the policy of health plans affiliated with Centene Corporation that there is insufficient scientific evidence to support effectiveness for the following:
  - **A.** Photodynamic endometrial ablation procedures;
  - **B.** Endometrial ablation for the treatment of all other conditions than those specified above.

#### **Background**

Menstrual disorders are among the most prevalent gynecological health problems in the United States, and abnormal menstrual bleeding affects up to 30% of people at some time during their



reproductive years.<sup>5</sup> Traditionally, medication therapy has been the initial treatment of choice, followed by hysterectomy, when medication does not provide the desired outcome. The levonorgestrel (LNG)-releasing intrauterine device (e.g., Mirena or Liletta; referred to as LNG 52) is an option in patients who do not desire pregnancy. Both the LNG 52 IUD and endometrial ablation are effective in reducing menstrual blood loss. The decision to use the LNG 52 or endometrial ablation depends on a patient's preferences regarding treatment factors, such as plans for fertility and contraception, convenience, and risks of anesthesia.<sup>22,25</sup> Endometrial ablation can offer an alternative to the more invasive hysterectomy treatment option.<sup>10</sup> Endometrial ablation is a minimally invasive surgical procedure used to treat premenopausal, abnormal uterine bleeding.

Endometrial ablation can also be used to treat residual menstrual bleeding in transgender men.  $^{24}$  Generally, masculinizing hormones cause cessation of menses within 2-6 months of initiation.  $^{18}$  The addition of a progestational agent or endometrial ablation may be considered for those wishing to completely cease menses.  $^{18}$ 

Endometrial ablation encompasses several techniques of targeted destruction of the endothelial surface of the uterine cavity through a vast array of energy sources. While hysterectomies provide permanent relief from abnormal uterine bleeding, they are associated with longer recovery times, higher rates of postoperative complications, substantial convalescent time and morbidity. 9,10 Although endometrial ablation has a high success rate, there are specific cases of endometrial ablation failures in which the patient will return for repeat care, often for a hysterectomy. 10 Among patients who return for hysterectomy after failure of endometrial ablation, endometriosis is the most common contributing diagnosis. 21

Pregnancy following endometrial ablation can occur, and premenopausal patients should be counseled that an appropriate contraception method should be used. Endometrial ablation is predominately indicated for patients who have no desire for future fertility. Post-operative complications from endometrial ablation include: (1) pregnancy after endometrial ablation; (2) pain-related to obstructed menses (hematometra, post ablation tubal sterilization syndrome); (3) failure to control menses; (4) risk from preexisting conditions (endometrial neoplasia, cesarean section; and (5) infection. Uterine perforation has been reported in 0.3 percent of non-resectoscopic endometrial ablation procedures and 1.3 percent of resectoscopic ablations or resections. 22

Table 1: FDA-Approved Techniques Approved For Endometrial Ablation

Procedure <sup>1,2,3</sup>	System <sup>1,2,13</sup>	Device	Treatment	Amenorrhea
Resectoscopic Ablation				
Laser Vaporization				37%
Electrosurgical Rollerball				25-60%
Transcervical resection of endometrium				26-40%
Radiofrequency Vaporization				N/A
Non-Resectoscopic Ablation				
Cryotherapy	Her Option	4.5	10–18	53%
Heated Free Fluid	Hydro ThermAblator	7.8	$\sim$ 14 $^*$	71%



Procedure <sup>1,2,3</sup>	System <sup>1,2,13</sup>	Device	Treatment	Amenorrhea
Microwave (no longer available in		8.5	2.5-4.5	61%
U.S.)				
Vapor ablation	Mara		2.0	
Radiofrequency Electricity	NovaSure	7.2	1.5	41%
Thermal Balloon	ThermaChoice	5.5	8.0	
Combined thermal and bipolar radiofrequency ablation device	Minerva		2.0	

<sup>\* 3</sup> minutes to heat the fluid to 90°C, 10 minutes to maintain that temperature to ablate the endometrium, and approximately 1 minute for the fluid to cool down allowing the device to be removed.

## **Coding Implications**

This clinical policy references Current Procedural Terminology (CPT®). CPT® is a registered trademark of the American Medical Association. All CPT codes and descriptions are copyrighted 2020 American Medical Association. All rights reserved. CPT codes and CPT descriptions are from the current manuals and those included herein are not intended to be all-inclusive and are included for informational purposes only. Inclusion or exclusion of any codes does not guarantee coverage. Providers should reference the most up-to-date sources of professional coding guidance prior to the submission of claims for reimbursement of covered services.

