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(Original Signature of Member)

118TH CONGRESS
2D SESSION

H. R. _____

To provide for a comprehensive Federal response to Long COVID, including research, education, and support for affected individuals, to direct the National Institutes of Health to establish a Long COVID research program, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

Ms. OMAR introduced the following bill; which was referred to the Committee
on _____

A BILL

To provide for a comprehensive Federal response to Long COVID, including research, education, and support for affected individuals, to direct the National Institutes of Health to establish a Long COVID research program, 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 “Long COVID Research
5 Moonshot Act”.

1 **TITLE I—LONG COVID**
2 **BIOMEDICAL RESEARCH**

3 **SEC. 101. ESTABLISHMENT OF LONG COVID RESEARCH**
4 **PROGRAM.**

5 Title IV of the Public Health Service Act (42 U.S.C.
6 281 et seq.) is amended by adding at the end the fol-
7 lowing:

8 **“PART K—LONG COVID PROGRAMS**

9 **“SEC. 499B. ESTABLISHMENT OF LONG COVID RESEARCH**
10 **PROGRAM.**

11 “(a) IN GENERAL.—There is established within the
12 Office of the Director of the National Institutes of Health
13 a research program, to be known as the Long COVID Re-
14 search Program (referred to in this part as the ‘Pro-
15 gram’), for purposes of expediting research to identify new
16 ways to prevent, detect, manage, and treat symptoms as-
17 sociated with Long COVID.

18 “(b) DIRECTOR.—

19 “(1) APPOINTMENT.—

20 “(A) IN GENERAL.—The Program shall be
21 headed by a Director, appointed by the Sec-
22 retary, in consultation with the Director of
23 NIH, who has—

24 “(i) experience managing clinical or
25 research programs focused on pathogenic

1 mechanisms and biological pathways re-
2 lated to Long COVID; and

3 “(ii) demonstrated commitment to ad-
4 dressing Long COVID and other infection-
5 associated chronic conditions, such as
6 myalgic encephalomyelitis/chronic fatigue
7 syndrome, postural orthostatic tachycardia
8 syndrome, and post-treatment Lyme dis-
9 ease syndrome/persistent Lyme disease.

10 “(B) CONSULTATION.—In appointing the
11 Director under subparagraph (A), the Secretary
12 shall consult with independent, patient-led orga-
13 nizations or advocacy groups representing Long
14 COVID patients and their families.

15 “(2) RESPONSIBILITIES.—The Director of the
16 Program shall—

17 “(A) act as the primary Federal official
18 with responsibility for coordinating all Long
19 COVID research conducted or supported by the
20 National Institutes of Health;

21 “(B) represent the National Institutes of
22 Health Long COVID Research Program at all
23 relevant Executive branch task force meetings
24 and committees; and

1 “(C) maintain communication with all rel-
2 evant Federal departments and agencies to en-
3 sure the timely transmission of information con-
4 cerning advances in Long COVID research and
5 the clinical treatment of Long COVID and
6 other infection-associated chronic conditions be-
7 tween such departments and agencies, and for
8 dissemination to affected communities and
9 health care providers.

10 “(c) ACTIVITIES.—The Program shall—

11 “(1) investigate the etiology, pathophysiology,
12 risk factors, and pathology of Long COVID in
13 adults and children;

14 “(2) explore the best ways to prevent, detect,
15 monitor, manage, and treat Long COVID in adults
16 and children;

17 “(3) contribute knowledge to the under-
18 standing, prevention, mitigation, management, and
19 treatment of Long COVID;

20 “(4) develop and facilitate programs on Long
21 COVID, within the National Institutes of Health
22 and in other settings;

23 “(5) conduct comparative research to under-
24 stand the similarities and differences between Long
25 COVID and other infection-associated chronic condi-

1 tions with similar phenotypes, such as myalgic
2 encephalomyelitis/chronic fatigue syndrome, postural
3 orthostatic tachycardia syndrome, and post-treat-
4 ment Lyme disease syndrome/persistent Lyme dis-
5 ease, and how activities funded by the Program
6 could improve understanding of such other condi-
7 tions; and

8 “(6) conduct comparative research to under-
9 stand the similarities and differences between Long
10 COVID and severe, long-term effects from COVID-
11 19 vaccinations.

