AN ACT

To amend sections 121.22, 2925.01, 2925.61, 4723.50, 4723.52, 4729.01, 4729.29, 4729.44, 4729.45, 4729.51, 4729.514, 4729.541, 4729.553, 4729.80, 4730.56, and 4731.83; to amend, for the purpose of adopting new section numbers as indicated in parentheses, sections 4723.486 (4723.488), 4723.488 (4723.484), and 4730.431 (4730.434); and to enact new section 4723.486 and sections 4723.485, 4729.515, 4730.435, and 4730.436 of the Revised Code regarding the administration of addiction treatment drugs, federal agency access to the Ohio Automated Rx Reporting System, the Board of Pharmacy's exemption from open meetings requirements, the occasional sale of certain drugs at wholesale, and naloxone access and education.

Section 1. That sections 121.22, 2925.01, 2925.61, 4723.50, 4723.52, 4729.01, 4729.29, 4729.44, 4729.45, 4729.51, 4729.514, 4729.541, 4729.553, 4729.80, 4730.56, and 4731.83 be amended; sections 4723.486 (4723.488), 4723.488 (4723.484), and 4730.431 (4730.434) be amended for the purpose of adopting new section numbers as indicated in parentheses; and new section 4723.486 and sections 4723.485, 4729.515, 4730.435, and 4730.436 of the Revised Code be enacted to read as follows:

Sec. 121.22. (A) This section shall be liberally construed to require public officials to take official action and to conduct all deliberations upon official business only in open meetings unless the subject matter is specifically excepted by law.

(B) As used in this section:

(1) "Public body" means any of the following:

(a) Any board, commission, committee, council, or similar decision-making body of a state agency, institution, or authority, and any legislative authority or board, commission, committee, council, agency, authority, or similar decision-making body of any county, township, municipal corporation, school district, or other political subdivision or local public institution;

(b) Any committee or subcommittee of a body described in division (B)(1)(a) of this section;

(c) A court of jurisdiction of a sanitary district organized wholly for the purpose of providing a water supply for domestic, municipal, and public use when meeting for the purpose of the appointment, removal, or reappointment of a member of the board of directors of such a district pursuant to section 6115.10 of the Revised Code, if applicable, or for any other matter related to such a district other than litigation involving the district. As used in division (B)(1)(c) of this section, "court of jurisdiction" has the same meaning as "court" in section 6115.01 of the Revised Code.

(2) "Meeting" means any prearranged discussion of the public business of the public body by
a majority of its members.

(3) "Regulated individual" means either of the following:
   (a) A student in a state or local public educational institution;
   (b) A person who is, voluntarily or involuntarily, an inmate, patient, or resident of a state or
       local institution because of criminal behavior, mental illness, an intellectual disability, disease,
       disability, age, or other condition requiring custodial care.

(4) "Public office" has the same meaning as in section 149.011 of the Revised Code.

(C) All meetings of any public body are declared to be public meetings open to the public at
all times. A member of a public body shall be present in person at a meeting open to the public to be
considered present or to vote at the meeting and for purposes of determining whether a quorum is
present at the meeting.

The minutes of a regular or special meeting of any public body shall be promptly prepared,
filed, and maintained and shall be open to public inspection. The minutes need only reflect the
general subject matter of discussions in executive sessions authorized under division (G) or (J) of this
section.

(D) This section does not apply to any of the following:
   (1) A grand jury;
   (2) An audit conference conducted by the auditor of state or independent certified public
       accountants with officials of the public office that is the subject of the audit;
   (3) The adult parole authority when its hearings are conducted at a correctional institution for
       the sole purpose of interviewing inmates to determine parole or pardon and the department of
       rehabilitation and correction when its hearings are conducted at a correctional institution for the sole
       purpose of making determinations under section 2967.271 of the Revised Code regarding the release
       or maintained incarceration of an offender to whom that section applies;
   (4) The organized crime investigations commission established under section 177.01 of the
       Revised Code;
   (5) Meetings of a child fatality review board established under section 307.621 of the
       Revised Code, meetings related to a review conducted pursuant to guidelines established by the
       director of health under section 3701.70 of the Revised Code, and meetings conducted pursuant to
       sections 5153.171 to 5153.173 of the Revised Code;
   (6) The state medical board when determining whether to suspend a license or certificate
       without a prior hearing pursuant to division (G) of either section 4730.25 or 4731.22 of the Revised
       Code;
   (7) The board of nursing when determining whether to suspend a license or certificate
       without a prior hearing pursuant to division (B) of section 4723.281 of the Revised Code;
   (8) The state board of pharmacy when determining whether to suspend or do either of the
       following:
       (a) Suspend a license, certification, or registration without a prior hearing, including during
           meetings conducted by telephone conference, pursuant to division (D) of section 4729.16 Chapters
           3719., 3796., 4729., and 4752. of the Revised Code and rules adopted thereunder; or
       (b) Restrict a person from obtaining further information from the drug database established in
           section 4729.75 of the Revised Code without a prior hearing pursuant to division (C) of section
Am. Sub. H. B. No. 341

133rd G.A.

4729.86 of the Revised Code.

(9) The state chiropractic board when determining whether to suspend a license without a hearing pursuant to section 4734.37 of the Revised Code;

(10) The executive committee of the emergency response commission when determining whether to issue an enforcement order or request that a civil action, civil penalty action, or criminal action be brought to enforce Chapter 3750. of the Revised Code;

(11) The board of directors of the nonprofit corporation formed under section 187.01 of the Revised Code or any committee thereof, and the board of directors of any subsidiary of that corporation or a committee thereof;

(12) An audit conference conducted by the audit staff of the department of job and family services with officials of the public office that is the subject of that audit under section 5101.37 of the Revised Code;

(13) The occupational therapy section of the occupational therapy, physical therapy, and athletic trainers board when determining whether to suspend a license or limited permit without a hearing pursuant to division (D) of section 4755.11 of the Revised Code;

(14) The physical therapy section of the occupational therapy, physical therapy, and athletic trainers board when determining whether to suspend a license without a hearing pursuant to division (E) of section 4755.47 of the Revised Code;

(15) The athletic trainers section of the occupational therapy, physical therapy, and athletic trainers board when determining whether to suspend a license without a hearing pursuant to division (D) of section 4755.64 of the Revised Code;

(16) Meetings of the pregnancy-associated mortality review board established under section 3738.01 of the Revised Code;

(17) Meetings of a fetal-infant mortality review board established under section 3707.71 of the Revised Code.

(E) The controlling board, the tax credit authority, or the minority development financing advisory board, when meeting to consider granting assistance pursuant to Chapter 122. or 166. of the Revised Code, in order to protect the interest of the applicant or the possible investment of public funds, by unanimous vote of all board or authority members present, may close the meeting during consideration of the following information confidentially received by the authority or board from the applicant:

(1) Marketing plans;

(2) Specific business strategy;

(3) Production techniques and trade secrets;

(4) Financial projections;

(5) Personal financial statements of the applicant or members of the applicant's immediate family, including, but not limited to, tax records or other similar information not open to public inspection.

The vote by the authority or board to accept or reject the application, as well as all proceedings of the authority or board not subject to this division, shall be open to the public and governed by this section.

(F) Every public body, by rule, shall establish a reasonable method whereby any person may
determine the time and place of all regularly scheduled meetings and the time, place, and purpose of all special meetings. A public body shall not hold a special meeting unless it gives at least twenty-four hours’ advance notice to the news media that have requested notification, except in the event of an emergency requiring immediate official action. In the event of an emergency, the member or members calling the meeting shall notify the news media that have requested notification immediately of the time, place, and purpose of the meeting.

The rule shall provide that any person, upon request and payment of a reasonable fee, may obtain reasonable advance notification of all meetings at which any specific type of public business is to be discussed. Provisions for advance notification may include, but are not limited to, mailing the agenda of meetings to all subscribers on a mailing list or mailing notices in self-addressed, stamped envelopes provided by the person.

(G) Except as provided in divisions (G)(8) and (J) of this section, the members of a public body may hold an executive session only after a majority of a quorum of the public body determines, by a roll call vote, to hold an executive session and only at a regular or special meeting for the sole purpose of the consideration of any of the following matters:

(1) To consider the appointment, employment, dismissal, discipline, promotion, demotion, or compensation of a public employee or official, or the investigation of charges or complaints against a public employee, official, licensee, or regulated individual, unless the public employee, official, licensee, or regulated individual requests a public hearing. Except as otherwise provided by law, no public body shall hold an executive session for the discipline of an elected official for conduct related to the performance of the elected official's official duties or for the elected official's removal from office. If a public body holds an executive session pursuant to division (G)(1) of this section, the motion and vote to hold that executive session shall state which one or more of the approved purposes listed in division (G)(1) of this section are the purposes for which the executive session is to be held, but need not include the name of any person to be considered at the meeting.

(2) To consider the purchase of property for public purposes, the sale of property at competitive bidding, or the sale or other disposition of unneeded, obsolete, or unfit-for-use property in accordance with section 505.10 of the Revised Code, if premature disclosure of information would give an unfair competitive or bargaining advantage to a person whose personal, private interest is adverse to the general public interest. No member of a public body shall use division (G)(2) of this section as a subterfuge for providing covert information to prospective buyers or sellers. A purchase or sale of public property is void if the seller or buyer of the public property has received covert information from a member of a public body that has not been disclosed to the general public in sufficient time for other prospective buyers and sellers to prepare and submit offers.

If the minutes of the public body show that all meetings and deliberations of the public body have been conducted in compliance with this section, any instrument executed by the public body purporting to convey, lease, or otherwise dispose of any right, title, or interest in any public property shall be conclusively presumed to have been executed in compliance with this section insofar as title or other interest of any bona fide purchasers, lessees, or transferees of the property is concerned.

(3) Conferences with an attorney for the public body concerning disputes involving the public body that are the subject of pending or imminent court action;

(4) Preparing for, conducting, or reviewing negotiations or bargaining sessions with public
employees concerning their compensation or other terms and conditions of their employment;
  (5) Matters required to be kept confidential by federal law or regulations or state statutes;
  (6) Details relative to the security arrangements and emergency response protocols for a
public body or a public office, if disclosure of the matters discussed could reasonably be expected to
jeopardize the security of the public body or public office;
  (7) In the case of a county hospital operated pursuant to Chapter 339. of the Revised Code, a
joint township hospital operated pursuant to Chapter 513. of the Revised Code, or a municipal
hospital operated pursuant to Chapter 749. of the Revised Code, to consider trade secrets, as defined
in section 1333.61 of the Revised Code;
  (8) To consider confidential information related to the marketing plans, specific business
strategy, production techniques, trade secrets, or personal financial statements of an applicant for
economic development assistance, or to negotiations with other political subdivisions respecting
requests for economic development assistance, provided that both of the following conditions apply:
  (a) The information is directly related to a request for economic development assistance that
is to be provided or administered under any provision of Chapter 715., 725., 1724., or 1728. or
sections 701.07, 3735.67 to 3735.70, 5709.40 to 5709.43, 5709.61 to 5709.69, 5709.73 to 5709.75, or
5709.77 to 5709.81 of the Revised Code, or that involves public infrastructure improvements or the
extension of utility services that are directly related to an economic development project.
  (b) A unanimous quorum of the public body determines, by a roll call vote, that the executive
session is necessary to protect the interests of the applicant or the possible investment or expenditure
of public funds to be made in connection with the economic development project.

If a public body holds an executive session to consider any of the matters listed in divisions
(G)(2) to (8) of this section, the motion and vote to hold that executive session shall state which one
or more of the approved matters listed in those divisions are to be considered at the executive
session.

A public body specified in division (B)(1)(c) of this section shall not hold an executive
session when meeting for the purposes specified in that division.

(H) A resolution, rule, or formal action of any kind is invalid unless adopted in an open
meeting of the public body. A resolution, rule, or formal action adopted in an open meeting that
results from deliberations in a meeting not open to the public is invalid unless the deliberations were
for a purpose specifically authorized in division (G) or (J) of this section and conducted at an
executive session held in compliance with this section. A resolution, rule, or formal action adopted in
an open meeting is invalid if the public body that adopted the resolution, rule, or formal action
violated division (F) of this section.

(I)(1) Any person may bring an action to enforce this section. An action under division (I)(1)
of this section shall be brought within two years after the date of the alleged violation or threatened
violation. Upon proof of a violation or threatened violation of this section in an action brought by any
person, the court of common pleas shall issue an injunction to compel the members of the public
body to comply with its provisions.

(2)(a) If the court of common pleas issues an injunction pursuant to division (I)(1) of this
section, the court shall order the public body that it enjoins to pay a civil forfeiture of five hundred
dollars to the party that sought the injunction and shall award to that party all court costs and, subject
to reduction as described in division (I)(2) of this section, reasonable attorney's fees. The court, in its discretion, may reduce an award of attorney's fees to the party that sought the injunction or not award attorney's fees to that party if the court determines both of the following:

(i) That, based on the ordinary application of statutory law and case law as it existed at the time of violation or threatened violation that was the basis of the injunction, a well-informed public body reasonably would believe that the public body was not violating or threatening to violate this section;

(ii) That a well-informed public body reasonably would believe that the conduct or threatened conduct that was the basis of the injunction would serve the public policy that underlies the authority that is asserted as permitting that conduct or threatened conduct.

