§ 285. Purpose of Institute

The general purpose of the National Cancer Institute (hereafter in this subpart referred to as the "Institute") is the conduct and support of research, training, health information dissemination, and other programs with respect to the cause, diagnosis, prevention, and treatment of cancer, rehabilitation from cancer, and the continuing care of cancer patients and the families of cancer patients.


AMENDMENTS


§ 285a. National Cancer Program

The National Cancer Program shall consist of (1) an expanded, intensified, and coordinated cancer research program encompassing the research programs conducted and supported by the Institute and the related research programs of the other national research institutes, including an expanded and intensified research program for the prevention of cancer caused by occupational or environmental exposure to carcinogens, and (2) the other programs and activities of the Institute.

(July 1, 1944, ch. 373, title IV, § 411, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 832.)

§ 285a–1. Cancer control programs

The Director of the Institute shall establish and support demonstration, education, and other programs for the detection, diagnosis, prevention, and treatment of cancer and for rehabilitation and counseling respecting cancer. Programs established and supported under this section shall include—

(A) the detection, diagnosis, prevention, and treatment of cancer,
(B) the continuing care of cancer patients and the families of cancer patients, and
(C) rehabilitation and counseling respecting cancer, to physicians and other health professionals who provide care to individuals who have cancer;
(D) the demonstration of and the education of students of the health professions and health professionals in—
   (A) effective methods for the prevention and early detection of cancer and the identification of individuals with a high risk of developing cancer, and
   (B) improved methods of patient referral to appropriate centers for early diagnosis and treatment of cancer; and
   (C) the demonstration of new methods for the dissemination of information to the general public concerning the prevention, early detection, diagnosis, and treatment and control of cancer and information concerning unapproved and ineffective methods, drugs, and devices for the diagnosis, prevention, treatment, and control of cancer.

(July 1, 1944, ch. 373, title IV, § 412, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 832.)

§ 285a–2. Special authorities of Director

(a) Information and education program

(1) The Director of the Institute shall establish an information and education program to collect, identify, analyze, and disseminate on a timely basis, through publications and other appropriate means, to cancer patients and their families, physicians and other health professionals, and the general public, information on cancer research, diagnosis, prevention, and treatment (including information respecting nutrition programs for cancer patients and the relationship between nutrition and cancer). The Director of the Institute may take such action as may be necessary to ensure that all channels for the dissemination and exchange of scientific knowledge and information are maintained between the Institute and the public and between the Institute and other scientific, medical, and biomedical disciplines and organizations nationally and internationally.

(2) In carrying out paragraph (1), the Director of the Institute shall—

(A) provide public and patient information and education programs, providing information that will help individuals take personal steps to reduce their risk of cancer, to make them aware of early detection techniques and to motivate appropriate utilization of those techniques, to help individuals deal with cancer if it strikes, and to provide information to improve long-term survival;

(B) continue and expand programs to provide physicians and the public with state-of-the-art information on the treatment of particular forms of cancers, and to identify those clinical trials that might benefit patients while advancing knowledge of cancer treatment;

(C) assess the incorporation of state-of-the-art cancer treatments into clinical practice and the extent to which cancer patients receive such treatments and include the results of such assessments in the biennial reports required under section 284b of this title;

(D) maintain and operate the International Cancer Research Data Bank, which shall collect, catalog, store, and disseminate information insofar as feasible the results of cancer research and treatment undertaken in any country for the use of any person involved in cancer research and treatment in any country; and

(E) to the extent practicable, in disseminating the results of such cancer research and treatment, utilize information systems available to the public.

(b) National Cancer Program

The Director of the Institute in carrying out the National Cancer Program—

1 See References in Text note below.
(1) shall establish or support the large-scale production or distribution of specialized biological materials and other therapeutic substances for cancer research and set standards of safety and care for persons using such materials;

(2) shall, in consultation with the advisory council for the Institute, support (A) research in the cancer field outside the United States by highly qualified foreign nationals which can be expected to benefit the American people, (B) collaborative research involving American and foreign participants, and (C) the training of American scientists abroad and foreign scientists in the United States;

(3) shall, in consultation with the advisory council for the Institute, support appropriate programs of education and training (including continuing education and laboratory and clinical research training);

(4) shall encourage and coordinate cancer research by industrial concerns where such concerns evidence a particular capability for such research;

(5) may obtain (after consultation with the advisory council for the Institute and in accordance with section 3109 of title 5, but without regard to the limitation in such section on the period of service) the services of not more than one hundred and fifty-one experts or consultants who have scientific or professional qualifications;

(6)(A) may, in consultation with the advisory council for the Institute, acquire, construct, improve, repair, operate, and maintain laboratories, other research facilities, equipment, and such other real or personal property as the Director determines necessary;

(B) may, in consultation with the advisory council for the Institute, make grants for construction or renovation of facilities; and

(C) may, in consultation with the advisory council for the Institute, acquire, without regard to section 8141 of title 40, by lease or otherwise through the Administrator of General Services, buildings or parts of buildings in the District of Columbia or communities located adjacent to the District of Columbia for the use of the Institute for a period not to exceed ten years;

(7) may, in consultation with the advisory council for the Institute, appoint one or more advisory committees composed of such private citizens and officials of Federal, State, and local governments to advise the Director with respect to the Director's functions;

(8) may, subject to section 284(b)(2) of this title and without regard to section 3324 of title 31 and section 6101 of title 41, enter into such contracts, leases, cooperative agreements, as may be necessary in the conduct of functions of the Director, with any public agency, or with any person, firm, association, corporation, or educational institution; and

(9) shall, notwithstanding section 284(a) of this title, prepare and submit, directly to the President for review and transmittal to Congress, an annual budget estimate (including an estimate of the number and type of personnel needs for the Institute) for the National Cancer Program, after reasonable opportunity for comment (but without change) by the Secretary, the Director of NIH, and the Institute's advisory council.

Except as otherwise provided, experts and consultants whose services are obtained under paragraph (5) shall be paid or reimbursed, in accordance with title 5 for their travel to and from their place of service and for other expenses associated with their assignment. Such expenses shall not be allowed in connection with the assignment of an expert or consultant whose services are obtained under paragraph (5) unless the expert or consultant has agreed in writing to complete the entire period of the assignment or one year of the assignment, whichever is shorter, unless separated or reassigned for reasons which are beyond the control of the expert or consultant and which are acceptable to the Director of the Institute. If the expert or consultant violates the agreement, the money spent by the United States for such expenses is recoverable from the expert or consultant as a debt due the United States. The Secretary may waive in whole or in part a right of recovery under the preceding sentence.

(c) Pre-clinical models to evaluate promising pediatric cancer therapies

(1) Expansion and coordination of activities

The Director of the National Cancer Institute shall expand, intensify, and coordinate the activities of the Institute with respect to research on the development of preclinical models to evaluate which therapies are likely to be effective for treating pediatric cancer.

(2) Coordination with other institutes

The Director of the Institute shall coordinate the activities under paragraph (1) with similar activities conducted by other national research institutes and agencies of the National Institutes of Health to the extent that those Institutes and agencies have responsibilities that are related to pediatric cancer.


References in Text


Codification


Amendments


1993—Subsec. (b)(9). Pub. L. 103–43 struck out subpar. (A) designation and subpar. (B) which permitted Direc-
tor to receive from President and Office of Management and Budget directly all funds appropriated for Congress for obligation and expenditure by Institute.

§ 285a–3. National cancer research and demonstration centers

(a) Cooperative agreements and grants for establishing and supporting

(1) The Director of the Institute may enter into cooperative agreements with and make grants to public or private nonprofit entities to pay all or part of the cost of planning, establishing, or strengthening, and providing basic operating support for centers for basic and clinical research into, training in, and demonstration of advanced diagnostic, prevention, control, and treatment methods for cancer.

(2) A cooperative agreement or grant under paragraph (1) shall be entered into in accordance with policies established by the Director of NIH and after consultation with the Institute’s advisory council.

(b) Uses for Federal payments under cooperative agreements or grants

Federal payments made under a cooperative agreement or grant under subsection (a) of this section may be used for—

(1) construction (notwithstanding any limitation under section 289e of this title);

(2) staffing and other basic operating costs, including such patient care costs as are required for research;

(3) clinical training, including training for allied health professionals, continuing education for health professionals and allied health professions personnel, and information programs for the public respecting cancer; and

(4) demonstration purposes.

As used in this paragraph, the term “construction” does not include the acquisition of land, and the term “training” does not include research training for which Ruth L. Kirschstein National Research Service Awards may be provided under section 288 of this title.

(c) Period of support; additional periods

Support of a center under subsection (a) of this section may be for a period of not to exceed five years. Such period may be extended by the Director for additional periods of not more than five years each if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director and if such group has recommended to the Director that such period should be extended.

(d) Construction

Research centers under this section may not be considered centers of excellence for purposes of section 282(b)(10) of this title.


AMENDMENTS


EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

§ 285a–4. President’s Cancer Panel; establishment, membership, etc., functions

(a)(1) The President’s Cancer Panel (hereafter in this section referred to as the “Panel”) shall be composed of three persons appointed by the President who by virtue of their training, experience, and background are exceptionally qualified to appraise the National Cancer Program. At least two members of the Panel shall be distinguished scientists or physicians.

(2)(A) Members of the Panel shall be appointed for three-year terms, except that (i) any member appointed to fill a vacancy occurring prior to the expiration of the term for which the member’s predecessor was appointed shall be appointed only for the remainder of such term, and (ii) a member may serve until the member’s successor has taken office. If a vacancy occurs in the Panel, the President shall make an appointment to fill the vacancy not later than 90 days after the date the vacancy occurred.

(B) The President shall designate one of the members to serve as the chairman of the Panel for a term of one year.

(C) Members of the Panel shall each be entitled to receive the daily equivalent of the annual rate of basic pay in effect for grade GS–18 of the General Schedule for each day (including traveltime) during which they are engaged in the actual performance of duties as members of the Panel and shall be paid or reimbursed, in accordance with title 5, for their travel to and from their place of service and for other expenses associated with their assignment.

(3) The Panel shall meet at the call of the chairman, but not less often than four times a year. A transcript shall be kept of the proceedings of each meeting of the Panel, and the chairman shall make such transcript available to the public.

(b) The Panel shall monitor the development and execution of the activities of the National Cancer Program, and shall report directly to the President. Any delays or blockages in rapid execution of the Program shall immediately be
brought to the attention of the President. The Panel shall submit to the President periodic progress reports on the National Cancer Program and shall submit to the President, the Secretary, and the Congress an annual evaluation of the efficacy of the Program and suggestions for improvements, and shall submit such other reports as the President shall direct.

(July 1, 1944, ch. 373, title IV, § 415, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 835.)

TERMINATION OF REPORTING REQUIREMENTS

For termination, effective May 15, 2000, of provisions in subsec. (b) of this section relating to the requirement that the Panel submit to Congress an annual evaluation of the efficacy of the Program and suggestions for improvements, see section 3003 of Pub. L. 101–509, set out in a note under section 5376 of Title 5.

TERMINATION OF ADVISORY PANELS

Advisory panels established after Jan. 5, 1973, to terminate not later than the expiration of the 2-year period beginning on the date of their establishment, unless, in the case of a panel established by the President or an officer of the Federal Government, such panel is renewed by appropriate action prior to the expiration of such 2-year period, or in the case of a panel established by the Congress, its duration is otherwise provided by law. See sections 3(2) and 14 of Pub. L. 92–463, Oct. 6, 1972, 86 Stat. 770, 776, set out in the Appendix to Title 5.

TERMINATION OF ADVISORY PANELS

Advisory panels established after Jan. 5, 1973, to terminate not later than the expiration of the 2-year period beginning on the date of their establishment, unless, in the case of a panel established by the President or an officer of the Federal Government, such panel is renewed by appropriate action prior to the expiration of such 2-year period, or in the case of a panel established by the Congress, its duration is otherwise provided by law. See sections 3(2) and 14 of Pub. L. 92–463, Oct. 6, 1972, 86 Stat. 770, 776, set out in the Appendix to Title 5, Government Organization and Employees.


REFERENCES IN OTHER LAWS TO GS–16, 17, OR 18 PAY RATES

References in laws to the rates of pay for GS–16, 17, or 18, or to maximum rates of pay under the General Schedule, to be considered references to rates payable under specified sections of Title 5, Government Organization and Employees, see section 529 (title 1, § 181(c)(1)) of Pub. L. 101–509, set out in a note under section 5376 of Title 5.

§ 285a–5. Associate Director for Prevention; appointment; function

(a) There shall be in the Institute an Associate Director for Prevention to coordinate and promote the programs in the Institute concerning the prevention of cancer. The Associate Director shall be appointed by the Director of the Institute from individuals who because of their professional training or experience are experts in public health or preventive medicine.

(b) The Associate Director for Prevention shall prepare for inclusion in the biennial report made under section 284b of this title a description of the prevention activities of the Institute, including a description of the staff and resources allocated to those activities.

(July 1, 1944, ch. 373, title IV, § 416, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 836.)

REFERENCES IN TEXT


§ 285a–6. Breast and gynecological cancers

(a) Expansion and coordination of activities

The Director of the Institute, in consultation with the National Cancer Advisory Board, shall expand, intensify, and coordinate the activities of the Institute with respect to research on breast cancer, ovarian cancer, and other cancers of the reproductive system of women.

(b) Coordination with other institutes

The Director of the Institute shall coordinate the activities of the Director under subsection (a) of this section with similar activities conducted by other national research institutes and agencies of the National Institutes of Health to the extent that such Institutes1 and agencies have responsibilities that are related to breast cancer and other cancers of the reproductive system of women.

(c) Programs for breast cancer

(1) In general

In carrying out subsection (a) of this section, the Director of the Institute shall conduct or support research to expand the understanding of the cause of, and to find a cure for, breast cancer. Activities under such subsection shall provide for an expansion and intensification of the conduct and support of—

(A) basic research concerning the etiology and causes of breast cancer;

(B) clinical research and related activities concerning the causes, prevention, detection and treatment of breast cancer;

(C) control programs with respect to breast cancer in accordance with section 285a–1 of this title, including community-based programs designed to assist women who are members of medically underserved populations, low-income populations, or minority groups;

(D) information and education programs with respect to breast cancer in accordance with section 285a–2 of this title; and

(E) research and demonstration centers with respect to breast cancer in accordance with section 285a–3 of this title, including the development and operation of centers for breast cancer research to bring together basic and clinical, biomedical and behavioral scientists to conduct basic, clinical, epidemiological, psychosocial, prevention and treatment research and related activities on breast cancer.

Not less than six centers shall be operated under subparagraph (E). Activities of such centers should include supporting new and innovative research and training programs for new researchers. Such centers shall give priority to expediting the transfer of research advances to clinical applications.

(2) Implementation of plan for programs

(A) The Director of the Institute shall ensure that the research programs described in paragraph (1) are implemented in accordance with a plan for the programs. Such plan shall include comments and recommendations that

1 See References in Text note below.
the Director of the Institute considers appropriate, with due consideration provided to the professional judgment needs of the Institute as expressed in the annual budget estimate prepared in accordance with section 285a–2(9) of this title. The Director of the Institute, in consultation with the National Cancer Advisory Board, shall periodically review and revise such plan.

(B) Not later than October 1, 1993, the Director of the Institute shall submit a copy of the plan to the President’s Cancer Panel, the Secretary, and the Director of NIH.

(C) The Director of the Institute shall submit any revisions of the plan to the President’s Cancer Panel, the Secretary, and the Director of NIH.

(D) The Secretary shall provide a copy of the plan submitted under subparagraph (A), and any revisions submitted under subparagraph (C), to the Committee on Energy and Commerce of the House of Representatives and the Committee on Labor and Human Resources of the Senate.

(d) Other cancers

In carrying out subsection (a) of this section, the Director of the Institute shall conduct or support research on ovarian cancer and other cancers of the reproductive system of women. Activities under such subsection shall provide for the conduct and support of—

(1) basic research concerning the etiology and causes of ovarian cancer and other cancers of the reproductive system of women;

(2) clinical research and related activities into the causes, prevention, detection and treatment of ovarian cancer and other cancers of the reproductive system of women;

(3) control programs with respect to ovarian cancer and other cancers of the reproductive system of women in accordance with section 285a–1 of this title;

(4) information and education programs with respect to ovarian cancer and other cancers of the reproductive system of women in accordance with section 285a–2 of this title; and

(5) research and demonstration centers with respect to ovarian cancer and cancers of the reproductive system in accordance with section 285a–3 of this title.

(e) Report

The Director of the Institute shall prepare, for inclusion in the biennial report submitted under section 284h of this title, a report that describes the activities of the National Cancer Institute under the research programs referred to in subsection (a) of this section, that shall include—

(1) a description of the research plan with respect to breast cancer prepared under subsection (c) of this section;

(2) an assessment of the development, revision, and implementation of such plan;

(3) a description and evaluation of the progress made, during the period for which such report is prepared, in the research programs on breast cancer and cancers of the reproductive system of women;

(4) a summary and analysis of expenditures made, during the period for which such report is made, for activities with respect to breast cancer and cancers of the reproductive system of women conducted and supported by the National Institutes of Health; and

(5) such comments and recommendations as the Director considers appropriate.

(July 1, 1944, ch. 373, title IV, § 417, as added Pub. L. 103–43, title IV, § 401, June 10, 1993, 107 Stat. 153.)

REFERENCES IN TEXT


CHANGE OF NAME

Committee on Labor and Human Resources of Senate changed to Committee on Health, Education, Labor, and Pensions of Senate by Senate Resolution No. 20, One Hundred Sixth Congress, Jan. 19, 1999.


§ 285a–7. Prostate cancer

(a) Expansion and coordination of activities

The Director of the Institute, in consultation with the National Cancer Advisory Board, shall expand, intensify, and coordinate the activities of the Institute with respect to research on prostate cancer.

(b) Coordination with other institutes

The Director of the Institute shall coordinate the activities of the Director under subsection (a) of this section with similar activities conducted by other national research institutes and agencies of the National Institutes of Health to the extent that such Institutes¹ and agencies have responsibilities that are related to prostate cancer.

(c) Programs

(1) In general

In carrying out subsection (a) of this section, the Director of the Institute shall conduct or support research to expand the understanding of the cause of, and to find a cure for, prostate cancer. Activities under such subsection shall provide for an expansion and intensification of the conduct and support of—

(A) basic research concerning the etiology and causes of prostate cancer;

(B) clinical research and related activities concerning the causes, prevention, detection and treatment of prostate cancer;

(C) prevention and control and early detection programs with respect to prostate cancer in accordance with section 285a–1 of this

¹ So in original. Probably should not be capitalized.
title, particularly as it relates to intensifying research on the role of prostate specific antigen for the screening and early detection of prostate cancer;

(D) an Inter-Institute Task Force, under the direction of the Director of the Institute, to provide coordination between relevant National Institutes of Health components of research efforts on prostate cancer;

(E) control programs with respect to prostate cancer in accordance with section 285a-1 of this title;

(F) information and education programs with respect to prostate cancer in accordance with section 285a-2 of this title; and

(G) research and demonstration centers with respect to prostate cancer in accordance with section 285a-3 of this title, including the development and operation of centers for prostate cancer research to bring together basic and clinical, biomedical and behavioral scientists to conduct basic, clinical, epidemiological, psychosocial, prevention and control, treatment, research, and related activities on prostate cancer.

Not less than six centers shall be operated under subparagraph (G). Activities of such centers should include supporting new and innovative research and training programs for new researchers. Such centers shall give priority to expediting the transfer of research advances to clinical applications.

(2) Implementation of plan for programs

(A) The Director of the Institute shall ensure that the research programs described in paragraph (1) are implemented in accordance with a plan for the programs. Such plan shall include any revisions submitted under subparagraph (C), to the Committee on Energy and Commerce of the House of Representatives and the Committee on Commerce of the Senate.

(B) Not later than October 1, 1993, the Director of the Institute shall submit a copy of the plan to the President’s Cancer Panel, the Secretary, and the Director of NIH.

(C) The Director of the Institute shall submit any revisions of the plan to the President’s Cancer Panel, the Secretary, and the Director of NIH.

(D) The Secretary shall provide a copy of the plan submitted under subparagraph (A), and any revisions submitted under subparagraph (C), to the Committee on Energy and Commerce of the House of Representatives and the Committee on Labor and Human Resources of the Senate.

(July 1, 1944, ch. 373, title IV, §417A, as added Pub. L. 103-43, title IV, §402, June 10, 1993, 107 Stat. 155.)

CHANGE OF NAME

Committee on Labor and Human Resources of Senate changed to Committee on Health, Education, Labor, and Pensions of Senate by Senate Resolution No. 20, One Hundred Sixth Congress, Jan. 19, 1999.

Committee on Energy and Commerce of House of Representatives treated as referring to Committee on Commerce of House of Representatives by section 1(a) of Pub. L. 104-14, set out as a note preceding section 21 of Title 2. The Congress Committee on Commerce of House of Representatives changed to Committee on Energy and Commerce of House of Representatives, and jurisdiction over matters relating to securities and exchanges and insurance generally transferred to Committee on Financial Services of House of Representatives by House Resolution No. 5, One Hundred Seventh Congress, Jan. 3, 2001.


EFFECTIVE DATE OF REPEAL

Repeal applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as an Effective Date of 2007 Amendment note under section 261 of this title.

§ 285a–9. Grants for education, prevention, and early detection of radiogenic cancers and diseases

(a) Definition

In this section the term “entity” means any—

(1) National Cancer Institute-designated cancer center;

(2) Department of Veterans Affairs hospital or medical center;

(3) Federally Qualified Health Center, community health center, or hospital;

(4) agency of any State or local government, including any State department of health; or

(5) nonprofit organization.

(b) In general

The Secretary, acting through the Administrator of the Health Resources and Services Administration in consultation with the Director of the National Institutes of Health and the Director of the Indian Health Service, may make competitive grants to any entity for the purpose of carrying out programs to—

(1) screen individuals described under section 4(a)(1)(A)(i) or 5(a)(1)(A) of the Radiation Exposure Compensation Act (42 U.S.C. 2210 note) for cancer as a preventative health measure;

(2) provide appropriate referrals for medical treatment of individuals screened under paragraph (1) and to ensure, to the extent practicable, the provision of appropriate follow-up services;

(3) develop and disseminate public information and education programs for the detection, prevention, and treatment of radiogenic cancers and diseases; and

(4) facilitate putative applicants in the documentation of claims as described in section 5(a) of the Radiation Exposure Compensation Act (42 U.S.C. 2210 note).

(c) Indian Health Service

The programs under subsection (a) of this section shall include programs provided through
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the Indian Health Service or through tribal contracts, compacts, grants, or cooperative agreements with the Indian Health Service and which are determined appropriate to raising the health status of Indians.

(d) Grant and contract authority

Entities receiving a grant under subsection (b) of this section may expend the grant to carry out the purpose described in such subsection.

(e) Health coverage unaffected

Nothing in this section shall be construed to affect any coverage obligation of a governmental or private health plan or program relating to an individual referred to under subsection (b)(1) of this section.


REFERENCES IN TEXT

Sections 4 and 5 of the Radiation Exposure Compensation Act, referred to in subsec. (b)(1) and (4), are sections 4 and 5 of Pub. L. 101–426, which are set out as a note under section 2210 of this title.

AMENDMENTS

2007—Subsec. (f). Pub. L. 109–482, §104(b)(1)(F), struck out heading and text of subsec. (f). Text read as follows: “Beginning on October 1 of the year following the date on which amounts are first appropriated to carry out this section and annually on each October 1 thereafter, the Secretary shall submit a report to the Committee on the Judiciary and the Committee on Health, Education, Labor, and Pensions of the Senate and to the Committee on the Judiciary and the Committee on Commerce of the House of Representatives. Each report shall summarize the expenditures and programs funded under this section as the Secretary determines to be appropriate.”

Subsec. (g). Pub. L. 109–482, §103(b)(16), struck out heading and text of subsec. (g). Text read as follows: “There are authorized to be appropriated for the purpose of carrying out this section $20,000,000 for fiscal year 1999 and such sums as may be necessary for each of the fiscal years 2000 through 2009.”

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

§ 285a–10. Research, information, and education with respect to blood cancer

(a) Joe Moakley Research Excellence Program

(1) In general

The Director of NIH shall expand, intensify, and coordinate programs for the conduct and support of research with respect to blood cancer, and particularly with respect to leukemia, lymphoma, and multiple myeloma.

(2) Administration

The Director of NIH shall carry out this subsection through the Director of the National Cancer Institute and in collaboration with any other agencies that the Director determines to be appropriate.

(b) Geraldine Ferraro Cancer Education Program

(1) In general

The Secretary shall direct the appropriate agency within the Department of Health and Human Services, in collaboration with the Director of NIH, to establish and carry out a program to provide information and education for patients and the general public with respect to blood cancer, and particularly with respect to the treatment of leukemia, lymphoma, and multiple myeloma.

(2) Administration

The Agency determined by the Secretary under paragraph (1) shall carry out this subsection in collaboration with private health organizations that have national education and patient assistance programs on blood-related cancers.


CODIFICATION

Section 3 of Pub. L. 107–172, which directed that section 417D (this section) be inserted after section 419C of part C of title IV of the Public Health Service Act, was executed by adding section 417D to part C of title IV of the Public Health Service Act, to reflect the probable intent of Congress, notwithstanding that part C does not contain a section 419C.

AMENDMENTS

2007—Subsec. (a)(3). Pub. L. 109–482, §103(b)(17)(A), struck out heading and text of par. (3). Text read as follows: “For the purpose of carrying out this subsection, there is authorized to be appropriated such sums as may be necessary for fiscal year 2002 and each subsequent fiscal year. Such authorizations of appropriations are in addition to other authorizations of appropriations that are available for such purpose.”

Subsec. (b)(3). Pub. L. 109–482, §103(b)(17)(B), struck out heading and text of par. (3). Text read as follows: “For the purpose of carrying out this subsection, there is authorized to be appropriated such sums as may be necessary for fiscal year 2002 and each subsequent fiscal year. Such authorizations of appropriations are in addition to other authorizations of appropriations that are available for such purpose.”

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

CONGRESSIONAL FINDINGS


“(1) An estimated 109,500 people in the United States will be diagnosed with leukemia, lymphoma, and multiple myeloma in 2001.

“(2) New cases of the blood cancers described in paragraph (1) account for 8.6 percent of new cancer cases.

“(3) Those devastating blood cancers will cause the deaths of an estimated 60,300 persons in the United States in 2001. Every 9 minutes, a person in the United States dies from leukemia, lymphoma, or multiple myeloma.

“(4) While less than 5 percent of Federal funds for cancer research are spent on those blood cancers, those blood cancers cause 11 percent of all cancer deaths in the United States.

“(5) Increased Federal support of research into leukemia, lymphoma, and multiple myeloma has resulted and will continue to result in significant advances in the treatment, and ultimately the cure, of those blood cancers as well as other cancers.”
§ 285a–11. Pediatric cancer research and awareness

(a) Pediatric cancer research

(1) Programs of research excellence in pediatric cancer

The Secretary, in collaboration with the Director of NIH and other Federal agencies with interest in prevention and treatment of pediatric cancer, shall continue to enhance, expand, and intensify pediatric cancer research and other activities related to pediatric cancer, including therapeutically applicable research to generate effective treatments, pediatric preclinical testing, and pediatric clinical trials through National Cancer Institute-supported pediatric cancer clinical trial groups and their member institutions. In enhancing, expanding, and intensifying such research and other activities, the Secretary is encouraged to take into consideration the application of such research and other activities for minority, health disparity, and medically underserved communities. For purposes of this section, the term “pediatric cancer research” means research on the causes, prevention, diagnosis, recognition, treatment, and long-term effects of pediatric cancer.

(2) Peer review requirements

All grants awarded under this subsection shall be awarded in accordance with section 289a of this title.

(b) Public awareness of pediatric cancers and available treatments and research

(1) In general

The Secretary may award grants to childhood cancer professional and direct service organizations for the expansion and widespread implementation of—

(A) activities that provide available information on treatment protocols to ensure early access to the best available therapies and clinical trials for pediatric cancers;

(B) activities that provide available information on the late effects of pediatric cancer treatment to ensure access to necessary long-term medical and psychological care; and

(C) direct resource services such as educational outreach for parents, peer-to-peer and parent-to-parent support networks, information on school re-entry and post-secondary education, and resource directories or referral services for financial assistance, psychological counseling, and other support services.

