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## HOUSE OF REPRESENTATIVES - FLOOR VERSION

STATE OF OKLAHOMA

1st Session of the 58th Legislature (2021)

ENGROSSED SENATE BILL NO. 779

By: Daniels, Bullard, Stephens,
David, Taylor and **Jett** of
the Senate

and

Lepak of the House

An Act relating to abortion; creating the Oklahoma Abortion-Inducing Drug Certification Program Act; defining terms; specifying applicability of act; directing creation of certification program; limiting provision of abortion-inducing drugs to certain practitioners and procedures; authorizing certain fees and contracts; directing State Board of Pharmacy to establish certain requirements for manufacturers, distributors and physicians; providing certification systems and requirements for manufacturers, distributors and physicians; requiring physician to maintain hospital admitting privileges or enter into certain written agreement; stating conditions of agreement; requiring Board to adopt certain reporting system; stating criteria of reporting system; requiring certain reporting of physicians; providing for reporting of adverse events; providing criminal penalties; providing for certain civil remedies, disciplinary sanctions and injunctive relief; specifying certain judicial procedures; directing Board to develop certain enforcement scheme; specifying criteria of enforcement scheme; providing for certain restitution; directing creation of certain public portal; requiring portal to list certain names and allow for certain complaints; providing for disposition of complaints; providing for confidentiality of complaints; providing certain construction and intent; authorizing certain intervention; providing severability; amending 59

1 O.S. 2011, Section 353.7, as last amended by Section 4, Chapter 106, O.S.L. 2018 (59 O.S. Supp. 2020, 2 Section 353.7), which relates to powers and duties of the Board; broadening allowed uses of fees; providing 3 for codification; and providing an effective date. 4 5 BE IT ENACTED BY THE PEOPLE OF THE STATE OF OKLAHOMA: 6 A new section of law to be codified 7 SECTION 1. NEW LAW in the Oklahoma Statutes as Section 1-757.1 of Title 63, unless 8 9 there is created a duplication in numbering, reads as follows: Sections 1 through 16 of this act shall be known and may be 10 11 cited as the "Oklahoma Abortion-Inducing Drug Certification Program 12 Act". SECTION 2. A new section of law to be codified 13 NEW LAW in the Oklahoma Statutes as Section 1-757.2 of Title 63, unless 14 15 there is created a duplication in numbering, reads as follows: As used in this act: 16 "Abortion" means the act of using or prescribing any 17 instrument, medicine, drug or any other substance, device or means 18 with the intent to terminate the pregnancy of a woman known to be 19 pregnant, with knowledge that the termination by those means will 20

with reasonable likelihood cause the death of the unborn child.

Such use, prescription or means is not an abortion if done with the intent to: 23

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- a. save the life or preserve the health of the unborn child.
  - b. remove a dead unborn child caused by spontaneous abortion, accidental trauma or a criminal assault on the pregnant woman or her unborn child,
  - c. remove an ectopic pregnancy, or

- d. treat a maternal disease or illness for which the prescribed drug is indicated;
- 2. "Abortion-inducing drug" means a medicine, drug or any other substance prescribed or dispensed with the intent of terminating the pregnancy of a woman known to be pregnant, with knowledge that the termination will with reasonable likelihood cause the death of the unborn child. This includes the off-label use of drugs known to have abortion-inducing properties, which are prescribed specifically with the intent of causing an abortion, such as mifepristone (Mifeprex), misoprostol (Cytotec) and methotrexate. This definition does not apply to drugs that may be known to cause an abortion, but which are prescribed for other medical indications, such as chemotherapeutic agents and diagnostic drugs. The use of such drugs to induce abortion is also known as "medical", "medication", "RU-486", "chemical", "Mifeprex regimen" or "drug-induced" abortion;
- 3. "Adverse Event", according to the Food and Drug

