ARTICLE 26. ABORTION CLINICS

Rule 0.5. Applicability

410 IAC 26-0.5-1 Applicability
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-19-3; IC 16-21-1

Sec. 1. This article applies to abortion clinics that perform surgical abortion procedures. An abortion clinic that provides an abortion inducing drug for the purpose of inducing an abortion must comply with 410 IAC 26.5. (Indiana Department of Health; 410 IAC 26-0.5-1; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

Rule 1. Definitions

410 IAC 26-1-1 Applicability
Authority: IC 16-21-1-7; IC 16-21-1-9
Affected: IC 16-19-3; IC 16-21-1

Sec. 1. The definitions in this rule apply throughout this article except as otherwise indicated. (Indiana Department of Health; 410 IAC 26-1-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3355; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-2 "Abortion" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-18-2-1; IC 16-21-1; IC 16-21-2

Sec. 2. "Abortion" has the meaning set forth in IC 16-18-2-1. (Indiana Department of Health; 410 IAC 26-1-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3355; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-3 "Abortion clinic" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-18-2-163; IC 16-21-1; IC 16-21-2

Sec. 3. (a) "Abortion clinic" means a health care provider (as defined in IC 16-18-2-163(d)(1)) that performs surgical abortion procedures.
(b) The term does not include the following:
(1) A hospital that is licensed as a hospital under IC 16-21-2.
(2) An ambulatory outpatient surgical center that is licensed as an ambulatory outpatient surgical center under IC 16-21-2.
(3) A health care provider that provides, prescribes, administers, or dispenses an abortion inducing drug to fewer than five (5) patients per year for the purposes of inducing an abortion. (Indiana Department of Health; 410 IAC 26-1-3; filed May 11, 2006, 9:36 a.m.: 29 IR 3355; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

410 IAC 26-1-3.2 "Affiliate" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-18-2-9.4; IC 16-21-1; IC 16-21-2

Sec. 3.2. "Affiliate" has the meaning set forth in IC 16-18-2-9.4. (Indiana Department of Health; 410 IAC 26-1-3.2; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)
410 IAC 26-1-3.5 "ASA Class I patient" defined
Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 3.5. "ASA Class I patient" means a normal, healthy patient. (Indiana Department of Health; 410 IAC 26-1-3.5; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-4 "Authenticate" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 4. "Authenticate" means the author or responsible individual has reviewed the clinical content of the order and validated an entry in the record by:
(1) a full signature, including first initial, last name, and discipline;
(2) written initials if full signature appears on the same page; or
(3) a unique identifier such as a number or computer key.
(Indiana Department of Health; 410 IAC 26-1-4; filed May 11, 2006, 9:36 a.m.: 29 IR 3355; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-4.6 "Biologics" defined
Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 4.6. "Biologics" means a biological product, such as:
(1) a globulin;
(2) a serum;
(3) a vaccine;
(4) an antitoxin;
(5) blood; or
(6) an antigen;
used in the prevention or treatment of disease. (Indiana Department of Health; 410 IAC 26-1-4.6; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-4.8 "Burn" defined
Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 4.8. "Burn" means any injury or damage to the tissues of the body caused by exposure to any of the following:
(1) Fire.
(2) Heat.
(3) Chemicals.
(4) Electricity.
(5) Radiation.
(6) Gases.
(Indiana Department of Health; 410 IAC 26-1-4.8; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)
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410 IAC 26-1-5 "Clinic" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 5. "Clinic" means an abortion clinic. (Indiana Department of Health; 410 IAC 26-1-5; filed May 11, 2006, 9:36 a.m.: 29 IR 3355; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-6 "Commissioner" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 6. "Commissioner" means the state health commissioner or the state health commissioner's designee. (Indiana Department of Health; 410 IAC 26-1-6; filed May 11, 2006, 9:36 a.m.: 29 IR 3355; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-7 "Council" defined (Repealed)

Sec. 7. (Repealed by Indiana Department of Health; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

410 IAC 26-1-8 "Department" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2


410 IAC 26-1-9 "Division" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 9. "Division" means the division of acute care of the department. (Indiana Department of Health; 410 IAC 26-1-9; filed May 11, 2006, 9:36 a.m.: 29 IR 3355; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-9.5 "Elopement" defined
Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 9.5. "Elopement" means any situation in which a registered or admitted patient, excluding events involving adults with decision making capacity, leaves the clinic without staff being aware that the patient has done so. (Indiana Department of Health; 410 IAC 26-1-9.5; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-10 "Governing body" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 10. "Governing body" means:
(1) board of trustees;
(2) governing board;
(3) board of directors; or
(4) other body or individual responsible for governing an abortion clinic.

410 IAC 26-1-11 "Health care provider" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-18-2-163; IC 16-21-1; IC 16-21-2

Sec. 11. "Health care provider" has the meaning set forth in IC 16-18-2-163.

410 IAC 26-1-12 "Health care worker" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 12. "Health care worker" means a person who provides services whether as an individual health care provider, volunteer, or student at or employee of a clinic.

410 IAC 26-1-12.4 "Hypoglycemia" defined
Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 12.4. "Hypoglycemia" means a physiologic state in which:
(1) the blood sugar falls below sixty (60) mg/dl (forty (40) mg/dl in neonates); and
(2) physiological or neurological, or both, dysfunction begins.

410 IAC 26-1-12.5 "Immediately postoperative" defined
Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 12.5. "Immediately postoperative" means within twenty-four (24) hours after either of the following:
(1) Administration of anesthesia (if surgery or other invasive procedure is not completed).
(2) Completion of surgery or other invasive procedure.

410 IAC 26-1-12.6 "Informed consent" defined
Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 12.6. "Informed consent" means a patient's authorization or agreement to undergo surgery or other invasive procedure.
that is based upon communication between a patient and his or her physician regarding the surgery or other invasive procedure. (Indiana Department of Health; 410 IAC 26-1-12.6; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-12.7 "Intended use" defined (Repealed)

Sec. 12.7. (Repealed by Indiana Department of Health; filed Oct 7, 2008, 10:26 a.m.: 20081105-IR-410080061FRA)

410 IAC 26-1-12.8 "Joint movement therapy" defined

Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-7

Sec. 12.8. "Joint movement therapy" means all types of manual techniques, to include:
(1) mobilization (movement of the spine or a joint within its physiologic range of motion);
(2) manipulation (movement of the spine or a joint beyond its normal voluntary physiologic range of motion); or
(3) any other type of manual musculoskeletal therapy;
regardless of their precise anatomic and physiologic focus or their discipline of origin. (Indiana Department of Health; 410 IAC 26-1-12.8; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-12.9 "Kernicterus" defined

Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 12.9. "Kernicterus" means the medical condition in which elevated levels of bilirubin cause brain damage. (Indiana Department of Health; 410 IAC 26-1-12.9; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-13 "Licensed health professional" defined

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2; IC 25-23-1-27.1

Sec. 13. "Licensed health professional" has the meaning set forth in IC 25-23-1-27.1. (Indiana Department of Health; 410 IAC 26-1-13; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-13.5 "Low-risk pregnancy" defined

Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 13.5. "Low-risk pregnancy" means a woman sixteen (16) to thirty-nine (39) years of age, with no previous diagnosis of any of the following:
(1) Essential hypertension.
(2) Renal disease.
(3) Collagen-vascular disease.
(4) Liver disease.
(5) Preeclampsia.
(6) Cardiovascular disease.
(7) Placenta previa.
(8) Multiple gestation.
(9) Intrauterine growth retardation.
(10) Smoking.
(11) Pregnancy-induced hypertension.
(12) Premature rupture of membranes.
(13) Other previously documented condition that poses a high risk of pregnancy-related mortality.

(Indiana Department of Health; 410 IAC 26-1-13.5; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196FRA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328FRA)

410 IAC 26-1-14 "Medical staff" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 14. "Medical staff" means physicians appointed by the governing body or contracted with by the governing body and responsible to the governing body for the following:
(1) The clinical and scientific work of the clinic.
(2) Advice regarding professional matters and policies.
(3) Review of the professional practices in the clinic for the purposes of reducing morbidity and mortality and for the improvement of the care of patients in the clinic, including the following:
   (A) The quality and necessity of care provided.
   (B) The preventability of complications and deaths occurring in the clinic.

(Indiana Department of Health; 410 IAC 26-1-14; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196FRA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328FRA)

410 IAC 26-1-14.3 "Minimal sedation" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 14.3. "Minimal sedation" means a drug induced state during which patients respond normally to verbal commands although cognitive function and coordination may be impaired. Ventilatory and cardiovascular functions are unaffected.

(Indiana Department of Health; 410 IAC 26-1-14.3; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

410 IAC 26-1-14.6 "Moderate sedation" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 14.6. "Moderate sedation" means a drug induced depression of consciousness during which patients respond purposefully to verbal commands, either alone or accompanied by light tactile stimulation. No interventions are required to maintain a patent airway, and spontaneous ventilation is adequate. Cardiovascular function is usually maintained.

(Indiana Department of Health; 410 IAC 26-1-14.6; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

410 IAC 26-1-15 "Pharmacist" defined
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2; IC 25-26-13

Sec. 15. "Pharmacist" means an individual licensed under IC 25-26-13.