12 “(d) DUTIES.—

13 “(1) INTERAGENCY COORDINATION OF LONG
14 COVID ACTIVITIES.—The Director of the Program
15 shall coordinate with the national research institutes
16 and national centers, as appropriate, on Long
17 COVID research. In carrying out this paragraph,
18 the Director of the Program shall evaluate the Long
19 COVID activities of each such institute or center
20 and shall provide for the periodic reevaluation of
21 such activities.

22 “(2) CONSULTATION.—The Director of the Pro-
23 gram shall carry out all duties, including the devel-
24 opment of the research plan under section 499B-1
25 in consultation with the heads of the national re-

1 search institutes and national centers, with the advi-
2 sory councils of such institutes and centers, and with
3 the Long COVID Research Program Advisory Board
4 established under section 499B-4.

5 “(e) NON-DUPLICATION OF EFFORT.—The Director
6 shall ensure that activities carried out under this section
7 do not unnecessarily duplicate the efforts of other Federal
8 departments or agencies.

9 **“SEC. 499B-1. LONG COVID RESEARCH PLAN.**

10 “(a) IN GENERAL.—Not later than 1 year after the
11 date of enactment of the Long COVID Research Moonshot
12 Act, the Director of the Program established under section
13 499B shall develop and make public a comprehensive re-
14 search plan for the conduct and support of all Long
15 COVID research activities of the national research insti-
16 tutes and national centers. The Director of the Program
17 shall update such plan annually.

18 “(b) CONTENTS.—The research plan developed under
19 subsection (a) shall—

20 “(1) identify current Long COVID research
21 conducted or supported by the national research in-
22 stitutes and national centers, opportunities and
23 needs for additional research, including among pa-
24 tients who face the highest disease burden and pedi-
25 atric patients, and priorities for such research;

1 “(2) evaluate the progress of Long COVID re-
2 search against strategic priorities, goals, and objec-
3 tives, identified in previous versions of the research
4 plan;

5 “(3) make recommendations for the coordina-
6 tion of such research conducted or supported by the
7 National Institutes of Health and other agencies of
8 the Federal Government; and

9 “(4) include goals and objectives of the Pro-
10 gram for conducting, supporting, and coordinating
11 Long COVID research.

12 “(c) REQUIREMENTS.—In developing the research
13 plan under subsection (a), the Director of the Program
14 shall—

15 “(1) ensure that the plan establishes priorities
16 among Long COVID research that the Program is
17 authorized to carry out;

18 “(2) ensure that the plan establishes objectives
19 regarding such research and describes the means for
20 achieving the objectives;

21 “(3) ensure that all amounts appropriated for
22 such research under section 499B–6 are expended in
23 accordance with the plan;

24 “(4) review the plan not less frequently than
25 annually, and revise the plan as appropriate to

1 prioritize funding and research relative to scientific
2 urgency;

3 “(5) ensure that the plan serves as a broad,
4 binding statement of policies regarding Long
5 COVID research of the National Institutes of
6 Health, but does not affect the responsibility of any
7 of the national research institutes or centers with re-
8 spect to the programs or projects of such institutes
9 and centers; and

10 “(6) annually prepare and submit to the Direc-
11 tor of NIH for review and transmittal by the Direc-
12 tor of NIH to the President and to Congress a
13 budget estimate for carrying out the plan for the up-
14 coming fiscal year.

