State of Arizona Senate Fifty-fifth Legislature First Regular Session 2021

SENATE BILL 1087

AN ACT

AMENDING SECTIONS 32-1901, 32-1901.01, 32-1904, 32-1922, 32-1924, 32-1925, 32-1930, 32-1931, 32-1937, 32-1941, 32-1967, 32-1974, 32-1982, 36-2602, 36-2604, 36-2607, 36-2608, 41-619.51, 41-1758 AND 41-1758.01, ARIZONA REVISED STATUTES; RELATING TO THE ARIZONA STATE BOARD OF PHARMACY.

(TEXT OF BILL BEGINS ON NEXT PAGE)

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read:

Be it enacted by the Legislature of the State of Arizona: Section 1. Section 32–1901, Arizona Revised Statutes, is amended to

32-1901. <u>Definitions</u>

In this chapter, unless the context otherwise requires:

- 1. "Administer" means the direct application of DIRECTLY APPLYING a controlled substance, prescription-only drug, dangerous drug or narcotic drug, whether by injection, inhalation, ingestion or any other means, to the body of a patient or research subject by a practitioner or by the practitioner's authorized agent or the patient or research subject at the direction of the practitioner.
- 2. "Advertisement" means all representations THAT ARE disseminated in any manner or by any means, other than by labeling, for the purpose of inducing, or that are likely to induce, directly or indirectly, the purchase of drugs, devices, poisons or hazardous substances.
- 3. "Advisory letter" means a nondisciplinary letter to notify a licensee or permittee that either:
- (a) While there is insufficient evidence to support disciplinary action, the board believes that continuation of the activities that led to the investigation may result in further board action against the licensee or permittee.
- (b) The violation is a minor or technical violation that is not of sufficient merit to warrant disciplinary action.
- (c) While the licensee or permittee has demonstrated substantial compliance through rehabilitation, remediation or reeducation that has mitigated the need for disciplinary action, the board believes that repetition of REPEATING the activities that led to the investigation may result in further board action against the licensee or permittee.
- 4. "Antiseptic", if a drug is represented as such on its label, means a representation that it is a germicide, except in the case of a drug purporting to be, or represented as, an antiseptic for inhibitory use as a wet dressing, ointment or dusting powder or other use that involves prolonged contact with the body.
- 5. "Authorized officers of the law" means legally empowered peace officers, compliance officers of the board of pharmacy and agents of the division of narcotics enforcement and criminal intelligence of the department of public safety.
- 6. "Automated prescription-dispensing kiosk" means a mechanical system that is operated as an extension of a pharmacy, that maintains all transaction information within the pharmacy operating system, that is separately permitted from the pharmacy and that performs operations that either:
- (a) Accept a prescription or refill order, store prepackaged or repackaged medications, label and dispense patient-specific prescriptions and provide counseling on new or refilled prescriptions.

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- (b) Dispense or deliver a prescription or refill that has been prepared by or on behalf of the pharmacy that oversees the automated prescription-dispensing kiosk.
- 7. "Board" or "board of pharmacy" means the Arizona state board of pharmacy.
- 8. "Certificate of composition" means a list of a product's ingredients.
- 9. "Certificate of free sale" means a document that authenticates a product that is generally and freely sold in domestic or international channels of trade.
 - 10. "Color additive" means a material that either:
- (a) Is any dye, pigment or other substance THAT IS made by a process of synthesis or similar artifice, or THAT IS extracted, isolated or otherwise derived, with or without intermediate or final change of identity, from any vegetable, animal, mineral or other source.
- (b) If added or applied to a drug, or to the human body or any part of the human body, is capable of imparting color, except that color additive does not include any material that has been or may be exempted under the federal act. Color includes black, white and intermediate grays.
- 11. "Compounding" means the preparation PREPARING, assembling, packaging or labeling of a drug by a pharmacist or an intern or pharmacy technician under the pharmacist's supervision, for the purpose of dispensing to a patient based on a valid prescription order. Compounding includes the preparation of PREPARING drugs in anticipation of prescription orders prepared on routine, regularly observed prescribing patterns and the preparation of PREPARING drugs as an incident to research, teaching or chemical analysis or for administration by a medical practitioner to the medical practitioner's patient and not for sale or dispensing. Compounding does not include the preparation of PREPARING commercially available products from bulk compounds or the preparation of PREPARING drugs for sale to pharmacies, practitioners or entities for the purpose of dispensing or distribution.
- 12. "Compressed medical gas distributor" means a person who THAT holds a current permit issued by the board to distribute compressed medical gases pursuant to a compressed medical gas suppliers and other entities that are registered, licensed or permitted to use, administer or distribute compressed medical gases.
- 13. "Compressed medical gases" means gases and liquid oxygen that a compressed medical gas distributor or manufacturer has labeled in compliance with federal law.
- 14. "Compressed medical gas order" means an order for compressed medical gases that is issued by a medical practitioner.
- 15. "Compressed medical gas supplier" means a person $\frac{1}{2}$ who THAT holds a current permit issued by the board to supply compressed medical gases

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pursuant to a compressed medical gas order and only to the consumer or the patient.

- 16. "Controlled substance" means a drug, substance or immediate precursor that is identified, defined or listed in title 36, chapter 27, article 2.
- 17. "Corrosive" means any substance that when it comes in contact with living tissue will cause destruction of THE tissue by chemical action.
- 18. "Counterfeit drug" means a drug that, or the container or labeling of which, without authorization, bears the trademark, trade name or other identifying mark, imprint, number or device, or any likeness of these, of a manufacturer, distributor or dispenser other than the person who THAT in fact manufactured, distributed or dispensed that drug.
- 19. "Dangerous drug" has the same meaning prescribed in section 13-3401.
 - 20. "Day" means a business day.
- 21. "Decree of censure" means an official action that is taken by the board and that may include a requirement for restitution of fees to a patient or consumer.
- 22. "Deliver" or "delivery" means the actual, constructive or attempted transfer from one person to another whether or not there is an agency relationship.
- 23. "Deputy director" means a pharmacist who is employed by the board and selected by the executive director to perform duties as prescribed by the executive director.
- 24. "Device", except as used in paragraph 18 of this section, section 32-1965, paragraph 4 and section 32-1967, subsection A, paragraph 15 and subsection C, means instruments AN INSTRUMENT, apparatuses and contrivances APPARATUS AND CONTRIVANCE, including their ITS components, parts and accessories, including all such items under the federal act, THAT IS intended either:
- (a) For use in the diagnosis, cure, mitigation, treatment or prevention of DIAGNOSING, CURING, MITIGATING, TREATING OR PREVENTING disease in the human body or other animals.
- (b) To affect the structure or any function of the human body or other animals.
- 25. "Director" means the director of the division of narcotics enforcement and criminal investigation of the department of public safety.
- 26. "Direct supervision of a pharmacist" means THAT the pharmacist is present. If relating to the sale of certain items, direct supervision of a pharmacist means that a pharmacist determines the legitimacy or advisability of a proposed purchase of those items.
- 27. "Dispense" means to deliver to an ultimate user or research subject by or pursuant to the lawful order of a practitioner, including

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 the prescribing, administering, packaging, labeling or compounding necessary to prepare for that delivery.

- 28. "Dispenser" means a practitioner who dispenses.
- 29. "Distribute" means to deliver, other than by administering or dispensing.
 - 30. "Distributor" means a person who distributes.
 - 31. "Drug" means:
- (a) Articles THAT ARE recognized, or for which standards or specifications are prescribed, in the official compendium.
- (b) Articles THAT ARE intended for use in the diagnosis, cure, mitigation, treatment or prevention of disease in the human body or other animals.
- (c) Articles other than food THAT ARE intended to affect the structure or any function of the human body or other animals.
- (d) Articles THAT ARE intended for use as a component of any articles specified in subdivision (a), (b) or (c) of this paragraph but does not include devices or their components, parts or accessories.
- 32. "Drug enforcement administration" means the drug enforcement administration of the United States department of justice or its successor agency.
- 33. "Drug or device manufacturing" means the production, preparation, propagation PRODUCING, PREPARING, PROPAGATING or processing of a drug or device, either directly or indirectly, by extraction from substances of natural origin or independently by means of chemical synthesis and includes any packaging or repackaging of substances or labeling or relabeling of its container and the promotion PROMOTING and marketing of the same. Drug or device manufacturing does not include compounding.
- 34. "DURABLE MEDICAL EQUIPMENT" MEANS TECHNOLOGICALLY SOPHISTICATED MEDICAL EQUIPMENT AS PRESCRIBED BY THE BOARD IN RULE THAT A PATIENT OR CONSUMER MAY USE IN A HOME OR RESIDENCE AND THAT MAY BE A PRESCRIPTION-ONLY DEVICE.
 - 35. "DURABLE MEDICAL EQUIPMENT DISTRIBUTOR":
- (a) MEANS A PERSON THAT STORES OR DISTRIBUTES DURABLE MEDICAL EQUIPMENT OTHER THAN TO THE PATIENT OR CONSUMER.
- (b) INCLUDES A VIRTUAL DURABLE MEDICAL EQUIPMENT DISTRIBUTOR AS PRESCRIBED IN RULE BY THE BOARD.
 - 36. "DURABLE MEDICAL EQUIPMENT SUPPLIER":
- (a) MEANS A PERSON THAT SELLS, LEASES OR SUPPLIES DURABLE MEDICAL EQUIPMENT TO THE PATIENT OR CONSUMER.
- (b) INCLUDES A VIRTUAL DURABLE MEDICAL EQUIPMENT SUPPLIER AS PRESCRIBED IN RULE BY THE BOARD.
- 34. 37. "Economic poison" means any substance that alone, in chemical combination with or in formulation with one or more other substances is a pesticide within the meaning of the laws of this state or

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the federal insecticide, fungicide and rodenticide act and that is used in the production, storage PRODUCING, STORING or transportation of TRANSPORTING raw agricultural commodities.

35. 38. "Enteral feeding" means nourishment THAT IS provided by means of a tube inserted into the stomach or intestine.

 $\frac{36.}{39.}$ "Established name", with respect to a drug or ingredient of a drug, means any of the following:

- (a) The applicable official name.
- (b) If there is no such name and the drug or ingredient is an article recognized in an official compendium, the official title in an official compendium.
- (c) If neither subdivision (a) nor (b) of this paragraph applies, the common or usual name of the drug.

 $\frac{37.}{}$ 40. "Executive director" means the executive director of the board of pharmacy.

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

39. 42. "Full service FULL-SERVICE wholesale permittee":

- (a) Means a permittee who may distribute prescription-only drugs and devices, controlled substances and over-the-counter drugs and devices to pharmacies or other legal outlets from a place devoted in whole or in part to wholesaling these items.
 - (b) Includes a virtual wholesaler as defined in rule by the board.
- 40. 43. "Good manufacturing practice" means a system for ensuring that products are consistently produced and controlled according to quality standards and covering all aspects of design, monitoring and control of manufacturing processes and facilities to ensure that products do not pose any risk to the consumer or public.

41. 44. "Highly toxic" means any substance that falls within any of the following categories:

- (a) Produces death within fourteen days in half or more than half of a group of ten or more laboratory white rats each weighing between two hundred and three hundred grams, at a single dose of fifty milligrams or less per kilogram of body weight, when orally administered.
- (b) Produces death within fourteen days in half or more than half of a group of ten or more laboratory white rats each weighing between two hundred and three hundred grams, if inhaled continuously for a period of one hour or less at an atmospheric concentration of two hundred parts per million by volume or less of gas or vapor or two milligrams per liter by volume or less of mist or dust, provided the concentration is likely to be encountered by humans if the substance is used in any reasonably foreseeable manner.

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 (c) Produces death within fourteen days in half or more than half of a group of ten or more rabbits tested in a dosage of two hundred milligrams or less per kilogram of body weight, if administered by continuous contact with the bare skin for twenty-four hours or less.

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

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

43. 46. "Intern" means a pharmacy intern.

44. 47. "Internship" means the practical, experiential, hands-on training of a pharmacy intern under the supervision of a preceptor.

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

46. 49. "Jurisprudence examination" means a board-approved pharmacy law examination that is written and administered in cooperation with the national association of boards of pharmacy or another board-approved pharmacy law examination.

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

 $\frac{48.}{100}$ 51. "Labeling" means all labels and other written, printed or graphic matter THAT either:

- (a) IS on any article or any of its containers or wrappers.
- (b) Accompanying ACCOMPANIES that article.

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

50. 53. "Limited service pharmacy" means a pharmacy that is approved by the board to practice a limited segment of pharmacy as indicated by the permit issued by the board.

51. 54. "Manufacture" or "manufacturer":

- (a) Means every person who prepares, derives, produces, compounds, processes, packages or repackages or labels any drug in a place, other than a pharmacy, that is devoted to manufacturing the drug.
- (b) Includes a virtual manufacturer as defined in rule by the board.

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 $\frac{52.}{13-3401.}$ 55. "Marijuana" has the same meaning prescribed in section 13-3401.

53. 56. "Medical practitioner" means any medical doctor, doctor of osteopathic medicine, dentist, podiatrist, veterinarian or other person who is licensed and authorized by law to use and prescribe drugs and devices for the treatment of TO TREAT sick and injured human beings or animals or for the diagnosis TO DIAGNOSE or prevention of PREVENT sickness in human beings or animals in this state or any state, territory or district of the United States.

