

State of Arizona  
Senate  
Fifty-second Legislature  
Second Regular Session  
2016

# SENATE BILL 1112

AN ACT

AMENDING SECTIONS 32-1901, 32-1970 AND 32-1974, ARIZONA REVISED STATUTES;  
RELATING TO THE ARIZONA STATE BOARD OF PHARMACY.

(TEXT OF BILL BEGINS ON NEXT PAGE)

1 Be it enacted by the Legislature of the State of Arizona:

2 Section 1. Section 32-1901, Arizona Revised Statutes, is amended to  
3 read:

4 32-1901. Definitions

5 In this chapter, unless the context otherwise requires:

6 1. "Administer" means the direct application of a controlled  
7 substance, prescription-only drug, dangerous drug or narcotic drug, whether  
8 by injection, inhalation, ingestion or any other means, to the body of a  
9 patient or research subject by a practitioner or by the practitioner's  
10 authorized agent or the patient or research subject at the direction of the  
11 practitioner.

12 2. "Advertisement" means all representations disseminated in any  
13 manner or by any means, other than by labeling, for the purpose of inducing,  
14 or that are likely to induce, directly or indirectly, the purchase of drugs,  
15 devices, poisons or hazardous substances.

16 3. "Advisory letter" means a nondisciplinary letter to notify a  
17 licensee or permittee that either:

18 (a) While there is insufficient evidence to support disciplinary  
19 action, the board believes that continuation of the activities that led to  
20 the investigation may result in further board action against the licensee or  
21 permittee.

22 (b) The violation is a minor or technical violation that is not of  
23 sufficient merit to warrant disciplinary action.

24 (c) While the licensee or permittee has demonstrated substantial  
25 compliance through rehabilitation, remediation or reeducation that has  
26 mitigated the need for disciplinary action, the board believes that  
27 repetition of the activities that led to the investigation may result in  
28 further board action against the licensee or permittee.

29 4. "Antiseptic", if a drug is represented as such on its label, means  
30 a representation that it is a germicide, except in the case of a drug  
31 purporting to be, or represented as, an antiseptic for inhibitory use as a  
32 wet dressing, ointment or dusting powder or other use that involves prolonged  
33 contact with the body.

34 5. "Authorized officers of the law" means legally empowered peace  
35 officers, compliance officers of the ~~state~~ board of pharmacy and agents of  
36 the division of narcotics enforcement and criminal intelligence of the  
37 department of public safety.

38 6. "Board" or "board of pharmacy" means the Arizona state board of  
39 pharmacy.

40 7. "Color additive" means a material that either:

41 (a) Is any dye, pigment or other substance made by a process of  
42 synthesis or similar artifice, or extracted, isolated or otherwise derived,  
43 with or without intermediate or final change of identity, from any vegetable,  
44 animal, mineral or other source.

1 (b) If added or applied to a drug, or to the human body or any part of  
2 the human body, is capable of imparting color, except that color additive  
3 does not include any material that has been or may be exempted under the  
4 federal act. Color includes black, white and intermediate grays.

5 8. "Compounding" means the preparation, mixing, assembling, packaging  
6 or labeling of a drug by a pharmacist or an intern or pharmacy technician  
7 under the pharmacist's supervision, for the purpose of dispensing to a  
8 patient based on a valid prescription order. Compounding includes the  
9 preparation of drugs in anticipation of prescription orders prepared on  
10 routine, regularly observed prescribing patterns and the preparation of drugs  
11 as an incident to research, teaching or chemical analysis or for  
12 administration by a medical practitioner to the medical practitioner's  
13 patient and not for sale or dispensing. Compounding does not include the  
14 preparation of commercially available products from bulk compounds or the  
15 preparation of drugs for sale to pharmacies, practitioners or entities for  
16 the purpose of dispensing or distribution.

17 9. "Compressed medical gas distributor" means a person who holds a  
18 current permit issued by the board to distribute compressed medical gases  
19 pursuant to a compressed medical gas order to compressed medical gas  
20 suppliers and other entities that are registered, licensed or permitted to  
21 use, administer or distribute compressed medical gases.

22 10. "Compressed medical gas order" means an order for compressed  
23 medical gases that is issued by a medical practitioner.

24 11. "Compressed medical gas supplier" means a person who holds a  
25 current permit issued by the board to supply compressed medical gases  
26 pursuant to a compressed medical gas order and only to the consumer or the  
27 patient.

28 12. "Compressed medical gases" means gases and liquid oxygen that a  
29 compressed medical gas distributor or manufacturer has labeled in compliance  
30 with federal law.

31 13. "Controlled substance" means a drug, substance or immediate  
32 precursor THAT IS identified, defined or listed in title 36, chapter 27,  
33 article 2.