  Administration, means any untoward medical occurrence associated

  with the use of a drug in humans, whether or not considered drug-

- related. It does not include an adverse event or suspected adverse reaction that, had it occurred in a more severe form, might have caused death;
- 4. "Associated physician" means a person licensed to practice medicine in the state including medical doctors and doctors of osteopathy, that has entered into an associated physician agreement;
- 5. "Complication" means any adverse physical or psychological condition arising from the performance of an abortion which includes, but is not limited to, uterine perforation, cervical perforation, infection, heavy or uncontrolled bleeding, hemorrhage, blood clots resulting in pulmonary embolism or deep vein thrombosis, failure to actually terminate the pregnancy, incomplete abortion (retained tissue), pelvic inflammatory disease, endometritis, missed ectopic pregnancy, cardiac arrest, respiratory arrest, renal failure, metabolic disorder, shock, embolism, coma, placenta previa in subsequent pregnancies, preterm delivery in subsequent pregnancies, free fluid in the abdomen, hemolytic reaction due to the administration of ABO-incompatible blood or blood products, adverse reactions to anesthesia and other drugs, subsequent development of breast cancer, psychological complications such as depression, suicidal ideation, anxiety, sleeping disorders, death and any other adverse event as defined by the Food and Drug Administration criteria provided in the Medwatch Reporting System;

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- 7. "Hospital" means an institution providing medical and surgical treatment and nursing care for sick or injured people, or institutions defined under Section 1-701 of Title 63 of the Oklahoma Statutes;
- 8. "Manufacturers and distributors" means individuals or entities that create, produce, supply, transport or sell drugs, which include:
  - a. any substances recognized by an official pharmacopoeia or formulary,
  - any substances intended for use in the diagnosis,
     cure, mitigation, treatment, or prevention of disease,
  - c. any substances other than food intended to affect the structure or any function of the body, or
  - d. any substances intended for use as a component of a medicine but not a device or a component, part or accessory of a device;
- 9. "Obstetrician/gynecologist", also known as OB/GYN, means a licensed physician who specializes in the care of women during pregnancy and childbirth and in the diagnosis and treatment of diseases of the female reproductive organs and specializes in other

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women's health issues such as menopause, hormone problems,
contraception or birth control, and infertility;

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- 10. "Physician" means any person licensed to practice medicine in this state. The term includes medical doctors and doctors of osteopathy;
- 11. "Pregnant" or "pregnancy" means that female reproductive condition of having an unborn child in the mother's uterus;
- 12. "Provide" or "provision" means, when used regarding abortion-inducing drugs, any act of giving, selling, dispensing, administering, transferring possession to or otherwise providing or prescribing an abortion-inducing drug; and
- 13. "Unborn child" means an individual organism of the species homo sapiens, beginning at fertilization, until the point of being born-alive as defined in Title 1 U.S.C., Section 8(b).
- SECTION 3. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.3 of Title 63, unless there is created a duplication in numbering, reads as follows:
- This act applies to any physician, health care provider or other person who is providing abortion-inducing drugs for use within this state, or any manufacturer or distributor providing abortion-inducing drugs within this state.
- 22 SECTION 4. NEW LAW A new section of law to be codified 23 in the Oklahoma Statutes as Section 1-757.4 of Title 63, unless 24 there is created a duplication in numbering, reads as follows:

- A. The State Board of Pharmacy shall promulgate rules to create
  a certification program to oversee and regulate the provision of
  abortion-inducing drugs. Abortion-inducing drugs shall be
  transported and provided in this state only by manufacturers or
  distributors certified to do so under this program. The drugs shall
  only be provided to patients by physicians certified to do so under
  this program.
  - B. The program shall be known as the Oklahoma Abortion-Inducing Drug Certification Program.
  - C. The Board may assess reasonable fees and enter into contracts with persons or entities to implement the Oklahoma Abortion-Inducing Drug Certification Program.
  - D. Abortion-inducing drugs shall not be provided directly to the patient through the mail, or otherwise outside of the parameters of the Oklahoma Abortion-Inducing Drug Certification Program.
  - SECTION 5. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.5 of Title 63, unless there is created a duplication in numbering, reads as follows:
  - A. The State Board of Pharmacy shall establish the following requirements for manufacturers and distributors of abortion-inducing drugs, at a minimum:
  - 1. Require completion of the certification process for physicians as described in Section 7 of this act, and for

