

FACT Book

FISCAL YEAR

2011



NATIONAL INSTITUTES OF HEALTH
NATIONAL HEART, LUNG, AND BLOOD INSTITUTE



FACT Book

FISCAL YEAR

2011

FEBRUARY 2012

FOR ADMINISTRATIVE USE

NATIONAL INSTITUTES

OF HEALTH

NATIONAL HEART, LUNG,

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NATIONAL INSTITUTES OF HEALTH
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1. Abbreviated Staff Directory*

Office of the Director	Bldg.	Room	Phone	MSC**†
Acting Director, Susan B. Shurin, M.D.	31	5A48	496-5166	2486
Acting Deputy Director, Carl A. Roth, Ph.D., LL.M.	31	5A07	496-6331	2482
Chief of Staff, Sheila Pohl	31	5A48	594-5355	2486
Administrative Officer, Cheryl R. Wagoner	31	5A16	496-5931	2490
Executive Officer, Kathleen B. O'Sullivan	31	5A48	496-2411	2490
Deputy Executive Officer, Timothy J. Wheeles	31	5A48	496-2411	2490
Administrative Officer, Cheryl R. Wagoner	31	5A16	496-5931	2490
Associate Director for Basic Research, Alan M. Michelson, M.D., Ph.D.	31	5A48	594-5353	2490
Center for Biomedical Informatics				
Acting Director, Kathleen B. O'Sullivan, Ph.D.	RKL1‡	6100	435-0119	7994
Administrative Officer, Kathleen D. Rechen	RKL2§	8095	435-6373	7921
Customer Support Branch				
Acting Chief, Brian Kotula	RKL1§	6106	435-0119	7994
Infrastructure Engineering Branch				
Chief, Christopher E. Olaes	RKL1	6212	435-0119	7994
Operation and Performance Management Branch				
Acting Chief, Matthew Raschka	RKL1	6112	435-0119	7994
Software Engineering Branch				
Acting Chief, Zeyad Mobassaleh	RKL1	6104	435-0119	7994
Center for Population Studies				
Director, Daniel Levy, M.D.			73 Mt. Wayte Avenue, Suite 2	
			Framingham, MA 01702-5827	
			508-935-3458	
Associate Director, Christopher J. O'Donnell, M.D., M.P.H.			73 Mt. Wayte Avenue, Suite 2	
			Framingham, MA 01702-5827	
			508-935-3435	
Ethics Office				
Director, Nancy O'Hanlon, J.D.	31	5A33	496-6471	2486
Lead Ethics Specialist, Kim Y. Brinson	31	5A33	496-6471	2486
Ethics Coordinator, Hedy S. Tam	31	5A33	496-6471	2486
Administrative Officer, Cheryl R. Wagoner	31	5A16	496-5931	2490

* Current as of September 30, 2011. For locating personnel not listed, the general information number is 301-496-4000. All listed phone numbers are in area code 301. The Personnel Directory, which is periodically updated throughout the year, is located on the NHLBI Home Page under About NHLBI.

** MSC—Mail Stop Code.

† Full mailing address formats are located at the end of this chapter.

‡ RKL1—Rockledge I Building.

§ RKL2—Rockledge II Building.

Office of the Director (continued)	Bldg.	Room	Phone	MSC
Office of Administrative Management				
Executive Officer, Kathleen B. O'Sullivan	31	5A48	496-2411	2490
Deputy Executive Officer, Timothy J. Wheeler	31	5A48	496-2411	2490
Administrative Officer, Cheryl R. Wagoner	31	5A16	496-5931	2490
Program Advisor, Marilyn G. Jackson	31	5A25	594-4690	2490
Administrative Officer, Tho-Van Tran	31	5A16	496-5931	2490
Extramural Administrative Management Branch				
Chief, Loretta L. Usilton	RKL2	8095	435-6373	7921
Financial Management Branch				
Chief, Sandra L. Gault	31	5A34	496-4653	2490
Freedom of Information and Privacy Act Branch				
Chief, Suzanne A. Freeman	RKL1	6070	496-9737	7957
Management Policy and Administrative Services Branch				
Chief, Alesha M. Holiday	31	5A16	496-5931	2490
Acting Deputy Chief, Jim Mitchel	31	5A16	496-5931	2490
Administrative Officer, Cheryl R. Wagoner	31	5A16	496-5931	2490
Office of Workforce Management				
Director, Gwen G. Platt	RKL1	6070	496-1763	7957
Office of Communications				
Director, Sally McDonough	31	4A31	496-5804	2490
Deputy Director, Susan G. Dambrauskas	31	4A31	496-4236	2480
Administrative Officer, Tho-Van Tran	31	5A16	496-5931	2490
Health Campaigns and Consumer Services Branch				
Chief, Ann M. Taubenheim, Ph.D.	31	4A31	496-4236	2480
Public Affairs Branch				
Chief, Diane E. Striar	31	4A31	496-4236	2480
Office of Global Health				
Director, Arun Chockalingam, Ph.D., M.S.	31	5A06	496-3620	2490
Deputy Director, Cristina Rabidan-Diehl, Ph.D., M.P.H.	31	5A06	496-3620	2490
Administrative Officer, Tho-Van Tran	31	5A16	496-5931	2490
Office of Research Training and Minority Health				
Director, Helena O. Mishoe, Ph.D., M.P.H.	RKL2	9093C	451-5081	7913
Deputy Director, Chitra Krishnamurti, Ph.D.	RKL2	9093C	451-5081	7913
Administrative Officer, Roy Rich	RKL2	8095	435-6373	7921
Office of Science and Technology				
Director, Carl A. Roth, Ph.D., LL.M.	31	5A07	496-6331	2482
Deputy Director, Barbara R. Marzetta, M.S.	31	5A07	496-9899	2482
Administrative Officer, Cheryl R. Wagoner	31	5A16	496-5931	2490
Program Studies and Reports Program				
Director, Carl A. Roth, Ph.D., LL.M.	31	5A07	496-6331	2482
Science and Special Issues Program				
Director, Barbara R. Marzetta, M.S.	31	5A07	496-9899	2482
Office of Public Liaison				
Coordinator, Hilary S. Leeds, J.D.	31	5A07	594-9869	2482

Office of the Director (continued)	Bldg.	Room	Phone	MSC
Office of Technology Transfer and Development				
Director, Alan H. Deutch	RKL1	6018	402-5579	7992
Administrative Officer, Terri B. Pike	RKL2	8095	435-6373	7921
Division of Cardiovascular Sciences				
Office of the Director				
Director, Michael S. Lauer, M.D.	RKL2	8128	435-0422	7959
Deputy Director, Sonia I. Skarlatos, Ph.D.	RKL2	8248	435-0466	7940
Administrative Officer, Lisa A. Freeny	RKL2	8095	435-6373	7921
Office of Special Projects				
Special Assistant for Clinical Studies, David J. Gordon, M.D., Ph.D.	RKL2	8134	435-0534	7940
Office of Biostatistics Research				
Director, Nancy L. Geller, Ph.D.	RKL2	9202	435-0434	7913
Office of Research Training and Career Development				
Director, Jane D. Scott, Sc.D., M.S.N.	RKL2	8138	435-0535	7940
Adult and Pediatric Cardiac Research Program				
Director, Gail D. Pearson, M.D., Sc.D.	RKL2	8104	435-0510	7940
Atherosclerosis and Coronary Artery Disease Branch				
Chief, Yves D. Rosenberg, M.D.	RKL2	8146	435-1292	7956
Heart Developmental and Structural Diseases Branch				
Chief, Jonathan R. Kaltman, M.D.	RKL2	8104	435-0510	7940
Heart Failure and Arrhythmias Branch				
Chief, David A. Lathrop, Ph.D.	RKL2	8170	435-0504	7956
Basic and Early Translational Research Program				
Director, Denis B. Buxton, Ph.D.	RKL2	8216	435-0513	7940
Advanced Technologies and Surgery Branch				
Chief, Marissa Miller, D.V.M., M.P.H.	RKL2	8214	435-0513	7940
Vascular Biology and Hypertension Branch				
Chief, Zorina S. Galis, Ph.D.	RKL2	8116	435-0560	7940
Prevention and Population Sciences Program				
Director, Diane E. Bild, M.D., M.P.H.	RKL2	10018	435-0457	7936
Clinical Applications and Prevention Branch				
Chief, Lawrence J. Fine, M.D.	RKL2	10216	435-0305	7936
Epidemiology Branch				
Chief, Paul D. Sorlie, Ph.D.	RKL2	10210	435-0456	7936
Women's Health Initiative Branch				
Chief, Jacques E. Rossouw, M.D.	RKL2	9192	402-2900	7913
Jackson Heart Study				
Director, (Vacant)		350 West Woodrow Wilson Drive Jackson, MS 39213 601-979-8754		
Division of Lung Diseases				
Office of the Director				
Director, James P. Kiley, Ph.D., M.S.	RKL2	10042	435-0233	7952
Deputy Director, Gail G. Weinmann, M.D.	RKL2	10042	435-0233	7952
Administrative Officer, Amy W. Sheetz	RKL2	8095	435-6373	7921

Division of Lung Diseases (continued)	Bldg.	Room	Phone	MSC
Airway Biology and Disease Branch				
Chief, Thomas L. Croxton, Ph.D., M.D.	RKL2	10042	435-0202	7952
Lung Biology and Disease Branch				
Chief, Dorothy B. Gail, Ph.D.	RKL2	10042	435-0222	7952
National Center on Sleep Disorders Research				
Director, Michael J. Twery, Ph.D.	RKL2	10042	435-0199	7952
Research Training Programs				
Leader, Sandra Colombini Hatch, M.D.	RLK2	10042	435-0222	7952
Leader, Ann E. Rothgeb	RLK2	10042	435-0202	7952

Division of Blood Diseases and Resources

Office of the Director				
Director, W. Keith Hoots, M.D.	RKL2	9136	435-0080	7950
Deputy Director, Donna M. DiMichele, M.D.	RKL2	9132	435-0080	7950
Administrative Officer, Amy W. Sheetz	RKL2	8095	435-6373	7921
Senior Program Analyst, Susan E. Pucie	RKL2	9138	435-0080	7950
Blood Diseases Branch				
Chief, Harvey S. Luksenburg, M.D.	RKL2	9160	435-0050	7950
Thrombosis and Hemostasis Branch				
Acting Chief, Donna M. DiMichele, M.D.	RKL2	9132	435-0080	7950
Transfusion Medicine and Cellular Therapeutics Branch				
Chief, Simone A. Glynn, M.D.	RKL2	9142	435-0065	7950
Research Training Programs				
Leader, Lis Welniak, Ph.D.	RKL2	9143	435-0065	7950
Leader, Rita Sarkar, Ph.D.	RKL2	9161	435-0070	7950
Leader, Ellen M. Werner, Ph.D.	RKL2	9162	435-0050	7950
Leader, Henry Chang, M.D.	RKL2	9176	435-0080	7950

Division for the Application of Research Discoveries

Director, Denise G. Simons-Morton, M.D, Ph.D.	31	4A11	496-5437	2480
Acting Deputy Director, Karen A. Donato, S.M.	31	4A11	496-1051	2480
Administrative Officer, Tho-Van Tran	31	5A16	496-5931	2490
Senior Manager for Program Operations				
Mishyelle Croom, M.A.	31	4A11	496-5437	2480
Enhanced Dissemination and Utilization Branch				
Acting Chief, Karen A. Donato, S.M.	31	4A11	496-1051	2480
Research Translation Branch				
Acting Chief, Denise G. Simons-Morton, M.D., Ph.D.	31	4A11	496-5437	2480

Division of Extramural Research Activities

Office of the Director				
Director, Stephen C. Mockrin, Ph.D.	RKL2	7100	435-0260	7922
Deputy Director, Jodi B. Black, Ph.D.	RKL2	7104	435-0260	7922
Chief of Staff, Marianna Mertts, Ph.D.	RKL2	7220	435-0260	7922
Administrative Officer, Veronica M. VanWagner	RKL2	8095	435-6373	7921
Office of Acquisitions				
Director, John C. Taylor	RKL2	6100	435-0330	7902
Deputy Director, Christopher J. Belt	RKL2	6106	435-6672	7902

Division of Extramural Research Activities (continued)	Bldg.	Room	Phone	MSC
Special Assistant to the Director				
Debra C. Hawkins	RKL2	6224	435-0361	7902
Consolidated Operations Acquisition Center Services Branch				
Chief, Joanna Magginas	RKL2	6150	435-0360	7902
NHLBI Extramural Contracts Branch				
Chief, Richard A. Phillips, M.B.A.	RKL2	6136	402-6462	7902
Office of Committee Management				
Director, Kathryn M. Valeda	RKL2	7110	435-0255	7922
Deputy Director & Scientific Review Evaluation Activities				
Administrator, David Alperin	RLK2	7118	435-0255	7922
Office of Extramural Policy and Review				
Director, Paul A. Velletri, Ph.D.	RKL2	7218	435-0569	7922
Office of Scientific Review				
Director, Valerie L. Prenger, Ph.D.	RKL2	7214	435-0270	7924
Referral Officer, Roy White, Ph.D.	RKL2	7176	435-0310	7924
Blood and Vascular Branch				
Chief, Jeffrey H. Hurst, Ph.D.	RKL2	7208	435-0303	7924
Cardiovascular and Pulmonary Branch				
Chief, William J. Johnson, Ph.D.	RKL2	7178	435-0725	7924
Clinical Studies and Training Branch				
Chief, Charles W. Joyce, Ph.D.	RKL2	7194	435-0288	7924
Office of Grants Management				
Director, Suzanne A. White	RKL2	7160	435-0166	7926
Deputy Director, Ryan C. Lombardi	RKL2	7172	435-0166	7926
Blood Diseases and Resources and Lung Diseases Branch				
Chief, Theresa R. Jarosik	RKL2	7156	435-0166	7926
Cardiovascular Sciences Branch				
Chief, Teresa F. Marquette	RKL2	7130	435-0166	7926
Office of Translational Alliances and Coordination				
Acting Director, Jodi B. Black, Ph.D.	RKL2	7104	435-0260	7922

Division of Intramural Research

Office of the Director				
Director, Robert S. Balaban, Ph.D.	10CRC*	4-1581	496-2116	1458
Office of the Scientific Director				
Director, Robert S. Balaban, Ph.D.	10CRC	4-1581	496-2116	1458
Deputy Director, L. Michelle Bennett, Ph.D.	10CRC	4-1581	496-2116	1458
Intramural Administrative Management Branch				
Chief, Gary Unger	10	7N214	451-0892	1686
Animal Program				
Director, James Hawkins, D.V.M.	14E	105C	451-6743	5570
Animal Care and Use Committee				
Program Coordinator, Kelly E. Cole	14E	106C	451-6459	5570
Laboratory of Animal Medicine and Surgery				
Chief, Robert Hoyt, D.V.M.	14E	105B	496-9673	5570
Imaging Probe Development Center/Road Map Initiative				
Chief, Gary L. Griffiths, Ph.D.	B**	3042	217-5770	3372

* 10CRC—Building 10 Clinical Research Center.

**B—B Building, off site.

Division of Intramural Research (continued)	Bldg.	Room	Phone	MSC
Office of Education				
Director, Herbert M. Geller, Ph.D.	10	6N248	451-9440	1754
Office of the Clinical Director				
Director, Richard O. Cannon III, M.D.	10CRC	5-3330	496-9895	1454
Deputy Director, Richard Childs, M.D.	10CRC	3-5330	451-7128	1230
Office of Clinical Affairs				
Associate Director, Melissa B. Bryant, M.S.	10CRC	6-5140	594-8378	1608
Hematology Branch				
Chief, Neal S. Young, M.D.	10CRC	3-5140	496-5093	1202
Flow Cytometry Core (FACS)				
Head, J. Philip McCoy, Ph.D.	10	8C104	451-8824	1357
Center for Molecular Medicine				
Chief, Toren Finkel, M.D., Ph.D.	10CRC	5-3330	402-4081	1454
Transgenic Core				
Head, Chengyu Liu, Ph.D.	50	3305	435-5034	8018
Human iPS Core				
Head, Guokai Chen, Ph.D.	10	6N240	594-4717	1754
Cardiovascular-Pulmonary Branch				
Chief, Andrew E. Arai, M.D.	10CRC	B1D416	496-3648	1061
MRI/Imaging Core				
Head, Stasia Anderson, Ph.D.	10	B1D416	402-0908	1061
Biochemistry and Biophysics Center				
Director, Nico Tjandra, Ph.D.	50	3503	402-3029	8013
Biophysics Core				
Head, Grzegorz Piszczek, Ph.D.	50	2341	435-8082	8012
Protein Analysis Core				
Head, Duck-Yeon Lee, Ph.D.	50	2339	435-8369	8012
Cell Biology and Physiology Center				
Director, Clare Waterman, Ph.D.	50	4535	435-2949	8019
Light Microscopy Core				
Head, Christian Combs, Ph.D.	10	6N309	496-3236	1623
Genetics and Development Biology Center				
Director, Alan M. Michelson, M.D., Ph.D.	31	5A48	594-5353	2490
Electron Microscopy Core				
Head, Mathew Daniels, Ph.D.	14E	111B	496-2898	5570
Pathology Core				
Acting Head, Zu-Xi Yu, Ph.D.	10	6D19	496-5035	1590
Genomics Core				
Head, Nalini Raghavachari, Ph.D.	10	8C103B	435-2304	1357
DNA Sequencing Core				
Head, Jun Zhu, Ph.D.	10	5N107	443-7927	1654
Immunology Center				
Director, Warren Leonard, M.D.	10	7B05	496-0098	1674
Systems Biology Center				
Director, Keji Zhao, Ph.D.	10	7B06A	496-2098	1674
Proteomics Core				
Head, Marjan Gucek, Ph.D.	10	8C103C	594-1060	1774

NIH Mailing Address Formats

NHLBI staff e-mail addresses can be found by using the NIH Directory and E-mail Forwarding Service located on the Internet at <http://directory.nih.gov>.

Please use the following formats for NIH mailing addresses:

Building 10	Full Name NHLBI, NIH Building 10, Room ____ 10 Center Drive MSC* ____ Bethesda, MD 20892-MSC**	Building B	Full Name NHLBI, NIH Building B, Room 3042 9800 Medical Center Drive MSC 3372 Bethesda, MD 20892-3372
Building 14E	Full Name NHLBI, NIH Building 14E, Room ____ 14 Service Road West MSC* ____ Bethesda, MD 20892-MSC**	Rockledge I Building	Full Name NHLBI, NIH One Rockledge Center, Room ____ 6705 Rockledge Drive MSC* ____ Bethesda, MD 20817-MSC**
Building 31	Full Name NHLBI, NIH Building 31, Room ____ 31 Center Drive MSC* ____ Bethesda, MD 20892-MSC**	Rockledge II Building	Full Name NHLBI, NIH Two Rockledge Center, Room ____ 6701 Rockledge Drive MSC* ____ Bethesda, MD 20817-MSC**
Building 50	Full Name NHLBI, NIH Building 50, Room ____ 50 South Drive MSC* ____ Bethesda, MD 20892-MSC**		

* Retain the letters MSC before adding the mail stop code number.

**Replace the letters MSC with the mail stop code number.



2. Program Overview

The National Heart Institute (NHI) was established in 1948 through the National Heart Act with a mission to support research and training in the prevention, diagnosis, and treatment of cardiovascular diseases (CVD). Twenty-four years later—through section 413 of the National Heart, Blood Vessel, Lung, and Blood Act (P.L. 92-423)—Congress mandated the Institute to expand and coordinate its activities in an accelerated attack against heart, blood vessel, lung, and blood diseases. The renamed National Heart, Lung, and Blood Institute (NHLBI) expanded its scientific areas of interest and intensified its efforts related to research on diseases within its purview. Over the years, the Institute's areas of interest have grown to encompass genetic, genomic, proteomic, and metabolomic research; systems biology; sleep disorders; and the Women's Health Initiative (WHI).

The NHLBI provides global leadership for research, training, and education programs to promote the prevention and treatment of heart, lung, and blood diseases and enhance the health of all individuals so that they can live longer and more fulfilling lives.

The NHLBI stimulates basic discoveries about the causes of disease, enables the translation of basic discoveries into clinical practice, fosters training and mentoring of emerging scientists and physicians, and communicates research advances to the public. It creates and supports a robust, collaborative research infrastructure in partnership with private and public organizations, including academic institutions, industry, and other government agencies. The Institute collaborates with patients, families, health care professionals, scientists, professional societies, patient advocacy groups, community organizations, and the media to promote the application of research results and leverage resources to address the health needs of the public. The NHLBI also collaborates with international organizations to help reduce the burden of heart, lung, and blood diseases worldwide.

Each year, the NHLBI assesses progress in the scientific areas for which it is responsible and updates its goals and objectives. As new opportunities are identified, the Institute expands and revises its areas of interest. Throughout the process, the approach used by the Institute is an orderly sequence of research activities that includes:

- Acquisition of knowledge
- Evaluation of knowledge
- Application of knowledge
- Dissemination of knowledge

NHLBI Programs

The programs of the NHLBI, as shown in the following table, are implemented through four extramural units:

- Division of Cardiovascular Sciences (DCVS)
- Division of Lung Diseases (DLD)
- Division of Blood Diseases and Resources (DBDR)
- Division for the Application of Research Discoveries (DARD)

and one intramural unit:

- Division of Intramural Research (DIR)

The extramural divisions use a variety of funding mechanisms, such as individual research project grants, cooperative agreements, program project grants, Small Business Innovation Research (SBIR) grants, Small Business Technology Transfer (STTR) grants, Specialized Centers of Clinically Oriented Research (SCCOR) grants, comprehensive center grants, contracts, and research training and career development grants.

In fiscal year (FY) 2010, the DCVS was created by combining two previously existing divisions—the Division of Cardiovascular Diseases and the Division

Programs Supported by the National Heart, Lung, and Blood Institute

Cardiovascular Diseases

Advanced Technologies and Surgery

Diagnostics Development
Emerging Therapeutics
Enabling Technologies
Surgical Advances

Atherothrombosis and Coronary Artery Disease

Acute and Chronic Coronary Syndromes
Acute and Silent Ischemia
Angina
Atherothrombosis
Coronary Artery Disease
Myocardial Infarction
Revascularization

Clinical Applications and Prevention

Behavioral Medicine
Prevention of Cardiovascular Disorders
Obesity Health Outcomes

Epidemiology

Analytical Resources
Field Studies and Clinical Epidemiology
Genetic Epidemiology

Heart Development and Structural Disease

Adult Congenital Disease
Cardiac Immunology and Infection
Cardiovascular Development
Heart Transplantation
Pediatric Cardiovascular Disease
Valvular Heart Disease

Heart Failure and Arrhythmias

Arrhythmias
Heart Failure
Myocardial Protection
Resuscitation
Sudden Cardiac Death

Vascular Biology and Hypertension

Aneurysms
Cerebrovascular Disease
Hypertension
Lymphatic Diseases
Peripheral Vascular Disease
Renal Vascular Disease
Vascular Biology
Vascular Development and Angiogenesis
Venous Disease

Cardiovascular Diseases (continued)

Women's Health Initiative

Hormone Therapy Trial
Dietary Modification Trial
Calcium and Vitamin D Trial
Observational Study
Memory Study

Lung Diseases

Airway Biology and Disease

Asthma
Chronic Obstructive Pulmonary Disease (COPD) and Environmental Lung Diseases
Cystic Fibrosis (CF)
Genetics, Genomics, and Biotechnology

Lung Biology and Disease

Acquired Immunodeficiency Syndrome (AIDS) and Tuberculosis (TB)
Critical Care and Acute Lung Injury
Developmental Biology and Pediatric Lung Disease
Immunology and Fibrosis
Lung Cell and Vascular Biology

National Center on Sleep Disorders Research

Neurobiology and Sleep
Sleep Disorders Medicine

Blood Diseases and Resources

Blood Diseases

Anemias
Erythropoiesis
Malaria
Red Cells
Sickle Cell Disease (SCD)
Thalassemia

Thrombosis and Hemostasis

Hematologic Immune Disorders
Hemophilia and Other Bleeding Disorders
Hemostasis
Immunity and Inflammation
Thrombosis

Blood Diseases and Resources (continued)

Transfusion Medicine and Cellular Therapeutics

Hematopoietic Stem Cell Transplantation
Immune Deficiencies, Reconstitution, Response, and Tolerance
Myelodysplasia, Marrow Failure, and Myeloproliferative Disorders
Novel Cellular Therapies for Repair and Regeneration
Stem Cell Biology
Transfusion Medicine Use, Safety, and Availability of Blood and Blood Components

Application of Research Discoveries

Research Translation Branch

Clinical Practice Guidelines
Development for Heart, Lung, and Blood Topics
Implementation for Heart, Lung, and Blood Topics
Knowledge Exchange Networks

Enhanced Dissemination and Utilization Branch

Community Health Promotion
Childhood Obesity Prevention
Health Disparities Reduction
Program Evaluations

Intramural Research

Clinical Research

Cardiothoracic Surgery
Hematology
Pulmonary and Vascular Medicine
Translational Medicine

Laboratory Research

Biochemistry and Biophysics
Cell Biology and Physiology
Genetics and Development Biology
Immunology

of Prevention and Population Sciences—so that the administrative structure would better match the dynamic interaction that exists among basic, clinical, and population sciences. Because the areas addressed by the two previous divisions are closely linked, the Institute believed that merging the two Divisions would stimulate the collaborative efforts that are needed to advance cardiovascular research.

Descriptions of the Divisions follow.

Division of Cardiovascular Sciences

The DCVS supports basic, clinical, population, and health services research on the causes, prevention, and treatment of CVD and technology development for its diagnosis and treatment. The Division fosters research in atherosclerosis, coronary artery disease, myocardial infarction and ischemia, heart failure, arrhythmia, sudden cardiac death, adult and pediatric congenital heart disease, high blood pressure, stroke, cardiovascular complications of diabetes and obesity, and other cardiovascular disorders. A SCCOR supports clinical collaborative research in vascular injury, repair, and remodeling, and a Centers Program supports cardiac translational research associated with preventing and treating heart failure and arrhythmias.

The Division's research portfolio includes a number of well-known epidemiological cohort studies that describe disease and risk factor patterns in populations; clinical trials of interventions to prevent disease and to reduce or eliminate risk factors; studies of the influence of genetic, behavioral, sociocultural, environmental, and health systems factors on disease risk and outcomes; and studies of the application of prevention and treatment strategies to determine ways to improve clinical care and public health. The Division also supports research training and career development in these areas.

In addition to the Office of the Director, the Division is organized into three Programs, eight Branches, and three Offices, which are described below.

Basic and Early Translational Research Program

The Basic and Early Translational Research Program supports research and research training and career development in vascular biology and hypertension, cardiovascular

surgery, and development of advanced technologies for the diagnosis and treatment of CVD. The portfolio includes an integrated basic and clinical research program to study the biological basis for vascular diseases and hypertension and their diagnosis, treatment, and prevention. Research on cardiovascular surgery includes both basic and preclinical research on surgical approaches and clinical trials to establish evidence-based surgical therapies. The development of diagnostics encompasses research on biosensors, imaging technologies, and the application of “omic” methodologies. Therapeutic development includes drug and nucleic acid delivery technologies, regenerative and reparative medicine, gene therapy, and device development.

The Program is divided into the two branches described below.

Advanced Technologies and Surgery Branch

The Advanced Technologies and Surgery Branch supports integrated basic, translational, and clinical research to develop technologies for the diagnosis, prevention, and treatment of CVD. Research on diagnostics focuses on proteomic, genomic, and other biomarker technologies and on imaging modalities and agents. Therapeutics research focuses on tissue-, cell-, and gene-based therapies; regenerative and reparative medicine; image-guided therapies; and cardiac and circulatory support and repair devices. Research related to surgery addresses improved surgical and image-guided therapies and the translation of cardiovascular surgical advances into clinical practice. Enabling technologies research includes bioinformatics, computational and systems biology, bioengineering, nanotechnology, materials research, and personalized medicine.

Vascular Biology and Hypertension Branch

The Vascular Biology and Hypertension Branch supports integrated basic, translational, and clinical research on the etiology, pathogenesis, prevention, diagnosis, and treatment of vascular diseases and hypertension. Vascular biology focuses on angiogenesis; development and repair of arteries, veins, lymphatics, and microcirculation; and biology of the endothelium and other vascular wall components. Vascular disease research focuses on diseases affecting peripheral (non-coronary) arteries, including the aorta and cerebral, renal, and limb vessels; veins; and lymphatics. Hypertension research focuses on the study of blood pressure regulation—including

central, renal, and vascular control—and end organ damage resulting from high blood pressure.

Adult and Pediatric Cardiac Research Program

The Adult and Pediatric Cardiac Research Program supports and provides leadership for basic, translational, and clinical research on development, maturation, and functioning of the heart throughout all stages of life. Areas of research include cardiac development and maturation, myocyte structure and function, myocardial energetics and metabolism, cardiac electrophysiology, coronary artery structure and function, the failing heart, valvular heart disease, exercise physiology, nutrition and the heart, congenital heart disease from birth through adulthood, the intrauterine environment and cardiovascular risk, cardiomyopathy, atherothrombosis, and coronary artery disease. A major function of the Program is to provide collaborative leadership for systematic oversight of clinical research across the Division, including clinical research information technology and standard but flexible operating procedures.

