A BILL FOR AN ACT

RELATING TO ACCESS TO TREATMENT FOR TERMINALLY ILL PATIENTS.

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

- 1 SECTION 1. The legislature finds that the process of
- 2 approval for investigational drugs and biological products in
- 3 the United States protects future patients from premature,
- 4 ineffective, and unsafe medications and treatments over the long
- 5 run, but the process often takes many years. Patients who have
- 6 a terminal illness can be severely restricted in care options
- 7 until an investigational drug or biological product receives
- 8 final approval from the United States Food and Drug
- 9 Administration.
- 10 The legislature further finds that because patients who
- 11 have a terminal illness may often not have the time to wait for
- 12 a potentially lifesaving investigational drug or biological
- 13 product to gain final approval from the United States Food and
- 14 Drug Administration, the federal government and forty-one states
- 15 have enacted "right-to-try" legislation that makes available
- 16 experimental drugs without Food and Drug Administration approval

1	to terminally ill patients with no other medication or treatment
2	options.
3	The purpose of this Act is to grant patients with terminal
4	illnesses access to potentially life-saving investigational
5	drugs and biological products that have not received final
6	approval from the United States Food and Drug Administration.
7	SECTION 2. Chapter 321, Hawaii Revised Statutes, is
8	amended by adding a new section to be appropriately designated
9	and to read as follows:
10	"§321- Access to investigational drugs and biological
11	products for terminally ill patients. (a) Notwithstanding
12	section 328-17, beginning January 1, 2024, a manufacturer of an
13	investigational drug or biological product may make available
14	the manufacturer's investigational drug or biological product to
15	eligible patients pursuant to this section. This section does
16	not require that a manufacturer make available an
17	investigational drug or biological product to an eligible
18	patient. A manufacturer may:
19	(1) Provide an investigational drug or biological product
20	to an eligible patient without receiving compensation;
21	or

1	(2)	Require an eligible patient to pay the costs of, or
2		the costs associated with, the manufacture of the
3		investigational drug or biological product.
4	(b)	A health insurance carrier may, but is not required
5	to, provi	de coverage for the cost of an investigational drug or
6	biologica	l product.
7	<u>(c)</u>	An insurer may deny coverage to an eligible patient
8	from the	time the eligible patient begins use of the
9	investiga	tional drug or biological product through a period not
10	to exceed	six months from the time the investigational drug or
11	biologica	l product is no longer used by the eligible patient;
12	provided	that coverage may not be denied for a preexisting
13	condition	and for coverage for benefits that commence prior to
14	the time	the eligible patient begins use of the investigational
15	drug or b	piological product.
16	(d)	If a patient dies while being treated by an
17	investiga	tional drug or biological product, the patient's heirs
18	shall not	be liable for any outstanding debt related to the
19	treatment	or lack of insurance due to the treatment.
20	(e)	Notwithstanding any law to the contrary, a licensing
21	board may	not revoke, fail to renew, suspend, or take any action

- 1 against a health care provider's license based on the health
- 2 care provider's recommendations to an eligible patient regarding
- 3 access to or treatment with an investigational drug or
- 4 biological product that is being developed to treat the type of
- 5 terminal illness that afflicts the patient, any physical or
- 6 psychological symptoms of the patient's terminal illness, or for
- 7 palliative care. Action against a health care provider's
- 8 medicare certification based on the health care provider's
- 9 recommendation that a patient have access to an investigational
- 10 drug or biological product that is being developed to treat the
- 11 type of terminal illness that afflicts the patient, any physical
- 12 or psychological symptoms of the patient's terminal illness, or
- 13 for palliative care shall be prohibited.
- 14 (f) An official, employee, or agent of the State shall not
- 15 block or attempt to block an eligible patient's access to an
- 16 investigational drug or biological product. Counseling, advice,
- 17 or a recommendation consistent with medical standards of care
- 18 from a licensed health care provider shall not constitute a
- 19 violation of this section.
- 20 (g) This section does not create a private cause of action
- 21 against a manufacturer of an investigational drug or biological