(b) If the court of common pleas does not issue an injunction pursuant to division (I)(1) of this section and the court determines at that time that the bringing of the action was frivolous conduct, as defined in division (A) of section 2323.51 of the Revised Code, the court shall award to the public body all court costs and reasonable attorney's fees, as determined by the court.

(3) Irreparable harm and prejudice to the party that sought the injunction shall be conclusively and irrebuttably presumed upon proof of a violation or threatened violation of this section.

(4) A member of a public body who knowingly violates an injunction issued pursuant to division (I)(1) of this section may be removed from office by an action brought in the court of common pleas for that purpose by the prosecuting attorney or the attorney general.

(J)(1) Pursuant to division (C) of section 5901.09 of the Revised Code, a veterans service commission shall hold an executive session for one or more of the following purposes unless an applicant requests a public hearing:

(a) Interviewing an applicant for financial assistance under sections 5901.01 to 5901.15 of the Revised Code;

(b) Discussing applications, statements, and other documents described in division (B) of section 5901.09 of the Revised Code;

(c) Reviewing matters relating to an applicant's request for financial assistance under sections 5901.01 to 5901.15 of the Revised Code.

(2) A veterans service commission shall not exclude an applicant for, recipient of, or former recipient of financial assistance under sections 5901.01 to 5901.15 of the Revised Code, and shall not exclude representatives selected by the applicant, recipient, or former recipient, from a meeting that the commission conducts as an executive session that pertains to the applicant's, recipient's, or former recipient's application for financial assistance.

(3) A veterans service commission shall vote on the grant or denial of financial assistance under sections 5901.01 to 5901.15 of the Revised Code only in an open meeting of the commission. The minutes of the meeting shall indicate the name, address, and occupation of the applicant, whether the assistance was granted or denied, the amount of the assistance if assistance is granted, and the votes for and against the granting of assistance.

Sec. 2925.01. As used in this chapter:

"pharmacy," "sale," "schedule I," "schedule II," "schedule III," "schedule IV," "schedule V," and "wholesaler" have the same meanings as in section 3719.01 of the Revised Code.

(B) "Drug dependent person" and "drug of abuse" have the same meanings as in section 3719.011 of the Revised Code.

(C) "Drug," "dangerous drug," "licensed health professional authorized to prescribe drugs," and "prescription" have the same meanings as in section 4729.01 of the Revised Code.

(D) "Bulk amount" of a controlled substance means any of the following:

(1) For any compound, mixture, preparation, or substance included in schedule I, schedule II, or schedule III, with the exception of any controlled substance analog, marihuana, cocaine, L.S.D., heroin, any fentanyl-related compound, and hashish and except as provided in division (D)(2), (5), or (6) of this section, whichever of the following is applicable:

(a) An amount equal to or exceeding ten grams or twenty-five unit doses of a compound, mixture, preparation, or substance that is or contains any amount of a schedule I opiate or opium derivative;

(b) An amount equal to or exceeding ten grams of a compound, mixture, preparation, or substance that is or contains any amount of raw or gum opium;

(c) An amount equal to or exceeding thirty grams or ten unit doses of a compound, mixture, preparation, or substance that is or contains any amount of a schedule I hallucinogen other than tetrahydrocannabinol or lysergic acid amide, or a schedule I stimulant or depressant;

(d) An amount equal to or exceeding twenty grams or five times the maximum daily dose in the usual dose range specified in a standard pharmaceutical reference manual of a compound, mixture, preparation, or substance that is or contains any amount of a schedule II opiate or opium derivative;

(e) An amount equal to or exceeding five grams or ten unit doses of a compound, mixture, preparation, or substance that is or contains any amount of phencyclidine;

(f) An amount equal to or exceeding one hundred twenty grams or thirty times the maximum daily dose in the usual dose range specified in a standard pharmaceutical reference manual of a compound, mixture, preparation, or substance that is or contains any amount of a schedule II stimulant that is in a final dosage form manufactured by a person authorized by the "Federal Food, Drug, and Cosmetic Act," 52 Stat. 1040 (1938), 21 U.S.C.A. 301, as amended, and the federal drug abuse control laws, as defined in section 3719.01 of the Revised Code, that is or contains any amount of a schedule II depressant substance or a schedule II hallucinogenic substance;

(g) An amount equal to or exceeding three grams of a compound, mixture, preparation, or substance that is or contains any amount of a schedule II stimulant, or any of its salts or isomers, that is not in a final dosage form manufactured by a person authorized by the Federal Food, Drug, and Cosmetic Act and the federal drug abuse control laws.

(2) An amount equal to or exceeding one hundred twenty grams or thirty times the maximum daily dose in the usual dose range specified in a standard pharmaceutical reference manual of a compound, mixture, preparation, or substance that is or contains any amount of a schedule III or IV substance other than an anabolic steroid or a schedule III opiate or opium derivative;

(3) An amount equal to or exceeding twenty grams or five times the maximum daily dose in the usual dose range specified in a standard pharmaceutical reference manual of a compound,
mixture, preparation, or substance that is or contains any amount of a schedule III opiate or opium derivative;

(4) An amount equal to or exceeding two hundred fifty milliliters or two hundred fifty grams of a compound, mixture, preparation, or substance that is or contains any amount of a schedule V substance;

(5) An amount equal to or exceeding two hundred solid dosage units, sixteen grams, or sixteen milliliters of a compound, mixture, preparation, or substance that is or contains any amount of a schedule III anabolic steroid;

(6) For any compound, mixture, preparation, or substance that is a combination of a fentanyl-related compound and any other compound, mixture, preparation, or substance included in schedule III, schedule IV, or schedule V, if the defendant is charged with a violation of section 2925.11 of the Revised Code and the sentencing provisions set forth in divisions (C)(10)(b) and (C)(11) of that section will not apply regarding the defendant and the violation, the bulk amount of the controlled substance for purposes of the violation is the amount specified in division (D)(1), (2), (3), (4), or (5) of this section for the other schedule III, IV, or V controlled substance that is combined with the fentanyl-related compound.

(E) "Unit dose" means an amount or unit of a compound, mixture, or preparation containing a controlled substance that is separately identifiable and in a form that indicates that it is the amount or unit by which the controlled substance is separately administered to or taken by an individual.

(F) "Cultivate" includes planting, watering, fertilizing, or tilling.

(G) "Drug abuse offense" means any of the following:

(1) A violation of division (A) of section 2913.02 that constitutes theft of drugs, or a violation of section 2925.02, 2925.03, 2925.04, 2925.041, 2925.05, 2925.06, 2925.11, 2925.12, 2925.13, 2925.22, 2925.23, 2925.24, 2925.31, 2925.32, 2925.36, or 2925.37 of the Revised Code;

(2) A violation of an existing or former law of this or any other state or of the United States that is substantially equivalent to any section listed in division (G)(1) of this section;

(3) An offense under an existing or former law of this or any other state, or of the United States, of which planting, cultivating, harvesting, processing, making, manufacturing, producing, shipping, transporting, delivering, acquiring, possessing, storing, distributing, dispensing, selling, inducing another to use, administering to another, using, or otherwise dealing with a controlled substance is an element;

(4) A conspiracy to commit, attempt to commit, or complicity in committing or attempting to commit any offense under division (G)(1), (2), or (3) of this section.

(H) "Felony drug abuse offense" means any drug abuse offense that would constitute a felony under the laws of this state, any other state, or the United States.

(I) "Harmful intoxicant" does not include beer or intoxicating liquor but means any of the following:

(1) Any compound, mixture, preparation, or substance the gas, fumes, or vapor of which when inhaled can induce intoxication, excitement, giddiness, irrational behavior, depression, stupor, paralysis, unconsciousness, asphyxiation, or other harmful physiological effects, and includes, but is not limited to, any of the following:

(a) Any volatile organic solvent, plastic cement, model cement, fingernail polish remover,
lacquer thinner, cleaning fluid, gasoline, or other preparation containing a volatile organic solvent;
(b) Any aerosol propellant;
(c) Any fluorocarbon refrigerant;
(d) Any anesthetic gas.
(2) Gamma Butyrolactone;
(3) 1,4 Butanediol.
(J) "Manufacture" means to plant, cultivate, harvest, process, make, prepare, or otherwise engage in any part of the production of a drug, by propagation, extraction, chemical synthesis, or compounding, or any combination of the same, and includes packaging, repackaging, labeling, and other activities incident to production.
(K) "Possess" or "possession" means having control over a thing or substance, but may not be inferred solely from mere access to the thing or substance through ownership or occupation of the premises upon which the thing or substance is found.
(L) "Sample drug" means a drug or pharmaceutical preparation that would be hazardous to health or safety if used without the supervision of a licensed health professional authorized to prescribe drugs, or a drug of abuse, and that, at one time, had been placed in a container plainly marked as a sample by a manufacturer.
(M) "Standard pharmaceutical reference manual" means the current edition, with cumulative changes if any, of references that are approved by the state board of pharmacy.
(N) "Juvenile" means a person under eighteen years of age.
(O) "Counterfeit controlled substance" means any of the following:
(1) Any drug that bears, or whose container or label bears, a trademark, trade name, or other identifying mark used without authorization of the owner of rights to that trademark, trade name, or identifying mark;
(2) Any unmarked or unlabeled substance that is represented to be a controlled substance manufactured, processed, packed, or distributed by a person other than the person that manufactured, processed, packed, or distributed it;
(3) Any substance that is represented to be a controlled substance but is not a controlled substance or is a different controlled substance;
(4) Any substance other than a controlled substance that a reasonable person would believe to be a controlled substance because of its similarity in shape, size, and color, or its markings, labeling, packaging, distribution, or the price for which it is sold or offered for sale.
(P) An offense is "committed in the vicinity of a school" if the offender commits the offense on school premises, in a school building, or within one thousand feet of the boundaries of any school premises, regardless of whether the offender knows the offense is being committed on school premises, in a school building, or within one thousand feet of the boundaries of any school premises.
(Q) "School" means any school operated by a board of education, any community school established under Chapter 3314. of the Revised Code, or any nonpublic school for which the state board of education prescribes minimum standards under section 3301.07 of the Revised Code, whether or not any instruction, extracurricular activities, or training provided by the school is being conducted at the time a criminal offense is committed.
(R) "School premises" means either of the following:
(1) The parcel of real property on which any school is situated, whether or not any instruction, extracurricular activities, or training provided by the school is being conducted on the premises at the time a criminal offense is committed;

(2) Any other parcel of real property that is owned or leased by a board of education of a school, the governing authority of a community school established under Chapter 3314. of the Revised Code, or the governing body of a nonpublic school for which the state board of education prescribes minimum standards under section 3301.07 of the Revised Code and on which some of the instruction, extracurricular activities, or training of the school is conducted, whether or not any instruction, extracurricular activities, or training provided by the school is being conducted on the parcel of real property at the time a criminal offense is committed.

(S) "School building" means any building in which any of the instruction, extracurricular activities, or training provided by a school is conducted, whether or not any instruction, extracurricular activities, or training provided by the school is being conducted in the school building at the time a criminal offense is committed.

(T) "Disciplinary counsel" means the disciplinary counsel appointed by the board of commissioners on grievances and discipline of the supreme court under the Rules for the Government of the Bar of Ohio.

(U) "Certified grievance committee" means a duly constituted and organized committee of the Ohio state bar association or of one or more local bar associations of the state of Ohio that complies with the criteria set forth in Rule V, section 6 of the Rules for the Government of the Bar of Ohio.

(V) "Professional license" means any license, permit, certificate, registration, qualification, admission, temporary license, temporary permit, temporary certificate, or temporary registration that is described in divisions (W)(1) to (37) of this section and that qualifies a person as a professionally licensed person.