The Program is organized into the three branches described below.

Atherothrombosis and Coronary Artery Disease Branch

The Atherothrombosis and Coronary Artery Disease Branch conducts and manages an integrated basic and clinical research program to study the etiology, pathogenesis, prevention, diagnosis, and treatment of coronary artery disease and atherothrombosis. Research on coronary artery disease focuses on acute and chronic coronary syndromes, including myocardial infarction; acute ischemia, angina, and silent ischemia; and percutaneous and surgical revascularization of stenotic and restenotic coronary lesions. Atherothrombosis research investigates atherosclerotic lesions in coronary arteries and other arterial beds; lipid fractions and interactions with the arterial wall; lesion instability, vulnerable plaques, and thrombosis; and biomarker and imaging diagnostics to quantify plaque and atherosclerosis progression. Atherothrombosis research also includes studies of diet, exercise, diabetes, obesity, and other metabolic conditions related to atherothrombosis.

Heart Development and Structural Diseases Branch

The Heart Development and Structural Diseases Branch supports integrated basic and clinical research on

normal and abnormal cardiovascular development and the etiology, pathogenesis, prevention, diagnosis, and treatment of pediatric and adult structural heart disease. Research areas in heart development include normal and abnormal development, molecular and genetic etiology of cardiovascular malformations, cardiomyogenic differentiation of stem cells, and gene–environment interactions in the development of congenital heart disease. Structural disease research includes the investigation of congenital heart disease, from embryology through adulthood, and the associated exercise physiology and neurodevelopmental outcomes; valve disease; pediatric cardiomyopathy and heart transplantation; and pediatric cardiac inflammation and infection.

Heart Failure and Arrhythmias Branch

The Heart Failure and Arrhythmias Branch supports integrated basic and clinical research on normal and abnormal cardiac function to improve diagnosis, treatment, and prevention of heart failure and arrhythmias and to protect the myocardium and manage resuscitation. Heart failure research addresses the pathogenesis and treatment of heart failure and cardiomyopathies, including use of devices, medical treatments, and cell-based therapies. Arrhythmias research investigates the etiology of rare and common arrhythmias, sudden cardiac death, and arrhythmogenesis and explores the genetic and environmental bases of normal cardiac electrical activity. Myocardium protection research focuses on stunning and hibernation, ischemic/reperfusion injury, and preconditioning. Resuscitation research includes the study of whole-body oxygen deprivation; organ preservation; and cell, tissue, and organ protection during cardiac arrest and traumatic shock.

Prevention and Population Sciences Program

The Prevention and Population Sciences Program supports and provides leadership for population- and clinic-based research on the causes, prevention, and clinical care of cardiovascular, lung, and blood diseases and sleep disorders; it also supports research training and career development in these areas. Areas of research include epidemiological studies to describe disease and risk factor patterns in populations and identify risk factors for disease; clinical trials of interventions to prevent disease; genetic, behavioral, sociocultural, and environmental influences on disease risk and outcomes; and application of prevention and treatment strategies to determine ways to improve clinical care and public health.

The Program is organized into the three branches and three offices described below.

Clinical Applications and Prevention Branch

The Clinical Applications and Prevention Branch supports, designs, and conducts research on behavioral, environmental, clinical, and health care approaches to reduce the occurrence and consequences of CVD. Prevention research examines the effectiveness of interventions to slow or halt risk factor or disease development or progression. Interventions—many of which focus on high-risk individuals and populations—include medications, behavioral strategies, and environmental change. Studies to examine lifestyle, nutrition and exercise, psychological and sociocultural factors, and environmental and genetic influences are relevant to prevention and are supported. Clinical application research examines approaches to improve health care delivery and patient outcomes in clinical and community trials and observational studies.

Epidemiology Branch

The Epidemiology Branch supports, designs, and conducts research on the epidemiology of cardiovascular, lung, and blood diseases and sleep disorders. Studies are conducted to identify temporal trends and population patterns in the prevalence, incidence, morbidity, and mortality from the diseases and include single- and multicenter observational epidemiologic studies of development, progression, and treatment of cardiovascular, lung, and blood diseases and sleep disorders. Areas of emphasis include environmental, lifestyle, physiological, and genetic risk factors for disease and risk factor development including characterization of gene–gene and gene–environment interactions. Large cohorts of minority participants, such as Hispanics and blacks, have been assembled to explore health disparities in minorities. The Branch also distributes data from eligible NHLBI studies to researchers through a process that adheres to guidelines for the protection of participant privacy and confidentiality.

Women's Health Initiative Branch

The Women's Health Initiative Branch—in collaboration with the National Cancer Institute (NCI), the National Institute of Arthritis and Musculoskeletal and Skin Diseases (NIAMS), the National Institute on Aging (NIA), the National Institute of Neurological Disorders and Stroke (NINDS), and the Office of Research on Women's Health (ORWH)—supports clinical trials and

observational studies to improve the understanding of the causes and prevention of major diseases affecting the health of women. Studies focus on CVD, cancer, and fractures. Large multicenter observational studies seek to identify risk markers for disease or to better quantify known markers using questionnaires, clinical examinations, and laboratory data. The large and long-term multicenter clinical trials tested promising but unproven interventions—such as hormone therapy, diet, and supplements—to prevent major diseases and evaluate overall effects on health. Currently, the program is determining the long-term effects of prior hormone therapy on the cohort that participated in the clinical trials of hormone therapy. The Branch has established an infrastructure to support the use of data and blood samples from the studies by the scientific community.

The Women's Health Initiative Memory Study (WHIMS), an ancillary study to the WHI, was designed to test whether hormone therapy prevents the development and progression of dementia symptoms in postmenopausal women.

Office of Biostatistics Research

The Office of Biostatistics Research (OBR) provides statistical expertise to the Institute and performs diverse functions in planning, designing, implementing, and analyzing NHLBI-sponsored studies. Its primary responsibility is to provide objective, statistically sound, and medically relevant solutions to problems. The OBR is expected to provide a new and valid statistical solution when presented with a problem for which techniques are not yet available. Its methodological interests concern survival analysis; longitudinal data analysis; and efficient study designs, including the monitoring of ongoing clinical studies for efficacy and safety. The OBR has recently made contributions to statistical genetics and has extended its expertise to bioinformatics.

Office of Research Training and Career Development

The Office of Research Training and Career Development supports training and career development programs in cardiovascular research for individuals at all educational levels, from high school students to faculty. It collaborates with the scientific community and professional organizations to ensure that its programs meet the needs of young scientists from diverse backgrounds. Activities include institutional and individual research training programs and fellowships; diversity supplements to provide mentored experiences with established

research scientists; the Pathway to Independence Program, which allows recipients to bridge the gap between a career development award and a research award; and career development programs designed for clinical research.

Office of Special Projects

The Office of Special Projects represents the DCVS on NHLBI and NIH policy committees; oversees and works with Division leadership on selected activities of the DCVS clinical studies portfolio; fosters communication within DCVS by developing and coordinating Division-wide and Institute-wide interest groups on various topics; develops and implements specific cross-cutting projects; and provides expert consultation as needed for large-scale projects or initiative development.

Division of Lung Diseases

The DLD supports research on the causes, diagnosis, treatment, and prevention of lung diseases and sleep disorders. Research is funded through investigator- and Institute-initiated grants and contracts in such disease areas as asthma, bronchopulmonary dysplasia, COPD, CF, sleep-disordered breathing, critical care and acute lung injury, developmental biology and pediatric pulmonary diseases, immunologic and fibrotic pulmonary disease, rare lung disorders, pulmonary vascular disease, and pulmonary complications of AIDS and tuberculosis. SCCORs support collaborative studies on COPD and pulmonary vascular disease. A Centers Program supports research on advanced diagnostics and experimental therapeutics in lung diseases.

The Division also supports demonstration and dissemination projects to transfer basic research and clinical findings to health care professionals and patients, and training and career development programs for individuals interested in furthering their professional abilities in lung disease research. The DLD, through the National Center on Sleep Disorders Research, coordinates sleep research activities across the NIH, other Federal Agencies, and outside organizations.

The Division is organized into the three Branches described below.

Airway Biology and Disease Branch

The Airway Biology and Disease Branch supports basic and clinical research and research training in

asthma, COPD, CF, and airway function in health and disease. The Branch supports innovative genetics, genomics, and biotechnology programs to advance discovery of lung disease risk factors, mechanisms, and treatment. It also funds applied studies to develop new methods of lung imaging. Health education research and demonstration and education projects for the management of asthma and COPD are additional areas of focus.

Asthma research investigates the origins, pathogenesis, and management of asthma, including the role of immunologic and nonimmunologic events and inflammation in its pathogenesis; genetics of asthma and atopy; airway remodeling and repair in asthma; mechanisms of severe asthma; and regulation of mucous hypersecretion and mucous cell metaplasia.

Research on COPD and other diseases of the lung related to smoking or environmental exposures explores pathogenetic mechanisms involved in the development and progression of COPD, emphysema, and lung disease associated with alpha-1-antitrypsin deficiency; genetic determinants of lung disease; treatment of COPD; and health effects of air pollution.

Research on CF focuses on the function of the CF transmembrane conductance regulator and its role in lung disease. Areas of interest include airway epithelial ion transport, airway surface liquids, animal and cellular models for CF, signaling pathways in airway cells, regulation of mucin expression and secretion, development and clinical testing of treatments, and mechanisms underlying the infectious and inflammatory aspects of CF lung disease.

Lung Biology and Disease Branch

The Lung Biology and Disease Branch supports basic, translational, and clinical research and research training programs in pulmonary conditions associated with human immunodeficiency virus (HIV)/AIDS, tuberculosis, acute lung injury and critical care medicine, lung development and pediatric lung diseases, lung immunobiology and interstitial lung diseases, lymphangioleiomyomatosis, and lung cell and vascular biology. In addition, it supports the development of tuberculosis curricula for medical schools.

AIDS and tuberculosis research focuses on the pathogenesis and course of pulmonary manifestations of HIV infection and tuberculosis and host lung defenses against

them and HIV-associated opportunistic infections. Emphasis is on identifying and understanding the pathogenesis of lung complications associated with HIV infection and characterizing the lung microbiome in HIV-infected and HIV-uninfected individuals.

Research on acute lung injury and critical care medicine explores the pathogenesis, treatment, and prevention of acute lung injury and acute respiratory distress syndrome (ARDS). The Branch supports development of new diagnostic tools for detection of acute lung injury and development of an artificial lung and oversees clinical studies of therapies for ARDS, including the ARDS Network.

Research in developmental biology and pediatric pulmonary diseases investigates the regulation of lung development, growth, and repair and focuses on pediatric pulmonary diseases in infants and children, including bronchopulmonary dysplasia, congenital and acquired upper airway abnormalities, and persistent pulmonary hypertension of the newborn. Research also focuses on identifying and determining the cell fate of lung progenitor stem cells, understanding lung regeneration, and exploring cell-based therapy for lung injury and disease.

Research on immunology and fibrosis includes studies of interstitial pulmonary fibrosis, sarcoidosis, occupational and environmental lung diseases, and the role of immune response and inflammation in these diseases. The Branch also supports research on lung immunobiology, lung transplantation, and pathogenesis of lymphangioleiomyomatosis.

Lung cell and vascular biology research investigates lung cell biology and function and pulmonary vascular disease, including pulmonary arterial hypertension and pulmonary embolism diagnosis. Research focuses on pulmonary alveolar epithelial cells, vascular endothelial cells, and the lung surfactant system. The Branch also performs research on the regulation of barrier function of pulmonary endothelial cells and regulation of lung permeability.

National Center on Sleep Disorders Research

The National Center on Sleep Disorders Research (NCSDR) supports research, health education, and research training related to sleep-disordered breathing and the fundamental function of sleep and circadian rhythms. Specific areas of interest include neurobiology

of ventilatory control, respiratory rhythmogenesis, chemosensitivity, basic neurobiology of sleep-wake regulation, circadian-coupled cellular function, and effects of sleep deprivation. The NCSDR also stewards several forums, including the Sleep Disorders Research Advisory Board and the Trans-NIH Sleep Research Coordinating Committee, which facilitate the coordination of sleep research across the NIH and with other Federal Agencies and outside organizations. The Center participates in translation of new sleep research findings for dissemination to health care professionals and the public.

Division of Blood Diseases and Resources

The DBDR supports research and research training on the causes, diagnosis, treatment, and prevention of non-malignant blood diseases, including anemias, SCD, and thalassemia; premalignant processes, such as myelodysplasia and myeloproliferative disorders; hemophilia and other abnormalities of hemostasis and thrombosis; and immune dysfunction. It supports specialized centers that focus on research in basic and translational research in SCD.

The Division also supports research in transfusion medicine and blood banking, stem cell biology and disease, hematopoiesis, clinical cellular medicine, and blood supply adequacy and safety. It provides biospecimens and cellular resources to the scientific community.

The Division is organized into the three Branches described below.

Blood Diseases Branch

The Blood Diseases Branch supports research and research training in blood diseases, including SCD, thalassemia, Fanconi anemia, Diamond-Blackfan anemia, and other aplastic anemias and malaria. Additionally, it supports outcomes-related research. Research in SCD and thalassemia focuses on elucidating the etiology and pathophysiology of the diseases and improving disease treatment and management. Areas of emphasis include genetics, regulation of hemoglobin synthesis, iron chelation, development of drugs to increase fetal hemoglobin production, hematopoietic transplantation, and gene therapy. Basic and translational red cell research are also areas of interest.

Thrombosis and Hemostasis Branch

The Thrombosis and Hemostasis Branch supports research and research training in hemostasis,

thrombosis, and endothelial cell biology, including basic research, clinical studies, and technology development. Areas of interest include hemophilia; von Willebrand disease; and such immune disorders as idiopathic thrombocytopenic purpura, thrombotic thrombocytopenic purpura, and systemic lupus erythematosus. Research on bleeding disorders focuses on identifying effective treatments. Emerging areas of interest are gene transfer; clinical proteomics; glycomics; inflammation related to vascular injury from trauma and sepsis; thrombosis; stroke; coagulation activation; autoimmune disease; and thrombotic complications of obesity, diabetes, and cancer.

The Branch also supports research on the pathogenesis of arterial and venous thrombosis to improve the diagnosis, prevention, and treatment of thrombosis in heart attack, stroke, and peripheral vascular diseases. A major goal is to find additional platelet inhibitors, anticoagulants, and fibrinolytic agents to treat thrombotic and thromboembolic disorders with better specificity and fewer side effects than those currently used for treatment.

Transfusion Medicine and Cellular Therapeutics Branch

The Transfusion Medicine and Cellular Therapeutics Branch supports research and research training in transfusion medicine, stem cell biology and disease, hematopoiesis, clinical cellular medicine, and blood supply adequacy and safety. Research focuses on the use, safety, and availability of blood and blood components for transfusion and cellular therapies. Research areas include transmission of disease, noninfectious complications of transfusions, immunobiology, cell biology and disease, novel cell-based therapies, hematopoietic stem cell transplantation, and overall product availability. The Branch develops programs for basic and clinical research related to normal and abnormal cellular biology and pathology. It also collaborates with governmental, private sector, and international organizations to improve the safety and availability of the global supply of blood and blood components. The Branch also supports major NHLBI resource programs that provide cellular therapeutic products and biospecimens to the NHLBI scientific community.

Division of Intramural Research

The DIR conducts laboratory and clinical research in heart, vascular, lung, blood, and kidney diseases and

develops technology related to cardiovascular and pulmonary diseases. Areas of interest include the biologic basis of arteriosclerosis and its manifestations; pathophysiology of hypertensive vascular disease; functions of the lung; clinical and experimental studies on physiologic and pharmacologic aspects of heart, lung, and blood diseases; and a broad program of other basic research and technical developments related to them.

The DIR is organized into the four Centers and three Branches described below.

Biochemistry and Biophysics Center

The Biochemistry and Biophysics Center develops a global view of the molecular basis of structure–function relationships of proteins and biologically relevant molecules. It performs state-of-the-art nuclear magnetic resonance spectroscopy studies of protein structure and functional interactions, develops mathematical tools for generating theoretical models of protein structure–function relationships, elucidates the mechanisms of enzyme function, and investigates the relationship between protein structure–function and cell signaling pathways.

Cell Biology and Physiology Center

The Cell Biology and Physiology Center develops a global view of the mechanisms that regulate cellular function and physiology. It evaluates the mechanisms that control different molecular machines within the cytosol, including those involved in muscle contraction and cytosolic and membrane transport processes. The Center studies cellular signaling events associated with hormone action, cytosolic trafficking, and energy metabolism; investigates the role of cellular processes on function and adaptation in whole-animal model systems; and develops unique measuring devices for studying biochemical and physiological processes in intact cells, whole animals, and clinical situations.

Genetics and Development Biology Center

The Genetics and Development Biology Center develops a global view of the mechanisms that regulate cardiovascular development and the etiology of congenital heart anomalies and CVD. It evaluates the function of specific genes and transcription factors in the development of the heart and other tissues, develops techniques and approaches for gene delivery and gene therapy in model systems, and works toward a better understanding of

basic processes involved in regulating and interpreting the genetic code in development and disease.

Immunology Center

The Immunology Center develops a global view of the molecular basis of immune processes. It studies intracellular and signaling processes involved in activation of lymphocytes and mast cells, investigates mechanisms by which drugs and other agents result in allergic-autoimmune reactions, and relates the results to the development of new diagnostic and therapeutic approaches in humans.

Translational Medicine Branch

The Translational Medicine Branch conducts biomedical research directed at defining normal and abnormal biologic function at the molecular level. It develops diagnostic and therapeutic modalities for the treatment and understanding of CVD and implements mechanism-based clinical studies centered on innovative discoveries and observations from inside and outside the Branch.

Hematology Branch

The Hematology Branch conducts basic and clinical research on normal and abnormal hematopoiesis. Areas of interest include bone marrow failure, viral infections of hematopoietic cells, gene therapy of hematologic and malignant diseases, bone marrow transplantation, and mechanisms of immunologically mediated syndromes, such as graft-versus-host disease and autoimmune diseases.

Pulmonary and Vascular Medicine Branch

The Pulmonary and Vascular Medicine Branch conducts research on the lung, heart, and systemic vasculature directed at defining normal physiological function and novel mechanisms of disease at the molecular, biochemical, and functional levels. It conducts research on emerging diseases of the lung characterized by unknown etiology and molecular pathogenesis. Areas of interest include lung diseases in blacks, such as SCD and sarcoidosis; the role of nitric oxide, nitrite, gender, preconditioning, and mitochondrial function on the modulation of ischemia and reperfusion injury of the heart and lung; and translational study and drug development for therapeutic modulation of vascular, pulmonary, and cardiac cellular and molecular dysfunction in diseases of the lung and heart.

Division for the Application of Research Discoveries

The DARD supports efforts to advance the application of scientific discoveries for preventing, detecting, and treating cardiovascular, lung, blood, and sleep diseases and conditions to improve the health of all Americans. It focuses on translating scientific evidence into clinical guidelines for physicians to implement in their practice and into community health promotion or education programs for communities to disseminate to the public. The Division uses several channels of communications, including communities of practice, knowledge networks, social media, Web sites, conferences, and symposia. DARD programs reach out to people in high-risk, low-income communities to improve health and reduce health disparities. DARD activities promote communication and collaboration among researchers, clinical and public health practitioners, patients, and the general public. They also focus on identifying gaps in knowledge that can be addressed by future research.

The Division is organized into the two Branches described below.

Research Translation Branch

The Research Translation Branch interprets research findings into effective approaches for practice. The Branch synthesizes and organizes evidence around priority diseases or conditions and leads the effort to develop both evidence-based systematic literature reviews and guidelines for clinical practice. The Branch also develops clinical decision support systems and other innovative applications for use in clinical and public health practice settings, and it facilitates knowledge exchange opportunities for researchers and practitioners around issues of research applicability and relevance to practice. Branch activities also identify knowledge gaps to inform future research.

Enhanced Dissemination and Utilization Branch

The Enhanced Dissemination and Utilization Branch collects, synthesizes, and communicates evidence-based findings on the determinants of population health to maintain and improve the health of diverse populations and reduce health disparities in underserved groups. The Branch translates research into effective community health promotion programs, establishes effective partnerships to improve health and reduce health disparities, and builds communication among organizations and

communities to ensure their personal involvement in improving community health. Results are achieved by providing technical assistance and information resources to diverse audiences, including high-risk and underserved groups in a variety of community practice settings. The

Branch identifies appropriate health outcomes for assessing successful implementation and conducts evaluation activities to ensure continuous improvement and inform program planning.



3. Important Events

June 16, 1948. President Harry S. Truman signs the National Heart Act, creating the NHI in the Public Health Service (PHS), with the National Advisory Heart Council as its advisory body.

July 7, 1948. Dr. Paul Dudley White is selected to be “Executive Director of the National Advisory Heart Council and Chief Medical Advisor to the National Heart Institute” under section 4b of the National Heart Act.

August 1, 1948. The NHI is established as an institute of the NIH by Surgeon General Leonard A. Scheele. As legislated in the National Heart Act, the NHI assumes responsibility for heart research, training, and administration. Intramural research projects in CVD and gerontology conducted elsewhere in the NIH are transferred to the NHI. The Director of the NHI assumes all leadership for the total PHS heart program. Dr. Cassius J. Van Slyke is appointed as the first Director of the NHI.

August 29, 1948. Surgeon General Scheele announces the membership of the first National Advisory Heart Council. Varying terms of membership for the 16-member Council commence September 1.

September 8, 1948. The National Advisory Heart Council holds its first meeting.

January 1949. Cooperative Research Units are established at four institutions: the University of California, the University of Minnesota, Tulane University, and Massachusetts General Hospital. Pending completion of the NHI’s own research organization and facilities, the Units are jointly financed by the NIH and the institutions.

July 1, 1949. The NHI Intramural Research Program is established and organized on three general research levels consisting of three laboratory sections, five laboratory-clinical sections, and four clinical sections. The Heart Disease Epidemiology Study at Framingham, Massachusetts, is transferred from the Bureau of State Services, PHS, to the NHI.

January 18–20, 1950. The NHI and the American Heart Association jointly sponsor the first National Conference on Cardiovascular Diseases to summarize current knowledge and to make recommendations concerning further progress against heart and blood vessel diseases.

December 1, 1952. Dr. James Watt is appointed Director of the NHI, succeeding Dr. Van Slyke, who is appointed Associate Director of the NIH.

July 6, 1953. The Clinical Center admits its first patient for heart disease research.

July 1, 1957. The first members of the NHI Board of Scientific Counselors begin their terms. The Board was established in 1956 “to provide advice on matters of general policy, particularly from a long-range viewpoint, as they relate to the intramural research program.”

February 19, 1959. The American Heart Association and the NHI present a report to the Nation—*A Decade of Progress Against Cardiovascular Disease*.

April 21, 1961. The President’s Conference on Heart Disease and Cancer, whose participants on March 15 were requested by President John F. Kennedy to assist “in charting the Government’s further role in a national attack on these diseases,” convenes at the White House and submits its report.

September 11, 1961. Dr. Ralph E. Knutti is appointed Director of the NHI, succeeding Dr. Watt, who becomes head of international activities for the PHS.

December 30, 1963. February is designated as “American Heart Month” by a unanimous joint resolution of Congress with approval from President Lyndon B. Johnson.

November 22–24, 1964. The Second National Conference on Cardiovascular Diseases—cosponsored by the American Heart Association, the NHI, and the Heart Disease Control Program of the PHS—is held to evaluate progress since the 1950

Conference and to assess needs and goals for continued and accelerated growth against heart and blood vessel diseases.

December 9, 1964. The President's Commission on Heart Disease, Cancer, and Stroke—appointed by President Johnson on March 7, 1964—submits its report to “recommend steps that can be taken to reduce the burden and incidence of these diseases.”

August 1, 1965. Dr. William H. Stewart assumes the Directorship of the NHI upon Dr. Knutti’s retirement.

September 24, 1965. Dr. William H. Stewart, NHI Director, is named Surgeon General of the PHS.

October 6, 1965. In FY 1966, Supplemental Appropriations Act (P.L. 89–199) allocates funds to implement the recommendations of the President's Commission on Heart Disease, Cancer, and Stroke that are within existing legislative authorities. The NHI is given \$5.05 million for new clinical training programs, additional graduate training grants, cardiovascular clinical research centers on cerebrovascular disease and thrombotic and hemorrhagic disorders, and planning grants for future specialized cardiovascular centers.

March 8, 1966. Dr. Robert P. Grant succeeds Dr. Stewart as Director of the NHI. Dr. Grant serves until his death on August 15, 1966.

November 6, 1966. Dr. Donald S. Fredrickson is appointed Director of the NHI.

March 15, 1968. Dr. Theodore Cooper succeeds Dr. Fredrickson as Director of the NHI, the latter electing to return to research activities with the Institute.

October 16, 1968. Dr. Marshall W. Nirenberg is awarded a Nobel Prize in Physiology or Medicine for discovering the key to deciphering the genetic code. Dr. Nirenberg, chief of the NHI Laboratory of Biochemical Genetics, is the first Nobel Laureate at the NIH and the first Federal employee to receive a Nobel Prize.

October 26, 1968. The NHI receives the National Hemophilia Foundation's Research and Scientific Achievement Award for its “medical leadership . . . , tremendous stimulation and support of research activities directly related to the study and treatment of hemophilia.”

November 14, 1968. The 20th anniversary of the NHI is commemorated at the White House under the auspices of President Johnson and other distinguished guests.

August 12, 1969. A major NHI reorganization plan creates five program branches along disease category lines in extramural programs (arteriosclerotic disease, cardiac disease, pulmonary disease, hypertension and kidney diseases, and thrombotic and hemorrhagic diseases); a Therapeutic Evaluations Branch and an Epidemiology Branch under the Associate Director for Clinical Applications; and three offices in the Office of the Director (heart information, program planning, and administrative management).

November 10, 1969. The NHI is redesignated by the Secretary, Health, Education, and Welfare (HEW), as the National Heart and Lung Institute (NHLI), reflecting a broadening scope of its functions.

February 18, 1971. President Richard M. Nixon's Health Message to Congress identifies sickle cell anemia as a high-priority disease and calls for increased Federal expenditures. The Assistant Secretary for Health and Scientific Affairs, HEW, is assigned lead-Agency responsibility for coordination of the National Sickle Cell Disease Program at the NIH and NHLI.

June 1971. The Task Force on Arteriosclerosis, convened by Dr. Cooper, presents its report. Volume I addresses general aspects of the problem and presents the major conclusions and recommendations in nontechnical language. Volume II contains technical information on the state of knowledge and conclusions and recommendations in each of the following areas: atherogenesis, presymptomatic atherosclerosis, overt atherosclerosis, and rehabilitation.

May 16, 1972. The National Sickle Cell Anemia Control Act (P.L. 92–294) provides for a national diagnosis, control, treatment, and research program. The Act does not mention the NHLI but has special pertinence because the Institute has been designated to coordinate the National Sickle Cell Disease Program.

June 12, 1972. Elliot Richardson, Secretary, HEW, approves a nationwide program for high blood pressure information and education and appoints two committees to implement the program: the Hypertension Information and Education Advisory Committee, chaired by the Director, NIH, and the Interagency Working Group, chaired by the Director, NHLI. A High Blood Pressure

Information Center is established within the NHLI Office of Information to collect and disseminate public and professional information about the disease.

July 1972. The NHLI launches its National High Blood Pressure Education Program (NHBPEP), a program of patient and professional education that has as its goal to reduce death and disability related to high blood pressure.

July 14, 1972. Secretary Richardson approves reorganization of the NHLI, with the Institute elevated to Bureau status within the NIH and comprising seven division-level components: Office of the Director, Division of Heart and Vascular Diseases (DHVD), DLD, DBDR, DIR, Division of Technological Applications, and Division of Extramural Affairs (DEA).

September 19, 1972. The National Heart, Blood Vessel, Lung, and Blood Act of 1972 (P.L. 92-423) expands the authority of the Institute to advance the national attack on the diseases within its mandate. The act calls for intensified and coordinated Institute activities to be planned by the Director and reviewed by the National Heart and Lung Advisory Council.

July 24, 1973. The first Five-Year Plan for the National Heart, Blood Vessel, Lung, and Blood Program is transmitted to the President and to Congress.

December 17, 1973. The National Heart and Lung Advisory Council completes its First Annual Report on the National Program.

February 13, 1974. The Director of the NHLI forwards his First Annual Report on the National Program to the President for transmittal to Congress.

April 5, 1974. The Assistant Secretary for Health, HEW, authorizes release of the Report to the President by the President's Advisory Panel on Heart Disease. The report of the 20-member panel, chaired by Dr. John S. Millis, includes a survey of the problem of heart and blood vessel disorders and panel recommendations to reduce illness and death from them.

August 2, 1974. The Secretary, HEW, approves regulations governing the establishment, support, and operation of National Research and Demonstration Centers for heart, blood vessel, lung, and blood diseases, which implement section 415(b) of the PHS Act, as amended by the National Heart, Blood Vessel, Lung, and Blood Act of 1972: (1) to carry out basic and clinical research on

heart, blood vessel, lung, and blood diseases; (2) to provide demonstrations of advanced methods of prevention, diagnosis, and treatment; and (3) to supply a training source for scientists and physicians concerned with the diseases.

