

103^D CONGRESS
2^D SESSION

H. R. 4370

To establish the AIDS Cure Project.

IN THE HOUSE OF REPRESENTATIVES

MAY 10, 1994

Mr. NADLER (for himself, Mr. DELLUMS, Ms. VELÁZQUEZ, Mr. OWENS, and Mr. MILLER of California) introduced the following bill; which was referred to the Committee on Energy and Commerce

A BILL

To establish the AIDS Cure Project.

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 “AIDS Cure Act”.

5 **SEC. 2. ESTABLISHMENT OF PROJECT TO CURE AIDS.**

6 (a) IN GENERAL.—The President shall in accordance
7 with this Act direct the Secretary of Health and Human
8 Services to establish a project for the purpose of develop-
9 ing a cure for acquired immune deficiency syndrome (in
10 this Act referred to as “AIDS”). The program may not
11 be administered by any officer or employee of the National

1 Institutes of Health. Subject to the preceding sentence,
2 the Governing Council shall designate an individual to
3 serve as a liaison between the Governing Council of the
4 project (established under section 4(a)), the Secretary,
5 and the President.

6 (b) DEFINITION.—For purposes of this Act, the term
7 “cure”, with respect to AIDS, means any and all ap-
8 proaches which will ensure a well-functioning immune sys-
9 tem and a normal life span with a reasonable quality of
10 life.

11 (c) CERTAIN REQUIREMENTS.—The Governing
12 Council, in carrying out the project under subsection (a),
13 shall ensure that the following requirements are met:

14 (1) The project shall pursue comprehensive
15 basic science investigations, based on diverse theo-
16 ries and schools of thought which elucidate the
17 pathogenesis of AIDS.

18 (2) The project shall identify, based on this
19 work, all promising curatives and oversee their time-
20 ly and adequate testing through the extraordinary
21 powers detailed in section 5.

22 **SEC. 3. OPEN AND PRODUCTIVE RESEARCH PATHS.**

23 The Governing Council, in carrying out the project
24 under section 2, shall ensure that the following require-
25 ments are met:

1 (1)(A) Thorough consideration shall be given to
2 both conventional and other medical approaches and
3 scientific theories, and researchers representing di-
4 vergent approaches shall be on the primary research
5 staff as well as be contributing researchers.

6 (B) The project shall aggressively pursue re-
7 search into all areas of AIDS pathogenesis, includ-
8 ing but not limited to—

9 (i) virological/immunological theories about
10 how immune system damage occurs, including
11 understudied approaches; and

12 (ii) theories about co-factors which may
13 precede, activate or even substitute for HIV in
14 the process of immune system damage leading
15 to AIDS.

16 (C) Examination shall be given to the full spec-
17 trum of pathogenesis theories, from those maintain-
18 ing that HIV is the sole and sufficient cause to
19 those considering HIV a primary cause together
20 with co-factors to those believing that HIV does not
21 necessarily play a causative role.

22 (D) Further work shall be done on the potential
23 role of recreational drugs and environmental toxins
24 in progression. The role of nutrition, exercise, and
25 adequate primary health care must also be re-

1 searched in the project. Psychoneuroimmunology
2 and the connections between psychological stress and
3 immune compromise shall also be studied.

4 (E) A diversity of theories should be developed
5 and tested through both laboratory experiments and
6 epidemiological research, including careful examina-
7 tion of cases of people with HIV and AIDS and
8 interviews with people with HIV and AIDS and their
9 care providers including but not limited to primary
10 care clinicians, gynecologists, nutritionists, alter-
11 native or holistic practitioners, and mental health
12 workers.

13 (F) All research funded in this Act shall be con-
14 ducted in conformity with the ethics and privacy of
15 current medical research.

16 (G) Researchers shall study epidemiological and
17 blood studies of long-term survivors from diverse
18 populations to attempt to isolate the factors that
19 have sustained them.

20 (H) Consideration shall be given to the
21 hypotheses and results obtained in other countries.
22 Researchers from other countries shall be invited to
23 participate in the project. This may include agree-
24 ments with another country to reassign particular
25 researchers to the project for an indefinite commit-

1 ment. The project's progress shall not await the con-
2 clusion of such international agreements.

