# 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 a nationwide drug
2	epidemic exists related to prescription pain relieving drugs
3	that are causing alarming rates of addiction, overdose, and
4	death. According to the National Institute on Drug Abuse, an
5	estimated 2,100,000 people in the United States suffer from
6	substance use disorders related to prescription opioid pain
7	relievers. Society is facing the devastating consequences of
8	this epidemic. The number of unintentional overdose deaths from
9	prescription pain relievers has more than quadrupled since 1999
10	According to data provided by the Pew Charitable Trusts, opioid
11	pain relievers killed nearly 20,000 Americans in 2014.
12	According to the National Institute on Drug Abuse, in terms
13	of abuse and mortality, opioids account for the greatest
14	proportion of the prescription drug abuse problem. The rise of
15	prescription opioids started in the beginning of the twenty-
16	first century and by 2002 prescription opioids caused more
17	deaths than heroin or cocaine. The National Institute on Drug

Abuse reports that the increase in the availability of opioid

- 1 pain relievers is the result of a drastic increase in the number
- 2 of prescriptions written and dispensed, greater social
- 3 acceptability for using medications for different purposes, and
- 4 aggressive marketing by pharmaceutical companies. As a result
- 5 of the staggering number of people suffering from substance use
- 6 disorders related to prescription opioid pain relievers, the
- 7 United States Centers for Disease Control and Prevention,
- 8 national and state legislators, and many others are trying to
- 9 curb this epidemic through public education and limits on opioid
- 10 prescribing practices.
- 11 The legislature also finds that informed consent is an
- 12 effective process between a provider and patient that relates to
- 13 a specific medication or a form of treatment such as safe opioid
- 14 therapy. The informed consent process allows the patient to
- 15 better understand the goals of treatment, potential benefits of
- 16 treatment, realistic outcomes, potential risks, how to use the
- 17 medication, and alternative treatment options. The informed
- 18 consent process is one approach to begin addressing the
- 19 nationwide opioid epidemic.
- The purpose of this Act is to reduce addiction, overdose,
- 21 and death related to the use of opioids by:

1	(1)	Requiring execution of an opioid therapy informed
2		consent process agreement between a patient and a
3		prescriber of opioids in circumstances that may carry
4		an elevated risk of causing dependency; and
5	(2)	Limiting initial concurrent prescriptions for opioids
6		and benzodiazepines to a maximum of seven consecutive
7		days, except for treatment of specified conditions.
8	SECT	ION 2. Chapter 329, Hawaii Revised Statutes, is
9	amended b	y adding a new section to be appropriately designated
10	and to re	ad as follows:
11	" <u>§</u> 32	9- Opioid therapy; informed consent process;
11 12		9- Opioid therapy; informed consent process; nt for written policies. (a) Beginning on July 1,
	requireme	
12	requireme	nt for written policies. (a) Beginning on July 1,
12 13	requireme 2018, any and maint	nt for written policies. (a) Beginning on July 1, provider authorized to prescribe opioids shall adopt
12 13 14	requireme 2018, any and maint of a writ	nt for written policies. (a) Beginning on July 1, provider authorized to prescribe opioids shall adopt ain written policy or policies that include execution
12 13 14 15	requireme 2018, any and maint of a writ	nt for written policies. (a) Beginning on July 1,  provider authorized to prescribe opioids shall adopt  ain written policy or policies that include execution  ten agreement to engage in an informed consent process
12 13 14 15 16	requireme 2018, any and maint of a writ between t	nt for written policies. (a) Beginning on July 1,  provider authorized to prescribe opioids shall adopt  ain written policy or policies that include execution  ten agreement to engage in an informed consent process
12 13 14 15 16 17	requireme  2018, any and maint of a writ between t  patient. (b)	nt for written policies. (a) Beginning on July 1,  provider authorized to prescribe opioids shall adopt ain written policy or policies that include execution ten agreement to engage in an informed consent process he prescribing provider and qualifying opioid therapy