September 16, 1975. Dr. Robert I. Levy is appointed Director of the NHLI, succeeding Dr. Theodore Cooper, who was appointed Deputy Assistant Secretary for Health, HEW, on April 19, 1974.

June 25, 1976. Legislation amending the PHS Act (P.L. 94-278) changes the name of the NHLI to the National Heart, Lung, and Blood Institute (NHLBI) and provides for an expansion in blood-related activities within the Institute and throughout the National Heart, Blood Vessel, Lung, and Blood Program.

August 1, 1977. The Biomedical Research Extension Act of 1977 (P.L. 95-83) reauthorizes the programs of the NHLBI, with continued emphasis on both the national program and related prevention and dissemination activities.

February 1978. The NHLBI and the American Heart Association jointly celebrate their 30th anniversaries.

September 1979. The Task Force on Hypertension, established in September 1975 to assess the state of hypertension research, completes its in-depth survey and recommendations for improved prevention, treatment, and control in 14 major areas. The recommendations are intended to guide the NHLBI in its future efforts.

November 1979. The results of the Hypertension Detection and Follow-Up Program (HDFP), a major clinical trial started in 1971, provide evidence that tens of thousands of lives are being saved through treatment of mild hypertension and that perhaps thousands more could be saved annually if all people with mild hypertension were under treatment.

November 21, 1980. The Albert Lasker Special Public Health Award is presented to the NHLBI for its HDFP, "which stands alone among clinical studies in its profound potential benefit to millions of people."

December 17, 1980. The Health Programs Extension Act of 1980 (P.L. 96-538) reauthorizes the NHLBI, with continued emphasis on both the national program and related prevention programs.

September 8, 1981. The Working Group on Arteriosclerosis—convened in 1978 to assess present understanding, highlight unresolved problems, and emphasize opportunities for future research in arteriosclerosis—completes its report. Volume I presents conclusions and recommendations in nontechnical language. Volume II provides an in-depth substantive basis for the conclusions and recommendations contained in Volume I.

October 2, 1981. The Beta-Blocker Heart Attack Trial (BHAT) demonstrates benefits to those in the trial who received the drug propranolol compared with the control group.

July 6, 1982. Dr. Claude Lenfant is appointed Director of the NHLBI. He succeeds Dr. Levy.

September 1982. The results of the Multiple Risk Factor Intervention Trial are released. They support measures to reduce cigarette smoking and to lower blood cholesterol to prevent coronary heart disease (CHD) mortality but raise questions about optimal treatment of mild hypertension.

October 26, 1983. The Coronary Artery Surgery Study (CASS) results are released. They demonstrate that mildly symptomatic patients with coronary artery disease can safely defer coronary artery bypass surgery until symptoms worsen.

January 12, 1984. The results of the Lipid Research Clinics Coronary Primary Prevention Trial (LRC-CPPT) are released. They establish conclusively that reducing total blood cholesterol reduces the risk of CHD in men at increased risk because of elevated cholesterol levels. Each 1 percent decrease in cholesterol can be expected to reduce heart attack risk by 2 percent.

April–September 1984. The *Tenth Report of the Director, NHLBI*, commemorates the 10th anniversary of the passage of the National Heart, Blood Vessel, Lung, and Blood Act. The five-volume publication reviews 10 years of research progress and presents a 5-year research plan for the national program.

April 1984. The Division of Epidemiology and Clinical Applications (DECA) is created. It provides the Institute with a single focus on clinical trials; prevention, demonstration, and education programs; behavioral medicine; nutrition; epidemiology; and biometry. It also provides new opportunities to examine the interrelationships of cardiovascular, respiratory, and blood diseases.

November 1984. An NHLBI–NIH Clinical Center inter-Agency agreement for studies on the transmission of HIV from humans to chimpanzees leads to the first definitive evidence that the transmission is by blood transfusion.

April 1985. Results of Phase I of the Thrombolysis in Myocardial Infarction (TIMI) trial comparing streptokinase (SK) with recombinant tissue plasminogen activator (t-PA) are published. The new thrombolytic agent recombinant t-PA is approximately twice as effective as SK in opening thrombosed coronary arteries.

October 1985. The NHLBI Smoking Education Program is initiated to increase health care provider awareness about clinical opportunities for smoking cessation programs, techniques for use within health care settings, and resources for use within communities to expand and reinforce such efforts.

October 14, 1985. NHLBI-supported researchers Michael S. Brown and Joseph L. Goldstein are awarded the Nobel Prize in Physiology or Medicine for their discoveries concerning the regulation of cholesterol metabolism.

November 1985. The NHLBI inaugurates the National Cholesterol Education Program (NCEP) to increase awareness among health professionals and the public that elevated blood cholesterol is a cause of CHD and that reducing elevated blood cholesterol levels will contribute to the reduction of CHD.

June 1986. Results of the Prophylactic Penicillin Trial demonstrate the efficacy of prophylactic penicillin therapy in reducing the morbidity and mortality associated with pneumococcal infections in children with SCD.

September 18, 1986. The NHLBI sponsors events on the NIH campus in conjunction with the meeting of the X World Congress of Cardiology in Washington, DC. Activities include a special exhibit at the National Library of Medicine titled “American Contributions to Cardiovascular Medicine and Surgery” and two symposia—“New Dimensions in Cardiovascular Disease Research” and “Cardiovascular Nursing and Nursing Research.”

December 17, 1986. The citizens of Framingham, Massachusetts, are presented a tribute by the Assistant Secretary, HHS, for their participation in the Framingham Heart Study over the past 40 years.

September 1987. The NHLBI commemorates the centennial of the NIH and the 40th anniversary of the Institute's inception. Two publications prepared for the Institute's anniversary—*Forty Years of Achievement in Heart, Lung, and Blood Research* and *A Salute to the Past: A History of the National Heart, Lung, and Blood Institute*—document significant Institute contributions to research and summarize recollections about the Institute's 40-year history.

October 1987. The National Blood Resource Education Program is established to ensure an adequate supply of safe blood and blood components to meet the Nation's needs and to ensure that blood and blood components are transfused only when therapeutically appropriate.

April 1988. The NHLBI initiates its Minority Research Supplements program to provide supplemental funds to ongoing research grants for support of minority investigators added to research teams.

September 1988. AIDS research is added to the National Heart, Blood Vessel, Lung, and Blood Diseases and Blood Resources Program. It is the first area of research to be added since the Program was established in 1973.

September 1988. The NHLBI funds the first of its new Programs of Excellence in Molecular Biology, designed to foster the study of the organization, modification, and expression of the genome in areas of importance to the Institute and to encourage investigators to become skilled in the experimental strategies and techniques of modern molecular biology.

September 1988. The Strong Heart Study is initiated. It focuses on CVD morbidity and mortality rates and distribution of CVD risk factors in three geographically diverse American Indian groups.

October 1988. The National Marrow Donor Program is transferred from the Department of the Navy to the NHLBI. The Program, which serves as a focal point for bone marrow research, includes a national registry of volunteers who have offered to donate marrow for transplant to patients not having suitably matched relatives.

March 1989. The NHLBI initiates a National Asthma Education Program to raise awareness of asthma as a serious chronic disease and to promote more effective

management of asthma through patient and professional education.

May 1989. The NHLBI Minority Access to Research Careers (MARC) Summer Research Training Program is initiated to provide an opportunity for MARC Honors Scholars to work with researchers in the NHLBI intramural laboratories.

September 14, 1990. The first human gene therapy protocol in history is undertaken at the NIH. A team of scientists—led by W. French Anderson, NHLBI, and R. Michael Blaese, NCI—insert a normal gene into a patient's cells to compensate for a defective gene that left the patient's cells unable to produce an enzyme essential to the functioning of the body's immune system.

January 1991. The NHLBI Obesity Education Initiative (OEI) begins. Its objective is to make a concerted effort to educate the public and health professionals about obesity as an independent risk factor for CVD and its relationship to other risk factors, such as high blood pressure and high blood cholesterol.

February 1991. The expert panel of the National Asthma Education Program releases its report, *Guidelines for Diagnosis and Management of Asthma*, to educate physicians and other health care providers in asthma management.

April 8–10, 1991. The First National Conference on Cholesterol and Blood Pressure Control is attended by more than 1,800 health professionals.

May 1991. The Task Force on Hypertension, established in November 1989 to assess the state of hypertension research and to develop a plan for future NHLBI funding, presents its conclusions. The report outlines a set of scientific priorities and develops a comprehensive plan for support over the next several years.

June 11, 1991. The NHLBI initiates a National Heart Attack Alert Program (NHAAP) to reduce premature morbidity and mortality from acute myocardial infarction (AMI) and sudden death. The Program emphasizes rapid disease identification and treatment.

July 1991. Results of the Systolic Hypertension in the Elderly Program (SHEP) demonstrate that low-dose pharmacologic therapy of isolated systolic hypertension in those older than 60 years of age significantly reduces stroke and myocardial infarction.

August 1991. Results of the Studies of Left Ventricular Dysfunction (SOLVD) are released. They demonstrate that use of the angiotensin-converting enzyme (ACE) inhibitor enalapril causes a significant reduction in mortality and hospitalization for congestive heart failure in patients with symptomatic heart failure.

August 1991. The NHLBI sponsors “Physical Activity and Cardiovascular Health: Special Emphasis on Women and Youth,” the first national workshop to assess the current knowledge in the field and to develop scientific priorities and plans for support. Recommendations from the Working Groups are published in the supplemental issue of *Medicine and Science in Sports and Exercise*.

March 1992. The *International Consensus Report on Diagnosis and Management of Asthma* is released. It is to be used by asthma specialists and medical opinion leaders to provide a framework for discussion of asthma management pertinent to their respective countries.

March 1992. Results of the Trials of Hypertension Prevention Phase I are published. They demonstrate that both weight loss and reduction of dietary salt reduce blood pressure in adults with high-normal diastolic blood pressure and may reduce the incidence of primary hypertension.

June 26–27, 1992. The Fourth National Minority Forum on Cardiovascular Health, Pulmonary Disorders, and Blood Resources is attended by nearly 600 individuals.

October 11–13, 1992. The First National Conference on Asthma Management is attended by more than 900 individuals.

October 30, 1992. A celebration of the 20th anniversary of the NHBPEP is held in conjunction with the NHBPEP Coordinating Committee meeting. The *Fifth Report of the Joint National Committee on the Detection, Evaluation, and Treatment of High Blood Pressure* (JNC V) and the *NHBPEP Working Group Report on the Primary Prevention of Hypertension* are released.

June 10, 1993. The NIH Revitalization Act of 1993 (P.L. 103–43) establishes the NCSDR within the NHLBI.

June 15, 1993. The *Second Report of the Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults* (ATP II) is released to the

public at a press conference held in conjunction with the NCEP Coordinating Committee meeting.

January 30, 1995. Results of the Multicenter Study of Hydroxyurea (MSH) are released through a clinical alert. They demonstrate that hydroxyurea reduced the number of painful episodes by 50 percent in severely affected adults with SCD. This is the first effective treatment for adult patients with this disorder.

September 1995. The NHLBI funds a new Program of Specialized Centers of Research in Hematopoietic Stem Cell Biology, which is designed to advance our knowledge of stem cell biology and enhance our ability to achieve successful stem cell therapy to cure genetic and acquired diseases.

September 21, 1995. Results of the Bypass Angioplasty Revascularization Investigation are released through a clinical alert. They demonstrate that patients on drug treatment for diabetes who had blockages in two or more coronary arteries and were treated with coronary artery bypass graft (CABG) surgery had, at 5 years, a death rate markedly lower than that of similar patients treated with angioplasty. The clinical alert recommends CABG over standard angioplasty for patients on drug therapy for diabetes who have multiple coronary blockages and are first-time candidates for either procedure.

November 5–6, 1995. The first Conference on Socioeconomic Status (SES) and Cardiovascular Health and Disease is held to determine future opportunities and needs for research on SES factors and their relationships with cardiovascular health and disease.

December 4–5, 1995. A celebration of the 10th anniversary of the NCEP is held in conjunction with the NCEP Coordinating Committee meeting. Results of the 1995 Cholesterol Awareness Surveys of physicians and the public are released.

May 1996. The NHLBI announces results from the Framingham Heart Study that conclude earlier and more aggressive treatment of hypertension is vital to preventing congestive heart failure. The Treatment of Mild Hypertension Study (TOMHS) demonstrates that life-style changes—such as weight loss, a healthy eating plan, and physical activity—are crucial for reducing blood lipids in those treated for Stage I hypertension.

September 1996. Findings from the Asthma Clinical Research Network (ACRN) show that for people with

asthma, taking an inhaled beta-agonist at regularly scheduled times is safe but provides no greater benefit than taking the medication only when asthma symptoms occur. The recommendation to physicians who treat patients with mild asthma is to prescribe inhaled beta-agonists only on an as-needed basis.

November 13, 1996. The NHLBI releases findings from two studies, Dietary Approaches to Stop Hypertension (DASH) Trial and Trial of Nonpharmacologic Intervention in the Elderly (TONE). The DASH Trial demonstrates that a diet low in fat and high in vegetables, fruits, fiber, and low-fat dairy products significantly and quickly lowers blood pressure. The TONE shows that weight loss and reduction of dietary sodium safely reduce the need for antihypertensive medication in older patients while keeping their blood pressure under control.

January 1997. Definitive results from the Pathobiological Determinants of Atherosclerosis in Youth (PDAY) program are published. They show that atherosclerosis develops before age 20 and that the following risk factors affect the progression of atherosclerosis equally in women and men, regardless of race: low high-density lipoprotein (HDL) cholesterol, high low-density lipoprotein (LDL) cholesterol, and cigarette smoking.

February 24, 1997. The National Asthma Education and Prevention Program (NAEPP) releases the *Expert Panel Report 2, Guidelines for the Diagnosis and Management of Asthma* to the public at a press conference held in conjunction with a meeting of the American Academy of Allergy, Asthma, and Immunology in San Francisco.

May 8, 1997. Results of the Antiarrhythmic Versus Implantable Defibrillator (AVID) clinical trial are presented. They show that an implantable cardiac defibrillator reduces mortality compared to pharmacologic therapy in patients at high risk for sudden cardiac death.

September 1997. The Stroke Prevention Trial in Sickle Cell Anemia (STOP) is terminated early because prophylactic transfusion resulted in a 90 percent relative decrease in the stroke rate among children 2 to 16 years old.

September 1997. The Institute's National Sickle Cell Disease Program celebrates its 25th anniversary.

October 1997. The NHLBI commemorates the 50th anniversary of the Institute's inception. A publication prepared for the Institute's anniversary—*Vital Signs: Discoveries in Diseases of the Heart, Lungs, and Blood*—documents the remarkable research advances of the past 50 years.

October 1, 1997. The WHI, initiated in 1991, is transferred to the NHLBI.

November 6, 1997. The *Sixth Report of the Joint National Committee on the Prevention, Detection, Evaluation, and Treatment of High Blood Pressure* (JNC VI) is released at a press conference held in conjunction with the 25th anniversary meeting and celebration of the NHBPEP Coordinating Committee.

December 1997. Findings from the Trial To Reduce Alloimmunization to Platelets (TRAP) demonstrate that leucocyte reduction by filtration or ultraviolet B irradiation of platelets—both methods are equally effective—decreases development of lymphocytotoxic antibodies and alloimmune platelet refractoriness.

February 1998. The Task Force on Behavioral Research in Cardiovascular, Lung, and Blood Health and Disease—established in November 1995 to develop a plan for future NHLBI biobehavioral research in cardiovascular, lung, and blood diseases and sleep disorders—presents its recommendations. The report outlines a set of scientific priorities and develops a comprehensive plan for support over the next several years.

February 19–21, 1998. The NHLBI and cosponsors—California CVD Prevention Coalition; California Department of Health Services; CVD Outreach, Resources, and Epidemiology Program; and the University of California, San Francisco—hold Cardiovascular Health: Coming Together for the 21st Century, A National Conference, in San Francisco.

March 16, 1998. A special symposium is held at the annual meeting of the American Academy of Asthma, Allergy, and Immunology to celebrate 50 years of NHLBI-supported science.

June 17, 1998. The NHLBI, in cooperation with the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK), releases *Clinical Guidelines on the Identification, Treatment, and Evaluation of Overweight and Obesity in Adults: Evidence Report*.

December 11, 1998. World Asthma Day is established on this date. The NAEPP launches the Asthma Management Model System, an innovative Web-based information management tool.

March 1999. The ARDS Network Study of Ventilator Management in ARDS is stopped early so that critical care specialists can be alerted to the results. The study demonstrated that approximately 25 percent fewer deaths occurred among intensive care patients with ARDS receiving small, rather than large, breaths of air from a mechanical ventilator.

March 22, 1999. The NAEPP holds its 10th anniversary meeting and celebration to recognize a decade of progress and a continued commitment to the future.

August 1999. Results of the Early Revascularization for Cardiogenic Shock are released. They show improved survival at 6 months in patients treated with balloon angioplasty or coronary bypass surgery compared with patients who receive intensive medical care to stabilize their condition.

September 27–29, 1999. The NHLBI sponsors the National Conference on Cardiovascular Disease Prevention: Meeting the Healthy People 2010 Objectives for Cardiovascular Health.

November 2, 1999. The NAEPP convenes a Workshop on Strengthening Asthma Coalitions: Thinking Globally, Acting Locally to gather information from coalition representatives on ways the NAEPP could support their efforts.

November 2–3, 1999. The NHLBI sponsors a Workshop on Research Training and Career Development.

March 8, 2000. A part of the Antihypertensive and Lipid-Lowering Treatment To Prevent Heart Attack Trial (ALLHAT) is terminated early because one of the tested drugs, an alpha-adrenergic blocker, was found to be less effective than the more traditional diuretic in reducing some forms of CVD.

March 29, 2000. The NHLBI launches the Web-based Healthy People 2010 Gateway to provide information and resources on cardiovascular health, asthma, sleep, and minority populations.

April 25, 2000. The NHLBI sponsors a special expert meeting, Scientific Frontiers in Cardiothoracic Surgery, to discuss the future of cardiothoracic research.

September 2000. NHLBI-supported investigators identify a gene for primary pulmonary hypertension.

October 2000. Results from the Childhood Asthma Management Program (CAMP) demonstrate that inhaled corticosteroids are safe and effective for long-term treatment of children with mild-to-moderate asthma.

January 2001. Results of the DASH-Sodium Trial are released. They show that dietary sodium reduction substantially lowers blood pressure in persons with high blood pressure; the greatest effect occurs when sodium reduction is combined with the DASH diet.

February 2001. The NHLBI launches a sleep education program for children, using star sleeper Garfield the Cat.

February 1, 2001. The NHLBI—along with the HHS Office of Disease Prevention and Health Promotion, the Office of the Surgeon General, the Centers for Disease Control and Prevention (CDC), the NINDS, and the American Heart Association—signs a memorandum of understanding to focus and coordinate their efforts to meet the Healthy People 2010 objectives on cardiovascular health.

March 26–27, 2001. A strategy development workshop, “Women’s Heart Health: Developing a National Health Education Action Plan,” is held to develop an agenda for the NHLBI’s new heart health education effort directed at women.

April 2001. The NHLBI releases the international guidelines for diagnosis, management, and prevention of COPD.

April 2001. NHLBI-supported investigators identify genes that regulate human cholesterol levels.

May 2001. The NHLBI releases the NCEP’s *Third Report of the Expert Panel on Detection, Evaluation, and Treatment of High Blood Cholesterol in Adults* (ATP III).

June 2001. NHLBI-supported investigators find that human heart muscle cells regenerate after a heart attack.

July 2001. A self-contained artificial heart is implanted in a patient for the first time.

August 2001. Early results from the National Emphysema Treatment Trial (NETT) identify characteristics of patients at high risk for death following lung volume reduction surgery.

August 2001. Scientists from the NHLBI SCOR program at Yale University identify two genes responsible for pseudohypoaldosteronism type II, a rare Mendelian form of high blood pressure. These genes encode for protein kinases involved in a previously unknown pathway and may provide new targets for therapy.

September 10, 2001. The NHLBI, along with the American Heart Association and other partners, launches “Act in Time to Heart Attack Signs,” a national campaign to increase awareness of the signs of heart attack and the need for a fast response.

October 2001. NHLBI-supported scientists report that the drug, infliximab, increases risk of TB reactivation and dissemination. The drug is used to treat refractory rheumatoid arthritis and Crohn’s disease and is proposed as a treatment for several chronic lung diseases.

November 2001. Results of the Randomized Evaluation of Mechanical Assistance for the Treatment of Chronic Heart Failure Trial demonstrate that using an implanted left ventricular assist device can prolong survival and improve quality of life in severely ill patients who are not candidates for heart transplantation.

December 2001. For the first time, scientists correct SCD in mice using gene therapy.

April 10, 2002. The World Hypertension League (WHL) and the NHLBI hold an international symposium; subsequently they prepare an action plan at the WHL Council Conference to control hypertension and obesity.

April 11–13, 2002. The NHLBI and cosponsors—the HHS Office of Disease Prevention and Health Promotion, the CDC, the American Heart Association, the Centers for Medicare & Medicaid Services, and the Health Resources and Services Administration—hold a national conference, “Cardiovascular Health for All: Meeting the Challenge of Healthy People 2010.”

June 2002. The NAEPP issues an update of selected topics in the *Guidelines for the Diagnosis and Management of Asthma*.

June 2002. The fourth edition of *The Management of Sickle Cell Disease*, which describes the current approach to counseling SCD patients and managing many of the medical complications of SCD, is issued to coincide with the 30th anniversary of the NHLBI Sickle Cell Program.

July 9, 2002. The NHLBI stops early the trial of the estrogen plus progestin component of the WHI due to increased breast cancer risk and lack of overall benefits. The multicenter trial also found increases in CHD, stroke, and pulmonary embolism in participants on estrogen plus progestin compared to women taking placebo pills.

August 2002. NHLBI-supported scientists identify a gene variant that is associated with arrhythmia in blacks.

December 4, 2002. Results of the Atrial Fibrillation Follow-Up Investigation of Rhythm Management Trial (AFFIRM) indicate that rate control rather than rhythm control may be the preferred approach for patients with atrial fibrillation. The rate control strategy involves the use of less expensive drugs and results in fewer hospitalizations.

December 17, 2002. Results of the ALLHAT, the largest hypertension clinical trial ever conducted, show that less expensive traditional diuretics are at least as good as newer medicines (calcium channel blocker and ACE inhibitors) in treating high blood pressure and preventing some forms of heart disease.

January 23, 2002. An NHLBI-supported study demonstrates that magnetic resonance imaging can be used to detect heart attacks faster and more accurately than traditional methods in patients who arrive at the emergency room with chest pain.

February 24, 2002. The Prevention of Recurrent Venous Thromboembolism Trial is stopped early because treatment with low-dose warfarin to prevent recurrence of deep vein thrombosis and pulmonary embolism was so beneficial.

April 2003. Results of the MSH Patients’ Follow-Up Study show that the adult patients who took hydroxyurea over a 9-year period experienced a 40 percent reduction in deaths. Survival was related to fetal hemoglobin levels and frequency of vaso-occlusive events.

April 23, 2003. Results of the PREMIER trial of behavioral lifestyle interventions for blood pressure control show that individuals with prehypertension or stage I hypertension can lower their blood pressure by making multiple lifestyle changes.

May 14, 2003. The *Seventh Report of the Joint National Committee on the Prevention, Detection,*

Evaluation, and Treatment of High Blood Pressure (JNC VII) is released.

May 22, 2003. The NETT finds that lung volume reduction surgery (LVRS) benefits emphysema patients with certain clinical characteristics. The findings will be useful in the determination of Medicare coverage policy.

July 2003. The NHLBI and Gen-Probe Corporation succeed in developing a test to screen donated blood for the West Nile Virus.

August 2003. The NHLBI establishes a partnership with the Canadian Institutes of Health Research (CIHR) to advance research on cardiovascular, respiratory, and blood diseases.

November 2003. The Public Access Defibrillation Trial demonstrates that use of an automated external defibrillator and CPR by trained community volunteers can increase survival for victims of sudden cardiac arrest.

March 2004. The NIH stops the estrogen-alone component of the WHI early due to the increased risk of stroke and deep vein thrombosis. Estrogen does not appear to affect heart disease.

March 2004. Preliminary results of the Sudden Cardiac Death in Heart Failure Trial demonstrate that an implantable cardiac defibrillator can reduce death in heart failure patients.

July 2004. The NHLBI releases an update to the 2001 NCEP ATP III guidelines on the treatment of high blood cholesterol in adults.

August 2004. The NHBPEP Working Group on High Blood Pressure in Children and Adolescents releases the *Fourth Report on the Diagnosis, Evaluation, and Treatment of High Blood Pressure in Children and Adolescents*.

August 2004. An NHLBI-funded study shows that nucleic acid amplification testing for HIV-1 and hepatitis C virus (HCV) further safeguards the Nation's blood supply.

October 2004. Results from a new study of adults with mild asthma by researchers participating in the ACRN demonstrate that genes affect patient response, over time, to daily doses of inhaled albuterol, a drug used for relief of acute asthma symptoms. A few weeks of its regular use improves overall asthma control in

individuals with one form of the gene, but stopping all use of albuterol eventually improves asthma control in those with another form of the gene. The findings could lead to better ways to individualize asthma therapy.

November 2004. Results of the Prevention of Events With Angiotensin Converting Enzyme Inhibition (PEACE) demonstrate that many heart disease patients who are already receiving state-of-the-art therapy do not gain extra cardiovascular protection from ACE inhibitors.

December 2004. The NHLBI stops early the Stroke Prevention in Sickle Cell Anemia Trial II (STOP II) so that physicians who treat children with sickle cell anemia can be alerted to its findings. STOP II, which is a study to determine whether children with sickle cell anemia and at high risk for stroke could at some point safely stop receiving the periodic blood transfusions that prevent strokes, shows that children revert to high risk for stroke when transfusions are stopped.

January 2005. The NHLBI issues new guidelines for managing asthma during pregnancy.

January 2005. Results from Sudden Cardiac Death in Heart Failure (SCD-HeFT) show that patients with class II or class III heart failure and left ventricular ejection fraction of 35 percent or less have improved survival with implantable cardiac defibrillators. There is no benefit with amiodarone.

January 26, 2005. Dr. Elizabeth G. Nabel is appointed Director of the NHLBI. She succeeds Dr. Claude Lenfant.

February 2005. NHLBI-supported scientists identify two genetic mutations common in individuals of African descent that are associated with a 40 percent reduction in LDL cholesterol.

June 1, 2005. HHS Secretary Mike Leavitt announces the launch of We Can!, Ways to Enhance Children's Activity & Nutrition, a national education program from the NIH to prevent overweight and obesity among youth ages 8–13 years.

February 15, 2006. Results from the WHI Calcium and Vitamin D Trial show that calcium and vitamin D supplements in healthy postmenopausal women provide a modest improvement in bone mass preservation and prevent hip fractures in certain groups, including older women, but do not prevent other types of fractures or colorectal cancer.

May 10, 2006. Results from the Childhood Asthma Research and Education (CARE) Network show that daily treatment with inhaled corticosteroids can reduce breathing problems in preschool-aged children at high risk for asthma, but does not prevent them from developing persistent asthma.

May 31, 2006. The Prospective Investigation of Pulmonary Embolism Diagnosis (PIOPED) II finds that the ability to diagnose pulmonary embolism is improved when a commonly used imaging test of the chest to detect potentially deadly blood clots in the lung is complemented by an extension of the scan to the legs—where the clots typically originate—or by a standard clinical assessment.

June 6, 2006. Results from the Should We Emergently Revascularize Occluded Coronaries for Cardiogenic Shock (SHOCK) trial show that treating heart attack patients who have a life-threatening complication called cardiogenic shock with emergency angioplasty or bypass surgery greatly improves their long-term survival.

July 18, 2006. NHLBI scientists find that a hormone called brain natriuretic peptide or BNP, which can be detected in a simple blood test, can identify patients with SCD who have developed a life-threatening complication called pulmonary hypertension. The hormone is also a predictor of death in adult sickle cell patients.

July 26, 2006. Results from two randomized clinical trials demonstrate that inhaled nitric oxide administered within the first few weeks of life helps prevent chronic lung disease in some low birthweight premature infants. Moreover, when administered within 48 hours after birth, it appears to protect some premature newborns from brain injury.

September 19, 2006. The NHLBI launches a peripheral artery disease awareness and education campaign, “Stay in Circulation: Take Steps To Learn About P.A.D.” (peripheral artery disease).

January 18, 2007. The NHLBI launches the Learn More Breathe Better campaign to increase COPD awareness among primary care physicians and the public.

August 29, 2007. The NAEPP issues the *Expert Panel Report 3: Guidelines for the Diagnosis and Management of Asthma—Full Report 2007*, an update of the latest scientific evidence and recommendations for clinical practice on asthma care.

October 1, 2007. The NHLBI launches an open access dataset for researchers worldwide. Known as SNP Health Association Resource (SHARE), the Web-based dataset will enable qualified researchers to access data from large population-based studies, starting with the landmark Framingham Heart Study. It is expected to accelerate discoveries linking genes and health, thereby advancing understanding of the causes and prevention of CVD and other disorders.

October 8, 2007. Mario Capecchi and Oliver Smithies, who are researchers supported by the NHLBI, are awarded the Nobel Prize in Physiology or Medicine for their creation of a gene-targeting technique that allows scientists to create transgenic mice that are genetically modified to develop human diseases.

