

1 ENGROSSED HOUSE AMENDMENT
TO
2 ENGROSSED SENATE BILL NO. 1150 By: Standridge of the Senate
3 and
4 Cox of the House
5

6
7 An Act relating to pharmacy; amending 59 O.S. 2011,
8 Section 353.1, as last amended by Section 1, Chapter
9 230, O.S.L. 2015 (59 O.S. Supp. 2015, Section 353.1),
10 which relates to definitions; updating statutory
11 references; amending 59 O.S. 2011, Section 353.11, as
12 amended by Section 7, Chapter 230, O.S.L. 2015 (59
13 O.S. Supp. 2015, Section 353.11), which relates to
14 license renewal; amending Section 8, Chapter 230,
15 O.S.L. 2015 (59 O.S. Supp. 2015, Section 353.11a),
16 which relates to continuing education requirements;
17 amending 59 O.S. 2011, Section 353.18, as amended by
18 Section 11, Chapter 230, O.S.L. 2015 (59 O.S. Supp.
19 2015, Section 353.18), which relates to the sale,
20 manufacturing, and packaging of dangerous drugs;
21 amending 59 O.S. 2011, Section 353.24, as amended by
22 Section 16, Chapter 230, O.S.L. 2015 (59 O.S. Supp.
23 2015, Section 353.24), which relates to unlawful
24 acts; amending 59 O.S. 2011, Section 353.26, as
amended by Section 17, Chapter 230, O.S.L. 2015 (59
O.S. Supp. 2015, Section 353.26), which relates to
revocations or suspensions of licenses; clarifying
language; repealing 59 O.S. 2011, Sections 353.13,
353.29, 364, and 366, which relate to unlawful acts,
supportive personnel, renewal certifications, and
alternative methods of meeting certain requirements;
and providing an effective date.

22 AUTHOR: Remove Senator Standridge as principal Senate author and
23 substitute with Senator Yen.

24 Add the following Senate Coauthor: Standridge

1 AMENDMENT NO. 1. Strike the title, enacting clause and entire bill
2 and insert

3 "An Act relating to pharmacy; amending 59 O.S. 2011,
4 Section 353.1, as last amended by Section 1, Chapter
5 230, O.S.L. 2015 (59 O.S. Supp. 2015, Section
6 353.1), which relates to definitions; updating
7 statutory references; amending 59 O.S. 2011, Section
8 353.11, as amended by Section 7, Chapter 230, O.S.L.
9 2015 (59 O.S. Supp. 2015, Section 353.11), which
10 relates to license renewal; amending Section 8,
11 Chapter 230, O.S.L. 2015 (59 O.S. Supp. 2015,
12 Section 353.11a), which relates to continuing
13 education requirements; amending 59 O.S. 2011,
14 Section 353.18, as amended by Section 11, Chapter
15 230, O.S.L. 2015 (59 O.S. Supp. 2015, Section
16 353.18), which relates to the sale, manufacturing
17 and packaging of dangerous drugs; amending 59 O.S.
18 2011, Section 353.24, as amended by Section 16,
19 Chapter 230, O.S.L. 2015 (59 O.S. Supp. 2015,
20 Section 353.24), which relates to unlawful acts;
21 amending 59 O.S. 2011, Section 353.26, as amended by
22 Section 17, Chapter 230, O.S.L. 2015 (59 O.S. Supp.
23 2015, Section 353.26), which relates to revocations
24 or suspensions of licenses; clarifying language;
amending Sections 1 and 4, Chapter 263, O.S.L. 2014
(59 O.S. Supp. 2015, Sections 357 and 360), which
relate to pharmacy benefit plans; defining terms;
modifying prohibited acts; modifying administrative
appeals procedure; modifying requirements for
certain contracts; repealing 59 O.S. 2011, Sections
353.13, 353.29, 364 and 366, which relate to
unlawful acts, supportive personnel, renewal
certifications and alternative methods of meeting
certain requirements; and providing an effective
date.

22 BE IT ENACTED BY THE PEOPLE OF THE STATE OF OKLAHOMA:

1 SECTION 1. AMENDATORY 59 O.S. 2011, Section 353.1, as
2 last amended by Section 1, Chapter 230, O.S.L. 2015 (59 O.S. Supp.
3 2015, Section 353.1), is amended to read as follows:

4 Section 353.1 For the purposes of the Oklahoma Pharmacy Act:

- 5 1. "Accredited program" means those seminars, classes,
6 meetings, work projects, and other educational courses approved by
7 the Board for purposes of continuing professional education;
- 8 2. "Act" means the Oklahoma Pharmacy Act;
- 9 3. "Administer" means the direct application of a drug, whether
10 by injection, inhalation, ingestion or any other means, to the body
11 of a patient;
- 12 4. "Assistant pharmacist" means any person presently licensed
13 as an assistant pharmacist in the State of Oklahoma by the Board
14 pursuant to Section 353.10 of this title and for the purposes of the
15 Oklahoma Pharmacy Act shall be considered the same as a pharmacist,
16 except where otherwise specified;
- 17 5. "Board" or "State Board" means the State Board of Pharmacy;
- 18 6. "Certify" or "certification of a prescription" means the
19 review of a filled prescription by a licensed pharmacist or a
20 licensed practitioner with dispensing authority to confirm that the
21 medication, labeling and packaging of the filled prescription are
22 accurate and meet all requirements prescribed by state and federal
23 law. For the purposes of this paragraph, "licensed practitioner"
24 shall not include optometrists with dispensing authority;

1 7. "Chemical" means any medicinal substance, whether simple or
2 compound or obtained through the process of the science and art of
3 chemistry, whether of organic or inorganic origin;

4 8. "Compounding" means the combining, admixing, mixing,
5 diluting, pooling, reconstituting or otherwise altering of a drug or
6 bulk drug substance to create a drug. Compounding includes the
7 preparation of drugs or devices in anticipation of prescription drug
8 orders based on routine, regularly observed prescribing patterns;

9 9. "Continuing professional education" means professional,
10 pharmaceutical education in the general areas of the socioeconomic
11 and legal aspects of health care; the properties and actions of
12 drugs and dosage forms; and the etiology, characteristics and
13 therapeutics of the diseased state;

14 10. "Dangerous drug", "legend drug", "prescription drug" or "Rx
15 Only" means a drug:

- 16 a. for human use subject to 21 U.S.C. 353(b)(1) ~~†~~ or
17 b. is labeled "Prescription Only", or labeled with the
18 following statement: "Caution: Federal law restricts
19 this drug except for use by or on the order of a
20 licensed veterinarian".

