GENERAL ASSEMBLY OF NORTH CAROLINA SESSION 2023

H HOUSE BILL 1029

Short Title:	Right To Try Individualized Treatments.	(Public)	
Sponsors:	Representatives Chesser, Blackwell, Potts, and Reeder (Primary Sponsors). For a complete list of sponsors, refer to the North Carolina General Assembly web site.		
Referred to:	Health, if favorable, Appropriations, if favorable, Rules, Calendar, and Operations of the House		

May 7, 2024

A BILL TO BE ENTITLED

AN ACT TO PROVIDE ELIGIBLE PATIENTS THE RIGHT TO TRY INDIVIDUALIZED INVESTIGATIONAL DRUGS, BIOLOGICAL PRODUCTS, AND DEVICES TO TREAT LIFE-THREATENING OR SEVERELY DEBILITATING ILLNESSES AND TO APPROPRIATE FUNDS TO THE DEPARTMENT OF HEALTH AND HUMAN SERVICES.

The General Assembly of North Carolina enacts:

SECTION 1. Article 23A of Chapter 90 of the General Statutes is amended by adding a new Part to read:

"Part 3. Individualized Treatments.

"§ 90-325.30. Definitions.

The following definitions apply in this Part, unless the context requires otherwise:

- (1) Eligible facility. Any institution operating under Federalwide Assurance for the Protection of Human Subjects in accordance with 45 C.F.R. § 46 and 42 U.S.C. § 289(a).
- (2) Eligible patient. An individual who meets all of the following criteria:
 - <u>a.</u> <u>Has a life-threatening or severely debilitating illness, attested to by a treating physician.</u>
 - b. Has, in consultation with a treating physician, considered all other treatment options currently approved by the United States Food and Drug Administration.
 - c. Has received a recommendation from the treating physician for use of an individualized investigational drug, biological product, or device for treatment of the life-threatening or severely debilitating illness.
 - d. Has given informed consent in writing to use of the individualized investigational drug, biological product, or device for treatment of the life-threatening or severely debilitating illness or, if the individual is a minor or is otherwise incapable of providing informed consent, the parent or legal guardian has given informed consent in writing to use of the individualized investigational drug, biological product, or device.
 - e. Has documentation from the treating physician that the individual meets all of the criteria for this definition. This documentation shall include an attestation from the treating physician that the treating



1			physician was consulted in the creation of the written, informed
2			consent required under this Part.
3	<u>(3)</u>		lualized investigational drug, biological product, or device. – A drug,
4			ical product, or device that is unique and produced exclusively for use
5			individual patient, based on their own genetic profile, including
6		individ	lualized gene therapy antisense oligonucleotides and individualized
7		neoant	igen vaccines.
8	<u>(4)</u>	<u>Institut</u>	ion. – As defined in 45 C.F.R. § 46.102(f).
9	<u>(5)</u>	Life-th	reatening or severely debilitating illness As those terms are defined
10		<u>in 21 C</u>	C.F.R. § 312.81.
11	<u>(6)</u>	Writter	n, informed consent. – A written document that is signed by an eligible
12		patient	; or if the patient is a minor, by a parent or legal guardian; or if the
13		patient	is incapacitated, by a designated health care agent pursuant to a health
14		care po	ower of attorney, that at a minimum includes all of the following:
15		<u>a.</u>	An explanation of the currently approved products and treatments for
16		_	the eligible patient's life-threatening or severely debilitating illness.
17		<u>b.</u>	An attestation that the eligible patient concurs with the treating
18		_	physician in believing that all currently approved treatments are
19			unlikely to prolong the eligible patient's life.
20		<u>c.</u>	Clear identification of the specific individualized investigational drug,
21			biological product, or device proposed for treatment of the eligible
22			patient's terminal illness.
23		<u>d.</u>	A description of the potentially best and worst outcomes resulting
24		<u></u>	from use of the individualized investigational drug, biological product,
25			or device to treat the eligible patient's life-threatening or severely
26			debilitating illness, along with a realistic description of the most likely
27			outcome. The description shall be based on the treating physician's
28			knowledge of the proposed treatment in conjunction with an
29			awareness of the eligible patient's life-threatening or severely
30			debilitating illness and shall include a statement acknowledging that
31			new, unanticipated, different, or worse symptoms might result from,
32			and that death could be hastened by, the proposed treatment.
33		A	A statement that eligibility for hospice care may be withdrawn if the
34		<u>e.</u>	eligible patient begins treatment of the life-threatening or severely
35			
36			debilitating illness with an individualized investigational drug,
30 37			biological product, or device and that hospice care may be reinstated
38			if such treatment ends and the eligible patient meets hospice eligibility
39		£	requirements. A statement that the eligible nationals health benefit plan or third party.
		<u>f.</u>	A statement that the eligible patient's health benefit plan or third-party
40			administrator and provider are not obligated to pay for any care or
41			treatments consequent to the use of the individualized investigational
42			drug, biological product, or device, unless specifically required to do
43			so by law or contract.
44		<u>g.</u>	A statement that the eligible patient understands that he or she is liable
45			for all expenses consequent to the use of the individualized
46			investigational drug, biological product, or device and that this
47			liability extends to the eligible patient's estate, unless a contract
48			between the patient and the manufacturer of the drug, biological
49			product, or device states otherwise.
50		<u>h.</u>	A statement that the eligible patient or, for an eligible patient who is a
51			minor or lacks capacity to provide informed consent, that the parent or

