Obstetrical and Gynecological Services
# Revision History

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<th>Version</th>
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<tr>
<td>1.0</td>
<td>Policies and procedures as of October 1, 2015 Published: February 25, 2016</td>
<td>New document</td>
<td>FSSA and HPE</td>
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<td>1.1</td>
<td>Policies and procedures as of April 1, 2016 Published: September 27, 2016</td>
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| 3.0     | Policies and procedures as of July 1, 2018 Published: January 10, 2019 | Scheduled update:  
- Updated the note box at the beginning of the module with new standard wording  
- Reorganized and edited text as needed for clarity  
- Incorporated relevant information from the *Medical Policy Manual*  
- Updated links to the new IHCP website  
- Standardized *antepartum* and *antenatal* references to *prenatal*, for consistency  
- Added the *Gynecological Services* heading and introduction to group related subsections together  
- Removed specific USPTF guidelines in the *Cervical Cancer Screenings* section  
- Added the *Pelvic Examination* section  
- Clarified requirements and added a statement about retroactive eligibility in the *Informed Consent and Acknowledgement Statement for Hysterectomies* section  
- Removed the *Retroactive Eligibility for Hysterectomies* section | FSSA and DXC |
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<td>• Added the <em>Prior Authorization for Total or Partial Hysterectomy</em> section</td>
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<td>• Added a link to the related web page in the <em>Notification of Pregnancy</em> section</td>
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<td>• Updated the <em>Billing for Pregnancy-Related Services</em> section, including removing specific diagnosis codes for pregnancy-related services and adding a note box regarding copayments</td>
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<td>• Added a statement regarding national standards in the <em>Prenatal Care</em> section</td>
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<td>• Clarified information in the <em>Prenatal Visits</em> section and added a note box defining normal pregnancy</td>
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<td>• Added the <em>Prenatal Immunizations</em> section</td>
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<td>• Updated the <em>Injections for the Prevention of Preterm Delivery</em> section with new policies regarding Makena</td>
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<td>• Removed information about applicable settings in the <em>Placental Alpha Microglobulin-1 (PAMG-1) Test</em> section</td>
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<td>• Combined the sonography and echography sections under the single <em>Prenatal Ultrasounds (Sonography/Echography)</em> heading, and updating the text including:</td>
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<td>– Clarified terminology</td>
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<td>– Removed the specific diagnosis code for antenatal screening</td>
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<td>– Removed information about noncovered CPT code 59072</td>
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<td>– Removed reference to the <em>First-Trimester Fetal Nuchal Translucency Ultrasound Procedure Codes</em> table, which is being removed from <em>Obstetrical and Gynecological Codes</em></td>
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<td>• Reorganized and clarified the information in the General Billing Guidelines for Obstetrical Delivery section and its subsections, and also:</td>
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<td>– Added a note box regarding anesthesia</td>
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<td>– Clarified that vaginal deliveries that occur due to spontaneous labor do not require condition codes</td>
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<td>• Clarified information in the Postpartum Care section and removed references to the following code tables being removed from Obstetrical and Gynecological Codes:</td>
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<td>– Added source information for Table 1 – Approved Medical Indications for a Medically Necessary Delivery Prior to 39 Weeks and 0 Days</td>
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<td>– Added “members presenting in labor” to the table</td>
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<td>• Updated the Multiple Births section, including:</td>
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<td>– Added instructions for multiple births when all the births are by cesarean section</td>
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<td>– Added a reminder about trimester modifiers</td>
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<td>– Replaced assistant surgeon information with a reference to the Surgical Services module</td>
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<td>• Updated the Birthing Centers section and its subsections, including adding a list of service limitations</td>
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<td>• Removed unnecessary information from the Medical Abortion by Oral Ingestion of Medication section</td>
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Introduction

This module presents Indiana Health Coverage Programs (IHCP) coverage, billing, and reimbursement policies for gynecological and obstetrical services, including prenatal care, delivery, and postpartum care.

Gynecological Services

The IHCP covers gynecological services, including cervical cancer screenings, pelvic exams, and medically necessary hysterectomies, as described in the following sections. For information about contraception and sterilization, see the Family Planning Services module.

Cervical Cancer Screenings

The IHCP covers cervical cancer screening services, including cytology Pap smear and human papillomavirus (HPV) testing, as well as medically necessary services such as the collection of the samples, screening by a cytotechnologist, and a physician’s interpretation of the test results.

The IHCP follows the current recommendations for cervical cancer screening set by the U.S. Preventive Services Task Force (USPSTF) and the American Society for Colposcopy and Cervical Pathology (ASCCP). For repeat testing, cytological thresholds for further diagnostic testing (colposcopy) and treatments, and extended surveillance, the IHCP follows the recommendations of the ASCCP.

Detailed information regarding these recommendations can be found on the USPSTF website at uspreventativeservicestaskforce.org and the ASCCP website at asccp.org.

Pelvic Examination

The IHCP covers pelvic examinations (including breast examination) for female members.

A pelvic exam performed under anesthesia may be done as part of another gynecological surgical procedure or as a single procedure. The IHCP covers anesthesia/conscious sedation, when required for a member, to enable the practitioner to complete the exam. Based on accompanying documentation, medically necessary care provided prior to surgery will be reimbursed.
Hysterectomy

The IHCP covers medically necessary hysterectomies performed to treat an illness or injury. The IHCP does not cover hysterectomies performed solely to render a member permanently incapable of bearing children, whether performed as a primary or secondary procedure. For information about sterilization services, see the Family Planning Services module.

Hysterectomy procedures must comply with Code of Federal Regulations 42 CFR 441.250-441.259 and with 405 IAC 5-28-9. Hysterectomy is subject to prior authorization. In accordance with 42 CFR 441.255, the IHCP pays for hysterectomies performed during an individual’s retroactive eligibility if documentation requirements are met.

Informed Consent and Acknowledgement Statement for Hysterectomies

The IHCP covers hysterectomy only when medically necessary, and only when the member has given informed consent. The provider must have informed the member orally and in writing that the procedure will render the member permanently incapable of reproducing, and the member or member’s representative must have signed a written acknowledgement of receipt of that information, as shown in the example in Figure 1. Providers cannot use the Consent for Sterilization form for hysterectomy procedures under any circumstances.