21 11. "Director" means the Executive Director of the State Board
22 of Pharmacy unless context clearly indicates otherwise;

23 12. "Dispense" or "dispensing" means the interpretation,
24 evaluation, and implementation of a prescription drug order,

1 including the preparation and delivery of a drug or device to a
2 patient or a patient's agent in a suitable container appropriately
3 labeled for subsequent administration to, or use by, a patient.
4 Dispense includes sell, distribute, leave with, give away, dispose
5 of, deliver or supply;

6 13. "Dispenser" means a retail pharmacy, hospital pharmacy, a
7 group of chain pharmacies under common ownership and control that do
8 not act as a wholesale distributor, or any other person authorized
9 by law to dispense or administer prescription drugs, and the
10 affiliated warehouses or distributions of such entities under common
11 ownership and control that do not act as a wholesale distributor.
12 For the purposes of this paragraph, "dispenser" does not mean a
13 person who dispenses only products to be used in animals in
14 accordance with 21 U.S.C. 360b(a) (5);

15 14. "Distribute" or "distribution" means the sale, purchase,
16 trade, delivery, handling, storage, or receipt of a product, and
17 does not include the dispensing of a product pursuant to a
18 prescription executed in accordance with 21 U.S.C. 353(b) (1) or the
19 dispensing of a product approved under 21 U.S.C. 360b(b);

20 15. "Doctor of Pharmacy" means a person licensed by the Board
21 to engage in the practice of pharmacy. The terms "pharmacist",
22 "D.Ph.", and "Doctor of Pharmacy" shall be interchangeable and shall
23 have the same meaning wherever they appear in the Oklahoma Statutes
24 and the rules promulgated by the Board;

1 16. "Drug outlet" means all manufacturers, repackagers,
2 outsourcing facilities, wholesale distributors, third-party
3 logistics providers, pharmacies, and all other facilities which are
4 engaged in dispensing, delivery, distribution or storage of
5 dangerous drugs;

6 17. "Drugs" means all medicinal substances and preparations
7 recognized by the United States Pharmacopoeia and National
8 Formulary, or any revision thereof, and all substances and
9 preparations intended for external and/or internal use in the cure,
10 diagnosis, mitigation, treatment or prevention of disease in humans
11 or animals and all substances and preparations, other than food,
12 intended to affect the structure or any function of the body of a
13 human or animals;

14 18. "Drug sample" means a unit of a prescription drug packaged
15 under the authority and responsibility of the manufacturer that is
16 not intended to be sold and is intended to promote the sale of the
17 drug;

18 19. "Filled prescription" means a packaged prescription
19 medication to which a label has been affixed which contains such
20 information as is required by the Oklahoma Pharmacy Act;

21 20. "Hospital" means any institution licensed as a hospital by
22 this state for the care and treatment of patients, or a pharmacy
23 operated by the Oklahoma Department of Veterans Affairs;

24

1 21. "Licensed practitioner" means an allopathic physician,
2 osteopathic physician, podiatric physician, dentist, veterinarian or
3 optometrist licensed to practice and authorized to prescribe
4 dangerous drugs within the scope of practice of such practitioner;

5 22. "Manufacturer" or "virtual manufacturer" means with respect
6 to a product:

7 a. a person that holds an application approved under 21
8 U.S.C. 355 or a license issued under 42 U.S.C. 262 for
9 such product, or if such product is not the subject of
10 an approved application or license, the person who
11 manufactured the product,

12 b. a co-licensed partner of the person described in
13 subparagraph a that obtains the product directly from
14 a person described in this subparagraph or
15 subparagraph a, or

16 c. an affiliate of a person described in subparagraph a
17 or b who receives the product directly from a person
18 described in this subparagraph or in subparagraph a or
19 b;

20 23. "Manufacturing" means the production, preparation,
21 propagation, compounding, conversion or processing of a device or a
22 drug, either directly or indirectly by extraction from substances of
23 natural origin or independently by means of chemical or biological
24 synthesis and includes any packaging or repackaging of the

1 substances or labeling or relabeling of its container, and the
2 promotion and marketing of such drugs or devices. The term
3 "manufacturing" also includes the preparation and promotion of
4 commercially available products from bulk compounds for resale by
5 licensed pharmacies, licensed practitioners or other persons;

6 24. "Medical gas" means those gases including those in liquid
7 state upon which the manufacturer or distributor has placed one of
8 several cautions, such as "Rx Only", in compliance with federal law;

9 25. "Medical gas order" means an order for medical gas issued
10 by a licensed prescriber;

11 26. "Medical gas distributor" means a person licensed to
12 distribute, transfer, wholesale, deliver or sell medical gases on
13 drug orders to suppliers or other entities licensed to use,
14 administer or distribute medical gas and may also include a patient
15 or ultimate user;

16 27. "Medical gas supplier" means a person who dispenses medical
17 gases on drug orders only to a patient or ultimate user;

18 28. "Medicine" means any drug or combination of drugs which has
19 the property of curing, preventing, treating, diagnosing or
20 mitigating diseases, or which is used for that purpose;

21 29. "Nonprescription drugs" means medicines or drugs which are
22 sold without a prescription and which are prepackaged for use by the
23 consumer and labeled in accordance with the requirements of the
24 statutes and regulations of this state and the federal government.