December 3, 2007. The NHLBI announces a new strategic plan to guide its next decade of research, training, and education to reduce the national burden of cardiovascular, lung, and blood diseases and sleep disorders.

December 10, 2007. Results of the Occluded Artery Trial (OAT) are incorporated into practice guidelines: The American College of Cardiology/American Heart Association’s *2007 Focused Update of the 2004 Guidelines for the Management of Patients With ST-Elevation Myocardial Infarction*. The guidelines discourage percutaneous coronary intervention of a totally occluded artery late in the course of myocardial infarction in the absence of symptoms if patients are stable and do not have evidence of severe ischemia.

January 28, 2008. Results from the ALLHAT demonstrate that in people—especially blacks—who have high blood pressure as part of metabolic syndrome, diuretics offer greater protection against CVD, including heart failure, and are at least as effective for lowering blood pressure as newer, more expensive medications.

February 2008. The NHLBI stops one treatment arm of the Action to Control Cardiovascular Risk in Diabetes (ACCORD) clinical trial of adults who have type 2 diabetes at high risk for heart attack and stroke after a review of available data showed that participants following a medical strategy to lower blood glucose below current recommendations to near-normal levels increased the risk of death compared with a standard treatment strategy. All participants now follow a medical strategy to reach the standard blood sugar levels while the lipid and blood pressure components of the study continue.

February 2008. An independent panel convened by the NIH concludes that the use of hydroxyurea for treating SCD should be increased among adolescents and adults who have the disease.

February 29, 2008. The NHLBI issues the first U.S. guidelines for the diagnosis and management of von Willebrand Disease, the most common inherited bleeding disorder.

March 2008. The NHLBI announces a comprehensive restructuring of its SCD research program to take advantage of new scientific opportunities and make SCD resources more widely available.

March 4, 2008. The WHI Follow-up Study confirms that the health risks of long-term combination hormone therapy outweigh the benefits for postmenopausal women. Researchers report that about 3 years after women stopped taking combination hormone therapy, many of the health effects of hormones—such as increased risk of heart disease—are diminished but overall risks of stroke, blood clots, and cancer remain high.

March 5, 2008. Scientists report that they have identified the variants of the gene VKORC1 that determine a patient's initial response to treatment with the blood-thinning (anticoagulant) drug warfarin. The finding is expected to enhance the ability of physicians to tailor the dosage of warfarin for individual patients.

April 2008. NHLBI-supported researchers identify gene variants associated with increased susceptibility to asthma and reduced lung function in three study populations. Risk for developing asthma is linked to variants in a gene called CHI3L1, which can be measured by checking levels of an inherited blood protein that it regulates.

April 8, 2008. Results from the Stop Atherosclerosis in Native Diabetic Study (SANDS) show that aggressively lowering cholesterol and blood pressure levels below current targets in adults with type 2 diabetes may help to prevent, and possibly reverse, hardening of the arteries.

April 14, 2008. The NHLBI, along with the NCI and National Institute of General Medical Sciences (NIGMS), signs a letter of intent with the Center for Genomic Medicine in Japan to create a Global Alliance for Pharmacogenomics to identify genetic factors that contribute to individual responses to medicines, including rare and dangerous side effects. Research results

will eventually allow physicians to ensure the safety and optimize the effectiveness of drugs for each patient.

August 18, 2008. The NHLBI launches an educational Web site, "Children and Clinical Studies," which features documentary videos, text, and graphics designed to promote a better understanding of research in children for health care professionals and the public.

September 15, 2008. The Surgeon General's *Call to Action To Prevent Deep Vein Thrombosis and Pulmonary Embolism* is released. The *Call to Action*, which urges a coordinated, multifaceted plan to reduce the number of cases of deep vein thrombosis and pulmonary embolism nationwide, resulted from a Surgeon General's Workshop on Deep Vein Thrombosis co-sponsored by the NHLBI.

September 25, 2008. Researchers announce that they have developed a genetically altered animal model for CF that closely matches the characteristics of the disease in humans.

October 6, 2008. NIH scientists show that tipifarnib, an experimental anticancer drug, can prevent, and even reverse, potentially fatal cardiovascular damage in a mouse model of progeria (rare genetic disorder that causes the most dramatic form of human premature aging).

December 15, 2008. The NHLBI expands its open-access dataset of genetic and clinical data to include information collected from three NHLBI-funded asthma research networks: ACRN, CAMP, and CARE.

December 19, 2008. Researchers identify a gene that directly affects the production of a form of hemoglobin that is instrumental in modifying the severity of SCD and thalassemia.

March 29, 2009. Results from the Surgical Treatment for Ischemic Heart Failure (STICH) study show that surgery to reshape the scarred left ventricle, the main pumping chamber of the heart, often performed in conjunction with coronary bypass surgery, fails to reduce deaths and hospitalizations in heart failure patients and does not improve quality of life compared with bypass alone.

June 5, 2009. Results from the Bypass Angioplasty Revascularization in Type 2 Diabetics (BARI 2D) study in patients with diabetes and stable coronary artery disease indicate that while revascularization can be delayed for many patients receiving optimal medical therapy, patients with extensive coronary artery disease do better with prompt bypass surgery than with medical therapy alone.

June 10, 2009. The NHLBI joins with UnitedHealth Group's Chronic Disease Initiative to launch a worldwide network of research and training centers to build institutional and community capacity to prevent and control chronic diseases globally.

July 28, 2009. The NHLBI stops the Sildenafil for Sickle Cell Disease-Associated Pulmonary Hypertension study after an interim review of the safety data shows that participants who are taking sildenafil are significantly more likely to have serious medical problems (e.g., severe pain called sickle cell crises) compared with participants on placebo.

August 16, 2009. Results from the Exome Project demonstrate the feasibility and value of isolating and sequencing all exons for identifying relatively rare genetic variants that may cause or contribute to disease. By focusing on the exome, important information about an individual can be obtained at a much lower cost than sequencing a person's entire genome.

August 19, 2009. Results from Sleep Heart Health Study show that moderate to severe obstructive sleep apnea is associated with an increased risk of death in middle-aged adults, especially men.

October 2009. The Division of Cardiovascular Sciences is created by combining two previously existing divisions, the Division of Cardiovascular Diseases and the Division of Prevention and Population Sciences, so that the administrative structure better matches the dynamic interaction that exists among basic, clinical, and population sciences.

December 9, 2009. Scientists, using a modified blood adult stem-cell transplant regimen, reverse SCD in 9 of 10 adults who had been severely affected by the disease.

May 2010. The NHLBI launches the National Asthma Control Initiative to improve asthma control in patients by bringing asthma care in line with evidence-based recommendations from the *Expert Panel Report 3—Guidelines for the Diagnosis and Management of Asthma* and its companion document, *Guidelines Implementation Panel Report—Partners Putting Guidelines Into Action*.

October 20, 2010. Follow-up findings from the WHI study of hormone therapy in postmenopausal women show that, in addition to having a higher incidence of breast cancer, the group treated with estrogen plus progestin had nearly double the rate of mortality from breast cancer than the placebo group 5 years after the study drug was discontinued.

April 7, 2011. Results from the STICH study show that adding bypass surgery to medical therapy for selected patients with chronic heart failure reduced the combination of deaths and heart-related hospital stays compared with medical therapy alone.

May 12, 2011. Results from the Pediatric Hydroxyurea Phase III Clinical Trial (Baby HUG) show that hydroxyurea appears to be safe for treating SCD in children aged 8–19 months, and can reduce their pain episodes and improve key blood measurements.

August 24, 2011. Results from the COPD Clinical Research Network show that adding a common antibiotic to the usual daily treatment regimen for COPD reduced the occurrence of acute exacerbations and improved the quality of life of patients.

September 26, 2011. Results from a genome-wide association study show that asthma patients who have two copies of a specific gene variant responded only one-third as well to steroid inhalers used to treat asthma as those with two copies of the normal gene.



4. Disease Statistics

Cardiovascular, lung, and blood diseases constitute a large morbidity, mortality, and economic burden on individuals, families, and the Nation. Common forms are atherosclerosis, hypertension, COPD, and blood-clotting disorders—embolisms and thromboses. The most serious atherosclerotic diseases are CHD, as manifested by heart attack and angina pectoris, and cerebrovascular disease, as manifested by stroke.

In 2008, cardiovascular, lung, and blood diseases accounted for 1,052,000 deaths and 43 percent of all deaths in the United States (p. 35). The estimated economic cost in 2008 for these diseases was \$392 billion, 22 percent of the total economic costs of illness, injuries, and death (p. 51). Of all diseases, heart disease is the leading cause of death; chronic lower respiratory diseases (CLRD), which includes COPD and asthma, ranks third (behind cancer); and cerebrovascular disease is fourth (p. 38). Cardiovascular and lung diseases account for 3 of the 4 leading causes of death (p. 38) and 4 of the 10 leading causes of infant death (p. 44). Hypertension, asthma, CHD, and COPD are especially prevalent and account for substantial morbidity in Americans (p. 47).

The purpose of the biomedical research conducted by the NHLBI is to contribute to the prevention and treatment of cardiovascular, lung, and blood diseases and sleep disorders. National disease statistics show that by mid-century, morbidity and mortality from these diseases had reached record high levels. Since then, however, substantial improvements have been achieved, especially over the past 40 years, as shown by the significant decline in mortality rates. Because many of these diseases begin early in life, their early detection and control can reduce the risk of disability and can delay death. Although important advances have been made in the treatment and control of cardiovascular, lung, and blood diseases, these diseases continue to be a major burden on the Nation.

Mortality statistics in this chapter are for diseases or conditions classified as the underlying cause of death. Heart failure, however, is never truly an underlying cause even though 56,830 deaths in 2008

were nominally coded to it as the underlying cause. Therefore, in this chapter, mortality statistics attributed to any mention of heart failure represent it as either the underlying cause or a contributing cause of death.

Cardiovascular Diseases

- In 2008, CVD caused 812,000 deaths—33 percent of all deaths (p. 35).
- Heart disease is the leading cause of death; the main form, CHD, caused 405,000 deaths in 2008 (pp. 36, 38).
- The annual number of deaths from CVD increased substantially from 1900 to 1970 and remains high (p. 37).
- The death rate (not age-adjusted) for CVD increased from 1920 until it peaked in 1968. Since then, the trend has been downward. In 2008, the rate was below the all-time low in 1900 (p. 37).
- Cerebrovascular disease, the fourth leading cause of death, accounted for 134,000 deaths in 2008 (pp. 36, 38).
- Heart disease is second only to all cancers combined in years of potential life lost (p. 38).
- Heart disease is the leading cause of death in blacks and Hispanics, but second to cancer in Asians and American Indians. Stroke ranks as the third or fourth leading cause of death in the minority groups, except in American Indians, where it ranks seventh (p. 38).
- Deaths with heart failure as the underlying or contributing cause increased from 1970 to 1993 and then remained constant to 2008 (p. 39).
- From 1999 to 2008, death rates for CHD and stroke declined in males and females of all racial/ethnic groups. CHD mortality remained highest in the black population and lowest in the Asian population. Stroke mortality continues to be highest in the black population (p. 40).
- Because of the rapid decline in mortality from CHD since the peak in 1968, there were 1,174,000 fewer deaths from CHD in 2008 than would have occurred if there had been no decline (p. 41).

- Substantial improvements have been made in the treatment of CVD. Since 1975 or 1985, the percent of hospitalizations for AMI, stroke, heart failure, and cardiac dysrhythmias that were discharged dead declined appreciably (p. 41).
- The decline in CHD mortality began earlier in the United States than in most countries and outpaced that in most countries until the 1990s (only selected countries are shown) (p. 42).
- From 1999 to 2008, the percentage decline in death rates for CHD and stroke was fairly similar for whites and blacks (p. 43).
- In 2008, an estimated 82.6 million persons in the United States had CVD, including 76.4 million with hypertension and 16.3 million with CHD (p. 47).
- Since the 1960s, there has been a substantial reduction in the prevalence of CVD risk factors: hypertension, smoking, and high cholesterol, but not overweight. The large decline in prevalence of hypertension from 1976–1980 to 1988–1994 was followed by a slightly higher prevalence in 1999–2004 and 2005–2008 (p. 48).
- From 1976–1980 to 2005–2008, the percentage of persons with hypertension who were aware of their condition, on treatment for it, and having their blood pressure under control increased substantially (p. 49).
- A 2005–2008 national survey showed only about 48 percent of hypertensive patients (systolic BP \geq 140 mmHg or diastolic BP \geq 90 mmHg or on antihypertensive medication) had their condition under control (p. 49).
- Hospitalization rates for heart failure in those aged 45 to 64 years increased from 1971 to 1993 and remained stable to 2009. Rates for those aged 65 years and older increased from 1971 to 1998 and remained relatively stable until 2005; rates then declined through 2009 (p. 50).
- The estimated economic cost of CVD for 2008 was \$298 billion:
 - \$179 billion in direct health expenditures
 - \$118 billion in indirect cost of mortality (p. 51).

Lung Diseases

- Lung diseases, excluding lung cancer, caused an estimated 242,000 deaths in 2008 (p. 35).
- CLRD caused 141,000 deaths in 2008 and is the third leading cause of death (pp. 36, 38).
- From 1999 to 2008, death rates for asthma declined in both black and white males and females; death

- rates for COPD declined in both black and white males but rose in both black and white females (p. 43).
- From 1980 to 2008, infant death rates for various lung diseases declined markedly (p. 43).
- In 2008, of the 10 leading causes of infant mortality, 4 were lung diseases or had a lung disease component (p. 44). From 1998 to 2008, changes in mortality for the causes were:
 - Congenital malformations (-9 percent)
 - Disorders of short gestation (-3 percent)
 - Sudden infant death syndrome (-27 percent)
 - Respiratory distress syndrome (-51 percent).
- In 2008, approximately one in six deaths in children under 1 year of age was due to a lung disease (p. 44).
- From 1980 to 2008, the CLRD death rate for females in the United States increased appreciably compared with the rates in several other countries (p. 45).
- From 2000 to 2008, death rates for CLRD decreased slightly for Asian and Hispanic females but were stable for American Indian and non-Hispanic white and non-Hispanic black females. For males, the rates decreased in all racial/ethnic groups (p. 46).
- Among the sleep disorders, sleep apnea is increasingly being recognized as an important health problem, which can lead to serious consequences. From 2000 to 2009, physician office visits for sleep apnea increased from 2 to 3.7 million (p. 46).
- Asthma is a common chronic condition, particularly in children (pp. 47, 48, 50).
- The economic cost of asthma, COPD, and pneumonia was \$89 billion in 2008:
 - \$68 billion in direct health expenditures
 - \$21 billion in indirect cost of mortality (p. 51).

Blood Diseases

- Approximately 10,000 deaths were attributed to blood diseases in 2008 (p. 35). These include the following:
 - 5,000 due to anemias
 - 1,800 due to coagulation defects
 - 800 due to purpura
 - 2,400 due to other blood diseases.
- A large proportion of deaths from AMI, cerebrovascular disease, and peripheral artery disease involve blood-clotting problems (no estimate available).
- In 2008, anemias cost the Nation's economy \$6 billion:
 - \$5 billion in direct health expenditures
 - \$1 billion in indirect cost of mortality (p. 51).

Deaths From All Causes and Deaths From Cardiovascular, Lung, and Blood Diseases, U.S., 1988 and 2008

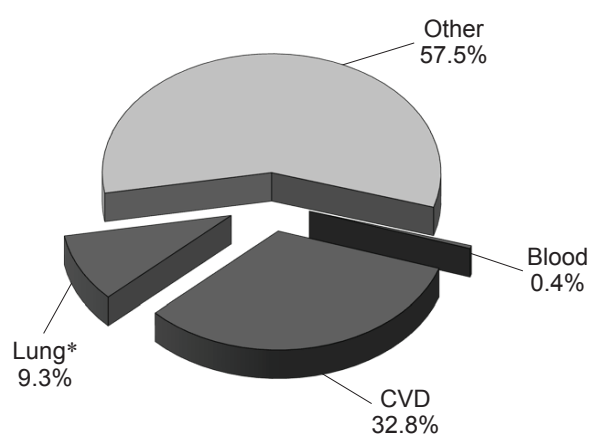
Cause of Death	1988		2008	
	Number of Deaths	Percent of Total	Number of Deaths	Percent of Total
All Causes	2,167,999	100	2,471,984	100
All Cardiovascular, Lung, and Blood Diseases	1,181,646	55	1,051,502	43
Cardiovascular	979,788	45	811,940	33
Blood	8,649	<1	10,066	<1
Lung	205,798*	9	242,350**	10
All Other Causes	986,353	45	1,420,482	57

* Includes 12,931 CVD deaths due to pulmonary heart disease.

** Includes 12,854 CVD deaths involving pulmonary heart disease.

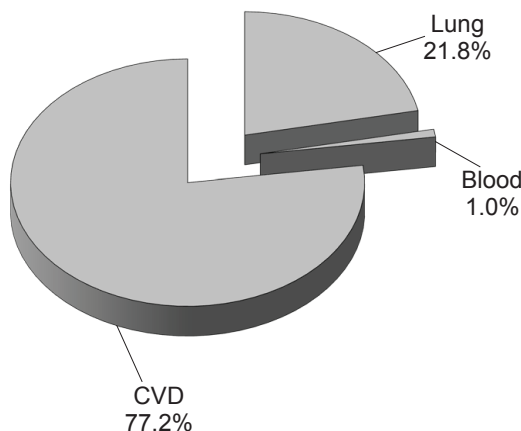
Source: Vital Statistics of the United States, National Center for Health Statistics (NCHS).

Deaths by Major Causes, U.S., 2008



■ Total Cardiovascular, Lung, and Blood Diseases 42.5%

Deaths From Cardiovascular, Lung, and Blood Diseases, U.S., 2008



* Excludes 12,854 deaths from pulmonary heart disease (0.5%).

Source: Vital Statistics of the United States, NCHS.

Deaths From Specific Cardiovascular, Lung, and Blood Diseases, U.S., 2008

Cause of Death	Deaths (Thousands)		
	Cardiovascular	Lung	Blood
Acute Myocardial Infarction	134	—	—*
Other Coronary Heart Disease	270	—	—
Cerebrovascular Diseases (Stroke)	134	—	—*
Other Atherosclerosis	28	—	—
Pulmonary Embolism	7	7**	—*
Deep Vein Thrombosis	2	—	—*
Other Cardiovascular Diseases	236	6**	—
Bleeding and Red Blood Cell Diseases [†]	—	—	10
Chronic Obstructive Pulmonary Disease	—	138	—
Asthma	—	3	—
Influenza and Pneumonia	—	56	—
Neonatal Pulmonary Disorders	—	4	—
Interstitial Lung Diseases	—	7	—
Lung Diseases Due to External Agents	—	18	—
Other Lung Diseases	—	3	—
Total	812[‡]	242	10[†]

* Deaths from pulmonary disorders also included as CVD.

** Deaths from anemias, coagulation defects, purpura, and other blood diseases. Deaths attributed to blood-clotting diseases classified to AMI, stroke, and peripheral artery disease are not included.

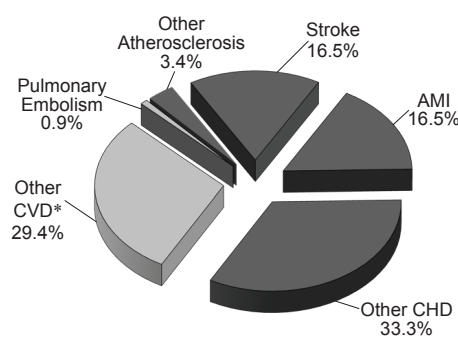
† Most deaths from this cardiovascular disease can be classified as a blood-clotting disease. No good estimate is available.

‡ Numbers do not sum to the total due to rounding.

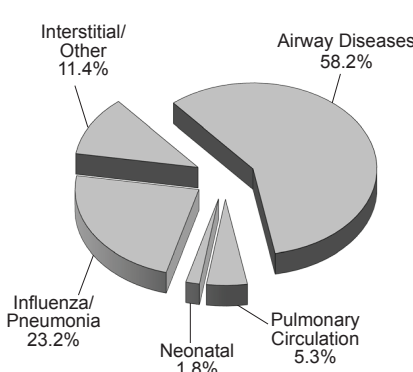
Note: Total, excluding overlap, is 1,051,502.

Source: Vital Statistics of the United States, NCHS.

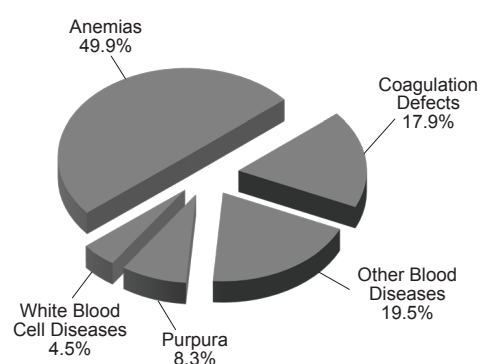
Deaths From Cardiovascular Diseases, U.S., 2008



Deaths From Lung Diseases, U.S., 2008



Deaths From Blood Diseases, U.S., 2008



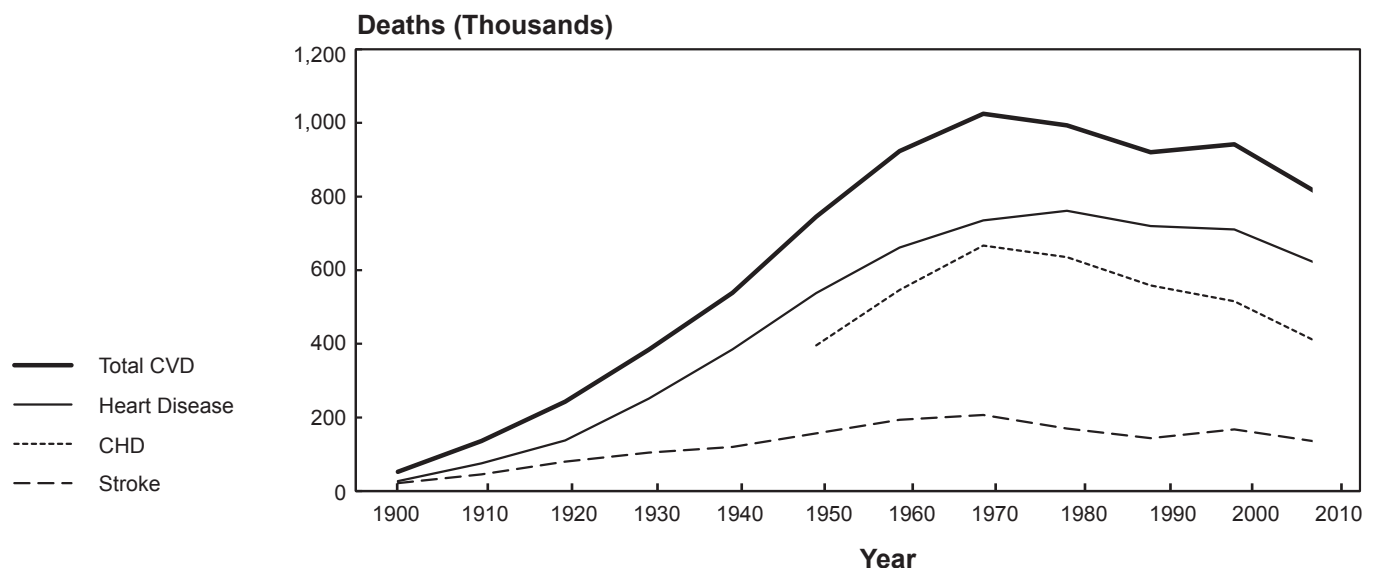
■ Atherosclerosis-related disease 69.7%

* Includes heart failure, cardiac dysrhythmias, hypertensive disease, deep vein thrombosis, and other heart and blood vessel diseases.

Note: Numbers do not sum to 100 percent due to rounding.

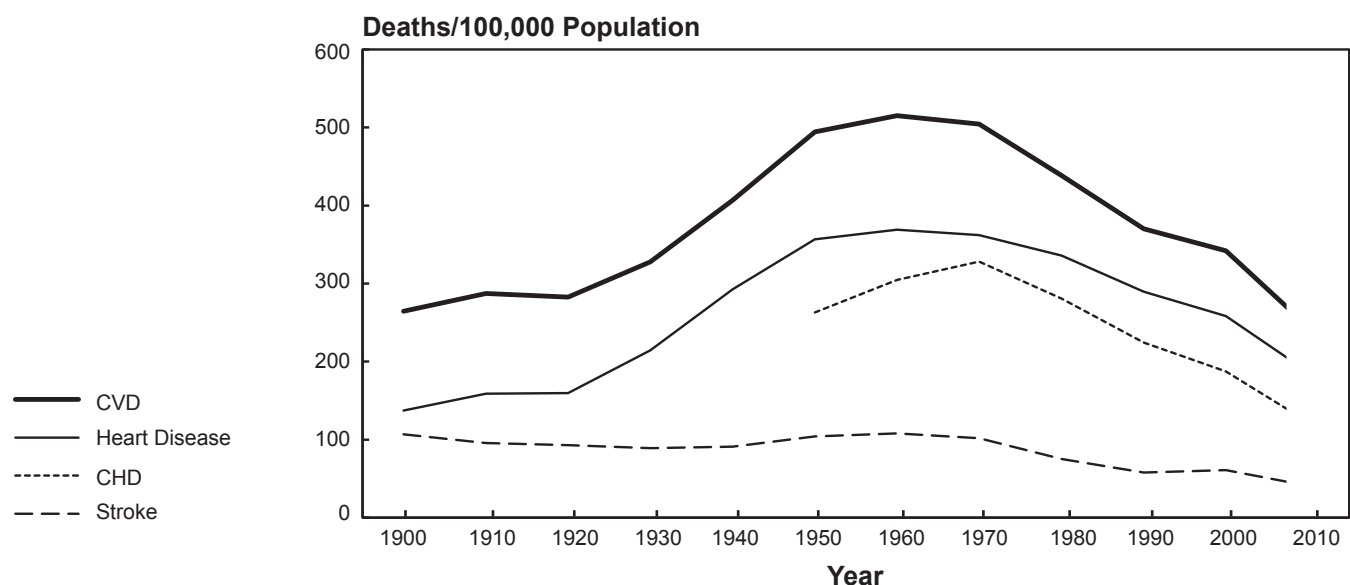
Source: Vital Statistics of the United States, NCHS.

Deaths From Cardiovascular Diseases, U.S., 1900–2008



Source: Vital Statistics of the United States, NCHS.

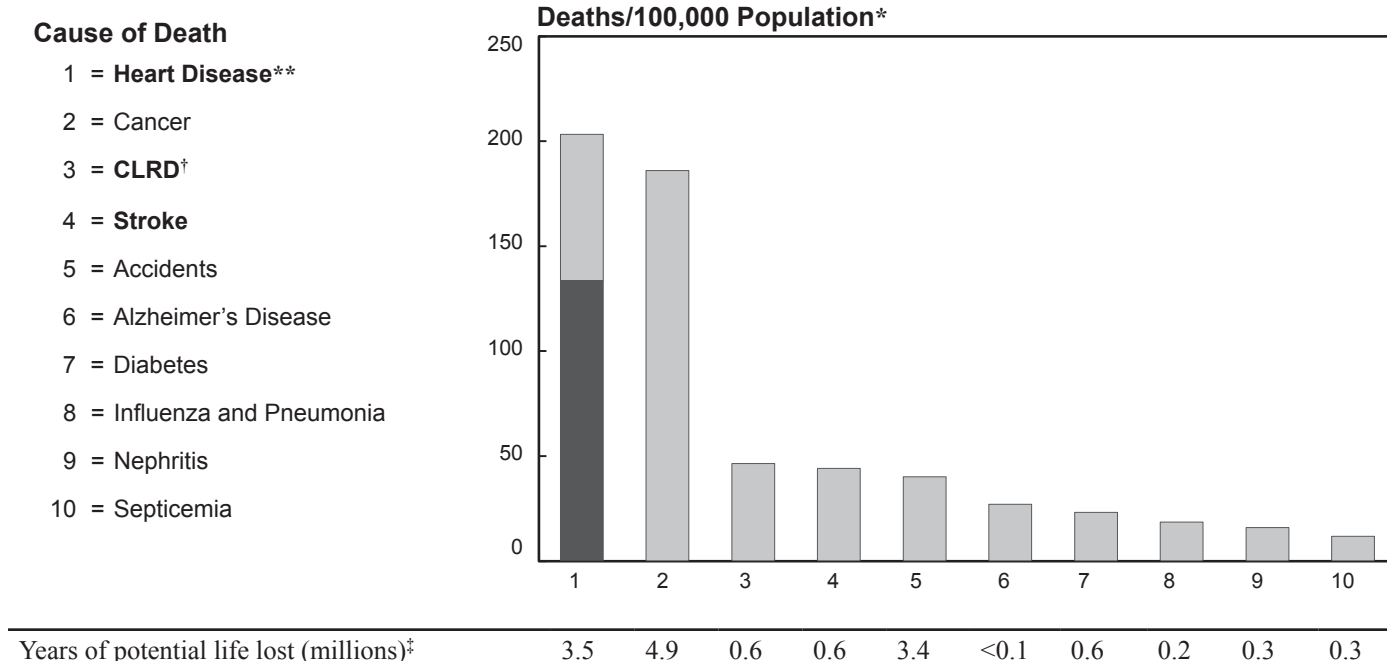
Death Rates* for Cardiovascular Diseases, U.S., 1900–2008



* Not age-adjusted.