34 14. "Corrosive" means any substance that when it comes in contact with  
35 living tissue will cause destruction of tissue by chemical action.

36 15. "Counterfeit drug" means a drug that, or the container or labeling  
37 of which, without authorization, bears the trademark, trade name or other  
38 identifying mark, imprint, number or device, or any likeness of these, of a  
39 manufacturer, distributor or dispenser other than the person who in fact  
40 manufactured, distributed or dispensed that drug.

41 16. "Dangerous drug" has the same meaning prescribed in section  
42 13-3401.

43 17. "Decree of censure" means an official action that is taken by the  
44 board and that may include a requirement for restitution of fees to a patient  
45 or consumer.

1           18. "Deliver" or "delivery" means the actual, constructive or attempted  
2 transfer from one person to another whether or not there is an agency  
3 relationship.

4           19. "Deputy director" means a pharmacist who is employed by the board  
5 and selected by the executive director to perform duties as prescribed by the  
6 executive director.

7           20. "Device", except as used in paragraph 15 of this section, section  
8 32-1965, paragraph 4 and section 32-1967, subsection A, paragraph 15 and  
9 subsection C, means instruments, apparatus and contrivances, including their  
10 components, parts and accessories, including all such items under the federal  
11 act, intended either:

12           (a) For use in the diagnosis, cure, mitigation, treatment or  
13 prevention of disease in the human body or other animals.

14           (b) To affect the structure or any function of the human body or other  
15 animals.

16           21. "Direct supervision of a pharmacist" means the pharmacist is  
17 present. If relating to the sale of certain items, direct supervision of a  
18 pharmacist means that a pharmacist determines the legitimacy or advisability  
19 of a proposed purchase of those items.

20           22. "Director" means the director of the division of narcotics  
21 enforcement and criminal investigation of the department of public safety.

22           23. "Dispense" means to deliver to an ultimate user or research subject  
23 by or pursuant to the lawful order of a practitioner, including the  
24 prescribing, administering, packaging, labeling or compounding necessary to  
25 prepare for that delivery.

26           24. "Dispenser" means a practitioner who dispenses.

27           25. "Distribute" means to deliver, other than by administering or  
28 dispensing.

29           26. "Distributor" means a person who distributes.

30           27. "Drug" means:

31           (a) Articles recognized, or for which standards or specifications are  
32 prescribed, in the official compendium.

33           (b) Articles intended for use in the diagnosis, cure, mitigation,  
34 treatment or prevention of disease in the human body or other animals.

35           (c) Articles other than food intended to affect the structure or any  
36 function of the human body or other animals.

37           (d) Articles intended for use as a component of any articles specified  
38 in subdivision (a), (b) or (c) of this paragraph but does not include devices  
39 or their components, parts or accessories.

40           28. "Drug enforcement administration" means the drug enforcement  
41 administration of the United States department of justice or its successor  
42 agency.

43           29. "Drug or device manufacturing" means the production, preparation,  
44 propagation or processing of a drug or device, either directly or indirectly,  
45 by extraction from substances of natural origin or independently by means of

1 chemical synthesis and includes any packaging or repackaging of substances or  
2 labeling or relabeling of its container and the promotion and marketing of  
3 the same. Drug or device manufacturing does not include compounding.

4 30. "Economic poison" means any substance that alone, in chemical  
5 combination or in formulation with one or more other substances is a  
6 pesticide within the meaning of the laws of this state or the federal  
7 insecticide, fungicide and rodenticide act and that is used in the  
8 production, storage or transportation of raw agricultural commodities.

9 31. "Established name", with respect to a drug or ingredient of a drug,  
10 means any of the following:

11 (a) The applicable official name.

12 (b) If there is no such name and the drug or ingredient is an article  
13 recognized in an official compendium, the official title in an official  
14 compendium.

15 (c) If neither subdivision (a) nor (b) of this paragraph applies, the  
16 common or usual name of such drug.

17 32. "Executive director" means the executive director of the board of  
18 pharmacy.

19 33. "Federal act" means the federal laws and regulations that pertain  
20 to drugs, devices, poisons and hazardous substances and that are official at  
21 the time any drug, device, poison or hazardous substance is affected by this  
22 chapter.

23 34. "Full service wholesale permittee" means a permittee who may  
24 distribute prescription-only drugs and devices, controlled substances and  
25 over-the-counter drugs and devices to pharmacies or other legal outlets from  
26 a place devoted in whole or in part to wholesaling these items.

27 35. "Graduate intern" means a person who has graduated from a college,  
28 school or program of pharmacy approved by the board and who meets the  
29 qualifications and experience for a pharmacy intern as provided in section  
30 32-1923.