54. 57. "Medication order" means a written or verbal order from a medical practitioner or that person's authorized agent to administer a drug or device.

 $\frac{55.}{13-3401.}$ "Narcotic drug" has the same meaning prescribed in section 13-3401.

56. "New drug" means either:

- (a) Any drug OF WHICH the composition of which is such that the drug is not generally recognized among experts qualified by scientific training and experience to evaluate the safety and effectiveness of drugs as safe and effective for use under the conditions prescribed, recommended or suggested in the labeling.
- (b) Any drug OF WHICH the composition of which is such that the drug, as a result of investigations to determine its safety and effectiveness for use under such conditions, has become so recognized, but that has not, other than in the investigations, been used to a material extent or for a material time under those conditions.
- 57. 60. "Nonprescription drug" or "over-the-counter drug" means any nonnarcotic medicine or drug that may be sold without a prescription and that is prepackaged and labeled for use by the consumer in accordance with the requirements of the laws of this state and federal law. Nonprescription drug does not include:
- (a) A drug that is primarily advertised and promoted professionally to medical practitioners and pharmacists by manufacturers or primary distributors.
 - (b) A controlled substance.
 - (c) A drug that is required to bear a label that states "Rx only".
 - (d) A drug that is intended for human use by hypodermic injection.
 - 58. 61. "Nonprescription drug wholesale permittee":
- (a) Means a permittee who may distribute only over-the-counter drugs and devices to pharmacies or other lawful outlets from a place devoted in whole or in part to wholesaling these items.
 - (b) Includes a virtual wholesaler as defined in rule by the board.
- 59. 62. "Notice" means personal service or the mailing of a copy of the notice by certified mail AND EMAIL addressed either to the person at the person's latest address of record in the board office or to the

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 PERSON AND THE person's attorney USING THE MOST RECENT INFORMATION PROVIDED TO THE BOARD IN THE BOARD'S LICENSING DATABASE.

60. 63. "Nutritional supplementation" means vitamins, minerals and caloric supplementation. Nutritional supplementation does not include medication or drugs.

61. 64. "Official compendium" means the latest revision of the United States pharmacopeia and the national formulary or any current supplement.

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

63. 66. "Package" means a receptacle THAT IS defined or described in the United States pharmacopeia and the national formulary as adopted by the board.

64. 67. "Packaging" means the act or process of placing a drug item or device in a container for the purpose or intent of dispensing or distributing the item or device to another.

65. 68. "Parenteral nutrition" means intravenous feeding that provides a person AN INDIVIDUAL with fluids and essential nutrients the person INDIVIDUAL needs while the person INDIVIDUAL is unable to receive adequate fluids or feedings by mouth or by enteral feeding.

66. 69. "Person" means an individual, partnership, corporation and association, and their duly authorized agents.

67. 70. "Pharmaceutical care" means the provision of drug therapy and other pharmaceutical patient care services.

68. 71. "Pharmacist" means an individual who is currently licensed by the board to practice the profession of pharmacy in this state.

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

70. 73. "Pharmacist licensure examination" means a board-approved examination that is written and administered in cooperation with the national association of boards of pharmacy or any other board-approved pharmacist licensure examination.

71. 74. "Pharmacy":

- (a) Means:
- (i) Any place where drugs, devices, poisons or related hazardous substances are offered for sale at retail.
- (ii) Any place in which the profession of pharmacy is practiced or where prescription orders are compounded and dispensed.
- (iii) Any place that has displayed on it or in it the words "pharmacist", "pharmaceutical chemist", "apothecary", "druggist", "pharmacy", "drugstore", "drugs" or "drug sundries" or any of these words

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or combinations of these words, or words of similar import either in English or any other language, or that is advertised by any sign containing any of these words.

- (iv) Any place where the characteristic symbols of pharmacy or the characteristic prescription sign "Rx" is exhibited.
- (v) Any place or a portion of any building or structure that is leased, used or controlled by the permittee to conduct the business authorized by the board at the address for which the permit was issued and that is enclosed and secured when a pharmacist is not in attendance.
- (vi) A remote dispensing site pharmacy where a pharmacy technician or pharmacy intern prepares, compounds or dispenses prescription medications under remote supervision by a pharmacist.
- (vii) A REMOTE HOSPITAL SITE PHARMACY, AS DEFINED BY THE BOARD IN RULE, THAT OPERATES UNDER DIRECT OR REMOTE SUPERVISION BY A PHARMACIST PURSUANT TO RULES ADOPTED BY THE BOARD.
 - (b) Includes a satellite pharmacy.
- 72. 75. "Pharmacy intern" means a person who has all of the qualifications and experience prescribed in section 32-1923.
- 73. 76. "Pharmacy technician" means a person who is licensed pursuant to this chapter.
- 74. 77. "Pharmacy technician trainee" means a person who is licensed pursuant to this chapter.
- 75. 78. "Poison" or "hazardous substance" includes, but is not limited to, any of the following if intended and suitable for household use or use by children:
- (a) Any substance that, according to standard works on medicine, pharmacology, pharmacognosy or toxicology, if applied to, introduced into or developed within the body in relatively small quantities by its inherent action uniformly produces serious bodily injury, disease or death.
 - (b) A toxic substance.
 - (c) A highly toxic substance.
 - (d) A corrosive substance.
 - (e) An irritant.
 - (f) A strong sensitizer.
- (g) A mixture of any of the substances described in this paragraph, if the substance or mixture of substances may cause substantial personal injury or substantial illness during or as a proximate result of any customary or reasonably foreseeable handling or use, including reasonably foreseeable ingestion by children.
- (h) A substance that is designated by the board to be a poison or hazardous substance. This subdivision does not apply to radioactive substances, economic poisons subject to the federal insecticide, fungicide and rodenticide act or the state pesticide act, foods, drugs and cosmetics subject to state laws or the federal act or substances intended for use as

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fuels when stored in containers and used in the heating, cooking or refrigeration system of a house. This subdivision applies to any substance or article that is not itself an economic poison within the meaning of the federal insecticide, fungicide and rodenticide act or the state pesticide act, but that is a poison or hazardous substance within the meaning of this paragraph by reason of bearing or containing an economic poison or hazardous substance.

76. 79. "Practice of pharmacy":

- (a) Means furnishing the following health care services as a medical professional:
- (i) Interpreting, evaluating and dispensing prescription orders in the patient's best interests.
- (ii) Compounding drugs pursuant to or in anticipation of a prescription order.
- (iii) Labeling drugs and devices in compliance with state and federal requirements.
- (iv) Participating in drug selection and drug utilization reviews, drug administration, drug or drug-related research and drug therapy monitoring or management.
- (v) Providing patient counseling necessary to provide pharmaceutical care.
- (vi) Properly and safely storing drugs and devices in anticipation of dispensing.
 - (vii) Maintaining required records of drugs and devices.
- (viii) Offering or performing acts, services, operations or transactions THAT ARE necessary in the TO conduct, operation, management OPERATE, MANAGE and control of a pharmacy.
- (ix) Initiating, monitoring and modifying drug therapy pursuant to a protocol-based drug therapy agreement with a provider as outlined in section 32-1970.
- (x) Initiating and administering immunizations or vaccines pursuant to section 32-1974.
- (b) Does not include initiating a prescription order for any medication, drug or other substance used to induce or cause a medication abortion as defined in section 36-2151.
- 77. 80. "Practitioner" means any physician, dentist, veterinarian, scientific investigator or other person who is licensed, registered or otherwise permitted to distribute, dispense, conduct research with respect to or administer a controlled substance in the course of professional practice or research in this state, or any pharmacy, hospital or other institution that is licensed, registered or otherwise permitted to distribute, dispense, conduct research with respect to or administer a controlled substance in the course of professional practice or research in this state.

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 $\frac{78.}{100}$ 81. "Preceptor" means a pharmacist who is serving as the practical instructor of an intern and WHO complies with section 32-1923.

79. 82. "Precursor chemical" means a substance that is:

- (a) The principal compound that is commonly used or that is produced primarily for use and that is an immediate chemical intermediary used or likely to be used in the manufacture of a controlled substance, the control of which is necessary to prevent, curtail or limit manufacture.
 - (b) Listed in section 13-3401, paragraph 26 or 27.

80. 83. "Prescription" means either a prescription order or a prescription medication.

81. 84. "Prescription medication" means any drug, including label and container according to context, that is dispensed pursuant to a prescription order.

85. "Prescription-only device" includes:

- (a) Any device that is limited by the federal act to use under the supervision of a medical practitioner.
- (b) Any device required by the federal act to bear on its label essentially the legend "Rx only".

83. 86. "Prescription-only drug" does not include a controlled substance but does include:

- (a) Any drug that because of its toxicity or other potentiality for harmful effect, the method of its use, or the collateral measures necessary to its use is not generally recognized among experts, qualified by scientific training and experience to evaluate its safety and efficacy, as safe for use except by or under the supervision of a medical practitioner.
- (b) Any drug that is limited by an approved new drug application under the federal act or section 32-1962 to use under the supervision of a medical practitioner.
- (c) Every potentially harmful drug, the labeling of which does not bear or contain full and adequate directions for use by the consumer.
- (d) Any drug, other than a controlled substance, THAT IS required by the federal act to bear on its label the legend "Rx only".

87. "Prescription order" means any of the following:

- (a) An order to a pharmacist for drugs or devices THAT IS issued and signed by a duly licensed medical practitioner in the authorized course of the practitioner's professional practice.
- (b) An order THAT IS transmitted to a pharmacist through word of mouth, telephone or other means of communication directed by that medical practitioner. Prescription orders received by word of mouth, telephone or other means of communication shall be maintained by the pharmacist pursuant to section 32-1964, and the record so made by the pharmacist constitutes the original prescription order to be dispensed by the

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pharmacist. This paragraph does not alter or affect laws of this state or any federal act requiring a written prescription order.

- (c) An order THAT IS initiated by a pharmacist pursuant to a protocol-based drug therapy agreement with a provider as outlined in section 32-1970, or immunizations or vaccines administered by a pharmacist pursuant to section 32-1974.
- (d) A diet order or an order for enteral feeding, nutritional supplementation or parenteral nutrition that is initiated by a registered dietitian or other qualified nutrition professional in a hospital pursuant to section 36-416.

85. 88. "Professionally incompetent" means:

- (a) Incompetence based on a variety of factors, including a lack of sufficient pharmaceutical knowledge or skills or experience to a degree likely to endanger the health of patients.
- (b) When considered with other indications of professional incompetence, a pharmacist or pharmacy intern who fails to obtain a passing score on a board-approved pharmacist licensure examination or a pharmacy technician or pharmacy technician trainee who fails to obtain a passing score on a board-approved pharmacy technician licensure examination.

86. 89. "Radioactive substance" means a substance that emits ionizing radiation.

87. 90. "Remote dispensing site pharmacy" means a pharmacy where a pharmacy technician or pharmacy intern prepares, compounds or dispenses prescription medications under remote supervision by a pharmacist.

88. 91. "Remote supervision by a pharmacist" means that a pharmacist directs and controls the actions of pharmacy technicians and pharmacy interns through the use of audio and visual technology.

89. 92. "Revocation" or "revoke" means the official cancellation of a license, permit, registration or other approval authorized by the board for a period of two years unless otherwise specified by the board. A request or new application for reinstatement may be presented to the board for review before the conclusion of the specified revocation period upon review of the executive director.

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

91. 94. "Satellite pharmacy" means a work area located within a hospital or on a hospital campus that is not separated by other commercial property or residential property, that is under the direction of a pharmacist, that is a remote extension of a centrally licensed hospital pharmacy, and that is owned by and dependent on the centrally licensed

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 hospital pharmacy for administrative control, staffing and drug procurement and that is not required to be separately permitted.

92. 95. "Symbol" means the characteristic symbols that have historically identified pharmacy, including show globes and mortar and pestle, and the sign "Rx".

93. 96. "Third-party logistics provider" means an entity that provides or coordinates warehousing or other logistics services for a prescription or over-the-counter dangerous drug or dangerous device in intrastate or interstate commerce on behalf of a manufacturer, wholesaler or dispenser of the prescription or over-the-counter dangerous drug or dangerous device THE FOLLOWING ITEMS, but that does not take ownership of the prescription or over-the-counter dangerous drug or dangerous device or have responsibility to direct its sale or disposition THE ITEMS, AND THAT DISTRIBUTES THOSE ITEMS AS DIRECTED BY A MANUFACTURER, WHOLESALER, DISPENSER OR DURABLE MEDICAL EQUIPMENT SUPPLIER THAT IS PERMITTED BY THE BOARD:

- (a) NARCOTIC DRUGS OR OTHER CONTROLLED SUBSTANCES.
- (b) DANGEROUS DRUGS AS DEFINED IN SECTION 13-3401.
- (c) PRESCRIPTION-ONLY DRUGS AND DEVICES.
- (d) NONPRESCRIPTION DRUGS AND DEVICES.
- (e) PRECURSOR CHEMICALS.
- (f) REGULATED CHEMICALS AS DEFINED IN SECTION 13-3401.

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

 $95.\,$ 98. "Ultimate user" means a person who lawfully possesses a drug or controlled substance for that person's own use, for the use of a member of that person's household or for administering to an animal owned by that person or by a member of that person's household.