In awarding grants under this paragraph, the Secretary is encouraged to take into consideration the extent to which an entity would use such grant for purposes of making activities and services described in this paragraph available to minority, health disparity, and medically underserved communities.

(2) Performance measurement, transparency, and accountability

For each grant awarded under this subsection, the Secretary shall develop and implement metrics-based performance measures to assess the effectiveness of activities funded under such grant.

(3) Informational requirements

Any information made available pursuant to a grant awarded under paragraph (1) shall be—

(A) culturally and linguistically appropriate as needed by patients and families affected by childhood cancer; and

(B) approved by the Secretary.

(c) Rule of construction

Nothing in this section shall be construed as being inconsistent with the goals and purposes of the Minority Health and Health Disparities Research and Education Act of 2000 (42 U.S.C. 202 note).

(d) Authorization of appropriations

For purposes of carrying out this section and section 280e–3a of this title, there are authorized to be appropriated $30,000,000 for each of fiscal years 2009 through 2013. Such authorization of appropriations is in addition to the authorization of appropriations established in section 282a of this title with respect to such purpose. Funds appropriated under this subsection shall remain available until expended.

(1) Establishment

Not later than 6 months after October 8, 2008, the Secretary shall establish a committee, to be known as the Interagency Breast Cancer and Environmental Research Coordinating Committee (in this section referred to as the “Committee”).

(2) Duties

The Committee shall—

(A) share and coordinate information on existing research activities, and make recommendations to the National Institutes of Health and other Federal agencies regarding how to improve existing research programs, that are related to breast cancer research;

(B) develop a comprehensive strategy and advise the National Institutes of Health and other Federal agencies in the solicitation of proposals for collaborative, multidisciplinary research, including proposals to evaluate environmental and genomic factors that may be related to the etiology of breast cancer that would—

(i) result in innovative approaches to study emerging scientific opportunities or

References in Text


§ 285a–12. Interagency Breast Cancer and Environmental Research Coordinating Committee

(a) Interagency Breast Cancer and Environmental Research Coordinating Committee

(1) Establishment

Not later than 6 months after October 8, 2008, the Secretary shall establish a committee, to be known as the Interagency Breast Cancer and Environmental Research Coordinating Committee (in this section referred to as the “Committee”).

(2) Duties

The Committee shall—

(A) share and coordinate information on existing research activities, and make recommendations to the National Institutes of Health and other Federal agencies regarding how to improve existing research programs, that are related to breast cancer research;

(B) develop a comprehensive strategy and advise the National Institutes of Health and other Federal agencies in the solicitation of proposals for collaborative, multidisciplinary research, including proposals to evaluate environmental and genomic factors that may be related to the etiology of breast cancer that would—

(i) result in innovative approaches to study emerging scientific opportunities or

1 So in original. See References in Text note below.
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eliminate knowledge gaps in research to improve the research portfolio;
(ii) outline key research questions, methodologies, and knowledge gaps;
(iii) expand the number of research proposals that involve collaboration between 2 or more national research institutes or national centers, including proposals for Common Fund research described in section 282(b)(7) of this title to improve the research portfolio; and
(iv) expand the number of collaborative, multidisciplinary, and multi-institutional research grants;
(C) develop a summary of advances in breast cancer research supported or conducted by Federal agencies relevant to the diagnosis, prevention, and treatment of cancer and other diseases and disorders; and
(D) not later than 2 years after the date of the establishment of the Committee, make recommendations to the Secretary—
(i) regarding any appropriate changes to research activities, including recommendations to improve the research portfolio of the National Institutes of Health to ensure that scientifically-based strategic planning is implemented in support of research priorities that impact breast cancer research activities;
(ii) to ensure that the activities of the National Institutes of Health and other Federal agencies, including the Department of Defense, are free of unnecessary duplication of effort;
(iii) regarding public participation in decisions relating to breast cancer research to increase the involvement of patient advocacy and community organizations representing a broad geographical area;
(iv) on how best to disseminate information on breast cancer research progress; and
(v) on how to expand partnerships between public entities, including Federal agencies, and private entities to expand collaborative, cross-cutting research.

(3) Rule of construction

For the purposes of the Committee, when focusing on research to evaluate environmental and genomic factors that may be related to the etiology of breast cancer, nothing in this section shall be construed to restrict the Secretary from including other forms of cancer, as appropriate, when doing so may advance research in breast cancer or advance research in other forms of cancer.

(4) Membership

(A) In general

The Committee shall be composed of the following voting members:
(i) Not more than 7 voting Federal representatives as follows:
(I) The Director of the Centers for Disease Control and Prevention.
(II) The Director of the National Institutes of Health and the directors of such national research institutes and national centers (which may include the National Institute of Environmental Health Sciences) as the Secretary determines appropriate.
(III) One representative from the National Cancer Institute Board of Scientific Advisors, appointed by the Director of the National Cancer Institute.
(IV) The heads of such other agencies of the Department of Health and Human Services as the Secretary determines appropriate.
(V) Representatives of other Federal agencies that conduct or support cancer research, including the Department of Defense.
(ii) 2 or more national research institutes or national centers (which may include the National Institutes of Health and the directors of such national centers, including proposals for Common Fund research described in section 282(b)(7) of this title to improve the research portfolio; and
(B) Additional members

The Committee shall include additional voting members appointed by the Secretary as follows:
(i) 6 members shall be appointed from among scientists, physicians, and other health professionals, who—
(I) are not officers or employees of the United States;
(II) represent multiple disciplines, including clinical, basic, and public health sciences;
(III) represent different geographical regions of the United States;
(IV) are from practice settings, academia, or other research settings; and
(V) are experienced in scientific peer review process.
(ii) 6 members shall be appointed from members of the general public, who represent individuals with breast cancer.
(C) Nonvoting members

The Committee shall include such nonvoting members as the Secretary determines to be appropriate.

(5) Chairperson

The voting members of the Committee shall select a chairperson from among such members. The selection of a chairperson shall be subject to the approval of the Director of NIH.

(6) Meetings

The Committee shall meet at the call of the chairperson of the Committee or upon the request of the Director of NIH, but in no case less often than once each year.

(b) Review

The Secretary shall review the necessity of the Committee in calendar year 2011 and, thereafter, at least once every 2 years.


SUBPART 2—NATIONAL HEART, LUNG, AND BLOOD INSTITUTE

§ 285b. Purpose of Institute

The general purpose of the National Heart, Lung, and Blood Institute (hereafter in this subpart referred to as the “Institute”) is the con-
duct and support of research, training, health information dissemination, and other programs with respect to heart, blood vessel, lung, and blood diseases and with respect to the use of blood and blood products and the management of blood resources.

(July 1, 1944, ch. 373, title IV, §418, as added Pub. L. 99–158, §2, Nov. 20, 1985, 99 Stat. 836.)

§ 285b–1. Heart, blood vessel, lung, and blood disease prevention and control programs

(a) The Director of the Institute shall conduct and support programs for the prevention and control of heart, blood vessel, lung, and blood diseases. Such programs shall include community-based and population-based programs carried out in cooperation with other Federal agencies, with public health agencies of State or local governments, with nonprofit private entities that are community-based health agencies, or with other appropriate public or nonprofit private entities.

(b) In carrying out programs under subsection (a) of this section, the Director of the Institute shall give special consideration to the prevention and control of heart, blood vessel, lung, and blood diseases in children, and in populations that are at increased risk with respect to such diseases.


AMENDMENTS

1993—Pub. L. 103–43 substituted subsecs. (a) and (b) for former subsec. (a) which read as follows: ‘‘The Director of the Institute, under policies established by the Director of NIH and after consultation with the advisory council for the Institute, shall establish programs as necessary for cooperation with other Federal health agencies, State, local, and regional public health agencies, and nonprofit private health agencies in the diagnosis, prevention, and treatment (including the provision of emergency medical services) of heart, blood vessel, lung, and blood diseases, appropriately emphasizing the prevention, diagnosis, and treatment of such diseases of children.’’

§ 285b–2. Information and education

The Director of the Institute shall collect, identify, analyze, and disseminate on a timely basis, through publications and other appropriate means, to patients, families of patients, physicians and other health professionals, and the general public, information on research, prevention, diagnosis, and treatment of heart, blood vessel, lung, and blood diseases, the maintenance of health to reduce the incidence of such diseases, and on the use of blood and blood products and the management of blood resources. In carrying out this section, the Director of the Institute shall place special emphasis upon the utilization of collaborative efforts with both the public and private sectors to—

(1) increase the awareness and knowledge of health care professionals and the public regarding the prevention of heart and blood vessel, lung, and blood diseases and the utilization of blood resources; and

(2) develop and disseminate to health professionals, patients and patient families, and the public information designed to encourage adults and children to adopt healthful practices concerning the prevention of such diseases.


AMENDMENTS

1988—Pub. L. 100–607 amended second sentence generally. Prior to amendment, second sentence read as follows: ‘‘In carrying out this section the Director of the Institute shall place special emphasis upon—

(1) the dissemination of information regarding diet and nutrition, environmental pollutants, exercise, stress, hypertension, cigarette smoking, weight control, and other factors affecting the prevention of arteriosclerosis and other cardiovascular diseases and of pulmonary and blood diseases; and

(2) the dissemination of information designed to encourage children to adopt healthful habits respecting the risk factors related to the prevention of such diseases.’’

§ 285b–3. National Heart, Blood Vessel, Lung, and Blood Diseases and Blood Resources Program; administrative provisions

(a)(1) The National Heart, Blood Vessel, Lung, and Blood Diseases and Blood Resources Program (hereafter in this subpart referred to as the ‘‘Program’’) may provide for—

(A) investigation into the epidemiology, etiology, and prevention of all forms and aspects of heart, blood vessel, lung, and blood diseases, including investigations into the social, environmental, behavioral, nutritional, biological, and genetic determinants and influences involved in the epidemiology, etiology, and prevention of such diseases;

(B) studies and research into the basic biological processes and mechanisms involved in the underlying normal and abnormal heart, blood vessel, lung, and blood phenomena;

(C) research into the development, trial, and evaluation of techniques, drugs, and devices (including computers) used in, and approaches to, the diagnosis, treatment (including the provision of emergency medical services), and prevention of heart, blood vessel, lung, and blood diseases and the rehabilitation of patients suffering from such diseases;

(D) establishment of programs that will focus and apply scientific and technological efforts involving the biological, physical, and engineering sciences to all facets of heart, blood vessel, lung, and blood diseases with emphasis on the refinement, development, and evaluation of technological devices that will assist, replace, or monitor vital organs and improve instrumentation for detection, diagnosis, and treatment of and rehabilitation from such diseases;

(E) establishment of programs for the conduct and direction of field studies, large-scale testing and evaluation, and demonstration of preventive, diagnostic, therapeutic, and rehabilitative approaches to, and emergency medical services for, such diseases;

(F) studies and research into blood diseases and blood, and into the use of blood for clinical purposes and all aspects of the manage-
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eral Services, buildings or parts of buildings in the District of Columbia or communities located adjacent to the District of Columbia for the use of the Institute for a period not to exceed ten years;

(3) subject to section 284(b)(2) of this title and without regard to section 3324 of title 31 and section 6101 of title 41, may enter into such contracts, leases, cooperative agreements, or other transactions, as may be necessary in the conduct of the Director's functions, with any public agency, or with any person, firm, association, corporation, or educational institutions;

(4) may make grants to public and nonprofit private entities to assist in meeting the cost of the care of patients in hospitals, clinics, and related facilities who are participating in research projects; and

(5) shall, in consultation with the advisory council for the Institute, conduct appropriate intramural training and education programs, including continuing education and laboratory and clinical research training programs.

Except as otherwise provided, experts and consultants whose services are obtained under paragraph (1) shall be paid or reimbursed, in accordance with title 5, for their travel to and from their place of service and for other expenses associated with their assignment. Such expenses shall not be allowed in connection with the assignment of an expert or consultant whose services are obtained under paragraph (1) unless the expert or consultant has agreed in writing to complete the entire period of the assignment or one year of the assignment, whichever is shorter, unless separated or reassigned for reasons which are beyond the control of the expert or consultant and which are acceptable to the Director of the Institute. If the expert or consultant violates the agreement, the money spent by the United States for such expenses is recoverable from the expert or consultant as a debt due the United States. The Secretary may waive in whole or in part a right of recovery under the preceding sentence.


CONSIDERATION


AMENDMENTS


Subsec. (b)(1). Pub. L. 100–607, § 127(2), substituted “after consultation with” for “, after approval of”.

ment of blood resources in the United States, including the collection, preservation, fractionation, and distribution of blood and blood products;

(G) the education (including continuing education) and training of scientists, clinical investigators, and educators, in fields and specialties (including computer sciences) requisite to the conduct of clinical programs respecting heart, blood vessel, lung, and blood diseases and blood resources;

(H) public and professional education relating to all aspects of such diseases, including the prevention of such diseases, and the use of blood and blood products and the management of blood resources;

(I) establishment of programs for study and research into heart, blood vessel, lung, and blood diseases of children (including cystic fibrosis, hyaline membrane, hemolytic diseases such as sickle cell anemia and Cooley's anemia, and hemophilic diseases) and for the development and demonstration of diagnostic, treatment, and preventive approaches to such diseases; and

(J) establishment of programs for study, research, development, demonstrations and evaluation of emergency medical services for people who become critically ill in connection with heart, blood vessel, lung, or blood diseases.

(2) The Program shall be coordinated with other national research institutes to the extent that they have responsibilities respecting such diseases and shall give special emphasis to the continued development in the Institute of programs related to the causes of stroke and to effective coordination of such programs with related stroke programs in the National Institute of Neurological and Communicative Disorders and Stroke. The Director of the Institute, with the advice of the advisory council for the Institute, shall revise annually the plan for the Program and shall carry out the Program in accordance with such plan.

(b) In carrying out the Program, the Director of the Institute, under policies established by the Director of NIH—

(1) may, after consultation with the advisory council for the Institute, obtain (in accordance with section 3109 of title 5, but without regard to the limitation in such section on the period of such service) the services of not more than one hundred experts or consultants who have scientific or professional qualifications;

(2)(A) may, in consultation with the advisory council for the Institute, acquire and construct, improve, repair, operate, alter, renovate, and maintain, heart, blood vessel, lung, and blood disease and blood resource laboratories, research, training, and other facilities, equipment, and such other real or personal property as the Director determines necessary;

(B) may, in consultation with the advisory council for the Institute, make grants for construction or renovation of facilities; and

(C) may, in consultation with the advisory council for the Institute, acquire, without regard to section 8141 of title 40, by lease or otherwise, through the Administrator of General Services, buildings or parts of buildings in the District of Columbia or communities located adjacent to the District of Columbia for the use of the Institute for a period not to exceed ten years;

(a) Heart, blood vessel, lung, blood diseases, and blood resources; utilization of centers for prevention programs

(1) The Director of the Institute may provide, in accordance with subsection (c) of this section, for the development of—
   (A) ten centers for basic and clinical research into, training in, and demonstration of, advanced diagnostic, prevention, and treatment and rehabilitation methods (including methods of providing emergency medical services) for heart and blood vessel diseases;
   (B) ten centers for basic and clinical research into, training in, and demonstration of, advanced diagnostic, prevention, and treatment and rehabilitation methods (including methods of providing emergency medical services) for lung diseases (including bronchitis, emphysema, asthma, cystic fibrosis, and other lung diseases of children);
   (C) ten centers for basic and clinical research into, training in, and demonstration of, advanced diagnostic, prevention, and treatment methods (including genetic studies, intrauterine environment studies, postnatal studies, heart arrhythmias, and acquired heart disease and preventive cardiology) for cardiovascular diseases in children.

(2) The centers developed under paragraph (1) shall, in addition to being utilized for research, training, and demonstrations, be utilized for the following prevention programs for cardiovascular, pulmonary, and blood diseases:
   (A) Programs to develop improved methods of detecting individuals with a high risk of developing cardiovascular, pulmonary, and blood diseases.
   (B) Programs to develop improved methods of intervention against those factors which cause individuals to have a high risk of developing such diseases.
   (C) Programs to develop health professions and allied health professions personnel highly skilled in the prevention of such diseases.
   (D) Programs to develop improved methods of providing emergency medical services for persons with such diseases.
   (E) Programs of continuing education for health and allied health professionals in the diagnosis, prevention, and treatment of such diseases and the maintenance of health to reduce the incidence of such diseases and information programs for the public respecting the prevention and early diagnosis and treatment of such diseases and the maintenance of health.

(3) The research, training, and demonstration activities carried out through any such center may relate to any one or more of the diseases referred to in paragraph (1) of this subsection.

(b) Sickle cell anemia

The Director of the Institute shall provide, in accordance with subsection (c) of this section, for the development of ten centers for basic and clinical research into the diagnosis, treatment, and control of sickle cell anemia.

(c) Cooperative agreements and grants for establishing and supporting; uses for Federal payments; period of support, additional periods

(1) The Director of the Institute may enter into cooperative agreements with and make grants to public or private nonprofit entities to pay all or part of the cost of planning, establishing, or strengthening, and providing basic operating support for centers for basic and clinical research into, training in, and demonstration of the management of blood resources and advanced diagnostic, prevention, and treatment methods for heart, blood vessel, lung, or blood diseases.

(2) A cooperative agreement or grant under paragraph (1) shall be entered into in accordance with policies established by the Director of NIH and after consultation with the Institute’s advisory council.

(3) Federal payments made under a cooperative agreement or grant under paragraph (1) may be used for—
   (A) construction (notwithstanding any limitation under section 289e of this title);
   (B) staffing and other basic operating costs, including such patient care costs as are required for research;
   (C) training, including training for allied health professionals; and
   (D) demonstration purposes.

As used in this subsection, the term "construction" does not include the acquisition of land, and the term "training" does not include research training for which Ruth L. Kirschstein National Research Service Awards may be provided under section 289e of this title.

(4) Support of a center under paragraph (1) may be for a period of not to exceed five years. Such period may be extended by the Director for additional periods of not more than five years each if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director and if such group has recommended to the Director that such period should be extended.


Amendments


Section, act July 1, 1944, ch. 373, title IV, § 423, as added Nov. 20, 1985, Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 841, directed Secretary to establish an Interagency Technical Committee on Heart, Blood Vessel, Lung, and Blood Diseases and Blood Resources.

§ 285b-6. Associate Director for Prevention; appointment; function

(a) There shall be in the Institute an Associate Director for Prevention to coordinate and promote the programs in the Institute concerning the prevention of heart, blood vessel, lung, and blood diseases. The Associate Director shall be appointed by the Director of the Institute from individuals who because of their professional training or experience are experts in public health or preventive medicine.

(b) The Associate Director for Prevention shall prepare for inclusion in the biennial report made under section 284b of this title a description of the prevention activities of the Institute, including a description of the staff and resources allocated to those activities.


REFERENCES IN TEXT
Section 284b of this title, referred to in subsec. (b), was repealed by Pub. L. 100–607, title I, § 104(b)(1)(C), Nov. 4, 1988, 102 Stat. 3055.

PRIOR PROVISIONS
A prior section 423 of act July 1, 1944, was classified to section 285b-5 of this title prior to repeal by Pub. L. 100–607.

§ 285b-7. National Center on Sleep Disorders Research

(a) Establishment

Not later than 1 year after June 10, 1993, the Director of the Institute shall establish the National Center on Sleep Disorders Research (in this section referred to as the “Center”). The Center shall be headed by a director, who shall be appointed by the Director of the Institute.

(b) Purpose

The general purpose of the Center is—

(1) the conduct and support of research, training, health information dissemination, and other activities with respect to sleep disorders, including biological and circadian rhythm research, basic understanding of sleep, chronobiological and other sleep related research; and

(2) to coordinate the activities of the Center with similar activities of other Federal agencies, including the other agencies of the National Institutes of Health, and similar activities of other public entities and nonprofit entities.

(c) Sleep Disorders Research Advisory Board

(1) The Director of the National Institutes of Health shall establish a board to be known as the Sleep Disorders Research Advisory Board (in this section referred to as the “Advisory Board”).

(2) The Advisory Board shall advise, assist, consult with, and make recommendations to the Director of the National Institutes of Health, through the Director of the Institute, and the Director of the Center concerning matters relating to the scientific activities carried out by and through the Center and the policies respecting such activities, including recommendations with respect to the plan required in subsection (c) of this section.

(3) (A) The Director of the National Institutes of Health shall appoint to the Advisory Board 12 appropriately qualified representatives of the public who are not officers or employees of the Federal Government. Of such members, eight shall be representatives of health and scientific disciplines with respect to sleep disorders and four shall be individuals representing the interests of individuals with or undergoing treatment for sleep disorders.

(B) The following officials shall serve as ex officio members of the Advisory Board:

(i) The Director of the National Institutes of Health.

(ii) The Director of the National Heart, Lung and Blood Institute.

(iii) The Director of the National Institute of Mental Health.

(v) The Director of the National Institute on Aging.

(vi) The Director of the Eunice Kennedy Shriver National Institute of Child Health and Human Development.

(vii) The Director of the National Institute of Neurological Disorders and Stroke.

(viii) The Assistant Secretary for Health.

(ix) The Assistant Secretary of Defense (Health Affairs).

(x) The Chief Medical Director of the Veterans Administration.

(4) The members of the Advisory Board shall, from among the members of the Advisory Board, designate an individual to serve as the chair of the Advisory Board.

(5) Except as inconsistent with, or inapplicable to, this section, the provisions of section 284a of this title shall apply to the advisory board established under this section in the same manner as such provisions apply to any advisory council established under such section.

(d) Development of comprehensive research plan; revision

(1) After consultation with the Director of the Center and the advisory board established under subsection (c) of this section, the Director of the National Institutes of Health shall develop a comprehensive plan for the conduct and support of sleep disorders research.

(2) The plan developed under paragraph (1) shall identify priorities with respect to such research and shall provide for the coordination of such research conducted or supported by the agencies of the National Institutes of Health.

1 See References in Text note below.

2 So in original. Probably should be capitalized.
(3) The Director of the National Institutes of Health (after consultation with the Director of the Center and the advisory board established under subsection (c) of this section) shall revise the plan developed under paragraph (1) as appropriate.

(e) Collection and dissemination of information

The Director of the Center, in cooperation with the Centers for Disease Control and Prevention, is authorized to coordinate activities with the Department of Transportation, the Department of Defense, the Department of Education, the Department of Labor, and the Department of Commerce to collect data, conduct studies, and disseminate public information concerning the impact of sleep disorders and sleep deprivation.


AMENDMENTS

CHANGE OF NAME
Reference to Chief Medical Director of Department of Veterans Affairs deemed to refer to Under Secretary for Health of Department of Veterans Affairs pursuant to section 302(e) of Pub. L. 102–405, set out as a note under section 217a of this title, provided that an advisory committee established pursuant to the Public Health Service Act shall terminate at such time as may be specifically prescribed by an Act of Congress enacted after Jan. 4, 1975.

§ 285b–7a. Heart attack, stroke, and other cardiovascular diseases in women

(a) In general

The Director of the Institute shall expand, intensify, and coordinate research and related activities of the Institute with respect to heart attack, stroke, and other cardiovascular diseases in women.

(b) Coordination with other institutes

The Director of the Institute shall coordinate activities under subsection (a) of this section with similar activities conducted by the other national research institutes and agencies of the National Institutes of Health to the extent that such Institutes and agencies have responsibilities that are related to heart attack, stroke, and other cardiovascular diseases in women.

(c) Certain programs

In carrying out subsection (a) of this section, the Director of the Institute shall conduct or support research to expand the understanding of the causes of, and to develop methods for preventing, cardiovascular diseases in women. Activities under such subsection shall include conducting and supporting the following:

(1) Research to determine the reasons underlying the prevalence of heart attack, stroke, and other cardiovascular diseases in women, including African-American women and other women who are members of racial or ethnic minority groups.

(2) Basic research concerning the etiology and causes of cardiovascular diseases in women.

(3) Epidemiological studies to address the frequency and natural history of such diseases and the differences among men and women, and among racial and ethnic groups, with respect to such diseases.

(4) The development of safe, efficient, and cost-effective diagnostic approaches to evaluating women with suspected ischemic heart disease.

(5) Clinical research for the development and evaluation of new treatments for women, including rehabilitation.

(6) Studies to gain a better understanding of methods of preventing cardiovascular diseases in women, including applications of effective methods for the control of blood pressure, lipids, and obesity.

(7) Information and education programs for patients and health care providers on risk factors associated with heart attack, stroke, and other cardiovascular diseases in women, and on the importance of the prevention or control of such risk factors and timely referral with appropriate diagnosis and treatment. Such programs shall include information and education on health-related behaviors that can improve such important risk factors as smoking, obesity, high blood cholesterol, and lack of exercise.


AMENDMENTS
2007—Subsec. (d). Pub. L. 109–482 struck out heading and text of subsec. (d). Text read as follows: “For the purpose of carrying out this section, there are authorized to be appropriated such sums as may be necessary for each of the fiscal years 1999 through 2003. The authorization of appropriations established in the preceding sentence is in addition to any other authorization of appropriation that is available for such purpose.”

EFFECTIVE DATE OF 2007 AMENDMENT
Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.
§ 285b–7b. Coordination of Federal asthma activities

(a) In general

The Director of the Institute shall, through the National Asthma Education Prevention Program Coordinating Committee—

(1) identify all Federal programs that carry out asthma-related activities; and

(2) develop, in consultation with appropriate Federal agencies and professional and voluntary health organizations, a Federal plan for responding to asthma.

(b) Representation of the Department of Housing and Urban Development

A representative of the Department of Housing and Urban Development shall be included on the National Asthma Education Prevention Program Coordinating Committee for the purpose of performing the tasks described in subsection (a) of this section.


AMENDMENTS

2007—Subsec. (a). Pub. L. 109–482, § 104(b)(1)(G), inserted “and” at end of par. (1), substituted a period for “; and” at end of par. (2), and struck out par. (3) which read as follows: “not later than 12 months after October 17, 2000, submit recommendations to the appropriate committees of the Congress on ways to strengthen and improve the coordination of asthma-related activities of the Federal Government.”

Subsec. (c). Pub. L. 109–482, § 103(b)(19), struck out heading and text of subsec. (c). Text read as follows: “For the purpose of carrying out this section, there are authorized to be appropriated such sums as may be necessary for each of the fiscal years 2003 through 2005.”

§ 285b–7c. Tuberculosis

(a) In general

The Director of the National Institutes of Health may expand, intensify, and coordinate research and related activities of the Institute with respect to tuberculosis, including activities toward the goal of eliminating such disease.

(b) Certain activities

Activities under subsection (a) may include—

(1) enhancing basic and clinical research on tuberculosis, including drug resistant tuberculosis;

(2) expanding research on the relationship between such disease and the human immunodeficiency virus; and

(3) developing new tools for the elimination of tuberculosis, including public health interventions and methods to enhance detection and response to outbreaks of tuberculosis, including multidrug resistant tuberculosis.


§ 285b–8. Congenital heart disease

(a) In general

The Director of the Institute may expand, intensify, and coordinate research and related activities of the Institute with respect to congenital heart disease, which may include congenital heart disease research with respect to—

(1) causation of congenital heart disease, including genetic causes;

(2) long-term outcomes in individuals with congenital heart disease, including infants, children, teenagers, adults, and elderly individuals;

(3) diagnosis, treatment, and prevention;

(4) studies using longitudinal data and retrospective analysis to identify effective treatments and outcomes for individuals with congenital heart disease; and

(5) identifying barriers to life-long care for individuals with congenital heart disease.

(b) Coordination of research activities

The Director of the Institute may coordinate research efforts related to congenital heart disease among multiple research institutions and may develop research networks.

(c) Minority and medically underserved communities

In carrying out the activities described in this section, the Director of the Institute shall consider the application of such research and other activities to minority and medically underserved communities.


PRIOR PROVISIONS


SUBPART 3—NATIONAL INSTITUTE OF DIABETES AND DIGESTIVE AND KIDNEY DISEASES

§ 285c. Purpose of Institute

The general purpose of the National Institute of Diabetes and Digestive and Kidney Diseases (hereafter in this subpart referred to as the “Institute”) is the conduct and support of research, training, health information dissemination, and other programs with respect to diabetes mellitus and endocrine and metabolic diseases, digestive diseases and nutritional disorders, and kidney, urologic, and hematologic diseases.

