
A BILL FOR AN ACT

RELATING TO HEALTH.

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF HAWAII:

1 SECTION 1. The legislature finds that the nationwide
2 opioid epidemic continues to result in an alarming number of
3 opioid overdose deaths. According to the Centers for Disease
4 Control and Prevention, opioid overdose fatalities have
5 increased from 33,000 in 2015 to 53,000 in 2016. Unintentional
6 drug poisonings, commonly referred to as drug overdoses, are one
7 of the leading causes of injury-related mortality in Hawaii.
8 Furthermore, an average of four hundred non-fatal overdoses
9 occur in Hawaii per year, and opioid related overdoses resulted
10 in about \$9,800,000 in hospital costs in 2016.

11 The legislature further finds that deaths caused by opioids
12 are often preventable via timely administration of an opioid
13 antagonist, such as naloxone. Studies have found that providing
14 opioid overdose training and naloxone kits can help people
15 identify signs of an opioid-related drug overdose and can help
16 reduce opioid overdose mortality. Thus, there is a need for
17 increased public access to health care professionals who can



1 safely provide naloxone and related education about the risks of
2 opioid misuse.

3 The legislature also finds that pharmacists are well
4 situated to provide education and access to naloxone and assist
5 with the prevention and health care burden of addressing opioid
6 overdose in Hawaii. A good example of how pharmacists can
7 positively impact the overall public health continuum and reduce
8 health care costs is seen with pharmacists providing
9 immunizations. Pharmacists now immunize more patients than any
10 other health care professionals, and immunization rates have
11 grown, reducing disease and morbidity in the overall population.

12 The legislature notes that there is significant precedent
13 in Hawaii law that supports expanded access to opioid
14 antagonists and the role of registered pharmacists in the
15 administration, dispensing, and prescription of opioid
16 antagonists, such as in Act 66, Session Laws of Hawaii 2017, Act
17 68, Session Laws of Hawaii 2016, and Act 217, Session Laws of
18 Hawaii 2015.

19 Accordingly, the purpose of this Act is to expand the scope
20 of registered pharmacists' practices by allowing registered
21 pharmacists to prescribe, dispense, and provide related



1 education of opioid antagonists without the need for a written,
2 approved collaborative agreement.

3 SECTION 2. Chapter 461, Hawaii Revised Statutes, is
4 amended by adding a new section to be appropriately designated
5 and to read as follows:

6 "§461- Opioid antagonist; authority to prescribe and
7 dispense; requirements. (a) A pharmacist may prescribe and
8 dispense an opioid antagonist to an individual who is at risk
9 for an opioid overdose or family member of an individual who is
10 at risk for an opioid overdose regardless of whether the
11 individual has evidence of a previous prescription for an opioid
12 antagonist from a practitioner authorized to prescribe opioids.
13 The opioid antagonist prescribed and dispensed for a family
14 member of an individual who is at risk for an opioid overdose
15 shall be prescribed and dispensed in the name of the individual
16 who is requesting the opioid antagonist.

17 (b) A pharmacist who prescribes and dispenses opioid
18 antagonists pursuant to subsection (a) shall:

19 (1) Complete a training program related to prescribing
20 opioid antagonists that is approved by the
21 Accreditation Council for Pharmacy Education (ACPE), a



curriculum-based program from an ACPE-accredited
college of pharmacy, a state or local health
department program, or a program recognized by the
board;

(2) Provide the individual who is receiving the opioid
antagonist with information and written educational
material on risk factors of opioid overdose, signs of
an overdose, overdose response steps, and the use of
the opioid antagonist;

(3) Obtain a signed acknowledgment by the individual
receiving the opioid antagonist. For opioid
antagonists that are prescribed to the individual at
risk for an opioid overdose, the practitioner who
authorized the original opioid prescription shall be
notified if applicable; and

(4) Dispense the opioid antagonist to the individual who
is at risk of an opioid overdose or family member as
soon as practicable after the pharmacist issues the
prescription.

(c) A pharmacist who prescribes an opioid antagonist
pursuant to subsection (a) shall not require the individual who



1 is at risk for an opioid overdose or family member to schedule
2 an appointment with the pharmacist for the prescribing or
3 dispensing of the opioid antagonist."