1 Such items shall also include medical and dental supplies and
2 bottled or nonbulk chemicals which are sold or offered for sale to
3 the general public if such articles or preparations meet the
4 requirements of the Federal Food, Drug and Cosmetic Act, 21
5 U.S.C.A., Section 321 et seq.;

6 30. "Outsourcing facility", including "virtual outsourcing
7 facility" means a facility at one geographic location or address
8 that:

- 9 a. is engaged in the compounding of sterile drugs,
- 10 b. has elected to register as an outsourcing facility,
- 11 and
- 12 c. complies with all requirements of 21 U.S.C. 353b;

13 31. "Package" means the smallest individual saleable unit of
14 product for distribution by a manufacturer or repackager that is
15 intended by the manufacturer for ultimate sale to the dispenser of
16 such product. For the purposes of this paragraph, "individual
17 saleable unit" means the smallest container of a product introduced
18 into commerce by the manufacturer or repackager that is intended by
19 the manufacturer or repackager for individual sale to a dispenser;

20 32. "Person" means an individual, partnership, limited
21 liability company, corporation or association, unless the context
22 otherwise requires;

23 33. "Pharmacist-in-charge" or "PIC" means the pharmacist
24 licensed in this state responsible for the management control of a

1 pharmacy and all other aspects of the practice of pharmacy in a
2 licensed pharmacy as defined by Section 353.18 of this title;

3 34. "Pharmacy" means a place regularly licensed by the Board of
4 Pharmacy in which prescriptions, drugs, medicines, chemicals and
5 poisons are compounded or dispensed or such place where pharmacists
6 practice the profession of pharmacy, or a pharmacy operated by the
7 Oklahoma Department of Veterans Affairs;

8 35. "Pharmacy technician", "technician", "Rx tech", or "tech"
9 means a person issued a Technician permit by the State Board of
10 Pharmacy to assist the pharmacist and perform nonjudgmental,
11 technical, manipulative, non-discretionary functions in the
12 prescription department under the immediate and direct supervision
13 of a pharmacist;

14 36. "Poison" means any substance which when introduced into the
15 body, either directly or by absorption, produces violent, morbid or
16 fatal changes, or which destroys living tissue with which such
17 substance comes into contact;

18 37. "Practice of pharmacy" means:

- 19 a. the interpretation and evaluation of prescription
20 orders,
21 b. the compounding, dispensing, administering and
22 labeling of drugs and devices, except labeling by a
23 manufacturer, repackager or distributor of
24

1 nonprescription drugs and commercially packaged legend
2 drugs and devices,

3 c. the participation in drug selection and drug
4 utilization reviews,

5 d. the proper and safe storage of drugs and devices and
6 the maintenance of proper records thereof,

7 e. the responsibility for advising by counseling and
8 providing information, where professionally necessary
9 or where regulated, of therapeutic values, content,
10 hazards and use of drugs and devices,

11 f. the offering or performing of those acts, services,
12 operations or transactions necessary in the conduct,
13 operation, management and control of a pharmacy, or

14 g. the provision of those acts or services that are
15 necessary to provide pharmaceutical care;

16 38. "Preparation" means an article which may or may not contain
17 sterile products compounded in a licensed pharmacy pursuant to the
18 order of a licensed prescriber;

19 39. "Prescriber" means a person licensed in this state who is
20 authorized to prescribe dangerous drugs within the scope of practice
21 of the person's profession;

22 40. "Prescription" means and includes any order for drug or
23 medical supplies written or signed, or transmitted by word of mouth,
24 telephone or other means of communication:

- 1 a. by a licensed practitioner,
- 2 b. under the supervision of an Oklahoma licensed
- 3 practitioner, an Oklahoma licensed advanced practice
- 4 registered nurse or an Oklahoma licensed physician
- 5 assistant, or
- 6 c. by an Oklahoma licensed wholesaler or distributor as
- 7 authorized in Section ~~353.29~~ 353.29.1 of this title;

8 41. "Product" means a prescription drug in a finished dosage

9 form for administration to a patient without substantial further

10 manufacturing, such as capsules, tablets, and lyophilized products

11 before reconstitution. "Product" does not include blood components

12 intended for transfusion, radioactive drugs or biologics and medical

13 gas;

14 42. "Repackager", including "virtual repackager", means a

15 person who owns or operates an establishment that repacks and

16 relabels a product or package for further sale or distribution

17 without further transaction;

18 43. "Sterile drug" means a drug that is intended for parental

19 administration, an ophthalmic or oral inhalation drug in aqueous

20 format, or a drug that is required to be sterile under state and

21 federal law;

22 44. "Supervising physician" means an individual holding a

23 current license to practice as a physician from the State Board of

24 Medical Licensure and Supervision, pursuant to the provisions of the

1 Oklahoma Allopathic Medical and Surgical Licensure and Supervision
2 Act, or the State Board of Osteopathic Examiners, pursuant to the
3 provisions of the Oklahoma Osteopathic Medicine Act, who supervises
4 an advanced practice registered nurse as defined in Section 567.3a
5 of this title, and who is not in training as an intern, resident, or
6 fellow. To be eligible to supervise an advanced practice registered
7 nurse, such physician shall remain in compliance with the rules
8 promulgated by the State Board of Medical Licensure and Supervision
9 or the State Board of Osteopathic Examiners;