(Indiana Department of Health; 410 IAC 26-1-15; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196FRA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328FRA)
410 IAC 26-1-16 "Physician" defined  
Authority: IC 16-21-1-7; IC 16-21-2-2.5  
Affected: IC 16-21-1; IC 16-21-2; IC 25-22.5-5  
Sec. 16. "Physician" means an individual licensed under IC 25-22.5-5. (Indiana Department of Health; 410 IAC 26-1-16; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)  

410 IAC 26-1-17 "Registered nurse" defined  
Authority: IC 16-21-1-7; IC 16-21-2-2.5  
Affected: IC 16-21-1; IC 16-21-2; IC 25-23-1  
Sec. 17. "Registered nurse" means an individual licensed under IC 25-23-1. (Indiana Department of Health; 410 IAC 26-1-17; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)  

410 IAC 26-1-17.5 "Serious disability" defined  
Authority: IC 16-19-3-4; IC 16-21-1-7  
Affected: IC 16-19-3; IC 16-21-1  
Sec. 17.5. "Serious disability" means either of the following:  
(1) Significant loss of function including sensory, motor, physiologic, or intellectual impairment:  
(A) not present on admission and requiring continued treatment; or  
(B) for which there is a high probability of long term or permanent lifestyle change at discharge.  
(2) Unintended loss of a body part. (Indiana Department of Health; 410 IAC 26-1-17.5; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)  

410 IAC 26-1-17.6 "Sexual assault" defined  
Authority: IC 16-19-3-4; IC 16-21-1-7  
Affected: IC 16-19-3; IC 16-21-1; IC 35-42-4; IC 35-46-1-3  
Sec. 17.6. "Sexual assault" means a crime included under IC 35-42-4 or IC 35-46-1-3. (Indiana Department of Health; 410 IAC 26-1-17.6; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)  

410 IAC 26-1-17.8 "Surgery or other invasive procedure" defined  
Authority: IC 16-19-3-4; IC 16-21-1-7  
Affected: IC 16-19-3; IC 16-21-1  
Sec. 17.8. "Surgery or other invasive procedure", for purposes of 410 IAC 26-6-2, means surgical or other invasive procedures that involve a skin incision, puncture, or insertion of an instrument or foreign material into tissues, cavities, or organs. A procedure begins at the time of the skin incision, puncture, or insertion of an instrument or foreign material into tissues, cavities, or organs. A procedure ends when the surgical incision has been closed or operative devices such as probes have been removed. The procedures include, but are not limited to, the following:  
(1) Open or percutaneous surgical procedures.  
(2) Percutaneous aspiration.  
(3) Selected injections.  
(4) Biopsy.  
(5) Percutaneous cardiac and vascular diagnostic or interventional procedures.
(6) Laparoscopies.
(7) Endoscopies.
(8) Colonoscopies.

The term excludes intravenous therapy, venipuncture for phlebotomy, or diagnostic tests without intravenous contrast agents. (Indiana Department of Health; 410 IAC 26-1-17.8; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; filed Oct 7, 2008, 10:26 a.m.: 20081105-IR-410080061FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-18 "Surgical abortion" defined

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 18. "Surgical abortion" means the use of a:
(1) surgical instrument; or
(2) machine;

to perform an abortion. (Indiana Department of Health; 410 IAC 26-1-18; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-1-19 "Toxic substance" defined

Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1

Sec. 19. "Toxic substance" means chemicals that are present in sufficient concentration to pose a hazard to human health. (Indiana Department of Health; 410 IAC 26-1-19; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 2. License Requirements

410 IAC 26-2-1 License

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 4-21.5-3-5; IC 16-21-2

Sec. 1. (a) A license must be obtained from the commissioner under IC 4-21.5-3-5 before:
(1) establishing;
(2) conducting;
(3) operating; or
(4) maintaining;

an abortion clinic. An abortion clinic may not operate without a license issued by the commissioner.
(b) A license to operate an abortion clinic:
(1) expires one (1) year after the date of issuance;
(2) is not assignable or transferable;
(3) is issued only for the premises named in the application; and
(4) is issued only for the scope of procedures to be performed as indicated by the applicant on the application.
(c) A license is valid for only one (1) location. Multiple clinics may not be operated under one (1) license.
(d) Upon closure of the clinic, the license shall be returned to the division. (Indiana Department of Health; 410 IAC 26-2-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164RFA)
410 IAC 26-2-2 Preoccupancy inspection
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. (a) The department will not issue a provisional license to operate an abortion clinic until the clinic has passed a preoccupancy inspection by the department.

(b) Once a new construction, addition, or renovation of an abortion clinic is complete, the abortion clinic must notify the department that the clinic is ready for occupancy. The department will then schedule and perform a preoccupancy inspection. The preoccupancy inspection is to determine compliance of the abortion clinic with 410 IAC 26-17-1 and 410 IAC 26-17-2. (Indiana Department of Health; 410 IAC 26-2-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-2-3 Application for initial license
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 3. (a) To obtain a license to operate an abortion clinic, an application for a license to operate an abortion clinic must be submitted to the division. At the latest, however, the application must be received by the department at least forty-five (45) days before the anticipated opening of the clinic. At least forty-five (45) days before the opening of the clinic, the applicant must inform the division of the anticipated date of opening.

(b) The initial license application includes the following:
(1) An application for a license to operate an abortion clinic on a form prescribed by the division to include the selection of only one (1) of the following procedure classifications:
   (A) Surgical abortions only. The clinic is precluded from performing drug induced abortions.
   (B) Both drug induced abortions and surgical abortions. The clinic must comply with this article and 410 IAC 26.5.

(2) Documents required by the application for a license to operate an abortion clinic.
(3) The appropriate license fee.
(c) The application for an abortion clinic license must require the applicant to do the following:
(1) Disclose whether the applicant, or an owner or affiliate of the applicant, operated an abortion clinic that was closed as a direct result of patient health and safety concerns.
(2) Disclose whether a principal or clinic staff member was convicted of a felony.
(3) Disclose whether a principal or clinic staff member was ever employed by a facility owned or operated by the applicant that closed as a result of administrative or legal action.
(d) All changes in ownership, name, and address must be reported in writing to the division. Reapplication must be filed when a change of fifty percent (50%) or greater ownership occurs. (Indiana Department of Health; 410 IAC 26-2-3; filed May 11, 2006, 9:36 a.m.: 29 IR 3356; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

410 IAC 26-2-4 Review and approval of initial license application
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2-11

Sec. 4. (a) Upon receipt of a completed application for an abortion clinic license, the department will review the application and accompanying documentation to determine that the applicant has met the requirements of IC 16-21-2-11(a)(1) and IC 16-21-2-11(a)(2).

(b) Upon determination by the commissioner that the applicant has failed to comply with this article, the commissioner may:
(1) request additional information concerning the application;
(2) conduct a further investigation to determine whether a provisional license should be granted; or
(3) deny the application.
(c) Upon determination by the commissioner that the applicant has complied with the provisional licensing requirements for an abortion clinic under this article, the commissioner will:
(1) provisionally approve the application for an abortion clinic license; and
(2) issue a provisional license to operate an abortion clinic.
The provisional license expires ninety (90) days after issue.
(d) After the opening of the clinic and before the expiration of the provisional license, the department will conduct a licensing survey to ensure that the clinic is operating in compliance with this article.
(e) If the clinic is found on the initial licensing survey to be in compliance with this article, the commissioner will issue a full license to operate an abortion clinic. If the clinic is not found to be in compliance with this article after the extended provisional period, the commissioner may:
(1) request additional information concerning the application;
(2) conduct a further investigation to determine whether a provisional license should be granted; or
(3) deny the application.
(Indiana Department of Health; 410 IAC 26-2-4; filed May 11, 2006, 9:36 a.m.: 29 IR 3357; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-2-5 Denial of license

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2; IC 27-13-1

Sec. 5. The commissioner may deny a license to operate an abortion clinic for any of the following reasons:
(1) If the licensee or licensees are not of reputable and responsible character.
(2) If the abortion clinic is not in compliance with the minimum standards for an abortion clinic adopted under this article.
(3) For violation of any of the provisions of IC 16-21 or this article.
(4) For permitting, aiding, or abetting the commission of any illegal act in the clinic.
(5) For knowingly collecting or attempting to collect from:
(A) a subscriber (as defined in IC 27-13-1-32); or
(B) an enrollee (as defined in IC 27-13-1-12);
of a health maintenance organization (as defined in IC 27-13-1-19) any amounts that are owed by the health maintenance organization.
(6) If conduct or practices of the clinic are found to be detrimental to the patients of the abortion clinic.
(7) If the application for a license to operate an abortion clinic or supporting documentation provided inaccurate statements or information.
(Indiana Department of Health; 410 IAC 26-2-5; filed May 11, 2006, 9:36 a.m.: 29 IR 3357; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-2-6 Renewal of license

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 6. (a) In order to renew its abortion clinic license, the clinic shall file an application for the renewal of an abortion clinic license with the division at least one (1) month before the expiration of the current license.
(b) The renewal application includes the following:
(1) An application for the renewal of a license to operate an abortion clinic on a form prescribed by the division to include the
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selection of only one (1) of the following procedure classifications:

(A) Surgical abortions only. The clinic is precluded from performing drug induced abortions.

(B) Both drug induced abortions and surgical abortions. The clinic must comply with this article and 410 IAC 26.5.

(2) Documents required by the application for the renewal of a license to operate an abortion clinic.

(3) The appropriate license fee.