4 SECTION 3. Section 328-16, Hawaii Revised Statutes, is
5 amended as follows:

6 1. By amending subsections (a) to (c) to read:

7 "(a) A prescription drug shall be dispensed only if its
8 label bears the following:

9 (1) The name, business address, and telephone number of
10 the seller. The business address shall be the
11 physical location of the pharmacy or the dispensing
12 practitioner's office;

13 (2) Except as otherwise authorized for expedited partner
14 therapy in section 453-52[7] or an opioid antagonist
15 in section 461- , the name of the person for whom the
16 drug was prescribed or the name of the owner of the
17 animal for which the drug was prescribed;

18 (3) The serial number of the prescription;

19 (4) The date the prescription was prepared;

20 (5) The name of the practitioner if the seller is not the
21 practitioner;



- 1 (6) The name, strength, and quantity of the drug;
- 2 (7) The "use by" date for the drug, which shall be:
- 3 (A) The expiration date on the manufacturer's
- 4 container; or
- 5 (B) One year from the date the drug is dispensed,
- 6 whichever is earlier;
- 7 (8) The number of refills available, if any;
- 8 (9) In the case of the dispensing of an equivalent generic
- 9 drug product, the statement "same as (brand name of
- 10 the drug product prescribed or the referenced listed
- 11 drug name)", or words of similar meaning;
- 12 (10) In the case of the dispensing of an interchangeable
- 13 biological product, the statement "interchangeable
- 14 with (brand name of the biological product prescribed
- 15 or the referenced biological drug name)", or words of
- 16 similar meaning; and
- 17 (11) Specific directions for the drug's use; provided that
- 18 if the specific directions for use are too lengthy for
- 19 inclusion on the label, the notation "take according
- 20 to written instructions" may be used if separate
- 21 written instructions for use are actually issued with



1 the drug by the practitioner or the pharmacist, but in
2 no event shall the notation "take as directed",
3 referring to oral instructions, be considered
4 acceptable.

5 If any prescription for a drug does not indicate the number of
6 times it may be refilled, if any, the pharmacist shall not
7 refill that prescription unless subsequently authorized to do so
8 by the practitioner. The act of dispensing a prescription drug
9 other than a professional sample or medical oxygen contrary to
10 this subsection shall be deemed to be an act that results in a
11 drug being misbranded while held for sale.

12 (b) In addition to the requirements enumerated in
13 subsection (a), a prescription drug shall be dispensed only:

14 (1) By a pharmacist pursuant to a valid prescription[
15 ~~section 461-1, or section 453-52,~~], or section 453-52,
16 section 461-1, or section 461- ;

17 (2) By a medical oxygen distributor pursuant to a
18 prescription or certificate of medical necessity;
19 provided that the drug to be dispensed is medical
20 oxygen; or

21 (3) By a practitioner to an ultimate user; provided that:



1 (A) Except as otherwise authorized for expedited
2 partner therapy in section 453-52, the
3 practitioner shall inform the patient, prior to
4 dispensing any drug other than a professional
5 sample, that the patient may have a written,
6 orally ordered, or electronically transmitted or
7 conveyed prescription directed to a pharmacy or a
8 medical oxygen distributor of the patient's own
9 choice;

10 (B) The practitioner shall promptly record in the
11 practitioner's records:

12 (i) The prescription in full;

13 (ii) The name, strength, and quantity of the
14 drug, and specific directions for the drug's
15 use;

16 (iii) The date the drug was dispensed;

17 (iv) Except as otherwise authorized for expedited
18 partner therapy in section 453-52 [7] or for
19 an opioid antagonist in section 461- , the
20 name and address of the person for whom the
21 drug was prescribed or the name of the owner



1 of the animal for which the drug was
2 prescribed; and

3 (v) Prescription drugs dispensed or prescribed
4 for expedited partner therapy as authorized
5 under section 453-52[+] or for an opioid
6 antagonist in section 461- ;

7 (C) The records described in subparagraph (B) shall
8 be subject to the inspection of the department or
9 its agents at all times; and

10 (D) No undisclosed rebate, refund, commission,
11 preference, discount, or other consideration,
12 whether in the form of money or otherwise, has
13 been offered to the practitioner as compensation
14 or inducement to dispense or prescribe any
15 specific drug in preference to other drugs that
16 might be used for the identical therapeutic
17 indication.