Figure 1 – Example of Acknowledgement of Receipt of Hysterectomy Information

Acknowledgement of Receipt of Hysterectomy Information

Member Name: ____________________________
IHCP Member ID: __________________________
Physician Name: __________________________
NPI or IHCP Provider ID: ____________________
AMA Education Number: ____________________

It has been explained orally and in writing to __________________________ that the hysterectomy to be performed on her will render her permanently incapable of bearing children.

☐ Signed before surgery
☐ Signed after surgery (at the time of the hysterectomy, eligibility was not established).

(Member or Representative Signature) __________________________ (Date) __________________________

Physician Statement

The hysterectomy in the above case is being done for medically necessary reason(s), and the resulting sterilization is incidental and is not, at any time ever, the reason for this surgical operation.

Diagnosis(ses)

__________________________________________

(Physician Signature) __________________________ (Date) __________________________
The signed acknowledgement of receipt of hysterectomy information is required in all cases, except when the patient is already sterile or a life-threatening emergency exists for which the physician determines prior acknowledgement is not possible. The physician who performs the hysterectomy under these circumstances must complete one of the following certification requirements:

- Certify in writing that the individual was already sterile at the time the hysterectomy was performed. The certification must state the cause of the sterility at the time of the hysterectomy.
- Certify in writing that the hysterectomy was performed under a life-threatening emergency in which the physician determined that prior acknowledgement was not possible. The physician must also include a description of the nature of the emergency.

Claims billed with the CPT or ICD-10 procedure codes for hysterectomy services, shown in the CPT Procedure Codes for Hysterectomy and ICD-10 Procedure Codes for Hysterectomy tables in Obstetrical and Gynecological Services Codes, accessible from the Code Sets page at in.gov/medicaid/providers, require documentation necessary to satisfy requirements for informed consent or physician certification of preexisting sterility or life-threatening emergency.

Providers must attach the appropriate documentation to the paper claim form, upload it to the Portal claim, or send it separately as an attachment to the electronic claim transaction (as described in the Paper Attachments with Electronic Claims section of the Claim Submission and Processing module). Providers of all hysterectomy-related services must attach a copy of the appropriate acknowledgement or physician certification to the claim. This requirement extends to all providers, including attending physicians and surgeons, assistant surgeons, anesthesiologists, inpatient and outpatient hospital facilities, and other providers of related services. The primary service provider should forward copies of the acknowledgement or physician certification statement to the related service providers to ensure timely payment.

**Prior Authorization for Total or Partial Hysterectomy**

Prior authorization for total or partial hysterectomies will be granted for members with documentation supporting one of the following:

- Nonmalignant uterine tumor causing abnormal pressure or bleeding (lasting longer than 8 days for more than two cycles, requiring additional bleeding protection, defined as large clots and gushes, limiting activity)
- Nonmalignant uterine tumor causing one of the following:
  - Uterus of 12-week gestational size or larger, with ill-defined adnexa (less than 12-week gestational size, or less than 8 cm could have vaginal procedure)
  - Postmenopausal enlargement (more than 12-week gestational size necessitates abdominal procedure)
  - Rapid uterine growth over the last 6 months
  - Pressure on adjacent organs
- Cervical intraepithelial neoplasia (CIN) III, diagnosed by endocervical curettage, uncontrolled by conservative surgery, such as laser excision, loop electrosurgical excision procedure (LEEP), large loop excision of transformation zone (LLETZ), or loop surgical excision
- Fibroids in premenopausal woman with both of the following:
- Uterus greater than 12 weeks’ size or documentation of need for abdominal, rather than vaginal, approach; and one of the following:
  - Abnormal bleeding
  - Uterus size doubled within 1 year
- Ureteral compression by ultrasound or intravenous pyelogram (IVP)
- Other symptoms, such as pelvic or abdominal pain or discomfort without other explanation, urinary frequency or urgency, or dyspareunia

- Fibroids in postmenopausal woman with all of the following:
  - Uterus greater than 12 weeks’ size or documentation of need for abdominal rather than vaginal approach; and one of the following:
  - Uterus size doubled within any time period
  - Ureteral compression by ultrasound or IVP
  - Other symptoms, such as pelvic or abdominal pain or discomfort without other explanation, urinary frequency or urgency, or dyspareunia
  - Pap smear within 6 months

- Dysfunctional uterine bleeding with all of the following:
  - Premenopausal woman
  - Abnormal bleeding uncontrolled by conservative therapy, such as hormonal therapy
  - No evidence of cancer demonstrated by hysteroscopy, endometrial biopsy, dilation and curettage (D&C), or transvaginal ultrasound
  - Pap smear within 6 months

- Postmenopausal bleeding with all of the following:
  - Abnormal bleeding continued after change in or discontinuation of hormone replacement therapy
  - No evidence of cancer demonstrated by hysteroscopy, endometrial biopsy, D&C, or transvaginal ultrasound
  - Pap smear within 6 months

- Pelvic inflammatory disease (PID) with one of the following:
  - Suspected rupture or leakage of pelvic abscess
  - Unsuccessful management with antibiotics for 10 to 14 days
  - Surgery for residual, inactive but symptomatic disease, if conservative therapy is not possible.

- Chronic PID with both of the following:
  - Chronic pelvic pain
  - Adhesions, scarring, or hydrosalpinx

- Recurrent abnormal uterine bleeding (lasting longer than eight days for more than two cycles, requiring additional protection, defined as large clots and gushes, with limitations of normal activity) and benign endometrial biopsy after failed medication therapy – excluding members on birth control pills or those with intrauterine devices (IUDs).