Sec. 2. Section 32-1901.01, Arizona Revised Statutes, is amended to read:

32-1901.01. <u>Definition of unethical and unprofessional</u> <u>conduct; permittees; licensees</u>

A. In this chapter, unless the context otherwise requires, for the purposes of disciplining a permittee, "unethical conduct" means the following, whether occurring in this state or elsewhere:

- 1. Committing a felony, whether or not involving moral turpitude, or a misdemeanor involving moral turpitude or any drug-related offense. In either case, conviction by a court of competent jurisdiction or a plea of no contest is conclusive evidence of the commission.
- 2. Committing an act that is substantially related to the qualifications, functions or duties of a permittee and that demonstrates either a lack of good moral character or an actual or potential unfitness to hold a permit in light of the public's safety.

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- 3. Working under the influence of alcohol or other drugs.
- 4. Being addicted to the use of USING alcohol or other drugs to such a degree as to render the permittee unfit to perform the permittee's employment duties.
- 5. Violating a federal or state law or administrative rule relating to the manufacture, sale or distribution of drugs, devices, poisons, hazardous substances or precursor chemicals.
- 6. Violating a federal or state law or administrative rule relating to marijuana, prescription-only drugs, narcotics, dangerous drugs, controlled substances or precursor chemicals.
- 7. Violating state or federal reporting or recordkeeping requirements on transactions relating to precursor chemicals.
- 8. Failing to report in writing to the board any evidence that a pharmacist or pharmacy intern is or may be professionally incompetent, is or may be guilty of unprofessional conduct or is or may be mentally or physically unable safely to engage in the practice of pharmacy.
- 9. Failing to report in writing to the board any evidence that a pharmacy technician or pharmacy technician trainee is or may be professionally incompetent, is or may be guilty of unprofessional conduct or is or may be mentally or physically unable safely to engage in the permissible activities of a pharmacy technician or pharmacy technician trainee.
- 10. Failing to report in writing to the board any evidence that appears to show that a permittee or permittee's employee is or may be guilty of unethical conduct, is or may be mentally or physically unable safely to engage in employment duties related to manufacturing, selling, distributing or dispensing of drugs, devices, poisons, hazardous substances, controlled substances or precursor chemicals or is or may be in violation of VIOLATING this chapter or a rule adopted under this chapter.
- 11. Intending to sell, transfer or distribute, or to offer for sale, transfer or distribution, or selling, transferring, distributing or dispensing or offering for sale, transfer or distribution an imitation controlled substance, imitation over-the-counter drug or imitation prescription-only drug as defined in section 13-3451.
- 12. Having the permittee's permit to manufacture, sell, distribute or dispense drugs, devices, poisons, hazardous substances or precursor chemicals denied or disciplined in another jurisdiction.
- 13. Committing an offense in another jurisdiction that if committed in this state would be grounds for discipline.
- 14. Obtaining or attempting to obtain a permit or a permit renewal by fraud, by misrepresentation or by knowingly taking advantage of the mistake of another person or an agency.
- 15. Wilfully making a false report or record THAT IS required by this chapter, THAT IS required by federal or state laws pertaining to

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drugs, devices, poisons, hazardous substances or precursor chemicals or THAT IS required for the payment TO PAY for drugs, devices, poisons or hazardous substances or precursor chemicals or for services pertaining to such drugs or substances.

- 16. Knowingly filing with the board any application, renewal or other document that contains false or misleading information.
- 17. Providing false or misleading information or omitting material information in any communication to the board or the board's employees or agents.
- 18. Violating or attempting to violate, directly or indirectly, or assisting in or abetting the violation of, or conspiring to violate this chapter.
- 19. Violating a formal order, terms of probation, a consent agreement or a stipulation issued or entered into by the board or its executive director pursuant to this chapter.
- 20. Failing to comply with a board subpoena or failing to comply in a timely manner with a board subpoena without providing any explanation to the board for not complying with the subpoena.
- 21. Failing to provide the board or its employees or agents or an authorized federal or state official conducting a site investigation, inspection or audit with access to any place for which a permit has been issued or for which an application for a permit has been submitted.
- 22. Failing to notify the board of a change of ownership, management or pharmacist in charge.
- 23. Failing to promptly produce on the request of the official conducting a site investigation, inspection or audit any book, record or document.
- 24. Overruling or attempting to overrule a pharmacist in matters of pharmacy ethics or interpreting laws pertaining to the practice of pharmacy or the distribution of drugs or devices.
- $\,$ 25. Distributing premiums or rebates of any kind in connection with the sale of prescription medication, other than to the prescription medication recipient.
- 26. Failing to maintain effective controls against the diversion of controlled substances or precursor chemicals to unauthorized persons or entities.
 - 27. Fraudulently claiming to have performed a service.
 - 28. Fraudulently charging a fee for a service.
- 29. Advertising drugs or devices, or services pertaining to drugs or devices, in a manner that is untrue or misleading in any particular, and that is known, or that by the exercise of reasonable care should be known, to be untrue or misleading.
- B. In this chapter, unless the context otherwise requires, for the purposes of disciplining a pharmacist or pharmacy intern, "unprofessional

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conduct" means the following, whether occurring in this state or elsewhere:

- 1. Being addicted to the use of USING alcohol or other drugs to such a degree as to render the licensee unfit to practice the profession of pharmacy.
- 2. Violating any federal or state law, rule or regulation relating to the manufacture or distribution of drugs and devices or the practice of pharmacy.
- 3. Dispensing a different drug or brand of drug in place of the drug or brand of drug ordered or prescribed without the express permission in each case of the orderer, or in the case of a prescription order, the medical practitioner. The conduct prohibited by this paragraph does not apply to substitutions authorized pursuant to section 32-1963.01.
- 4. Obtaining or attempting to obtain a license to practice pharmacy or a license renewal by fraud, by misrepresentation or by knowingly taking advantage of the mistake of another person or an agency.
- 5. Having the licensee's license to practice pharmacy denied or disciplined in another jurisdiction.
- 6. Claiming professional superiority in compounding or dispensing prescription orders.
- 7. Failing to comply with the mandatory continuing professional pharmacy education requirements of sections 32-1936 and 32-1937 and rules adopted by the board.
- 8. Committing a felony, whether or not involving moral turpitude, or a misdemeanor involving moral turpitude or any drug-related offense. In either case, conviction by a court of competent jurisdiction or a plea of no contest is conclusive evidence of the commission.
 - 9. Working under the influence of alcohol or other drugs.
- 10. Violating a federal or state law or administrative rule relating to marijuana, prescription-only drugs, narcotics, dangerous drugs, controlled substances or precursor chemicals when determined by the board or by conviction in a federal or state court.
- 11. Knowingly dispensing a drug without a valid prescription order as required pursuant to section 32-1968, subsection A.
- 12. Knowingly dispensing a drug on a prescription order that was issued in the course of the conduct of business of dispensing drugs pursuant to diagnosis by mail or the internet, unless the order was any of the following:
- (a) Made by a physician who provides temporary patient supervision on behalf of the patient's regular treating licensed health care professional or provides a consultation requested by the patient's regular treating licensed health care professional.
- (b) Made in an emergency medical situation as defined in section 41-1831.
 - (c) Written to prepare a patient for a medical examination.

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- (d) Written or the prescription medications were issued for use by a county or tribal public health department for immunization programs or emergency treatment or in response to an infectious disease investigation, a public health emergency, an infectious disease outbreak or an act of bioterrorism. For the purposes of this subdivision, "bioterrorism" has the same meaning prescribed in section 36-781.
- (e) Written or antimicrobials were dispensed by the prescribing or dispensing physician to a contact as defined in section 36-661 who is believed to have had significant exposure risk as defined in section 36-661 with another person who has been diagnosed with a communicable disease as defined in section 36-661.
- (f) Written or the prescription medications were issued for administration of ADMINISTERING immunizations or vaccines listed in the United States centers for disease control and prevention's recommended immunization schedule to a household member of a patient.
- (g) For epinephrine auto-injectors that are written or dispensed for a school district or charter school and that are to be stocked for emergency use pursuant to section 15-157 or for an authorized entity to be stocked pursuant to section 36-2226.01.
- (h) Written by a licensee through a telemedicine program that is covered by the policies and procedures adopted by the administrator of a hospital or outpatient treatment center.
- (i) Written pursuant to a physical or mental health status examination that was conducted during a real-time telemedicine encounter with audio and video capability.
- (j) For naloxone hydrochloride or any other opioid antagonist approved by the United States food and drug administration and written or dispensed for use pursuant to section 36-2228 or 36-2266.
- 13. Failing to report in writing to the board any evidence that a pharmacist or pharmacy intern is or may be professionally incompetent, is or may be guilty of unprofessional conduct or is or may be mentally or physically unable to safely engage in the practice of pharmacy.
- 14. Failing to report in writing to the board any evidence that a pharmacy technician or pharmacy technician trainee is or may be professionally incompetent, is or may be guilty of unprofessional conduct or is or may be mentally or physically unable to safely engage in the permissible activities of a pharmacy technician or pharmacy technician trainee.
- 15. Failing to report in writing to the board any evidence that a permittee or a permittee's employee is or may be guilty of unethical conduct or is or may be $\frac{1}{100}$ violation of VIOLATING this chapter or a rule adopted under this chapter.
- 16. Committing an offense in another jurisdiction that if committed in this state would be grounds for discipline.

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- 17. Knowingly filing with the board any application, renewal or other document that contains false or misleading information.
- 18. Providing false or misleading information or omitting material information in any communication to the board or the board's employees or agents.
- 19. Violating or attempting to violate, directly or indirectly, or assisting in or abetting in the violation of, or conspiring to violate this chapter.
- 20. Violating a formal order, terms of probation, a consent agreement or a stipulation issued or entered into by the board or its executive director pursuant to this chapter.
- 21. Failing to comply with a board subpoena or failing to comply in a timely manner with a board subpoena without providing any explanation to the board for not complying with the subpoena.
- 22. Refusing without just cause to allow authorized agents of the board to examine documents that are required to be kept pursuant to this chapter or title 36.
- 23. Participating in an arrangement or agreement to allow a prescription order or a prescription medication to be left at, picked up from, accepted by or delivered to a place that is not licensed as a pharmacy. This paragraph does not prohibit a pharmacist or a pharmacy from using an employee or a common carrier to pick up prescription orders at or deliver prescription medications to the office or home of a medical practitioner, the residence of a patient or a patient's hospital.
- 24. Paying rebates or entering into an agreement for the payment of PAYING rebates to a medical practitioner or any other person in the health care field.
- 25. Providing or causing to be provided to a medical practitioner prescription order blanks or forms bearing the pharmacist's or pharmacy's name, address or other means of identification.
 - 26. Fraudulently claiming to have performed a professional service.
 - 27. Fraudulently charging a fee for a professional service.
- 28. Failing to report a change of the licensee's home address, contact information, employer or employer's address as required by section 32-1926.
- 29. Failing to report a change in the licensee's residency status as required by section 32-1926.01.
- 30. Failing to maintain effective controls against the diversion of controlled substances or precursor chemicals to unauthorized persons or entities.
- C. In this chapter, unless the context otherwise requires, for the purposes of disciplining a pharmacy technician or pharmacy technician trainee, "unprofessional conduct" means the following, whether occurring in this state or elsewhere:

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- 1. Being addicted to the use of USING alcohol or other drugs to such a degree as to render the licensee unfit to perform the licensee's employment duties.
- 2. Violating a federal or state law or administrative rule relating to the manufacture or distribution of drugs or devices.
- 3. Obtaining or attempting to obtain a pharmacy technician or pharmacy technician trainee license or a pharmacy technician license renewal by fraud, by misrepresentation or by knowingly taking advantage of the mistake of another person or an agency.
- 4. Having the licensee's license to practice as a pharmacy technician denied or disciplined in another jurisdiction.
- 5. Failing to comply with the mandatory continuing professional education requirements of section 32-1925, subsection H and rules adopted by the board.
- 6. Committing a felony, whether or not involving moral turpitude, or a misdemeanor involving moral turpitude or any drug-related offense. In either case, conviction by a court of competent jurisdiction or a plea of no contest is conclusive evidence of the commission.
 - 7. Working under the influence of alcohol or other drugs.
- 8. Violating a federal or state law or administrative rule relating to marijuana, prescription-only drugs, narcotics, dangerous drugs, controlled substances or precursor chemicals when determined by the board or by conviction in a federal or state court.
- 9. Failing to report in writing to the board any evidence that a pharmacist or pharmacy intern is or may be professionally incompetent, is or may be guilty of unprofessional conduct or is or may be mentally or physically unable to safely engage in the practice of pharmacy.
- 10. Failing to report in writing to the board any evidence that a pharmacy technician or pharmacy technician trainee is or may be professionally incompetent, is or may be guilty of unprofessional conduct or is or may be mentally or physically unable to safely engage in the permissible activities of a pharmacy technician or pharmacy technician trainee.
- 11. Failing to report in writing to the board any evidence that a permittee or a permittee's employee is or may be guilty of unethical conduct or is or may be $\frac{1}{100}$ violation of VIOLATING this chapter or a rule adopted under this chapter.
- 12. Committing an offense in another jurisdiction that if committed in this state would be grounds for discipline.
- 13. Knowingly filing with the board any application, renewal or other document that contains false or misleading information.
- 14. Providing false or misleading information or omitting material information in any communication to the board or the board's employees or agents.