(c) Upon determination by the commissioner that the applicant has met the licensing requirements for an abortion clinic, the commissioner shall approve the application for the renewal of a license to operate an abortion clinic and issue a license. (Indiana Department of Health; 410 IAC 26-2-6; filed May 11, 2006, 9:36 a.m.: 29 IR 3357; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

410 IAC 26-2-7 Posting of license
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 7. A license issued under this article must be conspicuously posted on the premises in an area open to patients. (Indiana Department of Health; 410 IAC 26-2-7; filed May 11, 2006, 9:36 a.m.: 29 IR 3358; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-2-8 Enforcement actions
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-2; IC 16-21-3; IC 27-13-1

Sec. 8. (a) The commissioner may take any of the following actions on any of the grounds listed in subsection (b):
(1) Issue a letter of correction.
(2) Issue a probationary license.
(3) Conduct a resurvey.
(4) Deny the renewal of a license.
(5) Revoke a license.
(6) Impose a civil penalty in an amount not to exceed ten thousand dollars ($10,000).
(b) The commissioner may take action under subsection (a) on any of the following grounds:
(1) Violation of any provision of state law.
(2) Permitting, aiding, or abetting the commission of any illegal act in an abortion clinic.
(3) Knowingly collecting or attempting to collect from:
    (A) a subscriber (as defined in IC 27-13-1-32); or
    (B) an enrollee (as defined in IC 27-13-1-12);
    of a health maintenance organization (as defined in IC 27-13-1-19) any amounts that are owed by the health maintenance organization.
(4) Conduct or practice found by the department to be detrimental to the welfare of the patients of an abortion clinic.
(Indiana Department of Health; 410 IAC 26-2-8; filed May 11, 2006, 9:36 a.m.: 29 IR 3358; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA; filed Apr 26, 2021, 12:32 p.m.: 20210519-IR-410200604FRA)

410 IAC 26-2-9 Probationary license
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 4-21.5; IC 16-21

Sec. 9. A probationary license may be:
(1) issued for a period of three (3) months; and
(2) reissued;
but no more than three (3) probationary licenses may be issued during a twelve (12) month period. The issuance of a probationary license results in the automatic expiration of any other license held under this article. (Indiana Department of Health; 410 IAC 26-2-9; filed May 11, 2006, 9:36 a.m.: 29 IR 3358; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 3. Surveys

410 IAC 26-3-1 Survey procedures

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2-6

Sec. 1. (a) The abortion clinic shall fully cooperate with surveys conducted by representatives of the department. Upon arrival of department surveyors at the clinic, the clinic may immediately contact the department to confirm the identity of the surveyors. Upon confirmation by the department of the survey and surveyors, the clinic shall:

1. immediately admit the surveyors to the clinic; and
2. not delay the survey.

(b) Documents, registers, reports, records, and minutes of the abortion clinic must be made available to the department upon request for inspection and copying.

(c) Before any information is copied for use by the department, the abortion clinic shall redact all information that identifies or could be used to identify a patient or staff member. Department surveyors may review the unredacted original in the abortion clinic.

(d) Documents, registers, reports, records, and minutes required to be maintained by the abortion clinic include, but are not limited to, the following:

1. Documents showing ownership and a copy of articles of incorporation (if incorporated).
2. All documents pertaining to quality assurance and improvement of patient care and medical care.
3. Personnel records.
4. Medical records relating to surgical abortions.
5. Reports under IC 16-21-2-6.
6. Policies and procedures of the abortion clinic.

(e) If the governing body of the clinic is an individual responsible for governing the abortion clinic, the clinic is not required to prepare and maintain the documents referenced in this subsection. If the governing body is not an individual with sole authority and responsibility for the clinic, the clinic must prepare and maintain the following documents, registers, reports, records, and minutes to include, but not be limited to:

1. The constitution and bylaws of the governing body.
2. Minutes of meetings of the governing body and committees thereof.
3. Documents, registers, reports, records, and minutes must be complete and up-to-date. (Indiana Department of Health; 410 IAC 26-3-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3358; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-3-2 Licensing surveys

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. (a) The department will conduct a licensing survey of each abortion clinic at least one (1) time per calendar year. The licensing survey is conducted to ensure that the abortion clinic is operating in compliance with state law.

(b) Licensing surveys will be conducted during normal business hours of the abortion clinic unless the abortion clinic requests that the survey be conducted during nonbusiness hours.

(c) The division will notify the clinic of the results of the licensing survey in writing. (Indiana Department of Health; 410 IAC 26-3-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3359; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA; filed Apr 26, 2021, 12:32 p.m.: 20210519-IR-410200604FRA)
410 IAC 26-3-3 Complaint surveys
  Authority:  IC 16-21-1-7; IC 16-21-2-2.5
  Affected:  IC 16-21-1; IC 16-21-2

Sec. 3. (a) In accordance with division policy, the division shall investigate credible complaints received by the division that allege noncompliance with this article.
  (b) Complaints will be assigned a priority for investigation in accordance with division policy.
  (c) A licensing survey may be conducted simultaneously with and in addition to a complaint survey.
  (d) The division shall notify the abortion clinic of the results of the complaint survey in writing. (Indiana Department of Health; 410 IAC 26-3-3; filed May 11, 2006, 9:36 a.m.: 29 IR 3359; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-3-4 Plan of correction
  Authority:  IC 16-21-1-7; IC 16-21-2-2.5
  Affected:  IC 16-21-1; IC 16-21-2

Sec. 4. (a) The abortion clinic must file an acceptable plan of correction with the division within ten (10) days of receipt of a survey report from the division that documents noncompliance with state law.
  (b) Unless the commissioner determines that there is a need for immediate release, the abortion clinic will have ten (10) days after notification of a noncompliance to submit to the division an acceptable plan of correction before the survey report is made available to the public.
  (c) The plan of correction shall contain, for each deficient practice cited on the survey report, at least the following:
    1) How the deficient practice will be corrected.
    2) How the deficient practice will be prevented from reoccurrence.
    3) Who will be responsible for correction and prevention.
    4) The month, day, and year that the corrective action will be completed, not to exceed thirty (30) days from receipt of the notice of noncompliance.
    5) If the nature of the corrective action requires more than thirty (30) days from the date of receipt of the notice of noncompliance, the clinic shall submit justification and a completion date to the division.
  (d) If the division determines all or part of the submitted plan is unacceptable, the clinic shall submit a revised plan of correction within five (5) days of receipt of the notice identifying the unacceptable plan or part thereof.
  (e) Failure to submit any required plan of correction or failure to implement a corrective action by the completion date may result in an enforcement action under 410 IAC 26-2-8. (Indiana Department of Health; 410 IAC 26-3-4; filed May 11, 2006, 9:36 a.m.: 29 IR 3359; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA; filed Apr 26, 2021, 12:32 p.m.: 20210519-IR-410200604FRA)

410 IAC 26-3-5 Confidentiality
  Authority:  IC 16-21-1-7; IC 16-21-2-2.5
  Affected:  IC 16-21-1; IC 16-21-2

Sec. 5. The department shall maintain the confidentiality of patient identities, patient information, and patient records. (Indiana Department of Health; 410 IAC 26-3-5; filed May 11, 2006, 9:36 a.m.: 29 IR 3359; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 4. Governing Body

410 IAC 26-4-1 Powers and duties
  Authority:  IC 16-21-1-7; IC 16-21-2-2.5
  Affected:  IC 16-21-1; IC 16-21-2; IC 16-34; IC 25-22.5
Sec. 1. (a) The governing body:
(1) shall function as the ultimate authority; and
(2) is responsible for the conduct and management;
of the abortion clinic.

(b) If the governing body is an individual who has sole authority and responsibility for the clinic, that individual may also serve
as the clinic administrator or medical director, or both, if qualified. A clinic administrator appointed by the governing body may also
serve as the medical director if qualified.

(c) The governing body shall do the following:
(1) Assume responsibility for:
   (A) determining;
   (B) implementing; and
   (C) monitoring;
policies governing the clinic's operation.

(2) Ensure that:
   (A) clinic policies are followed so as to provide quality health care in a safe environment; and
   (B) the clinic complies with:
      (i) this article;
      (ii) IC 16-21; and
      (iii) IC 16-34.

(3) Review, at least every six (6) months, reports of management operations, including, but not limited to, the following:
   (A) Quality assessment and improvement program.
   (B) Patient services provided.
   (C) Results attained.
   (D) Recommendations made.
   (E) Actions taken.
   (F) Follow-up.

(4) Maintain documents, registers, and reports that show the following:
   (A) Ownership.
   (B) Compliance with local, state, and federal laws and regulations.
   (C) Adherence to clinic bylaws (if applicable) and clinic policies.

(5) Approve all appointments to or contracts with medical staff.

(6) Ensure the following:
   (A) Maintenance of the physical plant.
   (B) That the clinic is:
      (i) equipped; and
      (ii) staffed;
to meet the needs of the patients.

(7) Ensure that clinic policies and procedures are:
   (A) updated as needed; and
   (B) reviewed at least triennially.

(8) Establish the following:
   (A) A policy and procedure for communication with physicians concerning a patient emergency.
   (B) A process for the following:
      (i) Reporting licensed health professionals who fail to comply with state professional licensing requirements as
          found in IC 25-22.5.
      (ii) Documenting actions against licensed health professionals who fail to comply with the clinic policies and
          procedures.
      (iii) Reporting information that statute requires the abortion clinic to report to a state agency or law enforcement
          agency.

(d) If the governing body is not an individual responsible for the governing of the clinic, the governing body must do the
following:

1. Adopt bylaws and operate in compliance with the bylaws.
2. Review the bylaws at least triennially.
3. If the governing body is not an individual who is also serving as the medical director, the governing body shall do the following:
   (1) Designate a medical director who has the responsibility for the direction of:
       (A) medical;
       (B) nursing; and
       (C) health-related;
   services to patients.
4. Maintain a liaison with the medical director.
5. If the governing body is not an individual who is also serving as the clinic administrator, the governing body shall do the following:
   (1) Designate a clinic administrator who has the responsibility and authority to carry out the day-to-day operation of the clinic.
   (2) Develop criteria, which include, but are not limited to, defining educational and experience requirements for the clinic administrator.
   (3) Delineate in writing the responsibility and authority of the clinic administrator.
   (4) Require the following:
       (A) That the clinic administrator or a designee:
           (i) attend meetings of the governing body and its committees; and
           (ii) act as its representative at medical staff meetings.
       (B) That the clinic administrator:
           (i) designate in writing an administrative officer to serve during his or her absence; and
           (ii) participate in the development and implementation of appropriate policies and programs.