- Chronic incapacitating pelvic pain, unresponsive to conservative therapy, such as analgesics, and evidence of normal gastrointestinal (GI)/genitourinary (GU) evaluations.
  - A four- to six-month failed trial of oral contraceptives, diuretics, anti-inflammatories, or induced amenorrhea
  - Negative examinations of UT, GI tract, and musculoskeletal
  - Psychological and psychosexual counseling reveals no etiology of pain

- Postmenopausal bleeding more than 1 year after LMP, with D&C or endometrial biopsy within past 6 months; positive cytology of cervix requires abdominal procedure (cervical intra-epithelial neoplasia including carcinoma in situ)

- Premalignant adenomatous hyperplasia or adenocarcinoma of the endometrium, confirmed by pathology report
- Benign or malignant ovarian tumor and/or cyst in postmenopausal (more than 1 year) women
- Abdominal procedure when associated with correction of urinary stress incontinence or vaginal repair of cystocele, rectocele, enterocele, or uterine prolapse
- Uncontrolled postpartum bleeding within six 6 of delivery, uncontrolled by drug therapy (for example, Pitocin, Methergine, or Prostaglandin therapy) or D&C
- Endometriosis uncontrolled by hormonal therapy (for example, depot medroxyprogesterone, oral contraceptives, Gonadotropin-releasing hormone [GnRH] agonist, or danazol), surgical ablation, or excision
- Tubo-ovarian abscess
- Urinary incontinence due to fistula into vagina, uterus, or perineum, and fistula demonstrated by cystoscopy, radiological examination, visual inspection, or probing
- Uterine prolapse, second or third degree, and one of the following:
  - Pain
  - Pelvic pressure
  - Stress incontinence
  - Ulceration of vaginal mucosa or cervix with bleeding or spotting
  - Vaginal splinting

**Notification of Pregnancy**

Early prenatal care can address potential health risks that contribute to poor birth outcomes. In addition, earlier enrollment of pregnant women in Medicaid case management programs is associated with better birth outcomes. The Family and Social Services Administration (FSSA) data shows that some low-income pregnant women do not seek prenatal services in the earliest stages of pregnancy, which often leads to untreated health risks. The FSSA Neonatal Quality Committee, made up of Indiana health professionals, has identified early prenatal care and the identification of health-risk factors of expectant mothers as an area of focus.

Within managed care programs, the FSSA uses the Notification of Pregnancy (NOP) form to improve the identification of health-risk factors of expectant mothers as early as the first trimester of pregnancy. NOPs can be completed at any time during the managed care member’s pregnancy, preferably during the initial visit, to document and monitor pregnancy conditions. If a managed care member’s normal pregnancy becomes high-risk (see the **High-Risk Pregnancy** section), providers should use the NOP to document the change.

Providers may receive $60 for one NOP per managed care member, per pregnancy. The following requirements must be met for a provider to be eligible for reimbursement for submitting an NOP:

- The NOP must be submitted via the Portal no more than 5 calendar days from the date of the office visit on which the NOP is based.
- The member’s pregnancy must be less than 30 weeks gestation at the time of the office visit on which the NOP is based.
- The member must be enrolled with a managed care entity (MCE), including pregnant women enrolled in an MCE through HIP, Hoosier Care Connect, or Hoosier Healthwise, as well as presumptively eligible pregnant women enrolled with an MCE.
- The NOP cannot be a duplicate of a previously submitted NOP.
Note:  Duplicate NOPs (those for the same woman and the same pregnancy) do not qualify for the $60 reimbursement. Only one NOP per member, per pregnancy is eligible for reimbursement. Recognized providers receive a systematic message if the NOP appears to be a duplicate.

For more information on NOP, see the Notification of Pregnancy page at in.gov/Medicaid/providers.

**Process for Completion of the Notification of Pregnancy**

Recognized providers complete and submit the NOP electronically using the Portal. After logging in, complete the following steps:

1. Select the **Eligibility** tab to verify the member’s eligibility.
2. In the **Eligibility Verification Request** panel, enter any of the following three search criteria for the member:
   - Member ID
   - Social Security number (SSN) and birth date
   - Last name, first name, and birth date
3. Enter a date or date range for the inquiry. If no date is entered in the Effective From field, the system defaults to the current date.
4. Click **Submit**.
5. In the Coverage column of the **Eligibility Verification Information** panel, click the link for the member’s managed care plan to access the **Coverage Details** page.
6. Click [-] to expand the **Managed Care Assignment Details** panel and then click **Enter NOP** to begin the process of completing the NOP online. (The option to print a blank NOP is also available; however, note that only NOPs submitted online are reimbursable.)
7. Complete all information on the NOP form. An asterisk (*) indicates a required field.
8. Click **Submit** to submit the NOP.
9. The Portal checks for potential duplicate NOPs. If a duplicate is identified, the recognized provider is asked to provide a reason why the new NOP is not a duplicate. The recognized provider can choose from three reasons related to the prior pregnancy:
   - Member abortion
   - Member preterm delivery
   - Member miscarriage
   
   The provider can continue the process without identifying a reason; however, the duplicate NOP will not be reimbursed.
10. After submitting the NOP, click **Print NOP** to print the completed NOP for documentation purposes, or click **Close** to close the window without printing.

Note: Submit the NOP within 5 calendar days from the date of the office visit. NOPs submitted more than 5 days from the date of the office visit are not reimbursed.
Billing for Submitted Notifications of Pregnancy

For NOP claims, bill using Current Procedural Terminology (CPT®) code 99354 with modifier TH:

- 99354 – Prolonged evaluation and management or psychotherapy service(s) (beyond the typical service time of the primary procedure) in the office or other outpatient setting requiring direct patient contact beyond the usual service, first hour (list separately)

- TH – Obstetrical treatment/services, prenatal or postpartum

The date of service on the NOP claim should be the date the provider completed the risk assessment during a visit with the pregnant woman.

**NOP claims should be submitted to the appropriate managed care entity.** Physicians can submit claims for NOP reimbursement using the CMS-1500 claim form or the 837P electronic transaction. Hospitals can submit claims for NOP reimbursement using the UB-04 claim form or the 837I electronic transaction. NOP claims from hospitals must be coded with revenue code 960 – Professional fees– General, in addition to CPT code 99354 with modifier TH.

Billing for Pregnancy-Related Services

Providers must indicate pregnancy and enter the date of last menstrual period (LMP) on all professional claims for pregnancy-related services. Providers must indicate pregnancy and include the LMP date as follows, depending on claim submission method:

- **CMS-1500 claim form** – Enter the LMP date in field 14. Enter the pregnancy indicator P in field 24H for each service detail.

- **Provider Healthcare Portal (Portal) professional claim** – During Step 1 of the claim submission process, in the Claim Information section, select Pregnancy as the Date Type and enter the LMP date in the Date of Current field.

- **837P electronic transaction** – Indicate pregnancy by submitting Y in PAT09 in the 2000 loop. Submit LMP information in the DTP segment in the 2300 loop with a qualifier of 484.