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- 15. Violating or attempting to violate, directly or indirectly, or assisting in or abetting in the violation of, or conspiring to violate this chapter.
- 16. Violating a formal order, terms of probation, a consent agreement or a stipulation issued or entered into by the board or its executive director pursuant to this chapter.
- 17. Failing to comply with a board subpoena or failing to comply in a timely manner with a board subpoena without providing any explanation to the board for not complying with the subpoena.
- 18. Failing to report a change of the licensee's home address, contact information, employer or employer's address as required by section 32-1926.
- 19. Failing to report a change in the licensee's residency status as required by section 32-1926.01.
- Sec. 3. Section 32-1904, Arizona Revised Statutes, is amended to read:

32-1904. Powers and duties of board; immunity

- A. The board shall:
- 1. Make bylaws and adopt rules that are necessary to protect the public and that pertain to the practice of pharmacy, the manufacturing, wholesaling or supplying of drugs, devices, poisons or hazardous substances, the use of pharmacy technicians and support personnel and the lawful performance of its duties.
- 2. Fix standards and requirements to register and reregister pharmacies, except as otherwise specified.
- 3. Investigate compliance as to the quality, label and labeling of all drugs, devices, poisons or hazardous substances and take action necessary to prevent the sale of these if they do not conform to the standards prescribed in this chapter, the official compendium or the federal act.
- 4. Enforce its rules. In so doing, the board or its agents have free access, during the hours reported with the board or the posted hours at the facility, to any pharmacy, manufacturer, wholesaler, third-party logistics provider, nonprescription drug permittee or other establishment in which drugs, devices, poisons or hazardous substances are manufactured, processed, packed or held, or to enter any vehicle being used to transport or hold such drugs, devices, poisons or hazardous substances for the purpose of:
- (a) Inspecting the establishment or vehicle to determine whether any provisions of this chapter or the federal act are being violated.
- (b) Securing samples or specimens of any drug, device, poison or hazardous substance after paying or offering to pay for the sample.
- (c) Detaining or embargoing a drug, device, poison or hazardous substance in accordance with section 32–1994.

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- 5. Examine and license as pharmacists and pharmacy interns all qualified applicants as provided by this chapter.
- 6. Require each applicant for an initial license to apply for a fingerprint clearance card pursuant to section 41-1758.03. If an applicant is issued a valid fingerprint clearance card, the applicant shall submit the valid fingerprint clearance card to the board with the completed application. If an applicant applies for a fingerprint clearance card and is denied, the applicant may request that the board consider the application for licensure notwithstanding the absence of a valid fingerprint clearance card. The board, in its discretion, may approve an application for licensure despite the denial of a valid fingerprint clearance card if the board determines that the applicant's criminal history information on which the denial was based does not alone disqualify the applicant from licensure.
- 7. Issue duplicates of lost or destroyed permits on the payment of a fee as prescribed by the board.
- 8. Adopt rules to rehabilitate pharmacists and pharmacy interns as provided by this chapter.
- 9. At least once every three months, notify pharmacies regulated pursuant to this chapter of any modifications on prescription writing privileges of podiatrists, dentists, doctors of medicine, registered nurse practitioners, osteopathic physicians, veterinarians, physician assistants, optometrists and homeopathic physicians of which it receives notification from the state board of podiatry examiners, state board of dental examiners, Arizona medical board, Arizona state board of nursing, Arizona board of osteopathic examiners in medicine and surgery, Arizona state veterinary medical examining board, Arizona regulatory board of physician assistants, state board of optometry or board of homeopathic and integrated medicine examiners.
- 10. Charge a permittee a fee, as determined by the board, for an inspection if the permittee requests the inspection.
 - 11. Issue only one active or open license per individual.
- 12. Allow a licensee to regress to a lower level license on written explanation and review by the board for discussion, determination and possible action.
- 13. OPEN AN INVESTIGATION ONLY IF THE IDENTIFYING INFORMATION REGARDING A COMPLAINANT IS PROVIDED OR THE INFORMATION PROVIDED IS SUFFICIENT TO CONDUCT AN INVESTIGATION.
- 14. PROVIDE NOTICE TO AN APPLICANT, LICENSEE OR PERMITTEE USING ONLY THE INFORMATION PROVIDED TO THE BOARD THROUGH THE BOARD'S LICENSING DATABASE.
 - B. The board may:
- 1. Employ chemists, compliance officers, clerical help and other employees subject to title 41, chapter 4, article 4 and provide laboratory facilities for the proper conduct of its business.

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- 2. Provide, by educating and informing the licensees and the public, assistance in curtailing abuse in the use of drugs, devices, poisons and hazardous substances.
- 3. Approve or reject the manner of storage and security of drugs, devices, poisons and hazardous substances.
- 4. Accept monies and services to assist in enforcing this chapter from other than licensees:
 - (a) For performing inspections and other board functions.
- (b) For the cost of copies of the pharmacy and controlled substances laws, the annual report of the board and other information from the board.
- 5. Adopt rules for professional conduct appropriate to the establishment and maintenance of a high standard of integrity and dignity in the profession of pharmacy.
- 6. Grant permission to deviate from a state requirement for experimentation and technological advances.
- 7. Adopt rules for the training and practice of pharmacy interns, pharmacy technicians and support personnel.
- 8. Investigate alleged violations of this chapter, conduct hearings in respect to violations, subpoena witnesses and take such action as it deems necessary to revoke or suspend a license or a permit, place a licensee or permittee on probation or warn a licensee or permittee under this chapter or to bring notice of violations to the county attorney of the county in which a violation took place or to the attorney general.
 - 9. By rule, approve colleges or schools of pharmacy.
- 10. By rule, approve programs of practical experience, clinical programs, internship training programs, programs of remedial academic work and preliminary equivalency examinations as provided by this chapter.
- 11. Assist in the continuing education of pharmacists and pharmacy interns.
 - 12. Issue inactive status licenses as provided by this chapter.
- $13.\ \,$ Accept monies and services from the federal government or others for educational, research or other purposes pertaining to the enforcement of this chapter.
- 14. By rule, except from the application of all or any part of this chapter any material, compound, mixture or preparation containing any stimulant or depressant substance included in section 13-3401, paragraph 6, subdivision (c) or (d) from the definition of dangerous drug if the material, compound, mixture or preparation contains one or more active medicinal ingredients not having a stimulant or depressant effect on the central nervous system, provided that such admixtures are included in such combinations, quantity, proportion or concentration as to vitiate the potential for abuse of the substances that do have a stimulant or depressant effect on the central nervous system.

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- 15. Adopt rules for the revocation, suspension or reinstatement of licenses or permits or the probation of licensees or permittees as provided by this chapter.
- 16. Issue a certificate of free sale to any person that is licensed by the board as a manufacturer for the purpose of manufacturing or distributing food supplements or dietary supplements as defined in rule by the board and that wants to sell food supplements or dietary supplements domestically or internationally. The application shall contain all of the following:
- (a) The applicant's name, address, $\frac{e-mail}{}$ EMAIL address, telephone and fax number.
 - (b) The product's full, common or usual name.
- (c) A copy of the label for each product listed. If the product is to be exported in bulk and a label is not available, the applicant shall include a certificate of composition.
 - (d) The country of export, if applicable.
 - (e) The number of certificates of free sale requested.
- 17. Establish an inspection process to issue certificates of free sale or good manufacturing practice certifications. The board shall establish in rule:
 - (a) A fee to issue certificates of free sale.
 - (b) A fee to issue good manufacturing practice certifications.
 - (c) An annual inspection fee.
 - 18. Delegate to the executive director the authority to:
- (a) Void a license or permit application and deem all fees forfeited by the applicant if the applicant provided inaccurate information on the application. The applicant shall have the opportunity to correct the inaccurate information within thirty days after the initial application was reviewed by board staff and the applicant was informed of the inaccuracy.
- (b) (a) If the president or vice president of the board concurs after reviewing the case, enter into an interim consent agreement with a licensee or permittee if there is evidence that a restriction against the license or permit is needed to mitigate danger to the public health and safety. The board may subsequently formally adopt the interim consent agreement with any modifications the board deems necessary.
- (c) (b) Take no action or dismiss a complaint that has insufficient evidence that a violation of statute or rule governing the practice of pharmacy occurred.
- (d) (c) Request an applicant or licensee to provide court documents and police reports if the applicant or licensee has been charged with or convicted of a criminal offense. The executive director may do either of the following if the applicant or licensee fails to provide the requested documents to the board within thirty business days after the request:

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- (i) Close the application, deem the application fee forfeited and not consider a new application complete unless the requested documents are submitted with the application.
- (ii) Notify the licensee of an opportunity for a hearing in accordance with section 41-1061 to consider suspension of the licensee.
- $\frac{\text{(e)}}{\text{(d)}}$ Pursuant to section 36-2604, subsection B, review prescription information collected pursuant to title 36, chapter 28, article 1.
- C. At each regularly scheduled board meeting, the executive director shall provide to the board a list of the executive director's actions taken pursuant to subsection B, paragraph 18, subdivisions (a), (c) and (d) of this section since the last board meeting.
- D. THE BOARD MAY ISSUE NONDISCIPLINARY CIVIL PENALTIES OR DELEGATE TO THE EXECUTIVE DIRECTOR THE AUTHORITY TO ISSUE NONDISCIPLINARY CIVIL PENALTIES. THE NONDISCIPLINARY CIVIL PENALTIES SHALL BE PRESCRIBED BY THE BOARD IN RULE AND ISSUED USING A BOARD-APPROVED FORM. IF A LICENSEE OR PERMITTEE FAILS TO PAY A NONDISCIPLINARY CIVIL PENALTY THAT THE BOARD HAS IMPOSED ON IT, THE BOARD SHALL HOLD A HEARING ON THE MATTER. IN ADDITION TO ANY OTHER NONDISCIPLINARY CIVIL PENALTY ADOPTED BY THE BOARD, EITHER OF THE FOLLOWING ACTS OR OMISSIONS THAT IS NOT AN IMMINENT THREAT TO THE PUBLIC HEALTH AND SAFETY IS SUBJECT TO A NONDISCIPLINARY CIVIL PENALTY:
 - 1. AN OCCURRENCE OF EITHER OF THE FOLLOWING:
- (a) FAILING TO SUBMIT A REMODEL APPLICATION BEFORE REMODELING A PERMITTED FACILITY.
 - (b) FAILING TO NOTIFY THE BOARD OF THE RELOCATION OF A BUSINESS.
- 2. THE OCCURRENCE OF ANY OF THE FOLLOWING VIOLATIONS OR ANY OF THE VIOLATIONS ADOPTED BY THE BOARD IN RULE, WITH THREE OR MORE VIOLATIONS BEING PRESENTED TO THE BOARD AS A COMPLAINT:
- (a) THE LICENSEE OR PERMITTEE FAILS TO UPDATE THE LICENSEE'S OR PERMITTEE'S ONLINE PROFILE WITHIN TEN DAYS AFTER A CHANGE IN CONTACT INFORMATION, ADDRESS, TELEPHONE NUMBER OR EMAIL ADDRESS.
- (b) THE LICENSEE FAILS TO UPDATE THE LICENSEE'S ONLINE PROFILE WITHIN TEN DAYS AFTER A CHANGE IN EMPLOYMENT.
- (c) THE LICENSEE FAILS TO COMPLETE THE REQUIRED CONTINUING EDUCATION FOR A LICENSE RENEWAL.
- (d) THE LICENSEE FAILS TO UPDATE THE LICENSEE'S ONLINE PROFILE TO REFLECT A NEW PHARMACIST IN CHARGE WITHIN FOURTEEN DAYS AFTER THE POSITION CHANGE.
- (e) THE PERMITTEE FAILS TO UPDATE THE PERMITTEE'S ONLINE PROFILE TO REFLECT A NEW DESIGNATED REPRESENTATIVE WITHIN TEN DAYS AFTER THE POSITION CHANGE.
- (f) THE LICENSEE OR PERMITTEE FAILS TO NOTIFY THE BOARD OF A NEW CRIMINAL CHARGE, ARREST OR CONVICTION AGAINST THE LICENSEE OR PERMITTEE IN THIS STATE OR ANY OTHER JURISDICTION.