1 manufacturers and distributors, as described in Section 6 of this act;

- 2. Notify manufacturers and distributors of physicians certified under the Oklahoma Abortion-Inducing Drug Certification Program;
- 3. Develop a reporting system as specified in Section 9 of this 6 7 act;
  - Prohibit shipment of abortion-inducing drugs to physicians who become de-certified from the Oklahoma Abortion-Inducing Drug Certification Program;
  - 5. Audit newly certified manufacturers and distributors within ninety (90) calendar days after the manufacturer or distributor is authorized, and annually thereafter, to ensure that all processes and procedures are in place and functioning to support the requirements of the Oklahoma Abortion-Inducing Drug Certification Program;
  - 6. If a manufacturer or distributor is found to be noncompliant, immediately suspend manufacturer's or distributor's certification until the manufacturer or distributor demonstrates full compliance; and
    - Enforce compliance according to Section 12 of this act.
- The State Board of Pharmacy shall establish the following 22 В. requirements for physicians providing abortion-inducing drugs, at a minimum:

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1. Require completion of the certification process;

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- 2. Audit newly certified physicians within ninety (90) calendar days after the physician is authorized, and annually thereafter, to ensure that all required processes and procedures are in place and functioning to support the requirements of the Oklahoma Abortion-Inducing Drug Certification Program;
- 3. If a physician is found to be non-compliant, immediately suspend the physician's certification until such time that the physician demonstrates full compliance; and

Enforce compliance according to Section 12 of this act.

SECTION 6. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.6 of Title 63, unless there is created a duplication in numbering, reads as follows:

The State Board of Pharmacy shall adopt a certification system for any manufacturer or distributor intending to provide abortion-inducing drugs in the state. To be eligible to be certified under this section, manufacturers and distributors shall:

- 1. Be licensed by the Board;
- 2. Only distribute to physicians certified under this act;
- 3. Record each serial number from pharmaceutical packages distributed to each certified physician;
  - 4. Abide by all applicable standards of the Utilization Review

    Accreditation Commission (URAC) or National Association of Boards of

    Pharmacy (NABP);

- 5. For online sales or orders, hold a current ".pharmacy" or ".pharma" domain and abide by all the standards required by the NABP to maintain the domain;
- 6. Follow all other applicable state or federal laws related to the distribution or delivery of legend drugs including abortion-inducing drugs; and
- 7. Follow all acceptable processes and procedures to maintain a distribution or delivery system that is secure, confidential and follows all processes and procedures including those for storage, handling, shipping, tracking package serial numbers, proof of delivery and controlled returns of abortion-inducing drugs.
- SECTION 7. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.7 of Title 63, unless there is created a duplication in numbering, reads as follows:
- The State Board of Pharmacy shall adopt a certification system for any physician intending to provide abortion-inducing drugs to patients in the state. Individuals or physicians providing abortion-inducing drugs in other states are not automatically certified in this state, and shall be fully certified under this law prior to providing any abortion-inducing drugs to any pregnant women in this state. To be eligible to be certified under this section physicians shall:
- 1. Be licensed to practice medicine and in good standing in the state;

