

SENATE FILE NO. SF0003

Right to try.

Sponsored by: Senator(s) Burns and Representative(s)
Barlow, Berger, Clem, Miller, Patton and
Throne

A BILL

for

1 AN ACT relating to public health and safety; authorizing
2 provision of certain investigational drugs, biological
3 products and devices by manufacturers; specifying
4 availability and costs of investigational drugs, biological
5 products and devices; specifying that no private cause of
6 action against manufacturers and other entities is created;
7 providing definitions; and providing for an effective date.

8

9 *Be It Enacted by the Legislature of the State of Wyoming:*

10

11 **Section 1.** W.S. 35-7-1801 through 35-7-1805 are
12 created to read:

13

14

ARTICLE 18

15

RIGHT TO TRY ACT

1

2 **35-7-1801. Short title.**

3

4 This article is known and may be cited as the "Right To Try
5 Act."

6

7 **35-7-1802. Definitions.**

8

9 (a) As used in this article:

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11 (i) "Eligible patient" means a person who has:

12

13 (A) A terminal illness;

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15 (B) Considered all other treatment options
16 currently approved by the United States food and drug
17 administration;

18

19 (C) Received a recommendation from a
20 physician for an investigational drug, biological product
21 or device;

22

1 (D) Given written, informed consent for the
2 use of the investigational drug, biological product or
3 device or, if the patient is a minor or lacks the mental
4 capacity to provide informed consent, a parent or legal
5 guardian has given written informed consent on the
6 patient's behalf; and

7

8 (E) Documentation from a physician that the
9 person meets the requirements of this paragraph.

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11 (ii) "Investigational drug, biological product
12 or device" means a drug, biological product or device that
13 has:

14

15 (A) Successfully completed phase two of a
16 clinical trial but has not yet been approved for general
17 use by the United States Food and Drug Administration and
18 remains under investigation in a clinical trial; or

19

20 (B) Has been approved for general use in
21 one (1) or more member countries of the Organisation for
22 Economic Co-operation and Development.

23

1 (iii) "Terminal illness" means a disease that,
2 without life-sustaining procedures, will soon result in
3 death or a state of permanent unconsciousness from which
4 recovery is unlikely.

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6 **35-7-1803. Availability of investigational drugs,**
7 **biological products or devices; costs; insurance coverage.**

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9 (a) A manufacturer of an investigational drug,
10 biological product or device may make the drug, product or
11 device available to eligible patients in accordance with
12 the provisions of this section. Nothing in this section
13 shall be construed to require a manufacturer to make
14 available any drug, product or device.

15

16 (b) A manufacturer may:

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18 (i) Provide an investigational drug, biological
19 product or device to an eligible patient without receiving
20 compensation; or

21

1 (ii) Require an eligible patient to pay the
2 costs of or associated with the manufacture of the
3 investigational drug, biological product or device.

4

5 (c) A health care insurer may, but is not required
6 to, provide coverage for the cost of an investigational
7 drug, biological product or device.

8

9 (d) Nothing in this section expands the coverage
10 provided in W.S. 26-20-301.

11

12 **35-7-1804. Access to investigational drugs,**
13 **biological products and devices.**

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15 An official, employee or agent of this state shall not
16 block or attempt to block an eligible patient's access to
17 an investigational drug, biological product or device.
18 Counseling, advice or a recommendation consistent with
19 medical standards of care from a licensed health care
20 provider is not a violation of this section.

21

22 **35-7-1805. No cause of action created.**

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1 This article does not create a private cause of action
2 against a manufacturer of an investigational drug,
3 biological product or device, or against any other person
4 or entity involved in the care of an eligible patient using
5 the investigational drug, biological product or device, so
6 long as the manufacturer or other person or entity is
7 complying in good faith with the terms of this article.

8

9 **Section 2.** This act is effective July 1, 2015.

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11

(END)