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- (g) THE LICENSEE OR PERMITTEE FAILS TO NOTIFY THE BOARD OF A DISCIPLINARY ACTION TAKEN AGAINST THE LICENSEE OR PERMITTEE BY ANOTHER REGULATING AGENCY IN THIS STATE OR ANY OTHER JURISDICTION.
- (h) A LICENSEE OR PERMITTEE FAILS TO RENEW A LICENSE OR PERMIT WITHIN SIXTY DAYS AFTER THE LICENSE OR PERMIT EXPIRES. IF MORE THAN SIXTY DAYS HAVE LAPSED AFTER THE EXPIRATION OF A LICENSE OR PERMIT, THE LICENSEE OR PERMITTEE SHALL APPEAR BEFORE THE BOARD.
- (i) A NEW PHARMACIST IN CHARGE FAILS TO CONDUCT A CONTROLLED SUBSTANCE INVENTORY WITHIN TEN DAYS AFTER STARTING THE POSITION.
- (j) A PERSON FAILS TO OBTAIN A PERMIT BEFORE SHIPPING INTO THIS STATE ANYTHING THAT REQUIRES A PERMIT PURSUANT TO THIS CHAPTER.
- (k) ANY OTHER VIOLATIONS OF STATUTE OR RULE THAT THE BOARD OR THE BOARD'S DESIGNEE DEEMS APPROPRIATE FOR A NONDISCIPLINARY CIVIL PENALTY.
- $rac{ extsf{D.}}{ extsf{C}}$ E. The board shall develop substantive policy statements pursuant to section 41-1091 for each specific licensing and regulatory authority the board delegates to the executive director.
- E. F. The executive director and other personnel or agents of the board are not subject to civil liability for any act done or proceeding undertaken or performed in good faith and in furtherance of the purposes of this chapter.
- Sec. 4. Section 32-1922, Arizona Revised Statutes, is amended to read:

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32-1922. Qualifications of applicant: reciprocity:

preliminary equivalency examination; honorary
certificate; fee
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- A. An applicant for licensure as a pharmacist shall:
- 1. Be of good moral character.
- 2. Be a graduate of a school or college of pharmacy or department of pharmacy of a university recognized by the board or the accreditation council for pharmacy education, or qualify under subsection D of this section.
- 3. Have successfully completed, as substantiated by proper affidavits, a program of practical experience under the direct supervision of a licensed pharmacist who is approved by the board.
- 4. Pass the pharmacist licensure examination and jurisprudence examination approved by the board. An applicant who fails an examination three times shall petition the board for permission before retaking the examination. The board shall evaluate the petition and determine whether to require additional educational training before approving each additional retake of the examination.
- 5. Pay an application fee prescribed by the board of not more than five hundred dollars \$500. An applicant for reciprocal licensure shall pay the fee prescribed in section 32-1924, subsection D.
- B. The board may license as a pharmacist, without a pharmacist licensure examination, a person who is licensed as a pharmacist by a

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 pharmacist licensure examination in some other jurisdiction if that person:

- 1. Produces satisfactory evidence to the board of having had the required secondary and professional education and training.
- 2. Is possessed of good morals as demanded of applicants for licensure and relicensure under this chapter.
- 3. Presents proof to the board's satisfaction that the person is licensed by a pharmacist licensure examination equivalent to the pharmacist licensure examination required by the board and that the person holds the license in good standing. If the applicant was examined after June 1, 1979, the applicant must present proof to the board's satisfaction of having passed the national association of boards of pharmacy licensure examination or the north American pharmacist licensure examination.
- 4. Presents proof to the board's satisfaction that any other license granted to the applicant by any other jurisdiction has not been suspended, revoked or otherwise restricted for any reason except nonrenewal or for failure to obtain the required continuing education credits in any jurisdiction where the applicant is currently licensed but not engaged in the practice of pharmacy.
 - 5. Passes a board-approved jurisprudence examination.
- C. Subsection B of this section applies only if the jurisdiction in which the person is licensed grants, under like conditions, reciprocal licensure as a pharmacist to a pharmacist who is licensed by examination in this state and the person holds a license in good standing issued by an active member board of the national association of boards of pharmacy.
- D. If an applicant for licensure is a graduate of a pharmacy degree program at a school or college of pharmacy that was not recognized by the board at the time of the person's graduation, the applicant shall pass a preliminary equivalency examination approved by the board in order to qualify to take the examinations prescribed in subsection A of this section.
- E. The preliminary equivalency examination required pursuant to subsection D of this section shall cover proficiency in English and academic areas the board deems essential to a satisfactory pharmacy curriculum.
- F. An applicant who fails the preliminary equivalency examination required pursuant to subsection D of this section shall not retake the preliminary equivalency examination until the applicant files written proof with the board that the applicant has completed additional remedial academic work previously approved by the board to correct deficiencies in the applicant's education that were indicated by the results of the applicant's last preliminary equivalency examination.
- G. A pharmacist who has been licensed in this state for at least fifty years shall be granted an honorary certificate of licensure by the board without the payment of the usual renewal fee, but that certificate

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 of licensure does not confer an exemption from any other requirement of this chapter.

- H. The board may require a pharmacist who has not been actively engaged in the practice of pharmacy for over one year to serve not more than four hundred hours in an internship training program approved by the board or its designee before the pharmacist may resume the active practice of pharmacy.
- I. An applicant must complete the application process within twelve months after submitting the application.
- Sec. 5. Section 32-1924, Arizona Revised Statutes, is amended to read:

32-1924. Licenses; fees; rules; signatures; online profiles

- A. An applicant for licensure as a pharmacist who passes the board-approved examinations shall pay the board an initial licensure fee of not more than five hundred dollars \$500.
- B. An applicant for licensure as a pharmacist, intern, OR pharmacy technician or pharmacy technician trainee shall pay a fee prescribed by the board that does not exceed fifty dollars \$50 for issuance of a wall license. On payment of a fee of not more than fifty dollars \$50, the board may issue a replacement wall license to a licensee who requests a replacement because the original was damaged or destroyed, because of a change of name or for other good cause as prescribed by the board.
- C. An applicant for licensure as an intern shall pay a fee of not more than seventy-five dollars \$75. A license issued pursuant to this subsection expires five years after it is issued. The board shall adopt rules to prescribe the requirements for the renewal of a license that expires before the pharmacy intern completes the education or training required for licensure as a pharmacist.
- D. An applicant for reciprocal licensure as a pharmacist shall pay a fee of not more than five hundred dollars \$500 for the application and expense of making an investigation of INVESTIGATING the applicant's character, general reputation and pharmaceutical standing in the jurisdiction in which the applicant is licensed.
- E. All pharmacist licenses shall bear the signatures of the executive director and a majority of the members of the board.
- F. An applicant for licensure as a pharmacy technician trainee shall submit with the application a fee prescribed by the board that does not exceed one hundred dollars \$100. A license issued pursuant to this subsection expires thirty-six months after it is issued. A pharmacy technician trainee license may not be renewed or reissued.
- G. An applicant for licensure as a pharmacy technician shall submit with the application a fee prescribed by the board that does not exceed one hundred dollars \$100.
- $\mbox{\rm H.}$ A licensee shall create an online profile using the board's licensing software.

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Sec. 6. Section 32-1925, Arizona Revised Statutes, is amended to read:

32-1925. Renewal of license of pharmacists, interns and pharmacy technicians; fees; expiration dates; penalty for failure to renew; continuing education

- A. Except for interns and pharmacy technician trainees, the board shall assign all persons who are licensed under this chapter to one of two license renewal groups. Except as provided in section 32-4301, a holder of a license certificate designated in the licensing database as even by way of verbiage or numerical value shall renew it biennially on or before November 1 of the even-numbered year, two years from AFTER the last renewal date. Except as provided in section 32-4301, a holder of a license certificate designated in the licensing database as odd by way of verbiage or numerical value shall renew it biennially on or before November 1 of the odd-numbered year, two years from AFTER the last renewal date. Failure to renew and pay all required fees on or before November 1 of the year in which the renewal is due suspends the license. The board shall vacate a suspension when the licensee pays all past due fees and REINSTATEMENT penalties. REINSTATEMENT penalties shall not exceed three hundred fifty dollars \$350. The board may waive collection of a fee or REINSTATEMENT penalty due after suspension under conditions established by a majority of the board.
- B. A person shall not apply for license renewal more than sixty days before the expiration date of the license.
- C. A person who is licensed as a pharmacist or a pharmacy technician and who has not renewed the license for five consecutive years shall furnish to the board satisfactory proof of fitness to be licensed as a pharmacist or a pharmacy technician. , in addition to the payment of all past due fees A PERSON WHOSE LICENSE HAS LAPSED FOR TWO OR MORE RENEWAL CYCLES SHALL PAY THE FEES FOR THE TWO MOST RECENT RENEWAL CYCLES and THE penalties before being reinstated.
 - D. Biennial renewal fees for licensure shall be not more than:
 - 1. For a pharmacist, two hundred fifty dollars \$250.
 - 2. For a pharmacy technician, one hundred dollars \$100.
 - 3. For a duplicate renewal license, twenty-five dollars \$25.
- E. Fees that are designated to be not more than a maximum amount shall be set by the board for the following two fiscal years beginning November 1. The board shall establish fees approximately proportionate to the maximum fee allowed to cover the board's anticipated expenditures for the following two fiscal years. Variation in a fee is not effective except at the expiration date of a license.
- F. The board shall not renew a license for a pharmacist unless the pharmacist has complied with the mandatory continuing professional pharmacy education requirements of sections 32-1936 and 32-1937.

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- G. The board shall prescribe intern licensure renewal fees that do not exceed seventy-five dollars \$75. The license of an intern who does not receive specific board approval to renew the intern license or who receives board approval to renew but who does not renew and pay all required fees before the license expiration date is suspended after the license expiration date. The board shall vacate a suspension if the licensee pays all past due fees and penalties. Penalties shall not exceed three hundred fifty dollars \$350. The board may waive collection of a fee or penalty due after suspension under conditions established by the board.
- H. The board shall not renew a license for a pharmacy technician unless that person has a current board-approved license and has complied continuing professional board-approved mandatory requirements. If a pharmacy technician prepares, compounds or dispenses prescription medications at a remote dispensing site pharmacy, the shall technician complete, in addition to board-approved mandatory continuing professional education requirements, a two-hour continuing education program on remote dispensing site pharmacy practices provided by an approved provider.
- Sec. 7. Section 32-1930, Arizona Revised Statutes, is amended to read:

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32-1930. Types of permits; restrictions on permits; discontinuance of pharmacy permit
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- 1. If approved by the board, a pharmacy, limited service pharmacy, automated prescription-dispensing kiosk, full service wholesale drug, third-party logistics provider, nonprescription drug wholesale and drug manufacturer's permit.
- 2. Drug packager or drug prepackager permit to an individual or establishment that is currently listed by the United States food and drug administration and has met the requirements of that agency to purchase, repackage, relabel or otherwise alter the manufacturer's original package of an approved drug product with the intent of reselling these items to persons or businesses authorized to possess or resell the repackaged, prepackaged or relabeled drug.
- 3. A DURABLE MEDICAL EQUIPMENT DISTRIBUTOR AND compressed medical gas distributor permit and a durable medical equipment SUPPLIER and compressed medical gas supplier permit.
- B. The board shall deny or revoke a pharmacy permit if a medical practitioner receives compensation, either directly or indirectly, from a pharmacy as a result of the practitioner's prescription orders. This does not include compensation to a medical practitioner who is the owner of a building where space is leased to a pharmacy at the prevailing rate, not resulting in a rebate to the medical practitioner.

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- C. If a pharmacy permanently discontinues operation, the permittee shall immediately surrender the permit to the executive director. The permittee shall remove all drug signs and symbols, either within or without the premises, and shall remove or destroy all drugs, devices, poisons and hazardous substances.
- D. An automated prescription-dispensing kiosk may not contain or dispense a controlled substance as defined in section 36-2501 and the controlled substances act (P.L. 91-513; 84 Stat. 1242; 21 United States Code section 802).
- Sec. 8. Section 32-1931, Arizona Revised Statutes, is amended to read:

32-1931. <u>Permit fees; issuance; expiration; renewals; online profiles</u>

- A. The board shall assign the permit of all persons or firms issued under this chapter to one of two permit renewal groups. Except as provided in section 32-4301, a holder of a permit designated in the licensing database as even by way of verbiage or numerical value shall renew it biennially on or before November 1 of the even-numbered year, two years from AFTER the last renewal date. Except as provided in section 32-4301, a holder of a permit designated in the licensing database as odd by way of verbiage or numerical value shall renew it biennially on or before November 1 of the odd-numbered year, two years from AFTER the last renewal date. Failure to renew and pay all required fees on or before November 1 of the year in which the renewal is due suspends the permit. The board shall vacate a suspension when the permittee pays penalties of not to exceed \$350 and all past due fees. The board may waive collection of a fee or penalty due after suspension under conditions established by a majority of the board.
- B. Permit fees that are designated to be not more than a maximum amount shall be set by the board for the following two fiscal years beginning November 1. The board shall establish the fees approximately proportionate to the maximum fee allowed to cover the board's anticipated expenditures for the following two fiscal years. Variation in a fee is not effective except at the expiration date of the permit.
- C. Applications for permits shall be accompanied by the following biennial fees as determined by PURSUANT TO subsection B of this section:
 - 1. A drug manufacturer's permit, not more than \$1,000.
 - 2. A pharmacy permit, not more than \$500.
- 3. A limited service pharmacy permit or an automated prescription-dispensing kiosk permit, not more than \$500.
- 4. A full service wholesale drug permit or a third-party logistics provider permit, not more than \$1,000.
 - 5. A nonprescription drug wholesale permit, not more than \$500.
 - 6. A drug repackager's permit, not more than \$1,000.

