117TH CONGRESS 1ST SESSION

H. R. 1730

To amend the Federal Food, Drug, and Cosmetic Act to accelerate development of therapies across the spectrum of rare diseases and conditions and facilitate patient access to such therapies, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

March 10, 2021

Mr. Bilirakis (for himself and Mr. Butterfield) introduced the following bill; which was referred to the Committee on Energy and Commerce

A BILL

- To amend the Federal Food, Drug, and Cosmetic Act to accelerate development of therapies across the spectrum of rare diseases and conditions and facilitate patient access to such therapies, and for other purposes.
 - 1 Be it enacted by the Senate and House of Representa-
 - 2 tives of the United States of America in Congress assembled,
 - 3 SECTION 1. SHORT TITLE.
 - 4 This Act may be cited as the "Speeding Therapy Ac-
 - 5 cess Today Act of 2021".
 - 6 SEC. 2. TABLE OF CONTENTS.
 - 7 The table of contents of this Act is as follows:
 - Sec. 1. Short title.
 - Sec. 2. Table of contents.
 - Sec. 3. Intercenter Institute on Rare Diseases and Conditions.

- Sec. 4. Rare Disease and Condition Drug Advisory Committee.
- Sec. 5. Grants and contracts for development of drugs for rare diseases and conditions.

1 SEC. 3. INTERCENTER INSTITUTE ON RARE DISEASES AND

- 2 **CONDITIONS.**
- 3 (a) Establishment Required.—The first sentence
- 4 of section 1014(a) of the Federal Food, Drug, and Cos-
- 5 metic Act (21 U.S.C. 399g(a)) is amended by inserting
- 6 ", at least one of which shall be focused on rare diseases
- 7 and conditions" before the period at the end of the sen-
- 8 tence.
- 9 (b) Timing of Establishment.—Subsection (c) of
- 10 section 1014 of the Federal Food, Drug, and Cosmetic
- 11 Act (21 U.S.C. 399g) is amended to read as follows:
- 12 "(c) Timing.—Not later than the date that is 1 year
- 13 after the date of enactment of the Speeding Therapy Ac-
- 14 cess Today Act of 2021, the Secretary shall establish, in
- 15 accordance with this section and section 529B, an Insti-
- 16 tute under subsection (a) focused on rare diseases and
- 17 conditions, to be known as the Intercenter Institute on
- 18 Rare Diseases and Conditions.".
- 19 (c) Responsibilities.—Subchapter B of chapter V
- 20 of the Federal Food, Drug, and Cosmetic Act (relating
- 21 to drugs for rare diseases or conditions) is amended by
- 22 inserting after section 529A of such Act (21 U.S.C. 360ff-
- 23 1) the following new section:

1	"SEC. 529B. INTERCENTER INSTITUTE ON RARE DISEASES
2	AND CONDITIONS.
3	"(a) Responsibilities.—In addition to carrying out
4	activities listed in section 1014(a), the Intercenter Insti-
5	tute on Rare Diseases and Conditions shall—
6	"(1) serve as the Food and Drug Administra-
7	tion's coordinating office for engagement with rare
8	disease and condition stakeholders, complementing
9	but not supplanting engagement activities between
10	stakeholders and the review divisions;
11	"(2) build, within the Food and Drug Adminis-
12	tration, knowledge and understanding associated
13	with the review of medical products to treat rare dis-
14	eases and conditions, including advancements in trial
15	design, statistical analysis, regulatory science, prod-
16	uct manufacturing, and other topics as determined
17	by the Secretary;
18	"(3) implement cross-center rare disease and
19	condition-focused meetings and policy development;
20	"(4) coordinate rare disease and condition-spe-
21	cific regulatory science initiatives;
22	"(5) facilitate stakeholder engagement to the
23	external community and international regulatory
24	agencies on rare disease and condition product devel-
25	opment;

1	"(6) establish and implement the Accelerating
2	Lifesavings Therapies in Treating Ultra-rare Dis-
3	ease Entities Program under subsection (b); and
4	"(7) establish and carry out the rare disease
5	and condition third-party payor program under sub-
6	section (d).
7	"(b) ALTITUDE PROGRAM.—
8	"(1) In General.—The Intercenter Institute
9	shall establish and implement a program, to be
10	known as the Accelerating Lifesavings Therapies in
11	Treating Ultra-rare Disease Entities Program, to
12	identify and make recommendations to address cur-
13	rent and emerging regulatory science and public pol-
14	icy challenges associated with developing medical
15	products to treat rare diseases or conditions in an
16	individual or very small populations.
17	"(2) Issues.—The program under paragraph
18	(1) shall focus on issues including—
19	"(A) manufacturing standards for thera-
20	pies described in such paragraph, including in
21	non-industry settings;
22	"(B) trial designs and metrics;
23	"(C) regulatory flexibilities for abbreviated
24	toxicology studies, overlapping animal studies,
25	and patient dosing;