(W) "Professionally licensed person" means any of the following:

(1) A person who has received a certificate or temporary certificate as a certified public accountant or who has registered as a public accountant under Chapter 4701. of the Revised Code and who holds an Ohio permit issued under that chapter;

(2) A person who holds a certificate of qualification to practice architecture issued or renewed and registered under Chapter 4703. of the Revised Code;

(3) A person who is registered as a landscape architect under Chapter 4703. of the Revised Code or who holds a permit as a landscape architect issued under that chapter;

(4) A person licensed under Chapter 4707. of the Revised Code;

(5) A person who has been issued a certificate of registration as a registered barber under Chapter 4709. of the Revised Code;

(6) A person licensed and regulated to engage in the business of a debt pooling company by a legislative authority, under authority of Chapter 4710. of the Revised Code;

(7) A person who has been issued a cosmetologist's license, hair designer's license, manicurist's license, esthetician's license, natural hair stylist's license, advanced cosmetologist's license, advanced hair designer's license, advanced manicurist's license, advanced esthetician's license, advanced natural hair stylist's license, cosmetology instructor's license, hair design
instructor's license, manicurist instructor's license, esthetics instructor's license, natural hair style instructor's license, independent contractor's license, or tanning facility permit under Chapter 4713. of the Revised Code;

(8) A person who has been issued a license to practice dentistry, a general anesthesia permit, a conscious sedation permit, a limited resident's license, a limited teaching license, a dental hygienist's license, or a dental hygienist's teacher's certificate under Chapter 4715. of the Revised Code;

(9) A person who has been issued an embalmer's license, a funeral director's license, a funeral home license, or a crematory license, or who has been registered for an embalmer's or funeral director's apprenticeship under Chapter 4717. of the Revised Code;

(10) A person who has been licensed as a registered nurse or practical nurse, or who has been issued a certificate for the practice of nurse-midwifery under Chapter 4723. of the Revised Code;

(11) A person who has been licensed to practice optometry or to engage in optical dispensing under Chapter 4725. of the Revised Code;

(12) A person licensed to act as a pawnbroker under Chapter 4727. of the Revised Code;

(13) A person licensed to act as a precious metals dealer under Chapter 4728. of the Revised Code;

(14) A person licensed under Chapter 4729. of the Revised Code as a pharmacist or pharmacy intern or registered under that chapter as a registered pharmacy technician, certified pharmacy technician, or pharmacy technician trainee;

(15) A person licensed under Chapter 4729. of the Revised Code as a manufacturer of dangerous drugs, outsourcing facility, third-party logistics provider, repackager of dangerous drugs, wholesale distributor of dangerous drugs, or terminal distributor of dangerous drugs;

(16) A person who is authorized to practice as a physician assistant under Chapter 4730. of the Revised Code;

(17) A person who has been issued a license to practice medicine and surgery, osteopathic medicine and surgery, or podiatric medicine and surgery under Chapter 4731. of the Revised Code or has been issued a certificate to practice a limited branch of medicine under that chapter;

(18) A person licensed as a psychologist or school psychologist under Chapter 4732. of the Revised Code;

(19) A person registered to practice the profession of engineering or surveying under Chapter 4733. of the Revised Code;

(20) A person who has been issued a license to practice chiropractic under Chapter 4734. of the Revised Code;

(21) A person licensed to act as a real estate broker or real estate salesperson under Chapter 4735. of the Revised Code;

(22) A person registered as a registered sanitarian under Chapter 4736. of the Revised Code;

(23) A person licensed to operate or maintain a junkyard under Chapter 4737. of the Revised Code;

(24) A person who has been issued a motor vehicle salvage dealer's license under Chapter 4738. of the Revised Code;

(25) A person who has been licensed to act as a steam engineer under Chapter 4739. of the Revised Code;
(26) A person who has been issued a license or temporary permit to practice veterinary medicine or any of its branches, or who is registered as a graduate animal technician under Chapter 4741. of the Revised Code;
(27) A person who has been issued a hearing aid dealer's or fitter's license or trainee permit under Chapter 4747. of the Revised Code;
(28) A person who has been issued a class A, class B, or class C license or who has been registered as an investigator or security guard employee under Chapter 4749. of the Revised Code;
(29) A person licensed to practice as a nursing home administrator under Chapter 4751. of the Revised Code;
(30) A person licensed to practice as a speech-language pathologist or audiologist under Chapter 4753. of the Revised Code;
(31) A person issued a license as an occupational therapist or physical therapist under Chapter 4755. of the Revised Code;
(32) A person who is licensed as a licensed professional clinical counselor, licensed professional counselor, social worker, independent social worker, independent marriage and family therapist, or marriage and family therapist, or registered as a social work assistant under Chapter 4757. of the Revised Code;
(33) A person issued a license to practice dietetics under Chapter 4759. of the Revised Code;
(34) A person who has been issued a license or limited permit to practice respiratory therapy under Chapter 4761. of the Revised Code;
(35) A person who has been issued a real estate appraiser certificate under Chapter 4763. of the Revised Code;
(36) A person who has been issued a home inspector license under Chapter 4764. of the Revised Code;
(37) A person who has been admitted to the bar by order of the supreme court in compliance with its prescribed and published rules.
(X) "Cocaine" means any of the following:
(1) A cocaine salt, isomer, or derivative, a salt of a cocaine isomer or derivative, or the base form of cocaine;
(2) Coca leaves or a salt, compound, derivative, or preparation of coca leaves, including eggonine, a salt, isomer, or derivative of eggonine, or a salt of an isomer or derivative of eggonine;
(3) A salt, compound, derivative, or preparation of a substance identified in division (X)(1) or (2) of this section that is chemically equivalent to or identical with any of those substances, except that the substances shall not include decocainized coca leaves or extraction of coca leaves if the extractions do not contain cocaine or eggonine.
(Y) "L.S.D." means lysergic acid diethylamide.
(Z) "Hashish" means the a resin or a preparation of the a resin to which both of the following apply:
(1) It is contained in *marihuana* or derived from any part of the plant of the genus cannabis, whether in solid form or in a liquid concentrate, liquid extract, or liquid distillate form;
(2) It has a delta-9 tetrahydrocannabinol concentration of more than three-tenths per cent. "Hashish" does not include a hemp byproduct in the possession of a licensed hemp processor.
under Chapter 928. of the Revised Code, provided that the hemp byproduct is being produced, stored, and disposed of in accordance with rules adopted under section 928.03 of the Revised Code.

(AA) "Marihuana" has the same meaning as in section 3719.01 of the Revised Code, except that it does not include hashish.

(BB) An offense is "committed in the vicinity of a juvenile" if the offender commits the offense within one hundred feet of a juvenile or within the view of a juvenile, regardless of whether the offender knows the age of the juvenile, whether the offender knows the offense is being committed within one hundred feet of or within view of the juvenile, or whether the juvenile actually views the commission of the offense.

(CC) "Presumption for a prison term" or "presumption that a prison term shall be imposed" means a presumption, as described in division (D) of section 2929.13 of the Revised Code, that a prison term is a necessary sanction for a felony in order to comply with the purposes and principles of sentencing under section 2929.11 of the Revised Code.

(DD) "Major drug offender" has the same meaning as in section 2929.01 of the Revised Code.

(EE) "Minor drug possession offense" means either of the following:
   (1) A violation of section 2925.11 of the Revised Code as it existed prior to July 1, 1996;
   (2) A violation of section 2925.11 of the Revised Code as it exists on and after July 1, 1996, that is a misdemeanor or a felony of the fifth degree.

(FF) "Mandatory prison term" has the same meaning as in section 2929.01 of the Revised Code.

(GG) "Adulterate" means to cause a drug to be adulterated as described in section 3715.63 of the Revised Code.

(HH) "Public premises" means any hotel, restaurant, tavern, store, arena, hall, or other place of public accommodation, business, amusement, or resort.

(II) "Methamphetamine" means methamphetamine, any salt, isomer, or salt of an isomer of methamphetamine, or any compound, mixture, preparation, or substance containing methamphetamine or any salt, isomer, or salt of an isomer of methamphetamine.

(JJ) "Deception" has the same meaning as in section 2913.01 of the Revised Code.

(KK) "Fentanyl-related compound" means any of the following:
   (1) Fentanyl;
   (2) Alpha-methylfentanyl (N-[1-(alpha-methyl-beta-phenyl)ethyl-4- piperidyl]propionanilide;
       1-(1-methyl-2-phenylethyl)-4-(N-propanilido) piperidine);
   (3) Alpha-methylthiofentanyl (N-[1-methyl-2-(2-thienyl)ethyl-4- piperidinyl]-N-phenylpropanamide);
   (4) Beta-hydroxyfentanyl (N-[1-(2-hydroxy-2-phenethyl-4-piperidinyl]-N-phenylpropanamide);
   (5) Beta-hydroxy-3-methylfentanyl (other name: N-[1-(2-hydroxy-2- phenethyl)-3-methyl-4- piperidinyl]-N-phenylpropanamide);
   (6) 3-methylfentanyl (N-[3-methyl-1-(2-phenylethyl)-4-piperidyl]-N-phenylpropanamide);
   (7) 3-methylthiofentanyl (N-[3-methyl-1-[2-(thienyl)ethyl]-4- piperidinyl]-N-phenylpropanamide);
(8) Para-fluorofentanyl (N-(4-fluorophenyl)-N-[1-(2-phenethyl)-4-piperidinyl]propanamide; 
(9) Thiofentanyl (N-phenyl-N-[1-(2-thienyl)ethyl-4-piperidinyl]-propanamide; 
(10) Alfentanil; 
(11) Carfentanil; 
(12) Remifentanil; 
(13) Sufentanil; 
(14) Acetyl-alpha-methylfentanyl (N-[1-(1-methyl-2-phenethyl)-4-piperidinyl]-N-phenylacetamide); and 
(15) Any compound that meets all of the following fentanyl pharmacophore requirements to bind at the mu receptor, as identified by a report from an established forensic laboratory, including acetylfentanyl, furanylfentanyl, valerylfentanyl, butyrylfentanyl, isobutyrylfentanyl, 4-methoxybutyrylfentanyl, para-fluorobutyrylfentanyl, acrylfentanyl, and ortho-fluorofentanyl: 
    (a) A chemical scaffold consisting of both of the following: 
        (i) A five, six, or seven member ring structure containing a nitrogen, whether or not further substituted; 
        (ii) An attached nitrogen to the ring, whether or not that nitrogen is enclosed in a ring structure, including an attached aromatic ring or other lipophilic group to that nitrogen. 
    (b) A polar functional group attached to the chemical scaffold, including but not limited to a hydroxyl, ketone, amide, or ester; 
    (c) An alkyl or aryl substitution off the ring nitrogen of the chemical scaffold; and 
    (d) The compound has not been approved for medical use by the United States food and drug administration. 

(LL) "First degree felony mandatory prison term" means one of the definite prison terms prescribed in division (A)(1)(b) of section 2929.14 of the Revised Code for a felony of the first degree, except that if the violation for which sentence is being imposed is committed on or after the effective date of this amendment March 22, 2019, it means one of the minimum prison terms prescribed in division (A)(1)(a) of that section for a felony of the first degree. 

(MM) "Second degree felony mandatory prison term" means one of the definite prison terms prescribed in division (A)(2)(b) of section 2929.14 of the Revised Code for a felony of the second degree, except that if the violation for which sentence is being imposed is committed on or after the effective date of this amendment March 22, 2019, it means one of the minimum prison terms prescribed in division (A)(2)(a) of that section for a felony of the second degree. 

(NN) "Maximum first degree felony mandatory prison term" means the maximum definite prison term prescribed in division (A)(1)(b) of section 2929.14 of the Revised Code for a felony of the first degree, except that if the violation for which sentence is being imposed is committed on or after the effective date of this amendment March 22, 2019, it means the longest minimum prison term prescribed in division (A)(1)(a) of that section for a felony of the first degree. 

(OO) "Maximum second degree felony mandatory prison term" means the maximum definite prison term prescribed in division (A)(2)(b) of section 2929.14 of the Revised Code for a felony of the second degree, except that if the violation for which sentence is being imposed is committed on or after the effective date of this amendment March 22, 2019, it means the longest minimum prison term prescribed in division (A)(2)(a) of that section for a felony of the second degree.
(PP) "Delta-9 tetrahydrocannabinol" has the same meaning as in section 928.01 of the Revised Code.

Sec. 2925.61. (A) As used in this section:
(1) "Law enforcement agency" means a government entity that employs peace officers to perform law enforcement duties.
(2) "Licensed health professional" means all of the following:
   (a) A physician;
   (b) A physician assistant who is licensed under Chapter 4730. of the Revised Code, holds a valid prescriber number issued by the state medical board, and has been granted physician-delegated prescriptive authority;
   (c) An advanced practice registered nurse who holds a current, valid license issued under Chapter 4723. of the Revised Code and is designated as a clinical nurse specialist, certified nurse-midwife, or certified nurse practitioner.
(3) "Peace officer" has the same meaning as in section 2921.51 of the Revised Code.
(4) "Physician" means an individual who is authorized under Chapter 4731. of the Revised Code to practice medicine and surgery, osteopathic medicine and surgery, or podiatric medicine and surgery.

(B) A family member, friend, or other individual who is in a position to assist an individual who is apparently experiencing or at risk of experiencing an opioid-related overdose, is not subject to criminal prosecution for a violation of section 4731.41 of the Revised Code if the individual, acting in good faith, does all of the following:
   (1) Obtains naloxone pursuant to a prescription issued by a licensed health professional, or obtains naloxone from one of the following:
      (a) A licensed health professional;
      (b) An individual who is authorized to personally furnish naloxone by either a
         (i) A physician under section 4731.941 of the Revised Code;
         (ii) An advanced practice registered nurse under section 4723.485 of the Revised Code;
         (iii) A physician assistant under section 4730.435 of the Revised Code;
         (iv) A board of health under section 3707.561 of the Revised Code
            to personally furnish naloxone;
      (c) A pharmacist or pharmacy intern who is authorized by a physician or board of health
         under section 4729.44 of the Revised Code to dispense naloxone without a prescription.
   (2) Administers the naloxone obtained as described in division (B)(1) of this section to an individual who is apparently experiencing an opioid-related overdose;
   (3) Attempts to summon emergency services as soon as practicable either before or after administering the naloxone.

(C) An individual who is an employee, volunteer, or contractor of a service entity, as defined in section 4729.514 of the Revised Code, and has been authorized under section 3707.562, 4723.486,
4730.436, or 4731.943 of the Revised Code to administer naloxone is not subject to criminal prosecution for a violation of section 4731.41 of the Revised Code or criminal prosecution under this chapter, if the individual, acting in good faith, does all of the following:

(1) Obtains naloxone from the service entity of which the individual is an employee, volunteer, or contractor;

(2) Administers the naloxone obtained to an individual who is apparently experiencing an opioid-related overdose;

(3) Attempts to summon emergency services as soon as practicable either before or after administering the naloxone.

(D) Divisions (B) and (C) of this section do not apply to a peace officer or to an emergency medical technician-basic, emergency medical technician-intermediate, or emergency medical technician-paramedic, as defined in section 4765.01 of the Revised Code.