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- 7. A DURABLE MEDICAL EQUIPMENT DISTRIBUTOR AND compressed medical gas distributor permit, not more than \$200.
- 8. A durable medical equipment SUPPLIER and compressed medical gas supplier permit, not more than \$100.
- D. If an applicant is found to be satisfactory to the board, the executive director shall issue to the applicant a permit for each pharmacy, manufacturer, wholesaler or other place of business in which drugs are sold, manufactured, compounded, dispensed, stocked, exposed or offered for sale, for which application is made.
 - E. Permits issued under this section are not transferable.
- F. If a permittee does not apply for renewal, the permit expires pursuant to subsection A of this section. A person may activate and renew an expired permit by filing the required application and fee. Renewal thirty days after the expiration date of a permit may be made only on payment of the required biennial renewal fee, all past due fees and a penalty of one-half of the amount of the applicable biennial renewal fee. The board may waive the collection of a fee or penalty due after suspension pursuant to conditions prescribed by the board.
- G. A permittee shall create an online profile using the board's licensing software.
- Sec. 9. Section 32-1937, Arizona Revised Statutes, is amended to read:

32-1937. Exceptions to continuing education requirements

- A. The requirements of continuing professional pharmacy education provided in section 32-1936 do not apply to licensees during the year of their graduation from an accredited college of pharmacy BEGINNING THE DATE OF INITIAL LICENSURE UNTIL THE DATE OF THE FIRST LICENSE RENEWAL.
- B. The board may make exceptions from the requirements of section 32-1936 in emergency or hardship cases or for good cause shown based on a written request for an exception from the requirements.
- C. Pharmacists who are exempted from the requirements of continuing professional pharmacy education pursuant to subsection B of this section shall satisfactorily pass a written examination approved by the board for such THAT purpose prior to BEFORE license renewal.
- Sec. 10. Section 32-1941, Arizona Revised Statutes, is amended to read:

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32-1941. Third-party logistics providers; permit required; designated representative; fingerprinting requirements
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- A. A third-party logistics provider that engages in the logistics services of prescription or over-the-counter dangerous drugs or dangerous devices into, within or from this state shall hold a third-party logistics provider permit in this state.
- B. A third-party logistics provider shall comply with storage practices, including all of the following:

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- 1. Maintain access to warehouse space of ${\sf A}$ suitable size to facilitate safe operations, including a suitable area to quarantine a suspect product.
 - 2. Maintain adequate security.
 - 3. Have written policies and procedures to:
- (a) Address the receipt, security, storage, inventory, shipment and distribution of a product.
- (b) Identify, record and report confirmed significant losses or thefts in the United States.
 - (c) Correct errors and inaccuracies in inventories.
 - (d) Provide support for manufacturer recalls.
- (e) Prepare for, protect against and address any reasonably foreseeable crisis that affects a facility's security or operation, such as an employee strike, A fire or A flood.
- (f) Ensure that any expired product is segregated from other products and returned to the manufacturer, repackager or agent of the manufacturer or repackager or is destroyed.
- (g) Maintain records reflecting the receipt and distribution of products and supplies and records of inventories.
- (h) Quarantine or destroy a suspect product if directed to do so by the respective manufacturer, wholesale distributor or dispenser or an authorized governmental agency.
- C. A third-party logistics provider shall make its facility available to the board for inspection during regular business hours to ensure compliance with this section.
- D. A third-party logistics provider shall have a designated representative at each facility who has not been convicted of any felony violation under any federal, state or local law relating to wholesale or retail prescription or over-the-counter dangerous drugs or dangerous devices distribution or the distribution of controlled substances.
- E. A third-party logistics provider shall provide the board on the board's request with a list of all manufacturers, wholesale distributors, and dispensers AND DURABLE MEDICAL EQUIPMENT SUPPLIERS for whom the third-party logistics provider provides services at a facility.
- F. A third-party logistics provider's designated representative shall have a valid fingerprint clearance card issued pursuant to title 41, chapter 12, article 3.1, which shall be submitted with the completed application. If the third-party logistics provider changes its designated representative, the new designated representative shall have a valid fingerprint clearance card issued pursuant to title 41, chapter 12, article 3.1 and submitted to the board before the change in representation is made.

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Sec. 11. Section 32-1967, Arizona Revised Statutes, is amended to read:

32-1967. Acts constituting misbranding of a drug or device; exceptions; interpretation of misleading label; definition

- A. A drug or device is misbranded:
- 1. If its labeling is false or misleading in any particular.
- 2. If in package form unless it bears a label containing both:
- (a) The name and place of business of the manufacturer, packer or distributor.
- (b) An accurate statement of the quantity of the contents in terms of weight, measure or numerical count.
- 3. If any word, statement or other information required by or under authority of this chapter to appear on the label or labeling is not prominently placed on the label or labeling. Compliance with the federal act shall be deemed compliance with this chapter except for compliance with paragraph 16 of this subsection.
- 4. If it is for use by humans and contains any quantity of the narcotic or hypnotic substance alpha-eucaine, barbituric acid, beta-eucaine, bromal, cannabis, carbromal, chloral, coca, cocaine, codeine, heroin, marijuana, morphine, opium, paraldehyde, peyote or sulfonmethane, or any chemical derivative of such substance, which derivative or other substance has been found to be habit-forming, unless its label bears the name and quantity or proportion of such substance or derivative.
- 5. If it is a drug unless its label bears, to the exclusion of any other nonproprietary name, both:
- (a) The established name of the drug, if there is an established name.
- (b) In case it is fabricated from two or more ingredients, the established name and quantity of each active ingredient, including the kind and quantity or proportion of any alcohol, and also including, whether active or not, the established name and quantity or proportion of any bromides, ether, chloroform, atropine, hyoscine, hyoscyamine, arsenic, digitalis, digitalis glycosides, mercury, strychnine or thyroid, or derivative or preparation of any such substances, provided that the requirements for stating the quantity of the active ingredients, other than those specifically named in this subdivision, apply only to prescription drugs.
 - 6. Unless its labeling bears both:
 - (a) Adequate directions for use.
- (b) Adequate warnings against use in those pathological conditions or by children where its use may be dangerous to health, or against unsafe dosage or methods or duration of administration or application, in a manner and form as are necessary for the protection of users.

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- 7. If it is recognized in an official compendium, unless it is packed and labeled as prescribed in such compendium, provided that the method of packing may be modified with the consent of the board.
- 8. If it has been found by the board to be a drug or device liable to deterioration, unless it is packaged in that form and manner, and its label bears a statement of such precautions, as the rules issued by the board require as necessary for the protection of public health.
- $9.\ \mbox{If its container}$ is so made, formed or filled as to be misleading.
 - 10. If it is an imitation of another drug or device.
- 11. If it is offered for sale under the name of another drug or device.
- 12. If it is dangerous to health when used in the dosage or manner or with the frequency or duration prescribed, recommended or suggested in the labeling of the drug or device.
- 13. If it is a color additive, the intended use of which in or on drugs or devices is for the purpose of coloring only, unless its packaging and labeling are in conformity with such packaging and labeling requirements applicable to such color additive in the federal act or board rule.
- 14. In the case of any prescription-only drug or controlled substance distributed or offered for sale in this state, unless the manufacturer, packer or distributor of such drug or substance includes in all advertisements and other printed matter with respect to that drug a true statement of:
 - (a) The established name.
 - (b) The formula showing quantitatively each ingredient.
- (c) Other information in brief summary relating to side effects, contraindications or effectiveness as required in board rules or the federal act.
- 15. If a trademark, trade name or other identifying mark, imprint or device of another drug or device or any likeness of another drug or device has been placed on the drug or device or on its container with intent to defraud.
- 16. In the case of any prescription-only drug or controlled substance, if in final dosage form unless it bears a label containing both:
- (a) The name and place of business of the manufacturer, and if different, the packer or distributor.
- (b) An accurate statement of the quantity of the contents in terms of weight, measure or numerical count.
- 17. In the case of any foreign dangerous drug, if it is not approved by the United States food and drug administration or is obtained outside of the licensed supply chain regulated by the United States food and drug administration, the board or the department of health services.

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44 45 This paragraph does not apply to a foreign dangerous drug that is authorized for use by a state law or that is imported lawfully under the FEDERAL food, drug, and cosmetic act (21 United States Code section 301, et seq.) or pursuant to an announcement by the United States food and drug exercise of enforcement administration of the discretion instances, including clinical research purposes, drug shortages. development of countermeasures against chemical, biological, radiological and nuclear terrorism agents, or pandemic influenza preparedness and response.

- B. Drugs and devices that are to be processed, labeled or repacked at establishments other than those where originally processed or packed are exempt from any labeling or packaging requirements of this chapter, provided that such drugs and devices are being delivered, manufactured, processed, labeled, repacked or otherwise held in compliance with board rules or under the federal act.
- C. If an article is alleged to be misbranded because the labeling is misleading, then in determining whether the labeling is misleading there shall be taken into account, among other things, not only representations made or suggested by statement, word, design, device or any combination of them, but also the extent to which the labeling fails to reveal facts material in the light of such representations, or material with respect to consequences which THAT may result from the use of the article to which the labeling relates under the conditions of use prescribed in the labeling or under such conditions of use as are customary or usual.
- D. A drug or device is not considered misbranded if it is either of the following:
- 1. Intended for the use in pharmaceutical compounding by a licensed pharmacist, physician, drug manufacturer or distributor or registered outsourcing facility in compliance with the requirements of THIS chapter 18 of this title and the FEDERAL food, drug, and cosmetic act (21 United States Code section 321a and 321b 321).
- 2. Mislabeled or incorrectly filled because of a filling error by a pharmacy or a pharmacist.
- E. This section does not apply to any drug or device, whether or not approved by the United States food and drug administration, that is manufactured, packed or distributed for use in pharmaceutical compounding by a licensed pharmacist, physician, drug manufacturer or distributor or registered outsourcing facility in compliance with the requirements of THIS chapter 18 of this title, and the FEDERAL food, drug, and cosmetic act (21 United States Code section 321a and 321b 321).
- F. For the purposes of this section, "dangerous drug" means any drug that is unsafe for self-use in humans or animals and includes:
- 1. Any drug that bears the legend: "Caution: federal law prohibits dispensing without prescription", "Rx only", or words of similar import.

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- 2. Any device that bears the statement: "Caution: federal law restricts this device to sale by or on the order of a _____", "Rx only", or words of similar import, the blank to be filled in with the designation of the practitioner licensed to use or order use of the device.
- 3. Any other drug or device that by federal or state law can be lawfully dispensed only on prescription.
- Sec. 12. Section 32-1974, Arizona Revised Statutes, is amended to read:

32-1974. Pharmacists; administration of immunizations, vaccines and emergency medications; certification; reporting requirements; advisory committee; definitions

- A. Except as prescribed pursuant to subsection I of this section, a pharmacist who is licensed pursuant to this chapter and who meets the requirements of this section may administer the following to adults without a prescription order pursuant to rules and protocols adopted by the board pursuant to this section:
- 1. Immunizations or vaccines recommended for adults by the United States centers for disease control and prevention.
- 2. Immunizations or vaccines recommended by the United States centers for disease control and prevention's health information for international travel.
- B. A pharmacist who is licensed pursuant to this chapter and who meets the requirements of this section may administer the following to minors without a prescription order pursuant to rules and protocols adopted by the board pursuant to this section:
- 1. Influenza immunizations or vaccines to a person who is at least three years of age.
- 2. Booster doses for the primary adolescent series as recommended by the United States centers for disease control and prevention.
- 3. Immunizations or vaccines recommended by the United States centers for disease control and prevention to a person who is at least thirteen years of age.
- C. Except as prescribed in subsection B of this section, a pharmacist who is licensed pursuant to this chapter and who meets the requirements of this section may administer immunizations and vaccines, including the first dose for the primary adolescent series, to a person who is at least six years of age but under thirteen years of age only with a prescription order and pursuant to rules and protocols adopted by the board pursuant to this section.
- D. A pharmacist who wishes to administer immunizations and vaccines pursuant to this section must be certified to do so by the board. The board shall issue a certificate to a pharmacist who meets board requirements for certification as prescribed by the board by rule.