1	"(D) regulatory science, chemistry, manu-
2	facturing, and other needs associated with de-
3	veloping such therapies; and
4	"(E) other issues as determined by the
5	Secretary.
6	"(c) Proposals for Amending Labels.—
7	"(1) Stakeholder group.—Not later than
8	180 days after the date of enactment of this section,
9	the Intercenter Institute shall convene a meeting of
10	stakeholders from the rare disease community, in-
11	cluding patients, caregivers, product manufacturers,
12	third-party payors, and others, to consider potential
13	amendments to labels for medical products to treat
14	rare diseases or conditions approved pursuant to a
15	pathway under section 506.
16	"(2) GUIDANCE.—Not later than 90 days after
17	the date of the meeting under paragraph (1), the
18	Secretary shall issue guidance to propose changes to
19	how the labels of medical products to treat rare dis-
20	eases or conditions demonstrate clinical benefits and
21	reflect relevant scientific data including surrogate
22	endpoints.
23	"(d) Rare Disease and Condition Third-Party
24	Payor Program.—

1	"(1) IN GENERAL.—The Intercenter Institute
2	shall establish and carry out a voluntary rare disease
3	and condition early third-party payor feedback pro-
4	gram—
5	"(A) to inform coverage policies for rare
6	disease therapies; and
7	"(B) to inform clinical trial design, patient
8	engagement, and other data collections.
9	"(2) Program requirements.—The program
10	under paragraph (1) shall—
11	"(A) facilitate voluntary communication
12	between sponsors of medical products to treat
13	rare diseases and conditions and third-party
14	payors; and
15	"(B) require participation of the Centers
16	for Medicare & Medicaid Services with rep-
17	resentation from—
18	"(i) the Center for Medicare; and
19	"(ii) the Center for Medicaid and
20	CHIP Services.
21	"(3) Annual Report.—The Intercenter Insti-
22	tute shall—
23	"(A) on an annual basis, submit a report
24	to that Congress on—

1	"(i) the participation within the pro-
2	gram under paragraph (1); and
3	"(ii) the impacts of the program
4	under paragraph (1); and
5	"(B) post each such report on the public
6	website of the Intercenter Institute.
7	"(4) Bulletin to medicaid directors.—
8	Following the approval, clearance, or authorization
9	by the Food and Drug Administration of a medical
10	product to treat a rare disease or condition, the Sec-
11	retary shall issue a bulletin to State Medicaid direc-
12	tors containing information to help inform coverage
13	decisions on the product by State Medicaid and Chil-
14	dren's Health Insurance programs.
15	"(e) Definition.—In this section, the terms 'Inter-
16	center Institute on Rare Diseases and Conditions' and
17	'Intercenter Institute' refer to the Intercenter Institute on
18	Rare Diseases and Conditions established pursuant to sec-
19	tion 1014.".
20	SEC. 4. RARE DISEASE AND CONDITION DRUG ADVISORY
21	COMMITTEE.
22	Subchapter B of chapter V of the Federal Food,
23	Drug, and Cosmetic Act is further amended by inserting
24	after section 529B of such Act, as inserted by section 3,
25	the following new section:

1	"SEC. 529C. RARE DISEASE AND CONDITION DRUG ADVI-
2	SORY COMMITTEE.
3	"(a) In General.—The Secretary shall establish
4	and maintain a committee, to be known as the Rare Dis-
5	ease and Condition Drug Advisory Committee (in this sec-
6	tion referred to as the 'Advisory Committee').
7	"(b) Duty of Committee.—The Advisory Com-
8	mittee shall advise the Secretary on issues associated with
9	development of therapies to treat rare diseases or condi-
10	tions.
11	"(c) Specific Issues.—In advising the Secretary,
12	the Advisory Committee may address issues including—
13	"(1) modified or new regulatory pathways to
14	support review of therapies;
15	"(2) clinical trial design needs, including devel-
16	opment of innovative approaches to clinical trials;
17	"(3) qualifications of biomarkers or other drug
18	development tools for use in reviews;
19	"(4) modified or new standards to support the
20	review of already marketed drugs being evaluated
21	for repurposing to treat a rare disease or condition;
22	and
23	"(5) issues—
24	"(A) that pertain to an application for ap-
25	proval of a therapy to treat a rare disease or
26	condition; and

1	"(B) with respect to which a review divi-
2	sion has requested that the Advisory Committee
3	provide advice.
4	"(d) Membership.—
5	"(1) In General.—The Advisory Committee
6	shall consist of—
7	"(A) not more than 15 members appointed
8	by the Secretary in accordance with paragraph
9	(2); and
10	"(B) the nonvoting ex officio members
11	under paragraph (3).
12	"(2) Appointed members.—
13	"(A) Special government employ-
14	EES.—Members of the Advisory Committee ap-
15	pointed pursuant to paragraph (1)(A) shall
16	serve as special Government employees (as de-
17	fined in section 202(a) of title 18, United
18	States Code).
19	"(B) Eligibility.—To be eligible for ap-
20	pointment pursuant to paragraph (1)(A), an in-
21	dividual shall—
22	"(i) be eligible to serve as special Gov-
23	ernment employee (as defined in section
24	202(a) of title 18, United States Code);
25	and