1	posted to the department of health's website no later than
2	December 31, 2017.
3	(c) For the purposes of this section, "qualifying opioid
4	therapy patient" means:
5	(1) A patient requiring opioid treatment for more than
6	three months;
7	(2) A patient who is prescribed benzodiazepines and
8	opioids together; or
9	(3) A patient who is prescribed a dose of opioids that
10	exceeds ninety morphine equivalent doses.
11	(d) A violation of this section shall not be subject to
12	the penalty provisions of part IV of chapter 329."
13	SECTION 3. Section 329-38, Hawaii Revised Statutes, is
14	amended to read as follows:
15	"§329-38 Prescriptions. (a) No controlled substance in
16	schedule II may be dispensed without a written prescription of
17	practitioner, except:
18	(1) In the case of an emergency situation, a pharmacist
19	may dispense a controlled substance listed in schedul
20	II upon receiving oral authorization from a
21	prescribing practitioner; provided that:

1	(A)	The quantity prescribed and dispensed is limited
2		to the amount adequate to treat the patient
3		during the emergency period (dispensing beyond
4		the emergency period shall be pursuant to a
5		written prescription signed by the prescribing
6		<pre>practitioner);</pre>
7	(B)	If the prescribing practitioner is not known to
8		the pharmacist, the pharmacist shall make a
9		reasonable effort to determine that the oral
10		authorization came from a registered
11		practitioner, which may include a callback to the
12		prescribing practitioner using the phone number
13		in the telephone directory or other good faith
14		efforts to identify the prescriber; and
15	(C)	Within seven days after authorizing an emergency
16		oral prescription, the prescribing practitioner
17		shall cause a written prescription for the
18		emergency quantity prescribed to be delivered to
19		the dispensing pharmacist. In addition to
20		conforming to the requirements of this
21		subsection, the prescription shall have written

1		on its face "Authorization for Emergency
2		Dispensing". The written prescription may be
3		delivered to the pharmacist in person or by mail,
4		and if by mail, the prescription shall be
5		postmarked within the seven-day period. Upon
6		receipt, the dispensing pharmacist shall attach
7		this prescription to the oral emergency
8		prescription, which had earlier been reduced to
9		writing. The pharmacist shall notify the
10		administrator if the prescribing practitioner
11		fails to deliver a written prescription to the
12		pharmacy within the allotted time. Failure of
13		the pharmacist to do so shall void the authority
14		conferred by this paragraph to dispense without a
15		written prescription of a prescribing individual
16		practitioner. Any practitioner who fails to
17		deliver a written prescription within the seven-
18		day period shall be in violation of section 329-
19		41(a)(1);
20	(2)	No schedule II narcotic controlled substance may be
21		prescribed or dispensed for more than a thirty-day

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1		supply, except where such substances come in a single
2		unit dose package that exceeds the thirty-day limit or
3		where a terminally ill patient is certified by a
4		physician to exceed the thirty-day limit;
5	(3)	When dispensed directly by a practitioner, other than
6		a pharmacist, to the ultimate user. The practitioner
7		in dispensing a controlled substance in schedule II
8		shall affix to the package a label showing:
9		(A) The date of dispensing;
10		(B) The name, strength, and quantity of the drug
11		dispensed;
12		(C) The dispensing practitioner's name and address;
13		(D) The name of the patient;
14		(E) The "use by" date for the drug, which shall be:
15		(i) The expiration date on the manufacturer's or
16	•	principal labeler's container; or
17		(ii) One year from the date the drug is
18		dispensed, whichever is earlier; and
19		(F) Directions for use, and cautionary statements, if
20		any, contained in the prescription or as required
21		by law.

1		A complete and accurate record of all schedule II
2		controlled substances ordered, administered,
3		prescribed, and dispensed shall be maintained for five
4		years. Prescriptions and records of dispensing shall
5		otherwise be retained in conformance with the
6		requirements of section 329-36. No prescription for a
7		controlled substance in schedule II may be refilled;
8		or
9	(4)	In the case of an electronic prescription, a
10		pharmacist may dispense a controlled substance listed
11		in schedule II upon receiving an electronic
12		prescription.
13	(b)	A schedule II controlled substance prescription shall:
14	(1)	Be filled within seven days following the date the
15		prescription was issued to the patient; and
16	(2)	Be supplied to a patient only if the prescription has
17		been filled and held by the pharmacy for not more than
18		seven days.
19	<u>(c)</u>	Initial concurrent prescriptions for opioids and
20	benzodiaz	epines shall not be for longer than seven consecutive