15 “(d) CONSULTATION.—In developing, implementing,
16 reviewing, and prioritizing elements of the research plan
17 under this section, the Director of the Program shall con-
18 sult, as appropriate with—

19 “(1) representatives of other Federal agencies
20 involved in Long COVID research, including the
21 Centers for Disease Control and Prevention, the
22 Agency for Healthcare Research and Quality, and
23 the Administration for Community Living;

24 “(2) the Long COVID Research Advisory
25 Board established under section 499B–4;

1 “(3) the Office of Long COVID Research and
2 Practice of the Department of Health and Human
3 Services;

4 “(4) leading scientific experts on Long COVID;
5 and

6 “(5) independent, patient-led organizations or
7 advocacy groups representing patients with Long
8 COVID and other infection-associated chronic condi-
9 tions with similar phenotypes, and the families of
10 such patients.

11 “(e) REPORT.—The Director of the Program shall
12 submit the research plan developed under subsection (a),
13 and updates to such plan, to—

14 “(1) the Committee on Health, Education,
15 Labor, and Pensions of the Senate;

16 “(2) the Committee on Energy and Commerce
17 of the House of Representatives;

18 “(3) the Secretary;

19 “(4) the Office of Long COVID Research and
20 Practice of the Department of Health and Human
21 Services; and

22 “(5) the Director of NIH, who shall post the
23 plan, and updates to the plan, on the website of the
24 National Institutes of Health.

1 **“SEC. 499B-2. EXPEDITED LONG COVID RESEARCH.**

2 “(a) IN GENERAL.—The Director of NIH shall estab-
3 lish a process to expedite the award of grants, contracts,
4 and cooperative agreements for research projects con-
5 ducted or supported by the National Institutes of Health
6 and relating to Long COVID.

7 “(b) REQUIREMENTS FOR MAKING EXTERNAL
8 FUNDING AVAILABLE.—With respect to programs of
9 grants, contracts, and cooperative agreements described in
10 subsection (a), the Director of NIH shall—

11 “(1) make publicly available the deadlines for
12 submitting applications for such programs, and en-
13 sure that such deadlines provide applicants with suf-
14 ficient time from the date of the announcement for
15 such grant, contract, and cooperative agreement to
16 submit an application;

17 “(2) ensure that applicants receive a final deci-
18 sion on their applications within 120 days of submis-
19 sion; and

20 “(3) with respect to applications that are de-
21 nied, provide a written explanation to the applicant
22 on the reasons for the denial.

23 “(c) EVALUATION OF GRANT APPLICATIONS.—In
24 making a determination to award a grant, contract, and
25 cooperative agreement for research projects described in
26 subsection (a), the Director of NIH shall—

1 “(1) give priority to research that—

2 “(A) tests the outcomes of existing drug
3 and device interventions in patients with Long
4 COVID;

5 “(B) focuses on identifying interventions
6 for pediatric patients with Long COVID;

7 “(C) aids in the development of new inter-
8 ventions that have evidence to suggest effective-
9 ness in treating or curing Long COVID; or

10 “(D) includes institutions that represent,
11 or have a successful track record of providing
12 equitable care or services to, historically under-
13 served communities;

14 “(2) consider research that has the ability to
15 begin interventions in a timely manner;

16 “(3) consider research that uses decentralized
17 trials or remote monitoring techniques for data col-
18 lection; and

19 “(4) consider research that includes patients
20 with other infection-associated chronic conditions
21 with similar phenotypes, such as myalgic
22 encephalomyelitis/chronic fatigue syndrome, postural
23 orthostatic tachycardia syndrome, and post-treat-
24 ment Lyme disease syndrome/persistent Lyme dis-
25 ease.

1 “(d) REASONABLE PRICING.—In awarding contracts,
2 grants, and cooperative agreements for research projects
3 described in subsection (a) that relates to the development
4 of a drug or device for the potential treatment or manage-
5 ment of Long COVID, or identifying a new indication or
6 use specific to the treatment or management of Long
7 COVID in a drug or device that is already approved or
8 cleared by the Food and Drug Administration, the Direc-
9 tor of NIH shall include terms and conditions requiring
10 that the price of such a drug or device for purposes of
11 procurement by the Federal Government or if sold on the
12 commercial market, whether procured from, or sold by, the
13 recipient of such Federal award or another person—