When billing for pregnancy-related services on the professional claim, providers must indicate a pregnancy-related diagnosis code as the primary diagnosis (the first diagnosis code entered on the claim) and for each service detail, using diagnosis pointers. The IHCP limits payment for pregnancy-related services subject to prior authorization restrictions and in accordance with Indiana Administrative Code (IAC). See Obstetrical and Gynecological Services Codes, accessible from the Code Sets page at in.gov/medicaid/providers, for a list of diagnosis codes for normal, low-risk pregnancy and for high-risk pregnancy.

Providers must enter the charged amount for each prenatal visit and for each postpartum visit. The charged amount is entered in field 24F ($ Charges) on the CMS-1500 claim form, the Charged Amount field in the Portal professional claim, or the Line Item Charge Amount field on the 837P electronic transaction.

Federal regulations allow providers to bill claims for certain prenatal services to the IHCP first, even if the member has insurance coverage through another carrier. For a list of relevant diagnosis codes, see Prenatal and Preventive Pediatric Care Diagnosis Codes That Bypass Cost Avoidance, accessible from the Code Sets page at in.gov/medicaid/providers. For more information, see the Third Party Liability module.

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Prenatal Care

The IHCP covers prenatal care delivered according to national standards as outlined by the American College of Obstetricians and Gynecologists (ACOG), American Academy of Pediatrics (AAP), and the Agency for Healthcare Research and Quality (AHRQ).

Prenatal Visits

The IHCP reimburses up to 14 visits for prenatal care during a normal pregnancy, as follows:

- Three visits in the first trimester
- Three visits in the second trimester
- Eight visits in the third trimester

Note: A normal pregnancy is defined as one in which the physician determines the pregnant woman is not at risk of a preterm birth or poor pregnancy outcome due to medical or psychosocial reasons. Additional prenatal care visits are allowed for members considered to have a high-risk pregnancy; see the High-Risk Pregnancy section.

To identify prenatal visits in each trimester, providers must bill the procedure code for the visit in conjunction with the appropriate U1, U2, or U3 modifier for each specific date of service:

- U1 – Trimester one – 0 through 14 weeks, 0 days
- U2 – Trimester two – 14 weeks, 1 day through 28 weeks, 0 days
- U3 – Trimester three – 28 weeks, 1 day through delivery

For the first prenatal visit, an evaluation and management (E/M) procedure code may be used to accommodate the greater amount of work involved. For subsequent visits, prenatal-care-only codes should be used. For additional details, see the Prenatal Visit Procedure Codes and Billing Instructions table in Obstetrical and Gynecological Services Codes, accessible from the Code Sets page at in.gov/medicaid/providers.

Note: Prenatal-care-only CPT procedure codes 59425 and 59426 are not subject to National Correct Coding Initiative (NCCI) Column I/II editing when billed with modifiers U1, U2, or U3 and billed on the same date of service as applicable laboratory procedure codes.

Providers should list each prenatal visit individually on the professional claim (CMS-1500 claim form or electronic equivalent) and submit claims after each individual visit or at the end of the respective trimester. Prenatal visits in the same trimester should be billed within 30 days of the end of the trimester. Providers should bill prenatal care for pregnant members separately from the delivery and postpartum visits, using the appropriate procedure codes for each service.

See the Billing Information for Pregnancy-Related Services section for general billing information applicable to all pregnancy-related services.
Office Visits during Pregnancy for Concurrent Medical Condition

With the exception of the first prenatal visit (which providers can bill using an appropriate E/M code and applicable trimester modifier, as indicated in the Prenatal Visits section), E/M procedure codes should not be used for office visits related to prenatal care. However, providers can bill E/M procedure codes 99211–99215 for office visits rendered to pregnant members if the service is related to a concurrent medical condition requiring medical care or consultative referral. Providers must identify the concurrent condition as a primary or secondary condition by a valid ICD diagnosis code and indicate the appropriate diagnosis code in the diagnosis pointer field for the service billed. Additionally, providers can bill the first prenatal visit with E/M codes 99201–99215, the appropriate trimester modifier, and the expected date of delivery all indicated on the claim.

Prenatal Tests and Screenings

In addition to the prenatal visits, the IHCP covers prenatal tests and screenings delivered according to standards established by the ACOG and the AAP.

Providers can bill prenatal tests and screenings along with the appropriate visit code on the same CMS-1500 claim form, Portal professional claim, or 837P electronic transaction.

Note: Providers are not allowed to bill separately for each component of the total obstetrical panel when all the tests listed in the panel are performed on the same date of service. For example, if the total panel of tests and screenings is performed on the same date of service, providers must bill the total obstetrical panel using the bundled laboratory procedure code 80055.

HIV Testing of Pregnant Women and Newborns

Indiana Code IC 16-41-6-8 requires, as a routine component of prenatal care, that physicians, advanced practice nurses, or the physician’s or advanced practice nurse’s designee explain the purpose, risks, and benefits of human immunodeficiency virus (HIV) testing and order HIV tests for pregnant women. The results of this test are confidential. Pregnant women have the right to refuse this test. A signed statement acknowledging the pregnant woman was counseled and provided the information necessary to make an informed decision regarding whether or not to be tested must be maintained in the medical records.

If the woman consents to an HIV test, and the test is positive for HIV infection, the provider must inform the pregnant woman of the test results and provide treatment and referral options available to her for HIV prevention, healthcare, and psychosocial services. The physician must also discuss risk reduction activities, including methods to reduce the risk of perinatal HIV transmission and HIV transmission through breast milk.

As required in IC 16-41-6-4, a physician overseeing the care of a newborn infant may offer the parent the option of a confidential HIV test for the newborn within the first 48 hours after birth under the following circumstances:

- The mother of the newborn has not been previously tested for HIV.
- The mother of the newborn has refused an HIV test for the newborn.
- The physician believes that testing the newborn is medically necessary for reasons other than those listed above.

If, for religious reasons, the parent objects, in writing, to testing the newborn, the newborn is exempt from the testing requirement.
The results of the HIV test must be released to the newborn’s mother. If the test results are positive, the individual who provides the test results must provide the mother with treatment or referral options available to the newborn.