Indiana Department of Health; 410 IAC 26-4-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3359; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-4-2 Appointment and conduct of medical staff

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. (a) The medical staff of the clinic:
1. consists of physicians appointed to or contracted with to provide medical services at the clinic; and
2. must be composed of at least one (1) physician.
(b) The medical director must be a physician licensed to practice in the state of Indiana.
(c) If the medical staff consists of more than one (1) physician:
   (1) the medical director shall serve as coordinator of the medical staff; and
   (2) a current roster of members of the medical staff shall be maintained.
(d) In appointing or contracting with medical staff, the governing body shall do the following:
   (1) Ensure that appointments to or contracts with medical staff are acted upon with the advice and recommendation of the medical director.
   (2) Examine credentials of candidates for appointment, reappointment, or contracting to the medical staff in accordance with the following:
       (A) Clinic policy.
       (B) Applicable state and federal law.
   (3) Ensure that criteria for selection for medical staff include the following:
       (A) Individual character.
       (B) Competence.
       (C) Education.
       (D) Training.
(E) Experience.
(F) Judgment.

(4) Maintain a reasonably accessible hard copy or electronic file for each member of the medical staff, which includes, but is not limited to, the following:
   (A) A completed, signed application.
   (B) The date and year of completion of all Accreditation Council for Graduate Medical Education (ACGME) accredited residency training programs, if applicable.
   (C) A current copy of the individual's credentials as follows:
      (i) An Indiana license showing date of licensure and number or available data provided by the Indiana professional licensing agency. A copy of practice restrictions, if any, must be attached to the license issued by the Indiana professional licensing agency through the appropriate licensing board.
      (ii) Indiana controlled substance registration showing number as applicable.
      (iii) Drug Enforcement Agency registration showing number as applicable.
      (iv) Documentation of experience in the practice of medicine.
      (v) Documentation of specialty board certification as applicable.

(e) The governing body is responsible for the conduct of the medical staff activities related to the abortion clinic. The governing body shall ensure the following:
   (1) That the medical staff is accountable and responsible to the governing body for the quality of care provided to patients.
   (2) That procedures are performed only by a physician approved by the governing body to perform such procedures.
   (3) That procedures performed in the clinic are limited to procedures authorized by the governing body.

(f) The governing body is responsible for assuring that quality patient care is provided. In accordance with clinic policy, the governing body shall ensure that a qualified licensed physician who is a member of the medical staff is responsible for the care and treatment of each patient.

(g) The governing body is responsible for services delivered in the clinic by contractors for medical services. The governing body shall ensure the following:
   (1) That a contractor of any service furnishes those services in such a manner as to permit the clinic to comply with all applicable statutes and rules.
   (2) That the services performed under a contract are:
      (A) provided in a safe and effective manner; and
      (B) included in the clinic's quality assessment and improvement program.
   (3) That the clinic maintains a list of all contracted services, including the scope and nature of the services provided.

(Indiana Department of Health; 410 IAC 26-4-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3360; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 5. Administration and Policies

410 IAC 26-5-1 Administration

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. The clinic administrator is responsible for day-to-day operations of the abortion clinic to include, but not be limited to, the following functions:
   (1) Employing qualified staff:
      (A) commensurate with assigned duties and responsibilities; and
      (B) in accordance with the employee's:
         (i) licensure;
         (ii) certification;
         (iii) experience; and
         (iv) competence.
   (2) Ensuring that sufficient staff is present to provide quality patient care.
(3) Implementation of internal and external disaster and emergency preparedness plans with documentation of outcome.

(Indiana Department of Health; 410 IAC 26-5-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3361; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-5-2 Required policies and procedures

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. (a) The clinic shall develop, implement, and maintain the following:

(1) Written medical staff policies.

(2) Written procedures for the following:

(A) Emergencies.

(B) Initial treatment.

(C) Transfer.

(b) The clinic shall provide immediate lifesaving measures, within the scope of service available, to all persons in the clinic, to include, but not be limited to, the following:

(1) Timely assessment.

(2) Basic life support.

(3) Appropriate transfer.

(c) The clinic shall develop, implement, and maintain the following:

(1) Policies that cover health care worker practice problems, including, but not limited to, the following:

(A) Impaired health care workers.

(B) Criminal history.

(C) Disciplinary action.

(2) A written policy to address the internal review of unusual occurrences and disasters. This policy must include, but not be limited to, the following:

(A) Patient injuries or marked deterioration of patient condition occurring under unanticipated or unexpected circumstances.

(B) Unexplained loss of or theft of a controlled substance.

(C) Deaths occurring within the clinic.

(Indiana Department of Health; 410 IAC 26-5-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3361; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 6. Quality Assessment and Improvement

410 IAC 26-6-1 Quality assessment and improvement

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) The abortion clinic must develop or adopt, implement, and maintain an effective, organized, clinic-wide, comprehensive quality assessment and improvement program in which all areas of the clinic involved in the provision of surgical abortion participate. The program shall be ongoing and have a written plan of implementation that evaluates, but is not limited to, the following:

(1) All services, including services furnished by a contractor.

(2) All functions, including, but not limited to, the following:

(A) Discharge.

(B) Transfer.

(C) Infection control.

(D) Response to patient emergencies.

(3) All services performed in the clinic with regard to the following:
(A) Appropriateness of diagnoses and treatments related to a standard of care.
(B) Anticipated or expected outcomes.
(4) Medical and medication errors.
(b) The clinic shall take appropriate action to address the opportunities for improvement found through the quality assessment and improvement program as follows:
(1) The action must be documented.
(2) The outcome of the action must be documented as to its effectiveness, continued follow-up, and impact on patient care.

(Indiana Department of Health; 410 IAC 26-6-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3362; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-6-2 Reportable events

Authority: IC 16-19-3-4; IC 16-21-1-7
Affected: IC 16-19-3; IC 16-21-1; IC 25

Sec. 2. (a) The clinic's quality assessment and improvement program under section 1 of this rule shall include the following:
(1) A process for determining the occurrence of the following reportable events within the clinic:
(A) The following surgical events:
   (i) Surgery performed on the wrong body part, defined as any surgery performed on a body part that is not consistent with the documented informed consent for that patient. Excluded are emergent situations:
       (AA) that occur in the course of surgery; or
       (BB) whose exigency precludes obtaining informed consent;
   or both.
   (ii) Surgery performed on the wrong patient, defined as any surgery on a patient that is not consistent with the documented informed consent for that patient.
   (iii) Wrong surgical procedure performed on a patient, defined as any procedure performed on a patient that is not consistent with the documented informed consent for that patient. Excluded are emergent situations:
       (AA) that occur in the course of surgery; or
       (BB) whose exigency precludes obtaining informed consent;
   or both.
   (iv) Retention of a foreign object in a patient after surgery or other invasive procedure. The following are excluded:
       (AA) Objects intentionally implanted as part of a planned intervention.
       (BB) Objects present before surgery that were intentionally retained.
       (CC) Objects not present prior to surgery that are intentionally left in when the risk of removal exceeds the risk of retention, such as microneedles or broken screws.
   (v) Intraoperative or immediately postoperative death in an ASA Class I patient. Included are all ASA Class I patient deaths in situations where anesthesia was administered; the planned surgical procedure may or may not have been carried out.
(B) The following product or device events:
   (i) Patient death or serious disability associated with the use of contaminated drugs, devices, or biologics provided by the clinic. Included are generally detectable contaminants in drugs, devices, or biologics regardless of the source of contamination or product.
   (ii) Patient death or serious disability associated with the use or function of a device in patient care in which the device is used or functions other than as intended. Included are, but not limited to, the following:
       (AA) Catheters.
       (BB) Drains and other specialized tubes.
       (CC) Infusion pumps.
       (DD) Ventilators.
   (iii) Patient death or serious disability associated with intravascular air embolism that occurs while being cared for in the clinic. Excluded are deaths or serious disability associated with neurosurgical procedures known to
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present a high risk of intravascular air embolism.

(C) The following patient protection events:
  (i) Infants discharged to the wrong person.
  (ii) Patient death or serious disability associated with patient elopement.
  (iii) Patient suicide or attempted suicide resulting in serious disability, while being cared for in the clinic, defined as events that result from patient actions after admission to the clinic. Excluded are deaths resulting from self-inflicted injuries that were the reason for admission to the clinic.

(D) The following care management events:
  (i) Patient death or serious disability associated with a medication error, for example, errors involving the wrong:
      (AA) drug;
      (BB) dose;
      (CC) patient;
      (DD) time;
      (EE) rate;
      (FF) preparation; or
      (GG) route of administration.
  Excluded are reasonable differences in clinical judgment on drug selection and dose. Includes administration of a medication to which a patient has a known allergy and drug-drug interactions for which there is known potential for death or serious disability.
  (ii) Patient death or serious disability associated with a hemolytic reaction due to the administration of ABO/HLA incompatible blood or blood products.
  (iii) Maternal death or serious disability associated with labor or delivery in a low-risk pregnancy while being cared for in the clinic. Included are events that occur within forty-two (42) days postdelivery. Excluded are deaths from any of the following:
      (AA) Pulmonary or amniotic fluid embolism.
      (BB) Acute fatty liver of pregnancy.
      (CC) Cardiomyopathy.
  (iv) Patient death or serious disability associated with hypoglycemia, the onset of which occurs while the patient is being cared for in the clinic.
  (v) Death or serious disability (kernicterus) associated with the failure to identify and treat hyperbilirubinemia in neonates.
  (vi) Stage 3 or 4 pressure ulcers acquired after admission to the clinic. Excluded is progression from Stage 2 or Stage 3 if the Stage 2 or Stage 3 pressure ulcer was recognized upon admission or unstageable because of the presence of eschar.
  (vii) Patient death or serious disability resulting from joint movement therapy performed in the clinic.
  (viii) Artificial insemination with the wrong donor sperm or wrong egg.