(E)(1) If a peace officer, acting in good faith, administers naloxone to an individual who is apparently experiencing an opioid-related overdose, both of the following apply:

(a) The peace officer is not subject to administrative action, criminal prosecution for a violation of section 4731.41 of the Revised Code, or criminal prosecution under this chapter.

(b) The peace officer is not liable for damages in a civil action for injury, death, or loss to person or property for an act or omission that allegedly arises from obtaining, maintaining, accessing, or administering the naloxone.

(2) Division (E)(1)(b) of this section does not eliminate, limit, or reduce any other immunity or defense that an entity or person may be entitled to under section 9.86 or Chapter 2744. of the Revised Code, any other provision of the Revised Code, or the common law of this state.

Sec. 4723.484. (A) Notwithstanding any provision of this chapter or rule adopted by the board of nursing, an advanced practice registered nurse who is designated as a clinical nurse specialist, certified nurse-midwife, or certified nurse practitioner may personally furnish a supply of naloxone, or issue a prescription for naloxone, without having examined the individual to whom it may be administered if both of the following conditions are met:

(1) The naloxone supply is furnished to, or the prescription is issued to and in the name of, a family member, friend, or other individual in a position to assist an individual who there is reason to believe is at risk of experiencing an opioid-related overdose.

(2) The advanced practice registered nurse instructs the individual receiving the naloxone supply or prescription to summon emergency services as soon as practicable either before or after administering naloxone to an individual apparently experiencing an opioid-related overdose.

(B) An advanced practice registered nurse who under division (A) of this section in good faith furnishes a supply of naloxone or issues a prescription for naloxone is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is furnished or the prescription is issued: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

Sec. 4723.485. (A)(1) An advanced practice registered nurse who is designated as a clinical nurse specialist, certified nurse-midwife, or certified nurse practitioner and who has established a protocol that meets the requirements of division (C) of this section may authorize one or more other individuals to personally furnish a supply of naloxone pursuant to the protocol to either of the
following:

(a) An individual who there is reason to believe is experiencing or at risk of experiencing an opioid-related overdose;

(b) A family member, friend, or other person in a position to assist an individual who there is reason to believe is at risk of experiencing an opioid-related overdose.

(2) An individual authorized under this section to personally furnish naloxone may do so without having examined the individual to whom it may be administered.

(B) An individual authorized by an advanced practice registered nurse under this section may personally furnish naloxone to an individual described in division (A)(1)(a) or (b) of this section if both of the following conditions are met:

(1) The authorized individual complies with the protocol established by the authorizing advanced practice registered nurse, including having completed the training required by the protocol.

(2) The authorized individual instructs the individual to whom naloxone is furnished to summon emergency services as soon as practicable either before or after administering naloxone.

(C) A protocol established by an advanced practice registered nurse for purposes of this section shall be established in writing and include all of the following:

(1) A description of the clinical pharmacology of naloxone;

(2) Precautions and contraindications concerning furnishing naloxone;

(3) Any limitations the advanced practice registered nurse specifies concerning the individuals to whom naloxone may be furnished;

(4) The naloxone dosage that may be furnished and any variation in the dosage based on circumstances specified in the protocol;

(5) Labeling, storage, record keeping, and administrative requirements;

(6) Training requirements that must be met before an individual will be authorized to furnish naloxone;

(7) Any instructions or training that the authorized individual must provide to an individual to whom naloxone is furnished.

(D) An advanced practice registered nurse who in good faith authorizes another individual to personally furnish naloxone in accordance with a protocol established by the advanced practice registered nurse under this section is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is furnished: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

An individual authorized under this section to personally furnish naloxone who does so in good faith is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is furnished: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

Sec. 4723.486. (A) As used in this section, "service entity" has the same meaning as in section 4729.514 of the Revised Code.

(B) An advanced practice registered nurse who is designated as a clinical nurse specialist, certified nurse-midwife, or certified nurse practitioner and who has established a protocol under division (D) of this section may authorize an individual who is an employee, volunteer, or contractor of a service entity to administer naloxone to an individual who is apparently experiencing an opioid-related overdose.
related overdose.

(C) An individual authorized by an advanced practice registered nurse under this section may administer naloxone to an individual who is apparently experiencing an opioid-related overdose if all of the following conditions are met:

1. The naloxone is obtained from a service entity of which the authorized individual is an employee, volunteer, or contractor.
2. The authorized individual complies with the protocol established by the authorizing advanced practice registered nurse.
3. The authorized individual summons emergency services as soon as practicable either before or after administering the naloxone.

(D) A protocol established by an advanced practice registered nurse for purposes of this section must be in writing and include all of the following:

1. A description of the clinical pharmacology of naloxone;
2. Precautions and contraindications concerning the administration of naloxone;
3. Any limitations the advanced practice registered nurse specifies concerning the individuals to whom naloxone may be administered;
4. The naloxone dosage that may be administered and any variation in the dosage based on circumstances specified in the protocol;
5. Labeling, storage, record keeping, and administrative requirements;
6. Training requirements that must be met before an individual can be authorized to administer naloxone.

(E) An advanced practice registered nurse who in good faith authorizes an individual to administer naloxone under this section is not liable for or subject to any of the following for any act or omission of the authorized individual: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

A service entity or an employee, volunteer, or contractor of a service entity is not liable for or subject to any of the following for injury, death, or loss to person or property that allegedly arises from an act or omission associated with procuring, maintaining, accessing, or administering naloxone under this section, unless the act or omission constitutes willful or wanton misconduct: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

This section does not eliminate, limit, or reduce any other immunity or defense that a service entity or an employee, volunteer, or contractor of a service entity may be entitled to under Chapter 2305. or any other provision of the Revised Code or under the common law of this state.

Sec. 4723.486 4723.488. (A) Except as provided in division (B) of this section, in the case of a license holder who is seeking renewal of a license to practice nursing as an advanced practice registered nurse and who prescribes opioid analgesics or benzodiazepines, as defined in section 3719.01 of the Revised Code, the holder shall certify to the board whether the holder has been granted access to the drug database established and maintained by the state board of pharmacy pursuant to section 4729.75 of the Revised Code.

(B) The requirement in division (A) of this section does not apply if any of the following is the case:

1. The state board of pharmacy notifies the board of nursing pursuant to section 4729.861 of
the Revised Code that the license holder has been restricted from obtaining further information from the drug database.

(2) The state board of pharmacy no longer maintains the drug database.

(3) The license holder does not practice nursing in this state.

(C) If a license holder certifies to the board of nursing that the holder has been granted access to the drug database and the board finds through an audit or other means that the holder has not been granted access, the board may take action under section 4723.28 of the Revised Code.

Sec. 4723.50. (A) As used in this section:

(1) "Controlled substance" has the same meaning as in section 3719.01 of the Revised Code.

(2) "Medication-assisted treatment" has the same meaning as in section 340.01 of the Revised Code.

(B) In accordance with Chapter 119. of the Revised Code, the board of nursing shall adopt rules as necessary to implement the provisions of this chapter pertaining to the authority of advanced practice registered nurses who are designated as clinical nurse specialists, certified nurse-midwives, and certified nurse practitioners to prescribe and furnish drugs and therapeutic devices.

The board shall adopt rules that are consistent with a recommended exclusionary formulary the board receives from the committee on prescriptive governance pursuant to section 4723.492 of the Revised Code. After reviewing a formulary submitted by the committee, the board may either adopt the formulary as a rule or ask the committee to reconsider and resubmit the formulary. The board shall not adopt any rule that does not conform to a formulary developed by the committee.

The exclusionary formulary shall permit, in a manner consistent with section 4723.481 of the Revised Code, the prescribing of controlled substances, including drugs that contain buprenorphine used in medication-assisted treatment and both oral and long-acting opioid antagonists. The formulary shall not permit the prescribing or furnishing of any of the following:

(1) A drug or device to perform or induce an abortion;

(2) A drug or device prohibited by federal or state law.

(C) In addition to the rules described in division (B) of this section, the board shall adopt rules under this section that do the following:

(1) Establish standards for board approval of the course of study in advanced pharmacology and related topics required by section 4723.482 of the Revised Code;

(2) Establish requirements for board approval of the two-hour course of instruction in the laws of this state as required under division (C)(1) of section 4723.482 of the Revised Code; and division (B)(2) of section 4723.484 of the Revised Code;

(3) Establish criteria for the components of the standard care arrangements described in section 4723.431 of the Revised Code that apply to the authority to prescribe, including the components that apply to the authority to prescribe schedule II controlled substances. The rules shall be consistent with that section and include all of the following:

(a) Quality assurance standards;

(b) Standards for periodic review by a collaborating physician or podiatrist of the records of patients treated by the clinical nurse specialist, certified nurse-midwife, or certified nurse practitioner;

(c) Acceptable travel time between the location at which the clinical nurse specialist, certified
nurse-midwife, or certified nurse practitioner is engaging in the prescribing components of the nurse's practice and the location of the nurse's collaborating physician or podiatrist;
(d) Any other criteria recommended by the committee on prescriptive governance.
Sec. 4723.52. (A) As used in this section:
(1) "Community addiction services provider" has the same meaning as in section 5119.01 of the Revised Code.
(2) "Medication-assisted treatment" has the same meaning as in section 340.01 of the Revised Code.
(B) An advanced practice registered nurse shall comply with section 3719.064 of the Revised Code and rules adopted under section 4723.51 of the Revised Code when treating a patient for addiction with medication-assisted treatment or proposing to initiate such treatment.
(C) An advanced practice registered nurse who fails to comply with this section shall treat not more than thirty patients at any one time with medication-assisted treatment even if the facility or location at which the treatment is provided is either of the following:
(1) Exempted by divisions (B)(2)(a) to (d) or (i) of section 4729.553 of the Revised Code from being required to possess a category III terminal distributor of dangerous drugs license with an office-based opioid treatment classification;
(2) A community addiction services provider that provides alcohol and drug addiction services that are certified by the department of mental health and addiction services under section 5119.36 of the Revised Code.
Sec. 4729.01. As used in this chapter:
(A) "Pharmacy," except when used in a context that refers to the practice of pharmacy, means any area, room, rooms, place of business, department, or portion of any of the foregoing where the practice of pharmacy is conducted.
(B) "Practice of pharmacy" means providing pharmacist care requiring specialized knowledge, judgment, and skill derived from the principles of biological, chemical, behavioral, social, pharmaceutical, and clinical sciences. As used in this division, "pharmacist care" includes the following:
(1) Interpreting prescriptions;
(2) Dispensing drugs and drug therapy related devices;
(3) Compounding drugs;
(4) Counseling individuals with regard to their drug therapy, recommending drug therapy related devices, and assisting in the selection of drugs and appliances for treatment of common diseases and injuries and providing instruction in the proper use of the drugs and appliances;
(5) Performing drug regimen reviews with individuals by discussing all of the drugs that the individual is taking and explaining the interactions of the drugs;
(6) Performing drug utilization reviews with licensed health professionals authorized to prescribe drugs when the pharmacist determines that an individual with a prescription has a drug regimen that warrants additional discussion with the prescriber;
(7) Advising an individual and the health care professionals treating an individual with regard to the individual's drug therapy;
(8) Acting pursuant to a consult agreement with one or more physicians authorized under
Chapter 4731. of the Revised Code to practice medicine and surgery or osteopathic medicine and surgery, if an agreement has been established;

(9) Engaging in the administration of immunizations to the extent authorized by section 4729.41 of the Revised Code;

(10) Engaging in the administration of drugs to the extent authorized by section 4729.45 of the Revised Code.

(C) "Compounding" means the preparation, mixing, assembling, packaging, and labeling of one or more drugs in any of the following circumstances:

(1) Pursuant to a prescription issued by a licensed health professional authorized to prescribe drugs;

(2) Pursuant to the modification of a prescription made in accordance with a consult agreement;

(3) As an incident to research, teaching activities, or chemical analysis;

(4) In anticipation of orders for drugs pursuant to prescriptions, based on routine, regularly observed dispensing patterns;

(5) Pursuant to a request made by a licensed health professional authorized to prescribe drugs for a drug that is to be used by the professional for the purpose of direct administration to patients in the course of the professional's practice, if all of the following apply:

(a) At the time the request is made, the drug is not commercially available regardless of the reason that the drug is not available, including the absence of a manufacturer for the drug or the lack of a readily available supply of the drug from a manufacturer.

(b) A limited quantity of the drug is compounded and provided to the professional.

(c) The drug is compounded and provided to the professional as an occasional exception to the normal practice of dispensing drugs pursuant to patient-specific prescriptions.

(D) "Consult agreement" means an agreement that has been entered into under section 4729.39 of the Revised Code.

(E) "Drug" means:

(1) Any article recognized in the United States pharmacopoeia and national formulary, or any supplement to them, intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in humans or animals;

(2) Any other article intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in humans or animals;

(3) Any article, other than food, intended to affect the structure or any function of the body of humans or animals;

(4) Any article intended for use as a component of any article specified in division (E)(1), (2), or (3) of this section; but does not include devices or their components, parts, or accessories.

"Drug" does not include "hemp" or a "hemp product" as those terms are defined in section 928.01 of the Revised Code.

(F) "Dangerous drug" means any of the following:

(1) Any drug to which either of the following applies:

(a) Under the "Federal Food, Drug, and Cosmetic Act," 52 Stat. 1040 (1938), 21 U.S.C.A. 301, as amended, the drug is required to bear a label containing the legend "Caution: Federal law
prohibits dispensing without prescription" or "Caution: Federal law restricts this drug to use by or on
the order of a licensed veterinarian" or any similar restrictive statement, or the drug may be dispensed
only upon a prescription;

(b) Under Chapter 3715. or 3719. of the Revised Code, the drug may be dispensed only upon
a prescription.