(July 1, 1944, ch. 373, title IV, § 426, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 841.)

STUDY ON METABOLIC DISORDERS


“(a) In General.—The Secretary of Health and Human Services (in this section referred to as the ‘Sec-
Secretary) shall, in consultation with relevant experts or through the Institute of Medicine, study issues related to treatment of PKU and other metabolic disorders for children, adolescents, and adults, and mechanisms to assure access to effective treatment, including special diets, for children and others with PKU and other metabolic disorders. Such mechanisms shall be evidence-based and reflect the best scientific knowledge regarding effective treatment and prevention of disease progression.

"(b) Dissemination of Results.—Upon completion of the study referred to in subsection (a), the Secretary shall disseminate and otherwise make available the results of the study to interested groups and organizations, including insurance commissioners, employers, private insurers, health care professionals, State and local public health agencies, and State agencies that carry out the Medicaid program under title XIX of the Social Security Act (section 1396 et seq. of this title) or the State children’s health insurance program under title XXI of such Act (section 1397aa et seq. of this title).

"(c) Authorization of Appropriations.—There are authorized to be appropriated to carry out this section such sums as may be necessary for each of the fiscal years 2001 through 2003."

\$ 285c–1. Data systems and information clearinghouses

(a) National Diabetes Data System and National Diabetes Clearinghouse

The Director of the Institute shall (1) establish the National Diabetes Data System for the collection, storage, analysis, retrieval, and dissemination of data derived from patient populations with diabetes, including, where possible, data involving general populations for the purpose of detection of individuals with a risk of developing diabetes, and (2) establish the National Diabetes Information Clearinghouse to facilitate and enhance knowledge and understanding of diabetes on the part of health professionals, patients, and the public through the effective dissemination of information.

(b) National Digestive Diseases Data System and National Digestive Diseases Information Clearinghouse

The Director of the Institute shall (1) establish the National Digestive Diseases Data System for the collection, storage, analysis, retrieval, and dissemination of data derived from patient populations with digestive diseases, including, where possible, data involving general populations for the purpose of detection of individuals with a risk of developing digestive diseases, and (2) establish the National Digestive Diseases Information Clearinghouse to facilitate and enhance knowledge and understanding of digestive diseases on the part of health professionals, patients, and the public through the effective dissemination of information.

(c) National Kidney and Urologic Diseases Data System and National Kidney and Urologic Diseases Information Clearinghouse

The Director of the Institute shall (1) establish the National Kidney and Urologic Diseases Data System for the collection, storage, analysis, retrieval, and dissemination of data derived from patient populations with kidney and urologic diseases, including, where possible, data involving general populations for the purpose of detection of individuals with a risk of developing kidney and urologic diseases, and (2) establish the National Kidney and Urologic Diseases Information Clearinghouse to facilitate and enhance knowledge and understanding of kidney and urologic diseases on the part of health professionals, patients, and the public through the effective dissemination of information.

(July 1, 1944, ch. 733, title IV, \$427, as added Pub. L. 99–158, \$2, Nov. 20, 1985, 99 Stat. 841.)

\$ 285c–2. Division Directors for Diabetes, Endocrinology, and Metabolic Diseases, Digestive Diseases and Nutrition, and Kidney, Urologic, and Hematologic Diseases; functions

(a)(1) In the Institute there shall be a Division Director for Diabetes, Endocrinology, and Metabolic Diseases, a Division Director for Digestive Diseases and Nutrition, and a Division Director for Kidney, Urologic, and Hematologic Diseases. Such Division Directors, under the supervision of the Director of the Institute, shall be responsible for—

(A) developing a coordinated plan (including recommendations for expenditures) for each of the national research institutes within the National Institutes of Health with respect to research and training concerning diabetes, endocrine and metabolic diseases, digestive diseases and nutrition, and kidney, urologic, and hematologic diseases;

(B) assessing the adequacy of management approaches for the activities within such institutes concerning such diseases and nutrition and developing improved approaches if needed;

(C) monitoring and reviewing expenditures by such institutes concerning such diseases and nutrition; and

(D) identifying research opportunities concerning such diseases and nutrition and recommending ways to utilize such opportunities.

(2) The Director of the Institute shall transmit to the Director of NIH the plans, recommendations, and reviews of the Division Directors under subparagraphs (A) through (D) of paragraph (1) together with such comments and recommendations as the Director of the Institute determines appropriate.

(b) The Director of the Institute, acting through the Division Director for Diabetes, Endocrinology, and Metabolic Diseases, the Division Director for Digestive Diseases and Nutrition, and the Division Director for Kidney, Urologic, and Hematologic Diseases, shall—

(1) carry out programs of support for research and training (other than training for which Ruth L. Kirschstein National Research Service Awards may be made under section 288 of this title) in the diagnosis, prevention, and
treatment of diabetes mellitus and endocrine and metabolic diseases, digestive diseases and nutritional disorders, and kidney, urologic, and hematologic diseases, including support for training in medical schools, graduate clinical training, graduate training in epidemiology, epidemiology studies, clinical trials, and interdisciplinary research programs; and

(2) establish programs of evaluation, planning, and dissemination of knowledge related to such research and training.


AMENDMENTS


1993—Subsec. (b). Pub. L. 103–43 substituted “the’’ for “‘the the’ before ‘Division Director for Diabetes’ in introductory provisions.

§ 285c–3. Interagency coordinating committees

(a) Establishment and purpose

For the purpose of—

(1) better coordination of the research activities of all the national research institutes relating to diabetes mellitus, digestive diseases, and kidney, urologic, and hematologic diseases; and

(2) coordinating those aspects of all Federal health programs and activities relating to such diseases to assure the adequacy and technical soundness of such programs and activities and to provide for the full communication and exchange of information necessary to maintain adequate coordination of such programs and activities;

the Secretary shall establish a Diabetes Mellitus Interagency Coordinating Committee, a Digestive Diseases Interagency Coordinating Committee, and a Kidney, Urologic, and Hematologic Diseases Coordinating Committee (hereafter in this section individually referred to as a “Committee’’).

(b) Membership; chairman; meetings

Each Committee shall be composed of the Directors of each of the national research institutes and divisions involved in research with respect to the diseases for which the Committee is established, the Division Director of the Institute for the diseases for which the Committee is established, the Under Secretary for Health of the Department of Veterans Affairs, and the Assistant Secretary of Defense for Health Affairs (or the designees of such officers) and shall in (or the designees of such officers) and shall in

AMENDMENTS

2007—Subsecs. (c), (d). Pub. L. 109–82 struck out subsecs. (c) and (d) which required an annual report detailing the work of the Committee in carrying out subsec. (a) and an annual assessment on Federal pancreatic islet cell transplantation, respectively.


1992—Subsec. (b). Pub. L. 102–405 substituted “Under Secretary for Health of the Department of Veterans Affairs’’ for “Chief Medical Director of the Department of Veterans Affairs’’.

1990—Subsec. (b). Pub. L. 102–405 substituted “Chief Medical Director of the Department of Veterans Affairs’’ for “Chief Medical Director of the Veterans’ Administration’’.

SCHEDULE FOR THE DISEASES

The Secretary shall establish the following committees:

Interagency Coordinating Committee, a Diabetes Mellitus Interagency Coordinating Committee, a Digestive Diseases Interagency Coordinating Committee, and a Kidney, Urologic, and Hematologic Diseases Coordinating Committee (hereafter in this section individually referred to as a “Committee’’).

(b) Membership; ex officio members

Each Advisory Board shall be composed of eighteen appointed members and nonvoting ex officio members as follows:

(1) The Secretary shall appoint—

(A) twelve members from individuals who are scientists, physicians, and other health professionals, who are not officers or employees of the United States, and who represent the specialties and disciplines relevant to the diseases with respect to which the Advisory Board is established; and

(B) six members from the general public who are knowledgeable with respect to such diseases, including at least one member who is a person who has such a disease and one member who is a parent of a person who has such a disease.

Of the appointed members at least five shall by virtue of training or experience be knowledgeable in the fields of health education, nursing, data systems, public information, and community program development.

(2)(A) The following shall be ex officio members of each Advisory Board:

(i) The Assistant Secretary for Health, the Director of NIH, the Director of the National Institute of Diabetes and Digestive and Kidney Diseases, the Director of the Centers for Disease Control and Prevention, the Under
Secretary for Health of the Department of Veterans Affairs, the Assistant Secretary of Defense for Health Affairs, and the Division Director of the National Institute of Diabetes and Digestive and Kidney Diseases for the diseases for which the Board is established (or the designees of such officers).

(ii) Such other officers and employees of the United States as the Secretary determines necessary for the Advisory Board to carry out its functions.

(B) In the case of the National Diabetes Advisory Board, the following shall also be ex officio members: The Director of the National Heart, Lung, and Blood Institute, the Director of the National Eye Institute, the Director of the Eunice Kennedy Shriver National Institute of Child Health and Human Development, and the Administrator of the Health Resources and Services Administration (or the designees of such officers).

(c) Compensation

Members of an Advisory Board who are officers or employees of the Federal Government shall serve as members of the Advisory Board without compensation in addition to that received in their regular public employment. Other members of the Board shall receive compensation at rates not to exceed the daily equivalent of the annual rate in effect for grade GS–18 of the General Schedule for each day (including traveltime) they are engaged in the performance of their duties as members of the Board.

(d) Term of office; vacancy

The term of office of an appointed member of an Advisory Board is four years, except that no term of office may extend beyond the expiration of the Advisory Board. Any member appointed to fill a vacancy for an unexpired term shall be appointed for the remainder of such term. A member may serve after the expiration of the member’s term until a successor has taken office. If a vacancy occurs in an Advisory Board, the Secretary shall make an appointment to fill the vacancy not later than 90 days from the date the vacancy occurred.

(e) Chairman

The members of each Advisory Board shall select a chairman from among the appointed members.

(f) Executive director; professional and clerical staff; administrative support services and facilities

The Secretary shall, after consultation with and consideration of the recommendations of an Advisory Board, provide the Advisory Board with an executive director and one other professional staff member. In addition, the Secretary shall, after consultation with and consideration of the recommendations of the Advisory Board, provide the Advisory Board with such additional professional staff members, such clerical staff members, such services of consultants, such information, and (through contracts or other arrangements) such administrative support services and facilities, as the Secretary determines are necessary for the Advisory Board to carry out its functions.

(g) Meetings

Each Advisory Board shall meet at the call of the chairman or upon request of the Director of the Institute, but not less often than four times a year.

(h) Functions of National Diabetes Advisory Board and National Digestive Diseases Advisory Board

The National Diabetes Advisory Board and the National Digestive Diseases Advisory Board shall—

1. review and evaluate the implementation of the plan (referred to in section 285c–7 of this title) respecting the diseases with respect to which the Advisory Board was established and periodically update the plan to ensure its continuing relevance;

2. for the purpose of assuring the most effective use and organization of resources respecting such diseases, advise and make recommendations to the Congress, the Secretary, the Director of NIH, the Director of the Institute, and the heads of other appropriate Federal agencies for the implementation and revision of such plan; and

3. maintain liaison with other advisory bodies related to Federal agencies involved in the implementation of such plan, the coordinating committee for such diseases, and with key non-Federal entities involved in activities affecting the control of such diseases.

(i) Subcommittees; establishment and membership

In carrying out its functions, each Advisory Board may establish subcommittees, convene workshops and conferences, and collect data. Such subcommittees may be composed of Advisory Board members and nonmember consultants with expertise in the particular area addressed by such subcommittees. The subcommittees may hold such meetings as are necessary to enable them to carry out their activities.

(j) Termination of predecessor boards; time within which to appoint members

The National Diabetes Advisory Board and the National Digestive Diseases Advisory Board in existence on November 20, 1985, shall terminate upon the appointment of a successor Board under subsection (a) of this section. The Secretary shall make appointments to the Advisory Boards established under subsection (a) of this section before the expiration of 90 days after November 20, 1985. The members of the Boards in existence on November 20, 1985, may be appointed, in accordance with subsections (b) and (d) of this section, to the Boards established under subsection (a) of this section for diabetes and digestive diseases, except that at least one-half of the members of the National Diabetes Advisory Board in existence on November 20, 1985, shall be appointed to the National Diabetes Advisory Board first established under subsection (a) of this section.
§ 285c-5. Research and training centers; development or expansion

(a) Diabetes mellitus and related endocrine and metabolic diseases

(1) Consistent with applicable recommendations of the National Commission on Diabetes, the Director of the Institute shall provide for the development or substantial expansion of centers for research and training in diabetes mellitus and related endocrine and metabolic diseases. Each center developed or expanded under this subsection shall—

(A) utilize the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such research and training qualifications as may be prescribed by the Secretary; and

(B) conduct—

(i) research in the diagnosis and treatment of diabetes mellitus and related endocrine and metabolic diseases and the complications resulting from such diseases;

(ii) training programs for physicians and allied health personnel in current methods of diagnosis and treatment of such diseases and complications, and in research in diabetes; and

(iii) information programs for physicians and allied health personnel who provide primary care for patients with such diseases or complications.

(2) A center may use funds provided under paragraph (1) to provide stipends for nurses and allied health professionals enrolled in research training programs described in paragraph (1)(B)(ii).

(b) Digestive diseases and related functional, congenital, metabolic disorders, and normal development of digestive tract

Consistent with applicable recommendations of the National Digestive Diseases Advisory Board, the Director shall provide for the development or substantial expansion of centers for research in digestive diseases and related functional, congenital, metabolic disorders, and normal development of the digestive tract. Each center developed or expanded under this subsection—

(1) shall utilize the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such research qualifications as may be prescribed by the Secretary;

(2) shall develop and conduct basic and clinical research into the cause, diagnosis, early detection, prevention, control, and treatment of digestive diseases and nutritional disorders and related functional, congenital, or metabolic complications resulting from such diseases or disorders;

(3) shall encourage research into and programs for—

(A) providing information for patients with such diseases and the families of such patients, physicians and others who care for such patients, and the general public;

(B) model programs for cost effective and preventive patient care; and

(C) training physicians and scientists in research on such diseases, disorders, and complications; and

(4) may perform research and participate in epidemiological studies and data collection relevant to digestive diseases and disorders and disseminate such research, studies, and data to the health care profession and to the public.
(c) Kidney and urologic diseases

The Director shall provide for the development or substantial expansion of centers for research in kidney and urologic diseases. Each center developed or expanded under this subsection—

(1) shall utilize the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such research qualifications as may be prescribed by the Secretary;

(2) shall develop and conduct basic and clinical research into the cause, diagnosis, early detection, prevention, control, and treatment of kidney and urologic diseases;

(3) shall encourage research into and programs for—

(A) providing information for patients with such diseases, disorders, and complications and the families of such patients, physicians and others who care for such patients, and the general public;

(B) model programs for cost effective and preventive patient care; and

(C) training physicians and scientists in research on such diseases; and

(4) may perform research and participate in epidemiological studies and data collection relevant to kidney and urologic diseases in order to disseminate such research, studies, and data to the health care profession and to the public.

(d) Nutritional disorders

(1) The Director of the Institute shall, subject to the extent of amounts made available in appropriations Acts, provide for the development or substantial expansion of centers for research and training regarding nutritional disorders, including obesity.

(2) The Director of the Institute shall carry out paragraph (1) in collaboration with the Director of the National Cancer Institute and with the Directors of such other agencies of the National Institutes of Health as the Director of NIH determines to be appropriate.

Each center developed or expanded under paragraph (1) shall—

(A) utilize the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such research and training qualifications as may be prescribed by the Director;

(B) conduct basic and clinical research into the cause, diagnosis, early detection, prevention, control and treatment of nutritional disorders, including obesity and the impact of nutrition and diet on child development;

(C) conduct training programs for physicians and allied health professionals in current methods of diagnosis and treatment of such diseases and complications, and in research in such disorders; and

(D) conduct information programs for physicians and allied health professionals who provide primary care for patients with such disorders or complications.

(e) Geographic distribution; period of support, additional periods

Insofar as practicable, centers developed or expanded under this section should be geographically dispersed throughout the United States and in environments with proven research capabilities. Support of a center under this section may be for a period of not to exceed five years and such period may be extended by the Director of the Institute for additional periods of not more than five years each if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director and if such group has recommended to the Director that such period should be extended.


AMENDMENTS

1993—Subsecs. (d), (e). Pub. L. 103–43 added subsec. (d) and redesignated former subsec. (d) as (e).

§ 285c–6. Advisory council subcommittees

There are established within the advisory council for the Institute appointed under section 284a of this title a subcommittee on diabetes and endocrine and metabolic diseases, a subcommittee on digestive diseases and nutrition, and a subcommittee on kidney, urologic, and hematologic diseases. The subcommittees shall be composed of members of the advisory council who are outstanding in the diagnosis, prevention, and treatment of the diseases for which the subcommittees are established and members of the advisory council who are leaders in the fields of education and public affairs. The subcommittees are authorized to review applications made to the Director of the Institute for grants for research and training projects relating to the diagnosis, prevention, and treatment of the diseases for which the subcommittees are established and shall recommend to the advisory council those applications and contracts that the subcommittees determine will best carry out the purposes of the Institute. The subcommittees shall also review and evaluate the diabetes and endocrine and metabolic diseases, digestive diseases and nutrition, and kidney, urologic, and hematologic diseases programs of the Institute and recommend to the advisory council such changes in the administration of such programs as the subcommittees determine are necessary.

(July 1, 1944, ch. 373, title IV, §432, as added Pub. L. 99–158, §2, Nov. 20, 1985, 99 Stat. 847.)


The Director of the Institute shall prepare for inclusion in the biennial report made under section 284b1 of this title a description of the Institute’s activities—

(1) under the current diabetes plan under the National Diabetes Mellitus Research and Education Act; and

(2) under the current digestive diseases plan formulated under the Arthritis, Diabetes, and Digestive Diseases Amendments of 1976.

The description submitted by the Director shall include an evaluation of the activities of the

1See References in Text note below.
centers supported under section 285c-5 of this title.
(July 1, 1944, ch. 373, title IV, § 433, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 848.)

REFERENCES IN TEXT

The National Diabetes Mellitus Research and Education Act, referred to in par. (1), is Pub. L. 93–334, July 23, 1974, 88 Stat. 373, as amended, which enacted former sections 289c–1a, 289c–2, and 289c–3 of this title, amended section 247b and former section 289c–1 of this title, and enacted provisions formerly set out as notes under section 289c–3 of this title. For complete classification of this Act to the Code, see Short Title of 1974 Amendments note set out under section 201 of this title and Tables.


§ 285c–8. Nutritional disorders program

(a) Establishment
The Director of the Institute, in consultation with the Director of NIH, shall establish a program of conducting and supporting research, training, health information dissemination, and other activities with respect to nutritional disorders, including obesity.

(b) Support of activities
In carrying out the program established under subsection (a) of this section, the Director of the Institute shall conduct and support each of the activities described in such subsection.

(c) Dissemination of information
In carrying out the program established under subsection (a) of this section, the Director of the Institute shall carry out activities to facilitate and enhance knowledge and understanding of nutritional disorders, including obesity, on the part of health professionals, patients, and the public through the effective dissemination of information.
(July 1, 1944, ch. 373, title IV, § 434, as added Pub. L. 103–43, title VI, §601[(a)], June 10, 1993, 107 Stat. 161.)

§ 285c–9. Juvenile diabetes

(a) Long-term epidemiology studies
The Director of the Institute shall conduct or support long-term epidemiology studies in which individuals with or at risk for type 1, or juvenile, diabetes are followed for 10 years or more. Such studies shall investigate the causes and characteristics of the disease and its complications.

(b) Clinical trial infrastructure/innovative treatments for juvenile diabetes
The Secretary, acting through the Director of the National Institutes of Health, shall support regional clinical research centers for the prevention, detection, treatment, and cure of juvenile diabetes.

(c) Prevention of type 1 diabetes
The Secretary, acting through the appropriate agencies, shall provide for a national effort to prevent type 1 diabetes. Such effort shall provide for a combination of increased efforts in research and development of prevention strategies, including consideration of vaccine development, coupled with appropriate ability to test the effectiveness of such strategies in large clinical trials of children and young adults.

AMENDMENTS
2007—Subsec. (d). Pub. L. 109–482 struck out heading and text of subsec. (d). Text read as follows: “For the purpose of carrying out this section, there are authorized to be appropriated such sums as may be necessary for each of the fiscal years 2001 through 2006.”

EFFECTIVE DATE OF 2007 AMENDMENT
Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 201 of this title.

SUBPART 4—NATIONAL INSTITUTE OF ARTHRITIS AND MUSCULOSKELETAL AND SKIN DISEASES

§ 285d. Purpose of Institute

The general purpose of the National Institute of Arthritis and Musculoskeletal and Skin Diseases (hereafter in this subpart referred to as the “Institute”) is the conduct and support of research and training, the dissemination of health information, and other programs with respect to arthritis and musculoskeletal and skin diseases (including sports-related disorders), with particular attention to the effect of these diseases on children.

AMENDMENTS
1993—Pub. L. 103–43 substituted “(including sports-related disorders), with particular attention to the effect of these diseases on children” for “, including sports-related disorders’’.

§ 285d–1. National arthritis and musculoskeletal and skin diseases program

(a) Plan to expand, intensify, and coordinate activities; submission; periodic review and revision
The Director of the Institute, with the advice of the Institute’s advisory council, shall prepare and transmit to the Director of NIH a plan for a national arthritis and musculoskeletal and skin diseases program to expand, intensify, and coordinate the activities of the Institute respecting arthritis and musculoskeletal and skin diseases. The plan shall include such comments and recommendations as the Director of the In-
stitute determines appropriate. The plan shall place particular emphasis upon expanding research into better understanding the causes and the development of effective treatments for arthritis affecting children. The Director of the Institute shall periodically review and revise such plan and shall transmit any revisions of such plan to the Director of NIH.

(b) Coordination of activities with other national research institutes; minimum activities under program

Activities under the national arthritis and musculoskeletal and skin diseases program shall be coordinated with the other national research institutes to the extent that such institutes have responsibilities respecting arthritis and musculoskeletal and skin diseases, and shall, at least, provide for—

(1) investigation into the epidemiology, etiology, and prevention of all forms of arthritis and musculoskeletal and skin diseases, including sports-related disorders, primarily through the support of basic research in such areas as immunology, genetics, biochemistry, microbiology, physiology, bioengineering, and any other scientific discipline which can contribute important knowledge to the treatment and understanding of arthritis and musculoskeletal and skin diseases;

(2) research into the development, trial, and evaluation of techniques, drugs, and devices used in the diagnosis, treatment, including medical rehabilitation, and prevention of arthritis and musculoskeletal and skin diseases;

(3) research on the refinement, development, and evaluation of technological devices that will replace or be a substitute for damaged bone, muscle, and joints and other supporting structures;

(4) the establishment of mechanisms to monitor the causes of athletic injuries and identify ways of preventing such injuries on scholastic athletic fields; and

(5) research into the causes of arthritis affecting children and the development, trial, and evaluation of techniques, drugs and devices used in the diagnosis, treatment (including medical rehabilitation), and prevention of arthritis in children.

(c) Program to be carried out in accordance with plan

The Director of the Institute shall carry out the national arthritis and musculoskeletal and skin diseases program in accordance with the plan prepared under subsection (a) of this section and any revisions of such plan made under such subsection.

The Secretary shall establish an Arthritis and Musculoskeletal Diseases Interagency Coordinating Committee and a Skin Diseases Interagency Coordinating Committee (hereafter in this section individually referred to as a “Committee”).

(b) Membership; chairman; meetings

Each Committee shall be composed of the Directors of each of the national research institutes and divisions involved in research regarding the diseases with respect to which the Committee is established, the Under Secretary for Health of the Department of Veterans Affairs, and the Assistant Secretary of Defense for Health Affairs (or the designees of such officers), and representatives of all other Federal departments and agencies (as determined by the Secretary) whose programs involve health functions or responsibilities relevant to arthritis and musculoskeletal diseases or skin diseases, as the case may be. Each Committee shall be chaired by the Director of NIH (or the designee of the Director). Each Committee shall meet at the call of the chairman, but not less often than four times a year.

(Provisions related to the establishment and operation of the Arthritis and Musculoskeletal Diseases Interagency Coordinating Committee and the Skin Diseases Interagency Coordinating Committee, including the establishment of subcommittees, the function of the committees, and the responsibilities of the committees.)

The Director shall provide for the standardization of patient data and recordkeeping.

The Director shall provide for the standardization of patient data and recordkeeping for the collection, storage, analysis, retrieval, and dissemination of such data in cooperation with projects assisted under this section, centers assisted under section 285d–6 of this title, and other persons engaged in arthritis and musculoskeletal disease programs.

§ 285d-6. Multipurpose arthritis and musculoskeletal diseases centers

(a) Development, modernization, and operation

The Director of the Institute shall, after consultation with the advisory council for the Institute, provide for the development, modernization, and operation (including staffing and other
operating costs such as the costs of patient care required for research) of new and existing centers for arthritis and musculoskeletal diseases. For purposes of this section, the term “modernization” means the alteration, remodeling, expansion, and repair of existing buildings and the provision of equipment for such buildings to the extent necessary to make them suitable for use as centers described in the preceding sentence.

(b) Duties and functions

Each center assisted under this section shall—
1. (A) use the facilities of a single institution or a consortium of cooperating institutions, and (B) meet such qualifications as may be prescribed by the Secretary; and
2. conduct—
   (A) basic and clinical research into the cause, diagnosis, early detection, prevention, control, and treatment of and rehabilitation from arthritis and musculoskeletal diseases and complications resulting from arthritis and musculoskeletal diseases, including research into implantable biomaterials and biomechanical and other orthopedic procedures;
   (B) training programs for physicians, scientists, and other health and allied health professionals;
   (C) information and continuing education programs for physicians and other health and allied health professionals who provide care for patients with arthritis and musculoskeletal diseases; and
   (D) programs for the dissemination to the general public of information—
      (i) on the importance of early detection of arthritis and musculoskeletal diseases, of seeking prompt treatment, and of following an appropriate regimen; and
      (ii) to discourage the promotion and use of unapproved and ineffective diagnostic, preventive, treatment, and control methods and unapproved and ineffective drugs and devices.

A center may use funds provided under subsection (a) of this section to provide stipends for health professionals enrolled in training programs described in paragraph (2)(B).

(c) Optional programs

Each center assisted under this section may conduct programs to—
1. establish the effectiveness of new and improved methods of detection, referral, and diagnosis of individuals with a risk of developing arthritis and musculoskeletal diseases;
2. disseminate the results of research, screening, and other activities, and develop means of standardizing patient data and recordkeeping; and
3. develop community consultative services to facilitate the referral of patients to centers for treatment.

(d) Geographical distribution

The Director of the Institute shall, insofar as practicable, provide for an equitable geographical distribution of centers assisted under this section. The Director shall give appropriate consideration to the need for centers especially suited to meeting the needs of children affected by arthritis and musculoskeletal diseases.

(e) Period of support; additional periods

Support of a center under this section may be for a period of not to exceed five years. Such period may be extended by the Director of the Institute for one or more additional periods of not more than five years if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director and if such group has recommended to the Director that such period should be extended.

(f) Treatment and rehabilitation of children

Not later than October 1, 1993, the Director shall establish a multipurpose arthritis and musculoskeletal disease center for the purpose of expanding the level of research into the causes, diagnosis, early detection, prevention, control, and treatment of, and rehabilitation of children with arthritis and musculoskeletal diseases.

(1) In general

The Director of the Institute shall expand and intensify research and related activities of the Institute with respect to lupus.

(b) Coordination with other institutes

The Director of the Institute shall coordinate the activities of the Director under subsection (a) of this section with similar activities conducted by the other national research institutes and agencies of the National Institutes of Health to the extent that such Institutes and agencies have responsibilities that are related to lupus.