3 (2) The project's study of AIDS pathogenesis
4 and manifestations must focus on all populations of
5 people with AIDS and HIV. Equal consideration
6 shall be given to the differences between these popu-
7 lations as to their similarities or norms. This in-
8 cludes, but is not limited to people of all age groups
9 (including children and seniors), gender, sexual ori-
10 entation, women (regardless of reproductive health
11 status), gay men, lesbians, people of color (of var-
12 ious affected national-cultural groups), injection
13 drug users, prison inmates, people with hemophilia
14 and people with inadequate medical care or nutrition
15 (or both), and persons with disabilities and chronic
16 conditions related to AIDS.

17 (3) Basic science investigations and therapeutic
18 results shall be geared to people at every point on
19 the spectrum of AIDS and HIV—from the sickest to
20 the healthiest. Saving people considered near death
21 must be considered as important as early interven-
22 tion.

23 (4) Information generated by the project shall
24 be made freely available worldwide to researchers,
25 health care providers, people with AIDS and HIV

1 and their advocates as soon as it is available, with-
2 out being inhibited by professional publication prac-
3 tices. Funds shall be available as needed for the dis-
4 semination and translation of project materials.

5 (5) Curatives and therapies ultimately released
6 due primarily to project research shall not result in
7 financial gain to any private organization, and shall
8 be made available to all affected people worldwide
9 regardless of ability to pay. The project shall be re-
10 sponsible for establishing a mechanism for inter-
11 national funding and distribution of any such
12 curatives.

13 **SEC. 4. EFFICIENT AND COOPERATIVE MANAGEMENT OF**
14 **PROJECT.**

15 (a) GOVERNING COUNCIL.—

16 (1) IN GENERAL.—The project under section 2
17 shall be governed by, not merely advised by, a coun-
18 cil composed of scientists and clinicians representing
19 divergent approaches, and people with AIDS and
20 HIV, and their advocates, from all affected commu-
21 nities. This council shall set policy and oversee re-
22 search priorities, ethical standards, conflict of inter-
23 est rules and hiring of researchers, and administra-
24 tors.

1 (2) CERTAIN AUTHORITIES.—The Secretary
2 shall ensure that the following requirements are met
3 with respect to the council under paragraph (1):

4 (A) The council shall be composed of sci-
5 entists representing divergent approaches, clini-
6 cians with both research and community-based
7 experience and people with AIDS and HIV and
8 their advocates.

9 (B) The council shall have at least 21
10 members in order to adequately represent di-
11 verse communities, opinions and disciplines.
12 People with AIDS and HIV from diverse com-
13 munities shall be in the majority to ensure that
14 the project staff are ultimately accountable to
15 people directly affected by the course and out-
16 come of the research. Council members shall
17 step down and be replaced by new members on
18 a regular basis.

19 (C) The Council shall set policy for and
20 oversee research priorities. It shall develop
21 guidelines for and oversee the hiring of primary
22 research staff, ensuring both high quality (sci-
23 entific credentials and experience) and a diver-
24 sity of disciplines and perspectives (including
25 alternative” or holistic approaches). Having

1 pursued specific AIDS theories shall not be a
2 necessary prerequisite for hiring. The Council
3 shall have the power to create new research po-
4 sitions when necessary and to remove scientists
5 from their positions after due process and ap-
6 propriate review of their work.

7 (D) The Council shall be charged with
8 evaluating the work of the project, as well as
9 the pace of the research, to ensure that it
10 matches the urgency of the epidemic. Initially,
11 and throughout the life of the project, the
12 Council, in cooperation with the primary re-
13 search staff, shall solicit and evaluate theories
14 developed outside the project. It shall direct the
15 project scientists to evaluate and respond to de-
16 serving proposals and to devise new research
17 plans where desirable.

18 (E) The Council shall adopt strict, detailed
19 codes governing medical ethics and conflicts of
20 interest and shall monitor compliance with
21 these codes. Project scientists shall report di-
22 rectly to the council about the progress of their
23 work in a manner to be determined by the
24 council. The Council shall report to the Presi-

1 dent through the liaison about the progress of
2 the project.

3 (F) Council meetings, including those at
4 which all decisions are made, shall be public
5 and shall be held at least quarterly, with time
6 allotted for public comment. In addition, the
7 Council shall hold an annual public hearing on
8 its priorities and progress. A complete report of
9 the project's goals and accomplishments shall
10 be updated by the Council, submitted to the
11 President and released to the public at least
12 once quarterly. The Council shall evaluate its
13 structure and process at least once per year and
14 make changes which allow it to function more
15 effectively.

16 (G) Members of the Governing Council will
17 not be paid employees of the project. However,
18 the Governing Council will be provided with an
19 operating budget, including but not limited to,
20 the following purposes: Support staff; creation
21 and dissemination of reports and other mate-
22 rials; funds for transportation and other per
23 diem expenses; and funds for convening public
24 meetings.