- 2. Examine any patient in person prior to providing abortion-2 inducing drugs;
  - 3. Sign an annual "Dispensing Agreement Form," to be developed and provided by the State Board of Pharmacy, before providing abortion-inducing drugs;
  - 4. Inform the patient of gestational age-specific risks of using abortion-inducing drugs;
  - 5. Assess for signs of domestic abuse, reproductive control, human trafficking and other signals of coerced abortion, per current state guidelines;
  - 6. Adequately inform the patient of gestational age-specific age risks of using abortion-inducing drugs;
  - 7. Inform the patient that she may see the remains of her unborn child in the process of completing the abortion;
  - 8. Inform the patient that studies show that babies born following the abortion reversal process have a rate of birth defects no higher than the general population;
  - 9. Inform the patient that studies show that following this reversal process or otherwise treating a woman with progesterone during pregnancy does not lead to increased mortality rates;
  - 10. Refrain from knowingly supplying abortion-inducing drugs to patients who present with any of the following:
    - a. absence of a pregnancy,

1	b.	bein	g post-seventy days gestation or post-ten weeks of	
2		preg	nancy, and	
3	С.	havi	ng risk factors associated with abortion-inducing	
4		drug	s including, but not limited to:	
5		(1)	ectopic pregnancies,	
6		(2)	problems with the adrenal glands near the	
7			kidneys,	
8		(3)	being treated with long-term corticosteroid	
9			therapy,	
10		(4)	allergic reactions to abortion-inducing drugs,	
11			mifepristone, misoprostol or similar drugs,	
12		(5)	bleeding problems or is taking anticoagulant drug	
13			products,	
14		(6)	has inherited porphyria,	
15		(7)	has an intrauterine device in place, or	
16		(8)	being Rh Negative, requiring administration of	
17			Rhogam before providing abortion-inducing drugs;	
18	11. Provide or refer for emergency surgical intervention in			
19	cases of incomplete abortion, severe bleeding or other medical			
20	complications, through maintaining hospital admitting privileges or			
21	entering into a written agreement with an associated physician as			
22	specified in Section 8 of this act;			
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- 13. Sign, and ensure that the patient signs, all legally required informed consent material, providing patient with a copy showing both signatures, and placing the original in the patient's medical record;
- 14. Record the serial number from each package of each abortion-inducing drug given to the patient in her medical record;
- 15. Submit a written protocol of how efforts will be made to schedule with the patient the medically indicted follow-up appointment within fourteen (14) days to assure a completed abortion;
- 16. Report to the State Board of Pharmacy, as well as the Food and Drug Administration, any death associated with abortion-inducing drugs with the following guidelines:
  - a. the patient shall be noted by a non-identifiable reference and the serial number from each package of abortion-inducing drug given, whether or not considered drug-related,
  - b. this shall be done as soon as possible but no later than fifteen (15) calendar days from the initial receipt of the information by the physician, and

1 this requirement does not affect the physician's other 2 reporting and follow-up requirements under the Oklahoma Abortion-Inducing Drug Certification Program 3 or any additional requirements by another department 4 that oversees the abortion industry in this state;

- Submit a written protocol of how complications will be 17. handled by the certified physician and submit a copy of a signed contract with an associated physician credentialed to handle certain complications as outlined in Section 8 of this act;
- 18. Abide by all applicable state and federal laws regarding medical records retention, confidentiality and privacy; and
- Agree to follow and document compliance with all other legally required conditions for performing abortion in the state where the patient presents for her appointment including, but not limited to, waiting periods, informed consent requirements, statistical reporting, parental consent or notification, and required inspections.
- A new section of law to be codified SECTION 8. NEW LAW in the Oklahoma Statutes as Section 1-757.8 of Title 63, unless there is created a duplication in numbering, reads as follows:
- The State Board of Pharmacy shall also require the following of 21 certified physicians: 22
  - Maintaining hospital admitting privileges at one or more hospitals in the county or contiquous county where the abortion-

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inducing drug was provided, and informing the patient of any hospital where the physician holds admitting privileges.