1	days unle	ess a supply of longer than seven days is determined to
2	be medica	lly necessary for the treatment of:
3	(1)	Pain experienced while the patient is in post-
4		operative care;
5	(2)	Chronic pain and pain management;
6	(3)	Substance abuse or opioid or opiate dependence;
7	(4)	Cancer;
8	(5)	Pain experienced while the patient is in palliative
9		care; or
10	(6)	Pain experienced while the patient is in hospice care;
11	provided	that if a prescribing practitioner issues a concurrent
12	prescript	ion for more than a seven-day supply of an opioid and
13	benzodiaz	epine, the practitioner shall document in the patient's
14	medical r	ecord the condition for which the practitioner issued
15	the presc	ription and that an alternative to the opioid and
16	benzodiaz	epine was not appropriate treatment for the condition.
17	<u>(d)</u>	After an initial concurrent prescription for opioids
18	and benzo	diazepines has been made, a prescribing practitioner
19	may autho	rize subsequent prescriptions through a telephone
20	consultat	ion with the patient when the prescribing practitioner
21	deems suc	h action to be medically necessary for post-operative

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1	and pain management patients; provided that a prescribing				
2	practitioner shall consult with a patient in person at least				
3	once every ninety days for the duration during which the				
4	practitioner concurrently prescribes opioids and benzodiazepines				
5	to the patient.				
6	$[rac{\langle e \rangle}{}]$ (e) The transfer of original prescription information				
7	for a controlled substance listed in schedule III, IV, or V for				
8	the purpose of dispensing is permissible between pharmacies on a				
9	one time basis only. However, pharmacies electronically sharing				
10	a real-time, online database may transfer up to the maximum				
11	refills permitted by law and the prescriber's authorization.				
12	Transfers are subject to the following requirements:				
13	(1) The transfer shall be communicated directly between				
14	two licensed pharmacists, and the transferring				
15	pharmacist shall:				
16	(A) Write or otherwise place the word "VOID" on the				
17	face of the invalidated prescription;				
18	(B) Record on the reverse of the invalidated				
19	prescription the name, address, and Drug				
20	Enforcement Administration registration number of				

the pharmacy to which it was transferred and the

1		name	of the pharmacist receiving the prescription
2		info	rmation; and
3		(C) Reco	rd the date of the transfer and the name of
4		the	pharmacist transferring the information;
5	(2)	The pharm	acist receiving the transferred prescription
6		informati	on shall reduce to writing the following:
7		(A) Writ	e or otherwise place the word "transfer" on
8		the	face of the transferred prescription;
9		(B) Reco	rd all information required to be on a
10		pres	cription, including:
11		(i)	The date of issuance of original
12			prescription;
13		(ii)	The original number of refills authorized on
14			original prescription;
15		(iii)	The date of original dispensing;
16		(iv)	The number of valid refills remaining and
17			dates and locations of previous refills;
18		(v)	The pharmacy's name, address, Drug
19			Enforcement Administration registration
20			number, and original prescription number

1			from which the prescription information was
2			transferred;
3		(vi)	The name of the transferor pharmacist; and
4		(vii)	The pharmacy's name, address, and Drug
5			Enforcement Administration registration
6			number, along with the prescription number
7			from which the prescription was originally
8			filled;
9	(3)	Both the	original and transferred prescription shall
10		be mainta	ined for a period of five years from the date
11		of last r	efill; and
12	(4)	Any pharm	acy electronically accessing a prescription
13		record sh	all satisfy all information requirements of a
14		manual mo	de prescription transferal.
15	Fail	ure to com	ply with this subsection shall void the
16	authority	of the ph	armacy to transfer prescriptions or receive a
17	transferre	ed prescri	ption to or from another pharmacy.
18	[ <del>-(d)</del> -]	(f) A p	harmacy and an authorized central fill
19	pharmacy m	may share	information for initial and refill
20	prescript	ions of sc	hedule III, IV, or V controlled substances.
21	The follow	wing requi	rements shall apply:

1	(1)	A pharmacy may electronically transmit, including by		
2		facsimile, prescriptions for controlled substances		
3		listed in schedule III, IV, or V to a central fill		
4		pharmacy. The pharmacy transmitting the prescription		
5		information shall:		
6		(A) Ensure that all information required to be on a		
7		prescription pursuant to subsection $\left[\frac{g}{g}\right]$ (i) is		
8		transmitted to the central fill pharmacy either		
9		on the face of the prescription or		
10		electronically; and		
11		(B) Keep a record of receipt of the filled		
12		prescription, including the date of receipt, the		
13		method of delivery (private, common, or contract		
14		carrier) and the identity of the pharmacy		
15		employee accepting delivery; and		
16	(2)	The central fill pharmacy receiving the transmitted		
17		prescription shall:		
18		(A) Keep for five years a copy of a prescription		
19		received by facsimile or an electronic record of		
20		all the information transmitted by the pharmacy,		

including the name, address, and Drug Enforcement

1		Administration registration number of the
2		pharmacy transmitting the prescription;
3	(B)	Keep a record of the date of receipt of the
4		transmitted prescription, the name of the
5		licensed pharmacists filling the prescription,
6		and the dates the prescription was filled or is
7		refilled; and
8	(C)	Keep a record of the date the filled prescription
9		was shipped to the pharmacy.
10	[ <del>(e)</del> ] <u>(g)</u>	No controlled substance in schedule III, IV, or
11	V may be dispen	sed without a written, facsimile of a written,
12	oral prescripti	on of a practitioner, or receipt of an electronic
13	prescription, e	xcept when a controlled substance is dispensed
14	directly by a p	ractitioner, other than a pharmacist, to an
15	ultimate user.	The practitioner, in dispensing a controlled
16	substance in sc	hedule III, IV, or V, shall affix to the package
17	a label showing	:
18	(1) The d	ate of dispensing;
19	(2) The n	ame, strength, and quantity issued of the drug;
20	(3) The d	ispensing practitioner's name and business
21	addre	ess;

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1 (4)The name of the patient; 2 (5) The "use by" date for the drug, which shall be: 3 (A) The expiration date on the manufacturer's or 4 principal labeler's container; or 5 (B) One year from the date the drug is dispensed, 6 whichever is earlier; 7 Directions for use; and (6) 8 Cautionary statements, if any, contained in the (7)9 prescription or as required by law. 10 A complete and accurate record of all schedule III, IV, and V 11 controlled substances administered, prescribed, and dispensed 12 shall be maintained for five years. Prescriptions and records 13 of dispensing shall be retained in conformance with the 14 requirements of section 329-36 unless otherwise provided by law. 15 Prescriptions may not be filled or refilled more than three 16 months after the date of the prescription or be refilled more 17 than two times after the date of the prescription, unless the 18 prescription is renewed by the practitioner. 19  $\left[\frac{f}{f}\right]$  (h) The effectiveness of a prescription for the 20 purposes of this section shall be determined as follows:

1	(1)	A prescription for a controlled substance shall be
2		issued for a legitimate medical purpose by an
3		individual practitioner acting in the usual course of
4		the practitioner's professional practice. The
5		responsibility for the proper prescribing and
6		dispensing of controlled substances shall be upon the
7		prescribing practitioner, but a corresponding
8		responsibility shall rest with the pharmacist who
9		fills the prescription. An order purporting to be a
10		prescription issued not in the usual course of
11		professional treatment or for legitimate and
12		authorized research shall not be deemed a prescription
13		within the meaning and intent of this section, and the
14		person who knowingly fills such a purported
15		prescription, as well as the person who issues the
16		prescription, shall be subject to the penalties
17		provided for violations of this chapter;
18	(2)	A prescription may not be issued to allow an
19		individual practitioner to obtain controlled
20		substances for supplying the individual practitioner
21		for the purpose of general dispensing to patients;

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(3)	A prescription may not be issued for the dispensing of
	narcotic drugs listed in any schedule for the purpose
	of "medically managed withdrawal", also known as
	"detoxification treatment", or "maintenance treatment"
	except as follows:

(A) The administering or dispensing directly (but not prescribing) of narcotic drugs listed in any schedule to a narcotic drug-dependent person for "medically managed withdrawal", also known as "detoxification treatment" $_{\underline{\prime}}$  or "maintenance treatment" shall be deemed to be "in the course of a practitioner's professional practice or research" so long as the practitioner is registered separately with the department and the federal Drug Enforcement Agency as required by section 329-32(e) and complies with Title 21 Code of Federal Regulations section 823(g) and any other federal or state regulatory standards relating to treatment qualification, security, records, and unsupervised use of drugs; and