Source: Vital Statistics of the United States, NCHS.

Ten Leading Causes of Death: Death Rates, U.S., 2008



* Not age-adjusted.

** Includes 133.3 deaths per 100,000 population from CHD.

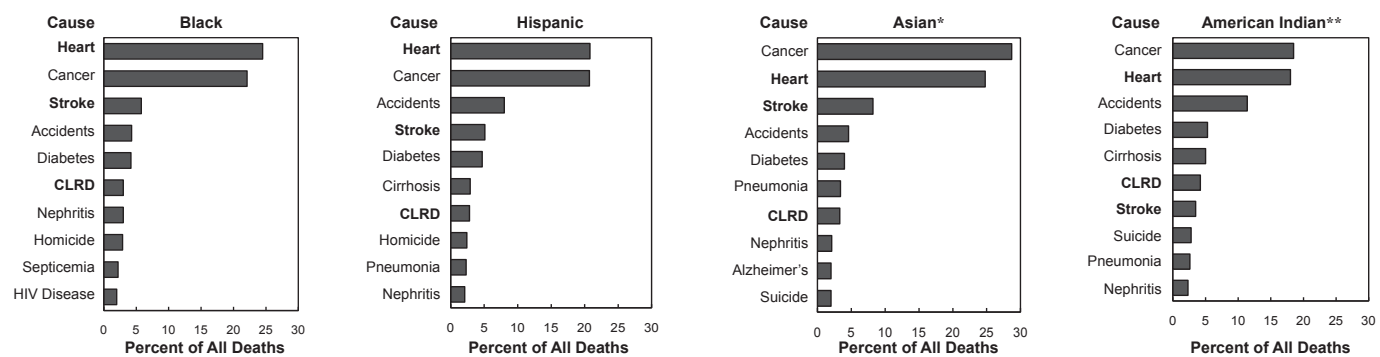
† CLRD is the term used in the ICD/10 for COPD and asthma.

‡ Based on the average remaining years of life up to age 77 years.

Note: Diseases shown in bold are those addressed in Institute programs.

Source: Vital Statistics of the United States, NCHS.

Ten Leading Causes of Death Among Minority Groups, U.S., 2008



* Includes deaths among individuals of Asian extraction and Asian-Pacific Islanders.

** Includes deaths among Aleuts and Eskimos.

Note: Causes of death shown in bold are those addressed in Institute programs.

Source: Vital Statistics of the United States, NCHS.

Age-Adjusted Death Rates for Cardiovascular and Noncardiovascular Diseases, U.S., 1963, 1988, and 2008

Cause of Death	Deaths/100,000 Population*			Percent Change	Percent Change
	1963	1988	2008	1963–2008	1988–2008
All Causes	1,346	975	760	-44	-22
Cardiovascular Diseases	805	450	245	-70	-46
Coronary Heart Disease	478	234	123	-74	-48
Stroke	174	74**	41	-77	-45
Other	153	142	81	-47	-43
Noncardiovascular Diseases	541	525	515	-5	-2
COPD and Asthma	16	38†	44	167	16
Other	524	487	471	-10	-3

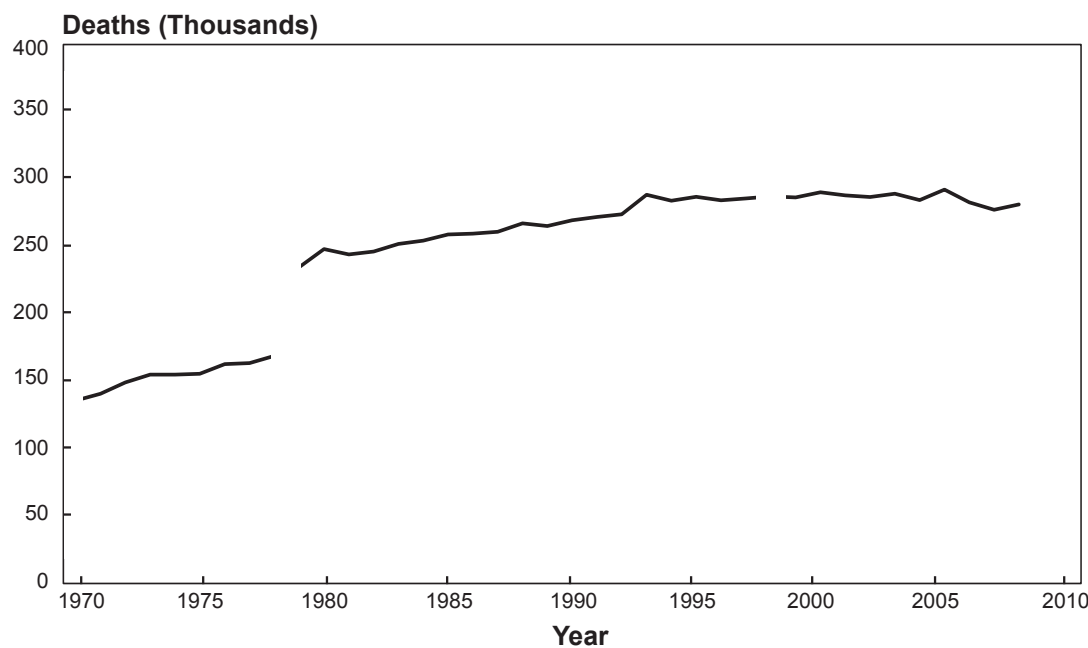
* Age-adjusted.

** ICD 10/9 comparability ratio (1.0502) applied.

† ICD 10/9 comparability ratio (1.0411) applied.

Source: Vital Statistics of the United States, NCHS.

Deaths Attributed to Heart Failure,* U.S., 1970–2008

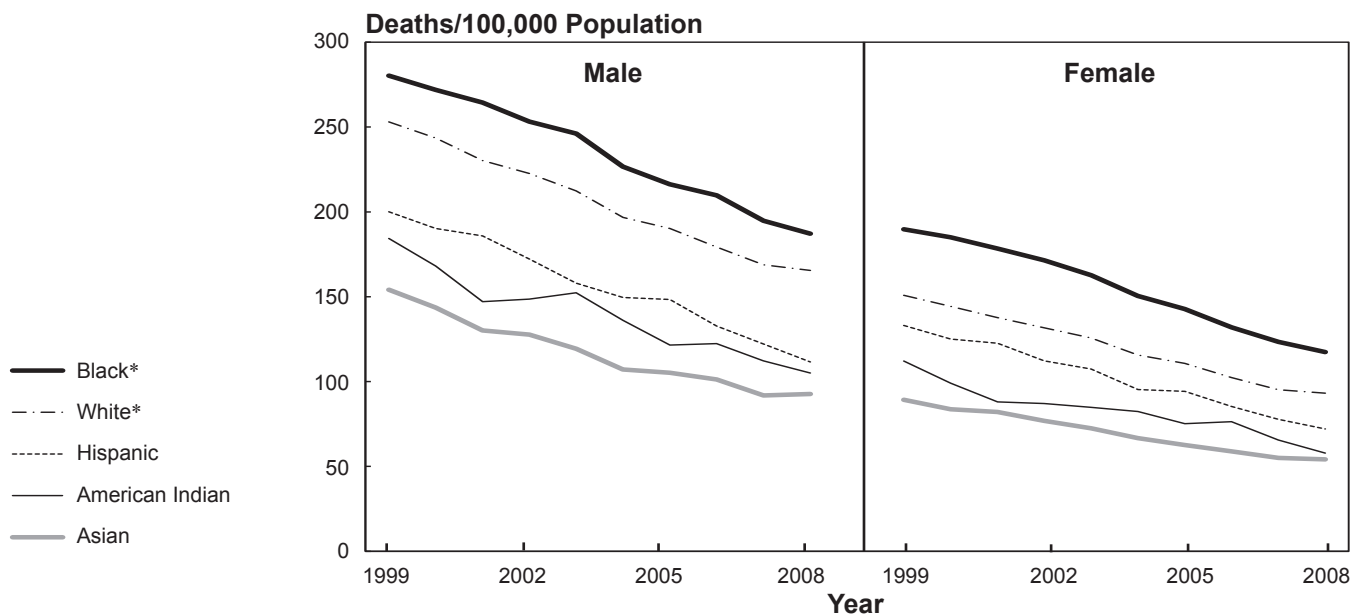


* Heart failure as the underlying cause of death or otherwise mentioned on the death certificate.

Note: Breaks in trend line indicate change in ICD codes.

Source: Vital Statistics of the United States, NCHS.

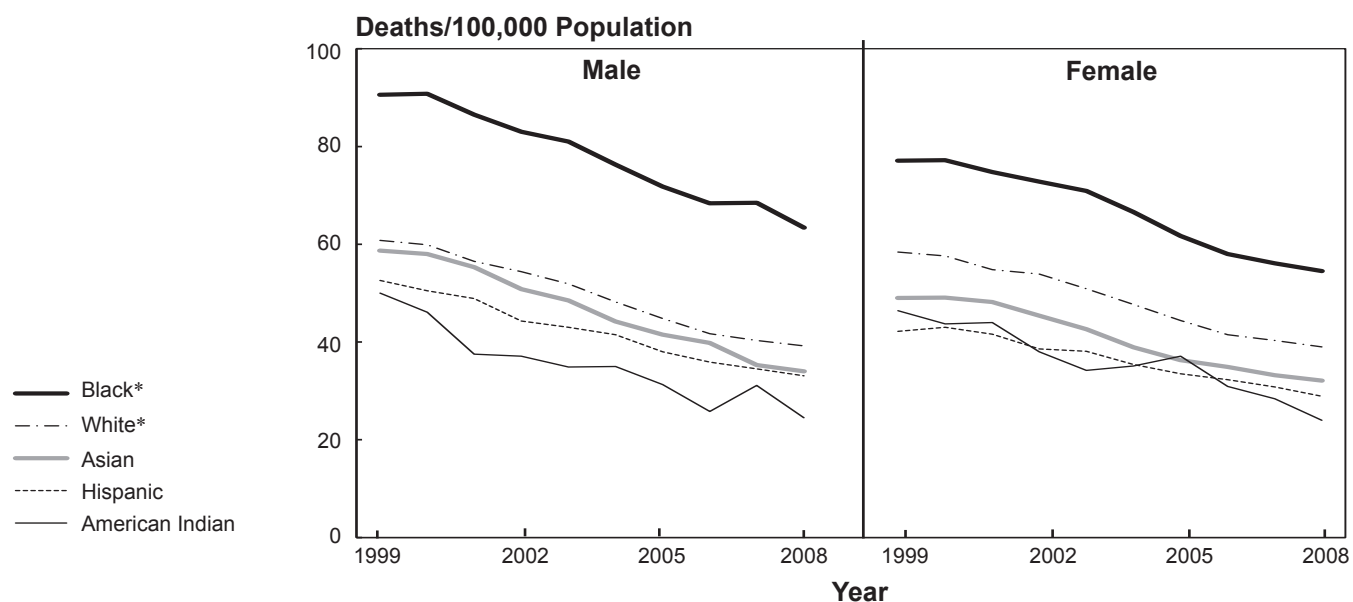
Age-Adjusted Death Rates for Coronary Heart Disease by Race/Ethnicity and Sex, U.S., 1999–2008



* Non-Hispanic.

Source: Vital Statistics of the United States, NCHS.

Age-Adjusted Death Rates for Stroke by Race/Ethnicity and Sex, U.S., 1999–2008

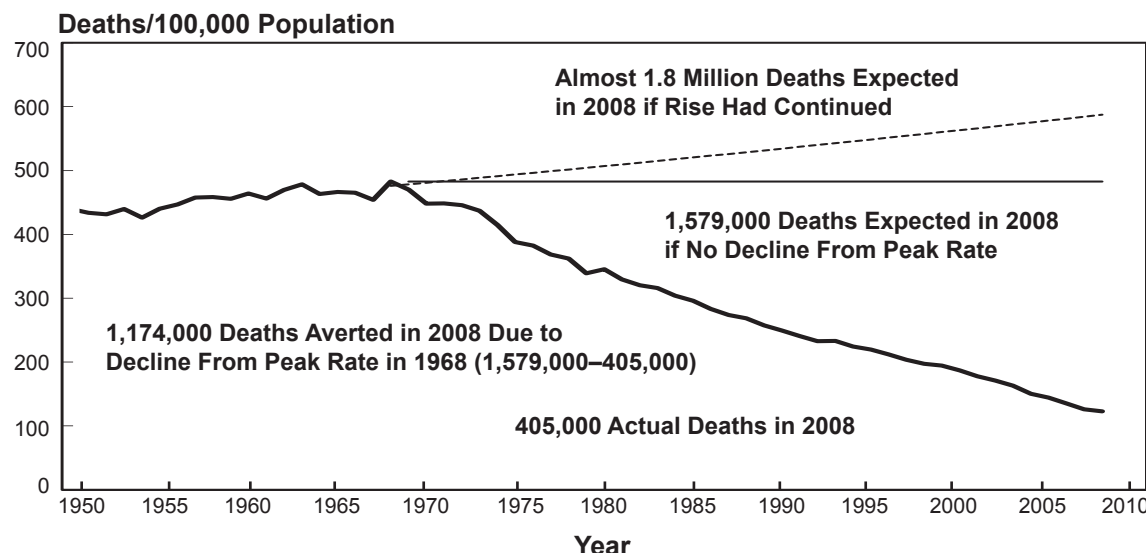


* Non-Hispanic.

Source: Vital Statistics of the United States, NCHS.

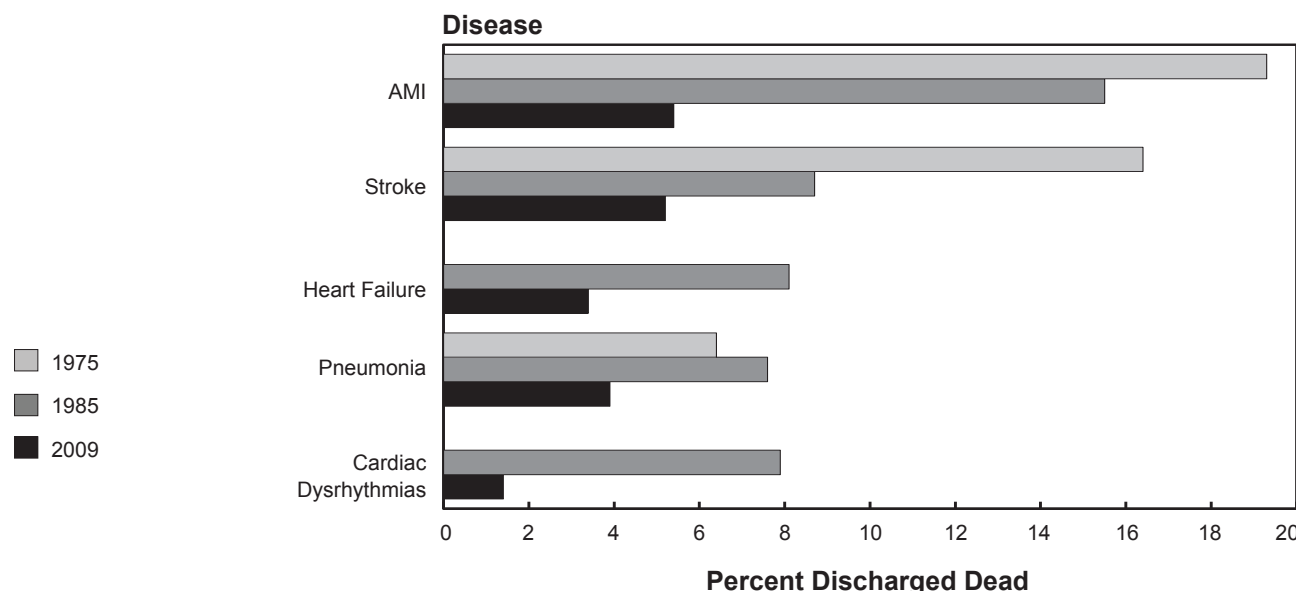
Age-Adjusted Death Rates for Coronary Heart Disease, U.S., 1950–2008

Actual Rate and Expected Rates if Rise Had Continued or Reached a Plateau



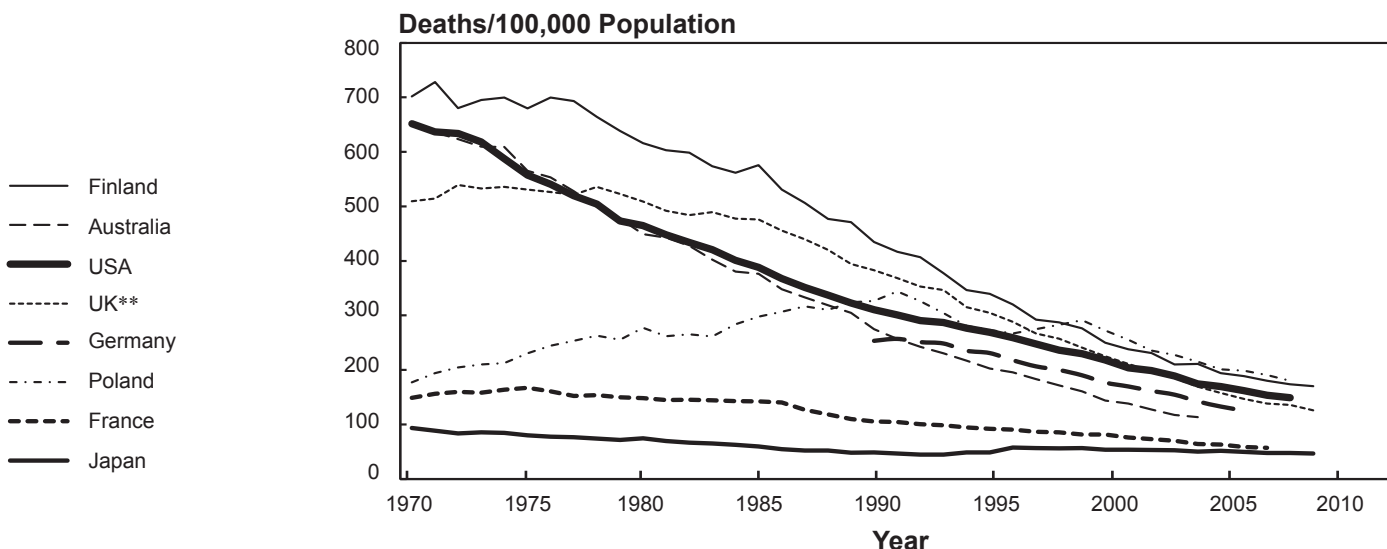
Source: Vital Statistics of the United States, NCHS.

Common Cardiovascular and Lung Diseases With High Percentage Discharged Dead From Hospitals, U.S., 1975, 1985, and 2009



Source: National Hospital Discharge Survey (NHDS), NCHS.

Death Rates* for Coronary Heart Disease in Males, Ages 35–74 Years, in Selected Countries, 1970–2009

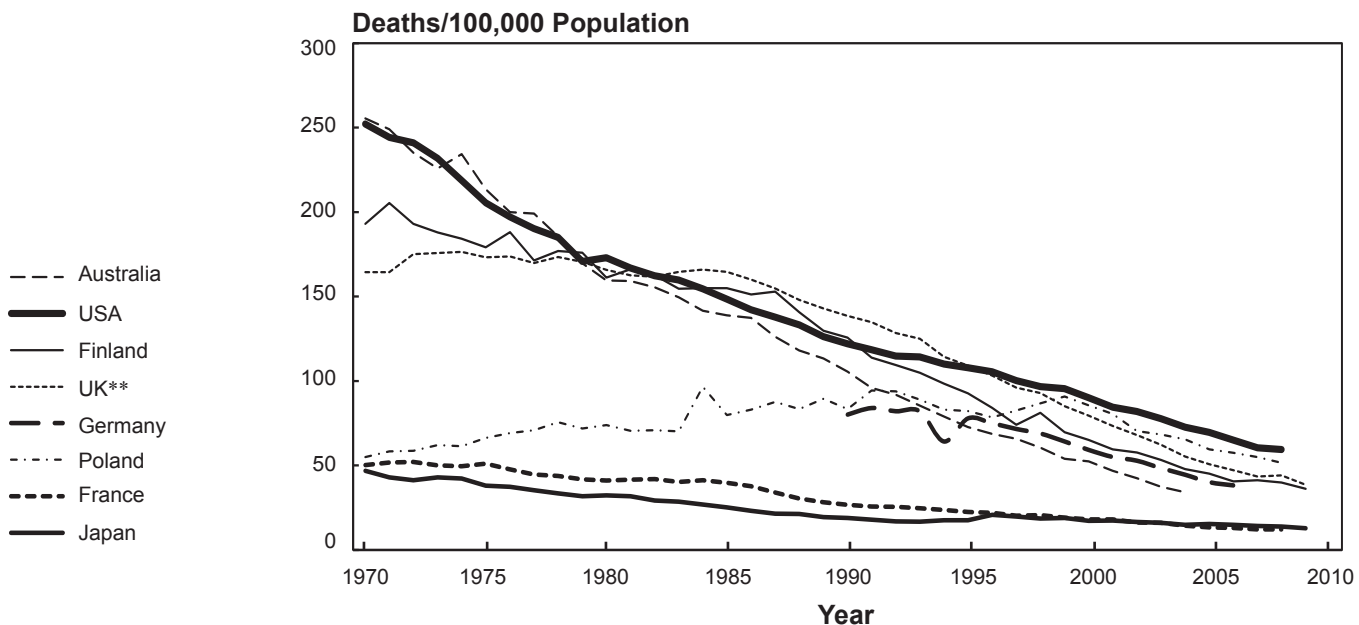


* Age adjusted to the European Standard Population.

**United Kingdom for 2008 and 2009; England and Wales for 1970–2007.

Source: World Health Organization (WHO) Mortality Database.

Death Rates* for Coronary Heart Disease in Females, Ages 35–74 Years, in Selected Countries, 1970–2009

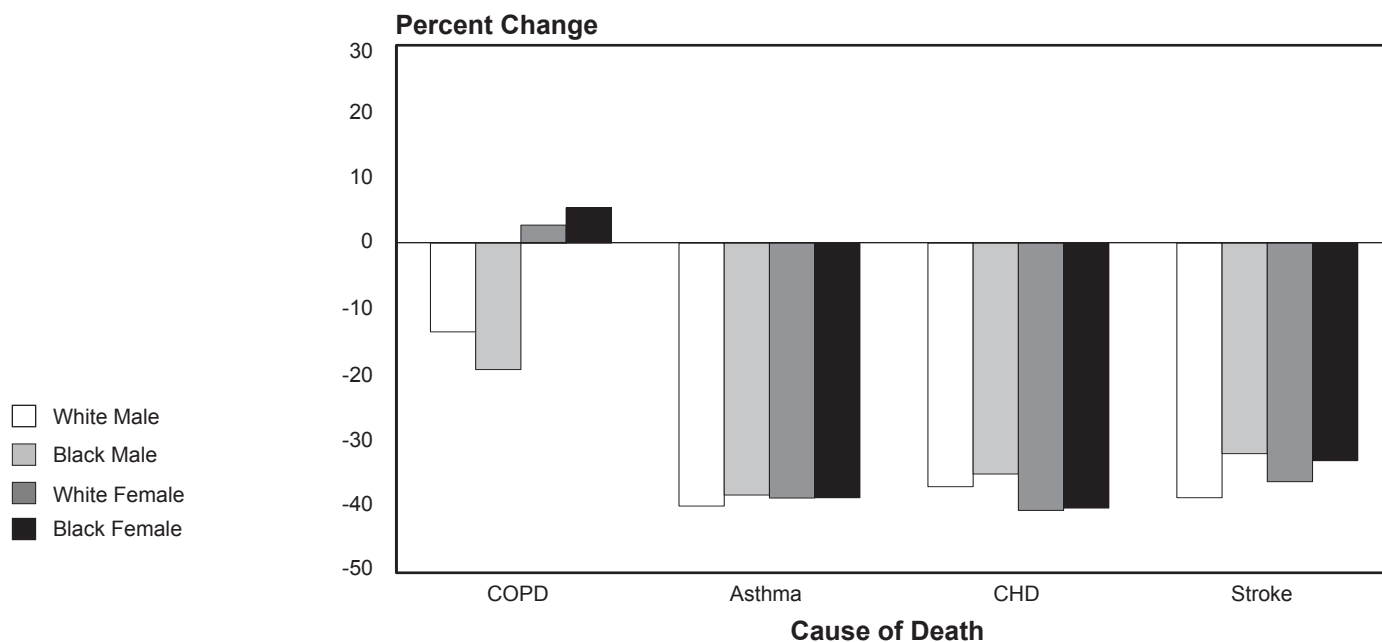


* Age adjusted to the European Standard Population.

**United Kingdom for 2008 and 2009; England and Wales for 1970–2007.

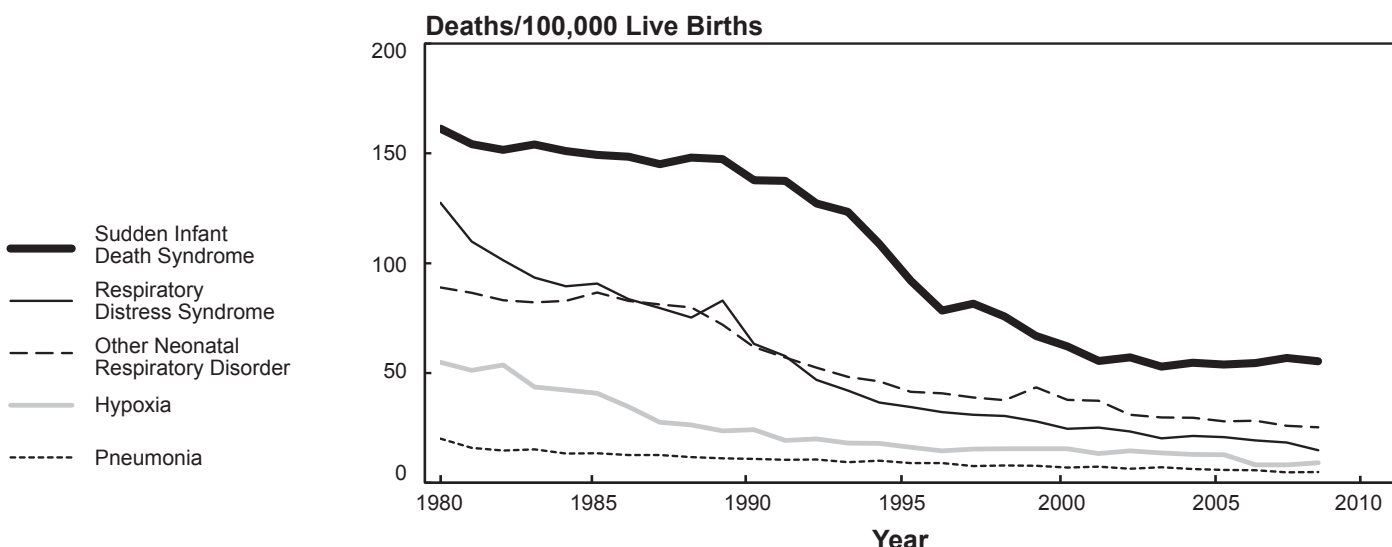
Source: WHO Mortality Database.

Percent Change in Age-Adjusted Death Rates for Selected Causes by Race and Sex, U.S., 1999–2008



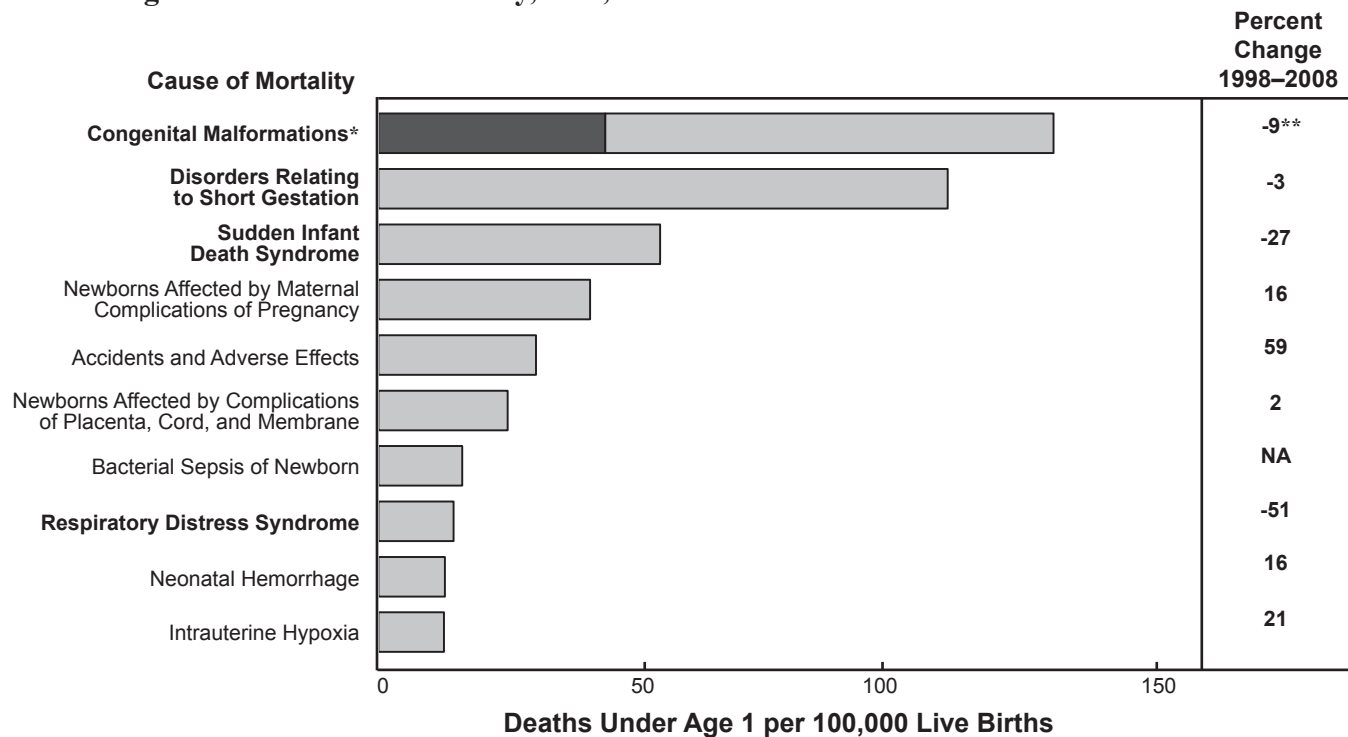
Source: Vital Statistics of the United States, NCHS.