(2) Any drug that contains a schedule V controlled substance and that is exempt from Chapter
3719. of the Revised Code or to which that chapter does not apply;

(3) Any drug intended for administration by injection into the human body other than through
a natural orifice of the human body;

(4) Any drug that is a biological product, as defined in section 3715.01 of the Revised Code.

(G) "Federal drug abuse control laws" has the same meaning as in section 3719.01 of the
Revised Code.

(H) "Prescription" means all of the following:

(1) A written, electronic, or oral order for drugs or combinations or mixtures of drugs to be
used by a particular individual or for treating a particular animal, issued by a licensed health
professional authorized to prescribe drugs;

(2) For purposes of sections 2925.61, 4723.488, 4723.484, 4730.434, 4730.433, 4730.434,
and 4731.94 of the Revised Code, a written, electronic, or oral order for naloxone issued to and in the name of a
family member, friend, or other individual in a position to assist an individual who there is reason to
believe is at risk of experiencing an opioid-related overdose.

(3) For purposes of section 4729.44 of the Revised Code, a written, electronic, or oral order
for naloxone issued to and in the name of either of the following:

(a) An individual who there is reason to believe is at risk of experiencing an opioid-related
overdose;

(b) A family member, friend, or other individual in a position to assist an individual who there
is reason to believe is at risk of experiencing an opioid-related overdose.

(4) For purposes of sections 4723.4810, 4729.282, 4730.432, and 4731.93 of the Revised
Code, a written, electronic, or oral order for a drug to treat chlamydia, gonorrhea, or trichomoniasis
issued to and in the name of a patient who is not the intended user of the drug but is the sexual
partner of the intended user;

(5) For purposes of sections 3313.7110, 3313.7111, 3314.143, 3326.28, 3328.29, 4723.483,
4729.88, 4730.433, 4731.96, and 5101.76 of the Revised Code, a written, electronic, or oral order for
an epinephrine autoinjector issued to and in the name of a school, school district, or camp;

(6) For purposes of Chapter 3728. and sections 4723.483, 4729.88, 4730.433, and 4731.96 of
the Revised Code, a written, electronic, or oral order for an epinephrine autoinjector issued to and in
the name of a qualified entity, as defined in section 3728.01 of the Revised Code.

(I) "Licensed health professional authorized to prescribe drugs" or "prescriber" means an
individual who is authorized by law to prescribe drugs or dangerous drugs or drug therapy related
devices in the course of the individual's professional practice, including only the following:

(1) A dentist licensed under Chapter 4715. of the Revised Code;

(2) A clinical nurse specialist, certified nurse-midwife, or certified nurse practitioner who
holds a current, valid license issued under Chapter 4723. of the Revised Code to practice nursing as
an advanced practice registered nurse;

(3) A certified registered nurse anesthetist who holds a current, valid license issued under Chapter 4723. of the Revised Code to practice nursing as an advanced practice registered nurse, but only to the extent of the nurse's authority under sections 4723.43 and 4723.434 of the Revised Code;

(4) An optometrist licensed under Chapter 4725. of the Revised Code to practice optometry under a therapeutic pharmaceutical agents certificate;

(5) A physician authorized under Chapter 4731. of the Revised Code to practice medicine and surgery, osteopathic medicine and surgery, or podiatric medicine and surgery;

(6) A physician assistant who holds a license to practice as a physician assistant issued under Chapter 4730. of the Revised Code, holds a valid prescriber number issued by the state medical board, and has been granted physician-delegated prescriptive authority;

(7) A veterinarian licensed under Chapter 4741. of the Revised Code.

(J) "Sale" or "sell" includes any transaction made by any person, whether as principal proprietor, agent, or employee, to do or offer to do any of the following: deliver, distribute, broker, exchange, gift or otherwise give away, or transfer, whether the transfer is by passage of title, physical movement, or both.

(K) "Wholesale sale" and "sale at wholesale" mean any sale in which the purpose of the purchaser is to resell the article purchased or received by the purchaser.

(L) "Retail sale" and "sale at retail" mean any sale other than a wholesale sale or sale at wholesale.

(M) "Retail seller" means any person that sells any dangerous drug to consumers without assuming control over and responsibility for its administration. Mere advice or instructions regarding administration do not constitute control or establish responsibility.

(N) "Price information" means the price charged for a prescription for a particular drug product and, in an easily understandable manner, all of the following:

(1) The proprietary name of the drug product;

(2) The established (generic) name of the drug product;

(3) The strength of the drug product if the product contains a single active ingredient or if the drug product contains more than one active ingredient and a relevant strength can be associated with the product without indicating each active ingredient. The established name and quantity of each active ingredient are required if such a relevant strength cannot be so associated with a drug product containing more than one ingredient.

(4) The dosage form;

(5) The price charged for a specific quantity of the drug product. The stated price shall include all charges to the consumer, including, but not limited to, the cost of the drug product, professional fees, handling fees, if any, and a statement identifying professional services routinely furnished by the pharmacy. Any mailing fees and delivery fees may be stated separately without repetition. The information shall not be false or misleading.

(O) "Wholesale distributor of dangerous drugs" or "wholesale distributor" means a person engaged in the sale of dangerous drugs at wholesale and includes any agent or employee of such a person authorized by the person to engage in the sale of dangerous drugs at wholesale.

(P) "Manufacturer of dangerous drugs" or "manufacturer" means a person, other than a
pharmacist or prescriber, who manufactures dangerous drugs and who is engaged in the sale of those dangerous drugs.

(Q) "Terminal distributor of dangerous drugs" or "terminal distributor" means a person who is engaged in the sale of dangerous drugs at retail, or any person, other than a manufacturer, repackager, outsourcing facility, third-party logistics provider, wholesale distributor, or pharmacist, who has possession, custody, or control of dangerous drugs for any purpose other than for that person's own use and consumption. "Terminal distributor" includes pharmacies, hospitals, nursing homes, and laboratories and all other persons who procure dangerous drugs for sale or other distribution by or under the supervision of a pharmacist, licensed health professional authorized to prescribe drugs, or other person authorized by the state board of pharmacy.

(R) "Promote to the public" means disseminating a representation to the public in any manner or by any means, other than by labeling, for the purpose of inducing, or that is likely to induce, directly or indirectly, the purchase of a dangerous drug at retail.

(S) "Person" includes any individual, partnership, association, limited liability company, or corporation, the state, any political subdivision of the state, and any district, department, or agency of the state or its political subdivisions.

(T) "Animal shelter" means a facility operated by a humane society or any society organized under Chapter 1717. of the Revised Code or a dog pound operated pursuant to Chapter 955. of the Revised Code.

(U) "Food" has the same meaning as in section 3715.01 of the Revised Code.

(V) "Pain management clinic" has the same meaning as in section 4731.054 of the Revised Code.

(W) "Investigational drug or product" means a drug or product that has successfully completed phase one of the United States food and drug administration clinical trials and remains under clinical trial, but has not been approved for general use by the United States food and drug administration. "Investigational drug or product" does not include controlled substances in schedule I, as defined in section 3719.01 of the Revised Code.

(X) "Product," when used in reference to an investigational drug or product, means a biological product, other than a drug, that is made from a natural human, animal, or microorganism source and is intended to treat a disease or medical condition.

(Y) "Third-party logistics provider" means a person that provides or coordinates warehousing or other logistics services pertaining to dangerous drugs including distribution, on behalf of a manufacturer, wholesale distributor, or terminal distributor of dangerous drugs, but does not take ownership of the drugs or have responsibility to direct the sale or disposition of the drugs.

(Z) "Repackager of dangerous drugs" or "repackager" means a person that repacks and relabels dangerous drugs for sale or distribution.

(AA) "Outsourcing facility" means a facility that is engaged in the compounding and sale of sterile drugs and is registered as an outsourcing facility with the United States food and drug administration.

(BB) "Laboratory" means a laboratory licensed under this chapter as a terminal distributor of dangerous drugs and entrusted to have custody of any of the following drugs and to use the drugs for scientific and clinical purposes and for purposes of instruction: dangerous drugs that are not
controlled substances, as defined in section 3719.01 of the Revised Code; dangerous drugs that are controlled substances, as defined in that section; and controlled substances in schedule I, as defined in that section.

Sec. 4729.29. Divisions (A) and (B) of section 4729.01 and section 4729.28 of the Revised Code do not do any of the following:

(A) Apply to a licensed health professional authorized to prescribe drugs who is acting within the prescriber's scope of professional practice;

(B) Prevent a prescriber from personally furnishing the prescriber's patients with drugs, within the prescriber's scope of professional practice, that seem proper to the prescriber, as long as the drugs are furnished in accordance with section 4729.291 of the Revised Code;

(C) Apply to an individual who personally furnishes a supply of naloxone under authority conferred by a physician under section 4723.485, 4730.435, or 4731.941 of the Revised Code or prevent that individual from personally furnishing the supply of naloxone in accordance with a protocol established by the physician under section 4723.485, 4730.435, or 4731.941 of the Revised Code;

(D) Apply to the sale of oxygen, the sale of peritoneal dialysis solutions, or the sale of drugs that are not dangerous drugs by a retail dealer, in original packages when labeled as required by the "Federal Food, Drug, and Cosmetic Act," 52 Stat. 1040 (1938), 21 U.S.C.A. 301, as amended.

Sec. 4729.44. (A) As used in this section:

(1) "Board of health" means a board of health of a city or general health district or an authority having the duties of a board of health under section 3709.05 of the Revised Code.

(B) "Physician" means an individual authorized under Chapter 4731. of the Revised Code to practice medicine and surgery, osteopathic medicine and surgery, or podiatric medicine and surgery.

(B) If use of the protocol developed pursuant to rules adopted under division (G) of this section has been authorized under section 3707.56 or 4731.942 of the Revised Code, a pharmacist or pharmacy intern may dispense naloxone without a prescription to either of the following in accordance with that protocol:

(1) An individual who there is reason to believe is experiencing or at risk of experiencing an opioid-related overdose;

(2) A family member, friend, or other individual in a position to assist an individual who there is reason to believe is at risk of experiencing an opioid-related overdose.

(C) A pharmacist or pharmacy intern who dispenses naloxone under this section shall instruct the individual to whom naloxone is dispensed to summon emergency services as soon as practicable either before or after administering naloxone.

(D) A pharmacist may document on a prescription form the dispensing of naloxone by the pharmacist or a pharmacy intern supervised by the pharmacist. The form may be assigned a number for record-keeping purposes.

(E) This section does not affect the authority of a pharmacist or pharmacy intern to fill or refill a prescription for naloxone.

(F) A board of health that in good faith authorizes a pharmacist or pharmacy intern to dispense naloxone without a prescription in accordance with a protocol developed pursuant to rules adopted under division (G) of this section is not liable for or subject to any of the following for any
action or omission of the individual to whom the naloxone is dispensed: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

A physician who in good faith authorizes a pharmacist or pharmacy intern to dispense naloxone without a prescription in accordance with a protocol developed pursuant to rules adopted under division (G) of this section is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is dispensed: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

A pharmacist or pharmacy intern authorized under this section to dispense naloxone without a prescription who does so in good faith is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is dispensed: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

(G) The state board of pharmacy shall, after consulting with the department of health and state medical board, adopt rules to implement this section. The rules shall specify a protocol under which pharmacists or pharmacy interns may dispense naloxone without a prescription.

All rules adopted under this section shall be adopted in accordance with Chapter 119. of the Revised Code.

(H)(1) The state board of pharmacy shall develop a program to educate all of the following about the authority of a pharmacist or pharmacy intern to dispense naloxone without a prescription:

(a) Holders of licenses issued under this chapter that engage in the sale or dispensing of naloxone pursuant to this section;
(b) Registered pharmacy technicians, certified pharmacy technicians, and pharmacy technician trainees registered under this chapter who engage in the sale of naloxone pursuant to this section;
(c) Individuals who are not licensed or registered under this chapter but are employed by license holders described in division (H)(1)(a) of this section.

(2) As part of the program, the board also shall educate the license holders, pharmacy technicians, and employees described in division (H)(1) of this section about maintaining an adequate supply of naloxone and methods for determining a pharmacy's stock of the drug.

(3) The board may use its web site to share information under the program.

Sec. 4729.45. (A) As used in this section, "physician" means an individual authorized under Chapter 4731. of the Revised Code to practice medicine and surgery or osteopathic medicine and surgery.

(B)(1) Subject to division (C) of this section, a pharmacist licensed under this chapter may administer by injection any of the following drugs as long as the drug that is to be administered has been prescribed by a physician and the individual to whom the drug was prescribed has an ongoing physician-patient relationship with the physician:

(a) An opioid antagonist used for treatment of drug addiction and an addiction treatment drug administered in a long-acting or extended-release form;
(b) An antipsychotic drug administered in a long-acting or extended-release form;
(c) Hydroxyprogesterone caproate;
(d) Medroxyprogesterone acetate;
(e) Cobalamin.
(2) As part of engaging in the administration of drugs by injection pursuant to this section, a pharmacist may administer epinephrine or diphenhydramine, or both, to an individual in an emergency situation resulting from an adverse reaction to a drug administered by the pharmacist.