**Prenatal Immunizations**

Refer to the following resources for guidelines for the recommended immunizations for pregnant women:

- The ACOG [Immunization](https://acog.org) web page at acog.org
- The AAP [Immunization Schedules](https://aap.org) web page at aap.org
- The Centers for Disease Control and Prevention (CDC) [Immunizations Schedules](https://cdc.gov) web page at cdc.gov

**High-Risk Pregnancy**

A pregnancy may be considered *high risk* if at least one medical or psychosocial complication is identified in the current pregnancy, or in the pregnant woman’s obstetrical history, that places her at risk for preterm birth or a poor pregnancy outcome.

The IHCP does not determine conditions that may or may not complicate a pregnancy. Therefore, if a physician determines that an illness or injury could complicate a pregnancy or have an adverse effect on the pregnancy’s outcome, the IHCP allows billing for covered services provided to treat the illness or injury.

**The IHCP reimburses high-risk-pregnancy care only when provided by physicians.** Nonphysician providers that render pregnancy-related services to pregnant IHCP members must refer members identified as having high-risk pregnancies only to appropriate physicians.

To document high-risk pregnancies for managed care members, providers may retain a copy of the submitted NOP in the patient’s record for retrospective review. If a normal pregnancy becomes high-risk at any time during the pregnancy, providers should use the NOP to document the change. See the Notification of Pregnancy section of this document for details.

**Increased Reimbursement and Additional Prenatal Visits for High-Risk Pregnancy**

Members with risk factors that may adversely affect the outcome of the pregnancy if not adequately treated are considered to have *high-risk pregnancies*. These complications, usually identified during the prenatal assessment, may place the member and the fetus in a high-risk pregnancy category that requires greater physician management.

Therefore, the IHCP reimburses physicians practicing obstetrics an additional $10 per prenatal visit procedure code (59425 and 59426) when the provider indicates a high-risk pregnancy diagnosis code on the submitted claim and documents the specific medical high-risk factors in the member’s medical records. Ensure that this information is easily identifiable on the medical record for audit purposes.

To be eligible for the higher reimbursement for prenatal office visits for patients who present with high-risk factors, providers must use a diagnosis code from the O09 series to signify high-risk pregnancy. (See the ICD-10 Diagnosis Codes for High-Risk Pregnancy table in Obstetrical and Gynecological Services Codes, accessible from the Code Sets page at in.gov/medicaid/providers.) These codes, when billed with prenatal office visit procedure codes 59425 and 59426, increase the maximum fee allowed for these services by $10 per visit.
Members identified as high-risk patients may receive additional prenatal care visits beyond the maximum of 14 allowed for a normal pregnancy. Claims for these additional visits must indicate one of the high-risk pregnancy diagnosis codes (from the O09 series), the LMP, the appropriate CPT code (procedure code 59425 for visits two through six, and procedure code 59426 for visits in excess of six), and the corresponding modifier.

**Injections for the Prevention of Preterm Delivery**

The IHCP covers compounded 17-alpha hydroxyprogesterone (17P) and Makena injections when medically necessary for the prevention of preterm delivery. Both drugs may be billed through the pharmacy benefit process, as described in the [Pharmacy Services](#) module.

Coverage for Makena is also included under the IHCP medical benefit. HCPCS code J1726 – Injection, hydroxyprogesterone caproate, (Makena), 10 mg may be billed on professional and institutional-outpatient claims. The product’s National Drug Code (NDC) must be included. Separate reimbursement is available in the outpatient setting when J1726 is billed with revenue code 636.

Makena is considered medically necessary for pregnant women who have a history of spontaneous preterm delivery. For dates of service on or after July 1, 2018, prior authorization is not required for the use of Makena.

**Placental Alpha Microglobulin-1 (PAMG-1) Test**

The IHCP reimburses for the placental alpha microglobulin-1 (PAMG-1) test when the test is considered medically necessary to confirm the diagnosis of premature rupture of membranes (PROM) or preterm premature rupture of membranes (PPROM). Prior authorization is not required for PAMG-1 testing; however, claims for this test are closely monitored for appropriateness of usage.

For reimbursement, providers should bill CPT code 84112 – Evaluation of cervicovaginal fluid for specific amniotic fluid protein(s) (eg, placental alpha macroglobulin-1 [PAMG-1], placental protein 12 [PP12], alpha-fetoprotein), qualitative, each specimen along with the appropriate trimester modifier (U1, U2, or U3). One PAMG-1 test equals one unit of service.

**Prenatal Ultrasounds (Sonography/Echography)**

The IHCP does not reimburse for routine ultrasounds or for ultrasounds performed for gender determination. A diagnosis of normal pregnancy does not explain the reason for the ultrasound. Prenatal ultrasounds performed when indicated for medical necessity should list an ICD-10 diagnosis code from the Z34 series (for normal pregnancy) or the O09 series (for high-risk pregnancy) as the primary diagnosis (see Obstetrical and Gynecological Services Codes, accessible from the [Code Sets](#) page at in.gov/medicaid/providers, for complete lists of these codes), and the appropriate antenatal screening diagnosis code as the secondary diagnosis.

Pregnancy-related ultrasounds billed without a secondary diagnosis to support medical necessity are subject to recoupment. Documentation in the patient’s medical record must substantiate the medical need for the ultrasound.

**Indications for Medical Necessity of Ultrasound**

The IHCP reimburses for ultrasounds performed during pregnancy when warranted by one or more of the following conditions:

- Early diagnosis of ectopic or molar pregnancy
- Placental localization associated with abnormal bleeding
When obstetrical and gynecological services are provided, the IHCP reimburses for the following:

- Fetal postmaturity syndrome
- Suspected multiple births
- Suspected congenital anomaly
- Polyhydramnios or oligohydramnios
- Guide for amniocentesis
- Fetal age determination, if necessitated by the following:
  - Discrepancy in size versus fetal age
  - Lack of fetal growth or suspected fetal death

First-Trimester Fetal Nuchal Translucency Ultrasound

The first-trimester fetal nuchal translucency ultrasound does not require prior authorization. However, the first-trimester fetal nuchal translucency ultrasound must be performed in conjunction with a maternal serum-free beta human chorionic gonadotropin (hCG) test and a pregnancy-associated plasma protein A (PAPP-A) test for the detection of chromosomal defects. The IHCP does not cover first-trimester fetal nuchal translucency testing when performed alone for the detection of chromosomal defects, as it is considered investigational.