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- E. A pharmacist who is certified to administer immunizations and vaccines pursuant to this section may administer without a prescription order:
- 1. Emergency medication to manage an acute allergic reaction to an immunization, vaccine or medication in accordance with the United States centers for disease control and prevention immunization guidelines.
- 2. Immunizations or vaccines to any person regardless of age during a public health emergency response of this state pursuant to section 36-787.
- F. A pharmacist who administers an immunization, vaccine or emergency medication pursuant to this section must:
- 1. Report the administration to the person's identified primary care provider or physician within forty-eight hours after administering the immunization, vaccine or emergency medication and as prescribed by the board by rule. Failure to report the administration of an immunization, vaccine or emergency medication pursuant to this section is a violation of section 32-1901.01, subsection B, paragraph 2. The pharmacist shall make a reasonable effort to identify the person's primary care provider or physician by one or more of the following methods:
- (a) Checking any adult immunization information system or vaccine registry established by the department of health services.
 - (b) Checking pharmacy records.
- (c) Requesting the information from the person or, in the case of a minor, the person's parent or guardian.
- 2. Report information to any adult immunization information system or vaccine registry established by the department of health services.
- 3. Maintain a record of the immunization pursuant to title 12, chapter 13, article 7.1 and as prescribed by the board by rule.
- 4. Report to the person's identified primary care provider or physician, within twenty-four hours of occurrence, any adverse reaction that is reported to or witnessed by the pharmacist and that is listed by the vaccine manufacturer as a contraindication to further doses of the vaccine.
- 5. Participate in any federal vaccine adverse event reporting system or successor database.
- G. This section does not establish a cause of action against a patient's primary care provider or physician for any adverse reaction, complication or negative outcome arising from the administration of any immunization, vaccine or emergency medication by a pharmacist to the patient pursuant to this section if it is administered without a prescription order written by the patient's primary care provider or physician.
- H. The board shall adopt rules for the administration of vaccines or immunizations pursuant to this section regarding:

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- 1. Protocols that are based on protocols approved by the United States centers for disease control and prevention and any advisory committee appointed by the board for the purpose of recommending protocols.
 - 2. Recordkeeping and reporting requirements.
- 3. Requirements and qualifications for pharmacist certification pursuant to this section.
- 4. Vaccine information and educational materials for those requesting vaccines and immunizations.
- 5. The administration of emergency medication pursuant to this section.
- The department of health services, by rule, shall establish and Ι. maintain a list of immunizations or vaccines that may be administered to adults by a pharmacist only pursuant to a prescription order. In adopting and maintaining this list, the department is exempt from the rulemaking requirements of title 41, chapter 6. The department shall adopt its initial rules within six months after receipt of the recommendations of the advisory committee appointed by the board and shall hold one public hearing before implementing the rules and any amendments to the rules. The list shall include those immunizations or vaccines listed in the United States centers for disease control and prevention's recommended adult immunization schedule or recommended by the United States centers for disease control and prevention's health information for international travel that have adverse reactions that could cause significant harm to a patient's health. A pharmacist may not administer immunizations or vaccines without a prescription order pursuant to this section before the department has established the list pursuant to this subsection. The board may not authorize a pharmacist to administer new immunizations or vaccines without a prescription order pursuant to this section until the department reviews the new immunizations and vaccines to determine if they should be added to the list established pursuant to this subsection.
- J. The board may appoint an advisory committee to assist the board in adopting and amending rules and developing protocols relating to the administration of immunizations, vaccines and emergency medications and certification requirements.
- K. A pharmacy intern who is certified by the board to administer immunizations and vaccines pursuant to this section may do so only in the presence and under the immediate personal supervision of a pharmacist who is certified as prescribed in this section.
- L. This section does not prevent a pharmacist who administers an immunization or vaccine from participating in the federal vaccines for children program.
- ${\sf M.}$ A pharmacist may not administer an immunization or vaccine to a minor without the consent of the minor's parent or guardian.
 - N. For the purposes of this section:

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- 1. "Emergency medication" means emergency epinephrine and antihistamines in accordance with the United States centers for disease control and prevention immunization guidelines.
- 2. "Primary adolescent series" means those immunizations or vaccines recommended by the United States centers for disease control and prevention for children starting at age eleven or twelve.
- Sec. 13. Section 32-1982, Arizona Revised Statutes, is amended to read:

32-1982. <u>Full-service wholesale permittees; bonds; designated</u> <u>representatives; fingerprinting requirements</u>

- A. A full service FULL-SERVICE wholesale permittee that engages in the wholesale distribution of prescription-only drugs into, within or from this state must maintain a bond AS REQUIRED BY FEDERAL LAW and have a designated representative. IF THE FULL-SERVICE WHOLESALE PERMITTEE CHANGES ITS DESIGNATED REPRESENTATIVE, THE NEW DESIGNATED REPRESENTATIVE MUST POSSESS AND SUBMIT A VALID FINGERPRINT CLEARANCE CARD BEFORE THE CHANGE IN REPRESENTATION IS MADE.
- B. The designated representative of a $\frac{\text{full service}}{\text{must:}}$ FULL-SERVICE wholesale permittee must:
 - 1. Be at least twenty-one years of age.
- 2. Have been employed full time for at least three years in a pharmacy or with a full service wholesale permittee in a capacity related to the dispensing and distribution of, and record keeping relating to, prescription-only drugs.
- 3. 2. Be employed by the $\frac{\text{full service}}{\text{permittee}}$ FULL-SERVICE wholesale permittee in a managerial level position.
- 4. 3. Be actively involved in the daily operation of the wholesale distribution of prescription-only drugs.
- 5. 4. Be physically present at the full service FULL-SERVICE wholesale permittee facility during regular business hours unless the absence of the designated representative is authorized.
- 6. 5. Serve as a designated representative for only one full service FULL-SERVICE wholesale permittee.
- 7. 6. Not have any criminal convictions under any federal, state or local laws relating to wholesale or retail prescription-only drug distribution or distribution of controlled substances.
- 7. POSSESS A VALID FINGERPRINT CLEARANCE CARD ISSUED PURSUANT TO TITLE 41, CHAPTER 12, ARTICLE 3.1.
- C. The board may require the applicant's designated representative to submit a full set of fingerprints to the board. The board shall submit the fingerprints to the department of public safety for the purpose of obtaining a state and federal criminal records check pursuant to section 41-1750 and Public Law 92-544. The department of public safety may exchange the fingerprint data with the federal bureau of investigation. The board may charge each applicant a fee determined by the department of

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public safety. The board shall forward this fee to the department of public safety.

D. The board shall require every full service wholesale permittee that is applying for an initial permit or renewal of a permit to submit a bond of at least one hundred thousand dollars or other equivalent means of security acceptable to the board. The board may use this bond to secure payment of any fines or penalties that are imposed by the board and any fees or costs that are incurred by the board regarding the permit authorized by law and that the permittee fails to pay within thirty days after the fine, penalty or cost becomes final. The bond must cover all permits held by the permittee in this state.

E. The board may waive the bond requirement if the full service wholesale permittee has previously obtained a comparable surety bond or other equivalent means of security for the purpose of licensure in another state where the full service wholesale permittee possesses a valid license in good standing.

F. C. For the purposes of this article, a full service FULL-SERVICE wholesale permittee does not include a hospital, chain pharmacy warehouse or third party THIRD-PARTY logistics provider.

Sec. 14. Section 36-2602, Arizona Revised Statutes, is amended to read:

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36-2602. <u>Controlled substances prescription monitoring program; contracts; retention and maintenance of records</u>
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- A. The board shall adopt rules to establish a controlled substances prescription monitoring program. The program shall:
 - 1. Be operated, monitored and maintained by the board.
 - 2. Be staffed by the board.
- 3. Include a computerized central database tracking system to track the prescribing, dispensing and consumption of schedule II, III, IV and V controlled substances that are dispensed by a medical practitioner or by a pharmacy that holds a valid license or permit issued pursuant to title 32. The database shall include data from the department of health services that identifies residents of this state who possess a registry identification card issued pursuant to chapter 28.1 of this title. The tracking system shall not interfere with the legal use of a controlled substance for the management of MANAGING severe or intractable pain.
- 4. Assist law enforcement to identify illegal activity related to the prescribing, dispensing and consumption of CONSUMING schedule II, III, IV and V controlled substances.
- 5. Provide information to patients, medical practitioners and pharmacists to help avoid the inappropriate use of schedule II, III, IV and V controlled substances.

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- 6. Be designed to minimize inconvenience to patients, prescribing medical practitioners and pharmacies while effectuating the collection and storage of information.
- B. The board may enter into private or public contracts, including intergovernmental agreements pursuant to title 11, chapter 7, article 3, to ensure the effective operation of the program. Each contractor must comply with the confidentiality requirements prescribed in this article and is subject to the criminal penalties prescribed in section 36-2610.
- C. The board shall maintain medical THE FOLLOWING records information in the program pursuant to the standards prescribed in section 12-2297 FOR THE FOLLOWING PERIODS OF TIME:
- 1. A RECORD OF DISPENSING A CONTROLLED SUBSTANCE FOR SEVEN YEARS AFTER THE DATE THE CONTROLLED SUBSTANCE WAS DISPENSED.
- 2. AFFIDAVITS FOR THE PURPOSE OF AN OPEN INVESTIGATION BY LAW ENFORCEMENT FOR TWO YEARS.
- 3. COURT ORDERS REQUESTING MEDICAL RECORD INFORMATION IN THE PROGRAM FOR TWO YEARS.
- 4. A PATIENT'S REQUEST OF THE PATIENT'S OWN PRESCRIPTION HISTORY FOR TWO YEARS.
 - 5. A PRESCRIBER REPORT FOR TWO YEARS.
- Sec. 15. Section 36-2604, Arizona Revised Statutes, is amended to read:

36-2604. <u>Use and release of confidential information:</u> definitions

- A. Except as otherwise provided in this section, prescription information submitted to the board pursuant to this article is confidential and is not subject to public inspection. The board shall establish procedures to ensure the privacy and confidentiality of patients and that patient information that is collected, recorded and transmitted pursuant to this article is not disclosed except as prescribed in this section.
- B. The board or its designee shall review the prescription information collected pursuant to this article. If the board or its designee has reason to believe an act of unprofessional or illegal conduct has occurred, the board or its designee shall notify the appropriate professional licensing board or law enforcement or criminal justice agency and provide the prescription information required for an investigation. The board may delegate the duties prescribed in this subsection to the executive director pursuant to section 32-1904.
- C. The board may release data collected by the program to the following:
- 1. A person who is authorized to prescribe or dispense a controlled substance SUBSTANCES, or a delegate who is authorized by the prescriber or dispenser, to assist that person to provide medical or pharmaceutical care to a patient or to evaluate a patient.

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- 2. An individual who requests the individual's own prescription monitoring information pursuant to section 12-2293.
- 3. A medical practitioner regulatory board established pursuant to title 32, chapter 7, 11, 13, 14, 15, 16, 17, 18, 25 or 29.
- 4. A local, state or federal law enforcement or criminal justice agency. Except as required pursuant to subsection B of this section, the board shall provide this information only if the requesting agency states in writing that the information is necessary for an open investigation or complaint.
- 5. The Arizona health care cost containment system administration and contractors regarding persons who are receiving services pursuant to chapters 29 and 34 of this title. Except as required pursuant to subsection B of this section, the board shall provide this information only if the administration or a contractor states in writing that the information is necessary for an open investigation or complaint, for performing a drug utilization review for controlled substances to help combat opioid overuse or abuse or for ensuring the continuity of care.
- 6. A person who is serving a lawful order of a court of competent jurisdiction.
- 7. A person who is authorized to prescribe or dispense a controlled substance SUBSTANCES and who performs an evaluation on an individual pursuant to section 23-1026.
- 8. A county medical examiner or alternate medical examiner who is directing an investigation into the circumstances surrounding a death as described in section 11-593 or a delegate who is authorized by the county medical examiner or alternate medical examiner.
- 9. The department of health services regarding persons who are receiving or prescribing controlled substances in order to implement a public health response to address opioid overuse or abuse, including a review pursuant to section 36-198. Except as required pursuant to subsection B of this section, the board shall provide this information only if the department states in writing that the information is necessary to implement a public health response to help combat opioid overuse or abuse.
- D. FOR A FEE DETERMINED BY THE BOARD, the board may provide data to public or private entities for statistical, research or educational purposes after removing information that could be used to identify individual patients or persons who received prescriptions from dispensers.
- E. A person who is authorized to prescribe or dispense a controlled substance SUBSTANCES or the chief medical officer of the administration or a contractor shall deactivate a delegate within five business days after an employment status change, the request of the delegate or the inappropriate use of the controlled substances prescription monitoring program's central database tracking system.
 - F. For the purposes of this section:

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- 1. "Administration" and "contractor" have the same meanings prescribed in section 36-2901.
 - 2. "Delegate" means any of the following:
- (a) A licensed health care professional who is employed in the office of or in a hospital with the prescriber or dispenser.
- (b) An unlicensed medical records technician, medical assistant or office manager who is employed in the office of or in a hospital with the prescriber or dispenser and who has received training regarding both the health insurance portability and accountability act privacy standards (45 Code of Federal Regulations part 164, subpart E) and security standards (45 Code of Federal Regulations part 164, subpart C).
- (c) A forensic pathologist, medical death investigator or other qualified person who is assigned duties in connection with a death investigation pursuant to section 11-594.
- (d) A licensed pharmacy technician trainee, pharmacy technician or pharmacy intern who works in a facility with the dispenser.
- (e) Any employee of the administration or a contractor who is authorized by the administration's or contractor's chief medical officer.
- Sec. 16. Section 36-2607, Arizona Revised Statutes, is amended to read:

36-2607. Disciplinary action

- A. The registrant's professional licensing board may revoke or suspend a registrant's registration or may place the registrant on probation for any of the following:
- 1. The registrant's professional licensing board determines that the registration was obtained by fraudulent means.
- 2. The registrant's professional licensing board takes action to revoke, suspend or place on probation the registrant's license, permit or registration to prescribe or dispense drugs.
 - 3. The registration was issued through error.
- 4. The registrant knowingly files with the board any application, renewal or other document that contains false or misleading information or the registrant gives false or misleading testimony to the board.
- 5. The registrant knowingly makes a false report or record required by this article.
- 6. A REGISTRANT THAT DISPENSES CONTROLLED SUBSTANCES DOES NOT RESOLVE DISCREPANCIES SUBMITTED TO THE PROGRAM'S CENTRAL DATABASE TRACKING SYSTEM WITHIN THIRTY BUSINESS DAYS AFTER BEING NOTIFIED OF THE ERROR BY THE BOARD.
- 7. A REGISTRANT THAT DISPENSES CONTROLLED SUBSTANCES DOES NOT RESOLVE A FAILED ATTEMPT OR MISSING TRANSMISSION TO THE PROGRAM'S CENTRAL DATABASE TRACKING SYSTEM WITHIN THIRTY BUSINESS DAYS AFTER THE OCCURRENCE.
- B. The board may deny a registration to an applicant for the grounds prescribed in subsection A OF THIS SECTION.