(C) To be authorized to administer drugs pursuant to this section, a pharmacist must do all of the following:

(1) Successfully complete a course in the administration of drugs that satisfies the requirements established by the state board of pharmacy in rules adopted under division (H)(1)(a) of this section;

(2) Receive and maintain certification to perform basic life-support procedures by successfully completing a basic life-support training course that is certified by the American red cross or American heart association or approved by the state board of pharmacy;

(3) Practice in accordance with a protocol that meets the requirements of division (F) of this section.

(D) Each time a pharmacist administers a drug pursuant to this section, the pharmacist shall do all of the following:

(1) Obtain permission in accordance with the procedures specified in rules adopted under division (H) of this section and comply with the following requirements:

(a) Except as provided in division (D)(1)(c) of this section, for each drug administered by a pharmacist to an individual who is eighteen years of age or older, the pharmacist shall obtain permission from the individual.

(b) For each drug administered by a pharmacist to an individual who is under eighteen years of age, the pharmacist shall obtain permission from the individual's parent or other person having care or charge of the individual.

(c) For each drug administered by a pharmacist to an individual who lacks the capacity to make informed health care decisions, the pharmacist shall obtain permission from the person authorized to make such decisions on the individual's behalf.

(2) In the case of an opioid antagonist—an addiction treatment drug described in division (B)(1)(a) of this section, obtain in accordance with division (E) of this section test results indicating that it is appropriate to administer the drug to the individual if either of the following is to be administered:

(a) The initial dose of the drug;

(b) Any subsequent dose, if the administration occurs more than thirty days after the previous dose of the drug was administered.

(3) Observe the individual to whom the drug is administered to determine whether the individual has an adverse reaction to the drug;

(4) Notify the physician who prescribed the drug that the drug has been administered to the individual.

(E) A pharmacist may obtain the test results described in division (D)(2) of this section in either of the following ways:

(1) From the physician;

(2) By ordering blood and urine tests for the individual to whom the opioid antagonist drug is to be administered.
If a pharmacist orders blood and urine tests, the pharmacist shall evaluate the results of the tests to determine whether they indicate that it is appropriate to administer the opioid antagonist drug. A pharmacist's authority to evaluate test results under this division does not authorize the pharmacist to make a diagnosis.

(F) All of the following apply with respect to the protocol required by division (C)(3) of this section:

1. The protocol must be established by a physician who has a scope of practice that includes treatment of the condition for which the individual has been prescribed the drug to be administered.
2. The protocol must satisfy the requirements established in rules adopted under division (H)(1)(b) of this section.
3. The protocol must do all of the following:
   a. Specify a definitive set of treatment guidelines;
   b. Specify the locations at which a pharmacist may engage in the administration of drugs pursuant to this section;
   c. Include provisions for implementing the requirements of division (D) of this section, including for purposes of division (D)(3) of this section provisions specifying the length of time and location at which a pharmacist must observe an individual who receives a drug to determine whether the individual has an adverse reaction to the drug;
   d. Specify procedures to be followed by a pharmacist when administering epinephrine, diphenhydramine, or both, to an individual who has an adverse reaction to a drug administered by the pharmacist.

(G) A pharmacist shall not do either of the following:
1. Engage in the administration of drugs pursuant to this section unless the requirements of division (C) of this section have been met;
2. Delegate to any person the pharmacist's authority to engage in the administration of drugs pursuant to this section.

(H)(1) The state board of pharmacy shall adopt rules to implement this section. The rules shall be adopted in accordance with Chapter 119. of the Revised Code and include all of the following:
   a. Requirements for courses in administration of drugs;
   b. Requirements for protocols to be followed by pharmacists in administering drugs pursuant to this section;
   c. Procedures to be followed by a pharmacist in obtaining permission to administer a drug to an individual.

2. The board shall consult with the state medical board before adopting rules regarding requirements for protocols under this section.

Sec. 4729.51. (A) No person other than a licensed manufacturer of dangerous drugs, outsourcing facility, third-party logistics provider, repackager of dangerous drugs, or wholesale distributor of dangerous drugs shall possess for sale, sell, distribute, or deliver, at wholesale, dangerous drugs or investigational drugs or products, except as follows:

1. A licensed terminal distributor of dangerous drugs that is a pharmacy may make occasional sales of dangerous drugs or investigational drugs or products at wholesale.
(2) A licensed terminal distributor of dangerous drugs having more than one licensed location may transfer or deliver dangerous drugs from one licensed location to another licensed location owned by the terminal distributor if the license issued for each location is in effect at the time of the transfer or delivery.

(3) A licensed terminal distributor of dangerous drugs that is not a pharmacy may make occasional sales of

naloxone

the following at wholesale:

(4) A licensed terminal distributor of dangerous drugs that is not a pharmacy may make occasional sales of dangerous:

(a) Naloxone;

(b) Dangerous drugs at wholesale if the drugs being sold are in shortage, as defined in rules adopted by the state board of pharmacy under section 4729.26 of the Revised Code;

(c) Dangerous drugs other than those described in divisions (A)(3)(a) and (b) of this section or investigational drugs or products if authorized by rules adopted under section 4729.26 of the Revised Code.

(B) No licensed manufacturer, outsourcing facility, third-party logistics provider, repackager, or wholesale distributor shall possess for sale, sell, or distribute, at wholesale, dangerous drugs or investigational drugs or products to any person other than the following:

(1) Subject to division (D) of this section, a licensed terminal distributor of dangerous drugs;

(2) Subject to division (C) of this section, any person exempt from licensure as a terminal distributor of dangerous drugs under section 4729.541 of the Revised Code;

(3) A licensed manufacturer, outsourcing facility, third-party logistics provider, repackager, or wholesale distributor;

(4) A terminal distributor, manufacturer, outsourcing facility, third-party logistics provider, repackager, or wholesale distributor that is located in another state, is not engaged in the sale of dangerous drugs within this state, and is actively licensed to engage in the sale of dangerous drugs by the state in which the distributor conducts business.

(C) No licensed manufacturer, outsourcing facility, third-party logistics provider, repackager, or wholesale distributor shall possess for sale, sell, or distribute, at wholesale, dangerous drugs or investigational drugs or products to either of the following:

(1) A prescriber who is employed by either of the following:

(a) A pain management clinic that is not licensed as a terminal distributor of dangerous drugs with a pain management clinic classification issued under section 4729.552 of the Revised Code;

(b) A facility, clinic, or other location that provides office-based opioid treatment but is not licensed as a terminal distributor of dangerous drugs with an office-based opioid treatment classification issued under section 4729.553 of the Revised Code if such a license is required by that section.

(2) A business entity described in division (A)(2) or (3) of section 4729.541 of the Revised Code that is, or is operating, either of the following:

(a) A pain management clinic without a license as a terminal distributor of dangerous drugs with a pain management clinic classification issued under section 4729.552 of the Revised Code;

(b) A facility, clinic, or other location that provides office-based opioid treatment without a license as a terminal distributor of dangerous drugs with an office-based opioid treatment
classification issued under section 4729.553 of the Revised Code if such a license is required by that
section.

(D) No licensed manufacturer, outsourcing facility, third-party logistics provider, repackager,
or wholesale distributor shall possess dangerous drugs or investigational drugs or products for sale at
wholesale, or sell or distribute such drugs at wholesale, to a licensed terminal distributor of
dangerous drugs, except as follows:

(1) In the case of a terminal distributor with a category II license, only dangerous drugs in
category II, as defined in division (A)(1) of section 4729.54 of the Revised Code;

(2) In the case of a terminal distributor with a category III license, dangerous drugs in
category II and category III, as defined in divisions (A)(1) and (2) of section 4729.54 of the Revised
Code;

(3) In the case of a terminal distributor with a limited category II or III license, only the
dangerous drugs specified in the license.

(E)(1) Except as provided in division (E)(2) of this section, no person shall do any of the
following:

(a) Sell or distribute, at retail, dangerous drugs;

(b) Possess for sale, at retail, dangerous drugs;

(c) Possess dangerous drugs.

(2)(a) Divisions (E)(1)(a), (b), and (c) of this section do not apply to any of the following:

(i) A licensed terminal distributor of dangerous drugs;

(ii) A person who possesses, or possesses for sale or sells, at retail, a dangerous drug in
accordance with Chapters 3719., 4715., 4723., 4725., 4729., 4730., 4731., and 4741. of the Revised
Code;

(iii) Any of the persons identified in divisions (A)(1) to (5) and (13) of section 4729.541 of
the Revised Code, but only to the extent specified in that section.

(b) Division (E)(1)(c) of this section does not apply to any of the following:

(i) A licensed manufacturer, outsourcing facility, third-party logistics provider, repackager, or
wholesale distributor;

(ii) Any of the persons identified in divisions (A)(6) to (12) of section 4729.541 of the
Revised Code, but only to the extent specified in that section.

(F) No licensed terminal distributor of dangerous drugs or person that is exempt from
licensure under section 4729.541 of the Revised Code shall purchase dangerous drugs or
investigational drugs or products from any person other than a licensed manufacturer, outsourcing
facility, third-party logistics provider, repackager, or wholesale distributor, except as follows:

(1) A licensed terminal distributor of dangerous drugs or person that is exempt from licensure
under section 4729.541 of the Revised Code may make occasional purchases of dangerous drugs or
investigational drugs or products that are sold in accordance with division (A)(1) or (3) of this
section.

(2) A licensed terminal distributor of dangerous drugs having more than one licensed location
may transfer or deliver dangerous drugs or investigational drugs or products from one licensed
location to another licensed location if the license issued for each location is in effect at the time of
the transfer or delivery.
(G) No licensed terminal distributor of dangerous drugs shall engage in the retail sale or other
distribution of dangerous drugs or investigational drugs or products or maintain possession, custody,
or control of dangerous drugs or investigational drugs or products for any purpose other than the
distributor's personal use or consumption, at any establishment or place other than that or those
described in the license issued by the state board of pharmacy to such terminal distributor.

(H) Nothing in this section shall be construed to interfere with the performance of official
duties by any law enforcement official authorized by municipal, county, state, or federal law to
collect samples of any drug, regardless of its nature or in whose possession it may be.

(I) Notwithstanding anything to the contrary in this section, the board of education of a city,
local, exempted village, or joint vocational school district may distribute epinephrine autoinjectors
for use in accordance with section 3313.7110 of the Revised Code and may distribute inhalers for use
in accordance with section 3313.7113 of the Revised Code.

Sec. 4729.514. (A) As used in this section, "service entity" means a public or private entity
that may provide services to or interact with individuals who there is reason to believe may be at risk
of experiencing an opioid-related overdose. "Service entity" includes a church or other place of
worship, college or university, school, local library, health department operated by the board of health
of a city or general health district, community addiction services provider, court, probation
department, halfway house, prison, jail, community residential center, homeless shelter, or similar
entity.

(B) A service entity may procure and maintain naloxone for either or both of the following
purposes:

(1) To use in emergency situations;
(2) To permit an employee, volunteer, or contractor of the service entity to personally furnish
a supply of naloxone pursuant to a protocol established under section 3707.561, 4723.485, 4730.435,
or 4731.941 of the Revised Code.

(C) A service entity or an employee, volunteer, or contractor of a service entity is not liable
for or subject to any of the following for injury, death, or loss to person or property that allegedly
arises from an act or omission associated with procuring, maintaining, accessing, or using, or
personally furnishing naloxone under this section, unless the act or omission constitutes willful or
wanton misconduct: damages in any civil action, prosecution in any criminal proceeding, or
professional disciplinary action.

This section does not eliminate, limit, or reduce any other immunity or defense that a service
entity or an employee, volunteer, or contractor of a service entity may be entitled to under Chapter
2305. or any other provision of the Revised Code or under the common law of this state.

Sec. 4729.515. (A) In accordance with divisions (B) and (C) of this section, a terminal
distributor of dangerous drugs may acquire and maintain a supply of naloxone for use in emergency
situations and for distribution through an automated mechanism. The naloxone may be maintained at
a location other than the location licensed as a terminal distributor of dangerous drugs.

(B) In the case of naloxone for use in emergency situations, a terminal distributor of
dangerous drugs shall do all of the following:

(1) Provide instructions regarding the emergency administration of naloxone to any
individual who accesses the naloxone, including a specific instruction to summon emergency
services as set forth in division (D) of this section;

(2) Specify a process to be used to notify the terminal distributor that the naloxone has been accessed within a reasonable time of its being accessed;

(3) Maintain the naloxone in accordance with the manufacturer's or distributor's instructions.

(C) In the case of naloxone for distribution through an automated mechanism, a terminal distributor of dangerous drugs shall comply with standards and procedures specified in rules adopted under division (F) of this section.

(D)(1) Notwithstanding any conflicting provision of the Revised Code, both of the following apply:

(a) Any individual may access naloxone maintained as provided in division (B) of this section and may administer it to an individual who there is reason to believe is experiencing an opioid-related overdose.

(b) Any individual may receive naloxone distributed through an automated system as provided in division (C) of this section and may administer it to an individual who there is reason to believe is experiencing an opioid-related overdose.

(2) An individual who administers naloxone as authorized by this section shall make a good faith effort to activate or have another individual activate an emergency medical services system as soon as possible, except that this requirement does not apply if the individual administering the naloxone is doing so as part of an emergency medical services system or at a hospital, as defined in section 3727.01 of the Revised Code.