14 “(1) is fair and reasonable, taking into ac-
15 count—

16 “(A) the value of the drug and device to
17 the public health, including the impact of the
18 price on access to the drug or device;

19 “(B) the costs incurred by the Federal
20 Government in research and development of the
21 drug or device;

22 “(C) the costs incurred by the recipient of
23 the award in research and development of the
24 drug or device, and the costs of manufacturing
25 such drug or device;

1 “(D) whether the drug or device provided
2 a significant improvement in health outcomes,
3 compared to other therapies available at the
4 time of its approval or authorization;

5 “(E) the cumulative expected global reve-
6 nues generated by the drug or device; and

7 “(F) other factors, as the Secretary deter-
8 mines appropriate; and

9 “(2) does not exceed the lowest price charged
10 for such drug or device, among Canada, France,
11 Germany, Italy, Japan, and the United Kingdom.

12 “(e) CONSULTATION.—In making a determination to
13 award a grant, contract, or cooperative agreement for re-
14 search projects relating to Long COVID, the Director of
15 NIH shall consult with the Long COVID Research Advi-
16 sory Board. Members of the Long COVID Research Advi-
17 sory Board shall provide a recommendation on any final
18 funding decisions. If the Director of NIH makes a decision
19 that is different than the recommendation, the Director
20 of NIH shall provide a written justification for the deci-
21 sion within 5 days.

22 **“SEC. 499B-3. SCIENTIFIC REVIEW GROUP.**

23 “(a) IN GENERAL.—In order to ensure high quality,
24 rigorous scientific review of applications for grants, con-
25 tracts, and cooperative agreements described in section

1 499B–2(a), consistent with section 492, the Director of
2 NIH shall establish a scientific review group on Long
3 COVID and other infection-associated chronic conditions,
4 and shall convene a group of leading scientific experts to
5 serve on such group, for terms of up to 5 years.

6 “(b) DUTIES.—The scientific research group shall
7 conduct an initial review of applications for grants, con-
8 tracts, and other cooperative agreements described in sec-
9 tion 499B–2(a), and submit a funding recommendation to
10 the Director of NIH for final determination.

11 **“SEC. 499B–4. LONG COVID RESEARCH PROGRAM ADVISORY**
12 **BOARD.**

13 “(a) IN GENERAL.—The Director of NIH shall estab-
14 lish the Long COVID Research Program Advisory Board
15 (referred to in this section as the ‘Advisory Board’).

16 “(b) MEMBERSHIP.—

17 “(1) IN GENERAL.—The Advisory Board shall
18 be comprised of 18 members, including appointed
19 members and nonvoting ex officio members, as fol-
20 lows:

21 “(A) The Secretary shall conduct a nomi-
22 nation process that allows for public input on
23 nominees. The Secretary shall appoint nomi-
24 nated individuals, giving particular consider-
25 ation to individuals from backgrounds that rep-

1 resent the diversity of the Long COVID popu-
2 lation, with an emphasis on patients who face
3 the highest disease burden. Individuals so ap-
4 pointed shall include the following:

5 “(i) 10 members who are scientists,
6 physicians, and other health care profes-
7 sionals, who are not officers or employees
8 of the Federal Government, and who have
9 primary expertise in Long COVID and
10 other infection-associated chronic condi-
11 tions, with consideration given to such in-
12 dividuals with expertise in pediatric popu-
13 lations.

14 “(ii) 5 members who live with Long
15 COVID.

16 “(iii) 1 member who is a caregiver to
17 an individual with Long COVID.

18 “(iv) 2 members who are employed by
19 the National Institutes of Health and have
20 expertise in Long COVID research.

21 “(B) The following shall be ex officio mem-
22 bers of the Advisory Board:

23 “(i) A representative of the Long
24 COVID Research Program established
25 under section 499.

1 “(ii) A representative of the National
2 Institutes of Health.

3 “(iii) A representative of the National
4 Institutes of Neurological Disorders and
5 Stroke.