1		(B) Nothing in this section shall prohibit a
2		physician or authorized hospital staff from
3		administering or dispensing, but not prescribing,
4	•	narcotic drugs in a hospital to maintain or
5		detoxify a person as an incidental adjunct to
6		medical or surgical treatment of conditions other
7		than addiction;
8	(4)	An individual practitioner shall not prescribe or
9		dispense a substance included in schedule II, III, IV,
10		or V for that individual practitioner's personal use,
11		except in a medical emergency; and
12	(5)	A pharmacist shall not dispense a substance included
13		in schedule II, III, IV, or V for the pharmacist's
14		personal use.
15	[ <del>-(g)</del>	(i) Prescriptions for controlled substances shall be
16	issued on	ly as follows:
17	(1)	All prescriptions for controlled substances shall
18		originate from within the State and be dated as of,
19		and signed on, the day when the prescriptions were
20		issued and shall contain:

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- (A) The first and last name and address of the patient; and
- (B) The drug name, strength, dosage form, quantity prescribed, and directions for use. Where a prescription is for gamma hydroxybutyric acid, methadone, or buprenorphine, the practitioner shall record as part of the directions for use, the medical need of the patient for the prescription.

Except for electronic prescriptions, controlled substance prescriptions shall be no larger than eight and one-half inches by eleven inches and no smaller than three inches by four inches. A practitioner may sign a prescription in the same manner as the practitioner would sign a check or legal document (e.g., J.H. Smith or John H. Smith) and shall use both words and figures (e.g., alphabetically and numerically as indications of quantity, such as five (5)), to indicate the amount of controlled substance to be dispensed. Where an oral order or electronic prescription is not permitted, prescriptions shall be

1	written with ink or indelible pencil or typed, shall
2	be manually signed by the practitioner, and shall
3	include the name, address, telephone number, and
4	registration number of the practitioner. The
5	prescriptions may be prepared by a secretary or agent
6	for the signature of the practitioner, but the
7	prescribing practitioner shall be responsible in case
8	the prescription does not conform in all essential
9	respects to this chapter and any rules adopted
10	pursuant to this chapter. In receiving an oral
11	prescription from a practitioner, a pharmacist shall
12	promptly reduce the oral prescription to writing,
13	which shall include the following information: the
14	drug name, strength, dosage form, quantity prescribed
15	in figures only, and directions for use; the date the
16	oral prescription was received; the full name, Drug
17	Enforcement Administration registration number, and
18	oral code number of the practitioner; and the name and
19	address of the person for whom the controlled
20	substance was prescribed or the name of the owner of

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the animal for which the controlled substance was prescribed.

A corresponding liability shall rest upon a pharmacist who fills a prescription not prepared in the form prescribed by this section. A pharmacist may add a patient's missing address or change a patient's address on all controlled substance prescriptions after verifying the patient's identification and noting the identification number on the back of the prescription document on file. The pharmacist shall not make changes to the patient's name, the controlled substance being prescribed, the quantity of the prescription, the practitioner's Drug Enforcement Administration number, the practitioner's name, the practitioner's electronic signature, or the practitioner's signature;

(2) An intern, resident, or foreign-trained physician, or a physician on the staff of a Department of Veterans

Affairs facility or other facility serving veterans, exempted from registration under this chapter, shall include on all prescriptions issued by the physician:

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1	(A) The registration number of the hospital or other
2	institution; and
3	(B) The special internal code number assigned to the
4	physician by the hospital or other institution in
5	lieu of the registration number of the
6	practitioner required by this section.
7	The hospital or other institution shall forward a copy
8	of this special internal code number list to the
9	department as often as necessary to update the
10	department with any additions or deletions. Failure
11	to comply with this paragraph shall result in the
12	suspension of that facility's privilege to fill
13	controlled substance prescriptions at pharmacies
14	outside of the hospital or other institution. Each
15	written prescription shall have the name of the
16	physician stamped, typed, or hand-printed on it, as
17	well as the signature of the physician;
18 (3)	An official exempted from registration shall include
19	on all prescriptions issued by the official:
20	(A) The official's branch of service or agency (e.g.,
21	"U.S. Army" or "Public Health Service"); and