Death Rates for Lung Diseases in Infants, U.S., 1980–2008



Source: Vital Statistics of the United States, NCHS.

Ten Leading Causes of Infant Mortality, U.S., 2008



* Congenital CVD and congenital respiratory diseases accounted for 44.6 deaths under age 1 per 100,000 live births (black bar), which is 34 percent of infant deaths due to all congenital malformations.

** From 1998 to 2008, congenital CVD declined 30 percent; congenital malformations of the respiratory system declined 47 percent; other congenital malformations increased 20 percent.

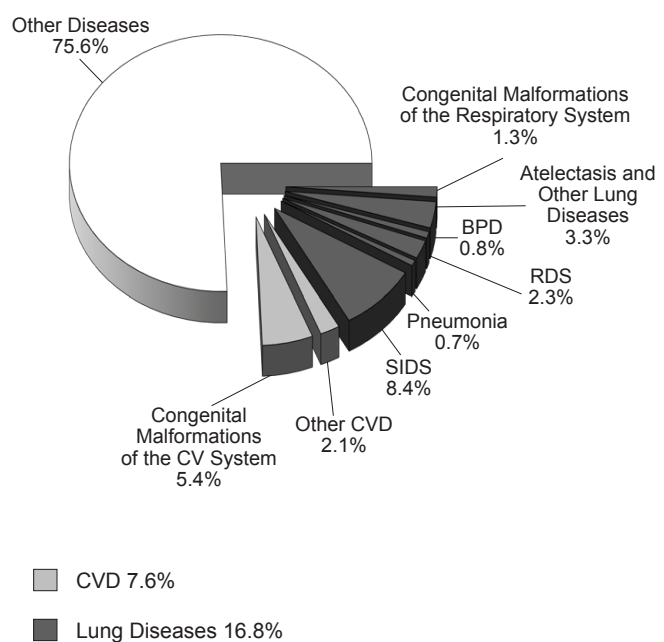
NA: Not available.

Note: Diseases shown in bold are those addressed in Institute programs.

Source: Vital Statistics of the United States, NCHS.

Deaths Under Age 1 Year Due to Cardiovascular and Lung Diseases, U.S., 2008

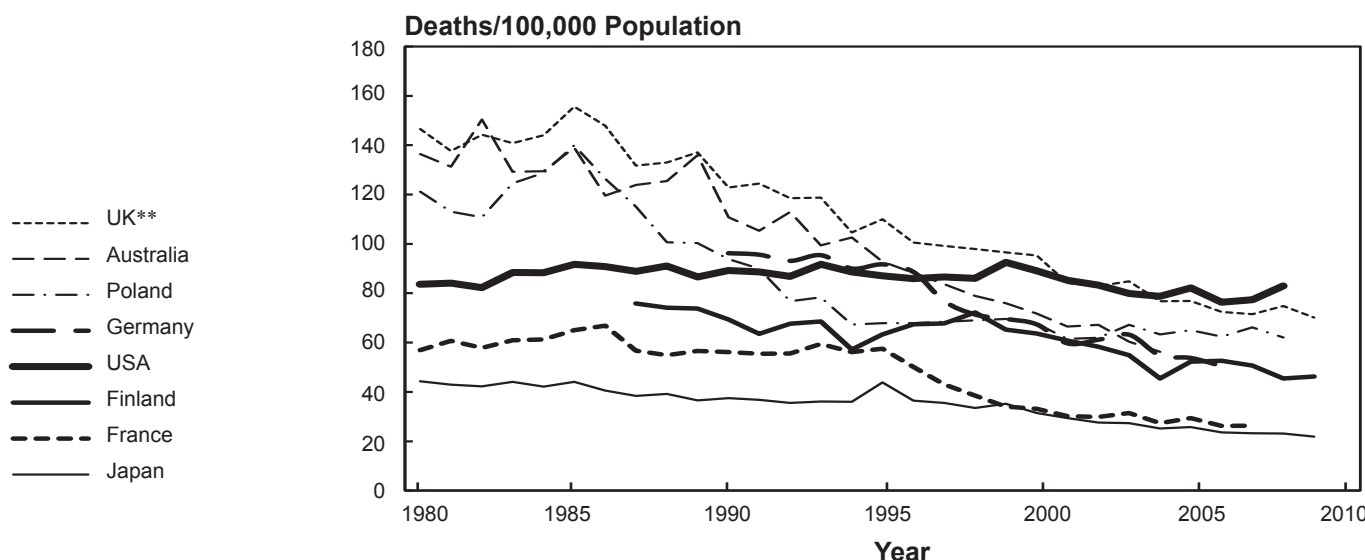
Cause of Death	Deaths Under Age 1
All Causes	28,059
Cardiovascular Diseases	2,121
Congenital Malformations	1,527
Other	594
Lung Diseases	4,718
Sudden Infant Death Syndrome	2,353
Respiratory Distress Syndrome	640
Pneumonia	210
Bronchopulmonary Dysplasia (BPD)	217
Atelectasis of Newborn	335
Congenital Malformations	371
Other Lung Diseases	592
Other Diseases	21,220



Note: Diseases shown in bold are those addressed in Institute programs.

Source: Vital Statistics of the United States, NCHS.

Death Rates* for Chronic Lower Respiratory Diseases in Males, Ages 35 Years and Older, in Selected Countries, 1980–2009

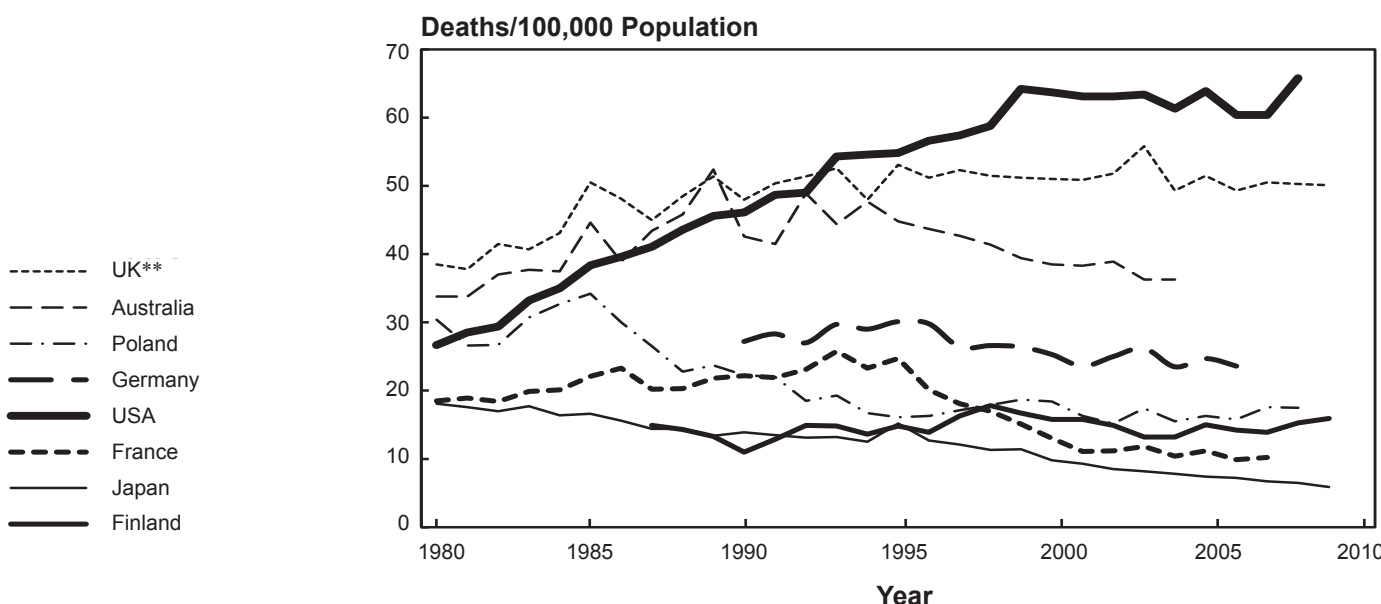


* Age adjusted to the European Standard Population.

** United Kingdom for 2008 and 2009; England and Wales for 1970–2007.

Source: WHO Mortality Database.

Death Rates* for Chronic Lower Respiratory Diseases in Females, Ages 35 Years and Older, in Selected Countries, 1980–2009

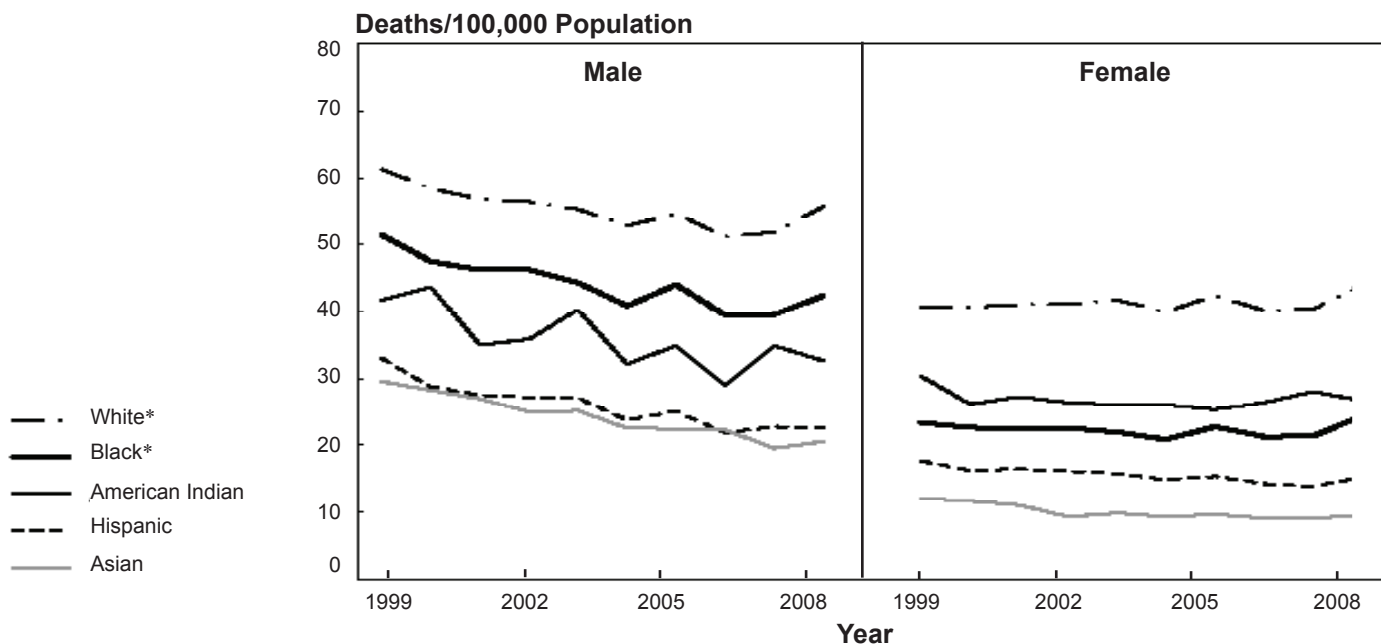


* Age adjusted to the European Standard Population.

** United Kingdom for 2008 and 2009; England and Wales for 1970–2007.

Source: WHO Mortality Database.

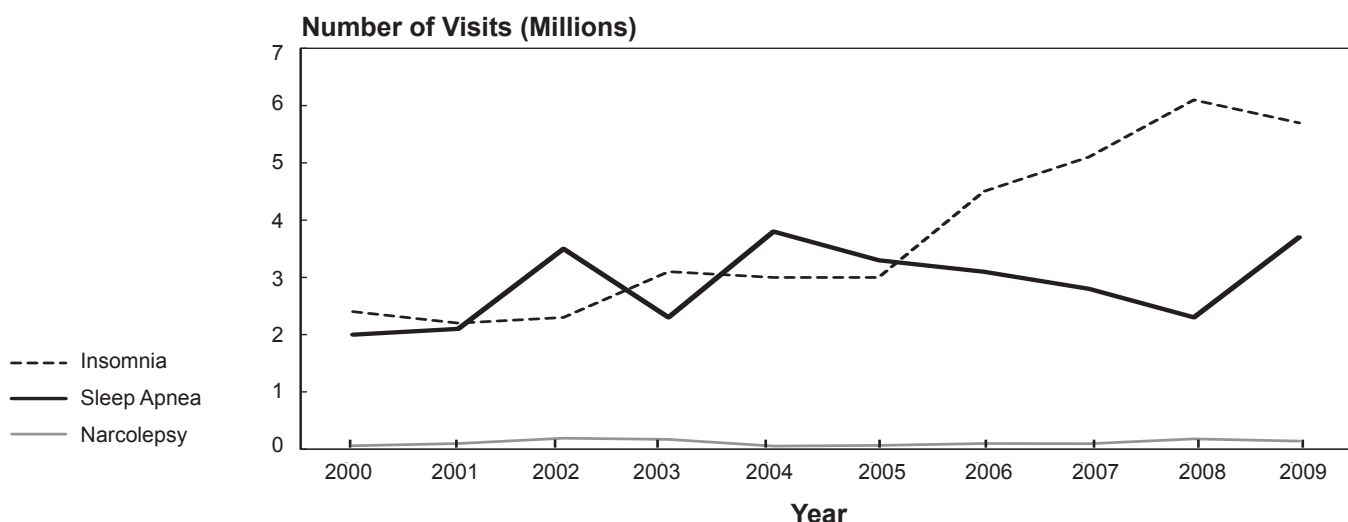
Age-Adjusted Death Rates for Chronic Lower Respiratory Diseases by Race/Ethnicity and Sex, U.S., 1999–2008



* Non-Hispanic.

Source: Vital Statistics of the United States, NCHS.

Physician Office Visits for Sleep Disorders, U.S., 2000–2009



Note: Primary and secondary diagnoses.

Source: National Ambulatory Medical Care Survey, NCHS.

Prevalence of Common Cardiovascular and Lung Diseases, U.S., 2005–2010

Disease	Number of Persons
Cardiovascular Diseases*	82,600,000
Hypertension**	76,400,000
Coronary Heart Disease	16,300,000
Heart Failure	5,700,000
Stroke	7,000,000
Congenital Heart Disease†	1,000,000
Asthma‡	39,190,000
COPD§	12,481,000

* Includes hypertension, CHD, stroke, or heart failure for ages 20 years and older.

** Hypertension is defined as systolic blood pressure ≥ 140 mmHg, or diastolic blood pressure ≥ 90 mmHg, or being on antihypertensive medication, or being told twice of having hypertension.

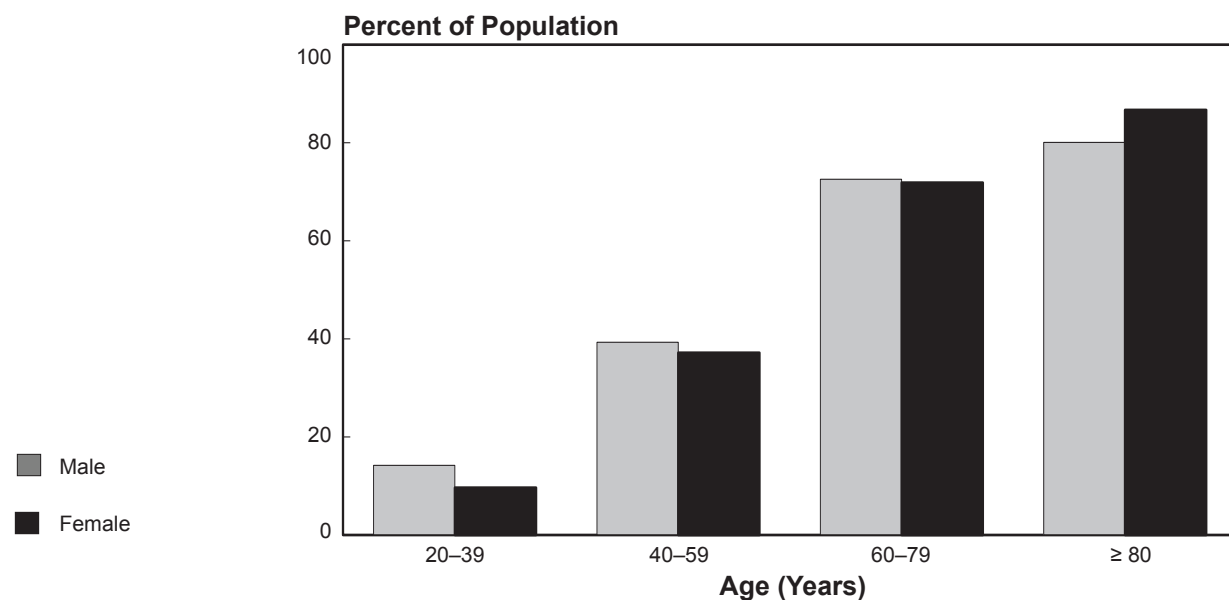
† Range from 650,000 to 1,300,000 for ages 18 years and older (Am Heart J 2004;147:425–439).

‡ 25,710,000 still have asthma and of those, 13,882,000 have had an attack in the past 12 months, for all ages.

§ An estimated 12,481,000 diagnosed (2010) and 12,000,000 undiagnosed (2006), for ages 18 years and older.

Sources: National Health and Nutrition Examination Survey (NHANES) 2005–2008, NCHS and National Health Interview Survey (NHIS) 2010, NCHS.

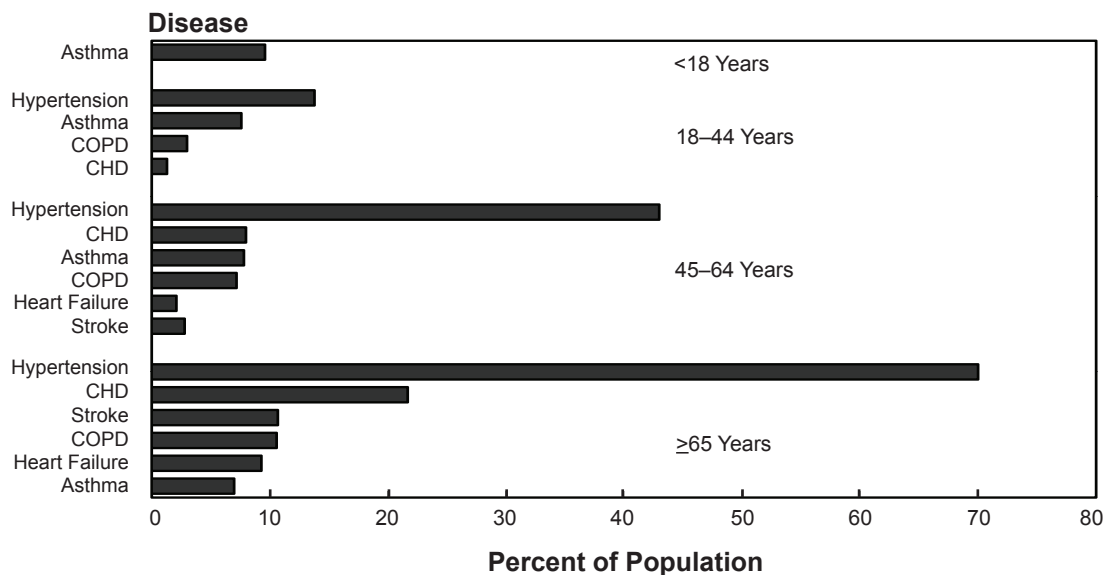
Prevalence of Cardiovascular Diseases* in Adults by Age and Sex, U.S., 2005–2008



* Hypertension, CHD, stroke, or heart failure. Hypertension is defined as systolic blood pressure ≥ 140 mmHg, or diastolic blood pressure ≥ 90 mmHg, or being on antihypertensive medication.

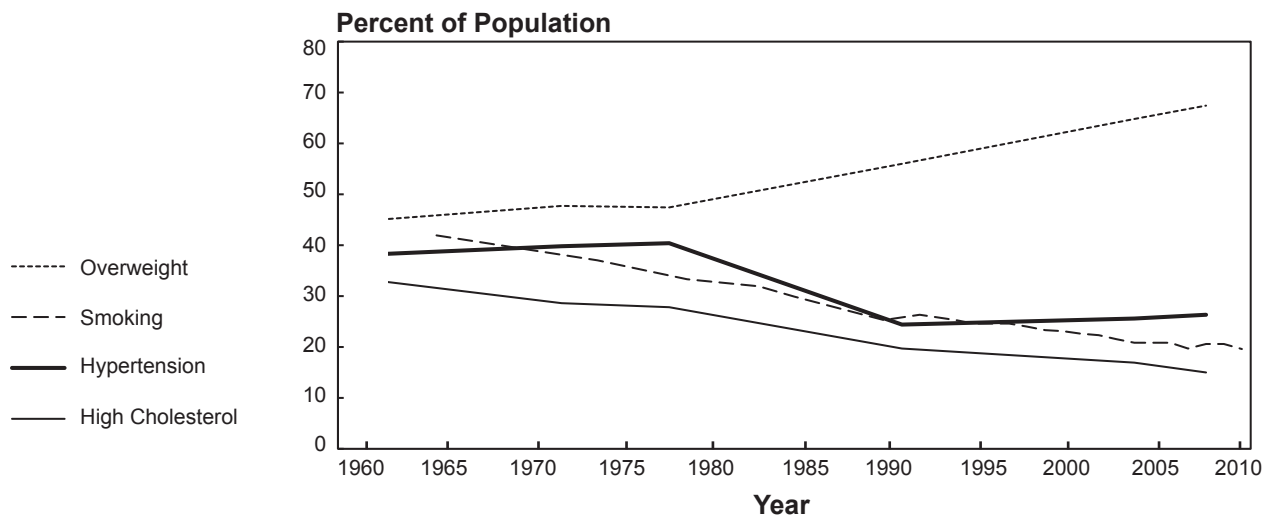
Source: NHANES, 2005–2008, NCHS.

Prevalence of Common Cardiovascular and Lung Diseases by Age, U.S., 2005–2010



Sources: NHIS and NHANES, NCHS.

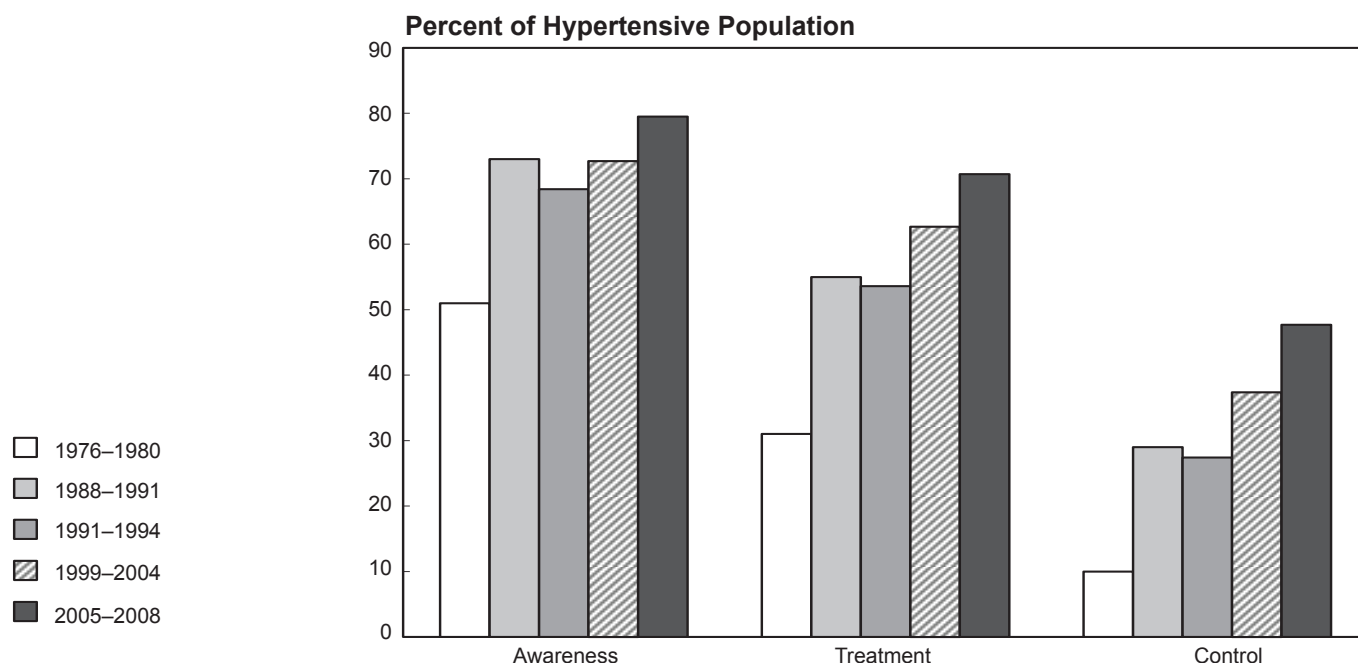
Age-Adjusted Prevalence of Cardiovascular Disease Risk Factors in Adults, U.S., 1961–2010



Notes: Hypertension is defined as systolic blood pressure ≥ 140 mmHg or diastolic blood pressure ≥ 90 mmHg, or being on antihypertensive medication. High cholesterol is ≥ 240 mg/dL. Overweight is BMI ≥ 25 kg/m 2 . Data were collected at six time periods: 1960–1961 (plotted at 1961), 1971–1974 (plotted at 1972), 1976–1980 (plotted at 1978), 1988–1994 (plotted at 1991), 1999–2004 (plotted at 2004), and 2005–2008 (plotted at 2008).

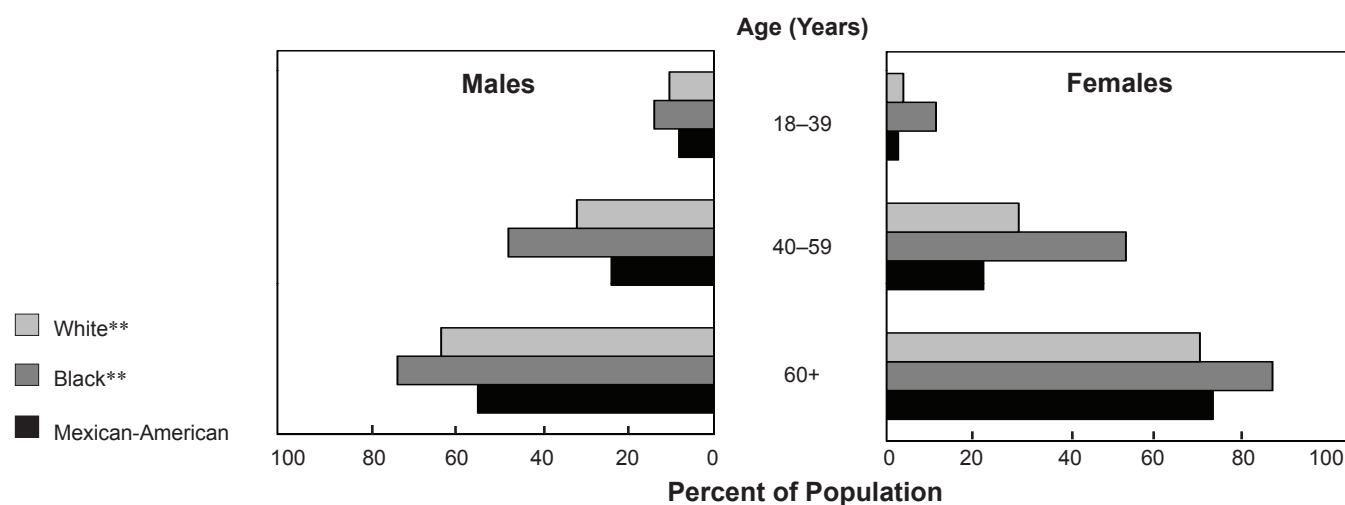
Sources: NHIS for smoking, ages ≥ 18 , NCHS; NHANES for the other risk factors, ages 20–74, NCHS.