31 36. "Highly toxic" means any substance that falls within any of the  
32 following categories:

33 (a) Produces death within fourteen days in half or more than half of a  
34 group of ten or more laboratory white rats each weighing between two hundred  
35 and three hundred grams, at a single dose of fifty milligrams or less per  
36 kilogram of body weight, when orally administered.

37 (b) Produces death within fourteen days in half or more than half of a  
38 group of ten or more laboratory white rats each weighing between two hundred  
39 and three hundred grams, if inhaled continuously for a period of one hour or  
40 less at an atmospheric concentration of two hundred parts per million by  
41 volume or less of gas or vapor or two milligrams per liter by volume or less  
42 of mist or dust, provided the concentration is likely to be encountered by  
43 humans if the substance is used in any reasonably foreseeable manner.

44 (c) Produces death within fourteen days in half or more than half of a  
45 group of ten or more rabbits tested in a dosage of two hundred milligrams or

1 less per kilogram of body weight, if administered by continuous contact with  
2 the bare skin for twenty-four hours or less.

3 If the board finds that available data on human experience with any substance  
4 indicate results different from those obtained on animals in the dosages or  
5 concentrations prescribed in this paragraph, the human data shall take  
6 precedence.

7 37. "Hospital" means any institution for the care and treatment of the  
8 sick and injured that is approved and licensed as a hospital by the  
9 department of health services.

10 38. "Intern" means a pharmacy intern and a graduate intern.

11 39. "Internship" means the practical, experiential, hands-on training  
12 of a pharmacy intern under the supervision of a preceptor.

13 40. "Irritant" means any substance, other than a corrosive, that on  
14 immediate, prolonged or repeated contact with normal living tissue will  
15 induce a local inflammatory reaction.

16 41. "Jurisprudence examination" means a ~~board-approved~~ BOARD-APPROVED  
17 pharmacy law examination that is written and administered in cooperation with  
18 the national association of boards of pharmacy or another ~~board-approved~~  
19 BOARD-APPROVED pharmacy law examination.

20 42. "Label" means a display of written, printed or graphic matter on  
21 the immediate container of any article that, unless easily legible through  
22 the outside wrapper or container, also appears on the outside wrapper or  
23 container of the article's retail package. For the purposes of this  
24 paragraph, the immediate container does not include package liners.

25 43. "Labeling" means all labels and other written, printed or graphic  
26 matter either:

27 (a) On any article or any of its containers or wrappers.

28 (b) Accompanying that article.

29 44. "Letter of reprimand" means a disciplinary letter that is a public  
30 document issued by the board and that informs a licensee or permittee that  
31 the licensee's or permittee's conduct violates state or federal law and may  
32 require the board to monitor the licensee or permittee.

33 45. "Limited service pharmacy" means a pharmacy that is approved by the  
34 board to practice a limited segment of pharmacy as indicated by the permit  
35 issued by the board.

36 46. "Manufacture" or "manufacturer" means every person who prepares,  
37 derives, produces, compounds, processes, packages or repackages or labels any  
38 drug in a place, other than a pharmacy, devoted to manufacturing the drug.

39 47. "Marijuana" has the same meaning prescribed in section 13-3401.

40 48. "Medical practitioner" means any medical doctor, doctor of  
41 osteopathy, dentist, podiatrist, veterinarian or other person WHO IS licensed  
42 and authorized by law to use and prescribe drugs and devices for the  
43 treatment of sick and injured human beings or animals or for the diagnosis or  
44 prevention of sickness in human beings or animals in this state or any state,  
45 territory or district of the United States.

1           49. "Medication order" means a written or verbal order from a medical  
2 practitioner or that person's authorized agent to administer a drug or  
3 device.

4           50. "Narcotic drug" has the same meaning prescribed in section 13-3401.

5           51. "New drug" means either:

6           (a) Any drug the composition of which is such that the drug is not  
7 generally recognized among experts qualified by scientific training and  
8 experience to evaluate the safety and effectiveness of drugs as safe and  
9 effective for use under the conditions prescribed, recommended or suggested  
10 in the labeling.

11           (b) Any drug the composition of which is such that the drug, as a  
12 result of investigations to determine its safety and effectiveness for use  
13 under such conditions, has become so recognized, but that has not, other than  
14 in the investigations, been used to a material extent or for a material time  
15 under those conditions.

16           52. "Nonprescription drug" or "over-the-counter drug" means any  
17 nonnarcotic medicine or drug that may be sold without a prescription and is  
18 prepackaged and labeled for use by the consumer in accordance with the  
19 requirements of the laws of this state and federal law. Nonprescription drug  
20 does not include:

21           (a) A drug that is primarily advertised and promoted professionally to  
22 medical practitioners and pharmacists by manufacturers or primary  
23 distributors.