(c) Programs for lupus

In carrying out subsection (a) of this section, the Director of the Institute shall conduct or support research to expand the understanding of the causes of, and to find a cure for, lupus. Activities under such subsection shall include conducting and supporting the following:
1. Research to determine the reasons underlying the elevated prevalence of lupus in women, including African-American women.
2. Basic research concerning the etiology and causes of the disease.
3. Epidemiological studies to address the frequency and natural history of the disease and the differences among the sexes and among racial and ethnic groups with respect to the disease.
4. The development of improved diagnostic techniques.
5. Clinical research for the development and evaluation of new treatments, including new biological agents.
The Secretary shall establish in the Institute the National Arthritis and Musculoskeletal and Skin Diseases Advisory Board (hereafter in this section referred to as the “Advisory Board”).

(b) Membership; ex officio members

The Advisory Board shall be composed of twenty appointed members and nonvoting, ex officio members, as follows:

(1) The Secretary shall appoint—

(A) twelve members from individuals who are scientists, physicians, and other health professionals, who are not officers or employees of the United States, and who represent the specialties and disciplines relevant to arthritis, musculoskeletal diseases, and skin diseases; and

(B) eight members from the general public who are knowledgeable with respect to such diseases, including one member who is a person who has such a disease, one person who is the parent of an adult with such a disease, and two members who are parents of children with arthritis.

Of the appointed members at least five shall by virtue of training or experience be knowledgeable in health education, nursing, data systems, public information, or community program development.

(2) The following shall be ex officio members of the Advisory Board:

(A) the Assistant Secretary for Health, the Director of the National Institute of Arthritis and Musculoskeletal and Skin Diseases, the Director of the Centers for Disease Control and Prevention, the Under Secretary for Health of the Department of Veterans Affairs, and the Assistant Secretary of Defense for Health Affairs (or the designees of such officers), and

(B) such other officers and employees of the United States as the Secretary determines necessary for the Advisory Board to carry out its functions.

(c) Compensation

Members of the Advisory Board who are officers or employees of the Federal Government shall serve as members of the Advisory Board without compensation in addition to that received in their regular public employment. Other members of the Advisory Board shall receive compensation at rates not to exceed the daily equivalent of the annual rate in effect for grade GS–18 of the General Schedule for each day (including traveltime) they are engaged in the performance of their duties as members of the Advisory Board.

(d) Term of office; vacancy

The term of office of an appointed member of the Advisory Board is four years. Any member appointed to fill a vacancy for an unexpired term shall be appointed for the remainder of such term. A member may serve after the expiration of the member’s term until a successor has taken office. If a vacancy occurs in the Advisory Board, the Secretary shall make an appointment to fill the vacancy not later than 90 days after the date the vacancy occurred.

(e) Chairman

The members of the Advisory Board shall select a chairman from among the appointed members.

(f) Executive director, professional and clerical staff; administrative support services and facilities

The Secretary shall, after consultation with and consideration of the recommendations of the Advisory Board, provide the Advisory Board with an executive director and one other professional staff member. In addition, the Secretary shall, after consultation with and consideration of the recommendations of the Advisory Board, provide the Advisory Board with such additional
professional staff members, such clerical staff members, and (through contracts or other arrangements) with such administrative support services and facilities, such information, and such services of consultants, as the Secretary determines are necessary for the Advisory Board to carry out its functions.

(g) Meetings
The Advisory Board shall meet at the call of the chairman or upon request of the Director of the Institute, but not less often than four times a year.

(h) Duties and functions
The Advisory Board shall—

(1) review and evaluate the implementation of the plan prepared under subsection (a) of this title and periodically update the plan to ensure its continuing relevance;

(2) for the purpose of assuring the most effective use and organization of resources respecting arthritis, musculoskeletal diseases, and skin diseases, advise and make recommendations to the Congress, the Secretary, the Director of NIH, the Director of the Institute, and the heads of other appropriate Federal agencies for the implementation and revision of such plan; and

(3) maintain liaison with other advisory bodies for Federal agencies involved in the implementation of such plan, the interagency coordinating committees for such diseases established under section 285d–4 of this title, and with key non-Federal entities involved in activities affecting the control of such diseases.

(i) Subcommittees; establishment and membership
In carrying out its functions, the Advisory Board may establish subcommittees, convene workshops and conferences, and collect data. Such subcommittees may be composed of Advisory Board members and nonmember consultants with expertise in the particular area addressed by such subcommittees. The subcommittees may hold such meetings as are necessary to enable them to carry out their activities.

(j) Termination of predecessor board; time within which to appoint members
The National Arthritis Advisory Board in existence on November 20, 1985, shall terminate upon the appointment of a successor Board under subsection (a) of this section. The Secretary shall make appointments to the Advisory Board established under subsection (a) of this section before the expiration of 90 days after November 20, 1985. The member of the Board in existence on November 20, 1985, may be appointed, in accordance with subsections (b) and (d) of this section, to the Advisory Board established under subsection (a) of this section.

(EMENDMENTS)

2007—Subsecs. (j), (k). Pub. L. 109–482 redesignated subsec. (k) as (j) and struck out former subsec. (j) which required the Advisory Board to prepare an annual report for the Secretary and set out the subjects for report.

1993—Subsec. (a). Pub. L. 102–531, § 312(d)(6), substituted “twenty” for “eighteen” in introductory provisions, “eight” for “six” and “including one member who is a person who has such a disease, one person who is the parent of an adult with such a disease, and two members who are parents of children with arthritis” for “including at least one member who is a person who has such a disease and one member who is a parent of a person who has such a disease, and two persons who have such a disease”.


1991—Subsec. (j). Pub. L. 102–405 substituted “Chief Medical Director” for “Chief Medical Director for Health”.

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

TERMINATION OF ADVISORY BOARDS

Advisory boards established after Jan. 5, 1973, to terminate not later than the expiration of the 2-year period beginning on the date of their establishment, unless, in the case of a board established by the President or an officer of the Federal Government, such board is renewed by appropriate action prior to the expiration of such 2-year period, or in the case of a board established by the Congress, its duration is otherwise provided by law. See sections 2(j)(1) and 22(d) of Pub. L. 92–584, Oct. 6, 1972, 88 Stat. 770, 776, set out in the Appendix to Title 5, Government Organization and Employees.

Pub. L. 92–584, § 6, Jan. 4, 1973, 88 Stat. 2275, set out as a note under section 217 of this title, provided that an advisory committee established pursuant to the Public Health Service Act shall terminate at such time as may be specifically prescribed by an Act of Congress enacted after Jan. 4, 1973.

REFERENCES IN OTHER LAWS TO GS–16, 17, OR 18 PAY RATES

References in laws to the rates of pay for GS–16, 17, or 18, to or maximum rates of pay under the General Schedule, to be considered references to rates payable under specified sections of Title 5, Government Organization and Employees, see section 529 (title 1, § 181) of Pub. L. 101–509, set out in a note under section 5376 of Title 5.

§ 285d–8. Juvenile arthritis and related conditions

(a) Expansion and coordination of activities
The Director of the Institute, in coordination with the Director of the National Institute of Allergy and Infectious Diseases, shall expand and intensify the programs of such Institutes with respect to research and related activities concerning juvenile arthritis and related conditions.

(b) Coordination
The Directors referred to in subsection (a) of this section shall jointly coordinate the pro-
grams referred to in such subsection and consult with the Arthritis and Musculoskeletal Diseases Interagency Coordinating Committee.


AMENDMENTS

2007—Subsec. (c). Pub. L. 109–482 struck out heading and text of subsec. (c). Text read as follows: “For the purpose of carrying out this section, there are authorized to be appropriated such sums as may be necessary for each of the fiscal years 2001 through 2005.”

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

SUBPART 5—NATIONAL INSTITUTE ON AGING

§ 285e. Purpose of Institute

The general purpose of the National Institute on Aging (hereafter in this subpart referred to as the ‘‘Institute’’) is the conduct and support of basic biomedical, social, and behavioral research, training, health information dissemination, and other programs with respect to the aging process and the diseases and other special problems and needs of the aged.

(July 1, 1944, ch. 373, title IV, §443, as added Pub. L. 99–158, §2, Nov. 20, 1985, 99 Stat. 854.)

STUDY OF MALNUTRITION IN ELDERLY

Pub. L. 103–43, title XIX, §1902, June 10, 1993, 107 Stat. 201, directed Secretary of Health and Human Services, acting through National Institute on Aging, to conduct a 3-year study on health benefits and cost-effectiveness of nutrition screening and intervention activities of the elderly, and a 3-year study to determine extent of malnutrition in elderly individuals in hospitals and long-term care facilities and in elderly individuals who are living independently, provided for creation of advisory panel to oversee studies, provided for submission to Congress of reports containing findings of such studies, and provided for termination of advisory panel 3 years after June 10, 1993.

STUDY OF PERSONNEL FOR HEALTH NEEDS OF ELDERLY

Section 8 of Pub. L. 99–158 directed Secretary to conduct a study on the adequacy and availability of personnel to meet the current and projected health needs (including needs for home and community-based care) of elderly Americans through the year 2020, and report the results of the study, with recommendations, to Congress by Mar. 1, 1987.

§ 285e–1. Special functions

(a) Education and training of adequate numbers of personnel

In carrying out the training responsibilities under this chapter or any other Act for health and allied health professions personnel, the Secretary shall take appropriate steps to insure the education and training of adequate numbers of allied health, nursing, and paramedical personnel in the field of health care for the aged.

(b) Scientific studies

The Director of the Institute shall conduct scientific studies to measure the impact on the biological, medical, social, and psychological aspects of aging of programs and activities assisted or conducted by the Department of Health and Human Services.

(c) Public information and education programs

The Director of the Institute shall carry out public information and education programs designed to disseminate as widely as possible the findings of research sponsored by the Institute, other relevant aging research and studies, and other information about the process of aging which may assist elderly and near-elderly persons in dealing with, and all Americans in understanding, the problems and processes associated with growing older.

(d) Grants for research relating to Alzheimer’s Disease

The Director of the Institute shall make grants to public and private nonprofit institutions to conduct research relating to Alzheimer’s Disease.

(July 1, 1944, ch. 373, title IV, §444, as added Pub. L. 99–158, §2, Nov. 20, 1985, 99 Stat. 854.)

§ 285e–2. Alzheimer’s Disease centers

(a) Cooperative agreements and grants for establishing and supporting

(1) The Director of the Institute may enter into cooperative agreements with and make grants to public or private nonprofit entities (including university medical centers) to pay all or part of the cost of planning, establishing, or strengthening, and providing basic operating support (including staffing) for centers for basic and clinical research (including multidisciplinary research) into, training in, and demonstration of advanced diagnostic, prevention, and treatment methods for Alzheimer’s disease.

(2) A cooperative agreement or grant under paragraph (1) shall be entered into in accordance with policies established by the Director of NIH and after consultation with the Institute’s advisory council.

(b) Use of Federal payments under cooperative agreement or grant

(1) Federal payments made under a cooperative agreement or grant under subsection (a) of this section may, with respect to Alzheimer’s disease, be used for—

(A) diagnostic examinations, patient assessments, patient care costs, and other costs necessary for conducting research;

(B) training, including training for allied health professionals;

(C) diagnostic and treatment clinics designed to meet the special needs of minority and rural populations and other underserved populations;

(D) activities to educate the public; and

(E) the dissemination of information.

(2) For purposes of paragraph (1), the term ‘‘training’’ does not include research training for which Ruth L. Kirschstein National Research Service Awards may be provided under section 288 of this title.

(c) Support period; additional periods

Support of a center under subsection (a) of this section may be for a period of not to exceed
five years. Such period may be extended by the Director for additional periods of not more than five years each if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director and if such group has recommended to the Director that such period should be extended. 


AMENDMENTS
1990—Subsec. (a)(1). Pub. L. 101–557, § 201(1), inserted “(including university medical centers)” after “nonprofit entities”, “(including staffing)” after “operating support”, and “(including multidisciplinary research)” after “clinical research” and substituted “Alzheimer’s disease” for “Alzheimer’s Disease”.
Subsec. (b). Pub. L. 101–557, § 201(2), amended subsec. (b) generally. Prior to amendment, subsec. (b) read as follows: “Federal payments made under a cooperative agreement or grant under subsection (a) of this section may be used for—

(1) construction (notwithstanding any limitation under section 289e of this title);

(2) staffing and other basic operating costs, including such patient care costs as are required for research;

(3) training, including training for allied health professionals; and

(4) demonstration purposes.

As used in this subsection, the term ‘construction’ does not include the acquisition of land, and the term ‘training’ does not include research training for which National Research Service Awards may be provided under section 288 of this title.”

ALZHEIMER’S DISEASE RESEARCH
Pub. L. 100–175, title III, Nov. 29, 1987, 101 Stat. 972, provided that:

“SEC. 301. REQUIREMENT FOR CLINICAL TRIALS.

“(a) IN GENERAL.—The Director of the National Institute on Aging shall provide for the conduct of clinical trials on the efficacy of the use of such promising therapeutic agents as have been or may be discovered and recommended for further scientific analysis by the National Institute on Aging and the Food and Drug Administration to treat individuals with Alzheimer’s disease, to retard the progression of symptoms of Alzheimer’s disease, or to improve the functioning of individuals with such disease. 

“(b) RULE OF CONSTRUCTION.—Nothing in this title shall be construed to affect adversely any research being conducted as of the date of the enactment of this Act (Nov. 29, 1987).

“SEC. 302. AUTHORIZATION OF APPROPRIATIONS.

“For the purpose of carrying out section 301, there is authorized to be appropriated $2,000,000 for fiscal year 1988.”

ALZHEIMER’S DISEASE REGISTRY
Section 12 of Pub. L. 99–158, which was formerly set out as a note under this section, was renumbered section 4462 of the Public Health Service Act by Pub. L. 103–43, title VIII, § 801(a), June 10, 1993, 107 Stat. 163, and is classified to section 258e–9 of this title.

§ 285e–3. Claude D. Pepper Older Americans Independence Centers

(a) Development and expansion of centers

The Director of the Institute shall enter into cooperative agreements with, and make grants to, public and private nonprofit entities for the development or expansion of not less than 10 centers of excellence in geriatric research and training of researchers. Each such center shall be known as a Claude D. Pepper Older Americans Independence Center.

(b) Functions of centers

Each center developed or expanded under this section shall—

(1) utilize the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such research and training qualifications as may be prescribed by the Director; and

(2) conduct—

(A) research into the aging processes and into the diagnosis and treatment of diseases, disorders, and complications related to aging, including menopause, which research includes research on such treatments, and

on medical devices and other medical interventions regarding such diseases, disorders, and complications, that can assist individuals in avoiding institutionalization and prolonged hospitalization and in otherwise increasing the independence of the individuals; and

(B) programs to develop individuals capable of conducting research described in subparagraph (A).

(c) Geographic distribution of centers

In making cooperative agreements and grants under this section for the development or expansion of centers, the Director of the Institute shall ensure that, to the extent practicable, any such centers are distributed equitably among the principal geographic regions of the United States.

(d) “Independence” defined

For purposes of this section, the term “independence”, with respect to diseases, disorders, and complications of aging, means the functional ability of individuals to perform activities of daily living without assistance or supervision. 


AMENDMENTS


American Public Health Association
§ 285e–4. Awards for leadership and excellence in Alzheimer’s disease and related dementias

(a) Senior researchers in biomedical research

The Director of the Institute shall make awards to senior researchers who have made distinguished achievements in biomedical research in areas relating to Alzheimer’s disease and related dementias. Awards under this section shall be used by the recipients to support research in areas relating to such disease and dementias, and may be used by the recipients to train junior researchers who demonstrate exceptional promise to conduct research in such areas.

(b) Eligible centers

The Director of the Institute may make awards under this section only to researchers at centers supported by the National Institute on Aging, other programs and centers in-volved in research on Alzheimer’s disease and related dementias supported by the National Institute of Mental Health on the psychopathology of the elderly, relevant activities of the Administration on Aging, other programs and centers involved in research on Alzheimer’s disease and related dementias supported by Federal agencies other than the Department, State or local agencies, community organizations, or private foundations.

(c) Required recommendation

The Director of the Institute shall make awards under this section only to researchers who have been recommended for such awards by the National Advisory Council on Aging.

(d) Selection procedures

The Director of the Institute shall establish procedures for the selection of the recipients of awards under this section.

(e) Term of award; renewal

Awards under this section shall be made for a one-year period, and may be renewed for not more than six additional consecutive one-year periods.

(1) Within 6 months after November 14, 1986, the Director of the Institute shall prepare and transmit to the Chairman of the Council on Alzheimer’s Disease (in this section referred to as the “Council”) a plan for the research to be conducted under subsection (a) of this section. The plan shall—

(A) provide for research concerning—

(i) the epidemiology of, and the identification of risk factors for, Alzheimer’s disease and related dementias; and

(ii) the development and evaluation of reliable and valid multidimensional diagnostic and assessment procedures and instruments; and

(B) ensure that research carried out under the plan is coordinated with, and uses, to the maximum extent feasible, resources of, other Federal programs relating to Alzheimer’s disease and related dementias, including centers supported under section 285e–2 of this title, centers supported by the National Institute of Mental Health on the psychopathology of the elderly, relevant activities of the Administration on Aging, other programs and centers involved in research on Alzheimer’s disease and related dementias supported by Federal agencies other than the Department, State or local agencies, community organizations, or private foundations.

(2) Within one year after transmitting the plan required under paragraph (1), and annually thereafter, the Director of the Institute shall prepare and transmit to the Chairman of the Council such revisions of such plan as the Director considers appropriate.

(c) Consultation for preparation and revision of plan

In preparing and revising the plan required by subsection (b) of this section, the Director of the Institute shall consult with the Chairman of the Council and the heads of agencies within the Department.

(d) Grants for promoting independence and preventing secondary disabilities

the Director of the Institute may develop, or make grants to develop—

(1) model techniques to—

1So in original. Probably should be capitalized.
(A) promote greater independence, including enhanced independence in performing activities of daily living and instrumental activities of daily living, for persons with Alzheimer’s disease and related disorders; and
(B) prevent or reduce the severity of secondary disabilities, including confusional episodes, falls, bladder and bowel incontinence, and adverse effects of prescription and over-the-counter medications, in such persons; and
(2) model curricula for health care professionals, health care paraprofessionals, and family caregivers, for training and application in the use of such techniques.

e) “Council on Alzheimer’s Disease” defined

For purposes of this section, the term “Council on Alzheimer’s Disease” means the council established in section 11211(a) of this title.


References in Text


Codification

Section was formerly classified to section 11241 of this title prior to renumbering by Pub. L. 100–607.

Amendments


1988—Pub. L. 100–607, §142(a), renumbered section 11241 of this title as this section.

Codification

Section was formerly classified to section 11241 of this title prior to renumbering by Pub. L. 100–607.

References in Text

Section 11211 of this title, referred to in subsec. (a)(3), inserted “on Alzheimer’s Disease (in this section referred to as the ‘Council’)” after “Council”.


Section was formerly classified to section 11241 of this title.

Codification

Section was formerly classified to section 11241 of this title prior to renumbering by Pub. L. 100–607.

Amendments

1988—Pub. L. 100–607, §142(a), renumbered section 11241 of this title as this section.

Fee for Information

The Clearinghouse may charge an appropriate fee for information provided through the toll-free telephone line established under subsection (a)(3).1

(c) Summaries of research findings from other agencies

The Director of the Institute, the Director of the National Institute of Mental Health, and the Director of the National Center for Health Services Research and Health Care Technology Assessment shall provide to the Clearinghouse summaries of the findings of research conducted under part D.


References in Text


1 See References in Text note below.

§285e–6. Dissemination of research results

The Director of the Institute shall disseminate the results of research conducted under section 285e–5 of this title and this section to appropriate professional entities and to the public.


Codification

Section was formerly classified to section 11242 of this title prior to renumbering by Pub. L. 100–607.

Amendments

1988—Pub. L. 100–607, §142(a), renumbered section 11242 of this title as this section.

Fee for Information

The Clearinghouse may charge an appropriate fee for information provided through the toll-free telephone line established under subsection (a)(3).1

(c) Summaries of research findings from other agencies

The Director of the Institute, the Director of the National Institute of Mental Health, and the Director of the National Center for Health Services Research and Health Care Technology Assessment shall provide to the Clearinghouse summaries of the findings of research conducted under part D.


References in Text


1 See References in Text note below.

So in original. No subsec. (a)(3) has been enacted.
§ 285e–8. Dissemination project

(a) Grant or contract for establishment

The Director of the Institute shall make a grant to, or enter into a contract with, a national organization representing individuals with Alzheimer's disease and related dementias for the conduct of the activities described in subsection (b) of this section.

(b) Project activities

The organization receiving a grant or contract under this section shall—

(1) establish a central computerized information system to—

(A) compile and disseminate information concerning initiatives by State and local governments and private entities to provide programs and services for individuals with Alzheimer's disease and related dementias; and

(B) translate scientific and technical information concerning such initiatives into information readily understandable by the general public, and make such information available upon request; and

(2) establish a national toll-free telephone line to make available the information described in paragraph (1), and information concerning Federal programs, services, and benefits for individuals with Alzheimer's disease and related dementias and their families.

c) Fees for information; exception

The organization receiving a grant or contract under this section may charge appropriate fees for information provided through the toll-free telephone line established under subsection (b)(2) of this section, and may make exceptions to such fees for individuals and organizations who are not financially able to pay such fees.

d) Application for grant or contract; contents

In order to receive a grant or contract under this section, an organization shall submit an application to the Director of the Institute. Such application shall contain—

(1) information demonstrating that such organization has a network of contacts which will enable such organization to receive information necessary to the operation of the central computerized information system described in subsection (b)(1) of this section;

(2) information demonstrating that, by the end of fiscal year 1991, such organization will be financially able to, and will, carry out the activities described in subsection (b) of this section without a grant or contract from the Federal Government; and

(3) such other information as the Director may prescribe.


CODIFICATION

Section was formerly classified to section 11282 of this title prior to renumbering by Pub. L. 100–607.

AMENDMENTS

1988—Pub. L. 100–607, § 142(a), renumbered section 11282 of this title as this section.

Subsecs. (a), (d). Pub. L. 100–607, § 142(d)(5), substituted “the Institute” for “the National Institute on Aging”.

§ 285e–9. Alzheimer's disease registry

(a) In general

The Director of the Institute may make a grant to develop a registry for the collection of epidemiological data about Alzheimer's disease and its incidence in the United States, to train personnel in the collection of such data, and for other matters respecting such disease.

(b) Qualifications

To qualify for a grant under subsection (a) of this section an applicant shall—

(1) be an accredited school of medicine or public health which has expertise in the collection of epidemiological data about individuals with Alzheimer's disease and in the development of disease registries, and

(2) have access to a large patient population, including a patient population representative of diverse ethnic backgrounds.


CODIFICATION

Section was formerly set out as a note under section 285e–2 of this title prior to renumbering by Pub. L. 103–43.

AMENDMENTS


Subsec. (a). Pub. L. 103–43, § 801(b)(1), substituted in heading “In general” for “Grant authority” and in text substituted “Director of the Institute” for “Director of the National Institute on Aging”.

Subsec. (c). Pub. L. 103–43, § 801(b)(2), struck out subsec. (c) which authorized appropriations of $2,500,000 for grants to remain available until expended or through fiscal year 1989, whichever occurred first.

§ 285e–10. Aging processes regarding women

The Director of the Institute, in addition to other special functions specified in section 285e–1 of this title and in cooperation with the Directors of the other national research institutes and agencies of the National Institutes of Health, shall conduct research into the aging processes of women, with particular emphasis given to the effects of menopause and the physiological and behavioral changes occurring during the transition from pre- to post-menopause, and into the diagnosis, disorders, and complications related to aging and loss of ovarian hormones in women.


§ 285f. Purpose of Institute

The general purpose of the National Institute of Allergy and Infectious Diseases is the conduct and support of research, training, health information dissemination, and other programs with respect to allergic and immunologic diseases and disorders and infectious diseases, including tropical diseases.


§ 285f–1. Research centers regarding chronic fatigue syndrome

(a) The Director of the Institute, after consultation with the advisory council for the Institute, may make grants to, or enter into contracts with, public or nonprofit private entities for the development and operation of centers to conduct basic and clinical research on chronic fatigue syndrome.

(b) Each center assisted under this section shall use the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such requirements as may be prescribed by the Director of the Institute.

(July 1, 1944, ch. 373, title IV, § 447, as added Pub. L. 103–43, title IX, § 902(a), June 10, 1993, 107 Stat. 164.)

Codification

Another section 447 of act July 1, 1944, was renumbered section 447A and is classified to section 285f–2 of this title.

§ 285f–2. Extramural study section

Section 902(b) of Pub. L. 103–43 provided that: “Not later than 6 months after the date of enactment of this Act [June 10, 1993], the Secretary of Health and Human Services shall establish an extramural study section for chronic fatigue syndrome research.”

§ 285f–3. Research activities on chronic fatigue syndrome

Section 1903 of Pub. L. 103–43 directed Secretary of Health and Human Services to, not later than Oct. 1, 1993, and annually thereafter for next 3 years, prepare
and submit to Congress a report that summarizes re-
search activities conducted or supported by National
Institutes of Health concerning chronic fatigue syn-
drome, with information concerning grants made, coop-
operative agreements or contracts entered into, intra-
mural activities, research priorities and needs, and
plan to address such priorities and needs.

§ 285f–2. Research and research training regard-
ing tuberculosis

In carrying out section 285f of this title, the
Director of the Institute shall conduct or support
research and research training regarding the cause,
diagnosis, early detection, prevention and treatment of tuberculosis.

(July 1, 1944, ch. 373, title IV, § 447A, as added

AMENDMENTS

2007—Pub. L. 109–482 struck out subsec. (a) designa-
tion before “In carrying out” and subsec. (b) which read as follows: “For the purpose of carrying out sub-
section (a) of this section, there are authorized to be appropriated $50,000,000 for fiscal year 1994, and such
sums as may be necessary for each of the fiscal years
1995 through 1998. Such authorization is in addition to
any other authorization of appropriations that is avail-
able for such purpose.”

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with
respect to amounts appropriated for fiscal year 2007 or
subsequent fiscal years, see section 109 of Pub. L.
109–482, set out as a note under section 281 of this title.

RESEARCH THROUGH FOOD AND DRUG ADMINISTRATION

Section 303 of Pub. L. 103–183 provided that: “The Secretary of Health and Human Services, acting through the Commissioner of Food and Drugs, shall im-
plement a tuberculosis drug and device research pro-
gram under which the Commissioner may—
“(1) provide assistance to other Federal agencies for the development of tuberculosis protocols;
“(2) review and evaluate medical devices designed for the diagnosis and control of airborne tuberculosis; and
“(3) conduct research concerning drugs or devices to be used in diagnosing, controlling and preventing tuberculosis.”

§ 285f–3. Sexually transmitted disease clinical re-
search and training awards

(a) In general

The Director of the Institute is authorized to estab-
lish and maintain a program to enhance and promote the translation of new scientific knowledge into clinical practice related to the diagnosis, care and treatment of individuals with sexually transmitted diseases.

(b) Support of promising clinicians

In order to foster the application of the most current developments in the etiology, patho-
genesis, diagnosis, prevention and treatment of sexually transmitted diseases, amounts made available under this section shall be directed to the support of promising clinicians through awards for research, study, and practice at centers of excellence in sexually transmitted disease research and treatment.

(c) Excellence in certain fields

Research shall be carried out under awards made under subsection (b) of this section in en-
vvironments of demonstrated excellence in the etiology and pathogenesis of sexually transmit-
ted diseases and shall foster innovation and inte-
gration of such disciplines or other environ-
ments determined suitable by the Director of the Institute.

(July 1, 1944, ch. 373, title IV, § 447B, as added
Pub. L. 106–505, title IX, § 901, Nov. 13, 2000, 114

AMENDMENTS

2007—Subsec. (d). Pub. L. 109–482 struck out heading and text of subsec. (d). Text read as follows: “For the purpose of carrying out this section, there are authorized to be appropriated $2,250,000 for fiscal year 2001, and such sums as may be necessary for each of fiscal years 2002 through 2005.”

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with
respect to amounts appropriated for fiscal year 2007 or
subsequent fiscal years, see section 109 of Pub. L.
109–482, set out as a note under section 281 of this title.

§ 285f–4. Microbicide research and development

The Director of the Institute, acting through the head of the Division of AIDS, shall, consist-
ent with the peer-review process of the National Institutes of Health, carry out research on, and
development of, safe and effective methods for use by women to prevent the transmission of the human immunoand other environments determined suitable by the Director of the Institute.