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legal guardian consents to the use of the individualized investigational drug, biological product, or device for treatment of the life-threatening or severely debilitating illness.

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"§ 90-325.31. Authorized access to and use of individualized investigational drugs, biological products, or devices.

- A manufacturer operating within an eligible facility and in accordance with all (a) applicable federal law may make available to an eligible patient, and an eligible patient may request, the manufacturer's individualized investigational drug, biological product, or device from an eligible facility or manufacturer operating within an eligible facility. However, nothing in this Part shall be construed to require a manufacturer of an individualized investigational drug, biological product, or device to make such individualized investigational drug, biological product, or device available to an eligible patient.
- A manufacturer of an individualized investigational drug, biological product, or (b) device may provide the individualized investigational drug, biological product, or device to an eligible patient without receiving compensation or may require the eligible patient to pay the costs of, or the costs associated with, the manufacture of the individualized investigational drug, biological product, or device.

"§ 90-325.32. No liability to heirs for outstanding debt related to use of individualized investigational drugs, biological products, or devices.

If an eligible patient dies while being treated with an individualized investigational drug, biological product, or device, the eligible patient's heirs are not liable for any outstanding debt related to the treatment, including any costs attributed to lack of insurance coverage for the treatment.

"§ 90-325.33. Sanctions against health care providers prohibited.

- A licensing board shall not revoke, fail to renew, suspend, or take any other disciplinary action against a health care provider licensed under this Chapter, based solely on the health care provider's recommendations to an eligible patient regarding access to or treatment with an individualized investigational drug, biological product, or device.
- An entity responsible for Medicare certification shall not take action against a health care provider's Medicare certification based solely on the health care provider's recommendation that a patient have access to an individualized investigational drug, biological product, or device. "§ 90-325.34. Prohibited conduct by State officials.

No official, employee, or agent of this State shall block or attempt to block an eligible patient's access to an individualized investigational drug, biological product, or device. Counseling, advice, or a recommendation consistent with medical standards of care from a licensed health care provider does not constitute a violation of this section.

No private right of action against manufacturers of individualized "\ 90-325.35. investigational drugs, biological products, or devices.

No private right of action may be brought against a manufacturer of an individualized investigational drug, biological product, or device, or against any other person or entity involved in the care of an eligible patient using an individualized investigational drug, biological product, or device, for any harm caused to the eligible patient resulting from use of the individualized investigational drug, biological product, or device as long as the manufacturer or other person or entity has made a good-faith effort to comply with the provisions of this Part and has exercised reasonable care in actions undertaken pursuant to this Part.

"§ 90-325.36. Insurance coverage of clinical trials.

Nothing in this Part shall be construed to affect a health benefit plan's obligation to provide coverage for an insured's participation in a clinical trial pursuant to G.S. 58-3-255."

SECTION 2. There is appropriated from the General Fund to the Department of Health and Human Services the nonrecurring sum of fifty thousand dollars (\$50,000) for the 2024-2025 fiscal year to implement the provisions of this act.

SECTION 3. Section 1 of this act becomes effective October 1, 2024. Section 2 of this act becomes effective July 1, 2024. The remainder of this act is effective when it becomes law.