1		(B) The official's service identification number, in
2		lieu of the registration number of the
3		practitioner required by this section. The
4		service identification number for a Public Health
5		Service employee shall be the employee's social
6		security or other government issued
7		identification number.
8		Each prescription shall have the name of the officer
9		stamped, typed, or handprinted on it, as well as the
10		signature of the officer; and
11	(4)	A physician assistant registered to prescribe
12		controlled substances under the authorization of a
13		supervising physician shall include on all controlled
14		substance prescriptions issued:
15		(A) The Drug Enforcement Administration registration
16		number of the supervising physician; and
17		(B) The Drug Enforcement Administration registration
18		number of the physician assistant.
19		Each written controlled substance prescription issued
20		shall include the printed, stamped, typed, or hand-

printed name, address, and phone number of both the

1	supervising physician and physician assistant, and
2	shall be signed by the physician assistant. The
3	medical record of each written controlled substance
4	prescription issued by a physician assistant shall be
5	reviewed and initialed by the physician assistant's
6	supervising physician within seven working days.
7	[ <del>(h)</del> ] <u>(j)</u> A prescription for controlled substances may
8	only be filled by a pharmacist acting in the usual course of the
9	pharmacist's professional practice and either registered
10	individually or employed in a registered pharmacy, central fill
11	pharmacy, or registered institutional practitioner. A central
12	fill pharmacy authorized to fill prescriptions on behalf of a
13	pharmacy shall have a contractual relationship with the pharmacy
14	that provides for this activity or shall share a common owner
15	with the pharmacy. A central fill pharmacy shall not prepare
16	prescriptions for any controlled substance listed in schedule
17	II.
18	$\left[\frac{(i)}{(k)}\right]$ Partial filling of controlled substance
19	prescriptions shall be determined as follows:
20	(1) The partial filling of a prescription for a controlled

substance listed in schedule II is permissible if the

1		pharmacist is unable to supply the full quantity
2		called for in a written, electronic prescription, or
3		emergency oral prescription and the pharmacist makes a
4		notation of the quantity supplied on the face of the
5		written prescription (or written record of the
6		electronic prescription or emergency oral
7		prescription). The remaining portion of the
8		prescription may be filled within seventy-two hours of
9		the first partial filling; provided that if the
10		remaining portion is not or cannot be filled within
11		the seventy-two-hour period, the pharmacist shall
12		notify the prescribing individual practitioner. No
13	~- ^	further quantity shall be supplied beyond seventy-two
14		hours without a new prescription;
15	(2)	The partial filling of a prescription for a controlled
16		substance listed in schedule III, IV, or V is
17		permissible; provided that:
18		(A) Each partial filling is recorded in the same
19		manner as a refilling;

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1		(B)	The total quantity dispensed in all partial
2			fillings does not exceed the total quantity
3			prescribed;
4		(C)	No dispensing occurs more than three months after
5			the date on which the prescription was issued;
6			and
7		(D)	The prescription is refilled no more than two
8			times after the initial date of the prescription,
9			unless the prescription is renewed by the
10			practitioner; and
11	(3)	A pr	escription for a schedule II controlled substance
12		issu	ed for a patient in a long-term care facility or
13		for	a patient with a medical diagnosis documenting a
14		term	inal illness may be filled in partial quantities
15		to i	nclude individual dosage units. If there is any
16		ques	tion whether a patient may be classified as having
17		a te	rminal illness, the pharmacist shall contact the
18		prac	titioner prior to partially filling the
19		pres	cription. Both the pharmacist and the prescribing
20		prac	titioner have a corresponding responsibility to

assure that the controlled substance is for a

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1	terminally ill patient. The pharmacist shall record
2	on the prescription document on file whether the
3	patient is "terminally ill" or a "long-term care
4	facility patient". For the purposes of this section,
5	"TI" means terminally ill and "LTCF" means long-term
6	care facility. A prescription that is partially
7	filled and does not contain the notation "TI" or "LTCF
8	patient" shall be deemed to have been filled in
9	violation of this section. For each partial filling,
0	the dispensing pharmacist shall record on the back of
.1	the prescription (or on another appropriate record,
2	uniformly maintained, and readily retrievable) the
3	date of the partial filling, quantity dispensed,
.4	remaining quantity authorized to be dispensed, and the
5	identification of the dispensing pharmacist. The
6	total quantity of schedule II controlled substances
.7	dispensed in all partial fillings shall not exceed the
8	total quantity prescribed, nor shall a prescription be
9	partially filled more than three times after the
20	initial date of the prescription. Schedule II
21	controlled substance prescriptions for patients in a