(July 1, 1944, ch. 373, title IV, § 447C, as added

SUBPART 7—EUNICE KENNEDY SHRIVER NATIONAL INSTITUTE OF CHILD HEALTH AND HUMAN DEVELOPMENT

AMENDMENTS


§ 285g. Purpose of Institute

The general purpose of the Eunice Kennedy Shriver National Institute of Child Health and Human Development (hereafter in this subpart referred to as the ‘‘Institute’’) is the conduct and support of research, training, health information dissemination, and other programs with respect to gynecologic health, maternal health, child health, intellectual disabilities, human growth and development, including prenatal develop-
ment, population research, and special health problems and requirements of mothers and children.

AMENDMENTS

CHANGE OF NAME
“Eunice Kennedy Shriver National Institute of Child Health and Human Development” substituted for “National Institute of Child Health and Development” in text, on authority of section 1(d) of Pub. L. 110–154, set out below.

EUNICE KENNEDY SHRIVER NATIONAL INSTITUTE OF CHILD HEALTH AND HUMAN DEVELOPMENT; FINDINGS

“(1) Since it was established by Congress in 1962 at the request of President John F. Kennedy, the National Institutes of Child Health and Human Development has achieved an outstanding record of achievement in catalyzing a concentrated attack on the unsolved health problems of children and of mother-infant relationships by fulfilling its mission to—

“(A) ensure that every individual is born healthy and wanted, that women suffer no harmful effects from reproductive processes, and that all children have the chance to achieve their full potential for healthy and productive lives, free from disease or disability; and

“(B) ensure the health, productivity, independence, and well-being of all individuals through optimal rehabilitation.

“(2) The National Institute of Child Health and Development has made unparalleled contributions to the advancement of child health and human development, including significant efforts to—

“(A) reduce dramatically the rates of Sudden Infant Death Syndrome, infant mortality, and maternal HIV transmission;

“(B) develop the Haemophilus Influenza B (Hib) vaccine, credited with nearly eliminating the incidence of intellectual disabilities; and

“(C) conduct intramural research, support extramural research, and train thousands of child health and human development researchers who have contributed greatly to dramatic gains in child health throughout the world.

“(3) The vision, drive, and tenacity of one woman, Eunice Kennedy Shriver, was instrumental in proposing, passing, and enacting legislation to establish the National Institutes of Child Health and Human Development (Public Law 87–838) (see Tables for classification) on October 17, 1962.

“(4) It is befitting and appropriate to recognize the substantial achievements of Eunice Kennedy Shriver, a tireless advocate for children with special needs, whose foresight in creating the National Institute of Child Health and Human Development gave life to the words of President Kennedy, who wished to ‘encourage imaginative research into the complex processes of human development from conception to old age.’:"

[For definition of “intellectual disabilities” in section 1(a) of Pub. L. 110–154, set out above, see Definitions note below.]

LONG-TERM CHILD DEVELOPMENT STUDY

“(a) PURPOSE.—It is the purpose of this section to authorize the Eunice Kennedy Shriver National Institute of Child Health and Human Development to conduct a national longitudinal study of environmental influences (including physical, chemical, biological, and psychosocial) on children’s health and development.

“(b) IN GENERAL.—The Director of the Eunice Kennedy Shriver National Institute of Child Health and Human Development shall establish a consortium of representatives from appropriate Federal agencies (including the Centers for Disease Control and Prevention, the Environmental Protection Agency, and the Department of Education) to—

“(1) plan, develop, and implement a prospective cohort study, from birth to adulthood, to evaluate the effects of both chronic and intermittent exposures on child health and human development; and

“(2) investigate basic mechanisms of developmental disorders and environmental factors, both risk and protective, that influence health and developmental processes.

“(c) REQUIREMENT.—The study under subsection (b) shall—

“(1) incorporate behavioral, emotional, educational, and contextual consequences to enable a complete assessment of the physical, chemical, biological, and psychosocial environmental influences on children’s well-being;

“(2) gather data on environmental influences and outcomes on diverse populations of children, which may include the consideration of prenatal exposures; and

“(3) consider health disparities among children which may include the consideration of prenatal exposures; and

“(4) be conducted in compliance with section 444 of the General Education Provisions Act (20 U.S.C. 1232j), including the requirement of prior parental consent for the disclosure of any education records, except without the use of authority or exceptions granted to authorized representatives of the Secretary of Education for the evaluation of Federally-supported education programs or in connection with the enforcement of the Federal legal requirements that relate to such programs.


NATIONAL COMMISSION TO PREVENT INFANT MORTALITY: COMPOSITION; VOLUNTARY SERVICES; DURATION
Pub. L. 100–436, title IV, Sept. 20, 1988, 102 Stat. 1709, provided that the National Commission to Prevent Infant Mortality was to be composed of sixteen members, including seven at large members, and that it had power to accept voluntary and uncompensated services, notwithstanding section 142 of title 31, and was to continue operating, notwithstanding sections 208 and 209 of Pub. L. 99–660 (formerly set out below).

NATIONAL COMMISSION TO PREVENT INFANT MORTALITY

DEFINITIONS
For meaning of references to an intellectual disability and to individuals with intellectual disabilities in
provisions amended by section 2 of Pub. L. 111–256, see section 2(k) of Pub. L. 111–256, set out as a note under section 1400 of Title 20, Education.

§ 285g–1. Sudden Infant Death Syndrome Research

The Director of the Institute shall conduct and support research which specifically relates to sudden infant death syndrome.

(July 1, 1944, ch. 373, title IV, § 449, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 856.)

§ 285g–2. Research on Intellectual Disabilities

The Director of the Institute shall conduct and support research and related activities into the causes, prevention, and treatment of intellectual disabilities.


AMENDMENTS
2010—Pub. L. 111–256 amended section generally. Prior to amendment, text read as follows: "The Director of the Institute shall conduct and support research and related activities into the causes, prevention, and treatment of mental retardation."

DEFINITIONS
For meaning of references to an intellectual disability and to individuals with intellectual disabilities in provisions amended by section 2 of Pub. L. 111–256, see section 2(k) of Pub. L. 111–256, set out as a note under section 1400 of Title 20, Education.

§ 285g–3. Associate Director for Prevention; Appointment; Function

There shall be in the Institute an Associate Director for Prevention to coordinate and promote the programs in the Institute concerning the prevention of health problems of mothers and children. The Associate Director shall be appointed by the Director of the Institute from individuals who because of their professional training or experience are experts in public health or preventive medicine.


AMENDMENTS
1998—Pub. L. 105–362 struck out subsec. (a) designation and struck out subsec. (b) which read as follows: "The Associate Director for Prevention shall prepare for inclusion in the biennial report made under section 284b of this title a description of the prevention activities of the Institute, including a description of the staff and resources allocated to those activities."

§ 285g–4. National Center for Medical Rehabilitation Research

(a) Establishment of Center

There shall be in the Institute an agency to be known as the National Center for Medical Rehabilitation Research (hereafter in this section referred to as the "Center"). The Director of the Institute shall appoint a qualified individual to serve as Director of the Center. The Director of the Center shall report directly to the Director of the Institute.

(b) Purpose

The general purpose of the Center is the conduct and support of research and research training (including research on the development of orthotic and prosthetic devices), the dissemination of health information, and other programs with respect to the rehabilitation of individuals with physical disabilities resulting from diseases or disorders of the neurological, musculoskeletal, cardiovascular, pulmonary, or any other physiological system (hereafter in this section referred to as "medical rehabilitation").

(c) Authority of Director

(1) In carrying out the purpose described in subsection (b) of this section, the Director of the Center may—

(A) provide for clinical trials regarding medical rehabilitation;

(B) provide for research regarding model systems of medical rehabilitation;

(C) coordinate the activities of the Center with similar activities of other agencies of the Federal Government, including the other agencies of the National Institutes of Health, and with similar activities of other public entities and of private entities;

(D) support multidisciplinary medical rehabilitation research conducted or supported by more than one such agency;

(E) in consultation with the advisory council for the Institute and with the approval of the Director of NIH—

(i) establish technical and scientific peer review groups in addition to those appointed under section 282(b)(16) of this title; and

(ii) appoint the members of peer review groups established under subparagraph (A); and

(F) support medical rehabilitation research and training centers.

The Federal Advisory Committee Act shall not apply to the duration of a peer review group appointed under subparagraph (E).

(2) In carrying out this section, the Director of the Center may make grants and enter into cooperative agreements and contracts.

(d) Research Plan

(1) In consultation with the Director of the Center, the coordinating committee established under subsection (e) of this section, and the advisory board established under subsection (f) of this section, the Director of the Institute shall develop a comprehensive plan for the conduct and support of medical rehabilitation research (hereafter in this section referred to as the "Research Plan").

(2) The Research Plan shall—

(A) identify current medical rehabilitation research activities conducted or supported by the Federal Government, opportunities and needs for additional research, and priorities for such research; and

(B) make recommendations for the coordination of such research conducted or supported by the National Institutes of Health and other agencies of the Federal Government.

(3)(A) Not later than 18 months after the date of the enactment of the National Institutes of
Health. Revitalization Amendments of 1990, the Director of the Institute shall transmit the Research Plan to the Director of NIH, who shall submit the Plan to the President and the Congress.

(3) Subparagraph (A) shall be carried out independently of the process of reporting that is required in sections 283 and 284b of this title.

(4) The Director of the Institute shall periodically revise and update the Research Plan as appropriate, after consultation with the Director of the Center, the coordinating committee established under subsection (e) of this section, and the advisory board established under subsection (f) of this section. A description of any revisions in the Research Plan shall be contained in each report prepared under section 284b of this title by the Director of the Institute.

(e) Medical Rehabilitation Coordinating Committee

(1) The Director of NIH shall establish a committee to be known as the Medical Rehabilitation Coordinating Committee (hereafter in this section referred to as the “Coordinating Committee”).

(2) The Coordinating Committee shall make recommendations to the Director of the Institute and the Director of the Center with respect to the content of the Research Plan and with respect to the activities of the Center that are carried out in conjunction with other agencies of the National Institutes of Health and with other agencies of the Federal Government.

(3) The Coordinating Committee shall be composed of the Director of the Institute, the Director of the National Institute on Aging, the National Institute of Arthritis and Musculoskeletal and Skin Diseases, the National Heart, Lung, and Blood Institute, the National Institute of Neurological Disorders and Stroke, and such other national research institutes and such representatives of the National Institute on Aging.

(f) National Advisory Board on Medical Rehabilitation Research

(1) Not later than 90 days after the date of the enactment of the National Institutes of Health Revitalization Amendments of 1990, the Director of NIH shall establish a National Advisory Board on Medical Rehabilitation Research (hereafter in this section referred to as the “Advisory Board”).

(2) The Advisory Board shall review and assess Federal research priorities, activities, and findings regarding medical rehabilitation research, and shall advise the Director of the Center and the Director of the Institute on the provisions of the Research Plan.

(3)(A) The Director of NIH shall appoint to the Advisory Board 18 qualified representatives of the public who are not officers or employees of the Federal Government. Of such members, 12 shall be representatives of health and scientific disciplines with respect to medical rehabilita-

1 See References in Text note below.
§ 285g–5. Research centers with respect to contraception and infertility

(a) Grants and contracts

The Director of the Institute, after consultation with the advisory council for the Institute, shall make grants to, or enter into contracts with, public or nonprofit private entities for the development and operation of centers to conduct activities for the purpose of improving methods of contraception and centers to conduct activities for the purpose of improving diagnosis and treatment of infertility.

(b) Number of centers

In carrying out subsection (a) of this section, the Director of the Institute shall, subject to the extent of amounts made available in appropriations Acts, provide for the establishment of three centers with respect to contraception and for two centers with respect to infertility.

(c) Duties

(1) Each center assisted under this section shall, in carrying out the purpose of the center involved—

(A) conduct clinical and other applied research, including—

(i) for centers with respect to contraception, clinical trials of new or improved drugs and devices for use by males and females (including barrier methods); and

(ii) for centers with respect to infertility, clinical trials of new or improved drugs and devices for the diagnosis and treatment of infertility in males and females;

(B) develop protocols for training physicians, scientists, nurses, and other health and allied health professionals;

(C) conduct training programs for such individuals;

(D) develop model continuing education programs for such professionals; and

(E) disseminate information to such professionals and the public.

(2) A center may use funds provided under subsection (a) of this section to provide stipends for health and allied health professionals enrolled in programs described in subparagraph (C) of paragraph (1), and to provide fees to individuals serving as subjects in clinical trials conducted under such paragraph.

(d) Coordination of information

The Director of the Institute shall, as appropriate, provide for the coordination of information among the centers assisted under this section.

(e) Facilities

Each center assisted under subsection (a) of this section shall use the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such requirements as may be prescribed by the Director of the Institute.

(f) Period of support

Support of a center under subsection (a) of this section may be for a period not exceeding 5 years. Such period may be extended for one or more additional periods not exceeding 5 years if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director and if such group has recommended to the Director that such period should be extended.

2007—Subsec. (g). Pub. L. 109–482 struck out subsec. (g) which read as follows: “For the purpose of carrying out this section, there are authorized to be appropriated $30,000,000 for fiscal year 1994, and such sums as may be necessary for each of the fiscal years 1995 and 1996.”

Amendments

2007—Subsec. (g). Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

§ 285g–6. Program regarding obstetrics and gynecology

The Director of the Institute shall establish and maintain within the Institute an intramural laboratory and clinical research program in obstetrics and gynecology.

2007—Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

§ 285g–7. Child health research centers

The Director of the Institute shall develop and support centers for conducting research with respect to child health. Such centers shall give priority to the expeditious transfer of advances from basic science to clinical applications and improving the care of infants and children.
§ 285g–8. Prospective longitudinal study on adolescent health

(a) In general

Not later than October 1, 1993, the Director of the Institute shall commence a study for the purpose of providing information on the general health and well-being of adolescents in the United States, including, with respect to such adolescents, information on—

(1) the behaviors that promote health and the behaviors that are detrimental to health; and

(2) the influence on health of factors particular to the communities in which the adolescents reside.

(b) Design of study

(1) In general

The study required in subsection (a) of this section shall be a longitudinal study in which a substantial number of adolescents participate as subjects. With respect to the purpose described in such subsection, the study shall monitor the subjects throughout the period of the study to determine the health status of the subjects and any change in such status over time.

(2) Population-specific analyses

The study required in subsection (a) of this section shall be conducted with respect to the population of adolescents who are female, the population of adolescents who are male, various socioeconomic populations of adolescents, and various racial and ethnic populations of adolescents. The study shall be designed and conducted in a manner sufficient to provide for a valid analysis of whether there are significant differences among such populations in health status and whether and to what extent any such differences are due to factors particular to the populations involved.

(c) Coordination with Women's Health Initiative

With respect to the national study of women being conducted by the Secretary and known as the Women's Health Initiative, the Secretary shall ensure that such study is coordinated with the component of the study required in subsection (a) of this section that concerns adolescents, including coordination in the design of the 2 studies.

§ 285g–9. Fragile X

(a) Expansion and coordination of research activities

The Director of the Institute, after consultation with the advisory council for the Institute, shall expand, intensify, and coordinate the activities of the Institute with respect to research on the disease known as fragile X.

(b) Research centers

(1) In general

The Director of the Institute shall make grants or enter into contracts for the development and operation of centers to conduct research for the purposes of improving the diagnosis and treatment of, and finding the cure for, fragile X.

(2) Number of centers

(A) In general

In carrying out paragraph (1), the Director of the Institute shall, to the extent that amounts are appropriated, and subject to subparagraph (B), provide for the establishment of at least three fragile X research centers.

(B) Peer review requirement

The Director of the Institute shall make a grant to, or enter into a contract with, an entity for purposes of establishing a center under paragraph (1) only if the grant or contract has been recommended after technical and scientific peer review required by regulations under section 289a of this title.

(3) Activities

The Director of the Institute shall, as appropriate, provide for the coordination of the activities of the centers assisted under this section, including providing for the exchange of information among the centers.

(4) Coordination among centers

The Director of the Institute shall, as appropriate, provide for the coordination of the activities of the centers established under paragraph (1) only if the grant or contract has been recommended after technical and scientific peer review required by regulations under section 289a of this title.

(5) Certain administrative requirements

Each center assisted under paragraph (1) shall use the facilities of a single institution, or be formed from a consortium of cooperating institutions, meeting such requirements as may be prescribed by the Director of the Institute.

(6) Duration of support

Support may be provided to a center under paragraph (1) for a period not exceeding 5 years. Such period may be extended for one or more additional periods, each of which may not exceed 5 years, if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director and if such group has recommended to the Director that such period be extended.

AMENDMENTS

2007—Subsec. (b)(7). Pub. L. 109–482 struck out heading and text of par. (7). Text read as follows: “For the purpose of carrying out this subsection, there are authorized to be appropriated such sums as may be necessary for each of the fiscal years 2001 through 2005.”

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.
§ 285g-10. Investment in tomorrow's pediatric researchers

In order to ensure the future supply of researchers dedicated to the care and research needs of children, the Director of the Institute, after consultation with the Administrator of the Health Resources and Services Administration, shall support activities to provide for—

(1) an increase in the number and size of institutional training grants to institutions supporting pediatric training; and

(2) an increase in the number of career development awards for health professionals who intend to build careers in pediatric basic and clinical research, including pediatric pharmacological research.


AMENDMENTS

2007—Pub. L. 109–482 struck out subsec. (a) designation and heading before “In order to” and struck out heading and text of subsec. (b). Text read as follows: “For the purpose of carrying out subsection (a) of this section, there are authorized to be appropriated such sums as may be necessary for each of the fiscal years 2006 through 2008.”

Par. (2). Pub. L. 110–85 inserted “including pediatric pharmacological research” before period at end.

Effective Date of 2007 Amendment

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

SUBPART 8—NATIONAL INSTITUTE OF DENTAL RESEARCH

§ 285h. Purpose of Institute

The general purpose of the National Institute of Dental Research is the conduct and support of research, training, health information dissemination, and other programs with respect to the cause, prevention, and methods of diagnosis and treatment of dental and oral diseases and conditions.

(July 1, 1944, ch. 373, title IV, § 453H, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 856.)

SUBPART 9—NATIONAL EYE INSTITUTE

§ 285i. Purpose of Institute

The general purpose of the National Eye Institute (hereafter in this subpart referred to as the “Institute”) is the conduct and support of research, training, health information dissemination, and other programs with respect to blinding eye diseases, visual disorders, mechanisms of visual function, preservation of sight, and the special health problems and requirements of the blind. Subject to section 285i–1 of this title, the Director of the Institute may carry out a program of grants for public and private nonprofit vision research facilities.


AMENDMENTS

1993—Pub. L. 103–43 substituted “Subject to section 285i–1 of this title, the Director” for “The Director” in second sentence.

§ 285i–1. Clinical research on eye care and diabetes

(a) Program of grants

The Director of the Institute, in consultation with the advisory council for the Institute, may award research grants to one or more Diabetes Eye Research Institutions for the support of programs in clinical or health services aimed at—

(1) providing comprehensive eye care services for people with diabetes, including a full complement of preventive, diagnostic and treatment procedures;

(2) developing new and improved techniques of patient care through basic and clinical research;

(3) assisting in translation of the latest research advances into clinical practice; and

(4) expanding the knowledge of the eye and diabetes through further research.

(b) Use of funds

Amounts received under a grant awarded under this section shall be used for the following:

(1) Establishing the biochemical, cellular, and genetic mechanisms associated with diabetic eye disease and the earlier detection of pending eye abnormalities. The focus of work under this paragraph shall require that ophthalmologists have training in the most up-to-date molecular and cell biological methods.

(2) Establishing new frontiers in technology, such as video-based diagnostic and research resources, to—

(A) provide improved patient care;

(B) provide for the evaluation of retinal physiology and its effect on diabetes; and

(C) provide for the assessment of risks for the development and progression of diabetic eye disease and a more immediate evaluation of various therapies aimed at preventing diabetic eye disease.

Such technologies shall be designed to permit evaluations to be performed both in humans and in animal models.

(3) The translation of the results of vision research into the improved care of patients with diabetic eye disease. Such translation shall require the application of institutional resources that encompass patient care, clinical research and basic laboratory research.

(4) The conduct of research concerning the outcomes of eye care treatments and eye health education programs as they relate to patients with diabetic eye disease, including the evaluation of regional approaches to such research.

(c) Authorized expenditures

The purposes for which a grant under subsection (a) of this section may be expended include equipment for the research described in such subsection.
(July 1, 1944, ch. 373, title IV, § 456, as added Pub. L. 103–43, title XI, §1101(a), June 10, 1993, 107 Stat. 168.)

SUBPART 10—NATIONAL INSTITUTE OF NEUROLOGICAL DISORDERS AND STROKE

AMENDMENTS

§ 285j. Purpose of Institute

The general purpose of the National Institute of Neurological Disorders and Stroke (hereafter in this subpart referred to as the “Institute”) is the conduct and support of research, training, health information dissemination, and other programs with respect to neurological disease and disorder and stroke.


AMENDMENTS
1988—Pub. L. 100–553 and Pub. L. 100–607 made identical amendments, substituting “Neurological Disorders” for “Neurological and Communicative Disorders” and “and disorder and stroke” for “disorder, stroke, and disorders of human communication”. Pub. L. 100–690 amended this section to read as if the amendments by Pub. L. 100–607 had not been enacted.

EFFECTIVE DATE OF 1988 AMENDMENT
For effective date of amendment by Pub. L. 100–690, see section 2613(b)(1) of Pub. L. 100–690, set out as an Effect of Enactment of Similar Provisions note under section 285m of this title.

§ 285j–1. Spinal cord regeneration research

The Director of the Institute shall conduct and support research into spinal cord regeneration.

(July 1, 1944, ch. 373, title IV, § 458, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 857.)

INTERAGENCY COMMITTEE ON SPINAL CORD INJURY
Section 7 of Pub. L. 99–158 provided that:
“(a) Establishment.—Within 90 days after the date of enactment of this Act [Nov. 20, 1985], the Secretary of Health and Human Services shall establish in the National Institute of Neurological and Communicative Diseases and Stroke an Interagency Committee on Spinal Cord Injury (hereafter in this section referred to as the ‘Interagency Committee’). The Interagency Committee shall plan, develop, coordinate, and implement comprehensive Federal initiatives in research on spinal cord injury and regeneration.
“(b) Committee Composition and Meetings.—(1) The Interagency Committee shall consist of representatives from—
“(A) the National Institute on Neurological and Communicative Disorders and Stroke; “(B) the Department of Defense; “(C) the Department of Education; “(D) the Veterans’ Administration; “(E) the Office of Science and Technology Policy; and “(F) the National Science Foundation; designated by the heads of such entities.
“(2) The Interagency Committee shall meet at least four times. The Secretary of Health and Human Services shall select the Chairman of the Interagency Committee from the members of the Interagency Committee.
“(c) Report.—Within the 18 months after the date of enactment of this Act [Nov. 20, 1985], the Interagency Committee shall prepare and transmit to the Congress a report concerning its activities under this section. The report shall include a description of research projects on spinal cord injury and regeneration conducted or supported by Federal agencies during such 18-month period, the nature and purpose of each such project, the amounts expended for each such project, and an identification of the entity which conducted the research under each such project.
“(d) Termination.—The Interagency Committee shall terminate 90 days after the date on which the Interagency Committee transmits the report required by subsection (c) to the Congress.”

§ 285j–2. Bioengineering research

The Director of the Institute shall make grants or enter into contracts for research on the means to overcome paralysis of the extremities through electrical stimulation and the use of computers.

(July 1, 1944, ch. 373, title IV, § 459, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 857.)


The Director of the Institute shall conduct and support research on multiple sclerosis, especially research on effects of genetics and hormonal changes on the progress of the disease.

(July 1, 1944, ch. 373, title IV, § 460, as added Pub. L. 103–43, title XII, §1201, June 10, 1993, 107 Stat. 169.)

SUBPART 11—NATIONAL INSTITUTE OF GENERAL MEDICAL SCIENCES

§ 285k. Purpose of Institute

The general purpose of the National Institute of General Medical Sciences is the conduct and support of research, training, and, as appropriate, health information dissemination, and other programs with respect to general or basic medical sciences and related natural or behavioral sciences which have significance for two or more other national research institutes or are outside the general area of responsibility of any other national research institute.

(July 1, 1944, ch. 373, title IV, § 461, as added Pub. L. 99–158, § 2, Nov. 20, 1985, 99 Stat. 857.)

SUBPART 12—NATIONAL INSTITUTE OF ENVIRONMENTAL HEALTH SCIENCES

§ 285l. Purpose of Institute

The general purpose of the National Institute of Environmental Health Sciences (in this subpart referred to as the “Institute”) is the conduct and support of research, training, health information dissemination, and other programs
with respect to factors in the environment that affect human health, directly or indirectly.


AMENDMENTS


§ 285–1. Applied Toxicological Research and Testing Program

(a) There is established within the Institute a program for conducting applied research and testing regarding toxicology, which program shall be known as the Applied Toxicological Research and Testing Program.

(b) In carrying out the program established under subsection (a) of this section, the Director of the Institute shall, with respect to toxicology, carry out activities—

    (1) to expand knowledge of the health effects of environmental agents;
    (2) to broaden the spectrum of toxicology information that is obtained on selected chemicals;
    (3) to develop and validate assays and protocols, including alternative methods that can reduce or eliminate the use of animals in acute or chronic safety testing;
    (4) to establish criteria for the validation and regulatory acceptance of alternative testing and to recommend a process through which scientifically validated alternative methods can be accepted for regulatory use;
    (5) to communicate the results of research to government agencies, to medical, scientific, and regulatory communities, and to the public; and
    (6) to integrate related activities of the Department of Health and Human Services.

(July 1, 1944, ch. 373, title IV, § 463A, as added Pub. L. 103–43, title XIII, § 1301(a), June 10, 1993, 107 Stat. 169.)

§ 285–2. Definitions

In sections 285–2 to 285–5 of this title:

(1) Alternative test method

The term “alternative test method” means a test method that—

    (A) includes any new or revised test method; and
    (B)(i) reduces the number of animals required;
    (ii) refines procedures to lessen or eliminate pain or distress to animals, or enhances animal well-being; or
    (iii) replaces animals with non-animal systems or one animal species with a phylogenetically lower animal species, such as replacing a mammal with an invertebrate.

(2) ICCVAM test recommendation

The term “ICCVAM test recommendation” means a summary report prepared by the ICCVAM characterizing the results of a scientific expert peer review of a test method.

§ 285–3. Interagency Coordinating Committee on the Validation of Alternative Methods

(a) In general

With respect to the interagency coordinating committee that is known as the Interagency Coordinating Committee on the Validation of Alternative Methods (referred to in sections 285–2 to 285–5 of this title as “ICCVAM”) and that was established by the Director of the National Institute of Environmental Health Sciences for purposes of section 285–1(b) of this title, the Director of the Institute shall designate such committee as a permanent interagency coordinating committee of the Institute under the National Toxicology Program Interagency Center for the Evaluation of Alternative Toxicological Methods. Sections 285–2 to 285–5 of this title may not be construed as affecting the authorities of such Director regarding ICCVAM that were in effect on the day before December 19, 2000, except to the extent inconsistent with sections 285–2 to 285–5 of this title.

(b) Purposes

The purposes of the ICCVAM shall be to—

    (1) increase the efficiency and effectiveness of Federal agency test method review;
    (2) eliminate unnecessary duplicative efforts and share experiences between Federal regulatory agencies;
    (3) optimize utilization of scientific expertise outside the Federal Government;
    (4) ensure that new and revised test methods are validated to meet the needs of Federal agencies; and
    (5) reduce, refine, or replace the use of animals in testing, where feasible.

(c) Composition

The ICCVAM shall be composed of the heads of the following Federal agencies (or their designees):

    (1) Agency for Toxic Substances and Disease Registry.
    (3) Department of Agriculture.
    (4) Department of Defense.
    (5) Department of Energy.
    (6) Department of the Interior.
    (7) Department of Transportation.
    (8) Environmental Protection Agency.
    (9) Food and Drug Administration.
    (10) National Institute for Occupational Safety and Health.
    (11) National Institutes of Health.
    (12) National Cancer Institute.
    (13) National Institute of Environmental Health Sciences.
    (14) National Library of Medicine.
    (15) Occupational Safety and Health Administration.
    (16) Any other agency that develops, or employs tests or test data using animals, or regulates on the basis of the use of animals in toxicity testing.
(d) **Scientific Advisory Committee**

(1) **Establishment**

The Director of the National Institute of Environmental Health Sciences shall establish a Scientific Advisory Committee (referred to in sections 285–2 to 285–5 of this title as the “SAC”) to advise ICCVAM and the National Toxictology Program Interagency Center for the Evaluation of Alternative Toxicological Methods regarding ICCVAM activities. The activities of the SAC shall be subject to provisions of the Federal Advisory Committee Act.