1	product o	r against another person or entity involved in the care
2	of an elig	gible patient using the investigational drug or
3	biologica	l product, for any harm done to the eligible patient
4	resulting	from the investigational drug or biological product;
5	provided	that the manufacturer or other person or entity is
6	complying	in good faith with the terms of this section; provided
7	further the	here was no failure to exercise reasonable care.
8	(h)	For the purposes of this section:
9	"Eli	gible patient" means a person who has:
10	(1)	A terminal illness, attested to by the patient's
11		treating physician;
12	(2)	Considered all other treatment options currently
13		approved by the United States Food and Drug
14		Administration;
15	(3)	Been unable to participate in a clinical trial for the
16		terminal illness within one hundred miles of the
17		patient's home address for the terminal illness, or
18		not been accepted to the clinical trial within one
19		week of completion of the clinical trial application
20		process;

1	(4)	Received a recommendation from the patient's physician
2		for an investigational drug or biological product in
3		order to treat the patient's terminal illness,
4		physical or psychological symptoms of the patient's
5		terminal illness, or for palliative care;
6	(5)	Given written, informed consent for the use of the
7		investigational drug or biological product or, if the
8		patient is a minor or lacks the mental capacity to
9		provide informed consent, a parent or legal guardian
10		has given written, informed consent on the patient's
11		behalf; and
12	(6)	Documentation from the patient's physician that the
13		patient meets the requirements of this definition.
14	"Eligible	patient" does not include a person being treated as an
15	inpatient	in an institution with an organized medical staff,
16	regulated	under section 321-11(10), or a health care facility
17	under cha	pter 323F.
18	"Inv	estigational drug or biological product" means a drug
19	or biolog	ical product that has successfully completed phase one
20	of a clin	ical trial but has not yet been approved for general
21	use by the	e United States Food and Drug Administration and

1	remains u	nder investigation in a united states rood and brug
2	Administra	ation-approved clinical trial.
3	<u>"Ter</u>	minal illness" means a disease that, without life-
4	sustainin	g procedures, will result in death or a state of
5	permanent	unconsciousness from which recovery is unlikely.
6	<u>"Wri</u>	tten, informed consent" means a written document signed
7	by the pa	tient and attested to by the patient's physician and a
8	witness t	hat, at a minimum:
9	(1)	Explains the currently approved products and
10		treatments for the disease or condition from which the
11		patient suffers;
12	(2)	Attests to the fact that the patient concurs with the
13		patient's physician in believing that all currently
14		approved and conventionally recognized treatments are
15		unlikely to prolong the patient's life;
16	<u>(3)</u>	Clearly identifies the specific proposed
17		investigational drug or biological product that the
18		patient is seeking to use;
19	(4)	Describes the potentially best and worst outcomes of
20		using the investigational drug or biological product
21		with a realistic description of the most likely

1		outcome, including the possibility that new,
2		unanticipated, different, or worse symptoms might
3		result, and that death could be hastened by the
4		proposed treatment, based on the physician's knowledge
5		of the proposed treatment in conjunction with an
6		awareness of the patient's condition;
7	<u>(5)</u>	Makes clear that the patient's health insurer and
8		provider are not obligated to pay for any care or
9		treatments consequent to the use of the
10		investigational drug or biological product;
11	(6)	Makes clear that the patient's eligibility for hospice
12		care may be withdrawn by the hospice care provider if
13		the patient begins curative treatment and care may be
14		reinstated if the curative treatment ends and the
15		patient meets hospice eligibility requirements;
16	(7)	Makes clear that in-home health care may be denied if
17		treatment begins; and
18	(8)	States that the patient understands that the patient
19		is liable for all expenses consequent to the use of
20		the investigational drug or biological product, and
21		that this liability extends to the patient's estate,

1	unless a contract between the patient and the
2	manufacturer of the investigational drug or biological
3	<pre>product states otherwise."</pre>
4	SECTION 3. New statutory material is underscored.
5	SECTION 4. This Act shall take effect upon its approval.
6	INTRODUCED BY: DULA A Belletti
	JAN 2 3 2023

Report Title:

Right-to-Try Act; Terminally Ill Patients; Investigational Drugs; Biological Products

Description:

Beginning on January 1, 2024, permits manufacturers of investigational drugs or biological products to make these drugs and products available to terminally ill patients under certain conditions.

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