Hypertensive* Population Aware, Treated, and Controlled, Ages 18 Years and Older, U.S., 1976–1980 to 2005–2008



* Hypertension is defined as systolic blood pressure ≥ 140 mmHg, or diastolic blood pressure ≥ 90 mmHg, or being on antihypertensive medication.
Source: NHANES, NCHS.

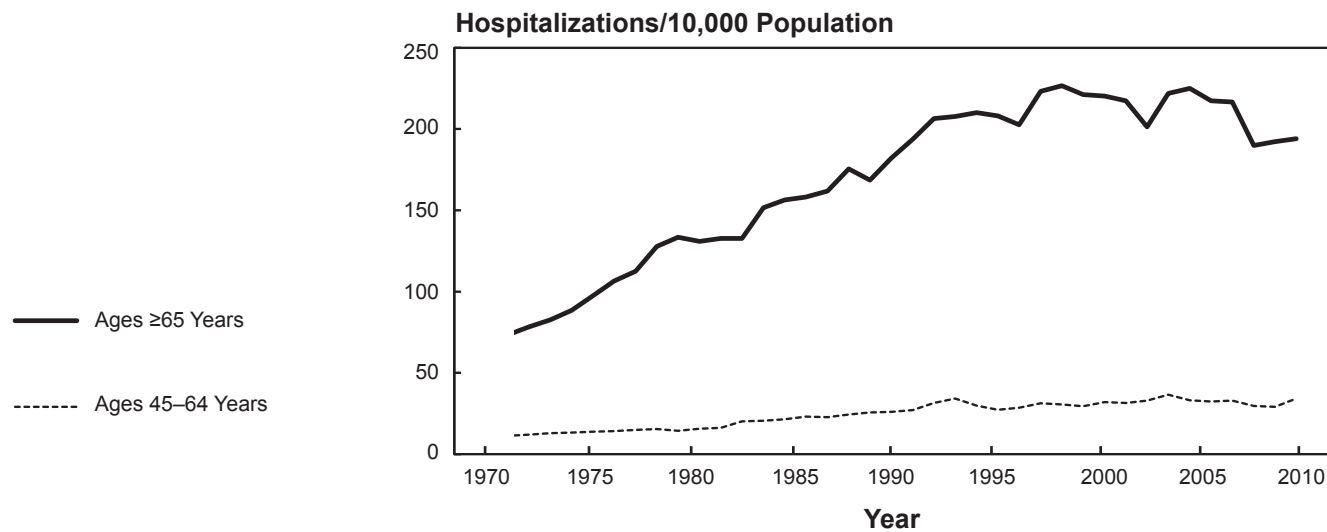
Adult Population With Hypertension* by Age, Race/Ethnicity, and Sex, U.S., 2005–2008



* Hypertension is systolic blood pressure ≥ 140 mm Hg, diastolic blood pressure ≥ 90 mmHg, or being on antihypertensive medication.
** Non-Hispanic.

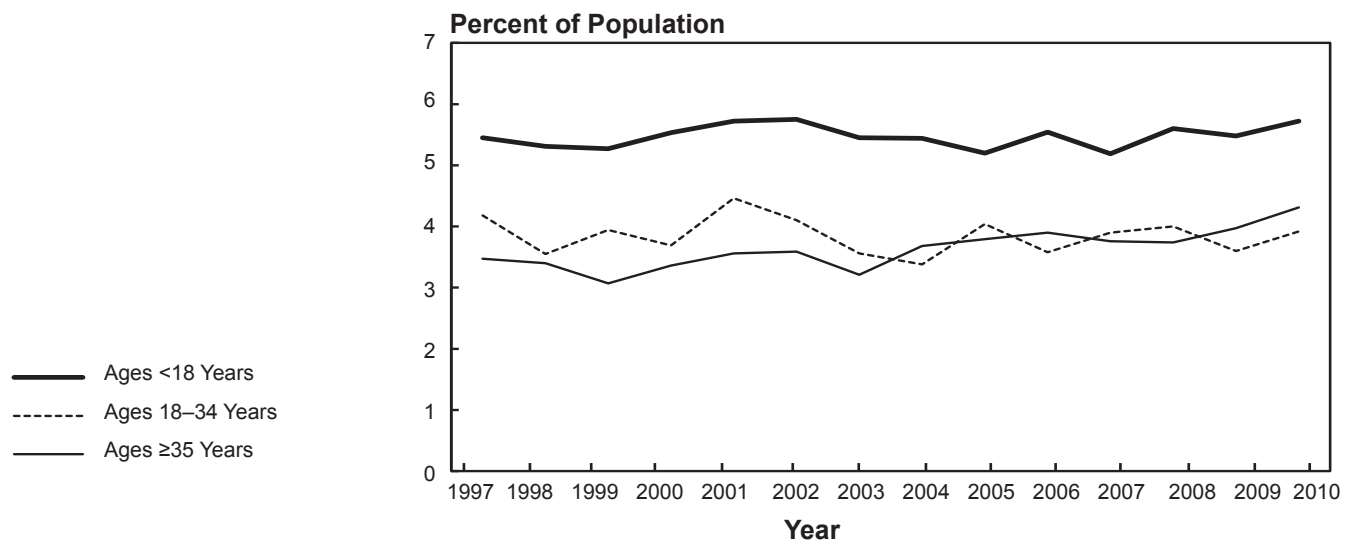
Source: NHANES, NCHS.

Hospitalization Rates for Heart Failure, Ages 45–64 Years and 65 Years and Older, U.S., 1971–2009



Source: NHDS, NCHS.

Prevalence of Asthma Episodes in Previous 12 Months by Age, U.S., 1997–2010



Source: NHIS, NCHS.

Direct and Indirect Economic Costs of Illness by Major Diagnosis, U.S., 2008

	Amount (Dollars in Billions)			Percent Distribution		
	Direct Cost*	Indirect Cost of Mortality**	Total	Direct Cost	Indirect Cost of Mortality	Total
Cardiovascular Diseases	\$179.3	\$118.4	\$297.7	15.6%	19.9%	17.1%
COPD, Asthma, Pneumonia	67.7	21.0	88.7	5.9	3.5	5.1
Anemias and Other Deficiencies	4.7	1.2	5.9	0.4	0.2	0.3
Subtotal	251.7	140.6	392.3	21.9	23.6	22.5
Neoplasms	77.4	124.0	201.5	6.7	20.9	11.5
Injury and Poisoning	77.7	101.6	179.3	6.8	17.1	10.3
Endocrine, Nutritional, and Metabolic Diseases	102.3	22.6	124.9	8.9	3.8	7.2
Diseases of the Digestive System	84.2	28.6	112.8	7.3	4.8	6.5
Diseases of the Respiratory System†	78.5	27.4	105.9	6.8	4.6	6.1
Diseases of the Musculoskeletal System	117.8	3.0	120.9	10.2	0.5	6.9
Diseases of the Nervous System	69.7	15.4	85.2	6.1	2.6	4.9
Mental Disorders	72.1	8.2	80.3	6.3	1.4	4.6
Diseases of the Genitourinary System	55.5	7.7	63.2	4.8	1.3	3.6
Infectious and Parasitic Diseases	16.2	23.9	40.1	1.4	4.0	2.3
Normal Live Birth	34.6	—	34.6	3.0	—	2.0
Diseases of the Skin	24.2	0.7	24.9	2.1	0.1	1.4
Other and Not Linked to Specific Condition	155.8	111.8	267.7	13.6	18.8	15.3
Total	\$1,150.0	\$594.8	\$1,744.8	100%	100%	100%

* Direct costs are personal health care expenditures for hospital and professional services care, prescribed medications, and home care reported by the Medical Expenditure Panel Survey (MEPS), Agency for Healthcare Research and Quality (AHRQ), by diagnosis, excluding nursing home care costs and costs due to comorbidities.

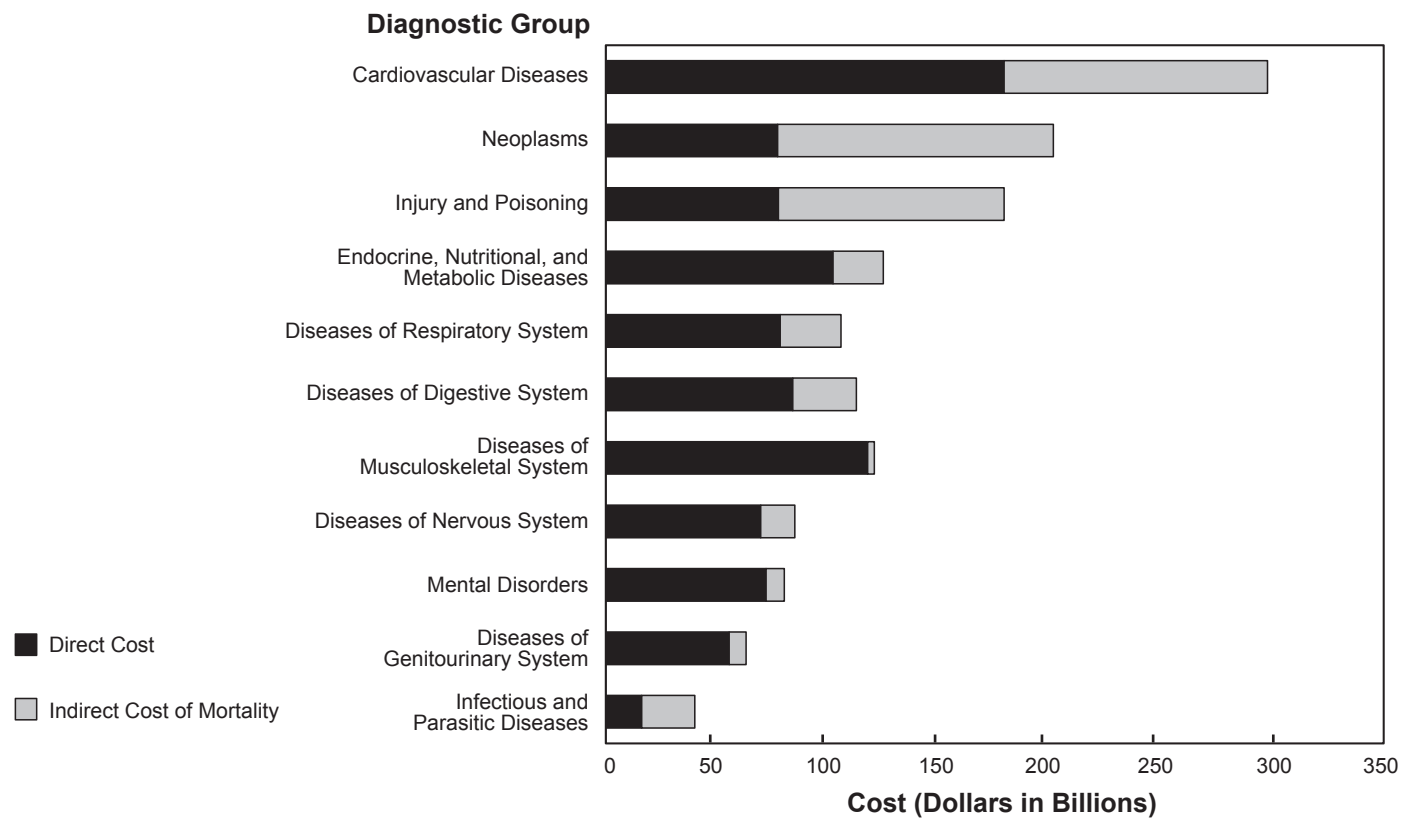
**The mortality cost for each disease group was estimated for 2008 by multiplying the number of deaths by age, sex, and cause of death in 2008 by the 2007 present value of lifetime earnings (latest available) discounted at 3 percent and projected to 2008 based on a 2008 inflation factor measured from mean earnings reported by the U.S. Census Bureau.

† Includes costs for COPD, asthma, and pneumonia.

Note: Estimates are not available for total lung diseases and blood clotting disorders.

Source: Prepared by NHLBI from direct costs on the MEPS Web site; numbers of deaths from Vital Statistics of the United States, NCHS; present value of lifetime earnings from the Institute for Health and Aging, University of California; and mean earnings from the U.S. Census Bureau.

Total Economic Cost of the Leading Diagnostic Groups, U.S., 2008



Source: MEPS, AHRQ.

Direct Economic Cost and Percent Distribution for Selected Conditions by Type of Service, U.S., 2008

Condition	Total Direct Cost (in Billions)	Percent Distribution by Type of Service				
		Hospital Outpatient or Office-Based Provider Visits	Hospital Inpatient Stays	Emergency Room Visits	Prescribed Medicines	Home Health
Heart Disease	\$95.6	18.0%	56.5%	7.6%	10.2%	8.0%
COPD, Asthma	53.7	24.6	24.4	5.7	38.0	7.4
Hypertension	47.4	27.5	13.1	3.6	45.0	10.8
Hyperlipidemia	38.6	23.4	3.5	0.3	70.3	2.6
Stroke	18.8	9.5	48.3	4.9	6.2	31.0
Other Circulatory Conditions	17.6	26.6	59.3	5.1	4.1	4.9
Pneumonia	14.0	5.4	85.3	4.0	1.9	3.4
Anemias	4.7	22.8	67.9	0.7	4.2	4.4

Source: MEPS, Household Component Summary Data Tables, AHRQ.



5. Institute-Initiated Programs Starting in FY 2011

More than two-thirds of the research supported by the NHLBI is initiated by individual investigators; the remainder is initiated by the Institute. Institute-initiated programs are developed in response to evolving national needs, Congressional mandates, and advances in scientific knowledge. Each initiative represents the outcome of extensive discussions and thorough reviews by representatives of the scientific community, Institute advisory committees, the Board of Extramural Experts (BEE), and the National Heart, Lung, and Blood Advisory Council (NHLBAC). The advisory committees and the BEE, together with professional societies and NHLBI staff, continually review the progress of research within the NHLBI program areas, assess newly acquired knowledge, and identify research topics that offer the best opportunities or constitute the greatest needs. This planning process contributes to policy development at the national level by setting priorities among programs and establishing budgets for individual programs and projects.

Initiatives generally emanate as Requests for Applications (RFAs) for grants, including cooperative agreements, or Requests for Proposals (RFPs) for contracts. Other initiatives take the form of Program Announcements (PAs) or Program Announcements with special receipt, referral, or review (PARs). Applications and proposals submitted in response to RFAs and RFPs compete among themselves for specific “set-aside” funds. Applications submitted in response to PAs or PARs generally compete with other investigator-initiated applications for funding.

RFA, RFP, and PA concepts prepared by the Institute are presented to the BEE, which reviews and prioritizes them. The concepts, along with the comments from the BEE, are then sent to the NHLBAC for review, comment, and concurrence. Initiatives that receive the concurrence of the NHLBAC are considered further by the NHLBI Director in the context of the Institute’s budget, program priorities, review workload, and proposed mechanisms. These considerations guide the Director’s subsequent decisions to approve initiatives for release. RFAs, RFPs, PAs, and PARs are

announced in the *NIH Guide to Grants and Contracts*.

Applications and proposals submitted in response to RFAs and RFPs are reviewed by peer review panels convened by the NHLBI. Applications submitted in response to PAs and PARs are reviewed by the NIH Center for Scientific Review.

Descriptions of new or competitively renewed Institute-initiated programs that began (i.e., were first funded) in FY 2011 are presented below according to NHLBI scientific programs. Also described are trans-NIH and trans-PHS initiatives in which the NHLBI participates.

Heart and Vascular Diseases Program

Initiatives Being Renewed

Action To Control Cardiovascular Risk in Diabetes Follow-Up Study (ACCORDION)

The purpose of this renewal is to support a post-trial follow-up study after termination of trial-assigned treatments in the ACCORD trial. The follow-up study will determine whether differences in mortality, CVD events, and microvascular diseases identified during the trial persist or change over time, and whether other differences emerge.

Antihypertensive and Lipid-Lowering Treatment To Prevent Heart Attack Trial (ALLHAT)—Extension

The purpose of this extension is to enable the ALLHAT Coordinating Center to support continued scientific contributions based on extensive ALLHAT data and to expand collaborations with relevant scientific communities.

Atherosclerosis Risk in Communities (ARIC)

The purpose of this renewal is to continue follow-up of the ARIC cohort and community surveillance for the study of CHD, stroke, and heart failure.

Cardiovascular Health Study (CHS): Core Support

The purpose of this renewal is to extend support for the CHS infrastructure to facilitate continued access to study resources and expertise by investigators who are new to the CHS; promote scientific productivity, collaboration, and mentorship of junior investigators; provide analytic support to existing CHS working groups; and assist in the establishment of a new health services research working group.

New Approaches to Arrhythmia Detection and Treatment

The purpose of this renewal is to improve the ability to detect, prevent, and treat all forms of cardiac arrhythmias. This initiative encourages small businesses to develop improved diagnostic and therapeutic tools or devices for cardiac arrhythmia monitoring, detection, and treatment.

Pediatric Heart Network

The purpose of this renewal is to evaluate, through multicenter clinical research, therapeutic and management strategies for children and adults with congenital heart defects and for children with inflammatory heart disease, heart muscle disease, and arrhythmias.

Women's Health Initiative (WHI)

The purpose of this renewal is to enhance the study's ability to investigate outcomes in subgroups and find rarer outcomes; determine long-term effects of clinical trial interventions; maintain and enhance the central database, data analysis capabilities, and biorepository; and conduct research on aging in older women.

New Initiatives

Clinical Research Career Development Programs in Emergency Medicine Research

The purpose of this RFA is to develop multidisciplinary clinical research training programs in emergency medicine that prepare physician-scientists for academic leadership roles and independent research careers in emergency medicine.

Phenotype Finder in Data Resources (PFINDR): A Tool To Support Cross-Study Data Discovery Among NHLBI Genomic Studies

The purpose of this RFA is to develop and apply advanced informatics approaches to categorize phenotypic measures in multiple datasets in data repositories

to help researchers identify potentially relevant genomic studies across cardiovascular, lung, blood, and sleep research domains.

Lung Diseases Program

Initiatives Being Renewed

Lung Tissue Research Consortium

The purpose of this renewal is to continue support for the Consortium to collect, process, and distribute lung tissue and associated clinical data for research that focuses primarily on COPD and idiopathic pulmonary fibrosis.

Severe Asthma Research Program

The purpose of renewing this RFA is to understand the evolution of severe asthma by defining it at the molecular and cellular levels longitudinally. Research findings will serve as a basis for designing mechanism-based diagnostic, prognostic, and treatment strategies for severe asthma in children and adults.

New Initiatives

Centers for Advanced Diagnostics and Experimental Therapeutics in Lung Diseases (CADET) Stage I

The purpose of this RFA is to develop agents for diagnosing and treating lung diseases and sleep-disordered breathing through the use of strategies based on fundamental pathobiologic processes.

Common Pathogenetic Mechanisms of Lung Cancer and COPD

The purpose of this RFA is to investigate common pathogenetic mechanisms of lung cancer and COPD. Scientists will study genotypic and phenotypic characteristics that determine individual susceptibility and shared biochemical, molecular, and immunological pathways that are involved in the origin and progression of both diseases.

Translational Programs in Lung Diseases

The purpose of this PAR is to support collaborative, translational research that will move mechanistic research to clinical applications to improve prevention, diagnosis, and treatment of lung diseases and sleep disorders.

Utilization of a Human Lung Tissue Resource for Vascular Research

The purpose of this RFA is to advance translational efforts in lung vascular disease. Investigators will use

human biospecimens collected by the Pulmonary Hypertension Breakthrough Initiative to determine the pathogenesis of pulmonary arterial hypertension.

Blood Diseases and Resources Program

Initiative Being Renewed

Recipient Epidemiology and Donor Evaluation Study-III (REDS-III)

The purpose of this renewal is to build on and extend the findings of the original REDS and REDS-II programs and address new research areas, including determining safe, effective blood banking and transfusion medicine practices through a comprehensive, multi-targeted strategy involving basic, translational, and clinical research to improve the benefits of transfusion while reducing its risks.

New Initiatives

Innovators in Hemoglobinopathies Care Career Development Award

The purpose of this RFA is to advance the research career development of physician-scientists in SCD and thalassemia. Individuals will address important questions about patient management and translate research results into practice through the use of multidisciplinary teams.

Planning Grants for Pivotal Clinical Trials in Hemoglobinopathies

The purpose of this RFA is to support pilot studies to obtain data that are critical for the design of robust clinical trials in the hemoglobinopathies, SCD, and thalassemias.

Selected Topics in Transfusion Medicine

The purpose of this PAR is to study topics aimed at improving the safety and availability of the blood supply and the practice of transfusion medicine.

Trans-NHLBI

Initiatives Being Renewed

Ancillary Studies in Clinical Trials

The purpose of this renewal is to support time-sensitive ancillary studies related to heart, lung, and blood diseases and sleep disorders in conjunction with ongoing NIH- and non-NIH-supported clinical trials.

Investigator-Initiated Multisite Clinical Trials

The purpose of this renewal is to support investigator-initiated, multisite, Phase II and Phase III randomized controlled clinical trials in areas relevant to the mission of the NHLBI.

Maintenance of NHLBI Biological Specimen Repository

The purpose of this renewal is to continue support for the repository of biologic specimens from NHLBI-sponsored epidemiological and clinical studies. Under the direction of the NHLBI, the repository makes specimens available to the scientific community for use in research related to the NHLBI's mission.

Research Dissemination and Implementation Grants

The purpose of this renewal is to develop and test strategies to accelerate the adoption and integration of efficacious or guideline-based treatments and effective prevention modalities for heart, lung, and blood diseases and sleep disorders in clinical, community, and other settings.

New Initiatives

Cross Organ Mechanism-Associated Phenotypes for Genetic Analyses of Heart, Lung, Blood, and Sleep Diseases (MAPGen for HLBS)

The purpose of this RFA is to identify and characterize common pathobiologic traits and mechanisms that cross organ systems and diseases with the goal of redefining heart, lung, and blood diseases and sleep disorders based on the findings.

MAPGen Knowledge Base and Coordination Center

The purpose of this RFA is to develop and implement a knowledge base and Coordinating Center for the MAPGen for HLBS (above).

Next Generation Genetic Association Studies

The purpose of this RFA is to build on existing genomic study findings with functional information gained by assessing cellular profiles that are surrogates for disease phenotypes.

NHLBI Clinical Trial Pilot Studies

The purpose of this PAR is to support pilot studies to fill gaps in scientific knowledge necessary to develop a competitive full-scale clinical trial to evaluate interventions for the treatment and prevention of heart, lung, and blood diseases and sleep disorders.

NHLBI Research Centers at Minority-Serving Institutions

The purpose of this RFA is to support minority-serving institutions in strengthening scientific faculty; augment predoctoral and postdoctoral science education and research training; and improve research infrastructure, capabilities, and resources in biomedical and behavioral research related to heart, lung, and blood diseases and sleep disorders.

Programs of Excellence in Glycosciences

The purpose of this RFA is to translate emerging discoveries in glycosciences into diagnostics and clinical applications and to build research capacity in glycosciences relevant to heart, lung, and blood diseases.

Science Moving TowArds Research Translation and Therapy Program (SMARTT)

The purpose of this RFP is to assist investigators with the processes needed to translate discoveries in the laboratory into potential new therapies for heart, lung, and blood diseases. The Program will provide manufacturing of synthetic, natural, or biologic products; pharmacology and toxicology testing; preclinical and early phase clinical study design support; and regulatory expertise.

Trans-NIH

Initiatives Being Renewed

Academic Research Enhancement Award (AREA)

The purpose of this renewal is to stimulate support for biomedical, behavioral, or clinical research in educational institutions that provide baccalaureate or advanced degrees for a significant number of the Nation's research scientists, but that have not been major recipients of NIH support.

Advancing Novel Science in Women's Health Research

The purpose of this renewal is to support research that will advance new concepts in women's health and the study of sex differences.

Bioengineering Research Grants

The purpose of this renewal is to support basic and applied multidisciplinary research that addresses biological, bioengineering, or medical research problems. This initiative differs from Bioengineering Research Partnerships (below) in that the research will be performed in a single laboratory by a single investigator or by a small group of investigators.

Bioengineering Research Partnerships

The purpose of this renewal is to support basic, applied, and translational multidisciplinary research that addresses biological, bioengineering, or medical research problems. A partnership of multidisciplinary teams will apply an integrative, systems approach to develop knowledge or methods to prevent, detect, diagnose, or treat disease or to understand health and behavior.

Blood and Marrow Transplant Clinical Research Network

The purpose of this renewal is to evaluate innovative treatment approaches and management strategies for children and adults who are undergoing hematopoietic stem cell transplantation. The focus of the initiative is to support multicenter Phase II and Phase III studies that provide information to improve hematopoietic stem cell transplantation therapy.

Diet Composition and Energy Balance

The purpose of this renewal is to investigate the role of diet composition in energy balance. Scientists are seeking to understand the impact of micro- or macronutrient composition on appetite, metabolism, and energy expenditure and to determine the efficacy of diets that differ in micro- or macronutrient composition, absorption, dietary variety, or energy density on weight loss and weight maintenance.

Dissemination and Implementation Research in Health

The purpose of this renewal is to support innovative approaches to overcoming barriers to adopting and integrating evidence-based interventions and guidelines that previous research has shown to be effective but where implementation has been limited or significantly delayed.

Exploratory Collaborations With National Centers for Biomedical Computing

The purpose of this renewal is to support investigators who are collaborating with NIH Roadmap National Centers for Biomedical Computing to engage in exploratory biomedical informatics and computational biology research.

Exploratory Innovations in Biomedical Computational Science and Technology

The purpose of this renewal is to promote research and development in biomedical informatics and computational

biology that will support rapid progress in areas of scientific opportunity in biomedical research.

Innovations in Biomedical Computational Science and Technology Initiative

The purpose of this renewal is to solicit STTR grant applications that propose innovative research in biomedical informatics and computational biology to promote the progress of biomedical research.

Methodology and Measurement in the Behavioral and Social Sciences

The purpose of this renewal is to improve the quality of data collected in the behavioral and social sciences relevant to the mission of the participating NIH Institutes.

Mouse Metabolic Phenotyping Centers Consortium

The purpose of this renewal is to provide the scientific community with standardized, high-quality metabolic and physiologic phenotyping services for mouse models of diabetes, diabetic complications, obesity, and related disorders.

Understanding and Promoting Health Literacy

The purpose of this renewal is to encourage research on health literacy concepts, theory, and interventions as they pertain to prevention, healthy living, chronic disease management, patient-based health care, cultural competence, and health disparities. Researchers will seek to develop health literacy assessment methods, measure association between health literacy and chronic illness, and evaluate health literacy interventions that are designed to improve health outcomes in heart, lung, and blood diseases and enhance understanding and use of genetic information in risk discussion.

New Initiatives

Biophysical and Biomechanical Aspects of Embryonic Development

The purpose of this PAR is to encourage innovative and high risk/impact research in the area of physics/mechanics of embryonic development in model organisms. The focus of the initiative is to promote research that will generate information about tissue mechanics relevant to vertebrate development and will improve the understanding of the basis for developmental disorders.

Deepwater Horizon Disaster Research Consortia

The purpose of this RFA is to investigate health outcomes from exposure to the Deepwater Horizon disaster, including (a) identifying those at greatest risk for adverse physical, mental, and behavioral effects as a consequence of the disaster and other stressors encountered by residents and (b) developing community-based approaches to respond to it and future disasters.

Development of Multifunctional Drug and Gene Delivery Systems

The purpose of this PAR is to develop multifunctional drug and gene delivery systems that can target therapies to particular cells and intracellular compartments, monitor delivery, and determine therapeutic efficacy through integration of advanced imaging or sensing technologies into the delivery system.

Knockout Mouse Phenotyping Program (KOMP2)

The purpose of this RFA is to support a systematic program to generate live mice from embryonic cells that contain individual, single-gene deletion of known and predicted protein-encoding genes in the mouse genome, which was successfully generated by the Knockout Mouse Project (KOMP) in collaboration with the International Knockout Mouse Consortium.

Lifestyle Interventions in Overweight and Obese Pregnant Women Consortium

The purpose of this RFA is to evaluate lifestyle interventions in overweight and obese pregnant women that are designed to improve weight and metabolic outcomes in both the pregnant women and their offspring.

New Strategies for Growing 3D Tissues

The purpose of this RFA is to understand how cells respond to their environment and develop assays and methods to understand how angiogenesis and organogenesis may instruct the creation of 3D-engineered cellular aggregates.

Safe and Effective Instruments and Devices for Use in the Neonatal Intensive Care Units

The purpose of this SBIR/STTR is to develop or improve safe and effective instruments for monitoring and treating newborn and small children.

Sleep and Social Environment: Basic Biopsychosocial Processes

The purpose of this RFA is to investigate the reciprocal interactions of the processes of sleep and circadian regulation and function with behavioral and social environment processes.

Social Network Analysis and Health

The purpose of this PAR is to advance and expand the utility of social network analysis and methods in studies of health and disease. The goal is to encourage basic research that can lead to new applications of social network methods and theory related to improving human health.

Trans-PHS

New Initiatives

Community-Based Partnerships for Childhood Obesity Prevention and Control Research

The purpose of this PA is to stimulate childhood obesity research through community-based partnerships that

include local, state, and regional teams of researchers, policymakers, and other relevant participants—such as community representatives, public health practitioners, and educators.

Obesity Policy Research: Evaluation and Measures

The purpose of this PA is to conduct evaluation research on obesity-related “natural experiments” (i.e., community and other population-level public policy interventions that may affect diet and physical activity behavior) and develop and validate relevant community-level measures (i.e., instruments and methodologies to assess the food and physical activity environments).