(2) **Membership**

(A) **In general**

The SAC shall be composed of the following voting members:

(1) At least one knowledgeable representative having a history of expertise, development, or evaluation of new or revised or alternative test methods from each of—

(I) the personal care, pharmaceutical, industrial chemicals, or agriculture industry;

(II) any other industry that is regulated by the Federal agencies specified in subsection (c) of this section; and

(III) a national animal protection organization established under section 501(c)(3) of title 26.

(ii) Representatives (selected by the Director of the National Institute of Environmental Health Sciences) from an academic institution, a State government agency, an international regulatory body, or any corporation developing or marketing new or revised or alternative test methodologies, including contract laboratories.

(B) **Nonvoting ex officio members**

The membership of the SAC shall, in addition to voting members under subparagraph (A), include as nonvoting ex officio members the agency heads specified in subsection (c) of this section (or their designees).

(e) **Duties**

The ICCVAM shall, consistent with the purposes described in subsection (b) of this section, carry out the following functions:

(1) Review and evaluate new or revised or alternative test methods, including batteries of tests and test screens, that may be acceptable for specific regulatory uses, including the coordination of technical reviews of proposed new or revised or alternative test methods of interagency interest.

(2) Facilitate appropriate interagency and international harmonization of acute or chronic toxicological test protocols that encourage the reduction, refinement, or replacement of animal test methods.

(3) Facilitate and provide guidance on the development of validation criteria, validation studies and processes for new or revised or alternative test methods and help facilitate the acceptance of such scientifically valid test methods and awareness of accepted test methods by Federal agencies and other stakeholders.

(4) Submit ICCVAM test recommendations for the test method reviewed by the ICCVAM, through expeditious transmittal by the Secretary of Health and Human Services (or the designee of the Secretary), to each appropriate Federal agency, along with the identification of specific agency guidelines, recommendations, or regulations for a test method, including batteries of tests and test screens, for chemicals or class of chemicals within a regulatory framework that may be appropriate for scientific improvement, while seeking to reduce, refine, or replace animal test methods.

(5) Consider for review and evaluation, petitions received from the public that—

(A) identify a specific regulation, recommendation, or guideline regarding a regulatory mandate; and

(B) recommend new or revised or alternative test methods and provide valid scientific evidence of the potential of the test method.

(6) Make available to the public final ICCVAM test recommendations to appropriate Federal agencies and the responses from the agencies regarding such recommendations.

(7) Prepare reports to be made available to the public on its progress under sections 285–2 to 285–5 of this title. The first report shall be completed not later than 12 months after December 19, 2000, and subsequent reports shall be completed biennially thereafter.


REFERENCES IN TEXT


CODIFICATION

Section was enacted as part of the ICCVAM Authorization Act of 2000, and not as part of the Public Health Service Act which comprises this chapter.

TERMINATION OF ADVISORY COMMITTEES

Advisory committees established after Jan. 5, 1973, to terminate not later than the expiration of the 2-year period beginning on the date of their establishment, unless, in the case of a committee established by the President or an officer of the Federal Government, such committee is renewed by appropriate action prior to the expiration of such 2-year period, or in the case of a committee established by the Congress, its duration is otherwise provided by law. See section 14 of Pub. L. 92–463, Oct. 6, 1972, 86 Stat. 776, set out in the Appendix to Title 5, Government Organization and Employees.

Pub. L. 93–641, § 6, Jan. 4, 1975, 88 Stat. 2275, set out as a note under section 217a of this title, provided that an advisory committee established pursuant to the Public Health Service Act shall terminate at such time as may be specifically prescribed by an Act of Congress enacted after Jan. 4, 1975.

§ 285–4. **Federal agency action**

(a) **Identification of tests**

With respect to each Federal agency carrying out a program that requires or recommends acute or chronic toxicological testing, such agency shall, not later than 180 days after receiving an ICCVAM test recommendation, identify and forward to the ICCVAM any relevant
§ 285l–5. Application

(a) Application

Sections 285l–2 to 285l–5 of this title shall not apply to research, including research performed using biotechnology techniques, or research related to the causes, diagnosis, treatment, control, or prevention of physical or mental diseases or impairments of humans or animals.

(b) Use of test methods

Nothing in sections 285l–2 to 285l–5 of this title shall prevent a Federal agency from retaining final authority for incorporating the test methods recommended by the ICCVAM in the manner determined to be appropriate by such Federal agency or regulatory body.

(c) Limitation

Nothing in sections 285l–2 to 285l–5 of this title shall be construed to require a manufacturer that is currently not required to perform animal testing to perform such tests. Nothing in sections 285l–2 to 285l–5 of this title shall be construed to require a manufacturer to perform redundant endpoint specific testing.

(d) Submission of tests and data

Nothing in sections 285l–2 to 285l–5 of this title precludes a party from submitting a test method or scientific data directly to a Federal agency for use in a regulatory program.


CODIFICATION

Section was enacted as part of the ICCVAM Authorization Act of 2000, and not as part of the Public Health Service Act which comprises this chapter.

§ 285l–6. Methods of controlling certain insect and vermin populations

The Director of the Institute shall conduct or support research to identify or develop methods of controlling insect and vermin populations that transmit to humans diseases that have significant adverse health consequences.


SUBPART 13—NATIONAL INSTITUTE ON DEAFNESS AND OTHER COMMUNICATION DISORDERS

§ 285m. Purpose of Institute

The general purpose of the National Institute on Deafness and Other Communication Disorders (hereafter referred to in this subpart as the “Institute”) is the conduct and support of research and training, the dissemination of health information, and other programs with respect to disorders of hearing and other communication processes, including diseases affecting hearing, balance, voice, speech, language, taste, and smell.


CODIFICATION

1988—Pub. L. 100–607 amended this section to read as if the amendments made by Pub. L. 100–607, which enacted this section, had not been enacted. See Codification note above.

SHORT TITLE OF 1988 AMENDMENT

For short title of Pub. L. 100–553 which enacted this subpart and amended sections 281 and 285j of this title as the “National Deafness and Other Communication Disorders Act of 1988”, see section 1 of Pub. L. 100–553, set out as a note under section 201 of this title.

EFFECT OF ENACTMENT OF SIMILAR PROVISIONS

Section 2613(b) of Pub. L. 100–607 provided that:

“(1) Paragraphs (2) and (3) shall take effect immediately after the enactment of both the bill, S. 1727, of the One Hundredth Congress, by subtitle A of title I of the Health Omnibus Programs Extension of 1988 [Pub. L. 100–607, approved Nov. 4, 1988].

“(2)(A) The provisions of the Public Health Service Act referred to in subparagraph (B), as similarly amended by the enactment of the bill, S. 1727, of the One Hundredth Congress, by subtitle A of title I of the Health Omnibus Programs Extension of 1988, and by subsection (a)(1) of this section, are amended to read as if the amendments made by such subtitle A and such subsection (a)(1) had not been enacted.

“(B) The provisions of the Public Health Service Act referred to in subparagraph (A) are—

“(A) sections 401(b)(1) and 457 [sections 281(b)(1) and 285j of this title];

“(B) part C of title IV [this part]; and

“(C) the heading for subpart 10 of such part C [42 U.S.C. prec. 285j].

“(3) Subsection (a)(2) of this section [set out below] is repealed.”

TRANSITIONAL AND SAVINGS PROVISIONS

Section 3 of Pub. L. 100–553 provided that:

“(a) TRANSFER OF PERSONNEL, ASSETS, AND LIABILITIES.—Personnel employed by the National Institutes of Health in connection with the functions vested under section 2 (enacting this subpart and amending sections 281 and 285j of this title) in the Director of the National Institute on Deafness and Other Communication Disorders, and assets, property, contracts, liabilities, records, unexpended balances of appropriations, authorizations, allocations, and other funds of the National Institutes of Health, arising from or employed, held, used, available to, or to be made available, in connection with such functions shall be transferred to the Director for appropriate allocation. Unexpended funds transferred under this subsection shall be used only for the purposes for which the funds were originally authorized and appropriated.

“(b) SAVINGS PROVISIONS.—With respect to functions vested under section 1 [probably means section 2, enacting this subpart and amending sections 281 and 285j of this title] in the Director of the National Institute on Deafness and Other Communication Disorders, all orders, rules, regulations, grants, contracts, certificates, licenses, privileges, and other determinations, actions, or official documents, that have been issued, made, granted, or allowed to become effective, and that are effective on the date of the enactment of this Act [Oct. 28, 1988], shall continue in effect according to their terms unless changed pursuant to law.”

Section 2613(a)(2) of Pub. L. 100–607, which enacted provisions that were substantially identical to the transitional and savings provisions above, was repealed by section 2613(b)(3) of Pub. L. 100–607.

§ 285m–1. National Deafness and Other Communication Disorders Program

(a) The Director of the Institute, with the advice of the Institute’s advisory council, shall establish a National Deafness and Other Communication Disorders Program (hereafter in this section referred to as the “Program”). The Director or the Institute shall, with respect to the Program, prepare and transmit to the Director of NIH a plan to initiate, expand, intensify and coordinate activities of the Institute respecting disorders of hearing (including tinnitus) and other communication processes, including diseases affecting hearing, balance, voice, speech, language, taste, and smell. The plan shall include such comments and recommendations as the Director of the Institute determines appropriate. The Director of the Institute shall periodically review and revise the plan and shall transmit any revisions of the plan to the Director of NIH.

(b) Activities under the Program shall include—

(1) investigation into the etiology, pathology, detection, treatment, and prevention of all forms of disorders of hearing and other communication processes, primarily through the support of basic research in such areas as anatomy, audiology, biochemistry, bioengineering, epidemiology, genetics, immunology, microbiology, molecular biology, the neurosciences, otolaryngology, psychology, pharmacology, physiology, speech and language pathology, and any other scientific disciplines that can contribute important knowledge to the understanding and elimination of disorders of hearing and other communication processes;

(2) research into the evaluation of techniques (including surgical, medical, and behavioral approaches) and devices (including hearing aids, implanted auditory and nonauditory prosthetic devices and other communication aids) used in diagnosis, treatment, rehabilitation, and prevention of disorders of hearing and other communication processes;

(3) research into prevention, and early detection and diagnosis, of hearing loss and speech and language disturbances (including stuttering) and research into preventing the effects of such disorders on learning and learning disabilities with extension of programs for appropriate referral and rehabilitation;

(4) research into the detection, treatment, and prevention of disorders of hearing and other communication processes in the growing elderly population with extension of rehabilitative programs to ensure continued effective communication skills in such population;

(5) research to expand knowledge of the effects of environmental agents that influence hearing or other communication processes; and

(6) developing and facilitating intramural programs on clinical and fundamental aspects of disorders of hearing and all other communication processes.


Footnote: 1 So in original. Probably should be “of”.
§ 285m–2. Data System and Information Clearinghouse

(a) The Director of the Institute shall establish a National Deafness and Other Communication Disorders Data System for the collection, storage, analysis, retrieval, and dissemination of data derived from patient populations with disorders of hearing or other communication processes, including where possible, data involving general populations for the purpose of identifying individuals at risk of developing such disorders.

(b) The Director of the Institute shall establish a National Deafness and Other Communication Disorders Information Clearinghouse to facilitate and enhance, through the effective dissemination of information, knowledge and understanding of disorders of hearing and other communication processes by health professionals, patients, industry, and the public.


§ 285m–3. Multipurpose deafness and other communication disorders center

(a) Development, modernization and operation; “modernization” defined

The Director of the Institute shall, after consultation with the advisory council for the Institute, provide for the development, modernization, and operation (including care required for research) of new and existing centers for studies of disorders of hearing and other communication processes. For purposes of this section, the term “modernization” means the alteration, remodel-
of children (particularly with respect to their education and training), affected by disorders of hearing or other communication processes.

(g) Period of support; recommended extensions of peer review group

Support of a center under this section may be for a period not to exceed seven years. Such period may be extended by the Director of the Institute for one or more additional periods of not more than five years if the operations of such center have been reviewed by an appropriate technical and scientific peer review group established by the Director, with the advice of the Institute’s advisory council, if such group has recommended to the Director that such period should be extended.


CODIFICATION


AMENDMENTS

1988—Pub. L. 100–690 amended this section to read as if the amendments made by Pub. L. 100–607, which enacted this section, had not been enacted. See Codification note above.

EFFECTIVE DATE OF 1988 AMENDMENT

For effective date of amendment by Pub. L. 100–690, see section 2613(b)(1) of Pub. L. 100–690, set out as an Effect of Enactment of Similar Provisions note under section 285m of this title.

§ 285m–4. National Institute on Deafness and Other Communication Disorders Advisory Board

(a) Establishment

The Secretary shall establish in the Institute the National Deafness and Other Communication Disorders Advisory Board (hereafter in this section referred to as the “Advisory Board”).

(b) Composition; qualifications; appointed and ex officio members

The Advisory Board shall be composed of eighteen appointed members and nonvoting ex officio members as follows:

1. The Secretary shall appoint—

   A. twelve members from individuals who are scientists, physicians, and other health and rehabilitation professionals, who are not officers or employees of the United States, and who represent the specialties and disciplines relevant to deafness and other communication disorders, including not less than two persons with a communication disorder; and

   B. six members from the general public who are knowledgeable with respect to such disorders, including not less than one person with a communication disorder and not less than one person who is a parent of an individual with such a disorder.

Of the appointed members, not less than five shall by virtue of training or experience be knowledgeable in diagnoses and rehabilitation of communication disorders, education of the hearing, speech, or language impaired, public health, public information, community program development, occupational hazards to communications senses, or the aging process.

(2) The following shall be ex officio members of each Advisory Board:

A. The Assistant Secretary for Health, the Director of NIH, the Director of the National Institute on Deafness and Other Communication Disorders, the Director of the Centers for Disease Control and Prevention, the Under Secretary for Health of the Department of Veterans Affairs, and the Assistant Secretary of Defense for Health Affairs (or the designees of such officers).

B. Such other officers and employees of the United States as the Secretary determines necessary for the Advisory Board to carry out its functions.

(c) Compensation

Members of an Advisory Board who are officers or employees of the Federal Government shall serve as members of the Advisory Board without compensation in addition to that received in their regular public employment. Other members of the Board shall receive compensation at rates not to exceed the daily equivalent of the annual rate in effect for grade GS–18 of the General Schedule for each day (including traveltime) they are engaged in the performance of their duties as members of the Board.

(d) Term of office; vacancies

The term of office of an appointed member of the Advisory Board is four years, except that no term of office may extend beyond the expiration of the Advisory Board. Any member appointed to fill a vacancy for an unexpired term shall be appointed for the remainder of such term. A member may serve after the expiration of the member’s term until a successor has taken office. If a vacancy occurs in the Advisory Board, the Secretary shall make an appointment to fill the vacancy not later than 90 days from the date the vacancy occurred.

(e) Chairman

The members of the Advisory Board shall select a chairman from among the appointed members.

(f) Personnel; executive director; professional and clerical staff members; consultants; information and administrative support services and facilities

The Secretary shall, after consultation with and consideration of the recommendations of the Advisory Board, provide the Advisory Board with an executive director and one other professional staff member. In addition, the Secretary shall, after consultation with and consideration of the recommendations of the Advisory Board, provide the Advisory Board with such additional professional staff members, such clerical staff members, such services of consultants, such information, and (through contracts or other arrangements) such administrative support services and facilities, as the Secretary determines are necessary for the Advisory Board to carry out its functions.
The Advisory Board shall meet at the call of the chairman or upon request of the Director of the Institute, but not less often than four times a year.

(h) Functions
The Advisory Board shall—

(1) review and evaluate the implementation of the plan prepared under section 285m–1(a) of this title and periodically update the plan to ensure its continuing relevance;

(2) for the purpose of assuring the most effective use and organization of resources respecting deafness and other communication disorders, advise and make recommendations to the Congress, the Secretary, the Director of NIH, the Director of the Institute, and the heads of other appropriate Federal agencies for the implementation and revision of such plan; and

(3) maintain liaison with other advisory bodies related to Federal agencies involved in the implementation of such plan and with key non-Federal entities involved in activities affecting the control of such disorders.

(i) Subcommittee activities; workshops and conferences; collection of data
In carrying out its functions, the Advisory Board may establish subcommittees, convene workshops and conferences, and collect data. Such subcommittees may be composed of Advisory Board members and nonmember consultants with expertise in the particular area addressed by such subcommittees. The subcommittees may hold such meetings as are necessary to enable them to carry out their activities.


(k) Commencement of existence
The National Deafness and Other Communication Disorders Advisory Board shall be established not later than April 1, 1989.


CODIFICATION

AMENDMENTS
2007—Subsec. (j). Pub. L. 109–482 struck out subsec. (j) which read as follows: “The Advisory Board shall prepare an annual report for the Secretary which—

“(1) describes the Advisory Board’s activities in the fiscal year for which the report is made;

“(2) describes and evaluates the progress made in such fiscal year in research, treatment, education, and training with respect to the deafness and other communication disorders;”

“(3) summarizes and analyzes expenditures made by the Federal Government for activities respecting such disorders in such fiscal year; and

“(4) contains the Advisory Board’s recommendations (if any) for changes in the plan prepared under section 285m–1(a) of this title.”


1989—Pub. L. 100–690, § 2613(b)(2), amended this section to read as if the amendments made by Pub. L. 100–690, § 2613(a)(1), which enacted this section, had not been enacted. See Codification note above.

EFFECTIVE DATE OF 2007 AMENDMENT
Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 198 of Pub. L. 109–482, set out as a note under section 281 of this title.

EFFECTIVE DATE OF 1988 AMENDMENT
For effective date of amendment by section 2613(b)(2) of Pub. L. 100–690, see section 2613(b)(1) of Pub. L. 100–690, set out as an Effect of Enactment of Similar Provisions note under section 265m of this title.

TERMINATION OF ADVISORY BOARDS
Advisory boards established after Jan. 5, 1973, to terminate not later than the expiration of the 2-year period beginning on the date of their establishment, unless, in the case of a board established by the President or an officer of the Federal Government, such board is renewed by appropriate action prior to the expiration of such 2-year period, or in the case of a board established by the Congress, its duration is otherwise provided by law. See sections 3(2) and 14 of Pub. L. 92–463, Oct. 6, 1972, 86 Stat. 770, 776, set out in the Appendix to Title 5, Government Organization and Employees.

Pub. L. 93–641, § 6, Jan. 4, 1975, 88 Stat. 2275, set out as a note under section 217a of this title, provided that an advisory committee established pursuant to the Public Health Service Act shall terminate at such time as may be specifically prescribed by an Act of Congress enacted after Jan. 4, 1975.

REFERENCES IN OTHER LAWS TO GS–16, 17, OR 18 PAY RATES
References in laws to the rates of pay for GS–16, 17, or 18 or to maximum rates of pay under the General Schedule, to be considered references to rates payable under specified sections of Title 5, Government Organization and Employees, see section 529 (title I, § 101(c)(1)) of Pub. L. 101–509, set out in a note under section 5376 of Title 5.

§ 285m–5. Interagency Coordinating Committee

(a) Establishment
The Secretary may establish a committee to be known as the Deafness and Other Communication Disorders Interagency Coordinating Committee (hereafter in this section referred to as the “Coordinating Committee”).

(b) Functions
The Coordinating Committee shall, with respect to deafness and other communication disorders—

(1) provide for the coordination of the activities of the national research institutes; and
(2) coordinate the aspects of all Federal health programs and activities relating to deafness and other communication disorders in order to assure the adequacy and technical soundness of such programs and activities and in order to provide for the full communication and exchange of information necessary to maintain adequate coordination of such programs and activities.

c) Composition

The Coordinating Committee shall be composed of the directors of each of the national research institutes and divisions involved in research with respect to deafness and other communication disorders and representatives of all other Federal departments and agencies whose programs involve health functions or responsibilities relevant to deafness and other communication disorders.

d) Chairman; meetings

The Coordinating Committee shall be chaired by the Director of NIH (or the designee of the Director). The Committee shall meet at the call of the chair, but not less often than four times a year.


Codification


Amendments

1993—Pub. L. 103–43 substituted “section 284c(a)(1)” for “section 284c(b)(1)”.

1988—Pub. L. 100–690, § 2613(b)(2), amended this section to read as if the amendments made by Pub. L. 100–690, § 2613(a)(1), which enacted this section, had not been enacted. See Codification note above.

Effective Date of 1988 Amendment

For effective date of amendment by section 2613(b)(2) of Pub. L. 100–690, see section 2613(b)(1) of Pub. L. 100–690, set out as an Effect of Enactment of Similar Provisions note under section 285m of this title.

§ 285m. Purpose of Institute

(a) In general

The general purpose of the National Institute on Alcohol Abuse and Alcoholism (hereafter in this subpart referred to as the “Institute”) is the conduct and support of biomedical and behavioral research, health services research, research training, and health information dissemination with respect to the prevention of alcohol abuse and the treatment of alcoholism.

(b) Research program

The research program established under this subpart shall encompass the social, behavioral, and biomedical etiology, mental and physical health consequences, and social and economic consequences of alcohol abuse and alcoholism. In carrying out the program, the Director of the Institute is authorized to—

1. collect and disseminate through publications and other appropriate means (including the development of curriculum materials), information as to, and the practical application of, the research and other activities under the program;
2. make available research facilities of the Public Health Service to appropriate public authorities, and to health officials and scientists engaged in special study;
3. make grants to universities, hospitals, laboratories, and other public or nonprofit institutions, and to individuals for such research projects as are recommended by the National Advisory Council on Alcohol Abuse and Alcoholism, giving special consideration to projects relating to—
   (A) the relationship between alcohol abuse and domestic violence,
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(B) the effects of alcohol use during pregnancy,
(C) the impact of alcoholism and alcohol abuse on the family, the workplace, and systems for the delivery of health services,
(D) the relationship between the abuse of alcohol and other drugs,
(E) the effect on the incidence of alcohol abuse and alcoholism of social pressures, legal requirements respecting the use of alcoholic beverages, the cost of such beverages, and the economic status and education of users of such beverages,
(F) the interrelationship between alcohol use and other health problems,
(G) the comparison of the cost and effectiveness of various treatment, methods for alcoholism and alcohol abuse and the effectiveness of prevention and intervention programs for alcoholism and alcohol abuse,
(H) alcoholism and alcohol abuse among women;

(4) secure from time to time and for such periods as he deems advisable, the assistance and advice of experts, scholars, and consultants from the United States or abroad;

(5) promote the coordination of research programs conducted by the Institute, and similar programs conducted by the National Institute of Drug Abuse and by other departments, agencies, organizations, and individuals, including all National Institutes of Health research activities which are or may be related to the problems of individuals suffering from alcoholism or alcohol abuse or those of their families or the impact of alcohol abuse on other health problems;

(6) conduct an intramural program of biomedical, behavioral, epidemiological, and social research, including research into the most effective means of treatment and service delivery, and including research involving human subjects, which is

(A) located in an institution capable of providing all necessary medical care for such human subjects, including complete 24-hour medical diagnostic services by or under the supervision of physicians, acute and intensive medical care, including 24-hour emergency care, psychiatric care, and such other care as is determined to be necessary for individuals suffering from alcoholism and alcohol abuse; and

(B) associated with an accredited medical or research training institution;

(7) for purposes of study, admit and treat at institutions, hospitals, and stations of the Public Health Service, persons not otherwise eligible for such treatment;

(8) provide to health officials, scientists, and appropriate public and other nonprofit institutions and organizations, technical advice and assistance on the application of statistical and other scientific research methods to experiments, studies, and surveys in health and medical fields;

(9) enter into contracts under this subchapter without regard to section 3324(a) and (b) of title 31 and section 6101 of title 41; and

(10) adopt, upon recommendation of the National Advisory Council on Alcohol Abuse and Alcoholism, such additional means as he deems necessary or appropriate to carry out the purposes of this section.

(c) Collaboration

The Director of the Institute shall collaborate with the Administrator of the Substance Abuse and Mental Health Services Administration in focusing the services research activities of the Institute and in disseminating the results of such research to health professionals and the general public.


CONDIFICATION


AMENDMENTS

2007—Subsec. (d). Pub. L. 109–482 struck out subsec. (d) which related to authorization of appropriations and allocation for health services research.


Subsec. (b). Pub. L. 102–321, §122(b)(1), (2)(A), transferred subsec. (b) of section 290bb of this title to subsec. (b) of this section, substituted “(b) RESEARCH PROGRAM.—The research program established under this subpart shall encompass the social, behavioral, and biomedical etiology, mental and physical health consequences, and social and economic consequences of alcohol abuse and alcoholism. In carrying out the program, the Director of the Institute is authorized” for “(b) In carrying out the program described in subsection (a) of this section, the Secretary, acting through the Institute, is authorized” in introductory provisions, and substituted a semicolon for period at end of par. (3)(H).

Subsecs. (c), (d). Pub. L. 102–321, §122(b)(2)(B), added subsecs. (c) and (d).

Effective Date of 2007 Amendment

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 and subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

Effective Date of 1992 Amendment

Section 3 of Pub. L. 102–352 provided that: “The amendments made by—

“(1) subsection (a) of section 2 (amending this section and sections 265n–2, 265o, 265o–2, 285p, 290aa–1, 290aa–3, 306x–7, 360z–27, 360z–31, 360z–53, and 309y of this title), shall take effect immediately upon the effectuation of the amendments made by titles I and II of the ADAMHA Reorganization Act [Pub. L. 102–321,
see Effective Date of 1992 Amendment note set out under section 236 of this title; and

"(2) subsections (b) and (c) of section 2 [amending sections 285cc–21, 285cc–28, and 285cc–30 of this title and provisions set out as notes under sections 285aa and 300x of this title], shall take effect on the date of enactment of this Act [Aug. 26, 1992]."

§ 285n–2. National Alcohol Research Centers; mandatory grant for research of effects of alcohol on elderly

(a) Designation; procedures applicable for approval of applications

The Secretary acting through the Institute may designate National Alcohol Research Centers for the purpose of interdisciplinary research relating to alcoholism and other biomedical, behavioral, and social issues related to alcoholism and alcohol abuse. No entity may be designated as a Center unless an application therefor has been submitted to, and approved by, the Secretary. Such an application shall be submitted in such manner and contain such information as the Secretary may reasonably require. The Secretary may not approve such an application unless—

(1) the application contains or is supported by reasonable assurances that—

(A) the applicant has the experience, or capability, to conduct, through biomedical, behavioral, social, and related disciplines, long-term research on alcoholism and other alcohol problems and to provide coordination of such research among such disciplines;

(B) the applicant has available to it sufficient facilities (including laboratory, reference, and data analysis facilities) to carry out the research plan contained in the application;

(C) the applicant has facilities and personnel to provide training in the prevention and treatment of alcoholism and other alcohol problems;

(D) the applicant has the capacity to train predoctoral and postdoctoral students for careers in research on alcoholism and other alcohol problems;

(E) the applicant has the capacity to conduct courses on alcohol problems and research on alcohol problems for undergraduate and graduate students, and for medical and osteopathic, nursing, social work, and other specialized graduate students; and

(F) the applicant has the capacity to conduct programs of continuing education in such medical, legal, and social service fields as the Secretary may require.1

1 See References in Text note below.

1 See in original. "The period probably should be " 20; and ".

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(2) the application contains a detailed five-year plan for research relating to alcoholism and other alcohol problems.

(b) Annual grants; amount; limitation on uses

The Secretary shall, under such conditions as the Secretary may reasonably require, make annual grants to Centers which have been designated under this section. No funds provided under a grant under this subsection may be used for the purchase of any land or the purchase, construction, preservation, or repair of any building. For the purposes of the preceding sentence, the term "construction" has the meaning given that term by section 292a(1).2 This title. The Secretary shall include in the grants made under this section for fiscal years beginning after September 30, 1981, a grant to a designated Center for research on the effects of alcohol on the elderly.