6 “(iv) A representative of the National
7 Heart, Lung, and Blood Institute.

8 “(v) A representative of the National
9 Institute of Allergy and Infectious Dis-
10 eases.

11 “(vi) A representative of the Office of
12 the Assistant Secretary for Health.

13 “(vii) A representative of the Centers
14 for Disease Control and Prevention.

15 “(viii) A representative of the Admin-
16 istration for Community Living.

17 “(ix) A representative of the Agency
18 for Healthcare Research and Quality.

19 “(x) Representatives of any other
20 agency or office of the Department of
21 Health and Human Services that the Sec-
22 retary determines appropriate for the Advi-
23 sory Board to carry out its function.

24 “(2) ENGAGEMENT WITH ORGANIZATIONS.—In
25 appointing individuals to the Advisory Board, the

1 Secretary shall engage with leading scientific experts
2 on Long COVID and independent, patient-led orga-
3 nizations of advocacy groups representing Long
4 COVID patients.

5 “(c) COMPENSATION.—Ex officio members of the Ad-
6 visory Board who are officers or employees of the Federal
7 Government shall not receive any compensation for service
8 on the Advisory Board. Non-Federal members of the Advi-
9 sory Board may receive, for each day (including travel
10 time) they are engaged in the performance of the functions
11 of the advisory committee, compensation at rates not to
12 exceed the daily equivalent to the annual rate of basic pay
13 for level III of the Executive Schedule under section 5314
14 of title 5, United States Code.

15 “(d) TERMS.—The term of office of an appointed
16 member of the Advisory Board is 5 years. Any member
17 appointed to fill a vacancy for an unexpired term shall
18 be appointed for the remainder of such term. A member
19 may serve after the expiration of the member’s term until
20 a successor has taken office. If a vacancy occurs in the
21 Advisory Board, the Secretary shall make an appointment
22 to fill the vacancy not later than 60 days from the date
23 the vacancy occurred.

1 “(e) CHAIR.—The members of the Advisory Board
2 shall select a chair from among the appointed members.
3 The term of the Office of Chair shall be 2 years.

4 “(f) MEETINGS.—

5 “(1) IN GENERAL.—The Advisory Board shall
6 meet at the call of the chairman or upon request of
7 the Director of the Program established under sec-
8 tion 499B, but not less often than monthly in the
9 first year after establishment, then not less often
10 than 6 times a year for each subsequent year. The
11 meetings of the Advisory Board may be held vir-
12 tually.

13 “(2) PURPOSE.—Of the meetings held, one or
14 more shall be held to address research priorities of
15 the National Institutes of Health relating to Long
16 COVID.

17 “(3) PUBLICATION OF SUMMARY.—For each
18 meeting held, the Director of NIH shall post on the
19 website of the National Institutes of Health a sum-
20 mary of the proceedings.

21 “(g) DUTIES.—The Advisory Board shall, subject to
22 the direction and supervision of the Director of NIH—

23 “(1) review, approve, and evaluate the imple-
24 mentation of the research plan issued under section
25 499B–1, and advise in updating the plan;

1 “(2) provide guidance to the Director of the
2 Program established under section 499B with re-
3 spect to appropriate research activities to be under-
4 taken regarding the clinical treatment of Long
5 COVID, which may include—

6 “(A) research on interventions for pre-
7 venting, treating, and understanding the mech-
8 anisms of Long COVID;

9 “(B) research on the effectiveness of treat-
10 ing Long COVID with drugs that are not yet
11 approved by the Food and Drug Administration
12 for the treatment of Long COVID;

13 “(C) reviewing ongoing publicly- and pri-
14 vately-supported research on treatments for
15 Long COVID;

16 “(D) issue and make available to health
17 care professionals and the public reports de-
18 scribing and evaluating research described in
19 subparagraphs (A), (B), and (C); and

20 “(E) convene accessible meetings for the
21 purpose of determining the recommendations
22 which may inform development of clinical guide-
23 lines by health care provider organizations; and

24 “(3) engage in other necessary activities to con-
25 tribute to the National Institutes of Health’s overall

1 research priorities related to Long COVID, and en-
2 sure accountability, transparency, and communica-
3 tion of results of the Program established under sec-
4 tion 499B.