(E) The following environmental events:
  (i) Patient death or serious disability associated with an electric shock while being cared for in the clinic. Excluded are events involving planned treatment, such as electrical countershock or elective cardioversion.
  (ii) Any incident in which a line designated for oxygen or other gas to be delivered to a patient:
      (AA) contains the wrong gas; or
      (BB) is contaminated by toxic substances.
  (iii) Patient death or serious disability associated with a burn incurred from any source while being cared for in the clinic.
  (iv) Patient death or serious disability associated with a fall while being cared for in the clinic.
  (v) Patient death or serious disability associated with the use of restraints or bed rails while being cared for in the clinic.

(F) The following criminal events:
  (i) Any instance of care ordered by or provided by someone impersonating a physician, nurse, pharmacist, or other licensed health care provider.
(ii) Abduction of a patient of any age.
(iii) Sexual assault on a patient within or on the grounds of the clinic.
(iv) Death or significant injury of a patient or staff member resulting from a physical assault, that is, battery, that occurs within or on the grounds of the clinic.

(2) A process for reporting to the department each reportable event listed in subdivision (1) that is determined by the clinic's quality assessment and improvement program to have occurred within the clinic.

(b) The process for determining the occurrence of the reportable events listed in subsection (a)(1) by the clinic's quality assessment and improvement program shall be designed by the clinic to accurately determine the occurrence of any of the reportable events listed in subsection (a)(1) within the clinic in a timely manner.

(c) The process for reporting the occurrence of a reportable event listed in subsection (a)(1) shall comply with the following:

(1) The report shall:
   (A) be made to the department;
   (B) be submitted not later than fifteen (15) working days after the reportable event is determined to have occurred by the clinic's quality assessment and improvement program;
   (C) be submitted not later than four (4) months after the potential reportable event is brought to the program's attention; and
   (D) identify the reportable event, the quarter of occurrence, and the clinic, but shall not include any identifying information for any:
      (i) patient;
      (ii) individual licensed under IC 25; or
      (iii) clinic employee involved;
      or any other information.

(2) A potential reportable event may be identified by a clinic that:
   (A) receives a patient as a transfer; or
   (B) admits a patient subsequent to discharge;
   from another health care facility subject to a reportable event requirement. In the event that a clinic identifies a potential reportable event originating from another health care facility subject to a reportable event requirement, the identifying clinic shall notify the originating health care facility as soon as they determine an event has potentially occurred for consideration by the originating health care facility's quality assessment and improvement program.

(3) The report, and any documents permitted under this section to accompany the report, shall be submitted in an electronic format, including a format for electronically affixed signatures.

(4) A quality assessment and improvement program may refrain from making a determination about the occurrence of a reportable event that involves a possible criminal act until criminal charges are filed in the applicable court of law.

(d) The clinic's report of a reportable event listed in subsection (a)(1) shall be used by the department for purposes of publicly reporting the type and number of reportable events occurring within each clinic. The department's public report will be issued annually. (Indiana Department of Health; 410 IAC 26-6-2; filed Nov 21, 2006, 7:20 a.m.: 20061220-IR-410050321FRA; filed Oct 7, 2008, 10:26 a.m.: 20081105-IR-410080061FRA; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)

Rule 7. Medical Records

410 IAC 26-7-1 Medical records, storage, and administration

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) The abortion clinic must do the following:

(1) Create and maintain a medical record on each surgical abortion patient.
(2) Have a written policy that ensures responsibility for and maintenance of surgical abortion records as follows:
   (A) The clinic must establish and implement the following:
      (i) Policies and procedures to assure that the care and services provided to each patient are appropriately
documented.

(ii) A system to assure that medical records are readily available in accordance with clinic policy and systematically organized to facilitate the compilation and retrieval of information.

(B) The policy must provide safeguards to assure protection of the medical records from the following:

(i) Fire.
(ii) Water.
(iii) Other sources of damage.

(C) All original medical records or legally reproduced medical records must be maintained by the clinic for a period of at least seven (7) years or the applicable statute of limitation, whichever is longer. Original medical records must be maintained in the clinic for at least two (2) years. Records over two (2) years old may be kept off-site but must be retrievable within forty-eight (48) business hours.

(b) A medical record must be maintained with documentation of service rendered for each surgical abortion patient of the clinic as follows:

(1) Medical records:
   (A) are documented accurately and in a timely manner;
   (B) are readily accessible; and
   (C) permit prompt retrieval of information.

(2) A unit record system of filing should be utilized. When this is not practicable, a system must be established by the clinic to retrieve, when necessary, all divergently located record components.

(3) The clinic shall use a system of author identification and record maintenance that:
   (A) ensures the integrity of the authentication; and
   (B) protects the security of all record entries.

Each entry must be authenticated in accordance with the clinic and medical staff policies.

(4) Medical records must be retained in their original or legally reproduced form as required by federal or state law.

(5) Plain paper facsimile orders, reports, and documents are acceptable for inclusion in the medical record if allowed by the clinic policies.

(6) The clinic shall have a system of coding and indexing medical records that allows for timely retrieval of records in order to support continuous quality assessment and improvement activities.

(7) The clinic shall ensure the confidentiality of patient records. The clinic must develop, implement, and maintain the following:

   (A) A procedure for releasing information or copies of records only to authorized individuals in accordance with federal and state laws.
   
   (B) A procedure that ensures that unauthorized individuals cannot gain access to medical records.

(c) A written or electronic register must be kept of all patients treated that provides the following:

(1) Identification data.
(2) Treatment rendered.
(3) Attending physician.
(4) Condition on discharge.
(5) Transfers to hospital facility.

(6) Other data deemed necessary by the clinic.

(Indiana Department of Health; 410 IAC 26-7-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3362; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-7-2 Content of the medical record

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2; IC 16-34-2-1.1

Sec. 2. (a) The medical record for surgical abortions must be accurate and contain sufficient information to do the following:

(1) Identify the patient to include name, age, and address.

(2) Document the following:
(A) Tests, examinations, and procedures performed.
(B) The course of the patient's stay in the clinic and the results.
(C) Evidence that the patient was provided the hotline telephone number for assistance to patients who are:
   (i) coerced into an abortion; or
   (ii) victims of sex trafficking.
(D) Evidence of appropriate informed consent for procedures and treatments as required by IC 16-34-2-1.1.
(E) Any report filed with a state agency or law enforcement agency pursuant to a statutory reporting requirement.

(b) Entries in the medical record must be as follows:
(1) Legible.
(2) Complete.
(3) Made by authorized individuals as specified in clinic and medical staff policies.
(4) Authenticated and dated in accordance with this article.
(c) Patient records for surgical abortions must document and contain, at a minimum, the following:
(1) Appropriate medical history.
(2) Results of the following:
   (A) A physical examination.
   (B) Diagnostic or laboratory studies, or both (if performed).
(3) Any allergies and abnormal drug reactions.
(4) Entries related to anesthesia administration.
(5) A report describing techniques, findings, and tissue removed or altered.
(6) Authentication of entries by the physician or physicians and health care workers who treated or cared for the patient.
(7) Condition on discharge, disposition of the patient, and time of discharge.
(8) Discharge entry to include instructions to the patient or patient's legal representative.
(9) A copy of the following:
   (A) The transfer form if the patient was referred to a hospital or other facility.
   (B) The terminated pregnancy report filed with the department.
   (C) Any document signed by the patient.
(d) An appropriate history and physical examination report must be in the patient's chart before a surgical abortion. The report shall include, but is not limited to, the following:
(1) Vital signs.
(2) Allergies.
(3) Any significant risk factors.
(4) The date written.

Rule 8. Personnel

410 IAC 26-8-1 Personnel policies and records
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) The abortion clinic shall maintain current and accurate personnel records for all employees. Personnel records shall:
(1) be maintained for each employee of the clinic; and
(2) include personal data to include:
   (A) education;
   (B) experience;
   (C) date of employment;
   (D) a copy of current license when required;
(E) evidence of participation in job-related educational and training activities; and
(F) health records of employees that relate to post offer and subsequent:
   (i) physical examinations;
   (ii) tests; and
   (iii) immunizations.

(b) If the clinical administrator is not the governing body, the clinic must establish employment criteria for the clinic administrator to include, but not be limited to, the following:
   (1) Educational requirements.
   (2) Experience requirements.
   (3) Professional certification, licensing, or registration requirements where appropriate.
(c) The clinic must do the following:
   (1) Maintain current job descriptions with reporting responsibilities for all personnel and annual performance evaluations, based on the job description, for each employee and contract and agency personnel.
   (2) Ensure that all health care workers, including contract and agency personnel, for whom a license, registration, or certification is required:
      (A) maintain current license, registration, or certification; and
      (B) keep documentation of same.

(Indiana Department of Health; 410 IAC 26-8-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3363; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-8-2 Employee health monitoring
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. The clinic shall do the following:
   (1) Develop, implement, and maintain a written policy for the control of communicable disease in compliance with applicable federal and state laws.
   (2) Monitor employee health in accordance with the clinic's infection control program.
   (3) Ensure that all employees, staff members, and contractors having direct patient contact are evaluated at least annually for tuberculosis as follows:
      (A) Any person with a negative history of tuberculosis or a negative test result must have a baseline two-step tuberculin skin test using the Mantoux method or a quantiferon-TB assay unless the individual has documentation that a tuberculin skin test has been applied at any time during the previous twelve (12) months and the result was negative.
      (B) The second step of a two-step tuberculin skin test using the Mantoux method must be administered one (1) to three (3) weeks after the first tuberculin skin test was administered.
      (C) Any person with a documented history of tuberculosis, documented previously positive test result for tuberculosis, documented completion of treatment for tuberculosis, or newly positive results to the tuberculin skin test must have one (1) chest radiograph to exclude a diagnosis of tuberculosis.
      (D) After baseline testing, tuberculosis screening must be completed annually and must include at a minimum a tuberculin skin test using the Mantoux method or a quantiferon-TB assay unless the individual was subject to clause (C).
      (E) Any person having a positive finding on a tuberculosis evaluation may not work in the abortion clinic or provide direct patient contact unless approved by a physician to work.
      (F) The abortion clinic must maintain documentation of tuberculosis evaluations showing that any person working for the abortion clinic or having direct patient contact has had a negative finding on a tuberculosis examination within the previous twelve (12) months.