1 (b) REQUIREMENTS.—The Governing Council, in car-
2 rying out the project under section 2, shall ensure that
3 the following requirements are met:

4 (1) The project shall establish a primary loca-
5 tion for its work, in an area with a high incidence
6 of AIDS. All primary research staff shall work at
7 that location; contributing researchers located
8 around the world shall interact via video teleconfer-
9 encing, an international computer network, and reg-
10 ularly scheduled face-to-face meetings.

11 (2) The National Institutes of Health's existing
12 AIDS research programs shall be maintained. All
13 National Institutes of Health basic science research
14 supplementary to that done by the project shall be
15 performed cooperatively with the project in coordina-
16 tion with the Office of AIDS Research.

17 (3)(A) All primary research staff and adminis-
18 trators shall be financially compensated only by the
19 project and may not have conflicts of interests with
20 private or public organizations (including but not
21 limited to universities, pharmaceutical companies,
22 and private research organizations).

23 (B) All primary research staff and administra-
24 tors shall be required to suspend their relationship
25 with any private or public organizations for the du-

1 ration of their association with the project. These re-
2 quirements shall include full-time, part-time, or con-
3 sultant positions with a private or public organiza-
4 tion or other government agencies, and the suspen-
5 sion would include employment, consulting or board
6 membership fees, and stock or business ownership.

7 (C) The Governing Council members shall be
8 required to suspend their relationship with for-profit
9 organizations which represent a conflict of interest.

10 (4) The project shall be funded by public, not
11 private monies. Appropriations for the project shall
12 not be diverted from other health research, health
13 care, or human service programs.

14 (5) The project shall, in addition to basic re-
15 search investigations, operate an on-site clinic to
16 conduct small scale research trials with human par-
17 ticipants if such trials are crucial for testing
18 hypotheses related to its basic research.

19 (c) COORDINATING COMMITTEE.—The Governing
20 Council shall ensure that a coordinating committee is es-
21 tablished for the project under section 2, in accordance
22 with the following:

23 (1) The community of scientists selected for the
24 project shall elect three of their members to serve as
25 the coordinating committee for the project, and de-

1 termine whether these positions should be perma-
2 nent or rotating.

3 (2) The coordinating committee shall be respon-
4 sible for facilitating communication among the dif-
5 ferent scientists working on the project, for evaluat-
6 ing the progress of its work, and for convening the
7 entire staff on a regular schedule (or when nec-
8 essary) to evaluate the progress of the project as a
9 whole, identify gaps in research, reevaluate the
10 project's direction, and to consider newly developed
11 theories emanating from both within and outside the
12 project.

13 (3) The coordinating committee shall also be re-
14 sponsible for keeping the Governing Council in-
15 formed of the progress of the project's work, at
16 times and in a manner to be determined by the Gov-
17 erning Council. The coordinating committee shall
18 also make decisions regarding the hiring of research
19 associates, technical staff, purchases of equipment
20 and other day-to-day needs.

21 (4) The first task of the coordinating committee
22 shall be to facilitate a preliminary review of all exist-
23 ing pathogenesis hypotheses, as well as other rel-
24 evant information about AIDS pathogenesis. At the
25 end of this review, which shall last no longer than

1 3 months, the primary research staff shall collec-
2 tively develop plans for evaluating and testing each
3 of the viable hypotheses, including timelines for eval-
4 uating the progress of this work, and submit these
5 plans to the Governing Council for review and com-
6 ments.

7 (d) SELECTION OF GOVERNING COUNCIL.—

8 (1) IN GENERAL.—The Secretary of Health and
9 Human Services (HHS) shall convene a national
10 AIDS congress to make recommendations to the
11 President for selecting the Governing Council. The
12 AIDS congress will meet only once, and for the sole
13 purpose of nominating the initial Governing Council.
14 The Secretary of HHS shall solicit nominations from
15 a wide variety of sources, including, but not limited
16 to, each of the following: AIDS activist groups;
17 health care providers; AIDS advocacy organizations;
18 AIDS service organizations; community-based AIDS
19 research organization; biomedical researchers and
20 nonmainstream (including alternative or holistic)
21 medical organizations; and health care planning
22 agencies, who shall send their nominations for the
23 Governing Council to the AIDS congress. The AIDS
24 congress will make recommendations to the Presi-
25 dent for the Governing Council. The President shall

1 select the Governing Council based on the rec-
2 ommendations of the AIDS congress. The AIDS
3 congress will consist of 2 representatives chosen by
4 each of the HIV health services planning councils
5 under section 2602(b) of the Public Health Service
6 Act, chosen in a forum that is open to the public by
7 each of the HIV planning councils. The President
8 shall widely publicize the request for nominations.