5 **“SEC. 499B-5. DATA SYSTEM AND CLEARINGHOUSE ON RE-**
6 **SEARCH INFORMATION.**

7 “(a) DATA SYSTEM.—

8 “(1) IN GENERAL.—The Director of the Na-
9 tional Institutes of Health, in consultation with the
10 Director of the Program established under section
11 499B and the Director of the National Library of
12 Medicine shall establish, maintain, and operate a
13 data system for the collection, storage, analysis, re-
14 trieval, and timely dissemination of primary data re-
15 garding research on Long COVID that is conducted
16 or supported by the Program. Information from the
17 data system shall be available through information
18 systems available to health care professionals and
19 providers, researchers, and members of the public.

20 “(2) REGISTRY.—

21 “(A) IN GENERAL.—The data system es-
22 tablished under paragraph (1) shall include a
23 registry of clinical trials of experimental treat-
24 ments that have been developed for research on
25 Long COVID. Such registry shall include infor-

1 mation on patient eligibility criteria, including
2 the definition of Long COVID, and, as applica-
3 ble, demographic information, including sex,
4 age, disability status, ethnicity, and race, and
5 the location of the trial site or sites.

6 “(B) SUBMISSION OF INFORMATION.—
7 Principal investigators of trials described in
8 subparagraph (A) shall provide such informa-
9 tion to the registry not later than 30 days after
10 public announcement of the clinical trial. Once
11 a trial has been completed, the principal investi-
12 gator shall provide the registry with information
13 pertaining to the results, including potential
14 toxicities or adverse effects associated with the
15 experimental treatment or treatments evalu-
16 ated.

17 “(C) PUBLIC AVAILABILITY.—The registry
18 described in this paragraph shall be made avail-
19 able to researchers and the general public, in a
20 machine-readable format.

21 “(b) CLEARINGHOUSE.—The Director of NIH, in
22 consultation with the Director of the Program and with
23 the National Library of Medicine, shall establish, main-
24 tain, and operate a program to provide information on re-

1 search and prevention activities of the national research
2 institutes that relate to research on Long COVID.

3 **“SEC. 499B–6. APPROPRIATIONS.**

4 “For purposes of carrying out this part, there are ap-
5 propriated, out of amounts in the Treasury not otherwise
6 appropriated, \$1,000,000,000 for each of fiscal years
7 2025 through 2034, to remain available until expended.”.

8 **TITLE II—PUBLIC HEALTH RE-**
9 **SEARCH, SURVEILLANCE AND**
10 **RELATED ACTIVITIES**

11 **SEC. 201. LONG COVID PROGRAMS.**

12 Title III of the Public Health Service Act (42 U.S.C.
13 241 et seq.) is amended by adding at the end the fol-
14 lowing:

15 **“PART X—LONG COVID ACTIVITIES**

16 **“SEC. 399PP. PUBLIC HEALTH SURVEILLANCE OF LONG**
17 **COVID AND INFECTION-ASSOCIATED CHRON-**
18 **IC CONDITIONS.**

19 “(a) **IN GENERAL.**—The Secretary, acting through
20 the Director of the Centers for Disease Control and Pre-
21 vention, shall establish or continue, as applicable, surveil-
22 lance activities to better understand the burden and sever-
23 ity of Long COVID and related infection-associated chron-
24 ic conditions, with specific consideration given to vulner-

1 able populations, such as children. In carrying out this
2 section, the Secretary shall—

3 “(1) collect data on the incidence, prevalence,
4 and severity of Long COVID and related infection-
5 associated chronic conditions;

6 “(2) monitor for Long COVID and Long
7 COVID-like conditions, as appropriate, to enable
8 early intervention and identification of factors asso-
9 ciated with severity of symptoms;

10 “(3) compile, and make publicly available, in
11 accessible formats, Long COVID data collected
12 under paragraph (1);

13 “(4) develop and disseminate best practices for
14 conducting surveillance for State, local, and Tribal
15 public health officials, and other relevant public
16 health stakeholders;

17 “(5) provide technical assistance to inter-
18 national organizations, as applicable, regarding the
19 monitoring of Long COVID; and

20 “(6) conduct additional surveillance activities,
21 as the Secretary determines appropriate, to better
22 understand the burden and severity of Long COVID.

