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SENATE BILL

51ST LEGISLATURE - STATE OF NEW MEXICO - SECOND SESSION, 2014

INTRODUCED BY

FOR THE LEGISLATIVE HEALTH AND HUMAN SERVICES COMMITTEE

AN ACT

RELATING TO HEALTH CARE; AMENDING THE PHARMACY ACT TO PROVIDE FOR THE MEMBERSHIP OF TWO PHARMACY TECHNICIANS ON THE BOARD OF PHARMACY; PROVIDING A NEW DEFINITION OF THE TERM "PHARMACY TECHNICIAN".

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF NEW MEXICO:

SECTION 1. Section 61-11-2 NMSA 1978 (being Laws 1969, Chapter 29, Section 2, as amended) is amended to read:

"61-11-2. DEFINITIONS.--As used in the Pharmacy Act:

A. "administer" means the direct application of a drug to the body of a patient or research subject by injection, inhalation, ingestion or any other means as a result of an order of a licensed practitioner;

B. "board" means the board of pharmacy;

C. "compounding" means preparing, mixing,

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1 assembling, packaging or labeling a drug or device as the
2 result of a licensed practitioner's prescription or for the
3 purpose of, or as an incident to, research, teaching or
4 chemical analysis and not for sale or dispensing.

5 "Compounding" also includes preparing drugs or devices in
6 anticipation of a prescription based on routine, regularly
7 observed prescribing patterns;

8 D. "confidential information" means information in
9 the patient's pharmacy records accessed, maintained by or
10 transmitted to the pharmacist or communicated to the patient as
11 part of patient counseling and may be released only to the
12 patient or as the patient directs; or to those licensed
13 practitioners and other authorized health care professionals as
14 defined by regulation of the board when, in the pharmacist's
15 professional judgment, such release is necessary to protect the
16 patient's health and well-being; or to [~~such~~] other persons
17 authorized by law to receive [~~such~~] the information, regardless
18 of whether [~~such~~] the information is on paper, preserved on
19 microfilm or stored on electronic media;

20 E. "consulting pharmacist" means a pharmacist whose
21 services are engaged on a routine basis by a hospital or other
22 health care facility and who is responsible for the
23 distribution, receipt and storage of drugs according to the
24 state and federal regulations;

25 F. "custodial care facility" means a nursing home,

1 retirement care, mental care or other facility that provides
2 extended health care;

3 G. "dangerous drug" means a drug that is required
4 by an applicable federal or state law or rule to be dispensed
5 pursuant to a prescription or is restricted to use by licensed
6 practitioners; or that is required by federal law to be labeled
7 with any of the following statements prior to being dispensed
8 or delivered:

9 (1) "Caution: federal law prohibits
10 dispensing without prescription.";

11 (2) "Caution: federal law restricts this drug
12 to use by or on the order of a licensed veterinarian."; or

13 (3) "RX only";

14 H. "device" means an instrument, apparatus,
15 implement, machine, contrivance, implant or similar or related
16 article, including a component part or accessory, that is
17 required by federal law to bear the label, "Caution: federal
18 or state law requires dispensing by or on the order of a
19 physician.";

20 I. "dispense" means the evaluation and
21 implementation of a prescription, including the preparation and
22 delivery of a drug or device to a patient or patient's agent in
23 a suitable container appropriately labeled for subsequent
24 administration to or use by a patient;

25 J. "distribute" means the delivery of a drug or

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1 device other than by administering or dispensing;

2 K. "drug" means:

3 (1) an article recognized as a drug in [~~any~~]
4 an official compendium or its supplement that is designated
5 from time to time by the board for use in the diagnosis, cure,
6 mitigation, treatment or prevention of disease in humans or
7 other animals;

8 (2) an article intended for use in the
9 diagnosis, cure, mitigation, treatment or prevention of
10 diseases in humans or other animals;

11 (3) an article, other than food, that affects
12 the structure or [~~any~~] a function of the body of humans or
13 other animals; and

14 (4) an article intended for use as a component
15 of an article described in Paragraph (1), (2) or (3) of this
16 subsection;

17 L. "drug regimen review" includes an evaluation of
18 a prescription and patient record for:

- 19 (1) known allergies;
- 20 (2) rational therapy contraindications;
- 21 (3) reasonable dose and route of
22 administration;
- 23 (4) reasonable directions for use;
- 24 (5) duplication of therapy;
- 25 (6) drug-drug interactions;