24           (b) A controlled substance.

25           (c) A drug that is required to bear a label that states "Rx only."

26           (d) A drug THAT IS intended for human use by hypodermic injection.

27           53. "Nonprescription drug wholesale permittee" means a permittee who  
28 may distribute only over-the-counter drugs and devices to pharmacies or other  
29 lawful outlets from a place devoted in whole or in part to wholesaling these  
30 items.

31           54. "Notice" means personal service or the mailing of a copy of the  
32 notice by certified mail addressed either to the person at the person's  
33 latest address of record in the board office or to the person's attorney.

34           55. "Official compendium" means the latest revision of the United  
35 States pharmacopeia and the national formulary or any current supplement.

36           56. "Other jurisdiction" means one of the other forty-nine states, the  
37 District of Columbia, the Commonwealth of Puerto Rico or a territory of the  
38 United States of America.

39           57. "Package" means a receptacle defined or described in the United  
40 States pharmacopeia and the national formulary as adopted by the board.

41           58. "Packaging" means the act or process of placing a drug item or  
42 device in a container for the purpose or intent of dispensing or distributing  
43 the item or device to another.

44           59. "Person" means an individual, partnership, corporation and  
45 association, and their duly authorized agents.

1           60. "Pharmaceutical care" means the provision of drug therapy and other  
2 pharmaceutical patient care services.

3           61. "Pharmacist" means an individual WHO IS currently licensed by the  
4 board to practice the profession of pharmacy in this state.

5           62. "Pharmacist in charge" means the pharmacist who is responsible to  
6 the board for a licensed establishment's compliance with the laws and  
7 administrative rules of this state and of the federal government pertaining  
8 to the practice of pharmacy, the manufacturing of drugs and the distribution  
9 of drugs and devices.

10          63. "Pharmacist licensure examination" means a ~~board-approved~~  
11 BOARD-APPROVED examination that is written and administered in cooperation  
12 with the national association of boards of pharmacy or any other ~~board~~  
13 ~~approved~~ BOARD-APPROVED pharmacist licensure examination.

14          64. "Pharmacy" means any place:

15           (a) Where drugs, devices, poisons or related hazardous substances are  
16 offered for sale at retail.

17           (b) In which the profession of pharmacy is practiced or where  
18 prescription orders are compounded and dispensed.

19           (c) That has displayed on it or in it the words "pharmacist,"  
20 "pharmaceutical chemist," "apothecary," "druggist," "pharmacy," "drugstore,"  
21 "drugs" or "drug sundries" or any of these words or combinations of these  
22 words, or words of similar import either in English or any other language, or  
23 that is advertised by any sign containing any of these words.

24           (d) Where the characteristic symbols of pharmacy or the characteristic  
25 prescription sign "Rx" is exhibited.

26           (e) Or a portion of any building or structure that is leased, used or  
27 controlled by the permittee to conduct the business authorized by the board  
28 at the address for which the permit was issued and that is enclosed and  
29 secured when a pharmacist is not in attendance.

30          65. "Pharmacy intern" means a person who has all of the qualifications  
31 and experience prescribed in section 32-1923.

32          66. "Pharmacy technician" means a person who is licensed pursuant to  
33 this chapter.

34          67. "Pharmacy technician trainee" means a person who is licensed  
35 pursuant to this chapter.

36          68. "Poison" or "hazardous substance" includes, but is not limited to,  
37 any of the following if intended and suitable for household use or use by  
38 children:

39           (a) Any substance that, according to standard works on medicine,  
40 pharmacology, pharmacognosy or toxicology, if applied to, introduced into or  
41 developed within the body in relatively small quantities by its inherent  
42 action uniformly produces serious bodily injury, disease or death.

43           (b) A toxic substance.

44           (c) A highly toxic substance.

45           (d) A corrosive substance.



1 (e) An irritant.

2 (f) A strong sensitizer.

3 (g) A mixture of any of the substances described in this paragraph, if  
4 the substance or mixture of substances may cause substantial personal injury  
5 or substantial illness during or as a proximate result of any customary or  
6 reasonably foreseeable handling or use, including reasonably foreseeable  
7 ingestion by children.

8 (h) A substance **THAT IS** designated by the board to be a poison or  
9 hazardous substance. This subdivision does not apply to radioactive  
10 substances, economic poisons subject to the federal insecticide, fungicide  
11 and rodenticide act or the state pesticide act, foods, drugs and cosmetics  
12 subject to state laws or the federal act or substances intended for use as  
13 fuels when stored in containers and used in the heating, cooking or  
14 refrigeration system of a house. This subdivision applies to any substance  
15 or article that is not itself an economic poison within the meaning of the  
16 federal insecticide, fungicide and rodenticide act or the state pesticide  
17 act, but that is a poison or hazardous substance within the meaning of this  
18 paragraph by reason of bearing or containing an economic poison or hazardous  
19 substance.