23 “(b) AUTHORIZATION OF APPROPRIATIONS.—For
24 purposes of carrying out this section, there are authorized

1 to be appropriated \$32,000,000 for each of fiscal years
2 2025 through 2034.

3 **“SEC. 399PP-1. PUBLIC HEALTH PROGRAMMING.**

4 “(a) IN GENERAL.—The Secretary, acting through
5 the Director of the Centers for Disease Control and Pre-
6 vention, shall make grants to State, local, and Tribal
7 health departments for the purpose of carrying out activi-
8 ties related to Long COVID.

9 “(b) USE OF FUNDS.—A State, local, or Tribal
10 health department that receives a grant under subsection
11 (a) may use funds received through such grant to—

12 “(1) provide training on the identification of
13 Long COVID to clinicians, public health experts,
14 and other relevant health care professionals;

15 “(2) link individuals with Long COVID to care,
16 as appropriate and applicable;

17 “(3) support the development and dissemination
18 of public information and educational materials on
19 Long COVID, including materials to address misin-
20 formation and disinformation;

21 “(4) support laboratory capacity for screening
22 and diagnosis of Long COVID and associated symp-
23 toms; and

1 “(5) build, maintain, and sustain jurisdiction-
2 level infrastructure related to preparedness for post-
3 infectious syndromes.

4 “(c) AUTHORIZATION OF APPROPRIATIONS.—For
5 purposes of carrying out this section, there are authorized
6 to be appropriated \$45,000,000 for each of fiscal years
7 2025 through 2034.

8 **“SEC. 399PP-2. NATIONAL PUBLIC EDUCATION CAMPAIGN**
9 **ON LONG COVID.**

10 “(a) IN GENERAL.—The Secretary, acting through
11 the Director of the Centers for Disease Control and Pre-
12 vention, and in collaboration with national, State, local,
13 and Tribal public health partners, shall develop a public
14 education campaign for patients, families, and caregivers
15 to educate and increase awareness about Long COVID in
16 children and adults. Such campaign shall include informa-
17 tion on—

18 “(1) the signs and symptoms of Long COVID;

19 “(2) how to prevent and seek treatment for
20 Long COVID;

21 “(3) self-management tools and support serv-
22 ices; and

23 “(4) other topics, as the Secretary determines
24 appropriate.

1 “(b) CONSULTATION.—In developing materials for
2 the campaign, the Secretary shall consult with inde-
3 pendent, patient-led organizations or advocacy groups rep-
4 resenting Long COVID patients and their families and
5 other relevant stakeholders.

6 “(c) ACCESSIBILITY.—The public education cam-
7 paign under this section shall be made available in mul-
8 tiple languages, including American Sign Language.

9 “(d) AUTHORIZATION OF APPROPRIATIONS.—For
10 purposes of carrying out this section, there are authorized
11 to be appropriated \$21,500,000 for each of fiscal years
12 2025 through 2029.

13 **“SEC. 399PP-3. PROVIDER EDUCATION.**

14 “(a) IN GENERAL.—The Secretary shall—

15 “(1) develop and make publicly available best
16 practices for coordinated, multidisciplinary care for
17 individuals with Long COVID;

18 “(2) develop, update, as appropriate, and make
19 publicly available clinical guidance and provider edu-
20 cation materials, including for providers working
21 with pediatric populations; and

22 “(3) facilitate provider education on Long
23 COVID signs, symptoms, maintenance, and treat-
24 ment, including through technology-enabled collabo-
25 rative learning.