(E) An individual is not liable for or subject to any of the following for injury, death, or loss to person or property that allegedly arises from an act or omission associated with any action authorized by this section, unless the act or omission constitutes willful or wanton misconduct: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

(F) The state board of pharmacy shall adopt rules establishing standards and procedures applicable to the distribution of naloxone through an automated mechanism. The rules shall be adopted in accordance with Chapter 119. of the Revised Code.

Sec. 4729.541. (A) Except as provided in divisions (B) to (D) of this section, all of the following are exempt from licensure as a terminal distributor of dangerous drugs:

(1) A licensed health professional authorized to prescribe drugs;

(2) A business entity that is a corporation formed under division (B) of section 1701.03 of the Revised Code, a limited liability company formed under Chapter 1705. of the Revised Code, or a professional association formed under Chapter 1785. of the Revised Code if the entity has a sole shareholder who is a prescriber and is authorized to provide the professional services being offered by the entity;

(3) A business entity that is a corporation formed under division (B) of section 1701.03 of the Revised Code, a limited liability company formed under Chapter 1705. of the Revised Code, a partnership or a limited liability partnership formed under Chapter 1775. of the Revised Code, or a professional association formed under Chapter 1785. of the Revised Code, if, to be a shareholder, member, or partner, an individual is required to be licensed, certified, or otherwise legally authorized under Title XLVII of the Revised Code to perform the professional service provided by the entity and
each such individual is a prescriber;

(4) An individual who holds a current license, certificate, or registration issued under Title XLVII of the Revised Code and has been certified to conduct diabetes education by a national certifying body specified in rules adopted by the state board of pharmacy under section 4729.68 of the Revised Code, but only with respect to insulin that will be used for the purpose of diabetes education and only if diabetes education is within the individual's scope of practice under statutes and rules regulating the individual's profession;

(5) An individual who holds a valid certificate issued by a nationally recognized S.C.U.B.A. diving certifying organization approved by the state board of pharmacy under rules adopted by the board, but only with respect to medical oxygen that will be used for the purpose of emergency care or treatment at the scene of a diving emergency;

(6) With respect to epinephrine autoinjectors that may be possessed under section 3313.7110, 3313.7111, 3314.143, 3326.28, or 3328.29 of the Revised Code, any of the following: the board of education of a city, local, exempted village, or joint vocational school district; a chartered or nonchartered nonpublic school; a community school established under Chapter 3314. of the Revised Code; a STEM school established under Chapter 3326. of the Revised Code; or a college-preparatory boarding school established under Chapter 3328. of the Revised Code;

(7) With respect to epinephrine autoinjectors that may be possessed under section 5101.76 of the Revised Code, any of the following: a residential camp, as defined in section 2151.011 of the Revised Code; a child day camp, as defined in section 5104.01 of the Revised Code; or a child day camp operated by any county, township, municipal corporation, township park district created under section 511.18 of the Revised Code, park district created under section 1545.04 of the Revised Code, or joint recreation district established under section 755.14 of the Revised Code;

(8) With respect to epinephrine autoinjectors that may be possessed under Chapter 3728. of the Revised Code, a qualified entity, as defined in section 3728.01 of the Revised Code;

(9) With respect to inhalers that may be possessed under section 3313.7113, 3313.7114, 3314.144, 3326.30, or 3328.30 of the Revised Code, any of the following: the board of education of a city, local, exempted village, or joint vocational school district; a chartered or nonchartered nonpublic school; a community school established under Chapter 3314. of the Revised Code; a STEM school established under Chapter 3326. of the Revised Code; or a college-preparatory boarding school established under Chapter 3328. of the Revised Code;

(10) With respect to inhalers that may be possessed under section 5101.77 of the Revised Code, any of the following: a residential camp, as defined in section 2151.011 of the Revised Code; a child day camp, as defined in section 5104.01 of the Revised Code; or a child day camp operated by any county, township, municipal corporation, township park district created under section 511.18 of the Revised Code, park district created under section 1545.04 of the Revised Code, or joint recreation district established under section 755.14 of the Revised Code;

(11) With respect to naloxone that may be possessed under section 2925.61 of the Revised Code, a law enforcement agency and its peace officers;

(12) With respect to naloxone that may be possessed under section 4729.514 of the Revised Code, for use in emergency situations or for personally furnishing supplies of naloxone, a service entity, as defined in that section;
(13) A facility that is owned and operated by the United States department of defense, the United States department of veterans affairs, or any other federal agency.

(B) If a person described in division (A) of this section is a pain management clinic or is operating a pain management clinic, the person shall hold a license as a terminal distributor of dangerous drugs with a pain management clinic classification issued under section 4729.552 of the Revised Code.

(C) If a person described in division (A) of this section is operating a facility, clinic, or other location described in division (B) of section 4729.553 of the Revised Code that must hold a category III terminal distributor of dangerous drugs license with an office-based opioid treatment classification, the person shall hold a license with that classification.

(D) Any of the persons described in divisions (A)(1) to (12) of this section shall hold a license as a terminal distributor of dangerous drugs in order to possess, have custody or control of, and distribute any of the following:

(1) Dangerous drugs that are compounded or used for the purpose of compounding;
(2) A schedule I, II, III, IV, or V controlled substance, as defined in section 3719.01 of the Revised Code.

Sec. 4729.553. (A) As used in this section:

(1) "Advanced practice registered nurse" has the same meaning as in section 4723.01 of the Revised Code.

(2) "Controlled substance" has the same meaning as in section 3719.01 of the Revised Code.

(3) "Hospital" means a hospital registered with the department of health under section 3701.07 of the Revised Code.

(4) "Office-based opioid treatment" means the treatment of opioid dependence or addiction using a controlled substance.

(5) "Physician" means an individual who is authorized under Chapter 4731. of the Revised Code to practice medicine and surgery or osteopathic medicine and surgery.

(6) "Physician assistant" means an individual who is licensed under Chapter 4730. of the Revised Code.

(B) (1) Except as provided in division (B)(2) and (3) of this section, no person shall knowingly operate a facility, clinic, or other location where a prescriber provides office-based opioid treatment to more than thirty patients or that meets any other identifying criteria established in rules adopted under this section without holding a category III terminal distributor of dangerous drugs license with an office-based opioid treatment classification.

(2) Division (B)(1) of this section does not apply to any of the following:

(a) A hospital;
(b) A facility for the treatment of opioid dependence or addiction that is operated by a hospital;
(c) A physician practice owned or controlled, in whole or in part, by a hospital or by an entity that owns or controls, in whole or in part, one or more hospitals;
(d) A facility that conducts only clinical research and uses controlled substances in studies approved by a hospital-based institutional review board or an institutional review board that is accredited by the association for the accreditation of human research protection programs, inc.;
(e) A facility that holds a category III terminal distributor of dangerous drugs license in accordance with section 4729.54 of the Revised Code for the purpose of treating drug dependence or addiction as part of an opioid treatment program and is the subject of a current, valid certification from the substance abuse and mental health services administration of the United States department of health and human services pursuant to 42 C.F.R. 8.11;

(f) A program or facility that holds a license or certification issued by the department of mental health and addiction services under Chapter 5119. of the Revised Code if the license or certification is approved by the state board of pharmacy;

(g) A federally qualified health center or federally qualified health center look-alike, as defined in section 3701.047 of the Revised Code;

(h) A state or local correctional facility, as defined in section 5163.45 of the Revised Code;

(i) A facility in which patients are treated on-site for opioid dependence or addiction exclusively through direct administration by a physician, physician assistant, or advanced practice registered nurse of drugs that are used for treatment of opioid dependence or addiction and are neither dispensed nor personally furnished to patients for off-site self-administration;

(j) Any other facility specified in rules adopted under this section.

(3) A patient who receives treatment on-site for opioid dependence or addiction through direct administration of a drug by a physician, physician assistant, or advanced practice registered nurse shall not be included in determining whether more than thirty patients are being provided office-based opioid treatment in a particular facility, clinic, or other location that is subject to division (B)(1) of this section.

(C) To be eligible to receive a license as a category III terminal distributor of dangerous drugs with an office-based opioid treatment classification, an applicant shall submit evidence satisfactory to the state board of pharmacy that the applicant's office-based opioid treatment will be operated in accordance with the requirements specified in division (D) of this section and that the applicant meets any other applicable requirements of this chapter.

If the board determines that an applicant meets all of the requirements, the board shall issue to the applicant a license as a category III terminal distributor of dangerous drugs with an office-based opioid treatment classification.

(D) The holder of a category III terminal distributor license with an office-based opioid treatment classification shall do all of the following:

(1) Be in control of a facility that is owned and operated solely by one or more physicians authorized under Chapter 4731. of the Revised Code to practice medicine and surgery or osteopathic medicine and surgery, unless the state board of pharmacy waives this requirement for the holder;

(2) Comply with the requirements for conducting office-based opioid treatment, as established by the state medical board in rules adopted under section 4731.056 of the Revised Code;

(3) Require any person with ownership of the facility to submit to a criminal records check in accordance with section 4776.02 of the Revised Code and send the results of the criminal records check directly to the state board of pharmacy for review and decision under section 4729.071 of the Revised Code;

(4) Require each person employed by or seeking employment with the facility to submit to a criminal records check in accordance with section 4776.02 of the Revised Code;
(5) Ensure that a person is not employed by the facility if the person, within the ten years immediately preceding the date the person applied for employment, was convicted of or pleaded guilty to either of the following, unless the state board of pharmacy permits the person to be employed by waiving this requirement for the facility:

(a) A theft offense, described in division (K)(3) of section 2913.01 of the Revised Code, that would constitute a felony under the laws of this state, any other state, or the United States;

(b) A felony drug offense, as defined in section 2925.01 of the Revised Code.

(6) Maintain a list of each person with ownership of the facility and notify the state board of pharmacy of any change to that list.

(E) No person subject to licensure as a category III terminal distributor of dangerous drugs with an office-based opioid treatment classification shall knowingly fail to remain in compliance with the requirements of division (D) of this section and any other applicable requirements of this chapter.

(F) The state board of pharmacy may impose a fine of not more than five thousand dollars on a person who violates division (B) or (E) of this section. A separate fine may be imposed for each day the violation continues. In imposing the fine, the board's actions shall be taken in accordance with Chapter 119. of the Revised Code.

(G) The state board of pharmacy shall adopt rules as it considers necessary to implement and administer this section. The rules shall be adopted in accordance with Chapter 119. of the Revised Code.

Sec. 4729.80. (A) If the state board of pharmacy establishes and maintains a drug database pursuant to section 4729.75 of the Revised Code, the board is authorized or required to provide information from the database only as follows:

(1) On receipt of a request from a designated representative of a government entity responsible for the licensure, regulation, or discipline of health care professionals with authority to prescribe, administer, or dispense drugs, the board may provide to the representative information from the database relating to the professional who is the subject of an active investigation being conducted by the government entity or relating to a professional who is acting as an expert witness for the government entity in such an investigation.

(2) On receipt of a request from a federal officer, or a state or local officer of this or any other state, whose duties include enforcing laws relating to drugs, the board shall provide to the officer information from the database relating to the person who is the subject of an active investigation of a drug abuse offense, as defined in section 2925.01 of the Revised Code, being conducted by the officer's employing government entity.

(3) Pursuant to a subpoena issued by a grand jury, the board shall provide to the grand jury information from the database relating to the person who is the subject of an investigation being conducted by the grand jury.

(4) Pursuant to a subpoena, search warrant, or court order in connection with the investigation or prosecution of a possible or alleged criminal offense, the board shall provide information from the database as necessary to comply with the subpoena, search warrant, or court order.

(5) On receipt of a request from a prescriber or the prescriber's delegate approved by the
board, the board shall provide to the prescriber a report of information from the database relating to a patient who is either a current patient of the prescriber or a potential patient of the prescriber based on a referral of the patient to the prescriber, if all of the following conditions are met:

(a) The prescriber certifies in a form specified by the board that it is for the purpose of providing medical treatment to the patient who is the subject of the request;

(b) The prescriber has not been denied access to the database by the board.

(6) On receipt of a request from a pharmacist or the pharmacist's delegate approved by the board, the board shall provide to the pharmacist information from the database relating to a current patient of the pharmacist, if the pharmacist certifies in a form specified by the board that it is for the purpose of the pharmacist's practice of pharmacy involving the patient who is the subject of the request and the pharmacist has not been denied access to the database by the board.

(7) On receipt of a request from an individual seeking the individual's own database information in accordance with the procedure established in rules adopted under section 4729.84 of the Revised Code, the board may provide to the individual the individual's own prescription history.

(8) On receipt of a request from a medical director or a pharmacy director of a managed care organization that has entered into a contract with the department of medicaid under section 5167.10 of the Revised Code and a data security agreement with the board required by section 5167.14 of the Revised Code, the board shall provide to the medical director or the pharmacy director information from the database relating to a medicaid recipient enrolled in the managed care organization, including information in the database related to prescriptions for the recipient that were not covered or reimbursed under a program administered by the department of medicaid.

(9) On receipt of a request from the medicaid director, the board shall provide to the director information from the database relating to a recipient of a program administered by the department of medicaid, including information in the database related to prescriptions for the recipient that were not covered or paid by a program administered by the department.