- 2. Alternatively, the physician may enter into a written agreement with an associated physician in the county or contiguous county where the abortion-inducing drug was provided. The written agreement shall meet these conditions:
  - a. a physician who provides an abortion-inducing drug shall notify the patient of the location of the hospital at which the associated physician has admitting privileges,
  - b. the physician shall keep, at the location of his or her practice, a copy of the written agreement,
  - c. the physician shall submit a copy of the written agreement to the State Department of Health as part of any required clinic licensure,
  - d. the State Department of Health shall verify the validity of the document, and shall remove any personal identifying information of the patient from the document before releasing the document in accordance with the following:
    - (1) the State Department of Health shall annually submit a copy of the written agreement described in this paragraph to each hospital located in the

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Age of patients served;

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- 4. County and state of residence of patients served;
- 5. If the patient resides outside the United States, city and country of residence;
  - 6. County and state of service;
  - 7. A list of staff attending patients including licensing numbers and evidence of other qualifications;
    - 8. Each medication used or provided per patient, by date;
  - 9. Any known complications or adverse events, and how they were addressed, by date; and
  - 10. Unresolved cases.
- B. This reporting system shall also be used by emergency
  department physicians and private physicians who treat post-abortion
  complications.
  - C. Physicians shall protect from disclosure any personally identifiable information of the patient in accordance with applicable federal and state law.
  - D. A certified physician shall also report to the State Board of Pharmacy, as well as the Medwatch Reporting System of the Food and Drug Administration (FDA), any complication or adverse event as defined according to the FDA criteria given in the Medwatch Reporting System.
- E. The State Board of Pharmacy shall develop a system of reporting adverse events from the use of abortion-inducing drugs for

- 1 this state. The system shall require reporting of complications and
- 2 | adverse events including, but not limited to:
- 3 1. Death;

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- 4 2. Blood loss including hemorrhage;
- 5 3. Infection including sepsis;
  - 4. Blood transfusions;
    - 5. Administer drug for an ectopic pregnancy; and
- 8 6. Other adverse effects requiring hospitalization or
- 9 additional medical care.
- F. The State Board of Pharmacy shall require the following providers and entities to report complications and adverse events in writing:
- 1. Physicians certified to provide abortion-inducing drugs;
- 2. Emergency room physicians;
- 3. Any doctor licensed in this state including an obstetrician/gynecologist who treats women with adverse events;
- 4. Provision of certification requires that the physician shall also report adverse events and any patient deaths to the FDA; and
- 5. Other individuals or entities as determined by the State Board of Pharmacy.
- 21 SECTION 10. NEW LAW A new section of law to be codified 22 in the Oklahoma Statutes as Section 1-757.10 of Title 63, unless 23 there is created a duplication in numbering, reads as follows:

- A. Individuals or entities not certified under the Oklahoma

  Abortion-Inducing Drug Certification Program that provide drugs for
  the purpose of inducing abortion are in violation of this act.
- B. Individuals or entities that provide abortion-inducing drugs to any person or entity that is not certified, or otherwise authorized, to provide abortion-inducing drugs under the Oklahoma Abortion-Inducing Drug Certification Program are in violation of this act.
- C. A person who intentionally, knowingly or recklessly violates any provision of this act is guilty of a misdemeanor.
- D. A person who intentionally, knowingly or recklessly violates any provision of this act by fraudulent use of an abortion-inducing drug, with or without the knowledge of the pregnant woman, is guilty of a felony.
- E. No civil or criminal penalty may be assessed against the pregnant woman upon whom the drug-induced abortion is attempted, induced or performed.
- SECTION 11. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.11 of Title 63, unless there is created a duplication in numbering, reads as follows:
- A. In addition to whatever remedies are available under the common or statutory law of this state, failure to comply with the requirements of this act shall:

- 1 l. Provide a basis for a civil malpractice action for actual and punitive damages;
  - 2. Provide a basis for a professional disciplinary action; and
  - 3. Provide a basis for recovery for the woman's survivors for the wrongful death of the woman.
  - B. When requested, the court shall allow a woman to proceed using solely her initials or a pseudonym and may close any proceedings in the case and enter other protective orders to preserve the privacy of the woman upon whom the drug-induced abortion was attempted, induced or performed.
  - C. If judgment is rendered in favor of the plaintiff, the court shall also render judgment for reasonable attorney's fees in favor of the plaintiff against the defendant.
  - D. If judgment is rendered in favor of the defendant and the court finds that the plaintiff's suit was frivolous and brought in bad faith, the court may render judgment for reasonable attorney's fees in favor of the defendant against the plaintiff.
  - E. A cause of action for injunctive relief against a person who has provided an abortion-inducing drug in violation of this act may be maintained by:
    - 1. A woman to whom such an abortion-inducing drug was provided;
- 22 2. A person who is the spouse, parent or guardian of, or a
  23 current or former licensed health care provider of, a woman to whom
  24 such an abortion-inducing drug was provided; or

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3. A prosecuting attorney with appropriate jurisdiction.

The injunction shall prevent the defendant from providing further abortion-inducing drugs in violation of this act.

SECTION 12. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.12 of Title 63, unless there is created a duplication in numbering, reads as follows:

- A. The State Board of Pharmacy shall develop an enforcement scheme to enforce this act, which includes:
- 1. When an individual or entity provides abortion-inducing drugs without first seeking certification under this act, the State Board of Pharmacy shall:
  - a. immediately report the illegal act to local law enforcement, or other applicable state and local agencies for investigation or other appropriate action, where appropriate,
  - b. impose a fine of no less than Five Million Dollars (\$5,000,000.00) for manufacturers or distributors and Two Hundred Fifty Thousand Dollars (\$250,000.00) for physicians;
- 2. When a certified manufacturer or distributor or physician is determined to be in non-compliance, suspend certification until compliance is proven to the satisfaction of the State Board of Pharmacy;

- 3. Where a current or previously certified manufacturer or distributer is found to have intentionally or knowingly violated this act, or refuses to bring operations into compliance within ninety (90) calendar days, remove certification and prohibit continued provision of abortion-inducing drugs by the manufacturer or distributor until compliance is demonstrated to the satisfaction of the State Board of Pharmacy;
  - 4. When a certified manufacturer, distributor or physician is in non-compliance, suspend all annual recertification until compliance is demonstrated to the satisfaction of the State Board of Pharmacy; and
  - 5. Where a current or previously certified manufacturer, distributer or physician is found to have intentionally or knowingly violated this act, or refuses to bring operations into compliance:
    - a. immediately suspend the manufacturer's, distributor's or physician's certification until full compliance is demonstrated,
    - b. for certified manufacturers or distributors, impose fines of not less than One Million Dollars (\$1,000,000.00) per offense,
    - c. for certified physicians, impose fines of not less than One Hundred Thousand Dollars (\$100,000.00) per offense,

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- d. permanently revoke the certification of the offender if offender fails to demonstrate compliance within ninety (90) calendar days,
- e. impose remedial actions, which may include additional education, additional reporting or other actions as required by the State Board of Pharmacy,
- f. in the case of a licensed manufacturer or distributor, recommend sanctioning to the appropriate disciplinary committee of the State Board of Pharmacy,
- g. in the case of a licensed physician, report the violation to the appropriate medical licensing board,
- h. publicly report any disciplinary actions, consistent with the practices of the State Board of Pharmacy,
- i. permanently revoke the certification of the offender,
- j. in the case of a licensed manufacturer or distributor, recommend permanent revocation of licensure,
- k. in the case of a licensed physician, recommend appropriate sanctioning to the appropriate medical licensing board, and
- publicly report any disciplinary actions consistent with the practices of the State Board of Pharmacy.
- B. Individuals have a Private Right of Action to seek restitution in any court of law with appropriate jurisdiction for any and all damages suffered due to a violation of this act.