(Indiana Department of Health; 410 IAC 26-8-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3364; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)
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410 IAC 26-8-3 Orientation and training requirements
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 3. (a) The clinic must do the following:
(1) Develop, implement, and maintain a policy and procedure for the orientation of new employees, contractors, and agency personnel providing direct care and services to patients.
(2) Orientate all new employees, including contract and agency personnel, to applicable clinic and personnel policies.
(b) The clinic shall ensure cardiopulmonary resuscitation (CPR) competence in accordance with current standards of practice and clinic policy for all health care workers including contract and agency personnel who provide direct patient care.
(c) The clinic shall ensure all employees receive annual training by law enforcement officers on identifying and assisting women who are:
(1) coerced into an abortion; or
(2) victims of sex trafficking.
(d) The clinic shall document in each employee's personnel file evidence of annual training provided by a law enforcement officer on identifying and assisting women who are:
(1) coerced into an abortion; or
(2) victims of sex trafficking.

Rule 9. Medical Staff

410 IAC 26-9-1 Medical staff services
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) The medical staff of the clinic is:
(1) accountable to the governing body of the clinic; and
(2) responsible to the governing board for the quality of medical care and services provided to patients.
(b) The medical director must do the following:
(1) Examine credentials of candidates for appointment, reappointment, or contracting to the medical staff.
(2) Make recommendations to the governing body on the appointment or reappointment of medical staff.
(c) The medical director must develop and maintain policies and procedures for the provision of medical services. The policies must provide for and the medical staff must ensure the following:
(1) An appropriate and timely medical history and physical examination is performed.
(2) All physician orders:
   (A) are in writing or acceptable computerized form;
   (B) must be authenticated by a responsible physician as allowed by clinic policies not to exceed thirty (30) days.
   (3) There is a provision for personnel authorized to take a verbal order.

Rule 10. Patient Care and Nursing Services

410 IAC 26-10-1 Patient care
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2
Sec. 1. (a) All patient care services must:
(1) meet the needs of the patient, within the scope of the service offered, in accordance with acceptable standards of practice;
(2) be under the direction of a qualified person or persons; and
(3) require that:
   (A) patient care services rendered are:
      (i) reviewed and analyzed at regular meetings of patient care personnel; and
      (ii) used as a basis for evaluating the quality of services provided; and
   (B) personnel with appropriate training are available at all times to handle possible emergencies involving patients of the clinic.

(b) Written patient care policies and procedures must be available to personnel and must include, but not be limited to, the following:
(1) A provision that a reliable method of patient identification must be used.
(2) A provision for instruction or instructions to be given to the patient or the patient's legal representative regarding follow-up care and transportation needed by the patient on discharge following a surgical abortion to include at least the following:
   (A) Signs and symptoms of possible complications.
   (B) Activities allowed and to be avoided.
   (C) Hygienic and other postdischarge procedures to be followed.
   (D) Clinic emergency phone numbers available on a twenty-four (24) hour basis for the patient to contact the clinic for a complication and be triaged for care and either:
      (i) be seen the same day by a practitioner; or
      (ii) be referred to an appropriate site of care.
   (E) Follow-up appointment, if indicated.
   (F) Counseling regarding Rh typing.
   (G) Administration of Rh immune globulin, if indicated, unless:
      (i) the patient signs a waiver refusing the administration; or
      (ii) other arrangements for administration are documented.
(3) A provision to maintain a written system of documentation of patients who report post-procedure complications and the clinic’s interventions. The interventions must be documented in the medical record.
(4) A provision that facilities, reusable equipment, and supplies must be thoroughly cleaned or sterilized following use according to clinic policies and procedures.
(5) A provision that all patients must be observed during the recovery period by qualified personnel.

410 IAC 26-10-2 Nursing services
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. If the clinic employs licensed nurses, the clinic must ensure the following:
(1) Registered nurses and licensed practical nurses are currently licensed in Indiana.
(2) Nursing personnel meet annual inservice requirements as established by clinic and federal and state requirements.

Rule 11. Infection Control Program

410 IAC 26-11-1 Infection control administration
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2
Sec. 1. (a) The clinic must do the following:
(1) Provide a safe and healthful environment that minimizes infection exposure and risk to the following:
   (A) Patients.
   (B) Health care workers.
   (C) Persons who accompany patients.
(2) Maintain a written infection control policy that provides for an active and effective clinic-wide infection control program.
The policy must include a system designed for the:
   (A) identification;
   (B) surveillance;
   (C) investigation;
   (D) control; and
   (E) prevention;
of infections and communicable diseases in patients and health care workers.
(b) The infection control program must identify and evaluate trends or clusters of clinic generated infections or communicable diseases.
(c) The clinic must designate a person qualified by training or experience as responsible for the following:
(1) Ongoing infection control activities.
(2) The development and implementation of policies governing control of infections and communicable diseases.
(d) The clinic administrator must do the following:
(1) Be responsible for the implementation of successful corrective action plans in affected problem areas and ensure that infection control policies are followed.
(2) Provide for appropriate infection control input into plans for renovation and new construction to ensure awareness of federal, state, and local rules that affect infection control practices as well as plan for appropriate protection of patients and employees during construction or renovation.
(e) The clinic must establish a committee to monitor and guide the infection control program in the clinic as follows:
(1) The infection control committee must meet at least quarterly. Membership must include, but is not limited to, the following:
   (A) The person directly responsible for management of the infection surveillance, prevention, and control program as established in subsection (c).
   (B) The medical director.
   (C) A representative from the nursing staff (if the clinic employs a licensed nurse).
   (D) Representatives from other appropriate services within the clinic as needed.
(2) The infection control committee responsibilities must include, but are not limited to, the following:
   (A) Establishing techniques and systems for:
      (i) identifying;
      (ii) reviewing; and
      (iii) reporting;
   infections in the clinic.
   (B) Recommending corrective action plans, reviewing outcomes, and assuring resolution of identified problems.
   (C) Reviewing employee exposure incidents and making appropriate recommendations to minimize risk.
   (D) Written reports of quarterly meetings.
   (E) Reviewing and recommending changes in procedures, policies, and programs that are pertinent to infection control.
   These include, but are not limited to, the following:
      (i) Sanitation, including proper disposal of removed tissue.
      (ii) Universal precautions, including infectious waste management.
      (iii) Cleaning, disinfection, and sterilization.
      (iv) Aseptic technique, invasive procedures, and equipment usage.
      (v) Reuse of disposables.
      (vi) A system for handling patients with communicable diseases.
      (vii) A system, which complies with state and federal law, to monitor the immune status of health care workers exposed to communicable diseases.
(viii) An employee health program to determine the communicable disease history of new personnel as well as an ongoing program for current personnel as required by state and federal agencies.
(ix) Requirements for personal hygiene and attire that meet acceptable standards of practice.
(x) A program of linen management.

(Indiana Department of Health; 410 IAC 26-11-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3365; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-11-2 Sterilization

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. (a) Sterilization of equipment and supplies must be provided, within the scope of the service offered, in accordance with acceptable standards of practice or manufacturer's recommendations and applicable state laws and rules (to include 410 IAC 1-4, Universal Precautions). Sterilization services must be directed by a qualified person or persons and must provide for the following:

(1) Biological indicators must be used to check sterilization processes at least monthly. Chemical sterilizing indicators must be used to check the sterilizing process of individual packs.

(2) Written policies and procedures must be available and followed by personnel responsible for sterilizing equipment and supplies, including, but not limited to, the following:

(A) Minimum time and temperature for processing various size bundles and packs.

(B) Instructions for:

(i) loading;
(ii) operating;
(iii) cleaning; and
(iv) maintaining;
sterilizers.

(C) Instructions for:

(i) cleaning;
(ii) packaging;
(iii) storing;
(iv) labeling; and
(v) dispensing of;
sterile supplies.

(D) The procedure for maintaining and recording the particular sterilizing cycle.

(E) Sterilization of heat labile reusable equipment.

(3) Records of results must be maintained and evaluated periodically to include, but not be limited to, the following:

(A) Records of recording thermometers or a daily record of the sterilizing cycle:

(i) date;
(ii) time;
(iii) temperature;
(iv) pressure; and
(v) contents;
for each sterilizer load.

(B) Results of biological indicators used in testing the sterilizing processes.

(b) Environmental surfaces and equipment not requiring sterilization that have been contaminated by blood or other potentially infectious materials must be cleaned then decontaminated in accordance with acceptable standards of practice and applicable state laws and rules (to include 410 IAC 1-4, Universal Precautions). (Indiana Department of Health; 410 IAC 26-11-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3366; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)
410 IAC 26-11-3 Laundry

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 3. The clinic, whether it operates its own laundry or uses outside laundry service, must ensure that the laundry process complies with a recognized laundry standard as follows:

1. Clean linen must be separated from soiled linen at all times.
2. Contaminated linens must be clearly identified and bagged.
3. Central clean linen storage space must be provided as follows:
   (A) If commercial laundry services are utilized:
   (i) a soiled linen collection area must be provided; and
   (ii) a hand washing facility is required in each area where unbagged soiled linen is handled.
   (B) If laundry is processed in the clinic:
   (i) a laundry processing area must be provided;
   (ii) clean linen storage and mending must be separated from soiled linen handling and storage; and
   (iii) employee hand washing facilities must be available in each room where clean or soiled linen is processed and handled.