20 69. "Practice of pharmacy" means furnishing the following health care  
21 services as a medical professional:

22 (a) Interpreting, evaluating and dispensing prescription orders in the  
23 patient's best interests.

24 (b) Compounding drugs pursuant to or in anticipation of a prescription  
25 order.

26 (c) Labeling of drugs and devices in compliance with state and federal  
27 requirements.

28 (d) Participating in drug selection and drug utilization reviews, drug  
29 administration, drug or drug-related research and drug therapy monitoring or  
30 management.

31 (e) Providing patient counseling necessary to provide pharmaceutical  
32 care.

33 (f) Properly and safely storing drugs and devices in anticipation of  
34 dispensing.

35 (g) Maintaining required records of drugs and devices.

36 (h) Offering or performing of acts, services, operations or  
37 transactions necessary in the conduct, operation, management and control of a  
38 pharmacy.

39 ~~Implementing~~ **INITIATING**, monitoring and modifying drug therapy  
40 pursuant to a protocol-based drug therapy agreement with a provider as  
41 outlined in section 32-1970.

42 (j) Initiating and administering immunizations or vaccines pursuant to  
43 section 32-1974.

44 70. "Practitioner" means any physician, dentist, veterinarian,  
45 scientific investigator or other person who is licensed, registered or

1 otherwise permitted to distribute, dispense, conduct research with respect to  
2 or administer a controlled substance in the course of professional practice  
3 or research in this state, or any pharmacy, hospital or other institution  
4 that is licensed, registered or otherwise permitted to distribute, dispense,  
5 conduct research with respect to or administer a controlled substance in the  
6 course of professional practice or research in this state.

7 71. "Preceptor" means a pharmacist who is serving as the practical  
8 instructor of an intern and complies with section 32-1923.

9 72. "Precursor chemical" means a substance that is:

10 (a) The principal compound that is commonly used or that is produced  
11 primarily for use and that is an immediate chemical intermediary used or  
12 likely to be used in the manufacture of a controlled substance, the control  
13 of which is necessary to prevent, curtail or limit manufacture.

14 (b) Listed in section 13-3401, paragraph 26 or 27.

15 73. "Prescription" means either a prescription order or a prescription  
16 medication.

17 74. "Prescription medication" means any drug, including label and  
18 container according to context, that is dispensed pursuant to a prescription  
19 order.

20 75. "Prescription-only device" includes:

21 (a) Any device that is limited by the federal act to use under the  
22 supervision of a medical practitioner.

23 (b) Any device required by the federal act to bear on its label  
24 essentially the legend "Rx only".

25 76. "Prescription-only drug" does not include a controlled substance  
26 but does include:

27 (a) Any drug that because of its toxicity or other potentiality for  
28 harmful effect, the method of its use, or the collateral measures necessary  
29 to its use is not generally recognized among experts, qualified by scientific  
30 training and experience to evaluate its safety and efficacy, as safe for use  
31 except by or under the supervision of a medical practitioner.

32 (b) Any drug that is limited by an approved new drug application under  
33 the federal act or section 32-1962 to use under the supervision of a medical  
34 practitioner.

35 (c) Every potentially harmful drug, the labeling of which does not  
36 bear or contain full and adequate directions for use by the consumer.

37 (d) Any drug, other than a controlled substance, required by the  
38 federal act to bear on its label the legend "Rx only".

39 77. "Prescription order" means any of the following:

40 (a) An order to a pharmacist for drugs or devices issued and signed by  
41 a duly licensed medical practitioner in the authorized course of the  
42 practitioner's professional practice.

43 (b) An order transmitted to a pharmacist through word of mouth,  
44 telephone or other means of communication directed by that medical  
45 practitioner. Prescription orders received by word of mouth, telephone or

1 other means of communication shall be maintained by the pharmacist pursuant  
2 to section 32-1964, and the record so made by the pharmacist constitutes the  
3 original prescription order to be dispensed by the pharmacist. This  
4 paragraph does not alter or affect laws of this state or any federal act  
5 requiring a written prescription order.

6 (c) An order initiated by a pharmacist pursuant to a protocol-based  
7 drug therapy agreement with a provider as outlined in section 32-1970, or  
8 immunizations or vaccines administered by a pharmacist pursuant to section  
9 32-1974.

10 78. "Professionally incompetent" means:

11 (a) Incompetence based on a variety of factors, including a lack of  
12 sufficient pharmaceutical knowledge or skills or experience to a degree  
13 likely to endanger the health of patients.