18 (c) A prescription may be communicated in writing, orally,
19 or by electronic transmission, and shall include the following
20 information:



1 (1) The authorization of the practitioner noted as
2 follows:

3 (A) Written prescriptions shall include the original
4 signature of the practitioner;

5 (B) Oral prescriptions shall be promptly recorded by
6 the pharmacist or medical oxygen distributor and
7 shall include the practitioner's oral code
8 designation; and

9 (C) Electronic prescriptions shall be irrefutably
10 traceable to the prescribing practitioner by a
11 recognizable and unique practitioner identifier
12 such as:

13 (i) A bitmap or graphic image of the
14 prescriber's handwritten signature and the
15 prescriber's oral code designation (or
16 license number or other identifier if the
17 prescriber is an out-of-state practitioner);

18 (ii) An electronic signature;

19 (iii) A digital signature; or

20 (iv) By other means as approved by the director;

21 (2) The date of issuance;



- 1 (3) The practitioner's name, business telephone number,
2 and business address, unless the practitioner is
3 otherwise uniquely identified and the pharmacy or
4 medical oxygen distributor dispensing the prescription
5 has the prescriber's contact information on file
6 accessible within the dispensing area;
- 7 (4) The name, strength, and quantity of the drug to be
8 dispensed, and specific directions for the drug's use;
- 9 (5) Except as otherwise authorized for expedited partner
10 therapy in section 453-52[7] or for an opioid
11 antagonist in section 461- , the name and address of
12 the person for whom the prescription was written or
13 the name of the owner of the animal for which the drug
14 was prescribed, unless the pharmacy or medical oxygen
15 distributor dispensing the prescription has the
16 address on file accessible within the dispensing area;
- 17 (6) The room number and route of administration, if the
18 patient is in an institutional facility; and
- 19 (7) The number of allowable refills, if the prescription
20 is refillable. If the number of refills authorized by
21 the practitioner is indicated using the terms "as



1 needed" or "prn", the prescription may be refilled up
2 to twelve months from the date the original
3 prescription was written. After the twelve-month
4 period, the "as needed" or "prn" prescription may be
5 refilled for a subsequent three-month period;
6 provided:

7 (A) The prescription is refilled only once during the
8 three-month period;

9 (B) The refill does not exceed a thirty-day supply of
10 the drug;

11 (C) The refill does not provide any amount of the
12 drug fifteen months beyond the date the original
13 prescription was written;

14 (D) In the case of medical oxygen, the duration of
15 therapy indicated on a certificate of medical
16 necessity shall supersede any limitations or
17 restrictions on refilling; and

18 (E) Subparagraphs (A) to (D) shall apply only to
19 pharmacies and medical oxygen distributors
20 practicing in the State."

21 2. By amending subsection (g) to read:



"(g) Any drug other than medical oxygen dispensed pursuant to a prescription shall be exempt from the requirements of section 328-15 (except paragraphs (1), (9), (11), and (12), and the packaging requirements of paragraphs (7) and (8)), if the drug bears a label containing:

(1) The name and address of the pharmacy;

(2) The serial number and the date of the prescription or of its filling;

(3) The name of the practitioner;

(4) Except as otherwise authorized for expedited partner therapy in section 453-52[7] or for an opioid antagonist in section 461- , the name of the patient;

(5) The directions for use; and

(6) Any cautionary statements contained in the prescription.

This exemption shall not apply to any drug dispensed in the course of the conduct of a business of dispensing drugs pursuant to diagnosis by mail, or to a drug dispensed in violation of subsection (a), (b), (c), or (d)."

SECTION 4. Section 328-17.6, Hawaii Revised Statutes, is amended as follows:



1 1. By amending subsections (c) and (d) to read:

2 "(c) Any pharmacist or medical oxygen distributor who
3 fills or refills a prescription from an out-of-state
4 practitioner shall:

5 (1) Note the following on the prescription record: the
6 out-of-state practitioner's full name, address, and
7 telephone number;

8 (2) Be responsible for validating and verifying the
9 practitioner's prescriptive authority by virtue of a
10 valid out-of-state license, a Drug Enforcement
11 Administration registration number, or other measures
12 as appropriate; and

13 (3) Except as otherwise authorized for expedited partner
14 therapy in section 453-52[7] or for an opioid
15 antagonist in section 461- , demand proper
16 identification from the person whose name appears on
17 the prescription prior to filling the prescription, in
18 addition to complying with any identification
19 procedures established by the department for filling
20 and refilling an out-of-state prescription.