CPT® Codes	Description
58353	Endometrial ablation, thermal, without hysteroscopic guidance
58356	Endometrial cryoablation with ultrasonic guidance, including
	endometrial curettage, when performed
58563	Hysteroscopy, surgical; with endometrial ablation (eg, endometrial
	resection, electrosurgical ablation, thermoablation)

ICD-10-CM Diagnosis Codes that Support Coverage Criteria

ICD-10-CM	Description
Code	
N92.0	Excessive and frequent menstruation with regular cycle
N92.1	Excessive and frequent menstruation with irregular cycle
N92.4	Excessive bleeding in the premenopausal period
N92.5	Other specified irregular menstruation
N92.6	Irregular menstruation, unspecified
N93.8	Other specified abnormal uterine and vaginal bleeding
N93.9	Abnormal uterine and vaginal bleeding, unspecified

Reviews, Revisions, and Approvals	Revision Date	Approval Date
Policy developed, reviewed by specialist	12/15	01/16
Language clarifications d/t confusion in criteria, no specific criteria change:	06/16	



Reviews, Revisions, and Approvals	Revision Date	Approval Date
I.C clarified that structural anomalies be limited to those requiring	Duce	Duce
surgery or are otherwise a contraindication to EA		
I.E language clarified		
I.F removed anatomic or pathologic conditions affecting the		
myometrium as this is similar to I.C.		
I.F.2 added "untreated" for disorders of hemostasis		
Changed active pelvic inflammatory disease to active pelvic infection	08/16	9/16
Removed postmenopausal women from contraindications as this is a		
relative, not absolute, contraindication.		
Added indication for residual menstrual bleeding in female to male	09/16	10/16
transgender persons after androgen therapy, no codes added as ICD-10		
codes would still be applicable for new indication.		
References reviewed and updated	08/17	09/17
Added "previous transmyometrial uterine surgery" in I.D. References	06/18	07/18
reviewed and updated.		
Added additional FDA approved devices (i.e., Mara, Minerva) to table	06/19	07/19
1. References reviewed and updated. Specialist review.		
Added "abnormal uterine bleeding" as an indication and combined this	10/19	11/19
with the residual menstrual bleeding after androgen therapy in a female		
to male transgender person indication. Removed reference to criteria in		
CP.MP.95 Gender Affirming Procedures. Added the following codes		
as medically necessary: N92.5, N92.6, N93.8, N93.9.		
References reviewed and updated.	07/20	07/20
Annual review completed. References reviewed and updated and	07/21	07/21
reformatted for AMA style. Changed "members" to		
"members/enrollees." Removed "experimental and investigation"		
from II, changing to "insufficient evidence." Changed "review date" in		
the header to "date of last revision" and "date" in the revision log		
header to "revision date." Specialty review completed. Added		
ThermaChoice to Table 1 per UpToDate reference "3".		
Annual review completed. Added "or HPV testing" to I.B. References	03/22	03/22
reviewed and updated. Background updated with no impact to criteria.		

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## **Important reminder**

This clinical policy has been developed by appropriately experienced and licensed health care professionals based on a review and consideration of currently available generally accepted standards of medical practice; peer-reviewed medical literature; government agency/program approval status; evidence-based guidelines and positions of leading national health professional organizations; views of physicians practicing in relevant clinical areas affected by this clinical policy; and other available clinical information. The Health Plan makes no representations and accepts no liability with respect to the content of any external information used or relied upon in developing this clinical policy. This clinical policy is consistent with standards of medical practice current at the time that this clinical policy was approved. "Health Plan" means a health plan that has adopted this clinical policy and that is operated or administered, in whole or in part, by Centene Management Company, LLC, or any of such health plan's affiliates, as applicable.

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**Note: For Medicaid members/enrollees**, when state Medicaid coverage provisions conflict with the coverage provisions in this clinical policy, state Medicaid coverage provisions take precedence. Please refer to the state Medicaid manual for any coverage provisions pertaining to this clinical policy.

**Note: For Medicare members/enrollees,** to ensure consistency with the Medicare National Coverage Determinations (NCD) and Local Coverage Determinations (LCD), all applicable NCDs, LCDs, and Medicare Coverage Articles should be reviewed <u>prior to</u> applying the criteria set forth in this clinical policy. Refer to the CMS website at <a href="http://www.cms.gov">http://www.cms.gov</a> for additional information.

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