14 (b) When considered with other indications of professional  
15 incompetence, a pharmacist, pharmacy intern or graduate intern who fails to  
16 obtain a passing score on a ~~board-approved~~ BOARD-APPROVED pharmacist  
17 licensure examination or a pharmacy technician or pharmacy technician trainee  
18 who fails to obtain a passing score on a ~~board-approved~~ BOARD-APPROVED  
19 pharmacy technician licensure examination.

20 79. "Radioactive substance" means a substance that emits ionizing  
21 radiation.

22 80. "Safely engage in employment duties" means that a permittee or the  
23 permittee's employee is able to safely engage in employment duties related to  
24 the manufacture, sale, distribution or dispensing of drugs, devices, poisons,  
25 hazardous substances, controlled substances or precursor chemicals.

26 81. "Symbol" means the characteristic symbols that have historically  
27 identified pharmacy, including ~~"show globes"~~, AND ~~"mortar and pestle,"~~  
28 and the sign "Rx".

29 82. "Toxic substance" means a substance, other than a radioactive  
30 substance, that has the capacity to produce injury or illness in humans  
31 through ingestion, inhalation or absorption through any body surface.

32 83. "Ultimate user" means a person who lawfully possesses a drug or  
33 controlled substance for that person's own use, for the use of a member of  
34 that person's household or for administering to an animal owned by that  
35 person or by a member of that person's household.

36 Sec. 2. Section 32-1970, Arizona Revised Statutes, is amended to read:  
37 32-1970. Initiating, monitoring and modifying drug therapy and  
38 use; conditions; definitions

39 A. A pharmacist licensed pursuant to this chapter may ~~implement~~  
40 INITIATE, monitor and modify drug therapy and use only under the following  
41 circumstances:

- 42 1. The patient's drug therapy and use are pursuant to a provider.
- 43 2. The pharmacist complies with rules adopted by the ~~state~~ board of  
44 pharmacy.

1           3. The pharmacist follows the written drug therapy management  
2 protocols prescribed by the provider who made the diagnosis and ~~implements~~  
3 ~~INITIATES~~, monitors or modifies a person's drug therapy and use only pursuant  
4 to those protocols. Each protocol developed pursuant to the drug therapy  
5 agreement shall contain detailed directions concerning the actions that the  
6 pharmacist may perform for that patient. The protocol shall specify, at a  
7 minimum, the specific drug or drugs to be managed by the pharmacist, the  
8 conditions and events for which the pharmacist must notify the provider and  
9 the laboratory tests that may be ordered. A provider who enters into a  
10 protocol-based drug therapy agreement must have a legitimate provider-patient  
11 relationship.

12           B. A licensee who violates this section commits an act of  
13 unprofessional conduct.

14           C. A pharmacist is responsible for the pharmacist's negligent acts  
15 that are the result of the pharmacist's change of medication or that relate  
16 to patient drug usage pursuant to drug therapy management protocols. This  
17 subsection does not limit a provider's liability for negligent acts that are  
18 not related to a pharmacist's change of medication pursuant to the protocols.

19           D. For the purposes of this section:

20           1. "~~Implement~~ INITIATE, monitor and modify" means that a pharmacist  
21 may perform specific acts as authorized by a provider pursuant to written  
22 guidelines and protocols. This does not include the selection of drug  
23 products not prescribed by the provider unless selection of the specific drug  
24 product is authorized by the written guidelines and protocols.

25           2. "Protocol" means a provider's written order, written standing  
26 medical order or other written order of protocol as defined by rules adopted  
27 by the Arizona medical board, ~~and~~ the ARIZONA board of osteopathic examiners  
28 in medicine and surgery AND THE ARIZONA STATE BOARD OF NURSING and that are  
29 patient, provider and pharmacist specific for prescriptions or orders given  
30 by the provider authorizing the written protocol.

31           3. "Provider" means a physician who is licensed pursuant to chapter 13  
32 or 17 of this title or a registered nurse practitioner who is licensed  
33 pursuant to chapter 15 of this title and who acts as a primary care  
34 practitioner.

35           Sec. 3. Section 32-1974, Arizona Revised Statutes, is amended to read:  
36 32-1974. Pharmacists; administration of immunizations, vaccines  
37 and emergency medications; certification; reporting  
38 requirements; advisory committee; definitions

39           A. Except as prescribed pursuant to subsection I of this section, a  
40 pharmacist who is licensed pursuant to this chapter and who meets the  
41 requirements of this section may administer the following to adults without a  
42 prescription order pursuant to rules and protocols adopted by the board  
43 pursuant to this section:

1           1. Immunizations or vaccines ~~listed in~~ RECOMMENDED FOR ADULTS BY the  
2 United States centers for disease control and ~~prevention's recommended adult~~  
3 ~~immunization schedule~~ PREVENTION.