(d) Before refilling a transferred out-of-state prescription, a pharmacist or medical oxygen distributor shall:

(1) Except as otherwise authorized for expedited partner therapy in section 453-52[7] or for an opioid antagonist in section 461- , advise the person whose name appears on the prescription that the prescription on file at the originating out-of-state pharmacy or medical oxygen distributor may be canceled; and

(2) Record all information required to be on a prescription, including:

(A) The date of issuance of the original prescription;

(B) The number of refills authorized on the original prescription;

(C) The date the original prescription was dispensed;

(D) The number of valid refills remaining and the date of the last refill;

(E) The out-of-state pharmacy's or out-of-state medical oxygen distributor's name, telephone number, and address, and the original



1 prescription number or control number from which
2 the prescription information was transferred; and
3 (F) The name of the transferor pharmacist or the
4 medical oxygen distributor's agent."

5 2. By amending subsection (f) to read:

6 "(f) An out-of-state prescription record shall state the
7 date of filling or refilling and, except as otherwise authorized
8 for expedited partner therapy in section 453-52[7] or for an
9 opioid antagonist in section 461- , the local address of the
10 person whose name appears on the prescription."

11 SECTION 5. Section 328-17.7, Hawaii Revised Statutes, is
12 amended by amending subsection (a) to read as follows:

13 "(a) Every practitioner, pharmacist, or medical oxygen
14 distributor who compounds, sells, or delivers any prescribed
15 drug to a patient or a patient's agent shall maintain records
16 that identify:

17 (1) The specific drug product dispensed, including:

18 (A) The product's national drug code (NDC) number; or

19 (B) The brand name or the established name and the
20 name or commonly accepted abbreviation of the



- 1 principal labeler of the drug product dispensed,
2 the product strength, and the dosage form;
- 3 (2) The quantity of the drug;
- 4 (3) Directions for use;
- 5 (4) The number of allowable refills;
- 6 (5) The date of initial dispensing and the dates of all
7 refilling;
- 8 (6) The date of any transfer of the prescription;
- 9 (7) The name, business address, and telephone number of
10 the recipient pharmacist or medical oxygen distributor
11 for any transfer of prescription;
- 12 (8) The prescribing practitioner, including name, business
13 address, and telephone number;
- 14 (9) The format (oral, written, or electronic) in which the
15 prescription was received;
- 16 (10) Except as otherwise authorized for expedited partner
17 therapy in section 453-52[7] or for an opioid
18 antagonist in section 461- , the patient, including
19 name, address, and telephone number;
- 20 (11) The date of prescribing; and



1 (12) The name of the practitioner, pharmacist, or medical
2 oxygen distributor dispensing the drug.
3 Every prescription dispensed shall have the name of the
4 pharmacist, dispensing practitioner, or medical oxygen
5 distributor responsible for the dispensing appended to the
6 prescription record, and every prescription record shall be
7 preserved and legible for a period of not less than five years.
8 The prescription records shall be subject at all times to the
9 inspection of the director of health or the director's agent."

10 SECTION 6. Section 461-1, Hawaii Revised Statutes, is
11 amended as follows:

12 1. By adding a new definition to be appropriately inserted
13 and to read:

14 "Family member" means an individual who can provide
15 assistance and is related to the individual at risk for an
16 opioid overdose."

17 2. By amending the definition of "practice of pharmacy" to
18 read:

19 "Practice of pharmacy" means:

20 (1) The interpretation and evaluation of prescription
21 orders; the compounding, dispensing, and labeling of



1 drugs and devices (except labeling by a manufacturer,
2 packer, or distributor of nonprescription drugs and
3 commercially legend drugs and devices); the
4 participation in drug selection and drug utilization
5 reviews; the proper and safe storage of drugs and
6 devices and the maintenance of proper records
7 therefor; the responsibility for advising when
8 necessary or where regulated, of therapeutic values,
9 content, hazards, and use of drugs and devices;

- 10 (2) Performing the following procedures or functions as
11 part of the care provided by and in concurrence with a
12 "health care facility" and "health care service" as
13 defined in section 323D-2, or a "pharmacy" or a
14 licensed physician or a licensed advanced practice
15 registered nurse with prescriptive authority, or a
16 "managed care plan" as defined in section 432E-1, in
17 accordance with policies, procedures, or protocols
18 developed collaboratively by health professionals,
19 including physicians and surgeons, pharmacists, and
20 registered nurses, and for which a pharmacist has