(Indiana Department of Health; 410 IAC 26-11-3; filed May 11, 2006, 9:36 a.m.: 29 IR 3367; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 12. Emergency Care

410 IAC 26-12-1 Emergency care

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) The clinic must have a readily accessible written protocol for the following:
1. Managing medical emergencies that occur within the clinic. The protocol must ensure physician coverage and provide for a timely response for emergencies.
2. The transfer of patients requiring further emergency care to a hospital.
(b) Patients not discharged from the clinic within twelve (12) hours following the conclusion of a surgical abortion procedure must be transferred to a hospital. (Indiana Department of Health; 410 IAC 26-12-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3367; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 13. Anesthesia and Surgical Services

410 IAC 26-13-1 Anesthesia services

Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2; IC 25

Sec. 1. (a) The clinic must provide adequate anesthesia services to meet the needs of the patient, within the scope of the services offered, in accordance with acceptable standards of practice, under the direction of a licensed physician with specialized training or experience in the administration of anesthetics.
(b) Anesthesia services must be provided in compliance with IC 25 and rules adopted under that title.
(c) Anesthesia services in a clinic are limited to the following:
   1. Minimal sedation.
   2. Moderate sedation.
   (d) The medical director shall adopt and implement policies and procedures that include, but are not limited to, the following:

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(1) Safety rules to be followed relating to the administration of anesthesia.
(2) Safety training required of personnel.
(e) Anesthesia must be administered by one (1) of the following:
   (1) A qualified physician with appropriate training and experience.
   (2) A registered nurse acting under the direction of and in the immediate presence of the operating physician or other physician and who holds a certificate of completion of a course in anesthesia approved by the:
       (A) American Association of Nurse Anesthetists; or
       (B) medical licensing board of Indiana.
(f) The clinic shall ensure the delineation of preanesthesia, intraoperative, and postanesthesia responsibilities as follows:
   (1) The completion, within forty-eight (48) hours before a surgical abortion, of a preanesthesia evaluation for each patient by an individual qualified to administer anesthesia. If completed more than forty-eight (48) hours before the surgical abortion, the preanesthesia evaluation shall be updated according to clinic policy.
   (2) When using moderate sedation, the patient shall be monitored by qualified personnel other than the physician performing the procedure that must include and document at five (5) minute intervals the following:
       (A) Pulse oximetry.
       (B) Observed pulmonary ventilation.
       (C) Heart rate.
       (D) Blood pressure.
       (E) Response to verbal commands.
   (3) The completion of a postanesthetic evaluation for proper anesthesia recovery of each patient before discharge in accordance with written policies and procedures approved by the medical staff.
   (4) The requirement that all postoperative patients must be discharged from the postanesthetic care unit by the physician responsible for the patient's care in accordance with clinic policy.

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(5) Completion of the abortion documented by:
(A) ultrasonography; or
(B) other clinical means.

(6) Provision of follow-up examination and services as indicated.

(Indiana Department of Health; 410 IAC 26-13-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3368; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-13-3 Equipment
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 3. (a) There must be sufficient patient care equipment to assure the safe, effective, and timely provision of the available services to patients.
(b) The following equipment and supplies must be available to the procedure and recovery areas:
(1) Emergency call system.
(2) Oxygen.
(3) Resuscitation equipment.
(c) The following equipment and supplies must be available to the procedure and recovery areas when using IV sedation:
(1) Defibrillator.
(2) Cardiac monitors.
(3) Pulse oximeter.
(4) Suction equipment.
(5) Other supplies and equipment specified by the medical staff.

(Indiana Department of Health; 410 IAC 26-13-3; filed May 11, 2006, 9:36 a.m.: 29 IR 3368; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 14. Dietary Services
410 IAC 26-14-1 Dietary services
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) If nourishment and other dietary needs of the patients are provided in the clinic, the clinic must comply with 410 IAC 7-24.
(b) If nourishments are to be prepared, a nourishment area with a hand washing lavatory and refrigeration must be provided.
(c) If prepackaged single-service nourishments are provided, refrigeration storage of nourishments and other food products must be separate from refrigeration storage for pharmaceuticals. (Indiana Department of Health; 410 IAC 26-14-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3368; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 15. Laboratory Services
410 IAC 26-15-1 Laboratory services
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) The clinic must provide, or make available, those pathology and medical laboratory services and consultations necessary to meet the needs of patients as determined by the medical staff.
(b) The laboratory performs tests, examines specimens, and reports the evaluation only upon the written request of individuals authorized by law.
(c) The clinic must assure that all laboratory services provided to its patients are performed in a facility possessing a valid certificate, in accordance with 42 CFR 493 (excluding Subparts F, R, Q, and T) authorizing the performance of testing in the specialty or subspecialty of service for level of complexity in which the test is categorized.

(d) Laboratory supervisory and testing personnel qualifications must be:
   (1) consistent with the work assignments; and
   (2) in compliance with 42 CFR 493.

(e) All nursing and other clinic personnel performing laboratory testing must have competency assessed annually with documentation of assessment maintained in the employee file for the procedures performed.

(f) The clinic must develop, implement, and maintain written quality control and quality assurance policies and procedures for complexity of testing performed that are consistent with and include all standards found in 42 CFR 493. (Indiana Department of Health; 410 IAC 26-15-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3368; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 16. Pharmaceutical Services

410 IAC 26-16-1 Pharmaceutical services
Authority:  IC 16-21-1-7; IC 16-21-2-2.5
Affected:  IC 16-21-1; IC 16-21-2

Sec. 1. The clinic must provide drugs and biologicals in a safe and effective manner in accordance with accepted professional practice. The clinic must have the following:
   (1) A:
      (A) designated professional person with prescriptive authority; or
      (B) pharmacist;
   who is responsible for the control of drug stocks in the clinic.
   (2) Records of stock supplies of all scheduled substances, including an accounting for all items purchased and dispensed.
   (3) Written policies and procedures developed, implemented, maintained, and made available to personnel, including, but not limited to, the following:
      (A) Drug:
         (i) handling;
         (ii) storing;
         (iii) labeling;
         (iv) dispensing; and
         (v) administration according to established clinic policies and acceptable standards of practice.
      (B) Reporting of adverse reactions and medication errors to the:
         (i) physician responsible for the patient; and
         (ii) appropriate committee;
   and documented in the patient's record.
   (C) Drugs must be accurately and clearly labeled and stored in specially designated, well-illuminated cabinets, closets, or storerooms and the following:
      (i) Drug cabinets must be accessible only to authorized personnel.
      (ii) Drug cabinets for storage of controlled drugs listed in Schedule II of the Comprehensive Drug Abuse Prevention and Control Act of 1976 and other drugs subject to abuse must be permanently affixed compartments that are separately locked.
      (iii) Drug carts, if used, with controlled drugs as designated in item (ii) must be securely affixed when not in use.
   (D) Instructions to the patient on the use of take home medication is the responsibility of the prescribing physician.
   (4) A formulary.
   (5) A list of available emergency drugs.
(Indiana Department of Health; 410 IAC 26-16-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3369; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)
Rule 17. Physical Plant; Maintenance; Equipment; Environment; Safety

410 IAC 26-17-1 Physical plant
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2; IC 22

Sec. 1. (a) The clinic must be constructed, arranged, and maintained to ensure the safety of the patient and to provide facilities for services authorized under the clinic license as follows:
(1) The plant operations and maintenance service, equipment maintenance, and environmental services must be as follows:
   (A) Staffed to meet the scope of the services provided.
   (B) Under the direction of a person or persons qualified by education, training, or experience according to clinic policy approved by the governing body.
(2) The clinic must provide a physical plant and equipment that meets the statutory requirements and regulatory provisions of the fire prevention and building safety commission (IC 22, 675 IAC 22), Indiana fire prevention codes (675 IAC 22), and Indiana building codes (675 IAC 13).
(b) Any full or partial replacement of the physical plant of a clinic, any addition or renovation to the physical plant of a clinic, or any acquisitions of additional buildings under the license of an existing abortion clinic shall:
   (1) comply with:
      (A) this article; and
      (B) all building, fire safety, and handicapped accessibility codes, and rules adopted and administered by the state building commission; and
   (2) be provided with water supply and sewage disposal services from municipal or community services.

410 IAC 26-17-2 Specifications of physical plant
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 2. (a) Building entrances used to reach the clinic shall be as follows:
(1) At grade level.
(2) Clearly marked.
(3) Located so that patients need not go through other activity areas.

When the abortion clinic is part of another facility, separation of and access to the clinic shall be maintained. Lobbies of multioccupancy buildings may be shared. The design of the clinic shall preclude unrelated traffic from the clinic.
(b) The clinic design shall ensure appropriate levels of patient:
   (1) audible and visual privacy; and
   (2) dignity;
throughout the care process.
(c) For common administration and authorized visitor areas, the clinic shall be able to accommodate wheelchairs and provide the following:
   (1) A reception and information counter. The reception and information counter or desk shall be as follows:
      (A) Located to provide visual control of the entrance to the clinic.
      (B) Immediately apparent from the entrance.
   (2) A waiting area. The waiting area shall be under staff control. The seating area shall contain not fewer than two (2) spaces for each examination and procedure room.
   (3) At least one (1) conveniently accessible toilet room containing a lavatory for hand washing.
   (4) Conveniently accessible drinking water.
   (5) Interview space for private interviews related, for example, to social services or credit.
   (6) General storage facilities for supplies and equipment needed for continuing operation.
(d) Requirements for clinical facilities are as follows:

(1) Procedure rooms shall be segregated and removed from general traffic flow and be a minimum of:
   (A) one hundred twenty (120) square feet, exclusive of vestibules, toilets, and closets for procedures requiring only local analgesia or nitrous oxide; and
   (B) two hundred fifty (250) square feet, exclusive of vestibules, toilets, or closets for procedures that require conscious sedation.