1	long-term care facility or patients with a medical
2	diagnosis documenting a terminal illness shall be
3	valid for a period not to exceed thirty days from the
4	issue date unless sooner terminated by the
5	discontinuance of medication.
6	$\left[\frac{(j)}{(j)}\right]$ A prescription for a schedule II controlled
7	substance may be transmitted by the practitioner or the
8	practitioner's agent to a pharmacy by facsimile equipment;
9	provided that the original written, signed prescription is
10	presented to the pharmacist for review prior to the actual
11	dispensing of the controlled substance, except as noted in
12	subsections $[\frac{(k)}{(l)}, \frac{(l)}{(l)}, \frac{(m)}{(l)}], \frac{(n)}{(l)}, \frac{(n)}{(l)}, \frac{(n)}{(l)}$ The original
13	prescription shall be maintained in accordance with section 329-
14	36. A prescription for a schedule III, IV, or V controlled
15	substance may be transmitted by the practitioner or the
16	practitioner's agent to a pharmacy by facsimile; provided that:
17	(1) The information shall be communicated only between the
18	prescribing practitioner or the prescriber's
19	authorized agent and the pharmacy of the patient's
20	choice. The original prescription shall be maintained
21	by the practitioner in accordance with section 329-36;

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1	(2)	The information shall be communicated in a
2		retrievable, recognizable format acceptable to the
3		intended recipient and shall include the physician's
4		oral code designation and the name of the recipient
5		pharmacy;
6	(3)	No electronic system, software, or other intervening

- (3) No electronic system, software, or other intervening mechanism or party shall alter the practitioner's prescription, order entry, selection, or intended selection without the practitioner's approval on a per prescription per order basis. Facsimile prescription information shall not be altered by any system, software, or other intervening mechanism or party prior to receipt by the intended pharmacy;
  - (4) The prescription information processing system shall provide for confidentiality safeguards required by federal or state law; and
- (5) Prescribing practitioners and pharmacists shall exercise prudent and professional judgment regarding the accuracy, validity, and authenticity of any facsimile prescription information. The facsimile shall serve as the original written prescription for

1	purposes of this section and shall be maintained in
2	accordance with section 329-36.
<b>3</b>	$\left[\frac{(k)}{m}\right]$ A prescription prepared in accordance with
4	subsection $\left[\frac{g}{g}\right]$ (i) written for a narcotic listed in schedule
5	II to be compounded for the direct administration to a patient
6	by parenteral, intravenous, intramuscular, subcutaneous, or
7	intraspinal infusion, but does not extend to the dispensing of
8	oral dosage units of controlled substances, may be transmitted
9	by the practitioner or the practitioner's agent to the pharmacy
10	by facsimile. The original prescription shall be maintained by
11	the practitioner in accordance with section 329-36. The
12	pharmacist shall note on the face of the facsimile prescription
13	in red ink "Home Infusion/IV" and this facsimile shall serve as
14	the original written prescription for purposes of this section
15	and it shall be maintained in accordance with section 329-36.
16	$\left[\frac{(1)}{(n)}\right]$ A prescription prepared in accordance with
17	subsection $\left[\frac{g}{g}\right]$ (i) written for a schedule II substance for a
18	patient enrolled in a hospice care program certified or paid for
19	by medicare under Title XVIII or a hospice program that is
20	licensed by the State may be transmitted by the practitioner or
21	the practitioner's agent to the dispensing pharmacy by

- 1 facsimile. The original prescription shall be maintained by the
- 2 practitioner in accordance with section 329-36. The
- 3 practitioner or practitioner's agent shall note on the
- 4 prescription that the patient is a hospice patient. The
- 5 pharmacist shall note on the face of the facsimile prescription
- 6 in red ink "HOSPICE" and this facsimile shall serve as the
- 7 original written prescription for purposes of this section and
- 8 it shall be maintained in accordance with section 329-36.
- 9 [-(m)] (o) A prescription prepared in accordance with
- 10 subsection  $\left[\frac{g}{g}\right]$  (i) written for a schedule II controlled
- 11 substance for a resident of a state-licensed long-term care
- 12 facility may be transmitted by the practitioner or the
- 13 practitioner's agent to the dispensing pharmacy by facsimile.
- 14 The original prescription shall be maintained by the
- 15 practitioner in accordance with section 329-36. The pharmacist
- 16 shall note on the face of the facsimile prescription in red ink
- 17 "LTCF" and this facsimile shall serve as the original written
- 18 prescription for purposes of this section and it shall be
- 19 maintained in accordance with section 329-36.