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1 (7) adverse drug reactions; and

2 (8) proper use and optimum therapeutic

3 outcomes;

4 M. "electronic transmission" means transmission of
5 information in electronic form or the transmission of the exact
6 visual image of a document by way of electronic equipment;

7 N. "hospital" means an institution that is licensed
8 as a hospital by the department of health;

9 O. "labeling" means the process of preparing and
10 affixing a label to ~~[any]~~ a drug container exclusive of the
11 labeling by a manufacturer, packer or distributor of a
12 nonprescription drug or commercially packaged prescription drug
13 or device; and which label includes all information required by
14 federal or state law or regulations adopted pursuant to federal
15 or state law;

16 P. "licensed practitioner" means a person engaged
17 in a profession licensed by ~~[any]~~ a state, territory or
18 possession of the United States who, within the limits of ~~[his]~~
19 the person's license, may lawfully prescribe, dispense or
20 administer drugs for the treatment of a patient's condition;

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

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

6 R. "nonprescription drugs" means nonnarcotic
7 medicines or drugs that may be sold without a prescription and
8 are prepackaged for use by a consumer and are labeled in
9 accordance with the laws and regulations of the state and
10 federal governments;

11 S. "nonresident pharmacy" means any pharmacy
12 located outside New Mexico that ships, mails or delivers, in
13 any manner, drugs into New Mexico;

14 T. "patient counseling" means the oral
15 communication by the pharmacist of information to a patient or
16 ~~[his]~~ the patient's agent or caregiver regarding proper use of
17 a drug or device;

18 U. "person" means an individual, corporation,
19 partnership, association or other legal entity;

20 V. "pharmaceutical care" means the provision of
21 drug therapy and other patient care services related to drug
22 therapy intended to achieve definite outcomes that improve a
23 patient's quality of life, including identifying potential and
24 actual drug-related problems, resolving actual drug-related
25 problems and preventing potential drug-related problems;

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1 W. "pharmacist" means a person who is licensed as a
2 pharmacist in this state;

3 X. "pharmacist in charge" means a pharmacist who
4 accepts responsibility for the operation of a pharmacy in
5 conformance with all laws and rules pertinent to the practice
6 of pharmacy and the distribution of drugs and who is personally
7 in full and actual charge of the pharmacy and its personnel;

8 Y. "pharmacy" means a licensed place of business
9 where drugs are compounded or dispensed and pharmaceutical care
10 is provided;

11 Z. "pharmacist intern" means a person licensed by
12 the board to train under a pharmacist;

13 AA. "pharmacy technician" means a person who is
14 registered with the board to perform repetitive tasks not
15 requiring the professional judgment of a pharmacist, including
16 technical activities associated with the preparation and
17 distribution of medication;

18 BB. "practice of pharmacy" means the evaluation and
19 implementation of a lawful order of a licensed practitioner;
20 the dispensing of prescriptions; the participation in drug and
21 device selection or drug administration that has been ordered
22 by a licensed practitioner, drug regimen reviews and drug or
23 drug-related research; the administering or prescribing of
24 dangerous drug therapy; the provision of patient counseling and
25 pharmaceutical care; the responsibility for compounding and

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1 labeling of drugs and devices; the proper and safe storage of
2 drugs and devices; and the maintenance of proper records;

3 CC. "prescription" means an order given
4 individually for the person for whom prescribed, either
5 directly from a licensed practitioner or [~~his~~] the licensed
6 practitioner's agent to the pharmacist, including electronic
7 transmission or indirectly by means of a written order signed
8 by the prescriber, that bears the name and address of the
9 prescriber, [~~his~~] the prescriber's license classification, the
10 name and address of the patient, the name and quantity of the
11 drug prescribed, directions for use and the date of issue;

12 DD. "significant adverse drug event" means a drug-
13 related incident that may result in harm, injury or death to
14 the patient; and

15 EE. "wholesale drug distributor" means a person
16 engaged in the wholesale distribution of prescription drugs,
17 including manufacturers, repackers, own-label distributors,
18 private-label distributors, jobbers, brokers, manufacturer's
19 warehouses, distributor's warehouses, chain drug warehouses,
20 wholesale drug warehouses, independent wholesale drug traders
21 and retail pharmacies that conduct wholesale distribution."