For optimal test results, the first-trimester fetal nuchal translucency ultrasound should be performed between 11 and 13 weeks of pregnancy. First-trimester fetal nuchal translucency ultrasounds are subject to the requirements found in 405 IAC 5-27-6.

The IHCP does not provide reimbursement for routine ultrasounds or ultrasounds performed for gender determination. The diagnosis of a normal pregnancy does not substantiate the medical necessity for an ultrasound to be performed. Documentation must be maintained in the patient’s medical record to support the medical need for an ultrasound.

Obstetrical Delivery and Postpartum Care

The IHCP provides reimbursement for obstetrical delivery and postpartum care when all coverage and billing requirements are met.

General Billing Guidelines for Obstetrical Delivery

The IHCP follows CPT guidelines for obstetrical delivery billing. Delivery services include the following:

- Admission to the hospital
- Admission history and physical examination
- Management of uncomplicated labor
- Vaginal delivery (with or without episiotomy, with or without forceps) or cesarean delivery
Medical problems complicating labor and delivery management may require additional resources, and
physicians should identify related services by using the codes in the Evaluation and Management Services
module, in addition to codes for maternity care.

**Note:** The IHCP covers anesthesia services for a vaginal or cesarean delivery. For additional information, see the Anesthesia Services module.

Professional claims (CMS-1500 claim form or electronic equivalent) for obstetrical delivery (CPT codes 59409, 59514, 59612, and 59620) must include one of the following modifiers:

- UA – Nonmedically necessary delivery prior to 39 weeks of gestation
- UB – Medically necessary delivery prior to 39 weeks of gestation
- UC – Delivery at 39 weeks of gestation or later

Institutional claims (UB-04 claim form or electronic equivalent) for obstetrical delivery services related to C-sections or inductions require one of the following condition codes (fields 18–24 of the UB-04 claim form), in addition to the appropriate revenue codes and ICD procedure codes:

- 81 – C-sections or inductions performed at less than 39 weeks’ gestation for medical necessity
- 82 – C-sections or inductions performed at less than 39 weeks’ gestation electively
- 83 – C-sections or inductions performed at 39 weeks’ gestation or greater

**Note:** Vaginal deliveries that occur due to spontaneous labor do not require condition codes on the institutional claim.

See the Early Deliveries section for information about the appropriate uses of these gestational-stage modifiers and condition codes. For additional billing information specific to deliveries performed in freestanding birthing centers, see the Birthing Centers section.

**Postpartum Care**

The IHCP provides reimbursement for inpatient or outpatient postpartum visits within 60 days after delivery. The IHCP does not cover CPT codes for combined delivery and postpartum care (59410, 59515, 59614, or 59622) on fee-for-service (FFS) claims. Providers are required to bill delivery services and postpartum care services separately using the appropriate procedure codes

Postpartum physician visits within 60 days after delivery are billed using CPT code 59430, which is for postpartum care only.

**Early Deliveries**

The IHCP does not cover early elective deliveries (EEDs), defined as deliveries performed prior to 39 weeks and 0 days gestation without medical indication. The IHCP does not reimburse for delivery CPT codes submitted with the UA modifier, signifying deliveries at less than 39 weeks of gestation that do not meet the IHCP’s stated guidelines for approved medically necessary deliveries. Additionally, the IHCP does not reimburse institutional claims submitted with condition code 82, signifying an elective C-section or induction performed at less than 39 weeks of gestation. This EED policy applies to all IHCP programs.

Deliveries that meet one of the approved medical indications for a medically necessary delivery prior to 39 weeks (listed in Table 1) are covered. The medical indications listed in Table 1 are compiled from lists released by the Indiana Perinatal Quality Improvement Collaborative (IPQIC), ACOG, and the Joint Commission. This comprehensive list of medical indications is intended to ensure that all medically indicated
deliveries prior to 39 weeks remain covered. The IHCP will continue to evaluate the list of approved medical indications to ensure that all medically necessary indications are covered.

For all early deliveries, documentation of the gestational age of the fetus and the medical indication for an early delivery must be completed and maintained in the member’s file. Suggested forms for documentation are the ACOG Patient Safety Checklists on the ACOG website at acog.org or the IPQIC Scheduling form on the ISDH website at in.gov.

### Table 1 – Approved Medical Indications for a Medically Necessary Delivery Prior to 39 Weeks and 0 Days

<table>
<thead>
<tr>
<th>Maternal Indications</th>
<th>Fetal Indications</th>
<th>Obstetric Indications</th>
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<tr>
<td>Antiphospholipid syndrome</td>
<td>Abo isoimmunization</td>
<td>Members presenting in labor</td>
</tr>
<tr>
<td>Chronic hypertension</td>
<td>Abnormal fetal heart rate</td>
<td>Abruptio placenta</td>
</tr>
<tr>
<td>Cardiovascular diseases</td>
<td>Chorioamnionitis</td>
<td>Abruption</td>
</tr>
<tr>
<td>Chronic pulmonary disease</td>
<td>Congenital heart defect/heart disease</td>
<td>Antepartum hemorrhage/bleeding</td>
</tr>
<tr>
<td>Coagulopathy defect</td>
<td>Fetal abnormality</td>
<td>Chronic hypertension with super imposed preeclampsia</td>
</tr>
<tr>
<td>Coagulopathy disorders</td>
<td>Fetal chromosomal anomaly</td>
<td>Chorioamnionitis</td>
</tr>
<tr>
<td>Congenital heart defect/heart disease</td>
<td>Fetal CNS anomaly</td>
<td>Gestational diabetes</td>
</tr>
<tr>
<td>Current cancer</td>
<td>Fetal damage due to disease</td>
<td>Gestational hypertension</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>Fetal damage due to drugs</td>
<td>Hypertensive disorder</td>
</tr>
<tr>
<td>Epilepsy/seizure disorder</td>
<td>Fetal damage due to radiation</td>
<td>Increta</td>
</tr>
<tr>
<td>Gastroenteric diseases/disorders</td>
<td>Fetal damage due to virus</td>
<td>Maternal/fetal hemorrhage</td>
</tr>
<tr>
<td>Hematological disorder</td>
<td>Fetal demise-singleton</td>
<td>Mild preeclampsia</td>
</tr>
<tr>
<td>HIV; asymptomatic HIV infection status</td>
<td>Fetal distress</td>
<td>Severe preeclampsia/HELLP/eclampsia</td>
</tr>
<tr>
<td>Hypertension nonspecified</td>
<td>Fetal/maternal hemorrhage</td>
<td>Multiple gestation/multiple gestation with loss</td>
</tr>
<tr>
<td>Liver disease</td>
<td>Intrauterine growth restriction</td>
<td>Oligohydramnios</td>
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<tr>
<td>Maternal/fetal hemorrhage</td>
<td>Nonreassuring fetal antepartum testing</td>
<td>Percreta</td>
</tr>
<tr>
<td>Previous stillborn</td>
<td>RH isoimmunization</td>
<td>Placenta accreta</td>
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<tr>
<td>Prior classical cesarean delivery</td>
<td></td>
<td>Placenta previa</td>
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<tr>
<td>Prior myomectomy entering endometrial cavity</td>
<td></td>
<td>Placental previa hemorrhage</td>
</tr>
<tr>
<td>Renal disease</td>
<td></td>
<td>Polyhydramnios</td>
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<tr>
<td></td>
<td></td>
<td>Premature rupture of membranes</td>
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<tr>
<td></td>
<td></td>
<td>Prolonged rupture of membranes</td>
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<tr>
<td></td>
<td></td>
<td>Ruptured membranes</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Unstable lie; multiple gestation with malpresentation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Vasa previa</td>
</tr>
</tbody>
</table>