4           2. Immunizations or vaccines recommended by the United States centers  
5 for disease control and prevention's health information for international  
6 travel.

7           B. A pharmacist who is licensed pursuant to this chapter and who meets  
8 the requirements of this section may administer the following to ~~a person who~~  
9 ~~is at least six years of age but under eighteen years of age~~ MINORS without a  
10 prescription order pursuant to rules and protocols adopted by the board  
11 pursuant to this section:

12           1. INFLUENZA immunizations or vaccines ~~for influenza~~ TO A PERSON WHO  
13 IS AT LEAST THREE YEARS OF AGE.

14           ~~2. Immunizations or vaccines in response to a public health emergency~~  
15 ~~declared by the governor pursuant to section 36-787.~~

16           2. BOOSTER DOSES FOR THE PRIMARY ADOLESCENT SERIES AS RECOMMENDED BY  
17 THE UNITED STATES CENTERS FOR DISEASE CONTROL AND PREVENTION.

18           3. IMMUNIZATIONS OR VACCINES RECOMMENDED BY THE UNITED STATES CENTERS  
19 FOR DISEASE CONTROL AND PREVENTION TO A PERSON WHO IS AT LEAST THIRTEEN YEARS  
20 OF AGE.

21           C. ~~Pursuant to a prescription order~~ EXCEPT AS PRESCRIBED IN SUBSECTION  
22 B OF THIS SECTION, a pharmacist who is licensed pursuant to this chapter and  
23 who meets the requirements of this section may administer immunizations and  
24 vaccines, INCLUDING THE FIRST DOSE FOR THE PRIMARY ADOLESCENT SERIES, to a  
25 person who is at least six years of age but under ~~eighteen~~ THIRTEEN years of  
26 age ONLY WITH A PRESCRIPTION ORDER AND pursuant to rules and protocols  
27 adopted by the board pursuant to this section.

28           D. A pharmacist who wishes to administer immunizations and vaccines  
29 pursuant to this section must be certified to do so by the board. The board  
30 shall issue a certificate to a pharmacist who meets board requirements for  
31 certification as prescribed by the board by rule.

32           E. A pharmacist who is certified to administer immunizations and  
33 vaccines pursuant to this section may administer WITHOUT A PRESCRIPTION  
34 ORDER:

35           1. Emergency medication to manage an acute allergic reaction to an  
36 immunization, ~~or~~ vaccine OR MEDICATION IN ACCORDANCE WITH THE UNITED STATES  
37 CENTERS FOR DISEASE CONTROL AND PREVENTION IMMUNIZATION GUIDELINES.

38           2. IMMUNIZATIONS OR VACCINES TO ANY PERSON REGARDLESS OF AGE DURING A  
39 PUBLIC HEALTH EMERGENCY RESPONSE OF THIS STATE PURSUANT TO SECTION 36-787.

40           F. A pharmacist who administers an immunization, vaccine or emergency  
41 medication pursuant to this section must:

42           1. Report the administration to the person's IDENTIFIED primary care  
43 provider or physician, ~~if the primary care provider or physician is~~  
44 ~~available~~, within forty-eight hours after administering the immunization,  
45 vaccine or emergency medication and as prescribed by the board by rule.

1 FAILURE TO REPORT THE ADMINISTRATION OF AN IMMUNIZATION, VACCINE OR EMERGENCY  
2 MEDICATION PURSUANT TO THIS SECTION IS A VIOLATION OF SECTION 32-1901.01,  
3 SUBSECTION B, PARAGRAPH 2. THE PHARMACIST SHALL MAKE A REASONABLE EFFORT TO  
4 IDENTIFY THE PERSON'S PRIMARY CARE PROVIDER OR PHYSICIAN BY ONE OR MORE OF  
5 THE FOLLOWING METHODS:

6 (a) CHECKING ANY ADULT IMMUNIZATION INFORMATION SYSTEM OR VACCINE  
7 REGISTRY ESTABLISHED BY THE DEPARTMENT OF HEALTH SERVICES.

8 (b) CHECKING PHARMACY RECORDS.

9 (c) REQUESTING THE INFORMATION FROM THE PERSON OR, IN THE CASE OF A  
10 MINOR, THE PERSON'S PARENT OR GUARDIAN.

11 2. Report information to any adult immunization information system or  
12 vaccine registry established by the department of health services.

13 3. Maintain a record of the immunization pursuant to title 12, chapter  
14 13, article 7.1 and as prescribed by the board by rule.