1	"(ii) have expertise in the fields of
2	public policy, law, regulatory policy, eco-
3	nomics, patient-focused product develop-
4	ment, or patient advocacy.
5	"(C) Composition.—Of the members of
6	the Advisory Committee appointed pursuant to
7	paragraph (1)(A)—
8	"(i) up to 10 shall be selected from
9	among experts in the disciplines relevant to
10	the activities of the Intercenter Institute
11	on Rare Diseases and Conditions, to in-
12	clude at least one expert in each of—
13	"(I) rare disease product develop-
14	ment;
15	"(II) conducting clinical trials
16	with respect to rare diseases and con-
17	ditions, including with respect to very
18	small patient populations;
19	"(III) rare disease and condition
20	natural history and related studies;
21	"(IV) health economics per-
22	taining to the development of medical
23	products for rare diseases or condi-
24	tions;

1	"(V) manufacturing and related
2	needs associated with medical prod-
3	ucts for rare diseases or conditions;
4	and
5	"(VI) patient experience data col-
6	lection; and
7	"(ii) up to 5 shall be selected from the
8	public, to include—
9	"(I) at least 4 individuals who
10	are representatives of the rare disease
11	patient community;
12	"(II) at least one individual who
13	is directly impacted by a rare disease
14	or condition; and
15	"(III) at least one person who
16	serves as a family caregiver to a per-
17	son diagnosed with a rare disease or
18	condition.
19	"(3) Nonvoting ex officio members.—The
20	nonvoting ex officio members of the Advisory Com-
21	mittee under paragraph (1)(B) shall consist of the
22	following:
23	"(A) The Secretary (or the Secretary's
24	designee).

1	"(B) The Director of the Intercenter Insti-
2	tute on Rare Diseases and Conditions.
3	"(C) The Director of the Center for Bio-
4	logics Evaluation and Research (or the Direc-
5	tor's designee).
6	"(D) The Director of the Center for Drug
7	Evaluation and Research (or the Director's des-
8	ignee).
9	"(E) The Director of the Center for De-
10	vices and Radiological Health (or the Director's
11	designee).
12	"(F) The Director of the National Center
13	for the Advancing Translational Sciences of the
14	National Institutes of Health (or the Director's
15	designee).
16	"(G) The Administrator of the Centers for
17	Medicare & Medicaid Services (or the Adminis-
18	trator's designee).
19	"(H) Any additional officers or employees
20	of the Department of Health and Human Serv-
21	ices as the Secretary determines necessary for
22	the Advisory Committee to effectively carry out
23	its functions.

"(4) CHAIR.—The Chair of the Advisory Com-1 2 mittee shall be the Director of the Intercenter Insti-3 tute for Rare Diseases and Conditions. "(5) Terms.— 4 "(A) Members.— 6 "(i) In general.—The term of a 7 member of the Advisory Committee ap-8 pointed pursuant to paragraph (1)(A) shall 9 be 4 years, except that any member ap-10 pointed to fill a vacancy in an unexpired 11 term shall be appointed for the remainder 12 of that term. 13 "(ii) Continued Service.—A mem-14 ber appointed pursuant to paragraph 15 (1)(A) may continue serving as a member of the Advisory Committee for up to 180 16 17 days after the expiration of that member's 18 term if a successor has not been appointed. 19 "(B) REAPPOINTMENT.—A member of the 20 Advisory Committee who has been appointed 21 pursuant to paragraph (1)(A) for a term of 4 22 years may not be reappointed to serve as a 23 member of the Advisory Committee before the 24 date that is 2 years after the date of expiration 25 of that member's term.

- 1 "(e) QUORUM.—A majority of the appointed mem-
- 2 bers of the Advisory Committee shall constitute a quorum
- 3 for the conduct of business.".
- 4 SEC. 5. GRANTS AND CONTRACTS FOR DEVELOPMENT OF
- 5 DRUGS FOR RARE DISEASES AND CONDI-
- 6 TIONS.
- 7 (a) AUTHORITY OF SECRETARY.—Section 5(a) of the
- 8 Orphan Drug Act (21 U.S.C. 360ee(a)) is amended—
- 9 (1) in paragraph (2), by striking "and" at the
- end; and
- 11 (2) by inserting before the period at the end ",
- and (4) developing practices pertaining to the chem-
- istry, manufacturing, regulatory approval of, and
- 14 controls of individualized therapies or therapies to
- treat very small populations".
- 16 (b) ALTITUDE PROGRAM.—In supporting grants
- 17 and contracts under section 5(a)(4) of the Orphan Drug
- 18 Act, as added by subsection (a), the Secretary of Health
- 19 and Human Services shall consult with the Director of the
- 20 Intercenter Institute on Rare Diseases and Conditions re-
- 21 garding the Accelerating Lifesavings Therapies in Treat-
- 22 ing Ultra-rare Disease Entities Program established under
- 23 section 529B(b) of the Federal Food, Drug, and Cosmetic
- 24 Act, as added by section 3(c) of this Act, to—

1	(1) identify the regulatory science and related
2	challenges and needs associated with developing indi-
3	vidualized therapies or therapies to treat very small
4	patient populations; and

(2) support research to address such challenges.

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