### Multiple Births

Multiple-birth deliveries are subject to multiple-surgery reimbursement. The reimbursement policy indicated in 405 IAC 5-28-1(g) for pricing multiple surgical procedures states that 100% of the global fee is reimbursed for the most expensive procedure. The second most expensive procedure is reimbursed at 50% of the global fee, and remaining procedures are reimbursed at 25% of the global fee.
The IHCP reimburses for only one cesarean procedure, regardless of the number of babies delivered during the cesarean section. Therefore, only one detail line with one unit of service is billed for cesarean delivery procedure codes. Multiple births should be billed as follows:

- When billing for multiple births and all the births are by cesarean section, the births are billed as a single detail with a single unit of either procedure code 59514 – Cesarean delivery only or 59620 – Cesarean delivery only, following attempted vaginal delivery after previous cesarean delivery.

- When billing for multiple births and all the births are vaginal deliveries, the first birth is billed using procedure code 59409 – Vaginal delivery only (with or without episiotomy and/or forceps) or 59612 – Vaginal delivery only, after previous cesarean delivery (with or without episiotomy and/or forceps). The second birth and any subsequent births are billed using procedure codes 59409 or 59612 with modifier 51 – Multiple procedures.

- When billing for one vaginal birth and one or more births by cesarean section, the cesarean birth is billed with procedure code 59514 or 59620, and the vaginal birth is billed using procedure code 59409 or 59612 with modifier 51.

- When billing for two or more vaginal births and one or more births by cesarean, the cesarean births are billed on one detail line with one unit of service using procedure code 59514 or 59620. The vaginal births are billed as separate details using procedure code 59409 or 59612 with modifier 51.

The appropriate UA, UB, or UC modifier is also required for all CPT delivery codes, as described in the General Billing Guidelines for Obstetrical Delivery section. For modifier requirements and reimbursement information related to assistant surgeon services during delivery, see the Surgical Services module.

**Reimbursement for Long-Acting Reversible Contraception Implanted During Delivery Stays**

The IHCP allows separate reimbursement for certain long-acting reversible contraception (LARC) devices implanted during an inpatient hospital or birthing center stay for a delivery. For applicable Healthcare Common Procedure Coding System (HCPCS) codes, see Obstetrical and Gynecological Services Codes, accessible from the Code Sets page at in.gov/medicaid/providers.

To receive separate reimbursement for LARC devices implanted during inpatient hospital or birthing center stays for delivery, the appropriate HCPCS code should be billed on a professional claim (CMS-1500 claim form or electronic equivalent). Separate reimbursement applies to the LARC devices only. Reimbursement for all other related services, procedures, supplies, and devices continue to be included in the inpatient hospital diagnosis-related group (DRG) or the birthing center all-inclusive reimbursement amount.

For more information about LARC devices, see the Family Planning Services module.

**Birthing Centers**

A freestanding birthing center, as defined by IC 16-18-2-36.5 and 410 IAC 27-1-3, is a licensed, freestanding entity that has the sole purpose of delivering a normal or uncomplicated (low-risk) pregnancy. This term does not include a hospital under IC 16-21-2, an ambulatory surgical center, or the residence of the woman giving birth.

Birthing centers are licensed to provide care during pregnancy, birth, and the immediate postpartum period to the low-risk expectant mother and her newborn. Each center shall admit and retain only low-risk expectant mothers anticipating a normal, full-term, spontaneous vaginal birth.
Services provided in a birthing center shall be limited in the following manner:

- Members receiving the services must be considered low-risk, or having a normal, uncomplicated pregnancy as defined in 410 IAC 27-1-15.5.
- Delivery shall be performed by one of the following professionals:
  - Certified nurse midwife
  - Physician
- Surgical services are limited to episiotomy and episiotomy repair, and shall not include operative obstetrics or cesarean sections.
- Labor shall not be inhibited, stimulated, or augmented with chemical agents during the first or second stage of labor.
- Systemic analgesia may be administered and local anesthesia for pudendal block and episiotomy repair may be performed.
- General and conductive anesthesia shall not be administered at birthing centers.
- Members shall not routinely remain in the facility in excess of 24 hours.

**Facility Billing and Reimbursement**

The IHCP created provider type 08 – Clinic and provider specialty code 088 – Birthing center to identify birthing centers. Birthing centers must be licensed by the Indiana State Department of Health (ISDH) before enrolling in the IHCP. Birthing centers are assigned to the limited risk category and are not required to pay an application fee during enrollment or revalidation (see the IHCP Provider Enrollment Risk Category and Application Fee Matrix at in.gov/medicaid/providers). Providers should refer to the IHCP Provider Type and Specialty Matrix at in.gov/medicaid/providers for other enrollment criteria.

Facility charges are billed on an institutional claim (UB-04 claim form or electronic equivalent). Birthing center claims must report billing provider taxonomy code 261QB0400X (birthing) on the claim.