10 45. "Supportive personnel" means technicians and auxiliary
11 supportive persons who are regularly paid employees of a pharmacy
12 who work and perform tasks in the pharmacy as authorized by Section
13 ~~353.19~~ 353.18A of this title;

14 46. "Third-party logistics provider", including "virtual third-
15 party logistics provider" means an entity that provides or
16 coordinates warehousing, or other logistics services of a product in
17 interstate commerce on behalf of a manufacturer, wholesale
18 distributor, or dispenser of a product but does not take ownership
19 of the product, nor have responsibility to direct the sale or
20 disposition of the product. For the purposes of this paragraph,
21 "third-party logistics provider" does not include shippers and the
22 United States Postal Service; and

23 47. "Wholesale distributor", including "virtual wholesale
24 distributor" means a person other than a manufacturer, a

1 manufacturer's co-licensed partner, a third-party logistics
2 provider, or repackager engaged in wholesale distribution as defined
3 by 21 U.S.C. 353(e) (4) as amended by the Drug Supply Chain Security
4 Act.

5 SECTION 2. AMENDATORY 59 O.S. 2011, Section 353.11, as
6 amended by Section 7, Chapter 230, O.S.L. 2015 (59 O.S. Supp. 2015,
7 Section 353.11), is amended to read as follows:

8 Section 353.11 A. 1. Every licensed pharmacist who desires to
9 continue in the profession of pharmacy in this state shall, on or
10 before the expiration date of the license, complete a renewal form
11 and remit to the State Board of Pharmacy a renewal fee to be fixed
12 by the Board. Upon compliance with the provisions of the Oklahoma
13 Pharmacy Act and payment of such renewal fee by a licensee in good
14 standing with the Board, a renewal certificate of licensure shall be
15 issued.

16 2. Every licensed pharmacist who fails to complete a renewal
17 form and remit the required renewal fee to the Board by the
18 fifteenth day after the expiration of the license shall pay a late
19 fee to be fixed by the Board.

20 B. If any pharmacist fails or neglects to procure the renewal
21 of his or her license, as herein required, the Board may, after the
22 expiration of thirty (30) days following the issue of the notice,
23 deprive the person of his or her license and all other privileges
24 conferred by the Oklahoma Pharmacy Act.

1 C. In order to regain licensure, the pharmacist shall apply in
2 writing to the Board requesting reinstatement. The pharmacist shall
3 pay ~~back~~ all back fees and provide proof of having obtained all
4 delinquent continuing education plus an additional fifteen (15)
5 hours of continuing education. The Board may require the pharmacist
6 to appear before the Board at a regular meeting. The Board may
7 require evidence of competency through examination or impose other
8 requirements for reinstatement.

9 SECTION 3. AMENDATORY Section 8, Chapter 230, O.S.L.
10 2015 (59 O.S. Supp. 2015, Section 353.11a), is amended to read as
11 follows:

12 Section 353.11a A. No annual renewal certificate shall be
13 issued to a pharmacist until such pharmacist has submitted proof to
14 the State Board of Pharmacy that the pharmacist has satisfactorily
15 completed no less than fifteen (15) clock hours of an accredited or
16 Board-approved program of continuing professional education during
17 the previous calendar year.

18 B. The Board may grant alternate methods of obtaining
19 continuing education hours to a pharmacist who meets all necessary
20 requirements for licensure except the continuing education
21 requirements.

22 C. 1. Any pharmacist who does not meet the requirements for
23 continuing education may obtain an inactive renewal certificate of
24 licensure.

1 2. The holder of an inactive renewal certificate of licensure
2 shall not engage in the practice of pharmacy in this state.

3 3. The holder of an inactive renewal certificate of licensure
4 may apply to the Board to ~~the~~ be removed from inactive status.

5 SECTION 4. AMENDATORY 59 O.S. 2011, Section 353.18, as
6 amended by Section 11, Chapter 230, O.S.L. 2015 (59 O.S. Supp. 2015,
7 Section 353.18), is amended to read as follows:

8 Section 353.18 A. 1. It shall be unlawful for any person,
9 including, but not limited to, Internet, website or online
10 pharmacies, to sell at retail or to offer for sale, dangerous drugs,
11 medicines, chemicals or poisons for the treatment of disease,
12 excluding agricultural chemicals and drugs, or to accept
13 prescriptions for same, without first procuring a license from the
14 State Board of Pharmacy. This licensure requirement applies whether
15 such sale, offer for sale or acceptance of prescriptions occurs in
16 this state, or such sale, offer for sale, or acceptance of
17 prescription occurs out of state and the dangerous drug, medicine,
18 chemical or poison is to be delivered, distributed or dispensed to
19 patients or customers in this state.

20 2. A pharmacy license shall be issued to such person as the
21 Board shall deem qualified upon evidence satisfactory to the Board
22 that:
23
24

- 1 a. the place for which the license is sought will be
2 conducted in full compliance with the law and the
3 rules of the Board,
4 b. the location and physical characteristics of the place
5 are reasonably consistent with the maintenance of
6 professional surroundings and constitute no known
7 danger to the public health and safety,
8 c. the place will be under the management and control of
9 a licensed pharmacist or pharmacist-in-charge who
10 shall be licensed as a pharmacist in Oklahoma, and
11 d. a licensed pharmacist shall be present and on duty at
12 all business hours; provided, however, the provisions
13 of this subparagraph shall not apply to hospital drug
14 rooms.

15 3. a. An application for an initial or renewal license
16 issued pursuant to the provisions of this subsection
17 shall:

18 (1) be submitted to the Board in writing,

19 (2) contain the name or names of persons owning the
20 pharmacy, and

21 (3) provide other such information deemed relevant by
22 the Board.