REFERENCES IN TEXT


 evacuated that subsec. (b) be amended by striking "or rental" before "of any land".

1986—Subsec. (b). Pub. L. 99–570, §4008(1), which directed that subsec. (b) be amended by striking "or rental" before "any land", could not be executed because "or rental" appeared before "of any land".

1983—Subsec. (a). Pub. L. 98–34, §2(b)(9)(B)(i), struck out direction that, insofar as practicable, the Secretary approve applications under this subsection in a manner resulting in an equitable geographic distribution of Centers.

2See References in Text note below.

Subsec. (b). Pub. L. 98–24, §2(b)(9)(B)(iii), struck out provision that no annual grant to any Center might exceed $1,500,000, and made a technical amendment to reference to section 292a of this title to reflect the transfer of this section to the Public Health Service Act.

Subsec. (c). Pub. L. 98–24, §2(b)(9)(B)(iv), struck out subsection (c) which authorized $6,000,000 for each of fiscal years ending Sept. 30, 1977, 1978, and 1979, $8,000,000 for fiscal year ending Sept. 30, 1980, and $9,000,000 for fiscal year ending Sept. 30, 1981.


1980—Subsec. (a). Pub. L. 96–180, §16(a), substituted: in first sentence "biomedical, behavioral, and social issues related to alcoholism and alcohol abuse" for "alcohol problems"; in par. (1)(B) "facilities (including laboratory, reference, and data analysis facilities) to carry out the research plan contained in the application" for "laboratory facilities and reference services (including reference services that will afford access to scientific alcohol literature)"; and in par. (1)(E) "medical and osteopathic, nursing, social work, and other specialized graduate students; and" for "medical and osteopathic students and physicians;" and added par. (3).

1979—Subsec. (a). Pub. L. 95–622 inserted provision following par. (2) relating to approval of applications under this subsection by the Secretary in a manner which results in equitable geographic distribution of Centers.

EFFECTIVE DATE OF 1992 AMENDMENTS


Amendment by Pub. L. 102–321 effective Oct. 1, 1992, with provision for programs providing financial assistance, see section 801(c), (d) of Pub. L. 102–321, set out as a note under section 236 of this title.

SUBPART 15—NATIONAL INSTITUTE ON DRUG ABUSE

§ 285o. Purpose of Institute

(a) In general

The general purpose of the National Institute on Drug Abuse (hereafter in this subpart referred to as the "Institute") is the conduct and support of biomedical and behavioral research, health services research, research training, and health information dissemination with respect to the prevention of drug abuse and the treatment of drug abusers.

(b) Research program

The research program established under this subpart shall encompass the social, behavioral, and biomedical etiology, mental and physical health consequences, and social and economic consequences of drug abuse. In carrying out the program, the Director of the Institute shall give special consideration to projects relating to drug abuse among women (particularly with respect to pregnant women).

(c) Collaboration

The Director of the Institute shall collaborate with the Substance Abuse and Mental Health Services Administration in focusing the services
research activities of the Institute and in disseminating the results of such research to health professionals and the general public.


AMENDMENTS

2007—Subsec. (d). Pub. L. 109–482 struck out subsec. (d) which related to authorization of appropriations and allocation for health services research.


EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

EFFECTIVE DATE OF 1992 AMENDMENT


EFFECTIVE DATE

Section effective Oct. 1, 1992, with provision for programs providing financial assistance, see section 801(c), (d) of Pub. L. 102–321, set out as an Effective Date of 1992 Amendment note under section 236 of this title.

§ 285o–1. Associate Director for Prevention

(a) In general

There shall be in the Institute an Associate Director for Prevention who shall be responsible for the full-time coordination and promotion of the programs in the Institute concerning the prevention of drug abuse. The Associate Director shall be appointed by the Director of the Institute from individuals who because of their professional training or expertise are experts in drug abuse and the prevention of such abuse.

(b) Report

The Associate Director for Prevention shall prepare for inclusion in the biennial report made under section 284b of this title a description of the prevention activities of the Institute, including a description of the staff and resources allocated to those activities.


REFERENCES IN TEXT


EFFECTIVE DATE

Section effective Oct. 1, 1992, with provision for programs providing financial assistance, see section 801(c), (d) of Pub. L. 102–321, set out as an Effective Date of 1992 Amendment note under section 236 of this title.

§ 285o–2. Drug Abuse Research Centers

(a) Authority

The Director of the Institute may designate National Drug Abuse Research Centers for the purpose of interdisciplinary research relating to drug abuse and other biomedical, behavioral, and social issues related to drug abuse. No entity may be designated as a Center unless an application therefore has been submitted to, and approved by, the Secretary. Such an application shall be submitted in such manner and contain such information as the Secretary may reasonably require. The Secretary may not approve such an application unless—

(1) the application contains or is supported by reasonable assurances that—

(A) the applicant has the experience, or capability, to conduct, through biomedical, behavioral, social, and related disciplines, long-term research on drug abuse and to provide coordination of such research among such disciplines;

(B) the applicant has available to it sufficient facilities (including laboratory, reference, and data analysis facilities) to carry out the research plan contained in the application;

(C) the applicant has facilities and personnel to provide training in the prevention and treatment of drug abuse;

(D) the applicant has the capacity to train predoctoral and postdoctoral students for careers in research on drug abuse;

(E) the applicant has the capacity to conduct courses on drug abuse problems and research on drug abuse for undergraduate and graduate students, and medical and osteopathic, nursing, social work, and other specialized graduate students; and

(F) the applicant has the capacity to conduct programs of continuing education in such medical, legal, and social service fields as the Secretary may require.\(^1\)

(2) the application contains a detailed five-year plan for research relating to drug abuse.

(b) Grants

The Director of the Institute shall, under such conditions as the Secretary may reasonably require, make annual grants to Centers which have been designated under this section. No funds provided under a grant under this subsection may be used for the purchase of any land or the purchase, construction, preservation, or repair of any building. For the purposes of the preceding sentence, the term “construction” has the meaning given that term by section 292a(1)\(^2\) of this title.

(c) Drug abuse and addiction research

(1) Grants or cooperative agreements

The Director of the Institute may make grants or enter into cooperative agreements to expand the current and ongoing interdisciplinary research and clinical trials with treatment centers of the National Drug Abuse Treatment Clinical Trials Network relating to drug abuse and addiction, including related biomedical, behavioral, and social issues.

(2) Use of funds

Amounts made available under a grant or cooperative agreement under paragraph (1) for

\(^1\) See References in Text note below.

\(^2\) See References in Text note below.
drug abuse and addiction may be used for research and clinical trials relating to—
   (A) the effects of drug abuse on the human body, including the brain;
   (B) the addictive nature of drugs and how such effects differ with respect to different individuals;
   (C) the connection between drug abuse and mental health;
   (D) the identification and evaluation of the most effective methods of prevention of drug abuse and addiction;
   (E) the identification and development of the most effective methods of treatment of drug addiction, including pharmacological treatments;
   (F) risk factors for drug abuse;
   (G) effects of drug abuse and addiction on pregnant women and their fetuses; and
   (H) cultural, social, behavioral, neurological, and psychological reasons that individuals abuse drugs, or refrain from abusing drugs.

(3) Research results

The Director shall promptly disseminate research results under this subsection to Federal, State, and local entities involved in combating drug abuse and addiction.


REFERENCES IN TEXT

Section 292a of this title, referred to in subsec. (b), was in the original a reference to section 701 of this title July 1, 1944. Section 701 of that Act was omitted in the general revision of subchapter V of this chapter by Pub. L. 102–408, title I, §102, Oct. 13, 1992, 106 Stat. 1994. Pub. L. 102–408 enacted a new section 701 of act July 1, 1944, relating to statement of purpose, and a new section 702, relating to scope and duration of loan insurance program, which are classified to sections 292 and 292a, respectively, of this title. For provisions relating to definitions, see sections 292d and 295p of this title.

AMENDMENTS

2007—Subsec. (c)(4). Pub. L. 109–482 struck out par. (4) which authorized appropriations and provided they were supplemental to other funding of research on drug abuse.

2002—Subsec. (c). Pub. L. 107–273 amended heading and text of subsec. (c) generally, substituting provisions relating to grants and cooperative agreements for research and clinical trials relating to drug abuse and addiction for similar provisions relating to grants or cooperative agreements for research and clinical trials relating to methamphetamine abuse and addiction.


1992—Subsec. (b). Pub. L. 102–352 substituted "292a(1)" for "292a(1)".

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 285n of this title.

EFFECTIVE DATE OF 1992 AMENDMENT


EFFECTIVE DATE

Section effective Oct. 1, 1992, with provision for programs providing financial assistance, see section 801(c).

§ 285o–3. Office on AIDS

The Director of the Institute shall establish within the Institute an Office on AIDS. The Office shall be responsible for the coordination of research and determining the direction of the Institute with respect to AIDS research related to—

(1) primary prevention of the spread of HIV, including transmission via drug abuse;
(2) drug abuse services research; and
(3) other matters determined appropriate by the Director.


EFFECTIVE DATE

Section effective Oct. 1, 1992, with provision for programs providing financial assistance, see section 801(c).

§ 285o–4. Medication Development Program

(a) Establishment

There is established in the Institute a Medication Development Program through which the Director of such Institute shall—

(1) conduct periodic meetings with the Commissioner of Food and Drugs to discuss measures that may facilitate the approval process of drug abuse treatments;
(2) encourage and promote (through grants, contracts, international collaboration, or otherwise) expanded research programs, investigations, experiments, community trials, and studies, into the development and use of medications to treat drug addiction;
(3) establish or provide for the establishment of research facilities;
(4) report on the activities of other relevant agencies relating to the development and use of pharmacotherapeutic treatments for drug addiction;
(5) collect, analyze, and disseminate data useful in the development and use of pharmacotherapeutic treatments for drug addiction and collect, catalog, analyze, and disseminate through international channels, the results of such research;
(6) directly or through grants, contracts, or cooperative agreements, support training in the fundamental sciences and clinical disciplines related to the pharmacotherapeutic treatment of drug abuse, including the use of training stipends, fellowships, and awards where appropriate; and

(7) coordinate the activities conducted under this section with related activities conducted within the National Institute on Alcohol Abuse and Alcoholism, the National Institute of Mental Health, and other appropriate institutes and shall consult with the Directors of such Institutes.

(b) Duties

In carrying out the activities described in subsection (a) of this section, the Director of the Institute—

(1) shall collect and disseminate through publications and other appropriate means, information pertaining to the research and other activities under this section;

(2) shall make grants to or enter into contracts and cooperative agreements with individuals and public and private entities to further the goals of the program;

(3) may, in accordance with section 289e of this title, and in consultation with the National Advisory Council on Drug Abuse, acquire, construct, improve, repair, operate, and maintain pharmacotherapeutic research centers, laboratories, and other necessary facilities and equipment, and such other real or personal property as the Director determines necessary, and may, in consultation with such Advisory Council, make grants for the construction or renovation of facilities to carry out the purposes of this section;

(4) may accept voluntary and uncompensated services;

(5) may accept gifts, or donations of services, money, or property, real, personal, or mixed, tangible or intangible; and

(6) shall take necessary action to ensure that all channels for the dissemination and exchange of scientific knowledge and information are maintained between the Institute and the other scientific, medical, and biomedical disciplines and organizations nationally and internationally.

(c) Report

(1) In general

Not later than December 31, 1992, and each December 31 thereafter, the Director of the Institute shall submit to the Office of National Drug Control Policy established under section 1501 of title 21 a report, in accordance with paragraph (3), that describes the objectives and activities of the program assisted under this section.

(2) National Drug Control Strategy

The Director of National Drug Control Policy shall incorporate, by reference or otherwise, each report submitted under this subsection in the National Drug Control Strategy submitted the following February 1 under section 1504 of title 21.

(d) "Pharmacotherapeutics" defined

For purposes of this section, the term "pharmacotherapeutics" means medications used to treat the symptoms and disease of drug abuse, including medications to—

(1) block the effects of abused drugs;

(2) reduce the craving for abused drugs;

(3) moderate or eliminate withdrawal symptoms;

(4) block or reverse the toxic effect of abused drugs; or

(5) prevent relapse in persons who have been detoxified from drugs of abuse.


REFERENCES IN TEXT

Sections 1501 and 1504 of title 21, referred to in subsec. (c), were repealed by Pub. L. 100–690, title I, §1009, Nov. 18, 1988, 102 Stat. 4188, as amended.

AMENDMENTS

2007—Subsec. (e). Pub. L. 109–482 struck out heading and text of subsec. (e). Text read as follows: "For the purpose of carrying out this section, there are authorized to be appropriated $85,000,000 for fiscal year 1993, and $95,000,000 for fiscal year 1994." 1993—Subsec. (b)(6). Pub. L. 103–43 substituted "Institute" for "Administration".

EFFECTIVE DATE OF 2007 AMENDMENT

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

EFFECTIVE DATE

Section effective Oct. 1, 1992, with provision for programs providing financial assistance, see section 801(c), (d) of Pub. L. 102–321, set out as an Effective Date of 1992 Amendment note under section 236 of this title.

REPORT BY INSTITUTE ON MEDICINE

Section 701 of Pub. L. 102–321 directed Secretary of Health and Human Services to enter into a contract with a public or nonprofit private entity to conduct a study concerning (1) role of the private sector in development of anti-addiction medications, including legislative proposals designed to encourage private sector development of such medications, (2) process by which anti-addiction medications receive marketing approval from Food and Drug Administration, including an assessment of feasibility of expediting marketing approval process in a manner consistent with maintaining safety and effectiveness of such medications, (3) with respect to pharmacotherapeutic treatments for drug addiction (A) recommendations with respect to a national strategy for developing such treatments and improvements in such strategy, (B) state of the scientific knowledge concerning such treatments, and (C) assessment of progress toward development of safe, effective pharmacological treatments for drug addiction, and (4) other related information determined appropriate by the authors of the study, and to submit to Congress a report of the results of such study not later than 18 months after July 10, 1992.

1 See References in Text note below.
§ 285p. Purpose of Institute

(a) In general

The general purpose of the National Institute of Mental Health (hereafter in this subpart referred to as the “Institute”) is the conduct and support of biomedical and behavioral research, health services research, research training, and health information dissemination with respect to the causes, diagnosis, treatment, control, and prevention of mental illness.

(b) Research program

The research program established under this subpart shall include support for biomedical and behavioral neuroscience and shall be designed to further the treatment and prevention of mental illness, the promotion of mental health, and the study of the psychological, social and legal factors that influence behavior.

(c) Collaboration

The Director of the Institute shall collaborate with the Administrator of the Substance Abuse and Mental Health Services Administration in focusing the services research activities of the Institute and in disseminating the results of such research to health professionals and the general public.

(d) Information with respect to suicide

(1) In general

The Director of the Institute shall—

(A) develop and publish information with respect to the causes of suicide and the means of preventing suicide; and

(B) make such information generally available to the public and to health professionals.

(2) Youth suicide

Information described in paragraph (1) shall especially relate to suicide among individuals under 24 years of age.

(e) Associate Director for Special Populations

(1) In general

The Director of the Institute shall designate an Associate Director for Special Populations.

(2) Duties

The Associate Director for Special Populations shall—

(A) develop and coordinate research policies and programs to assure increased emphasis on the mental health needs of women and minority populations;

(B) support programs of basic and applied social and behavioral research on the mental health problems of women and minority populations;

(C) study the effects of discrimination on institutions and individuals, including majority institutions and individuals;

(D) support and develop research designed to eliminate institutional discrimination; and

(E) provide increased emphasis on the concerns of women and minority populations in training programs, service delivery programs, and research endeavors of the Institute.


AMENDMENTS

2007—Subsec. (f). Pub. L. 109–482 struck out subsec. (f) which authorized appropriations and provided that at least 15% of the appropriated amounts were to carry out health services research relating to mental health.


Effective Date of 2007 Amendment

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

Effective Date of 1992 Amendment


§ 285p–1. Associate Director for Prevention

(a) In general

There shall be in the Institute an Associate Director for Prevention who shall be responsible for the full-time coordination and promotion of the programs in the Institute concerning the prevention of mental disorder. The Associate Director shall be appointed by the Director of the Institute from individuals who because of their professional training or expertise are experts in mental disorder and the prevention of such.

(b) Report

The Associate Director for Prevention shall prepare for inclusion in the biennial report made under section 284b 1 of this title a description of the prevention activities of the Institute, including a description of the staff and resources allocated to those activities.


REFERENCES IN TEXT


1 See References in Text note below.
§ 285p–2. Office of Rural Mental Health Research

(a) In general

There is established within the Institute an office to be known as the Office of Rural Mental Health Research (hereafter in this section referred to as the “Office”). The Office shall be headed by a director, who shall be appointed by the Director of such Institute from among individuals experienced or knowledgeable in the provision of mental health services in rural areas. The Secretary shall carry out the authorities established in this section acting through the Director of the Office.

(b) Coordination of activities

The Director of the Office, in consultation with the Director of the Institute and with the Director of the Office of Rural Health Policy, shall—

(1) coordinate the research activities of the Department of Health and Human Services as such activities relate to the mental health of residents of rural areas; and

(2) coordinate the activities of the Office with similar activities of public and nonprofit private entities.

(c) Research, demonstrations, evaluations, and dissemination

The Director of the Office may, with respect to the mental health of adults and children residing in rural areas—

(1) conduct research on conditions that are unique to the residents of rural areas, or more serious or prevalent in such residents; and

(2) conduct research on improving the delivery of services in such areas; and

(3) disseminate information to appropriate public and nonprofit private entities.

(d) Authority regarding grants and contracts

The Director of the Office may carry out the authorities established in subsection (c) of this section directly and through grants, cooperative agreements, or contracts with public or nonprofit private entities.

Effective Date: October 1, 1992.

§ 285q. Purpose of Institute

The general purpose of the National Institute of Nursing Research (in this subpart referred to as the “Institute”) is the conduct and support of, and dissemination of information respecting, basic and clinical nursing research, training, and other programs in patient care research.

Effective Date: October 1, 1992.
§ 285q–1. Specific authorities

To carry out section 285q of this title, the Director of the Institute may provide research training and instruction and establish, in the Institute and other nonprofit institutions, research traineeships and fellowships in the study and investigation of the prevention of disease, health promotion, and the nursing care of individuals with and the families of individuals with acute and chronic illnesses. The Director may make grants to nonprofit institutions to provide such training and instruction and traineeships and fellowships.


CODIFICATION

Section was formerly classified to section 287c-1 of this title prior to renumbering by Pub. L. 103–43.

AMENDMENTS


§ 285q–2. Advisory council

(a) Appointment; functions and duties; acceptance of conditional gifts; subcommittees

(1) The Secretary shall appoint an advisory council for the Institute which shall advise, assist, consult with, and make recommendations to the Secretary and the Director of the Institute on matters related to the activities carried out by and through the Institute and the policies respecting such activities.

(2) The advisory council for the Institute may recommend to the Secretary acceptance, in accordance with section 228 of this title, of conditional gifts for study, investigations, and research and for the acquisition of grounds or construction, equipping, or maintenance of facilities for the Institute.

(3) The advisory council for the Institute—

(A)(i) may make recommendations to the Director of the Institute respecting research conducted in the Institute,

(ii) may review applications for grants and cooperative agreements for research or training and recommend for approval applications for projects which show promise of making valuable contributions to human knowledge, and

(iii) may review any grant, contract, or cooperative agreement proposed to be made or entered into by the Institute;

(B) may collect, by correspondence or by personal investigation, information as to studies which are being carried on in the United States or any other country as to the diseases, disorders, or other aspects of human health with respect to which the Institute is concerned and with the approval of the Director of the Institute make available such information through appropriate publications for the benefit of public and private health entities and health professions personnel and scientists and for the information of the general public; and

(C) may appoint subcommittees and convene workshops and conferences.

(b) Membership; ex officio members; compensation

(1) The advisory council shall consist of ex officio members and not more than eighteen members appointed by the Secretary.

(2) The ex officio members of the advisory council shall consist of—

(A) the Secretary, the Director of NIH, the Director of the Institute, the chief nursing officer of the Department of Veterans Affairs, the Assistant Secretary of Defense for Health Affairs, the Director of the Division of Nursing of the Health Resources and Services Administration (or the designees of such officers), and

(B) such additional officers or employees of the United States as the Secretary determines necessary for the advisory council to effectively carry out its functions.

(3) The members of the advisory council who are not ex officio members shall be appointed as follows:

(A) Two-thirds of the members shall be appointed by the Secretary from among the leading representatives of the health and scientific disciplines (including public health and the behavioral or social sciences) relevant to the activities of the Institute. Of the members appointed pursuant to this subparagraph, at least seven shall be professional nurses who are recognized experts in the area of clinical practice, education, or research.

(B) One-third of the members shall be appointed by the Secretary from the general public and shall include leaders in fields of public policy, law, health policy, economics, and management.

(4) Members of the advisory council who are officers or employees of the United States shall not receive any compensation for service on the advisory council. The other members of the advisory council shall receive, for each day (including traveltime) they are engaged in the performance of the functions of the advisory council, compensation at rates not to exceed the daily equivalent of the annual rate in effect for grade GS-18 of the General Schedule.

(c) Term of office; vacancy; reappointment

The term of office of an appointed member of the advisory council is four years, except that any member appointed to fill a vacancy for an unexpired term shall be appointed for the remainder of such term and the Secretary shall make appointments to an advisory council in such a manner as to ensure that the terms of the members do not all expire in the same year. A member may serve after the expiration of the member’s term until a successor has taken office. A member who has been appointed for a term of four years may not be reappointed to an advisory council before two years from the date
of expiration of such term of office. If a vacancy occurs in the advisory council among the appointed members, the Secretary shall make an appointment to fill the vacancy within 90 days from the date the vacancy occurs.

(d) Chairman; selection; term of office

The chairman of the advisory council shall be selected by the Secretary from among the appointed members, except that the Secretary may select the Director of the Institute to be the chairman of the advisory council. The term of office of the chairman shall be two years.

(e) Meetings

The advisory council shall meet at the call of the chairman or upon the request of the Director of the Institute, but at least three times each fiscal year. The location of the meetings of the advisory council is subject to the approval of the Director of the Institute.

(f) Executive secretary; staff; orientation and training for new members

The Director of the Institute shall designate a member of the staff of the Institute to serve as the executive secretary of the advisory council. The Director of the Institute shall make available to the advisory council such staff, information, and other assistance as it may require to carry out its functions. The Director of the Institute shall provide orientation and training for new members of the advisory council to provide them with such information and training as may be appropriate for their effective participation in the functions of the advisory council.

(g) Material for inclusion in biennial report; additional reports

The advisory council may prepare, for inclusion in the biennial report made under section 285q–3 of this title, (1) comments respecting the activities of the advisory council in the fiscal years respecting which the report is prepared, (2) comments on the progress of the Institute in meeting its objectives, and (3) recommendations respecting the future directions and program policies of the Institute. The advisory council may prepare such additional reports as it may determine appropriate.


1990—Subsec. (a)(2). Pub. L. 101–381 made technical amendment to reference to section 300aa of this title to reflect renumbering of corresponding section of original act.

TERMINATION OF ADVISORY COUNCILS

Advisory councils established after Jan. 5, 1973, to terminate not later than the expiration of the 2-year period beginning on the date of their establishment, unless, in the case of a council established by the President or an officer of the Federal Government, such council is renewed by appropriate action prior to the expiration of such 2-year period, or in the case of a council established by the Congress, its duration is otherwise provided by law. See sections 3(2) and 14 of Pub. L. 92–463, Oct. 6, 1972, 86 Stat. 770, 776, set out in the Appendix to Title 5, Government Organization and Employees.

Pub. L. 93–641, §6, Jan. 4, 1975, 88 Stat. 2275, set out as a note under section 217a of this title, provided that an advisory committee established pursuant to the Public Health Service Act shall terminate at such time as may be specifically prescribed by an Act of Congress enacted after Jan. 4, 1975.

REFERENCES IN OTHER LAWS TO GS–16, 17, OR 18 PAY RATES

References in laws to the rates of pay for GS–16, 17, or 18, or to maximum rates of pay under the General Schedule, to be considered references to rates payable under specified sections of Title 5, Government Organization and Employees, see section 529 (title I, §101(c)(1)) of Pub. L. 101–509, set out in a note under section 5376 of Title 5.


The Director of the Institute after consultation with the advisory council for the Institute, shall prepare for inclusion in the biennial report made under section 285q of this title a biennial report which shall consist of a description of the activities of the Institute and program policies of the Director of the Institute in the fiscal years respecting which the report is prepared. The Director of the Institute may prepare such additional reports as the Director determines appropriate. The Director of the Institute shall provide the advisory council of the Institute an opportunity for the submission of the written comments referred to in section 285q–2(g) of this title.

(July 1, 1944, ch. 373, title IV, §464Y, formerly §486, as added Pub. L. 99-158, §2, Nov. 20, 1985, 99 Stat. 869; renumbered §485A, renumbered §464Y, §2008(b)(13), which directed the substitution of “Department of Veterans Affairs” for “Veterans’ Administration” in section 287c–2(b)(2)(A) of this title could not be transmitted because the words “Veterans’ Administration” do not appear in subsec. (b)(2)(A) of this section subsequent to amendment by Pub. L. 102–54 and because of the renumbering of this section. See Codification note above and 1991 Amendment note below.


Subsecs. (d) to (f), Pub. L. 103–43, §1511(a)(3)(C), substituted “Institute” for “Center” wherever appearing. Subsec. (g), Pub. L. 103–43, §1511(a)(3)(C), (b)(4)(B), substituted “section 285q–3” for “section 287c–3” and “Institute” for “Center” in two places.

CODEFICATION
Section was formerly classified to section 287c–3 of this title prior to renumbering by Pub. L. 103–43.

AMENDMENTS
1993—Pub. L. 103–43, §1511(a)(4), (b)(4)(C), substituted “Institute” for “Center” wherever appearing and “section 287c–2(g)” for “section 287c–2(g)”.

SUBPART 18—NATIONAL INSTITUTE OF BIOMEDICAL IMAGING AND BIOENGINEERING

§ 285r. Purpose of the Institute

(a) In general
The general purpose of the National Institute of Biomedical Imaging and Bioengineering (in this section referred to as the “Institute”) is the conduct and support of research, training, the dissemination of health information, and other programs with respect to biomedical imaging, biomedical engineering, and associated technologies and modalities with biomedical applications (in this section referred to as “biomedical imaging and bioengineering”).

(b) National Biomedical Imaging and Bioengineering Program
(1) The Director of the Institute, with the advice of the Institute’s advisory council, shall establish a National Biomedical Imaging and Bioengineering Program (in this section referred to as the “Program”).

(2) Activities under the Program shall include the following with respect to biomedical imaging and bioengineering:

(A) Research into the development of new techniques and devices.

(B) Related research in physics, engineering, mathematics, computer science, and other disciplines.

(C) Technology assessments and outcomes studies to evaluate the effectiveness of biologics, materials, processes, devices, procedures, and informatics.

(D) Research in screening for diseases and disorders.

(E) The advancement of existing imaging and bioengineering modalities, including imaging, biomaterials, and informatics.

(F) The development of target-specific agents to enhance images and to identify and delineate disease.

(G) The development of advanced engineering and imaging technologies and techniques for research from the molecular and genetic to the whole organ and body levels.

(H) The development of new techniques and devices for more effective interventional procedures (such as image-guided interventions).

(3) (A) With respect to the Program, the Director of the Institute shall prepare and transmit to the Secretary and the Director of NIH a plan to initiate, expand, intensify, and coordinate activities of the Institute with respect to biomedical imaging and bioengineering. The plan shall include such comments and recommendations as the Director of the Institute determines appropriate. The Director of the Institute shall periodically review and revise the plan and shall transmit any revisions of the plan to the Secretary and the Director of NIH.

(B) The plan under subparagraph (A) shall include the recommendations of the Director of the Institute with respect to the following:

(i) Where appropriate, the consolidation of programs of the National Institutes of Health for the express purpose of enhancing support of activities regarding basic biomedical imaging and bioengineering research.

(ii) The coordination of the activities of the Institute with related activities of the other agencies of the National Institutes of Health and with related activities of other Federal agencies.