(2) Rooms exclusively used for examinations shall be a minimum of eighty (80) square feet exclusive of vestibules, toilets, or closets.

(3) A hand washing station shall be included within each procedure room.

(4) Scrub facilities:
   (A) shall be provided near the entrance of procedure rooms;
   (B) may provide service to multiple procedure rooms if needed; and
   (C) shall be arranged to minimize splatter on nearby personnel or supply carts.

(5) A separate recovery room or area shall be included and provide for the following:
   (A) A minimum clear area of two (2) feet, six (6) inches around three (3) sides of each recovery cart or lounge chair for work and circulation.
   (B) A method of providing privacy for each patient in the room or area.
   (C) A work station with the following:
      (i) A countertop.
      (ii) Space for supplies.
      (iii) Provisions for charting.
      (iv) A communication system.

(6) A drug distribution station will be included. The station:
   (A) may be a part of the work station; and
   (B) shall include a:
      (i) work counter;
      (ii) sink;
      (iii) refrigerator (if needed); and
      (iv) locked storage for biologicals and drugs.

(7) A toilet room containing a lavatory for hand washing shall be accessible from all examination and procedure rooms. Where a clinic has no more than a total of three (3) examination and procedures rooms, the patient toilet may also serve as the toilet for the waiting area.

(e) Requirements for design standards are as follows:

(1) At least one (1) housekeeping room with:
   (A) a service sink; and
   (B) adequate storage for housekeeping supplies and equipment; shall be provided.

(2) Hand washing stations shall:
   (A) be located and arranged to meet the needs of the clinic; and
   (B) permit proper use and operation.

Provisions for hand drying shall be included at all hand washing stations except scrub sinks.

(3) There shall be an equipment room or rooms for:
   (A) heating;
   (B) air conditioning;
   (C) hot water;
   (D) other mechanical; and
   (E) electrical; equipment.

(4) Incinerators, if used, shall also conform to the building standards prescribed by area air pollution regulations.

(5) The minimum corridor width shall be forty-four (44) inches. Items such as drinking fountains, telephones, or vending
machines shall not:
   (A) restrict corridor traffic; or
   (B) reduce the corridor width below the required minimum.
(6) The minimum nominal door width for patient use shall be three (3) feet.
(7) Each building shall have at least two (2) exits that are remote from each other.
(8) An approved antiscald device shall be provided on the hot water supply to all hand washing facilities limiting the water temperature to a maximum of one hundred ten (110) degrees Fahrenheit (forty-three (43) degrees Celsius).

(Indiana Department of Health; 410 IAC 26-17-2; filed May 11, 2006, 9:36 a.m.: 29 IR 3370; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164RFA)

410 IAC 26-17-3 Maintenance of physical plant
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 3. The condition of the physical plant and the overall clinic environment must be developed and maintained in such a manner that the safety and well-being of patients is assured as follows:
(1) No condition in the clinic or on the grounds may be maintained that may be conducive to the harboring or breeding of:
   (A) insects;
   (B) rodents; or
   (C) other vermin.
(2) No condition may be created or maintained that may result in a hazard to:
   (A) patients;
   (B) authorized visitors; or
   (C) employees.
(3) Provision must be made for the periodic inspection, preventive maintenance, and repair of the physical plant and equipment by qualified personnel as follows:
   (A) Operation, maintenance, and spare parts manuals must be available, along with training or instruction, or both, of the appropriate clinic personnel, in the maintenance and operation of fixed and movable equipment.
   (B) All mechanical equipment (pneumatic, electric, sterilizing, or other) must be on a documented maintenance schedule of appropriate frequency in accordance with one (1) of the following:
      (i) Acceptable standards of practice.
      (ii) The manufacturer's recommended maintenance schedule.
   (C) Operational and maintenance control records must be as follows:
      (i) Established and analyzed at least triennially.
      (ii) Readily available on the premises.
   (D) Maintenance and repairs must be carried out in accordance with applicable codes, rules, standards, and requirements of the following:
      (i) Local jurisdictions.
      (ii) The state fire marshal.
      (iii) The department.

(Indiana Department of Health; 410 IAC 26-17-3; filed May 11, 2006, 9:36 a.m.: 29 IR 3371; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-17-4 Maintenance of equipment
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 4. All patient care equipment must be in good working order and regularly serviced and maintained as follows:
(1) All patient care equipment must be on a documented maintenance schedule of appropriate frequency in accordance with
one (1) of the following:
   (A) Acceptable standards of practice.
   (B) The manufacturer's recommended maintenance schedule.

(2) There must be evidence of preventive maintenance on all patient care equipment.

(3) Appropriate records must be:
   (A) kept pertaining to:
      (i) equipment maintenance;
      (ii) repairs; and
      (iii) electrical current leakage checks; and
   (B) analyzed at least triennially.

(4) Defibrillators must be discharged at least in accordance with manufacturers' recommendations, and a discharge log with initialed entries must be maintained.

(Indiana Department of Health; 410 IAC 26-17-4; filed May 11, 2006, 9:36 a.m.: 29 IR 3371; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-17-5 Environment

Sec. 5. The building or buildings, including fixtures, walls, floors, ceiling, and furnishings throughout, must be kept clean and orderly in accordance with current standards of practice, including the following:

(1) Environmental services must be provided in such a way as to guard against transmission of disease to patients, health care workers, the public, and visitors by using the current principles of the following:
   (A) Asepsis.
   (B) Cross-contamination prevention.
   (C) Safe practice.

(2) Refuse, biohazards, infectious waste, and garbage must be:
   (A) collected;
   (B) transported;
   (C) sorted; and
   (D) disposed of;

by methods that will minimize nuisances or hazards in compliance with federal, state, and local laws and rules.

(Indiana Department of Health; 410 IAC 26-17-5; filed May 11, 2006, 9:36 a.m.: 29 IR 3371; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

410 IAC 26-17-6 Safety

Sec. 6. (a) A safety management program must include, but not be limited to, the following:

(1) A review of safety functions.

(2) Development, implementation, and monitoring of a safety management program to include, but not be limited to, the following:
   (A) Periodic equipment inspections.
   (B) Insect, rodent, or other vermin control.
   (C) Instructions for operating and maintaining the building or building portion and equipment.
   (D) Chemical substance use and storage.
   (E) Surgical waste and similar material disposal.
   (F) General housekeeping precautions.

(3) An ongoing clinic-wide process to evaluate and collect information about hazards and safety practices.
(4) A safety program that includes, but is not limited to, the following:
   (A) Patient safety.
   (B) Health care worker safety.
   (C) Public and visitor safety.

(5) A written fire control plan that contains provisions for the following:
   (A) Prompt reporting of fires.
   (B) Extinguishing of fires.
   (C) Protection of the following:
      (i) Patients.
      (ii) Personnel.
      (iii) Guests.
   (D) Evacuation.
   (E) Cooperation with firefighting authorities.
   (F) Fire drills.

(6) Maintenance of written evidence of regular inspection and approval by state or local fire control agencies in accordance
    with the following:
   (A) Clinic policy.
   (B) State and local regulations.

(7) Emergency and disaster preparedness coordinated with appropriate community, state, and federal agencies.
(b) The clinic must maintain adequate battery-powered lighting and sufficient equipment needed to provide for the:
   (1) completion of services; and
   (2) safety of patients and staff;

in the event of a power loss. (Indiana Department of Health; 410 IAC 26-17-6; filed May 11, 2006, 9:36 a.m.: 29 IR 3372; readopted
filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA)

Rule 18. Other Services

410 IAC 26-18-1 Other services
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2

Sec. 1. (a) If the clinic provides other services not covered in specific sections of this article, the services must meet the needs
of the patients served, within the scope of the service offered, and in accordance with acceptable standards of practice.
   (b) The services must be as follows:
      (1) Under the direction of a qualified person or persons.
      (2) Staffed in accordance with written clinic policies and in compliance with the applicable state and federal rules.

(Indiana Department of Health; 410 IAC 26-18-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3372; readopted filed Jul 12, 2012, 12:08
p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25
p.m.: 20191002-IR-410190164FRA)

Rule 18.5. Informed Consent Brochures

410 IAC 26-18.5-1 Informed consent brochures
Authority: IC 16-21-1-7; IC 16-21-2-2.5
Affected: IC 16-21-1; IC 16-21-2; IC 16-34-2-1.5

Sec. 1. Abortion clinics must provide informed consent brochures, as described in IC 16-34-2-1.5, in English, Spanish, and
German, inside the abortion clinic. (Indiana Department of Health; 410 IAC 26-18.5-1; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)
Rule 19. Incorporation by Reference

410 IAC 26-19-1 Incorporation by reference
   Authority: IC 16-21-1-7; IC 16-21-2-2.5
   Affected: IC 16-21-1; IC 16-21-2

   Sec. 1. (a) 42 CFR 493 (October 1, 2017) is hereby incorporated by reference as part of this article.
   (b) Federal rules that have been incorporated by reference do not include any later amendments than those specified in the incorporated citation. The Code of Federal Regulations is found at https://www.gpo.gov/fdsys/browse/collectionCfr.action?collectionCode=CFR.
   (c) All incorporated material is available for public review at the department. (Indiana Department of Health; 410 IAC 26-19-1; filed May 11, 2006, 9:36 a.m.: 29 IR 3372; readopted filed Jul 12, 2012, 12:08 p.m.: 20120808-IR-410120196RFA; readopted filed Sep 26, 2018, 2:48 p.m.: 20181024-IR-410180328RFA; filed Sep 6, 2019, 3:25 p.m.: 20191002-IR-410190164FRA)