- SECTION 13. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.13 of Title 63, unless there is created a duplication in numbering, reads as follows:
  - A. The State Board of Pharmacy shall develop on its website a complaint portal for patients, pharmacy, nursing and medical professionals and the public to submit information about potential violations offered at no charge to the parties named in this subsection.
  - B. The portal shall list the names of manufacturers and distributors that are certified under the program, as well as the physicians that are certified under the program.
  - C. The portal shall allow the party to make a complaint anonymously.
  - D. The State Board of Pharmacy shall review each complaint and determine a disposition including referral to another appropriate state agency, within thirty (30) days.
  - E. Confidentiality of the originator of the complaint shall be protected at all times except for intra-state referrals for investigation.
  - SECTION 14. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.14 of Title 63, unless there is created a duplication in numbering, reads as follows:
- A. Nothing in this act shall be construed as creating or recognizing a right to abortion.

B. It is not the intention of this act to make lawful an abortion that is otherwise unlawful.

C. Nothing in this act repeals, replaces or otherwise invalidates existing federal or state laws, regulations or policies.

SECTION 15. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.15 of Title 63, unless there is created a duplication in numbering, reads as follows:

The Legislature, by joint resolution, may appoint one or more of its members, who sponsored or cosponsored this act in his or her official capacity, to intervene as a matter of right in any case in which the constitutionality of this act is challenged.

SECTION 16. NEW LAW A new section of law to be codified in the Oklahoma Statutes as Section 1-757.16 of Title 63, unless there is created a duplication in numbering, reads as follows:

If any one or more provisions, sections, subsections, sentences, clauses, phrases or words of this act or the application thereof to any person or circumstance is found to be unconstitutional, the same is hereby declared to be severable and the balance of this act shall remain effective notwithstanding such unconstitutionality. The Legislature hereby declares that it would have passed this act, and each provision, section, subsection, sentence, clause, phrase or word thereof, irrespective of the fact that any one or more provisions, sections, subsections, sentences, clauses, phrases or words be declared unconstitutional.

1 | SECTION 17. AMENDATORY 59 O.S. 2011, Section 353.7, as

2 | last amended by Section 4, Chapter 106, O.S.L. 2018 (59 O.S. Supp.

- 3 2020, Section 353.7), is amended to read as follows:
- 4 Section 353.7. The State Board of Pharmacy shall have the power
- 5 and duty to:

- 1. Regulate the practice of pharmacy;
- 7 | 2. Regulate the sale and distribution of drugs, medicines,
- 8 chemicals and poisons;
- 9 3. Regulate the dispensing of drugs and medicines in all places
- 10 where drugs and medicines are compounded and/or dispensed;
- 4. Examine and issue appropriate certificates of licensure as
- 12 | Doctor of Pharmacy to all applicants whom the Board deems qualified
- 13 | under the provisions of the Oklahoma Pharmacy Act;
- 14 5. Issue licenses to manufacturers, repackagers, outsourcing
- 15 | facilities, wholesale distributors, third-party logistics providers,
- 16 pharmacies, and other dispensers, medical gas suppliers, and medical
- 17 gas distributors;
- 18 6. Issue sterile compounding and drug supplier permits for
- 19 pharmacies at the fee set by the Board, with the expiration date of
- 20 | such permits to coincide with the pharmacy license annual expiration
- 21 date;
- 7. Prescribe minimum standards with respect to floor space and
- 23 other physical characteristics of pharmacies and hospital drug rooms
- 24 | as may be reasonably necessary for the maintenance of professional