1 “(b) AUTHORIZATION OF APPROPRIATIONS.—For the
2 purpose of carrying out this section, there are authorized
3 to be appropriated \$3,000,000 for each of fiscal years
4 2025 through 2034.”.

5 **SEC. 202. REHABILITATION RESEARCH AND TRAINING CEN-**
6 **TER ON LONG COVID AMONG PEOPLE WITH**
7 **DISABILITIES.**

8 (a) IN GENERAL.—Section 240(b)(2)(C) of the Reha-
9 bilitation Act of 1973 (29 U.S.C. 764(b)(2)(C)) is amend-
10 ed—

11 (1) in clause (v), by striking “; and” and insert-
12 ing a semicolon;

13 (2) in clause (vi), by striking the period and in-
14 serting “; and”; and

15 (3) by adding at the end the following:

16 “(vii) applied research regarding evi-
17 dence-based treatments, services, and sup-
18 ports for individuals with disabilities with
19 Long COVID or other infection-associated
20 chronic conditions.”.

21 (b) AUTHORIZATION OF APPROPRIATIONS.—To carry
22 out the amendment made by subsection (b), there are au-
23 thorized to be appropriated to the Director of the National
24 Institute on Disability, Independent Living, and Rehabili-

1 tation Research, \$10,000,000 for the period of fiscal years
2 2025 through 2029.

3 **SEC. 203. CLINICAL OUTCOMES ASSESSMENTS.**

4 (a) IN GENERAL.—The Secretary of Health and
5 Human Services, acting through the Commissioner of
6 Food and Drugs, shall establish or continue the develop-
7 ment and validation of clinical outcomes assessments to
8 support regulatory decision making for drugs, including
9 biological products, and devices used to treat Long
10 COVID.

11 (b) AUTHORIZATION OF APPROPRIATIONS.—For pur-
12 poses of carrying out this section, there are authorized to
13 be appropriated \$9,000,000 for each of fiscal years 2025
14 through 2034.

15 **SEC. 204. ELECTRONIC REPORTING FORM.**

16 (a) IN GENERAL.—The Secretary of Health and
17 Human Services, acting through the Commissioner of
18 Food and Drugs, shall establish or continue the develop-
19 ment, refinement, and maintenance of a Long COVID
20 electronic reporting form for patients to identify current
21 treatments and treatments under development for Long
22 COVID.

23 (b) AUTHORIZATION OF APPROPRIATIONS.—For pur-
24 poses of carrying out this section, there are authorized to

1 be appropriated \$16,600,000 for each of fiscal years 2025
2 through 2034.

3 **SEC. 205. LONG COVID CARE NETWORK.**

4 (a) IN GENERAL.—The Secretary of Health and
5 Human Services, acting through the Director of the Agen-
6 cy for Healthcare Research and Quality, shall develop, or
7 continue to support, multidisciplinary Long COVID clinics
8 to provide access to comprehensive, coordinated care for
9 individuals with Long COVID, particularly underserved
10 populations that are disproportionately impacted by the
11 effects of Long COVID.

12 (b) AUTHORIZATIONS OF APPROPRIATIONS.—For
13 purposes of carrying out this section, there are authorized
14 to be appropriated \$10,000,000 for each of fiscal years
15 2025 through 2034.

16 **SEC. 206. RESEARCH ON LONG COVID BEST PRACTICES.**

17 (a) IN GENERAL.—The Secretary of Health and
18 Human Services, in coordination with the Director of the
19 Agency for Healthcare Research and Quality, shall de-
20 velop, test, synthesize, and disseminate best practices and
21 decision support tools related to the clinical care organiza-
22 tion, delivery, and integration of clinical and social services
23 for Long COVID and other infection-associated chronic
24 conditions.

1 (b) AUTHORIZATION OF APPROPRIATIONS.—For the
2 purposes of carrying out this section, there are authorized
3 to be appropriated \$10,000,000 for each of fiscal years
4 2025 through 2034.