23 b. An application for an initial or renewal license shall
24 be accompanied by a licensing fee not to exceed Three

1 Hundred Dollars (\$300.00) for each period of one (1)
2 year. Prior to opening for business, all applicants
3 for an initial license or permit shall be inspected.
4 An initial licensure applicant shall pay an inspection
5 fee not to exceed Two Hundred Dollars (\$200.00);
6 provided, however, that no charge shall be made for
7 the licensing of any Federal Veterans Hospital in the
8 State of Oklahoma. Non-resident pharmacies shall
9 reimburse the Board for any actual expenses incurred
10 for inspections.

11 c. A license issued pursuant to the provisions of this
12 subsection shall be valid for a period set by the
13 Board and shall contain the name of the licensee and
14 the address of the place at which such business shall
15 be conducted.

16 4. A retail pharmacy that prepares sterile drugs shall obtain a
17 pharmacy license, and shall also obtain a sterile compounding permit
18 at a fee set by the Board, not to exceed Seventy-five Dollars
19 (\$75.00). Such pharmacy shall meet requirements set by the Board by
20 rule for sterile compounding permits.

21 5. An outsourcing facility desiring to dispense prescriptions
22 to patients must additionally license and meet the requirements of a
23 pharmacy.

1 B. 1. It shall be unlawful for any person to manufacture,
2 repackage, distribute, outsource, warehouse or ~~have an outsourcing~~
3 ~~facility, be a~~ third-party logistics provider, ~~or warehouse of~~ any
4 dangerous drugs, medicines, medical gases, chemicals, or poisons for
5 the treatment of disease, excluding agricultural chemicals ~~and~~
6 ~~drugs, or to sell or offer to sale at retail or wholesale medical~~
7 ~~gases~~ without first procuring a license from the Board. It shall be
8 unlawful to sell or offer for sale at retail or wholesale dangerous
9 drugs, medicines, medical gases, chemicals or poisons without first
10 procuring a license from the Board. This licensure requirement
11 shall apply when the manufacturing, repackaging, distributing,
12 outsourcing, warehousing, outsourcing facility or third-party
13 ~~logistics provider or facility sale or offer to sell~~ or provision of
14 third-party logistics occurs in this state or ~~when such dangerous~~
15 ~~drugs, medicines, chemicals or poisons are sold or offered to be~~
16 ~~sold~~ out of state for delivery, distribution, or dispensing to
17 patients or customers in this state.

18 2. A license shall be issued to such person as the Board shall
19 deem qualified upon satisfactory evidence to the Board that:

- 20 a. the place for which the license is sought will be
21 conducted in full compliance with the laws of this
22 state and the administrative rules of the Board,
- 23 b. the location and physical characteristics of the place
24 of business are reasonably consistent with the

1 maintenance of professional surroundings and
2 constitute no known danger to public health and
3 safety,

4 c. the place shall be under the management and control of
5 such persons as may be approved by the Board after a
6 review and determination of the persons'

7 qualifications, and

8 d. an outsourcing facility shall designate in writing on
9 a Board-approved form a person to serve as the
10 pharmacist-in-charge who is a pharmacist licensed by
11 the Board_{7.}

12 3. a. An application for an initial or renewal license
13 issued pursuant to the provisions of this subsection
14 shall:

15 (1) be submitted to the Board in writing,

16 (2) contain the name or names of the owners or the
17 applicants, and

18 (3) provide such other information deemed relevant by
19 the Board_{7.}

20 b. An application for an initial or renewal license shall
21 be accompanied by a licensing fee not to exceed Three
22 Hundred Dollars (\$300.00) for each period of one (1)
23 year. Prior to opening for business, all applicants
24 for initial or renewal license shall be inspected. An

1 initial licensure applicant shall pay an inspection
2 fee not to exceed Two Hundred Dollars (\$200.00). Non-
3 resident applicants shall reimburse the Board for any
4 actual expenses incurred for inspections.

5 c. A license issued pursuant to the provisions of this
6 subsection shall contain the name of the licensee and
7 the address of the place at which such business shall
8 be conducted and shall be valid for a period of time
9 set by the Board.

10 C. A licensee or permit holder who, pursuant to the provisions
11 of this section, fails to complete an application for a renewal
12 license or permit by the fifteenth day after the expiration of the
13 license or permit shall pay a late fee to be fixed by the Board.

14 D. 1. The Board shall promulgate rules regarding the issuance
15 and renewal of licenses and permits pursuant to the Oklahoma
16 Pharmacy Act which shall include, but need not be limited to
17 provisions for new or renewal application requirements for its
18 licensees and permit holders. Requirements for new and renewal
19 applications may include, but need not be limited to, the following:

- 20 a. type of ownership, whether individual, partnership,
21 limited liability company or corporation,
22 b. names and addresses of principal owners or officers
23 and their Social Security numbers, including
24 applicant's full name, all trade or business names

1 used, full business address, telephone numbers, and
2 email addresses,

3 c. names of designated representatives and facility
4 managers and their Social Security numbers and dates
5 of birth,

6 d. evidence of a criminal background check and
7 fingerprinting of the applicant, if a person, and all
8 of the applicant's designated representatives and
9 facility managers,

10 e. a copy of the license from the applicant's home state,
11 and if applicable, from the federal government,

12 f. bond requirements, and

13 g. any other information deemed by the Board to be
14 necessary to protect the public health and safety.

15 2. The Board shall be authorized to use an outside agency, such
16 as the National Association of Boards of Pharmacy (NABP) or the
17 Verified-Accredited Wholesale Distributors (VAWD), to accredit
18 wholesale distributors and repackagers.

19 E. The Oklahoma Pharmacy Act shall not be construed to prevent
20 the sale of nonprescription drugs in original manufacturer packages
21 by any merchant or dealer.