(10) On receipt of a request from a medical director of a managed care organization that has entered into a contract with the administrator of workers' compensation under division (B)(4) of section 4121.44 of the Revised Code and a data security agreement with the board required by section 4121.447 of the Revised Code, the board shall provide to the medical director information from the database relating to a claimant under Chapter 4121., 4123., 4127., or 4131. of the Revised Code assigned to the managed care organization, including information in the database related to prescriptions for the claimant that were not covered or reimbursed under Chapter 4121., 4123., 4127., or 4131. of the Revised Code, if the administrator of workers' compensation confirms, upon request from the board, that the claimant is assigned to the managed care organization.

(11) On receipt of a request from the administrator of workers' compensation, the board shall provide to the administrator information from the database relating to a claimant under Chapter 4121., 4123., 4127., or 4131. of the Revised Code, including information in the database related to prescriptions for the claimant that were not covered or reimbursed under Chapter 4121., 4123., 4127., or 4131. of the Revised Code.

(12) On receipt of a request from a prescriber or the prescriber's delegate approved by the board, the board shall provide to the prescriber information from the database relating to a patient's mother, if the prescriber certifies in a form specified by the board that it is for the purpose of
providing medical treatment to a newborn or infant patient diagnosed as opioid dependent and the prescriber has not been denied access to the database by the board.

(13) On receipt of a request from the director of health, the board shall provide to the director information from the database relating to the duties of the director or the department of health in implementing the Ohio violent death reporting system established under section 3701.93 of the Revised Code.

(14) On receipt of a request from a requestor described in division (A)(1), (2), (5), or (6) of this section who is from or participating with another state's prescription monitoring program, the board may provide to the requestor information from the database, but only if there is a written agreement under which the information is to be used and disseminated according to the laws of this state.

(15) On receipt of a request from a delegate of a retail dispensary licensed under Chapter 3796. of the Revised Code who is approved by the board to serve as the dispensary's delegate, the board shall provide to the delegate a report of information from the database pertaining only to a patient's use of medical marijuana, if both of the following conditions are met:

(a) The delegate certifies in a form specified by the board that it is for the purpose of dispensing medical marijuana for use in accordance with Chapter 3796. of the Revised Code.

(b) The retail dispensary or delegate has not been denied access to the database by the board.

(16) On receipt of a request from a judge of a program certified by the Ohio supreme court as a specialized docket program for drugs, the board shall provide to the judge, or an employee of the program who is designated by the judge to receive the information, information from the database that relates specifically to a current or prospective program participant.

(17) On receipt of a request from a coroner, deputy coroner, or coroner's delegate approved by the board, the board shall provide to the requestor information from the database relating to a deceased person about whom the coroner is conducting or has conducted an autopsy or investigation.

(18) On receipt of a request from a prescriber, the board may provide to the prescriber a summary of the prescriber's prescribing record if such a record is created by the board. Information in the summary is subject to the confidentiality requirements of this chapter.

(19)(a) On receipt of a request from a pharmacy's responsible person, the board may provide to the responsible person a summary of the pharmacy's dispensing record if such a record is created by the board. Information in the summary is subject to the confidentiality requirements of this chapter.

(b) As used in division (A)(19)(a) of this section, "responsible person" has the same meaning as in rules adopted by the board under section 4729.26 of the Revised Code.

(20) The board may provide information from the database without request to a prescriber or pharmacist who is authorized to use the database pursuant to this chapter.

(21)(a) On receipt of a request from a prescriber or pharmacist, or the prescriber's or pharmacist's delegate, who is a designated representative of a peer review committee, the board shall provide to the committee information from the database relating to a prescriber who is subject to the committee's evaluation, supervision, or discipline if the information is to be used for one of those purposes. The board shall provide only information that it determines, in accordance with rules adopted under section 4729.84 of the Revised Code, is appropriate to be provided to the committee.
(b) As used in division (A)(21)(a) of this section, "peer review committee" has the same meaning as in section 2305.25 of the Revised Code, except that it includes only a peer review committee of a hospital or a peer review committee of a nonprofit health care corporation that is a member of the hospital or of which the hospital is a member.

(22) On receipt of a request from a requestor described in division (A)(5) or (6) of this section who is from or participating with a prescription monitoring program that is operated by a federal agency and approved by the board, the board may provide to the requestor information from the database, but only if there is a written agreement under which the information is to be used and disseminated according to the laws of this state.

(23) Any personal health information submitted to the board pursuant to section 4729.772 of the Revised Code may be provided by the board only as authorized by the submitter of the information and in accordance with rules adopted under section 4729.84 of the Revised Code.

(B) The state board of pharmacy shall maintain a record of each individual or entity that requests information from the database pursuant to this section. In accordance with rules adopted under section 4729.84 of the Revised Code, the board may use the records to document and report statistics and law enforcement outcomes.

The board may provide records of an individual's requests for database information only to the following:

(1) A designated representative of a government entity that is responsible for the licensure, regulation, or discipline of health care professionals with authority to prescribe, administer, or dispense drugs who is involved in an active criminal or disciplinary investigation being conducted by the government entity of the individual who submitted the requests for database information;

(2) A federal officer, or a state or local officer of this or any other state, whose duties include enforcing laws relating to drugs and who is involved in an active investigation being conducted by the officer's employing government entity of the individual who submitted the requests for database information;

(3) A designated representative of the department of medicaid regarding a prescriber who is treating or has treated a recipient of a program administered by the department and who submitted the requests for database information.

(C) Information contained in the database and any information obtained from it is confidential and is not a public record. Information contained in the records of requests for information from the database is confidential and is not a public record. Information contained in the database that does not identify a person, including any licensee or registrant of the board or other entity, may be released in summary, statistical, or aggregate form.

(D) A pharmacist or prescriber shall not be held liable in damages to any person in any civil action for injury, death, or loss to person or property on the basis that the pharmacist or prescriber did or did not seek or obtain information from the database.

Sec. 4730.434 (A) Notwithstanding any provision of this chapter or rule adopted by the state medical board, a physician assistant who holds a valid prescriber number issued by the board and has been granted physician-delegated prescriptive authority may personally furnish a supply of naloxone, or issue a prescription for naloxone, without having examined the individual to whom it may be administered if both of the following conditions are met:
(1) The naloxone supply is furnished to, or the prescription is issued to and in the name of, a family member, friend, or other individual in a position to assist an individual who there is reason to believe is at risk of experiencing an opioid-related overdose.

(2) The physician assistant instructs the individual receiving the naloxone supply or prescription to summon emergency services as soon as practicable either before or after administering naloxone to an individual apparently experiencing an opioid-related overdose.

(B) A physician assistant who under division (A) of this section in good faith furnishes a supply of naloxone or issues a prescription for naloxone is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is furnished or the prescription is issued: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

Sec. 4730.435. (A) (1) A physician assistant who holds a valid prescriber number issued by the state medical board, who has been granted physician-delegated prescriptive authority, and who has established a protocol that meets the requirements of division (C) of this section may authorize one or more other individuals to personally furnish a supply of naloxone pursuant to the protocol to either of the following:

(a) An individual who there is reason to believe is experiencing or at risk of experiencing an opioid-related overdose;
(b) A family member, friend, or other person in a position to assist an individual who there is reason to believe is at risk of experiencing an opioid-related overdose.

(2) An individual authorized under this section to personally furnish naloxone may do so without having examined the individual to whom it may be administered.

(B) An individual authorized by a physician assistant under this section may personally furnish naloxone to an individual described in division (A)(1)(a) or (b) of this section if both of the following conditions are met:

(1) The authorized individual complies with the protocol established by the authorizing physician assistant, including having completed the training required by the protocol.
(2) The authorized individual instructs the individual to whom naloxone is furnished to summon emergency services as soon as practicable either before or after administering naloxone.

(C) A protocol established by a physician assistant for purposes of this section shall be established in writing and include all of the following:

(1) A description of the clinical pharmacology of naloxone;
(2) Precautions and contraindications concerning furnishing naloxone;
(3) Any limitations the physician assistant specifies concerning the individuals to whom naloxone may be furnished;
(4) The naloxone dosage that may be furnished and any variation in the dosage based on circumstances specified in the protocol;
(5) Labeling, storage, record keeping, and administrative requirements;
(6) Training requirements that must be met before an individual will be authorized to furnish naloxone;
(7) Any instructions or training that the authorized individual must provide to an individual to whom naloxone is furnished.
(D) A physician assistant who in good faith authorizes another individual to personally furnish naloxone in accordance with a protocol established by the physician assistant under this section is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is furnished: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

An individual authorized under this section to personally furnish naloxone who does so in good faith is not liable for or subject to any of the following for any action or omission of the individual to whom the naloxone is furnished: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

Sec. 4730.436. (A) As used in this section, "service entity" has the same meaning as in section 4729.514 of the Revised Code.

(B) A physician assistant who holds a valid prescriber number issued by the state medical board, who has been granted physician-delegated prescriptive authority, and who has established a protocol under division (D) of this section may authorize an individual who is an employee, volunteer, or contractor of a service entity to administer naloxone to an individual who is apparently experiencing an opioid-related overdose.

(C) An individual authorized by a physician assistant under this section may administer naloxone to an individual who is apparently experiencing an opioid-related overdose if all of the following conditions are met:

(1) The naloxone is obtained from a service entity of which the authorized individual is an employee, volunteer, or contractor.

(2) The authorized individual complies with the protocol established by the authorizing physician assistant.

(3) The authorized individual summons emergency services as soon as practicable either before or after administering the naloxone.

(D) A protocol established by a physician assistant for purposes of this section must be in writing and include all of the following:

(1) A description of the clinical pharmacology of naloxone;

(2) Precautions and contraindications concerning the administration of naloxone;

(3) Any limitations the physician assistant specifies concerning the individuals to whom naloxone may be administered;

(4) The naloxone dosage that may be administered and any variation in the dosage based on circumstances specified in the protocol;

(5) Labeling, storage, record keeping, and administrative requirements;

(6) Training requirements that must be met before an individual can be authorized to administer naloxone.

(E) A physician assistant who in good faith authorizes an individual to administer naloxone under this section is not liable for or subject to any of the following for any act or omission of the authorized individual: damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

A service entity or an employee, volunteer, or contractor of a service entity is not liable for or subject to any of the following for injury, death, or loss to person or property that allegedly arises
from an act or omission associated with procuring, maintaining, accessing, or administering naloxone under this section, unless the act or omission constitutes willful or wanton misconduct; damages in any civil action, prosecution in any criminal proceeding, or professional disciplinary action.

This section does not eliminate, limit, or reduce any other immunity or defense that a service entity or an employee, volunteer, or contractor of a service entity may be entitled to under Chapter 2305. or any other provision of the Revised Code or under the common law of this state.

Sec. 4730.56. (A) As used in this section:
(1) "Community addiction services provider" has the same meaning as in section 5119.01 of the Revised Code.
(2) "Medication-assisted treatment" has the same meaning as in section 340.01 of the Revised Code.

(B) A physician assistant shall comply with section 3719.064 of the Revised Code and rules adopted under section 4730.55 of the Revised Code when treating a patient with medication-assisted treatment or proposing to initiate such treatment.

(C) A physician assistant who fails to comply with this section shall treat not more than thirty patients at any one time with medication-assisted treatment even if the facility or location at which the treatment is provided is either of the following:
(1) Exempted by divisions (B)(2)(a) to (d) or (i) of section 4729.553 of the Revised Code from being required to possess a category III terminal distributor of dangerous drugs license with an office-based opioid treatment classification;
(2) A community addiction services provider that provides alcohol and drug addiction services that are certified by the department of mental health and addiction services under section 5119.36 of the Revised Code.

Sec. 4731.83. (A) As used in this section:
(1) "Medication-assisted treatment" has the same meaning as in section 340.01 of the Revised Code.
(2) "Physician" means an individual authorized by this chapter to practice medicine and surgery or osteopathic medicine and surgery.

(B) A physician shall comply with section 3719.064 of the Revised Code and rules adopted under section 4731.056 of the Revised Code when treating a patient with medication-assisted treatment or proposing to initiate such treatment.

(C) A physician who fails to comply with this section shall treat not more than thirty patients at any one time with medication-assisted treatment even if the facility or location at which the treatment is provided is either of the following:
(1) Exempted by divisions (B)(2)(a) to (d) or (i) of section 4729.553 of the Revised Code from being required to possess a category III terminal distributor of dangerous drugs license with an office-based opioid treatment classification;
(2) A community addiction services provider that provides alcohol and drug addiction services that are certified by the department of mental health and addiction services under section 5119.36 of the Revised Code.
SECTION 3. The General Assembly, applying the principle stated in division (B) of section 1.52 of the Revised Code that amendments are to be harmonized if reasonably capable of simultaneous operation, finds that the following sections, presented in this act as composites of the sections as amended by the acts indicated, are the resulting versions of the sections in effect prior to the effective date of the sections as presented in this act: Section 2925.61 of the Revised Code as amended by both H.B. 216 and S.B. 319 of the 131st General Assembly. Section 4729.553 of the Revised Code as amended by both H.B. 101 and S.B. 229 of the 132nd General Assembly. Section 4730.431 of the Revised Code as amended by both H.B. 4 and S.B. 110 of the 131st General Assembly.
Speaker ___________________ of the House of Representatives.

President ___________________ of the Senate.

Passed ________________________, 20____

Approved ________________________, 20____

Governor.
The section numbering of law of a general and permanent nature is complete and in conformity with the Revised Code.

________________________________________________________

Director, Legislative Service Commission.

Filed in the office of the Secretary of State at Columbus, Ohio, on the ____ day of __________, A. D. 20____.

________________________________________________________

Secretary of State.

File No. __________  Effective Date __________________________