1	( <del>n)</del>	<u>(p)</u> An electronic prescription for a schedule II,
2	III, IV,	or V controlled substance may be electronically
3	transmitt	ed by the practitioner to a pharmacy; provided that:
4	(1)	The information shall be communicated only between the
5		prescribing practitioner and the pharmacy of the
6		patient's choice. The electronic prescription shall
7		be maintained by the practitioner in accordance with
8		section 329-36;
9	(2)	The information shall be communicated in a
10		retrievable, recognizable format acceptable to the
11		intended recipient;
12	(3)	No electronic system, software, or other intervening
13		mechanism or party shall alter the practitioner's
14		prescription, order entry, selection, or intended
15		selection without the practitioner's approval on a
16		per-prescription, per-order basis. Transmitted
17		prescription information shall not be altered by any
18		electronic system, software, or other intervening
19		mechanism or party prior to receipt by the intended
20		pharmacy;

pharmacy;

1	(4)	The prescription information processing system shall
2		provide for confidentiality safeguards required by any
3		applicable federal or state law; and
4	(5)	Prescribing practitioners and pharmacists shall
5		exercise prudent and professional judgment regarding
6		the accuracy, validity, and authenticity of any
7		electronic prescription information."
8	SECT	ION 4. Section 457-12, Hawaii Revised Statutes, is
9	amended by	y amending subsection (a) to read as follows:
10	"(a)	In addition to any other actions authorized by law,
11	the board	shall have the power to deny, revoke, limit, or
12	suspend ar	ny license to practice nursing as a registered nurse or
13	as a lice	nsed practical nurse applied for or issued by the board
14	in accorda	ance with this chapter, and to fine or to otherwise
15	discipline	e a licensee for any cause authorized by law, including
16	but not 1:	imited to the following:
17	(1)	Fraud or deceit in procuring or attempting to procure
18		a license to practice nursing as a registered nurse or
19		as a licensed practical nurse;
20	(2)	Gross immorality;

1	(3)	Unfitness or incompetence by reason of negligence,
2		habits, or other causes;
3	(4)	Habitual intemperance, addiction to, or dependency on
4		alcohol or other habit-forming substances;
5	(5)	Mental incompetence;
6	(6)	Unprofessional conduct as defined by the board in
7		accordance with its own rules;
8	(7)	Wilful or repeated violation of any of the provisions
9		of this chapter or any rule adopted by the board;
10	(8)	Revocation, suspension, limitation, or other
11		disciplinary action by another state of a nursing
12		license;
13	(9)	Conviction, whether by nolo contendere or otherwise,
14		of a penal offense substantially related to the
15		qualifications, functions, or duties of a nurse,
16		notwithstanding any statutory provision to the
17		contrary;
18	(10)	Failure to report to the board any disciplinary action
19		taken against the licensee in another jurisdiction
20		within thirty days after the disciplinary action
21		becomes final;

1	(11)	Submitting to or filing with the board any notice,
2		statement, or other document required under this
3		chapter, which is false or untrue or contains any
4		material misstatement of fact, including a false
5		attestation of compliance with continuing competency
6		requirements; [ox]
7	(12)	Violation of the conditions or limitations upon which
8		any license is issued[+]; or
9	(13)	Violation of chapter 329, the uniform controlled
10		substances act, or any rule adopted thereunder except
1		as provided in section 329-122."
12	SECT	ION 5. Statutory material to be repealed is bracketed
13	and stric	ken. New statutory material is underscored.
14	SECT	ION 6. This Act shall take effect on July 1, 2017, and
15	shall be	repealed on June 30, 2023; provided that sections 329-
16	38 and 45	7-12(a), Hawaii Revised Statutes, shall be reenacted in
17	the form	in which they read on the day prior to the effective
18	date of t	his Act.

#### Report Title:

Opioid Therapy; Informed Consent; Prescription Limits; Nurses

#### Description:

Requires prescribing healthcare providers to adopt and maintain policies for informed consent to opioid therapy in circumstances that carry elevated risk of dependency. Establishes limits for concurrent opioid and benzodiazepine prescriptions. Clarifies Board of Nursing authority to enforce compliance with Uniform Controlled Substances Act. Repeals 6/30/2023. (CD1)

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