22 SECTION 5. AMENDATORY 59 O.S. 2011, Section 353.24, as
23 amended by Section 16, Chapter 230, O.S.L. 2015 (59 O.S. Supp. 2015,
24 Section 353.24), is amended to read as follows:

1 Section 353.24 A. It shall be unlawful for any licensee or
2 other person to:

3 1. Forge or increase the quantity of drug in any prescription,
4 or to present a prescription bearing forged, fictitious or altered
5 information or to possess any drug secured by such forged,
6 fictitious or altered prescription;

7 2. Sell, offer for sale, barter or give away any unused
8 quantity of drugs obtained by prescription, except through a program
9 pursuant to the Utilization of Unused Prescription Medications Act
10 or as otherwise provided by the State Board of Pharmacy;

11 3. Sell, offer for sale, barter or give away any drugs damaged
12 by fire, water, or other causes without first obtaining the written
13 approval of the Board or the State Department of Health;

14 4. ~~Enter into any arrangement whereby prescription orders are~~
15 ~~received, or prescriptions are delivered at a place other than the~~
16 ~~pharmacy in which they are filled, compounded or dispensed. No~~
17 person, firm or business establishment shall offer to the public, in
18 any manner, their services as a "pick-up station" or intermediary
19 for the purpose of having prescriptions filled or delivered, whether
20 for profit or gratuitously. Nor may the owner of any pharmacy or
21 drug store authorize any person, firm or business establishment to
22 act for them in this manner with these exceptions:

23 a. patient-specific filled prescriptions may be delivered
24 or shipped to a prescriber's clinic for pick-up by

1 those patients who the prescriber has individually
2 determined and documented do not have a permanent or
3 secure mailing address,

4 b. patient-specific filled prescriptions for drugs which
5 require special handling written by a prescriber may
6 be delivered or shipped to the prescriber's clinic for
7 administration or pick-up at the prescriber's office,

8 c. patient-specific filled prescriptions, including
9 sterile compounded drugs, may be delivered or shipped
10 to a prescriber's clinic where they shall be
11 administered,

12 d. patient-specific filled prescriptions for patients
13 under Medicare and/or Medicaid for End Stage Renal
14 Disease (ESRD) may be delivered or shipped to a
15 prescriber's clinic for administration or final
16 delivery to the patient, or

17 e. patient-specific filled prescriptions for
18 radiopharmaceuticals may be delivered or shipped to a
19 prescriber's clinic for administration or pick-up.

20 However, nothing in this paragraph shall prevent a pharmacist or
21 an employee of the pharmacy from personally receiving a prescription
22 or delivering a legally filled prescription to a residence, office
23 or place of employment of the patient for whom the prescription was
24 written. Provided further, the provisions of this paragraph shall

1 not apply to any Department of Mental Health and Substance Abuse
2 Services employee or any person whose facility contracts with the
3 Department of Mental Health and Substances Abuse Services whose
4 possession of any dangerous drug, as defined in Section 353.1 of
5 this title, is for the purpose of delivery of a mental health
6 consumer's medicine to the consumer's home or residence. Nothing in
7 this paragraph shall prevent veterinary prescription drugs from
8 being shipped directly from an Oklahoma licensed wholesaler or
9 distributor registered with the Oklahoma Board of Veterinary Medical
10 Examiners to a client; provided, such drugs may be dispensed only on
11 prescription of a licensed veterinarian and only when an existing
12 veterinary-client-patient relationship exists;

13 5. Sell, offer for sale or barter or buy any professional
14 samples except through a program pursuant to the Utilization of
15 Unused Prescription Medications Act;

16 6. Refuse to permit or otherwise prevent members of the Board
17 or such representatives thereof from entering and inspecting any and
18 all places, including premises, vehicles, equipment, contents, and
19 records, where drugs, medicine, chemicals or poisons are stored,
20 sold, vended, given away, compounded, dispensed, repackaged,
21 transported, or manufactured;

22 7. Interfere, refuse to participate in, impede or otherwise
23 obstruct any inspection, investigation or disciplinary proceeding
24 authorized by the Oklahoma Pharmacy Act;

1 8. Possess dangerous drugs without a valid prescription or a
2 valid license to possess such drugs; provided, however, this
3 provision shall not apply to any Department of Mental Health and
4 Substance Abuse Services employee or any person whose facility
5 contracts with the Department of Mental Health and Substances Abuse
6 Services whose possession of any dangerous drug, as defined in
7 Section 353.1 of this title, is for the purpose of delivery of a
8 mental health consumer's medicine to the consumer's home or
9 residence;

10 9. Fail to establish and maintain effective controls against
11 the diversion of drugs for any other purpose than legitimate
12 medical, scientific or industrial uses as provided by state, ~~and~~
13 federal, and local law;

14 10. Fail to have a written drug diversion detection and
15 prevention policy;

16 11. Possess, sell, offer for sale, barter or give away any
17 quantity of dangerous drugs not listed as a scheduled drug pursuant
18 to Sections 2-201 through 2-212 of Title 63 of the Oklahoma Statutes
19 when obtained by prescription bearing forged, fictitious or altered
20 information.

21 a. A first violation of this section shall constitute a
22 misdemeanor and upon conviction shall be punishable by
23 imprisonment in the county jail for a term not more
24

1 than one (1) year and a fine in an amount not more
2 than One Thousand Dollars (\$1,000.00).

3 b. A second violation of this section shall constitute a
4 felony and upon conviction shall be punishable by
5 imprisonment in the Department of Corrections for a
6 term not exceeding five (5) years and a fine in an
7 amount not more than Two Thousand Dollars (\$2,000.00);

8 12. Violate a Board order or agreed order;

9 13. Compromise the security of licensure examination materials;

10 or

11 14. Fail to notify the Board, in writing, within ten (10) days
12 of a licensee or permit holder's address change.

13 B. 1. It shall be unlawful for any person other than a
14 licensed pharmacist or physician to certify a prescription before
15 delivery to the patient or the patient's representative or
16 caregiver.