15 4. REPORT TO THE PERSON'S IDENTIFIED PRIMARY CARE PROVIDER OR  
16 PHYSICIAN, WITHIN TWENTY-FOUR HOURS OF OCCURRENCE, ANY ADVERSE REACTION  
17 LISTED BY THE VACCINE MANUFACTURER AS A CONTRAINDICATION TO FURTHER DOSES OF  
18 THE VACCINE.

19 ~~4.~~ 5. Participate in any federal vaccine adverse event reporting  
20 system or successor database.

21 G. This section does not establish a cause of action against a  
22 patient's primary care provider OR PHYSICIAN for any adverse reaction,  
23 complication or negative outcome arising from the administration of any  
24 immunization, vaccine or emergency medication by a pharmacist to ~~a~~ THE  
25 patient pursuant to this section if it is administered without a prescription  
26 ORDER written by the patient's primary care provider OR PHYSICIAN.

27 H. The board shall adopt rules for the administration of vaccines or  
28 immunizations pursuant to this section regarding:

29 1. Protocols that are based on protocols approved by the United States  
30 centers for disease control and prevention and any advisory committee  
31 appointed by the board for the purpose of recommending protocols.

32 2. ~~Record keeping~~ RECORDKEEPING and reporting requirements.

33 3. Requirements and qualifications for pharmacist certification  
34 pursuant to this section.

35 4. Vaccine information and educational materials for those requesting  
36 vaccines and immunizations.

37 5. The administration of emergency medication pursuant to this  
38 section.

39 I. The department of health services, by rule, shall establish and  
40 maintain a list of immunizations or vaccines that may be administered to  
41 adults by a pharmacist only pursuant to a prescription order. In adopting  
42 and maintaining this list, the department is exempt from the ~~rule-making~~  
43 RULEMAKING requirements of title 41, chapter 6. The department shall adopt  
44 its initial rules within six months after receipt of the recommendations of  
45 the advisory committee appointed by the board and shall hold one public

1 hearing before implementing the rules and any amendments to the rules. The  
2 list shall include those immunizations or vaccines listed in the United  
3 States centers for disease control and prevention's recommended adult  
4 immunization schedule or recommended by the United States centers for disease  
5 control and prevention's health information for international travel that  
6 have adverse reactions that could cause significant harm to a patient's  
7 health. A pharmacist may not administer immunizations or vaccines without a  
8 prescription order pursuant to this section before the department has  
9 established the list pursuant to this subsection. The board may not  
10 authorize a pharmacist to administer new immunizations or vaccines without a  
11 prescription order pursuant to this section until the department reviews the  
12 new immunizations and vaccines to determine if they should be added to the  
13 list established pursuant to this subsection.

14 J. The board may appoint an advisory committee to assist the board in  
15 adopting and amending rules and developing protocols relating to the  
16 administration of immunizations, vaccines and emergency medications and  
17 certification requirements.

18 K. A pharmacy intern who is certified by the board to administer  
19 immunizations and vaccines pursuant to this section may do so only in the  
20 presence and under the immediate personal supervision of a pharmacist WHO IS  
21 certified as prescribed in this section.

22 L. This section does not prevent a pharmacist who administers an  
23 immunization or vaccine from participating in the federal vaccines for  
24 children program.

25 M. A pharmacist may not administer an immunization or vaccine to a  
26 minor ~~pursuant to subsection B or C of this section~~ without the consent of  
27 the minor's parent or guardian.

28 N. For the purposes of this section: ~~—~~

29 1. "Emergency medication" means emergency epinephrine and  
30 ~~diphenhydramine~~ AND ANTIHISTAMINES IN ACCORDANCE WITH THE UNITED STATES  
31 CENTERS FOR DISEASE CONTROL AND PREVENTION IMMUNIZATION GUIDELINES.

32 2. "PRIMARY ADOLESCENT SERIES" MEANS THOSE IMMUNIZATIONS OR VACCINES  
33 RECOMMENDED BY THE UNITED STATES CENTERS FOR DISEASE CONTROL AND PREVENTION  
34 FOR CHILDREN STARTING AT AGE ELEVEN OR TWELVE.

35 Sec. 4. Legislative intent

36 It is the intent of the legislature that the changes made by this act  
37 to section 32-1974, Arizona Revised Statutes, allow families greater access  
38 to immunizations and vaccinations, maintain and enhance collaboration between  
39 pharmacists and primary care providers and affirm the importance of annual  
40 well-child visits in a medical home during critical developmental ages. This  
41 act recognizes the efficiencies and improved outcomes when care is delivered  
42 through a medical home where a primary care provider working in collaboration  
43 with the family and other providers oversees acute, chronic and preventative  
44 health needs in a coordinated and comprehensive fashion.