- surroundings and for the protection of the safety and welfare of the public, and to refuse the issuance of new or renewal licenses for failure to comply with such standards. Minimum standards for hospital drug rooms shall be consistent with the State Department of Health, Hospital Standards, as defined in OAC 310:667;
  - 8. Authorize its inspectors, compliance officers, and duly authorized representatives to enter and inspect any and all places, including premises, vehicles, equipment, contents and records, where drugs, medicines, chemicals, or poisons are stored, sold, vended, given away, compounded, dispensed, manufactured, repackaged or transported;
  - 9. Employ the number of inspectors and pharmacist compliance officers necessary in the investigation of criminal activity or preparation of administrative actions at an annual salary to be fixed by the Board, and to authorize necessary expenses. Any inspector certified as a peace officer by the Council of Enforcement Education and Training shall have statewide jurisdiction to perform the duties authorized by this section. In addition, the inspectors shall be considered peace officers and shall have the same powers and authority as that granted to peace officers. In addition, such inspectors or pharmacist compliance officers shall have the authority to take and copy records and the duty to confiscate all drugs, medicines, chemicals or poisons found to be stored, sold,

- vended, given away, compounded, dispensed or manufactured contrary
  to the provisions of the Oklahoma Pharmacy Act;
  - 10. Investigate complaints, subpoena witnesses and records, initiate prosecution, and hold hearings;
  - 11. Administer oaths in all manners pertaining to the affairs of the Board and to take evidence and compel the attendance of witnesses on questions pertaining to the enforcement of the Oklahoma Pharmacy Act;
  - 12. Reprimand, place on probation, suspend, revoke permanently and levy fines not to exceed Three Thousand Dollars (\$3,000.00) for each count for which any person charged with violating the Oklahoma Pharmacy Act or Oklahoma Board of Pharmacy administrative rules has been convicted in Board hearings. The Board also may take other disciplinary action. The Board may impose as part of any disciplinary action the payment of costs expended by the Board for any legal fees and costs, including, but not limited to, staff time, salary and travel expense, witness fees and attorney fees. The Board may also require additional continuing education, including attendance at a live continuing education program, and may require participation in a rehabilitation program for the impaired. The Board may take such actions singly or in combination, as the nature of the violation requires;
  - 13. Adopt and establish rules of professional conduct appropriate to the establishment and maintenance of a high standard

of integrity and dignity in the profession of pharmacy. Such rules shall be subject to amendment or repeal by the Board as the need may arise;

- 14. Make and publish rules such as may be necessary for carrying out and enforcing the provisions of the Oklahoma Pharmacy Act, Oklahoma drug laws and rules, federal drug laws and regulations, and make such other rules as in its discretion may be necessary to protect the health, safety, and welfare of the public;
- 15. Establish and collect appropriate fees for licenses, permits, inspections, and services provided; and such fees shall be nonrefundable. Such fees shall be promulgated to implement the provisions of the Oklahoma Pharmacy Act under the provisions of the Administrative Procedures Act and the Oklahoma Abortion-Inducing Drug Certification Program Act;

## 16. Regulate:

- a. personnel working in a pharmacy, such as interns and supportive personnel, including technicians, and issue pharmacy technician permits and intern licenses,
- b. interns, preceptors and training areas through which the training of applicants occurs for licensure as a pharmacist, and
- c. such persons regarding all aspects relating to the handling of drugs, medicines, chemicals, and poisons;

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17. Acquire by purchase, lease, gift, solicitation of gift or
by any other manner, and to maintain, use and operate or to contract
for the maintenance, use and operation of or lease of any and all
property of any kind, real, personal or mixed or any interest
therein unless otherwise provided by the Oklahoma Pharmacy Act;
provided, all contracts for real property shall be subject to the
provisions of Section 63 of Title 74 of the Oklahoma Statutes;

- 18. Perform other such duties, exercise other such powers and loy such personnel as the provisions and enforcement of the ahoma Pharmacy Act may require; and
- Approve pilot projects designed to utilize new or expanded hnology or processes and provide patients with better pharmacy ducts or provide pharmacy services in a more safe and efficient Such approvals may include provisions granting exemptions any rule adopted by the Board.
  - SECTION 18. This act shall become effective November 1, 2021.

MITTEE REPORT BY: COMMITTEE ON PUBLIC HEALTH, dated 04/07/2021 -DO PASS, As Coauthored.

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