17 2. It shall be unlawful for any person to institute or manage a
18 pharmacy unless such person is a licensed pharmacist or has placed a
19 licensed pharmacist in charge of such pharmacy7.

20 3. No licensed pharmacist shall manage, supervise or be in
21 charge of more than one pharmacy.

22 4. No pharmacist being requested to sell, furnish or compound
23 any drug, medicine, chemical or other pharmaceutical preparation, by
24 prescription or otherwise, shall substitute or cause to be

1 substituted for it, without authority of the prescriber ~~of~~ or
2 purchaser, any like drug, medicine, chemical or pharmaceutical
3 preparation.

4 5. No pharmacy, pharmacist-in-charge or other person shall
5 permit the practice of pharmacy except by a licensed pharmacist or
6 assistant pharmacist.

7 6. No person shall subvert the authority of the pharmacist-in-
8 charge of the pharmacy by impeding the management of the
9 prescription department to act in compliance with federal and state
10 law.

11 C. 1. It shall be unlawful for a pharmacy to resell dangerous
12 drugs to any wholesale distributor.

13 2. It shall be unlawful for a wholesale distributor to purchase
14 drugs from a pharmacy.

15 SECTION 6. AMENDATORY 59 O.S. 2011, Section 353.26, as
16 amended by Section 17, Chapter 230, O.S.L. 2015 (59 O.S. Supp. 2015,
17 Section 353.26), is amended to read as follows:

18 Section 353.26 A. The State Board of Pharmacy may:

19 1. Revoke permanently or suspend any certificate, license or
20 permit issued pursuant to the Oklahoma Pharmacy Act or reprimand or
21 place on probation any holder of a certificate, license, or permit
22 who:

23 a. violates any provision of the Oklahoma Pharmacy Act or
24 any other applicable state or federal law,

- b. violates any of the provisions of the Uniform Controlled Dangerous Substances Act,
- c. has been convicted of a felony or has pleaded guilty or no contest to a felony,
- d. engages in the practice of pharmacy while incapacitated or abuses intoxicating liquors or other chemical substances,
- e. conducts himself or herself in a manner likely to lower public esteem for the profession of pharmacy,
- f. has been disciplined by another State Board of Pharmacy or by another state or federal entity,
- g. has been legally adjudged to be not mentally competent, or
- h. exercises conduct and habits inconsistent with the rules of professional conduct established by the Board; and

2. Levy administrative fines not to exceed Three Thousand Dollars (\$3,000.00) for each count of which any holder of a certificate, license, or permit has been convicted in Board hearings.

B. 1. The Board, its employees, or other agents of the Board shall keep confidential information obtained during an investigation into violations of the Oklahoma Pharmacy Act; provided, however, such information may be introduced by the state in administrative

1 proceedings before the Board and the information then becomes a
2 public record.

3 To ensure the confidentiality of such information obtained
4 during the investigation but not introduced in administrative
5 proceedings, this information shall not be deemed to be a record as
6 that term is defined in the Oklahoma Open Records Act, nor shall the
7 information be subject to subpoena or discovery in any civil or
8 criminal proceedings, except that the Board may give such
9 information to law enforcement and other state agencies as necessary
10 and appropriate in the discharge of the duties of that agency and
11 only under circumstances that ensure against unauthorized access to
12 the information.

13 2. The respondent may acquire information obtained during an
14 investigation, unless the disclosure of the information is otherwise
15 prohibited, except for the investigative report, if the respondent
16 signs a protective order whereby the respondent agrees to use the
17 information solely for the purpose of defense in the Board
18 proceeding and in any appeal therefrom and agrees not to otherwise
19 disclose the information.

20 C. 1. The Board shall mail by certified mail to respondent at
21 the last address provided by respondent to the Board, postmarked at
22 least ten (10) days before the hearing, the sworn complaint filed
23 with its Executive Director against respondent and notice of the
24 date and place of a hearing thereon. Alternatively, at least ten

1 (10) days before the hearing, the Board may serve respondent
2 personally by any person appointed to make service by the Executive
3 Director of the Board and in any manner authorized by the law of
4 this state for the personal service of summonses in proceedings in a
5 state court. Such service shall be effective upon the personal
6 service or mailing of the complaint and notice, and shall constitute
7 good service. If the Board finds that the allegations of the
8 complaint are supported by the evidence rendered at the hearing, the
9 Board is hereby authorized and empowered to, by written order,
10 revoke permanently or suspend for a designated period, the
11 certificate, license or permit of the respondent and/or reprimand,
12 place on probation and/or fine the respondent.

13 2. A person whose certificate, license, or permit has been
14 revoked or suspended or who has been reprimanded or placed on
15 probation or fined may appeal such Board order pursuant to the
16 Administrative Procedures Act.

17 3. The Board's order shall constitute a judgment and may be
18 entered on the judgment docket of the district court in a county in
19 which the respondent has property and ~~execution~~ may be executed
20 thereon in the same manner as any other judgment of a court of
21 record, unless the fine is paid within thirty (30) days after the
22 appeal time has run.

23 D. A person, other than a pharmacy technician, whose license or
24 permit has been suspended by the Board or by operation of law shall

1 pay a reinstatement fee not to exceed One Hundred Fifty Dollars
2 (\$150.00) as a condition of reinstatement of the license.