1 received appropriate training required by these
2 policies, procedures, or protocols:

3 (A) Ordering or performing routine drug therapy
4 related patient assessment procedures;

5 (B) Ordering drug therapy related laboratory tests;

6 (C) Initiating emergency contraception oral drug
7 therapy in accordance with a written
8 collaborative agreement approved by the board,
9 between a licensed physician or advanced practice
10 registered nurse with prescriptive authority and
11 a pharmacist who has received appropriate
12 training that includes programs approved by the
13 American Council of Pharmaceutical Education
14 (ACPE), curriculum-based programs from an ACPE-
15 accredited college of pharmacy, state or local
16 health department programs, or programs
17 recognized by the board of pharmacy;

18 (D) Administering drugs orally, topically, by
19 intranasal delivery, or by injection, pursuant to
20 the order of the patient's licensed physician or
21 advanced practice registered nurse with



1 prescriptive authority, by a pharmacist having
2 appropriate training that includes programs
3 approved by the ACPE, curriculum-based programs
4 from an ACPE-accredited college of pharmacy,
5 state or local health department programs, or
6 programs recognized by the board of pharmacy;

7 (E) Administering:

8 (i) Immunizations orally, by injection, or by
9 intranasal delivery, to persons eighteen
10 years of age or older by a pharmacist having
11 appropriate training that includes programs
12 approved by the ACPE, curriculum-based
13 programs from an ACPE-accredited college of
14 pharmacy, state or local health department
15 programs, or programs recognized by the
16 board of pharmacy;

17 (ii) Vaccines to persons between fourteen and
18 seventeen years of age pursuant to section
19 461-11.4; and

20 (iii) Human papillomavirus, Tdap (tetanus,
21 diphtheria, pertussis), meningococcal, and



1 influenza vaccines to persons between eleven
2 and seventeen years of age pursuant to
3 section 461-11.4;

4 (F) As authorized by the written instructions of a
5 licensed physician or advanced practice
6 registered nurse with prescriptive authority,
7 initiating or adjusting the drug regimen of a
8 patient pursuant to an order or authorization
9 made by the patient's licensed physician or
10 advanced practice registered nurse with
11 prescriptive authority and related to the
12 condition for which the patient has been seen by
13 the licensed physician or advanced practice
14 registered nurse with prescriptive authority;
15 provided that the pharmacist shall issue written
16 notification to the patient's licensed physician
17 or advanced practice registered nurse with
18 prescriptive authority or enter the appropriate
19 information in an electronic patient record
20 system shared by the licensed physician or



1 advanced practice registered nurse with
2 prescriptive authority, within twenty-four hours;

3 (G) Transmitting a valid prescription to another
4 pharmacist for the purpose of filling or
5 dispensing;

6 (H) Providing consultation, information, or education
7 to patients and health care professionals based
8 on the pharmacist's training and for which no
9 other licensure is required; or

10 (I) ~~[Dispensing an opioid antagonist in accordance~~
11 ~~with a written collaborative agreement approved~~
12 ~~by the board, between a licensed physician and a~~
13 ~~pharmacist who has received appropriate training~~
14 ~~that includes programs approved by the ACPE,~~
15 ~~curriculum-based programs from an ACPE accredited~~
16 ~~college of pharmacy, state or local health~~
17 ~~department programs, or programs recognized by~~
18 ~~the board;]~~ Prescribing and dispensing an opioid
19 antagonist pursuant to section 461- ;



- 1 (3) The offering or performing of those acts, services,
2 operations, or transactions necessary in the conduct,
3 operation, management, and control of pharmacy; and
4 (4) Prescribing and dispensing contraceptive supplies
5 pursuant to section 461-11.6."

6 SECTION 7. Statutory material to be repealed is bracketed
7 and stricken. New statutory material is underscored.

8 SECTION 8. This Act shall take effect on July 1, 3000.



Report Title:

Pharmacists; Opioid Antagonist; Prescription

Description:

Authorizes a pharmacist to prescribe and dispense opioid antagonists without a written collaborative agreement; provided that the pharmacist meets certain qualification requirements and the individual receiving the prescription receives opioid antagonist education and signs an acknowledgement. (HB1924 HD1)

The summary description of legislation appearing on this page is for informational purposes only and is not legislation or evidence of legislative intent.