(c) Membership
The establishment under section 284a of this title of an advisory council for the Institute is subject to the following:

(1) The number of members appointed by the Secretary shall be 12.

(2) Of such members—

(A) six members shall be scientists, engineers, physicians, and other health professionals who represent disciplines in biomedical imaging and bioengineering and who are not officers or employees of the United States; and

(B) six members shall be scientists, engineers, physicians, and other health professionals who represent other disciplines and are knowledgeable about the applications of biomedical imaging and bioengineering in medicine, and who are not officers or employees of the United States.

(3) In addition to the ex officio members specified in section 284a(b)(2) of this title, the ex officio members of the advisory council shall include the Director of the Centers for Disease Control and Prevention, the Director of the National Science Foundation, and the Director of the National Institute of Standards and Technology (or the designee of such officers).

(4) The Secretary shall be 12.

(5) The number of members appointed by the Secretary shall be 12.

(6) The number of members appointed by the Secretary shall be 12.

(7) The number of members appointed by the Secretary shall be 12.”


AMENDMENTS

EFFECTIVE DATE OF 2007 AMENDMENT
Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

EFFECTIVE DATE
Pub. L. 106–580, §4, Dec. 29, 2000, 114 Stat. 3092, provided that: “This Act [enacting this subpart, amending section 281 of this title, and enacting provisions set out as notes under this section and section 281 of this title] takes effect October 1, 2000, or upon the date of the enactment of this Act [Dec. 29, 2000], whichever occurs later.”
FINDINGS
Pub. L. 106–580, § 2, Dec. 29, 2000, 114 Stat. 3088, provided that: "The Congress makes the following findings:

"(1) Basic research in imaging, bioengineering, computer science, informatics, and related fields is critical to improving health care but is fundamentally different from the research in molecular biology on which the current national research institutes at the National Institutes of Health (\n\n\n\nNIH\n\n\n\n) are based. To ensure the development of new techniques and technologies for the 21st century, these disciplines therefore require an identity and research home at the NIH that is independent of the existing institute structure.

"(2) Advances based on medical research promise new, more effective treatments for a wide variety of diseases, but the development of new, noninvasive imaging techniques for earlier detection and diagnosis of disease is essential to take full advantage of such new treatments and to promote the general improvement of health care.

"(3) The development of advanced genetic and molecular imaging techniques is necessary to continue the current rapid pace of discovery in molecular biology.

"(4) Advances in telemedicine, and teleradiology in particular, are increasingly important in the delivery of high-quality, reliable medical care to rural citizens and other underserved populations. To fulfill the promise of telemedicine and related technologies fully, a structure is needed at the NIH to support basic research focused on the acquisition, transmission, processing, and optimal display of images.

"(5) A number of Federal departments and agencies support imaging and engineering research with potential medical applications, but a central coordinating body, preferably housed at the NIH, is needed to coordinate these disparate efforts and facilitate the transfer of technologies with medical applications.

"(6) Several breakthrough imaging technologies, including magnetic resonance imaging ('MRI') and computed tomography ('CT'), have been developed primarily abroad, in large part because of the absence of a home at the NIH for basic research in imaging and related fields. The establishment of a central focus for imaging and bioengineering research at the NIH would promote both scientific advance and United States economic development.

"(7) At a time when a consensus exists to add significant resources to the NIH in coming years, it is appropriate to modernize the structure of the NIH to ensure that research dollars are expended more effectively and efficiently and that the fields of medical science that have contributed the most to the detection, diagnosis, and treatment of disease in recent years receive appropriate emphasis.

"(8) The establishment of a National Institute of Biomedical Imaging and Bioengineering at the NIH would accelerate the development of new technologies with clinical and research applications, improve coordination and efficiency at the NIH and throughout the Federal Government, reduce duplication and waste, lay the foundation for a new medical information age, promote economic development, and provide a structure to train the young researchers who will make the pathbreaking discoveries of the next century."

ESTABLISHMENT OF INSTITUTE AND ADVISORY COUNCIL
Pub. L. 106–580, § 3(b)–(d), Dec. 29, 2000, 114 Stat. 3091, provided that:

"(b) USE OF EXISTING RESOURCES.—In providing for the establishment of the National Institute of Biomedical Imaging and Bioengineering pursuant to the amendments made by subsection (a) [enacting this subpart], the Director of the National Institutes of Health (referred to in this subsection as 'NIH')—

"(1) may transfer to the National Institute of Biomedical Imaging and Bioengineering such personnel of NIH as the Director determines to be appropriate;

"(2) may, for quarters for such Institute, utilize such facilities of NIH as the Director determines to be appropriate; and

"(3) may obtain administrative support for the Institute from the other agencies of NIH, including the other national research institutes.

"(c) CONSTRUCTION OF FACILITIES.—None of the provisions of this Act [enacting this subpart, amending section 281 of this title, and enacting provisions set out as notes under this section and section 201 of this title] or the amendments made by the Act may be construed as authorizing the construction of facilities, or the acquisition of land, for purposes of the establishment or operation of the National Institute of Biomedical Imaging and Bioengineering.

"(d) DATE CERTAIN FOR ESTABLISHMENT OF ADVISORY COUNCIL.—Not later than 90 days after the effective date of this Act [Dec. 29, 2000] under section 4 [set out above], the Secretary of Health and Human Services shall complete the establishment of an advisory council for the National Institute of Biomedical Imaging and Bioengineering in accordance with section 406 of the Public Health Service Act [section 284a of this title] and in accordance with section 454h of such Act (as added by subsection (a) of this section) [this section]."

SUBPART 19—NATIONAL HUMAN GENOME RESEARCH INSTITUTE

AMENDMENTS

§ 285s. Purpose of Institute

(a) General purpose

The general purpose of the National Human Genome Research Institute (in this subpart referred to as the ‘‘Institute’’) is to characterize the structure and function of the human genome, including the mapping and sequencing of individual genes. Such purpose includes—

(1) planning and coordinating the research goal of the genome project;

(2) reviewing and funding research proposals;

(3) developing training programs;

(4) coordinating international genome research;

(5) communicating advances in genome science to the public; and

(6) reviewing and funding proposals to address the ethical and legal issues associated with the genome project (including legal issues regarding patents).

(b) Research training

The Director of the Institute may conduct and support research training—

(1) for which fellowship support is not provided under section 288 of this title; and

(2) that is not residency training of physicians or other health professionals.

(c) Amount available for ethical and legal issues

(1) Except as provided in paragraph (2), of the amounts appropriated to carry out subsection (a) of this section for a fiscal year, the Director of the Institute shall make available not less than 5 percent for carrying out paragraph (6) of such subsection.

(2) With respect to providing funds under subsection (a)(6) of this section for proposals to ad-
address the ethical issues associated with the genome project, paragraph (1) shall not apply for a fiscal year if the Director of the Institute certifies to the Committee on Energy and Commerce of the House of Representatives, and to the Committee on Labor and Human Resources of the Senate, that the Director has determined that an insufficient number of such proposals meet the applicable requirements of sections 289 and 289a of this title.


CODIFICATION
Section was formerly classified to section 287c of this title prior to renumbering by Pub. L. 109–482.

AMENDMENTS
2007—Pub. L. 109–482, § 101(c)(4)(C), substituted “Institute” for “Center” wherever appearing in section catchline and text.

CHANGE OF NAME
Committee on Labor and Human Resources of Senate changed to Committee on Health, Education, Labor, and Pensions of Senate by Senate Resolution No. 20, One Hundred Sixth Congress, Jan. 19, 1999.
Committee on Energy and Commerce of House of Representatives treated as referring to Committee on Commerce of House of Representatives by section 1(a) of Pub. L. 104–14, set out as a note under section 21 of Title 2, The Congress. Committee on Commerce of House of Representatives changed to Committee on Energy and Commerce of House of Representatives, and jurisdiction over matters relating to securities and exchanges and insurance generally transferred to Committee on Financial Services of House of Representatives by House Resolution No. 5, One Hundred Seventh Congress, Jan. 3, 2001.

EFFECTIVE DATE OF 2007 AMENDMENT
Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

SUBPART 20—NATIONAL INSTITUTE ON MINORITY HEALTH AND HEALTH DISPARITIES

AMENDMENTS

§ 285t. Purpose of Institute

(a) In general
The general purpose of the National Institute on Minority Health and Health Disparities (in this subpart referred to as the ‘‘Institute’’) is the conduct and support of research, training, dissemination of information, and other programs with respect to minority health conditions and other populations with health disparities.

(b) Priorities
The Director of the Institute shall in expending amounts appropriated under this subpart give priority to conducting and supporting minority health disparities research.

(c) Minority health disparities research
For purposes of this subpart:
(1) The term “minority health disparities research” means basic, clinical, and behavioral research on minority health conditions (as defined in paragraph (2)), including research to prevent, diagnose, and treat such conditions.
(2) The term “minority health conditions”, with respect to individuals who are members of minority groups, means all diseases, disorders, and conditions (including with respect to mental health and substance abuse)—
(A) unique to, more serious, or more prevalent in such individuals;
(B) for which the factors of medical risk or types of medical intervention may be different for such individuals, or for which it is unknown whether such factors or types are different for such individuals; or
(C) with respect to which there has been insufficient research involving such individuals as subjects or insufficient data on such individuals.
(3) The term “minority group” has the meaning given the term “racial and ethnic minority group” in section 300o–6 of this title.
(4) The terms “minority” and “minorities” refer to individuals from a minority group.

(d) Health disparity populations
For purposes of this subpart:
(1) A population is a health disparity population if, as determined by the Director of the Institute after consultation with the Director of the Agency for Healthcare Research and Quality, there is a significant disparity in the overall rate of disease incidence, prevalence, morbidity, mortality, or survival rates in the population as compared to the health status of the general population.
(2) The Director shall give priority consideration to determining whether minority groups qualify as health disparity populations under paragraph (1).
(3) The term “health disparities research” means basic, clinical, and behavioral research on health disparity populations (including individual members and communities of such populations) that relates to health disparities as defined under paragraph (1), including the causes of such disparities and methods to prevent, diagnose, and treat such disparities.

(e) Coordination of activities
The Director of the Institute shall act as the primary Federal official with responsibility for coordinating all minority health disparities research and other health disparities research conducted or supported by the National Institutes of Health, and—
(1) shall represent the health disparities research program of the National Institutes of Health, including the minority health disparities research program, at all relevant Executive branch task forces, committees and planning activities; and
(2) shall maintain communications with all relevant Public Health Service agencies, including the Indian Health Service, and various other departments of the Federal Government to ensure the timely transmission of information concerning advances in minority health disparities research and other health disparities research between these various agencies for dissemination to affected communities and health care providers.

(f) Collaborative comprehensive plan and budget

(1) In general

Subject to the provisions of this section and other applicable law, the Director of NIH, the Director of the Institute, and the directors of the other agencies of the National Institutes of Health in collaboration (and in consultation with the advisory council for the Institute) shall—

(A) establish a comprehensive plan and budget for the conduct and support of all minority health disparities research and other health disparities research activities of the agencies of the National Institutes of Health (which plan and budget shall be first established under this subsection not later than 12 months after November 22, 2000);

(B) ensure that the plan and budget establish priorities among the health disparities research activities that such agencies are authorized to carry out;

(C) ensure that the plan and budget establish objectives regarding such activities, describes the means for achieving the objectives, and designates the date by which the objectives are expected to be achieved;

(D) ensure that, with respect to amounts appropriated for activities of the Institute, the plan and budget give priority in the expenditure of funds to conducting and supporting minority health disparities research;

(E) ensure that all amounts appropriated for such activities are expended in accordance with the plan and budget;

(F) review the plan and budget not less than annually, and revise the plan and budget as appropriate;

(G) ensure that the plan and budget serve as a broad, binding statement of policies regarding minority health disparities research and supporting minority health disparities research activities of the agencies, but do not remove the responsibility of the heads of the agencies for the approval of specific programs or projects, or for other details of the daily administration of such activities, in accordance with the plan and budget; and

(H) promote coordination and collaboration among the agencies conducting or supporting minority health or other health disparities research.

(2) Certain components of plan and budget

With respect to health disparities research activities of the agencies of the National Institutes of Health, the Director of the Institute shall ensure that the plan and budget under paragraph (1) provide for—

(A) basic research and applied research, including research and development with respect to products;

(B) research that is conducted by the agencies;

(C) research that is supported by the agencies;

(D) proposals developed pursuant to solicitations by the agencies and for proposals developed independently of such solicitations; and

(E) behavioral research and social sciences research, which may include cultural and linguistic research in each of the agencies.

(3) Minority health disparities research

The plan and budget under paragraph (1) shall include a separate statement of the plan and budget for minority health disparities research.

(g) Participation in clinical research

The Director of the Institute shall work with the Director of NIH and the directors of the agencies of the National Institutes of Health to carry out the provisions of section 289a–2 of this title that relate to minority groups.

(h) Research endowments

(1) In general

The Director of the Institute may carry out a program to facilitate minority health disparities research and other health disparities research by providing for research endowments—

(1) at centers of excellence under section 293 of this title; and

(2) at centers of excellence under section 285t–1 of this title.

(2) Eligibility

The Director of the Institute may provide for a research endowment under paragraph (1) only if the institution involved meets the following conditions:

(A) The institution does not have an endowment that is worth in excess of an amount equal to 50 percent of the national median of endowment funds at institutions that conduct similar biomedical research or training of health professionals.

(B) The application of the institution under paragraph (1) regarding a research endowment has been recommended pursuant to technical and scientific peer review and has been approved by the advisory council under subsection (j) of this section.

(i) Certain activities

In carrying out subsection (a) of this section, the Director of the Institute—

(1) shall assist the Director of the National Institute for Research Resources in carrying out section 287a–1(c)(3) of this title and in committing resources for construction at Institutions of Emerging Excellence;

(2) shall establish projects to promote cooperation among Federal agencies, State, local, tribal, and regional public health agen-
cies, and private entities in health disparities research; and
(3) may utilize information from previous health initiatives concerning minorities and other health disparity populations.

(j) Advisory council

(1) In general

The Secretary shall, in accordance with section 294a of this title, establish an advisory council to advise, assist, consult with, and make recommendations to the Director of the Institute on matters relating to the activities described in subsection (a) of this section, and with respect to such activities to carry out any other functions described in section 294a of this title for advisory councils under such section. Functions under the preceding sentence shall include making recommendations on budgetary allocations made in the plan under subsection (f) of this section, and shall include reviewing reports under subsection (k) of this section before the reports are submitted under such subsection.

(2) Membership

With respect to the membership of the advisory council under paragraph (1), a majority of the members shall be individuals with demonstrated expertise regarding minority health disparity and other health disparity issues; representatives of communities impacted by minority and other health disparities shall be included; and a diversity of health professionals shall be represented. The membership shall in addition include a representative of the Office of Behavioral and Social Sciences Research under section 283c of this title.

(h) Interagency coordination

The Director of the Institute, as the primary Federal official with responsibility for coordinating all research and activities conducted or supported by the National Institutes of Health on minority health and health disparities, shall plan, coordinate, review and evaluate research and other activities conducted or supported by the Institutes and Centers of the National Institutes of Health.

(2007—Subsec. (k). Pub. L. 109–482, §104(h)(1)(N), struck out heading and text of subsec. (k). Text read as follows: “The Secretary shall, in accordance with section 294a of this title, establish an advisory council to advise, assist, consult with, and make recommendations to the Director of the Institute on matters relating to the activities described in subsection (a) of this section, and with respect to such activities to carry out any other functions described in section 294a of this title for advisory councils under such section. Functions under the preceding sentence shall include making recommendations on budgetary allocations made in the plan under subsection (f) of this section, and shall include reviewing reports under subsection (k) of this section before the reports are submitted under such subsection.”)

Codification

Section was formerly classified to section 287c–31 of this title prior to renumbering by Pub. L. 111–148.

Amendments


Subsec. (a). Pub. L. 111–148, §10334(c)(1)(D)(ii), (iii), substituted “National Institute on Minority Health and Health Disparities” for “National Center on Minority Health and Health Disparities” and “Institute” for “Center”.

Subsec. (b), (d) to (g). Pub. L. 111–148, §10334(c)(1)(D)(iii), substituted “Institute” for “Center” wherever appearing.


Subsec. (h)(1). Pub. L. 111–148, §10334(c)(2)(A), in par. (1) of subsec. (h) relating to research endowments, substituted “research endowments—” (1) at centers of excellence under section 293 of this title; and (2) at centers of excellence under section 285–1 of this title.” for “research endowments at centers of excellence under section 293 of this title.”

Pub. L. 111–148, §10334(c)(1)(D)(iii), in par. (1) of subsec. (h), relating to research endowments, substituted “Institute” for “Center”.

Subsec. (h)(2). Pub. L. 111–148, §10334(c)(1)(D)(iii), in par. (2) of subsec. (h) relating to research endowments, substituted “Institute” for “Center” wherever appearing.

2007—Subsec. (k). Pub. L. 109–482, §104(h)(1)(N), struck out heading and text of subsec. (k). Text read as follows: “The Secretary shall, in accordance with section 294a of this title, establish an advisory council to advise, assist, consult with, and make recommendations to the Director of the Institute on matters relating to the activities described in subsection (a) of this section, and with respect to such activities to carry out any other functions described in section 294a of this title for advisory councils under such section. Functions under the preceding sentence shall include making recommendations on budgetary allocations made in the plan under subsection (f) of this section, and shall include reviewing reports under subsection (k) of this section before the reports are submitted under such subsection.”

(2) Membership

With respect to the membership of the advisory council under paragraph (1), a majority of the members shall be individuals with demonstrated expertise regarding minority health disparity and other health disparity issues; representatives of communities impacted by minority and other health disparities shall be included; and a diversity of health professionals shall be represented. The membership shall in addition include a representative of the Office of Behavioral and Social Sciences Research under section 283c of this title.

(h) Interagency coordination

The Director of the Institute, as the primary Federal official with responsibility for coordinating all research and activities conducted or supported by the National Institutes of Health on minority health and health disparities, shall plan, coordinate, review and evaluate research and other activities conducted or supported by the Institutes and Centers of the National Institutes of Health.

(2007—Pub. L. 109–482, §103(h)(1)(N), struck out heading and text of subsec. (h). Text read as follows: “The Secretary shall, in accordance with section 294a of this title, establish an advisory council to advise, assist, consult with, and make recommendations to the Director of the Institute on matters relating to the activities described in subsection (a) of this section, and with respect to such activities to carry out any other functions described in section 294a of this title for advisory councils under such section. Functions under the preceding sentence shall include making recommendations on budgetary allocations made in the plan under subsection (f) of this section, and shall include reviewing reports under subsection (k) of this section before the reports are submitted under such subsection.”)

Effective Date of 2007 Amendment

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

Findings

Pub. L. 106–525, §2, Nov. 22, 2000, 114 Stat. 2495, provided that: “The Congress finds as follows:
(1) Despite notable progress in the overall health of the Nation, there are continuing disparities in the burden of illness and death experienced by African Americans, Hispanics, Native Americans, Alaska Native, and Asian Pacific Islanders, compared to the United States population as a whole.
(2) The largest numbers of the medically underserved are white individuals, and many of them have the same health care access problems as do members sold...
of minority groups. Nearly 20,000,000 white individuals live below the poverty line with many living in nonmetropolitan, rural areas such as Appalachia, whereas the high percentage of counties designated as health professional shortage areas (47 percent) and the high rate of poverty contribute to disparity outcomes. However, there is a higher proportion of racial and ethnic minorities in the United States represented among the medically underserved.

(3) There is a national need for minority scientists in the fields of biomedical, clinical, behavioral, and health services research. Ninety percent of minority physicians educated at Historically Black Medical Colleges live and serve in minority communities.

(4) Demographic trends inspire concern about the Nation’s ability to meet its future scientific, technological, and engineering workforce needs. Historically, non-Hispanic white males have made up the majority of the United States scientific, technological, and engineering workers.

(5) The Hispanic and Black population will increase significantly in the next 50 years. The scientific, technological, and engineering workforce may decrease if participation by underrepresented minorities remains the same.

(6) Increasing rates of Black and Hispanic workers can help ensure a strong scientific, technological, and engineering workforce.

(7) Individuals such as underrepresented minorities and women in the scientific, technological, and engineering workforce enable society to address its diverse needs.

(8) If there had not been a substantial increase in the number of science and engineering degrees awarded to women and underrepresented minorities over the past few decades, the United States would be facing even greater shortages in scientific, technological, and engineering workers.

(9) In order to effectively promote a diverse and strong 21st century scientific, technological, and engineering workforce, Federal agencies should expand or add programs that effectively overcome barriers such as educational transition from one level to the next and student requirements for financial resources.

(10) Federal agencies should work in concert with the private nonprofit sector to emphasize the recruitment and retention of qualified individuals from ethnic and gender groups that are currently underrepresented in the scientific, technological, and engineering workforce.

(11) Behavioral and social sciences research has increased awareness and understanding of factors associated with health care utilization and access, patient attitudes toward health services, and risk and protective behaviors that affect health and illness. These factors have the potential to then be modified to help close the health disparities gap among ethnic minority populations. In addition, there is a shortage of minority behavioral science researchers and behavioral health care professionals. According to the National Science Foundation, only 15.5 percent of behavioral research-oriented psychology doctorate degrees were awarded to minority students in 1997. In addition, only 17.9 percent of practice-oriented psychology doctorate degrees were awarded to ethnic minorities.

§ 285t–1. Centers of excellence for research education and training

(a) In general

The Director of the Institute shall make awards of grants or contracts to designated biomedical and behavioral research institutions under paragraph (1) of subsection (c) of this section, or to consortia under paragraph (2) of such subsection, for the purpose of assisting the institutions in supporting programs of excellence in biomedical and behavioral research training for individuals who are members of minority health disparity populations or other health disparity populations.

(b) Required use of funds

An award may be made under subsection (a) of this section only if the applicant involved agrees that the grant will be expended—

(1) to train members of minority health disparity populations or other health disparity populations as professionals in the area of biomedical or behavioral research or both; or

(2) to expand, remodel, renovate, or alter existing research facilities or construct new research facilities for the purpose of conducting minority health disparities research and other health disparities research.

(c) Centers of excellence

(1) In general

For purposes of this section, a designated biomedical and behavioral research institution is a biomedical and behavioral research institution that—
(A) has a significant number of members of minority health disparity populations or other health disparity populations enrolled as students in the institution (including individuals accepted for enrollment in the institution);  
(B) has been effective in assisting such students of the institution to complete the program of education or training and receive the degree involved;  
(C) has made significant efforts to recruit minority students to enroll in and graduate from the institution, which may include providing means-tested scholarships and other financial assistance as appropriate; and  
(D) has made significant recruitment efforts to increase the number of minority or other members of health disparity populations serving in faculty or administrative positions at the institution.  

(2) Consortium  
Any designated biomedical and behavioral research institution involved may, with other biomedical and behavioral institutions (designated or otherwise), including tribal health programs, form a consortium to receive an award under subsection (a) of this section.  

(3) Application of criteria to other programs  
In the case of any criteria established by the Director of the Institute for purposes of determining whether institutions meet the conditions described in paragraph (1), this section may not, with respect to minority health disparity populations or other health disparity populations, be construed to authorize, require, or prohibit the use of such criteria in any program other than the program established in this section.  

(d) Duration of grant  
The period during which payments are made under a grant under subsection (a) of this section may not exceed 5 years. Such payments shall be subject to annual approval by the Director of the Institute and to the availability of appropriations for the fiscal year involved to make the payments.  

(e) Maintenance of effort  
(1) In general  
With respect to activities for which an award under subsection (a) of this section is authorized to be expended, the Director of the Institute may make such an award only if the institutions involved agree that the institutions will, before expending the award, expend the Federal amounts obtained from sources other than the award.  

(f) Certain expenditures  
The Director of the Institute may authorize a designated biomedical and behavioral research institution to expend a portion of an award under subsection (a) of this section for research endowments.  

(g) Definitions  
For purposes of this section:  
(1) The term “designated biomedical and behavioral research institution” has the meaning indicated for such term in subsection (c)(1) of this section. Such term includes any health professions school receiving an award of a grant or contract under section 293 of this title.  
(2) The term “program of excellence” means any program carried out by a designated biomedical and behavioral research institution with an award under subsection (a) of this section, if the program is for purposes for which the institution involved is authorized in subsection (b) of this section to expend the grant.  


CODIFICATION  
Section was formerly classified to section 287c–32 of this title prior to renumbering by Pub. L. 111–148.  

AMENDMENTS  
2010—Subsecs. (a), (c)(3) to (f). Pub. L. 111–148, § 10334(c)(1)(D)(i), substituted “Institute” for “Center” wherever appearing.  
2007—Subsec. (b). Pub. L. 109–482 struck out heading and text of subsec. (b). Text read as follows: “For the purpose of making grants under subsection (a) of this section, there are authorized to be appropriated such sums as may be necessary for each of the fiscal years 2001 through 2005.”  

EFFECTIVE DATE OF 2007 AMENDMENT  
Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.  

§ 285t–2. Loan repayment program for minority health disparities research  
(a) In general  
The Director of the Institute shall establish a program of entering into contracts with qualified health professionals under which such health professionals agree to engage in minority health disparities research or other health disparities research in consideration of the Federal Government agreeing to repay, for each year of engaging in such research, not more than $35,000 of the principal and interest of the educational loans of such health professionals.
(b) Service provisions

The provisions of sections 254l–1, 254m, and 254o of this title shall, except as inconsistent with subsection (a) of this section, apply to the program established in such subsection to the same extent and in the same manner as such provisions apply to the National Health Service Corps Loan Repayment Program established in subpart III of part D of subchapter II of this chapter.

(c) Requirement regarding health disparity populations

The Director of the Institute shall ensure that not fewer than 50 percent of the contracts entered into under subsection (a) of this section are for appropriately qualified health professionals who are members of a health disparity population.

(d) Priority

With respect to minority health disparities research and other health disparities research under subsection (a) of this section, the Secretary shall ensure that priority is given to conducting projects of biomedical research.


CODIFICATION

Section was formerly classified to section 287c–33 of this title prior to renumbering by Pub. L. 111–148.

AMENDMENTS

2010—Pub. L. 111–148, §10334(c)(1)(D)(iii), substituted “Institute” for “Center” in section catchline and text.

2007—Pub. L. 109–482 struck out subsec. (a) designation and heading before “The Secretary” and struck out subsec. (b) which related to evaluation of this subpart not later than 5 years after Nov. 22, 2000, and report on such evaluation not later than 1 year after its commencement.

Effective Date of 2007 Amendment

Amendment by Pub. L. 109–482 applicable only with respect to amounts appropriated for fiscal year 2007 or subsequent fiscal years, see section 109 of Pub. L. 109–482, set out as a note under section 281 of this title.

PART D—NATIONAL LIBRARY OF MEDICINE

SUBPART 1—GENERAL PROVISIONS

§ 286. National Library of Medicine

(a) Purpose and establishment

In order to assist the advancement of medical and related sciences and to aid the dissemination and exchange of scientific and other information important to the progress of medicine and to the public health, there is established the National Library of Medicine (hereafter in this part referred to as the “Library”).

(b) Functions

The Secretary, through the Library and subject to subsection (d) of this section, shall—

(1) acquire and preserve books, periodicals, prints, films, recordings, and other library materials pertinent to medicine;

(2) organize the materials specified in paragraph (1) by appropriate cataloging, indexing, and bibliographical listings;

(3) publish and disseminate the catalogs, indexes, and bibliographies referred to in paragraph (2);

(4) make available, through loans, photographic or other copying procedures, or otherwise, such materials in the Library as the Secretary determines appropriate;

(5) provide reference and research assistance;

(6) publicize the availability from the Library of the products and services described in any of paragraphs (1) through (5);

(7) promote the use of computers and telecommunications by health professionals (including health professionals in rural areas) for the purpose of improving access to biomedical information for health care delivery and medical research;

(8) engage in such other activities as the Secretary determines appropriate and as the Library’s resources permit.

(c) Exchange, destruction, or disposal of materials not needed

The Secretary may exchange, destroy, or otherwise dispose of any books, periodicals, films, and other library materials not needed for the permanent use of the Library.

(d) Availability of publications, materials, facilities, or services; prescription of rules

(1) The Secretary may, after obtaining the advice and recommendations of the Board of Regents, prescribe rules under which the Library will—