3 SECTION 7. AMENDATORY Section 1, Chapter 263, O.S.L.
4 2014 (59 O.S. Supp. 2015, Section 357), is amended to read as
5 follows:

6 Section 357. As used in this act:

7 1. "Covered entity" means a nonprofit hospital or medical
8 service organization, insurer, health coverage plan or health
9 maintenance organization; a health program administered by the state
10 in the capacity of provider of health coverage; or an employer,
11 labor union, or other entity organized in the state that provides
12 health coverage to covered individuals who are employed or reside in
13 the state. This term does not include a health plan that provides
14 coverage only for accidental injury, specified disease, hospital
15 indemnity, disability income, or other limited benefit health
16 insurance policies and contracts that do not include prescription
17 drug coverage;

18 2. "Covered individual" means a member, participant, enrollee,
19 contract holder or policy holder or beneficiary of a covered entity
20 who is provided health coverage by the covered entity. A covered
21 individual includes any dependent or other person provided health
22 coverage through a policy, contract or plan for a covered
23 individual;

24 3. "Department" means the Oklahoma Insurance Department;

1 4. "Maximum allowable cost" or "MAC" means the list of drug
2 products delineating the maximum per-unit reimbursement for
3 multiple-source prescription drugs, medical product or device;

4 5. "Multisource drug product reimbursement" (reimbursement)
5 means the total amount paid to a pharmacy inclusive of any reduction
6 in payment to the pharmacy, excluding prescription dispense fees;

7 6. "Pharmacy benefits management" means a service provided to
8 covered entities to facilitate the provision of prescription drug
9 benefits to covered individuals within the state, including
10 negotiating pricing and other terms with drug manufacturers and
11 providers. Pharmacy benefits management may include any or all of
12 the following services:

- 13 a. claims processing, retail network management and
14 payment of claims to pharmacies for prescription drugs
15 dispensed to covered individuals,
- 16 b. clinical formulary development and management
17 services,
- 18 c. rebate contracting and administration,
- 19 d. certain patient compliance, therapeutic intervention
20 and generic substitution programs, or
- 21 e. disease management programs;

22 ~~6.~~ 7. "Pharmacy benefits manager" or "PBM" means a person,
23 business or other entity that performs pharmacy benefits management.
24 The term includes a person or entity acting for a PBM in a

1 contractual or employment relationship in the performance of
2 pharmacy benefits management for a managed care company, nonprofit
3 hospital, medical service organization, insurance company, third-
4 party payor, or a health program administered by an agency of this
5 state;

6 ~~7.~~ 8. "Plan sponsor" means the employers, insurance companies,
7 unions and health maintenance organizations or any other entity
8 responsible for establishing, maintaining, or administering a health
9 benefit plan on behalf of covered individuals; and

10 ~~8.~~ 9. "Provider" means a pharmacy licensed by the State Board
11 of Pharmacy, or an agent or representative of a pharmacy, including,
12 but not limited to, the pharmacy's contracting agent, which
13 dispenses prescription drugs or devices to covered individuals.

14 SECTION 8. AMENDATORY Section 4, Chapter 263, O.S.L.
15 2014 (59 O.S. Supp. 2015, Section 360), is amended to read as
16 follows:

17 Section 360. A. The pharmacy benefits manager shall, with
18 respect to contracts between a pharmacy benefits manager and a
19 provider:

20 1. Include in such contracts the sources utilized to determine
21 the maximum allowable cost (MAC) pricing of the pharmacy, update
22 ~~maximum allowable cost~~ MAC pricing at least every seven (7) calendar
23 days, and establish a process for providers to readily access the
24 MAC list specific to that provider;

1 2. In order to place a drug on the MAC list, ensure that the
2 drug is listed as "A" or "B" rated in the most recent version of the
3 FDA's Approved Drug Products with Therapeutic Equivalence
4 Evaluations, also known as the Orange Book, or has an "NR" or "NA"
5 rating or a similar rating by a nationally recognized reference, and
6 the drug is generally available for purchase by pharmacies in the
7 state from national or regional wholesalers and is not obsolete;

8 3. Ensure dispensing fees are not included in the calculation
9 of MAC price reimbursement to pharmacy providers;

10 4. Provide a reasonable administration appeals procedure to
11 allow a provider or a provider's representative to contest ~~maximum~~
12 ~~allowable cost rates~~ reimbursement amounts within ten (10) business
13 days of ~~prescription claim~~ the final adjusted payment date. The
14 pharmacy benefits manager must respond to a provider or provider's
15 representative who has contested a ~~maximum allowable cost rate~~
16 reimbursement amount through this procedure within ten (10) business
17 days. If a price update is warranted, the pharmacy benefits manager
18 shall make the change in the ~~MAC~~ reimbursement amount, permit the
19 challenging pharmacy to reverse and rebill the claim in question,
20 and make the ~~MAC~~ reimbursement amount change effective for each
21 similarly contracted Oklahoma provider; and

22 5. If the ~~MAC~~ reimbursement appeal is denied, the PBM shall
23 provide the reason for the denial, including the National Drug Code
24 number from national or regional wholesalers where the drug is

1 generally available for purchase by pharmacies in the state at or
2 below the PBM's ~~Maximum Allowable Cost~~ reimbursement.

3 B. The pharmacy benefits manager may not place a drug on a
4 ~~maximum allowable cost~~ MAC list, unless there are at least two
5 therapeutically equivalent, multiple-source drugs, or at least one
6 generic drug available from only one manufacturer, generally
7 available for purchase by network pharmacies from national or
8 regional wholesalers.

9 C. The pharmacy benefits manager shall not require
10 accreditation or licensing of providers other than by the State
11 Board of Pharmacy or other state or federal government entity.

12 SECTION 9. REPEALER 59 O.S. 2011, Sections 353.13,
13 353.29, 364 and 366, are hereby repealed.

14 SECTION 10. This act shall become effective November 1, 2016."

15 Passed the House of Representatives the 19th day of April, 2016.

16

17

18 _____
Presiding Officer of the House of
19 Representatives

20 Passed the Senate the ____ day of _____, 2016.

21

22

23 _____
Presiding Officer of the Senate

24