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C. In addition to any other law, a licensed or permitted medical practitioner, pharmacist or pharmacy that fails to comply with the requirements of this article is subject to disciplinary action by the medical practitioner's, pharmacist's or pharmacy's professional licensing board. The board of pharmacy shall report to the appropriate professional licensing board the failure of a licensed or permitted medical practitioner, pharmacist or pharmacy to comply with the requirements of this article.

Sec. 17. Section 36-2608, Arizona Revised Statutes, is amended to read:

36-2608. Reporting requirements; waiver; exceptions

- A. If a medical practitioner dispenses a controlled substance listed in section 36-2513, 36-2514, 36-2515 or 36-2516, or if a prescription for a controlled substance listed in any of those sections OR NALOXONE HYDROCHLORIDE OR ANY OTHER OPIOID ANTAGONIST THAT IS APPROVED BY THE UNITED STATES FOOD AND DRUG ADMINISTRATION is dispensed by a pharmacy in this state, a health care facility in this state for outpatient use or a board-permitted nonresident pharmacy for delivery to a person residing in this state, the medical practitioner, health care facility or pharmacy must report the following information as applicable and as prescribed by the board by rule:
- 1. The name, address, telephone number, prescription number and United States drug enforcement administration controlled substance registration number of the dispenser.
- 2. The name, address and date of birth of the person for whom the prescription is written.
- 3. The name, address, telephone number and United States drug enforcement administration controlled substance registration number of the prescribing medical practitioner.
- 4. The name, strength, quantity, dosage and national drug code number of the schedule II, III, IV or V controlled substance OR NALOXONE HYDROCHLORIDE OR OTHER OPIOID ANTAGONIST dispensed.
 - 5. The date the prescription was dispensed.
- 6. The number of refills, if any, authorized by the medical practitioner.
- B. Except as provided in subsection D of this section, a dispenser must use the September 28, 2011 version 4, release 2 standard implementation guide for prescription monitoring programs published by the American society for automation in pharmacy or any subsequent version or release of that guide to report the required information.
- C. The board shall allow the reporter to transmit the required information by electronic data transfer if feasible or, if not feasible, on reporting forms as prescribed by the board. The reporter shall submit the required information once each day.

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- D. A dispenser who does not have an automated recordkeeping system capable of producing an electronic report in the established format may request a waiver from electronic reporting by submitting a written request to the board. The board shall grant the request if the dispenser agrees in writing to report the data by submitting a completed universal claim form as prescribed by the board by rule.
- E. The board by rule may prescribe the prescription form to be used in prescribing a schedule II, III, IV or V controlled substance if the board determines that this would facilitate the reporting requirements of this section.
- F. The reporting requirements of this section do not apply to the following:
- 1. A controlled substance $\overline{\text{THAT}}$ IS administered directly to a patient.
- 2. A controlled substance THAT IS dispensed by a medical practitioner at a health care facility licensed by this state if the quantity dispensed is limited to an amount adequate to treat the patient for a maximum of seventy-two hours with not more than two seventy-two-hour cycles within any fifteen-day period.
 - 3. A controlled substance sample.
- 4. The wholesale distribution of a schedule II, III, IV or V controlled substance. For the purposes of this paragraph, "wholesale distribution" has the same meaning prescribed in section 32-1981.
- 5. A facility that is registered by the United States drug enforcement administration as a narcotic treatment program and that is subject to the recordkeeping provisions of 21 Code of Federal Regulations section 1304.24.
- G. A PHARMACIST WHO DISPENSES NALOXONE HYDROCHLORIDE OR ANOTHER OPIOID ANTAGONIST TO AN INDIVIDUAL PURSUANT TO SECTION 32-1979 SHALL REPORT THE INFORMATION LISTED IN SUBSECTION A, PARAGRAPHS 1, 2, 3 AND 5 OF THIS SECTION AND THE NAME, STRENGTH, QUANTITY, DOSAGE AND NATIONAL DRUG CODE NUMBER AS PRESCRIBED BY THE BOARD BY RULE PURSUANT TO SUBSECTION A OF THIS SECTION.
- H. NALOXONE HYDROCHLORIDE OR ANY OTHER OPIOID ANTAGONIST SHALL NOT BE VIEWABLE IN THE PATIENT UTILIZATION REPORT.
- Sec. 18. Section 41-619.51, Arizona Revised Statutes, is amended to read:

41-619.51. <u>Definitions</u>

In this article, unless the context otherwise requires:

1. "Agency" means the supreme court, the department of economic security, the department of child safety, the department of education, the department of health services, the department of juvenile corrections, the department of emergency and military affairs, the department of public safety, the department of transportation, the state real estate department, the department of insurance and financial institutions, the

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Arizona game and fish department, the Arizona department of agriculture, the board of examiners of nursing care institution administrators and assisted living facility managers, the state board of dental examiners, the Arizona state board of pharmacy, or the board of physical therapy or the state board of technical registration.

- 2. "Board" means the board of fingerprinting.
- registry exception" means "Central notification department of economic security, the department of child safety or the department of health services, as appropriate, pursuant to section 41-619.57 that the person is not disqualified because of a central registry check conducted pursuant to section 8-804.
- 4. "Expedited review" means an examination, in accordance with board rule, of the documents an applicant submits by the board or its hearing officer without the applicant being present.
- 5. "Good cause exception" means the issuance of a fingerprint clearance card to an employee pursuant to section 41-619.55.
- "Person" means a person who is required to be fingerprinted pursuant to this article or who is subject to a central registry check and any of the following:
 - (a) Section 3-314.
 - (b) Section 8-105.
 - (c) Section 8-322.
 - (d) Section 8-463.
 - (e) Section 8-509.
 - (f) Section 8-802.
- (g) Section 8-804.
 - (h) Section 15-183.
 - (i) Section 15-503.
- (j) Section 15-512.
- 29 30 (k) Section 15-534.
- 31
 - (1) Section 15-763.01.
- Section 15-782.02. 32 (m)
- 33 (n) Section 15-1330.
- (o) Section 15-1881. 34
- 35 (p) Section 17-215.
- 36 Section 28-3228. (p)
 - Section 28-3413. (r)
 - Section 32-122.02. (s)
 - (t) Section 32-122.05.
- 40 (u) Section 32-122.06.
- 41 (v) Section 32-1232.
- (w) Section 32-1276.01. 42
- 43 (x) Section 32-1284.
- Section 32-1297.01. 44 (y)
- 45 (z) Section 32-1904.

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(aa)

Section 32-1941.

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                        Section 36-3008.
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                        Section 41-619.53.
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                        Section 41-1964.
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                        Section 41-1967.01.
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           (xx)
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                        Section 41-1968.
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           <del>(yy)</del>
                  (zz) Section 41-1969.
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           (22) (aaa) Section 41-2814.
           (ada) (bbb) Section 46-141, subsection A or B.
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           (bbb) (ccc) Section 46-321.
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                      Section 41-1758, Arizona Revised Statutes, is amended to
           Sec. 19.
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     read:
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           41-1758. <u>Definitions</u>
            In this article, unless the context otherwise requires:
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                "Agency" means the supreme court, the department of economic
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     security, the department of child safety, the department of education, the
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     department of health services, the department of juvenile corrections, the
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     department of emergency and military affairs, the department of public
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     safety.
               the
                     department of transportation, the state
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     department, the department of insurance and financial institutions, the
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     board of fingerprinting, the Arizona game and fish department, the Arizona
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     department of agriculture, the board of examiners of nursing care
     institution administrators and assisted living facility managers, the
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     state board of dental examiners, the Arizona state board of pharmacy, or
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     the board
                  of
                       physical therapy or the state board of technical
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     registration.
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- 2. "Division" means the fingerprinting division in the department of public safety.
- "Electronic or internet-based fingerprinting services" means a secure system for digitizing applicant fingerprints and transmitting the applicant data and fingerprints of a person or entity submitting fingerprints to the department of public safety for any authorized purpose under this title. For the purposes of this paragraph, "secure system" means a system that complies with the information technology security policy approved by the department of public safety.
- 4. "Good cause exception" means the issuance of a fingerprint clearance card to an applicant pursuant to section 41-619.55.
- "Person" means a person who is required to be fingerprinted pursuant to any of the following:

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(a) Section 3-314.
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- (b) Section 8-105.
- (c) Section 8-322.
- (d) Section 8-463.
- - (e) Section 8-509.
- (f) Section 8-802.
- (g) Section 15-183.
 - (h) Section 15-503.
- 22 (i) Section 15-512.
 - (j) Section 15-534.
- 24 (k) Section 15-763.01.
- 25 (1) Section 15-782.02.
- 26 (m) Section 15-1330.
 - (n) Section 15-1881.
- (o) Section 17-215. 28
- 29
 - Section 28-3228. (p)
- 30 Section 28-3413. (p)
 - (r) Section 32-122.02.
 - Section 32-122.05. (s)
- Section 32-122.06. 33 (t)
- 34 (u) Section 32-1232.
 - (v) Section 32-1276.01.
 - (w) Section 32-1284.
 - Section 32-1297.01. (X)
- Section 32-1904. 38 (y)
- 39 (z) Section 32-1941.
- 40 (aa) SECTION 32-1982.
- 41 (da) (bb) Section 32-2022.
- (cc) Section 32-2108.01. 42 (bb)
- 43 (cc) (dd) Section 32-2123.
- (ee) Section 32-2371. 44 (dd)
- 45 (ee) (ff) Section 32-3620.

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                       Section 32-3669.
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                       Section 36-425.03.
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                       Section 36-897.01.
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                       Section 36-897.03.
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                       Section 41-619.53.
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           (VV)
                 (ww) Section 41-1964.
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           <del>(ww)</del>
                 (xx) Section 41-1967.01.
19
           (xx)
                 (yy) Section 41-1968.
20
           <del>(yy)</del>
                 (zz) Section 41-1969.
21
           (zz) (aaa) Section 41-2814.
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           (aaa) (bbb) Section 46-141, subsection A or B.
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           (bbb) (ccc) Section 46-321.
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           6. "Vulnerable adult" has the same meaning prescribed in section
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     13-3623.
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           Sec. 20. Section 41-1758.01, Arizona Revised Statutes, is amended
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     to read:
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           41-1758.01. Fingerprinting division: powers and duties
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           A. The fingerprinting division is established in the department of
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     public safety and shall:
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           1. Conduct fingerprint background checks for persons and applicants
     who are seeking licenses from state agencies, employment with licensees,
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     contract providers and state agencies or employment or educational
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     opportunities with agencies that require fingerprint background checks
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     pursuant to sections 3-314, 8-105, 8-322, 8-463, 8-509, 8-802, 15-183,
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     15-503, 15-512, 15-534, 15-763.01, 15-782.02, 15-1330, 15-1881, 17-215,
     28-3228, 28-3413, 32-122.02, 32-122.05, 32-122.06, 32-1232, 32-1276.01,
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     32-1284, 32-1297.01, 32-1904, 32-1941, 32-1982, 32-2022,
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                                                                       32-2108.01,
     32-2123, 32-2371, 32-3620, 32-3668, 32-3669, 36-113, 36-207, 36-411,
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     36-425.03, 36-446.04, 36-594.01, 36-594.02, 36-882, 36-883.02, 36-897.01,
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     36-897.03, 36-3008, 41-619.52, 41-619.53, 41-1964, 41-1967.01, 41-1968,
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     41-1969 and 41-2814, section 46-141, subsection A or B and section 46-321.
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           2. Issue fingerprint clearance cards. On issuance, a fingerprint
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     clearance card becomes the personal property of the cardholder and the
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cardholder shall retain possession of the fingerprint clearance card.

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- 3. On submission of an application for a fingerprint clearance card, collect the fees established by the board of fingerprinting pursuant to section 41-619.53 and deposit, pursuant to sections 35-146 and 35-147, the monies collected in the board of fingerprinting fund.
- 4. Inform in writing each person who submits fingerprints for a fingerprint background check of the right to petition the board of fingerprinting for a good cause exception pursuant to section 41-1758.03, 41-1758.04 or 41-1758.07.
- 5. If after conducting a state and federal criminal history records check the division determines that it is not authorized to issue a fingerprint clearance card to a person, inform the person in writing that the division is not authorized to issue a fingerprint clearance card. The notice shall include the criminal history information on which the denial was based. This criminal history information is subject to dissemination restrictions pursuant to section 41-1750 and Public Law 92-544.
- 6. Notify the person in writing if the division suspends, revokes or places a driving restriction notation on a fingerprint clearance card pursuant to section 41-1758.04. The notice shall include the criminal history information on which the suspension, revocation or placement of the driving restriction notation was based. This criminal history information is subject to dissemination restrictions pursuant to section 41-1750 and Public Law 92-544.
 - 7. Administer and enforce this article.
- B. The fingerprinting division may contract for electronic or internet-based fingerprinting services through an entity or entities for the acquisition and transmission of applicant fingerprint and data submissions to the department, including identity verified fingerprints pursuant to section 15-106. The entity or entities contracted by the department of public safety may charge the applicant a fee for services provided pursuant to this article. The entity or entities contracted by the department of public safety shall comply with:
- 1. All information privacy and security measures and submission standards established by the department of public safety.
- 2. The information technology security policy approved by the department of public safety.

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