22 SECTION 2. Section 61-11-4 NMSA 1978 (being Laws 1969,
23 Chapter 29, Section 3, as amended) is amended to read:

24 "61-11-4. BOARD CREATED--MEMBERS--QUALIFICATIONS--
25 TERMS--VACANCIES--REMOVAL.--

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1 A. There is created the "board of pharmacy". The
2 board shall be administratively attached to the regulation and
3 licensing department. The board consists of [~~nine~~] eleven
4 members, each of whom shall be a citizen of the United States
5 and a resident of New Mexico.

6 B. Five members shall be pharmacists appointed by
7 the governor for staggered terms of five years each from lists
8 submitted to the governor by the New Mexico pharmaceutical
9 association, which lists contain the names of two pharmacists
10 residing in each of the five pharmacy districts. Appointments
11 of pharmacist members shall be made for five years or less each
12 and made in such a manner that the term of one pharmacist
13 member expires on July 1 of each year. One pharmacist member
14 shall be appointed from each pharmacy district. A pharmacist
15 member of the board shall have been actively engaged in the
16 pharmaceutical profession in this state for at least three
17 years immediately prior to [~~his~~] appointment and shall have had
18 a minimum of eight years of practical experience as a
19 pharmacist. A vacancy shall be filled by appointment by the
20 governor for the unexpired term from lists submitted by the New
21 Mexico pharmaceutical association to the governor. Pharmacist
22 members shall reside in the district from which they are
23 appointed.

24 C. Two members shall be pharmacy technicians
25 appointed by the governor from a list submitted to the governor

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1 by the New Mexico pharmaceutical association. Initial
2 appointments of pharmacy technician members shall be made for
3 staggered terms of five years or less and made in such a manner
4 so that not more than one pharmacy technician's term shall
5 expire on July 1 of each year. A vacancy in a pharmacy
6 technician's term shall be filled by appointment by the
7 governor for the unexpired term.

8 ~~[G-]~~ D. Three members of the board shall be
9 appointed by the governor to represent the public. The public
10 members of the board shall not have been licensed as
11 pharmacists or have any significant financial interest, whether
12 direct or indirect, in the profession regulated. A vacancy in
13 a public member's term shall be filled by appointment by the
14 governor for the unexpired term. Initial appointments of
15 public members shall be made for staggered terms of five years
16 or less and made in such a manner that not more than two public
17 members' terms shall expire on July 1 of each year.

18 ~~[D-]~~ E. One member of the board shall be a
19 pharmacist appointed at large from a list submitted to the
20 governor by the New Mexico society of ~~[health systems]~~ health-
21 system pharmacists. The member shall be appointed by the
22 governor to a term of five years. A vacancy in the member's
23 term shall be filled by appointment by the governor for the
24 unexpired term from a list submitted to the governor by the New
25 Mexico society of ~~[health systems]~~ health-system pharmacists.

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1 ~~[E-]~~ F. There are created five pharmacy districts
2 as follows:

3 (1) northeast district, which shall be
4 composed of the counties of Colfax, Guadalupe, Harding, Los
5 Alamos, Mora, Quay, Rio Arriba, Sandoval, San Miguel, Santa Fe,
6 Taos, Tarrant and Union;

7 (2) northwest district, which shall be
8 composed of the counties of McKinley, San Juan, Valencia and
9 Cibola;

10 (3) central district, which shall be composed
11 of the county of Bernalillo;

12 (4) southeast district, which shall be
13 composed of the counties of Chaves, Curry, De Baca, Eddy, Lea
14 and Roosevelt; and

15 (5) southwest district, which shall be
16 composed of the counties of Catron, Dona Ana, Grant, Hidalgo,
17 Lincoln, Luna, Otero, Sierra and Socorro.

18 ~~[F-]~~ G. A board member shall not serve more than
19 two full terms, consecutive or otherwise.

20 ~~[G-]~~ H. A board member failing to attend three
21 consecutive regular meetings is automatically removed as a
22 member of the board.

23 ~~[H-]~~ I. The governor may remove a member of the
24 board for neglect of a duty required by law, for incompetency
25 or for unprofessional conduct and shall remove a board member

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who violates a provision of the Pharmacy Act."

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