9 (2) DATE CERTAIN FOR SELECTION.—In keep-
10 ing with the emergency nature of this project, the
11 nomination and selection process must be completed
12 within 3 months of the enactment of this Act.

13 **SEC. 5. EXTRAORDINARY POWERS.**

14 In carrying out the project under section 2, the Gov-
15 erning Council shall have extraordinary powers to carry
16 out the following:

17 (1)(A) Utilize, in cooperation with the agencies,
18 any and all existing United States Government fund-
19 ed research entities nationwide (including but not
20 limited to the AIDS Clinical Trial Group (ACTG),
21 the Community Program for Clinical Research on
22 AIDS (CPCRA)), and their facilities to clinically
23 test promising cures developed on the basis of its re-
24 search and to direct the manner in which such re-
25 search shall proceed, including staffing, participants,

1 location, and timing. Such research shall be funded
2 by the project.

3 (B) The project shall design its own protocols
4 and work with these existing clinical trial programs
5 to develop research designs and methods appropriate
6 to the project's goals, assuring that data gathered
7 by the NIH would accurately reflect the use of these
8 compounds in all populations and stages of illness.

9 (C) The project shall provide funding for these
10 clinical trials of its own compounds. In areas of con-
11 flict, the project shall have the power to implement
12 its goals.

13 (2) Exercise the right of eminent domain to
14 carry out the following:

15 (A)(i) Obtain from public and private orga-
16 nizations, with just compensation, samples of
17 all potential curatives and all data (excluding
18 medical records) regarding their development
19 (including safety and efficacy data) as well as
20 other information, materials, or products
21 deemed crucial to the project, but protect the
22 privacy of research subjects and the researcher
23 over which the project will be exercising the
24 right of eminent domain. The project shall use

1 its power of eminent domain only after reason-
2 able attempts at cooperation have failed.

3 (ii) To use eminent domain power, the
4 project must determine and identify whether
5 this research is essential information to the
6 project's research and that it is unavailable
7 other than through eminent domain.

8 (iii) Once obtaining through eminent do-
9 main the research of an outside researcher, the
10 project shall make all efforts to preserve appro-
11 priate recognition of the scientist's work. The
12 project shall also afford any researcher, whose
13 work is acquired through eminent domain, an
14 opportunity to participate in the project with
15 just compensation.

16 (B) If a drug company is found to be im-
17 peding or halting the development of a promis-
18 ing compound, the project shall first attempt to
19 work with the company to develop the needed
20 timetable for research and trials. A company
21 lacking the resources to develop a compound
22 shall have the option of selling the patent to the
23 project for just compensation, or allowing por-
24 tions of its development to be undertaken by
25 the project.

1 (C) If, however, a company refuses to co-
2 operate with the project by not releasing needed
3 data, or by withholding samples of requested
4 compounds, whether under development or not,
5 the project is authorized to use powers of emi-
6 nent domain to procure samples and data. The
7 project shall have the power to obtain the pat-
8 ents of such compounds if, after reasonable at-
9 tempts at cooperation, it finds that a company
10 will not develop a promising compound accord-
11 ing to an approved timeframe by the coordinat-
12 ing committee. After notification by the project
13 that this power will be used, a company shall
14 have 30 days in which to develop, for the
15 project's approval, a plan for accelerated devel-
16 opment of the compound to avoid losing exclu-
17 sive rights to the patent. Unless said companies
18 adhere to an approved timeframe and are forth-
19 coming with their data as such work proceeds,
20 then the project can implement clinical testing
21 for potential curatives by private companies.

22 (D) Use existing pharmaceutical company
23 facilities (with just compensation) for the pro-
24 duction of promising curatives to be utilized in
25 project research and, if effective, to produce

1 such curatives in sufficient amounts to be dis-
2 seminated to all people needing them.

3 **SEC. 6. PLANNING FUNDS.**

4 Funds shall be allocated immediately to be used for
5 planning of the project under section 2 (including creating
6 facilities, selection of staff, funding, structure, and sched-
7 ules), so that the project can begin functioning as soon
8 as is possible.

9 **SEC. 7. REAUTHORIZATION OF PROJECT.**

10 After 5 years of operation, the Congress shall have
11 the power to reauthorize the project under section 2.

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