Birthing centers are paid at an all-inclusive rate. The services are billed using revenue code 724 – Birthing center. Only vaginal deliveries should be billed with this revenue code. Reimbursement rates are based on the revenue code 724 when the member delivers. When labor occurs but does not result in delivery, providers should bill revenue code 724 along with HCPCS code S4005 – Interim labor facility global (labor occurring but not resulting in delivery).

**Professional Billing and Reimbursement**

Professional services rendered at birthing centers are billed on a professional claim (CMS-1500 claim form or electronic equivalent). Services rendered by the following providers are payable when performed at birthing centers:

- Certified nurse midwife (Provider type 09, specialty 095)
- Physician (Provider type 31, all specialties)

Professional charges are reimbursed directly to the practitioner at the applicable reimbursement rate. Other staff services, such as services provided by registered nurses (RNs) and licensed nurse practitioners (LPNs), are included in the delivery rate and not separately reimbursed.

Birthing center professional services are to be billed with place-of-service code 25 – Birthing center.
Abortion and Related Services

The IHCP uses the word abortion to describe the early termination of pregnancy. The IHCP does not consider termination of an ectopic pregnancy to be an abortion.

There are two types of abortion:

- **Spontaneous abortion (or missed abortion)** occurs for no apparent reason during early pregnancy and requires treatment to ensure the health of the mother. The IHCP reimburses for therapeutic treatment of spontaneous or missed abortion, and services relevant to this treatment, according to the IHCP-allowable amount. Providers should follow the coding guidelines included in this section.

- **Elective abortion** is an abortion that a doctor performs because the mother has chosen to terminate the pregnancy. IC 16-34-1-2 prohibits the State from making payment from any fund under its control for an elective abortion, unless the elective abortion is necessary to preserve the life of the pregnant woman or unless federal law requires the State to cover it, such as in the case of rape or incest. Elective abortions performed for any other reason are noncovered services according to 405 IAC 5-28-7. Providers must adhere to the procedures described in the following section to obtain payment for an elective abortion.

Documentation Requirements

For spontaneous abortions, the IHCP requires no documentation from providers billing with the appropriate treatment code and following the guidelines described in this document.

For elective abortions, the physician must specify in writing the physical condition of the patient leading to the professional judgment that the abortion was one of the following:

- Necessary to preserve the life of the pregnant woman
- Due to rape or incest

The documentation must contain the name and address of the member, dates of service, physician’s name, and physician’s signature. Providers must attach this documentation to the paper claim form, upload it to the Portal claim, or send it separately as an attachment to the electronic claim transaction (as described in the Paper Attachments with Electronic Claims section of the Claim Submission and Processing module). The IHCP must receive correct documentation with claims before it will make payment for the elective abortion or any directly related service. The primary service provider should forward copies of the physician certification to the related service provider to bill for these services.

If providers submit a claim with a diagnosis code or procedure code indicating that a possible elective abortion was performed, the IHCP requires documentation for claim payment consideration. The IHCP suspends these claims for review of medical documentation. For lists of diagnosis codes and procedure codes that suspend for appropriate documentation supporting medical necessity, see the following tables in Obstetrical and Gynecological Services Codes, accessible from the Code Sets page at in.gov/medicaid/providers:

- ICD-10 Abortion Diagnosis Codes That Suspend for Appropriate Documentation Supporting Medical Necessity
- CPT and HCPCS Abortion Procedure Codes That Suspend for Appropriate Documentation Supporting Medical Necessity
- ICD-10 Abortion Procedure Codes That Suspend for Appropriate Documentation Supporting Medical Necessity
Medical Abortion by Oral Ingestion of Medication

The IHCP reimburses for mifepristone and misoprostol for use in medical abortion procedures based on the same coverage criteria applicable to surgical abortions.

The IHCP reimburses only the FDA-approved regimen for medically induced abortions using orally administered mifepristone and misoprostol. The IHCP does not reimburse what is commonly known as the evidence-based regimen for medical abortion with mifepristone and misoprostol, which includes at-home or vaginal administration of misoprostol.

The FDA-approved regimen for these medications is as follows:

- **Recommended gestational age** – 49 days from last menstrual period
- **Mifepristone dose** – 600 mg orally administered on Day 1 office visit
- **Misoprostol dose** – 400 mcg orally administered on Day 3 office visit
- **Misoprostol timing** – 48 hours after receiving mifepristone

Medical abortion by oral ingestion of mifepristone and misoprostol requires three separate office visits to complete the procedure. Confirmation of pregnancy status must occur before the Day 1 office visit. The Day 1 office visit must occur after the 18-hour counseling and waiting period required by IC 16-34-2-1.1(a)(1).

The following list shows the billing guidelines for these office visits and the medications provided during the office visits. Providers must bill all claims for medical abortion by oral ingestion of mifepristone and misoprostol on the professional claim (CMS-1500 claim form or electronic equivalent).

- **Day 1:**
  - Member reviews and signs the Patient Agreement.
  - Provider orally administers three 200 mg tablets of mifepristone.
  - Provider bills HCPCS code S0190 – Mifepristone, oral, 200 mg, three units.
  - Provider bills the appropriate E/M code for the office visit.

- **Day 3:**
  - Provider checks pregnancy status with clinical examination or ultrasound exam.
  - If an ultrasound is performed, provider bills the appropriate code for the service provided.
  - Provider orally administers two 200 mcg tablets of misoprostol.
  - Provider bills HCPCS code S0191 – Misoprostol, oral, 200 mcg, two units.
  - Provider bills appropriate E/M code for the office visit.

- **Day 14:**
  - Provider verifies pregnancy termination with clinical examination or ultrasound exam.
  - If an ultrasound is performed, the provider bills the appropriate code for the service provided.
  - Provider bills appropriate E/M code for the office visit.

The IHCP suspends claims for Day 1 and Day 3 office visits pending submission of required documentation. To be reimbursed for services, the IHCP requires providers to submit all necessary documentation with claims for these office visits, as described in the Documentation Requirements section.

In addition, medical abortion by oral ingestion of mifepristone and misoprostol requires submission of the signed Prescriber’s Agreement and Patient Agreement. These agreements are available from Danco, and Danco requires their use. Providers must attach documentation to the paper claim form, upload it to the Portal claim, or send it separately as an attachment to the electronic claim transaction (as described in the
Paper Attachments with Electronic Claims section of the Claim Submission and Processing module). If providers fail to submit this documentation, the IHCP must deny the claims.