6. Institute Public Advisory Committees

National Heart, Lung, and Blood Advisory Council

Structure

Chair: Susan B. Shurin, M.D., Acting Director, NHLBI

Executive Secretary: Stephen C. Mockrin, Ph.D., Director, Division of Extramural Research Activities, NHLBI, National Institutes of Health, Bethesda, MD 20892; 301-435-0260

The Secretary of HHS appoints 18 members: 12 members are leading representatives of the health and scientific disciplines (including public health and behavioral or social sciences), and 6 are from the general public and are leaders in the fields of public policy, law, health policy, economics, and management.

Members are appointed for overlapping terms of 4 years.

The Council includes the following ex officio members:

- Secretary, HHS
- Director, NIH
- Director, NHLBI
- Chief Medical Director, Veterans Affairs, or Designee
- Assistant Secretary of Defense for Health Affairs, or Designee
- Designee, Centers for Disease Control and Prevention

Functions

The NHLBAC reviews applications for research grants, cooperative agreements, and training grants in heart, blood vessel, lung, and blood diseases;

sleep disorders; and blood resources, and recommends scientific projects that merit support to the Director, NHLBI.

The Council advises the Secretary, HHS; the Assistant Secretary for Health, HHS; and the Directors, NIH and NHLBI on matters relating to causes, prevention, diagnosis, and treatment of diseases and resources within the purview of the Institute. The Council also may review any grant, contract, or cooperative agreement proposed to be made or entered into by the Institute; may make recommendations to the Director of the Institute respecting research conducted at the Institute; may collect, by correspondence or by personal investigation, information as to studies that are being carried on in the United States or any other country with respect to the cause, prevention, diagnosis, and treatment of heart, blood vessel, lung, and blood diseases, and to the use of blood and blood products and the management of blood resources 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 may assemble ad hoc working groups, appoint subcommittees, and convene workshops and conferences.

The Council may also make recommendations to the Director, NIH and other authorized officials regarding the acceptance of conditional gifts pursuant to section 231 of the Public Health Service Act, as amended.

Meetings

The Chair convenes meetings not fewer than four times a year and approves the agenda.

National Heart, Lung, and Blood Advisory Council Membership*

Susan B. Shurin, M.D.

Chair

National Heart, Lung, and Blood Institute

Jonathan R. Alger, J.D. (2014)

Rutgers, The State University of New Jersey

C. Noel Bairey Merz, M.D. (2011)

Cedars-Sinai Medical Center

Coletta C. Barrett, R.N. (2014)

Mission for Our Lady of the Lake Regional Medical Center

Ivor J. Benjamin, M.D. (2014)

University of Utah School of Medicine

Ingrid B. Borecki, Ph.D. (2012)

Washington University in St. Louis

Barry S. Coller, M.D. (2012)

The Rockefeller University

Jack A. Elias, M.D. (2012)

Yale University School of Medicine

Gary H. Gibbons, M.D. (2013)

Morehouse School of Medicine

Beverly W. Hogan (2012)

Tougaloo College

Lanetta B. Jordan, M.D. (2013)

Memorial Healthcare System

Talmadge E. King, M.D. (2013)

University of California, San Francisco

Naomi L.C. Luban, M.D. (2014)

The George Washington University School of Medicine

Andrew R. Marks, M.D. (2011)

Columbia University

Michael S. Parmacek, M.D. (2012)

University of Pennsylvania School of Medicine

Polly E. Parsons, M.D. (2014)

University of Vermont

Marlene Rabinovitch, M.D. (2011)

Stanford University

Leslee J. Shaw, Ph.D. (2013)

Emory University

Gilbert C. White II, M.D. (2014)

Blood Center of Wisconsin

Ex Officio Members

Francis S. Collins, M.D., Ph.D.

National Institutes of Health

Robert L. Jesse, M.D., Ph.D.

Veterans Health Administration

Kathleen Sebelius, M.P.A.

Department of Health and Human Services

* Current as of October 2011. The current roster, containing full addresses for the NHLBI Advisory Council and Committees, can be obtained from the Internet at <http://www.nhlbi.nih.gov/meetings/nhlbac/roster.htm>.

Program Advisory and Review Committee

Sickle Cell Disease Advisory Committee

Chair: Edward J. Benz, M.D., Dana-Farber Cancer Institute

Executive Secretary: W. Keith Hoots, M.D., Director, Division of Blood Diseases and Resources, NHLBI, National Institutes of Health, Bethesda, MD 20892; 301–435–0080

The Sickle Cell Disease Advisory Committee advises the Directors of the NIH, the NHLBI, and the DBDR on matters related to planning, executing, conducting, supporting, and evaluating research in SCD.

Membership*

Thomas D. Coates, M.D. (2013)
University of Southern California

Elaine K. Gallin, Ph.D. (2012)
Doris Duke Charitable Foundation

Nigel S. Key, M.D. (2013)
University of North Carolina at Chapel Hill

Roberto F. Machado, M.D. (2012)
University of Illinois, Chicago

Punam Malik, M.D. (2012)
Cincinnati Children's Hospital Medical Center

Leslie V. Parise, Ph.D. (2012)
University of North Carolina

Ex Officio Members

Francis S. Collins, M.D., Ph.D.
National Institutes of Health

William Hannon, Ph.D.
Centers for Disease Control and Prevention

Marie Y. Mann, M.D.
Health Resources and Services Administration

David E. McCune, M.D.
Madigan Army Medical Center

Sleep Disorders Research Advisory Board

Chair: Michael V. Vitiello, Ph.D., University of Washington

Executive Secretary: Michael J. Twery, Ph.D., Director, National Center on Sleep Disorders Research, NHLBI, National Institutes of Health, Bethesda, MD 20892; 301–435–0202

The Sleep Disorders Research Advisory Board advises the Directors of the NIH, the NHLBI, and the NCSDR on matters related to planning, executing, conducting, supporting, and evaluating research in sleep disorders.

Membership*

Mercedes R. Carnethon, Ph.D., M.P.H. (2013)
Northwestern University

Ronald D. Chervin, M.D. (2012)
University of Michigan

Leszek K. Kubin, Ph.D. (2014)
University of Pennsylvania

Sairam Parthasarathy, M.D. (2014)
University of Arizona

R. Daniel Rudic, Ph.D. (2012)
Medical College of Georgia

Catherine Vena, Ph.D., R.N. (2013)
Emory University

Gagandeep Walia, M.B.A. (2013)
Visa, Inc.

Ex Officio Members

Thomas J. Balkin, Ph.D.
Walter Reed Army Institute of Research

Francis S. Collins, M.D., Ph.D.
National Institutes of Health

Robert W. Greene, M.D., Ph.D.
Veterans Administration, North Texas Medical Center

* Current as of October 2011.

Rosalind King, Ph.D.
NICHD, National Institutes of Health

Miroslav Mackiewicz, Ph.D.
NIA, National Institutes of Health

Merrill M. Mitler, Ph.D.
NINDS, National Institutes of Health

Michael J. Twery, Ph.D.
NHLBI, National Institutes of Health

Aleksandra Vicentic, Ph.D.
NIMH, National Institutes of Health

Heart, Lung, and Blood Initial Review Group

Scientific Review Officer: Jeffery H. Hurst, Ph.D.,
Scientific Review Officer, Division of Extramural
Research Activities, NHLBI, National Institutes of
Health, Bethesda, MD 20892; 301–435–0303

The Heart, Lung, and Blood Initial Review Group provides initial technical merit review for the NHLBAC and the Director, NHLBI. This group consists of three subcommittees: the Heart, Lung, and Blood Program Project Review Committee; the Clinical Trials Review Committee; and the NHLBI Institutional Training Mechanism Review Committee.

Heart, Lung, and Blood Program Project Review Committee

Chair: David J. Pinsky, M.D., University of Michigan

Scientific Review Officer: Jeffery H. Hurst, Ph.D.,
Scientific Review Officer, Division of Extramural
Research Activities, NHLBI, National Institutes of
Health, Bethesda, MD 20892; 301–435–0303

The Heart, Lung, and Blood Program Project Review Committee provides initial technical merit review for the NHLBAC and the Director, NHLBI on program project applications proposing research in the areas of heart, lung, and blood diseases and resources.

Membership*

David Atkinson, Ph.D. (2012)
Boston University School of Medicine

Peter M. Buttrick, M.D. (2014)
University of Colorado

Jennifer L. Hall, Ph.D. (2015)
University of Minnesota

Thomas H. Hintze, Ph.D. (2014)
New York Medical College

Pedro A. Jose, M.D., Ph.D. (2015)
Children's National Medical Center

Jay K. Kolls, M.D. (2014)
University of Pittsburgh School of Medicine

Monica Kraft, M.D. (2014)
Duke University Medical Center

Lucy Liaw, Ph.D. (2014)
Maine Medical Center Research Institute

Nigel Mackman, Ph.D. (2013)
University of North Carolina at Chapel Hill

Catherine S. Manno, M.D. (2013)
New York University School of Medicine

Fernando D. Martinez, M.D. (2013)
University of Arizona

Christine S. Moravec, Ph.D. (2012)
The Cleveland Clinic Foundation

Nanduri R. Prabhakar, Ph.D., D.Sc. (2012)
University of Chicago

Frank C. Sciurba, M.D. (2013)
University of Pittsburgh

Nancy S. Speck, Ph.D. (2013)
University of Pennsylvania School of Medicine

James C. Zimring, M.D., Ph.D. (2015)
Emory University School of Medicine

* Current as of October 2011.

Clinical Trials Review Committee

Chair: Julio A. Panza, M.D., Washington Hospital Center

Scientific Review Officer: Keary A. Cope, Ph.D., Scientific Review Officer, Division of Extramural Research Activities, NHLBI, National Institutes of Health, Bethesda, MD 20892; 301–435–2222

The Clinical Trials Review Committee provides initial technical merit review for the NHLBAC and the Director of the NHLBI on clinical trial applications for the support of studies to evaluate preventive or therapeutic measures of blood, cardiovascular, and lung diseases.

Membership*

Roberta A. Ballard, M.D. (2014)
University of California, San Francisco

Bruce A. Barton, Ph.D. (2014)
University of Massachusetts Medical School

Jeffrey L. Carson, M.D. (2014)
University of Medicine and Dentistry of New Jersey

Mina K. Chung, M.D. (2014)
Case Western Reserve University

Gerard J. Criner, M.D. (2014)
Temple University School of Medicine

Karen L. Margolis, M.D. (2012)
Health Partners Research Foundation

Lori J. Mosca, M.D., Ph.D. (2013)
Columbia University Medical Center

Steven Piantadosi, M.D., Ph.D. (2013)
Cedars-Sinai Medical Center

Arshed A. Quyyumi, M.D. (2013)
Emory University School of Medicine

Gary E. Raskob, Ph.D. (2015)
University of Oklahoma Health Sciences Center

Margaret M. Redfield, M.D. (2013)
Mayo Clinic

Madeline M. Rice, Ph.D. (2013)
The George Washington University

Amy D. Shapiro, M.D. (2012)
Indiana Hemophilia and Thrombosis Center, Inc.

Stanley J. Szefler, M.D. (2012)
National Jewish Health

Barbara C. Tilley, Ph.D. (2015)
University of Texas

O. Dale Williams, Ph.D. (2015)
Florida International University

NHLBI Institutional Training Mechanism Review Committee

Chair: William C. Balke, M.D., University of California, San Francisco

Scientific Review Officer: Charles Joyce, Ph.D., Scientific Review Officer, Division of Extramural Research Activities, NHLBI, National Institutes of Health, Bethesda, MD 20892; 301–435–0291

The NHLBI Institutional Training Mechanism Review Committee provides initial technical merit review for the NHLBAC and the Director of the NHLBI on training applications that provide predoctoral, postdoctoral, and short-term research training at academic institutions.

Membership*

Ifeanyi J. Arinze, Ph.D. (2012)
Meharry Medical College

Jennifer K. Barton, Ph.D. (2015)
University of Arizona

Neil Blumberg, M.D. (2015)
University of Rochester

Meredith Bond, Ph.D. (2013)
University of Maryland School of Medicine

Kathleen B. Brosnihan, Ph.D. (2015)
Wake Forest University

Linda J. Burns, M.D. (2013)
University of Minnesota

* Current as of October 2011.

David M. Center, M.D. (2013)
Boston University School of Medicine

Mark W. Geraci, M.D. (2015)
University of Colorado

David M. Guidot, M.D. (2015)
Emory University

Gerardo Heiss, M.D., Ph.D. (2014)
University of North Carolina at Chapel Hill

Kirk U. Knowlton, M.D. (2013)
University of California, San Diego

Robert J. Levy, M.D. (2014)
The Children's Hospital of Philadelphia

Alice H. Lichtenstein, D.Sc. (2013)
Tufts University

Russell V. Luepker, M.D. (2012)
University of Minnesota

Jonathan C. Makieleski, M.D. (2012)
University of Wisconsin Hospitals and Clinics

William H. Pearce, M.D. (2014)
Northwestern University

Michael I. Phillips, Ph.D., D.Sc. (2014)
Keck Graduate Institute

Josef T. Prchal, M.D. (2012)
University of Utah

Robin Shandas, Ph.D. (2012)
University of Colorado Health Sciences Center

Kingman P. Strohl, M.D. (2015)
University Hospitals of Cleveland

Mary I. Townsley, Ph.D. (2012)
University of South Alabama

Carolyn F. Whitsett, M.D. (2015)
Kings County Hospital

Jo Rae Wright, Ph.D. (2014)
Duke University

National Heart, Lung, and Blood Institute Special Emphasis Panel

The Institute established the NHLBI Special Emphasis Panel (SEP) to carry out the initial peer review of applications and proposals that were previously handled by ad hoc committees. Concept review, previously handled by divisional program advisory committees, has also been incorporated into the SEP system. The SEP, which has neither a fixed membership nor a set meeting schedule, is constituted to provide required peer review expertise at precisely the time that it is needed.

Board of Scientific Counselors

Chair: Michael I. Kotlikoff, V.M.D., Ph.D.,
Cornell University

Executive Secretary: Robert S. Balaban, Ph.D.,
Director, Division of Intramural Research, NHLBI,
National Institutes of Health, Bethesda, MD 20892;
301-496-2116

The Board of Scientific Counselors advises the Director and the Deputy Director for Intramural Research, NIH, and the Directors of NHLBI and the Division of Intramural Research, NHLBI, on the intramural research programs of the NHLBI.

Membership*

Grover C. Bagby, M.D. (2015)
Oregon Health Sciences University

Kenneth R. Chien, M.D., Ph.D. (2012)
Massachusetts General Hospital

Serpil C. Erzurum, M.D. (2015)
The Cleveland Clinic Foundation

James F. Greenleaf, Ph.D. (2015)
Mayo Clinic College of Medicine

Aldons J. Lusis, Ph.D. (2013)
University of California, Los Angeles

David J. Sahn, M.D. (2014)
Oregon Health and Science University

Douglas C. Wallace, Ph.D. (2014)
University of California, Irvine

* Current as of October 2011.



7. Fiscal Year 2011 Budget Overview

NHLBI Obligations by Funding Mechanism: Fiscal Year 2011

Funding Mechanism	Obligated Dollars* (Thousands)	Percent of Total NHLBI Budget
Research Project Grants**	\$2,079,920	67.8%
SCCORs and P50 Centers	34,365	1.5
Sickle Cell Centers	5,901	0.2
Centers for AIDS Research	3,334	0.1
Other Research Centers Grants	12,331	0.4
Other Research Grants	152,772	5.0
<i>Research Careers Programs</i> [†]	79,090	2.6
Training Programs	97,998	3.2
Research and Development Contracts	368,291	12.0
Intramural Laboratory and Clinical Research	192,083	6.1
Research Management and Support [‡]	122,554	4.0
Total Obligations	\$3,069,549	100.0%

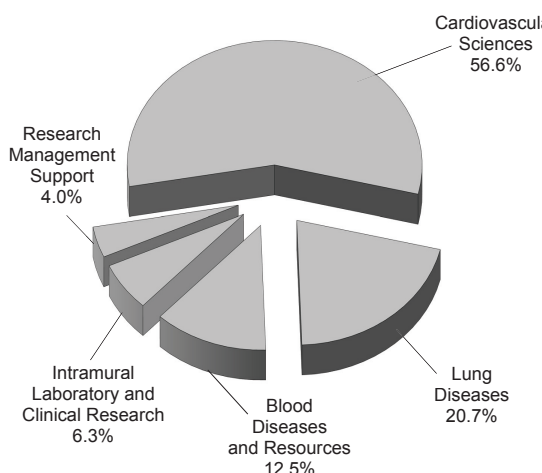
* Excludes funds provided by other Agencies by means of a reimbursable agreement.

** Includes \$74,935 for Small Business Innovation Research (SBIR) Grants/Small Business Technology Transfer (STTR) Grants.

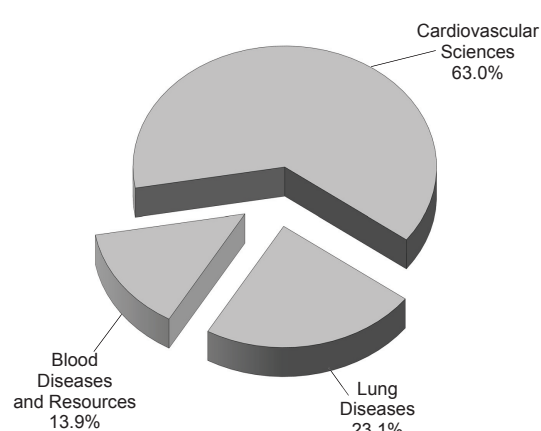
† Research Career Programs are a subset of Other Research Grants and are not added as a distinct funding mechanism.

‡ Excludes OD and DIR research contracts, which are included in R&D contracts.

NHLBI Total Obligations by Budget Category



NHLBI Extramural Obligations by Program



For detailed data on FY 2011:

- Research grants, see Chapters 9 and 11.
- Research and development contracts, see Chapters 10 and 11.
- Research training and career development, see Chapter 13.
- Geographic distribution of awards, see Chapter 14.

NHLBI Extramural Obligations by Program: Fiscal Year 2011

Program	Obligated Dollars (Thousands)	Percent of NHLBI Extramural Budget
Cardiovascular Sciences	\$1,736,223	63.0%
Lung Diseases	635,493	23.1
Blood Diseases and Resources	383,196	13.9
Total, Extramural Obligations	\$2,754,912	100%

NHLBI Cardiovascular Sciences Program

Obligations by Funding Mechanism: Fiscal Year 2011

Funding Mechanism	Obligated Dollars (Thousands)	Percent of Program Budget
Research Project Grants	\$1,281,756	73.8%
SCCOR and P50 Centers	11,928	0.7
Other Research Centers Grants	12,180	0.7
Other Research Grants	73,283	4.2
<i>Research Career Programs*</i>	39,949	2.3
Training Programs	57,876	3.3
Research and Development Contracts	299,200	17.2
Total, Cardiovascular Sciences	\$1,736,223	100%

NHLBI Lung Diseases Program

Obligations by Funding Mechanism: Fiscal Year 2011

Funding Mechanism	Obligated Dollars (Thousands)	Percent of Program Budget
Research Project Grants	\$504,070	79.3%
SCCORs and P50 Centers	22,437	3.5
Other Research Centers Grants	151	0.0
Other Research Grants	57,606	9.1
<i>Research Career Programs*</i>	25,844	4.1
Training Programs	25,891	4.1
Research and Development Contracts	25,338	4.0
Total, Lung Diseases	\$635,493	100%

NHLBI Blood Diseases and Resources Program

Obligations by Funding Mechanism: Fiscal Year 2011

Funding Mechanism	Obligated Dollars (Thousands)	Percent of Program Budget
Research Project Grants	\$294,094	76.7%
Sickle Cell Centers	5,901	1.5
Centers for AIDS Research	3,334	0.9
Other Research Grants	21,884	5.7
<i>Research Career Programs*</i>	13,297	3.5
Training Programs	14,231	3.7
Research and Development Contracts	43,752	11.4
Total, Blood Diseases and Resources	\$383,196	100%

* Research Career Programs are a subset of Other Research Grants and are not added as a distinct funding mechanism.



8. Long-Term Trends

Budget History of the NHLBI: Fiscal Years 1950–2011

Dollars (Thousands)

Fiscal Year	Budget Estimate to Congress	House Allowance	Senate Allowance	Appropriation	Obligations	Cumulative Fiscal Year Obligations
1950	\$ 34,630	\$ 11,575	\$ 29,117	\$ 16,075	\$ 15,768	\$ 15,768
1951	8,800	8,800	9,400	9,400	8,497	24,265
1952	10,237	10,074	10,156	10,083	9,850	34,115
1953	9,779	9,623	12,000	12,000	11,398	45,513
1954	11,040	12,000	15,418	15,168	14,952	60,465
1955	14,570	16,168	17,168	16,668	16,595	77,060
1956	17,454	17,398	23,976	18,808	18,838	95,898
1957	22,106	25,106	33,396	33,396	32,392	128,290
1958	33,436	33,436	38,784	35,936	35,973	164,263
1959	34,820	36,212	49,529	45,613	45,468	209,731
1960	45,594	52,744	89,500	62,237	61,565	271,296
1961	63,162	71,762	125,166	86,900	86,239	357,535
1962	97,073	105,723	160,000	132,912	110,849	468,384
1963	126,898	143,398	149,498	147,398	120,597	588,981
1964	130,108	129,325	130,545	132,404	117,551	706,532
1965	125,640	124,521	125,171	124,824	124,412	830,944
1966	141,412	146,212	143,462	141,462	141,171	972,115
1967	148,407	154,770	164,770	164,770	164,342	1,136,457
1968	167,954	167,954	177,954	167,954	162,134	1,298,591
1969	169,735	164,120	172,120	166,928	161,834	1,460,425
1970	160,513	160,513	182,000	171,257	160,433	1,620,858
1971	171,747	178,479	203,479	194,901	194,826	1,815,684
1972	195,492	211,624	252,590	232,627	232,577	2,048,261
1973	255,280	300,000	350,000	300,000	255,722	2,303,983
1974	265,000	281,415	320,000	302,915	327,270	2,631,253
1975	309,299	321,196	330,000	327,996	327,953	2,959,206
1976	324,934	329,079	379,059	370,096	368,648	3,327,854
TQ ^A	59,715	58,015	58,015	58,763	60,639	3,388,493
1977	342,855	380,661	420,661	396,661	396,857	3,785,350
1978	403,642	432,642	456,000	447,901	447,968	4,233,318
1979	454,336	485,584	485,584	510,134	510,080	4,743,398
1980	507,344	527,544	527,544	527,544	527,248	5,270,646
1981	532,799	560,264	565,264	549,693	550,072	5,820,718
1982	579,602	583,831	587,741	559,637	559,800	6,380,518
1983	577,143	620,947	624,542	624,259	624,260	7,004,778
1984	639,774	665,859	683,489	704,939	705,064	7,709,842
1985	718,852	764,135	807,149	805,269	803,810	8,513,652
1986	775,254	856,388	863,652	859,239	821,901	9,335,553
1987	785,697	921,410	921,502	930,001	929,982	10,265,535
1988	821,887	990,808	1,000,349	965,536	965,283	11,230,818
1989	1,054,503	1,018,983	1,056,003	1,045,985	1,045,508	12,276,326
1990	1,039,846	1,090,930	1,091,597	1,072,354	1,070,683	13,347,009

A TQ=Transition Quarter, July 1–September 30, 1976.

Budget History of the NHLBI: Fiscal Years 1950–2011 (Continued)

Dollars (Thousands)

Fiscal Year	Budget Estimate to Congress	House Allowance	Senate Allowance	Appropriation	Obligations	Cumulative Fiscal Year Obligations
1991	1,112,502	1,135,589	1,137,235	1,126,942	1,125,915	14,472,924
1992	1,209,924	1,202,398	1,190,396	1,191,500	1,190,070	15,662,994
1993	1,245,396	1,228,455	1,228,455	1,214,693	1,214,693	16,877,687
1994	1,198,402	1,277,880	1,277,880	1,277,880	1,277,852	18,155,539
1995	1,266,961	1,259,590	1,259,590	1,258,472	1,314,969	19,470,508
1996	1,337,021	1,355,866	1,320,254 ^B	1,355,866	1,351,422 ^C	20,821,930
1997	1,320,555 ^D	1,438,265	1,344,742 ^D	1,432,529 ^E	1,431,821	22,253,751
1998	1,467,189	1,513,004	1,531,898	1,531,061 ^F	1,526,276	23,780,027
1999	1,709,328 ^G	1,720,344	1,793,697	1,793,697 ^F	1,788,008	25,568,035
2000	1,759,806	1,937,404	2,001,185	2,040,291 ^F	2,027,286	27,595,321
2001	2,069,582	2,328,102	2,328,105	2,299,866 ^H	2,298,035	29,893,356
2002	2,567,429	2,547,675	2,618,966	2,576,125 ^I	2,569,794	32,463,150
2003	2,791,411	2,812,011	2,818,684	2,812,011 ^J	2,793,681	35,256,831
2004	2,867,995	2,867,995	2,897,595	2,882,715 ^K	2,882,601	38,139,432
2005	2,963,953	2,963,953	2,985,900	2,965,453	2,922,573 ^L	41,062,005
2006	2,951,270	2,951,270	3,023,381	2,951,270 ^J	2,893,527	43,955,532
2007	2,901,012	2,901,012	2,924,299	2,921,757	2,922,322 ^L	46,877,854
2008	2,894,341	2,965,775	2,992,197	2,974,900 ^E	2,937,333	49,815,187
2009	2,924,942	3,025,500	3,006,344	3,015,689	3,014,552	52,829,739
2010	3,050,356	3,123,403	3,066,827	3,096,916	3,093,501	55,923,240
2011	3,187,516	— ^M	3,182,524	3,069,723	3,069,550	58,992,790

B Senate Allowance reflects the Institute share of the Government-wide rescission and the HHS rescission.

G Includes Bioterrorism reduction.

C Obligations reflect the Institute share of the Government-wide rescission, the HHS rescission, and a transfer to other NIH Institutes through the NIH Director's 1 percent transfer authority.

H Excludes Office of Human Research Protection transfer, Secretary transfer, and rescission.

D Excludes funds for AIDS research activities consolidated in the NIH Office of AIDS Research (OAR).

I Excludes Government-wide rescission, Labor/HHS/Education rescission, from HHS to OMB rescission, and Secretary 1 percent transfer.

E Excludes enacted administrative reduction.

J Excludes Government-wide rescission.

F Excludes Director transfer, Secretary transfer, and rescission.

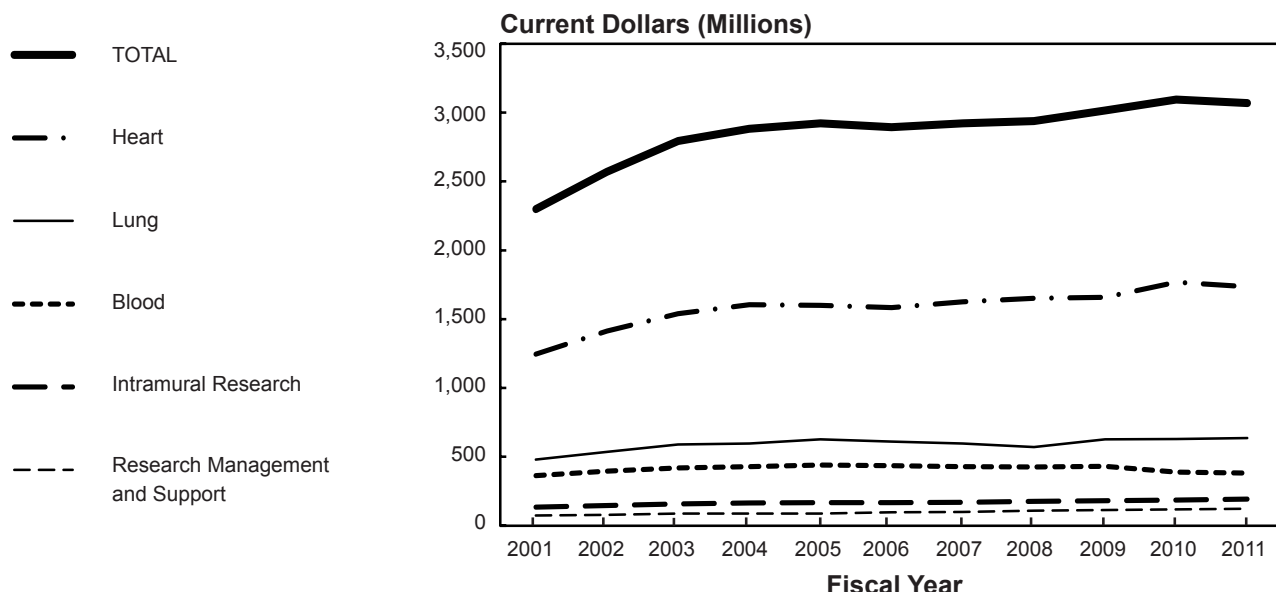
K Includes Roadmap adjustments.

L Includes Roadmap Transfer and Government-wide rescission.

M No House allowance provided in FY 2011.

NHLBI Total Obligations by Budget Category: Fiscal Years 2001–2011

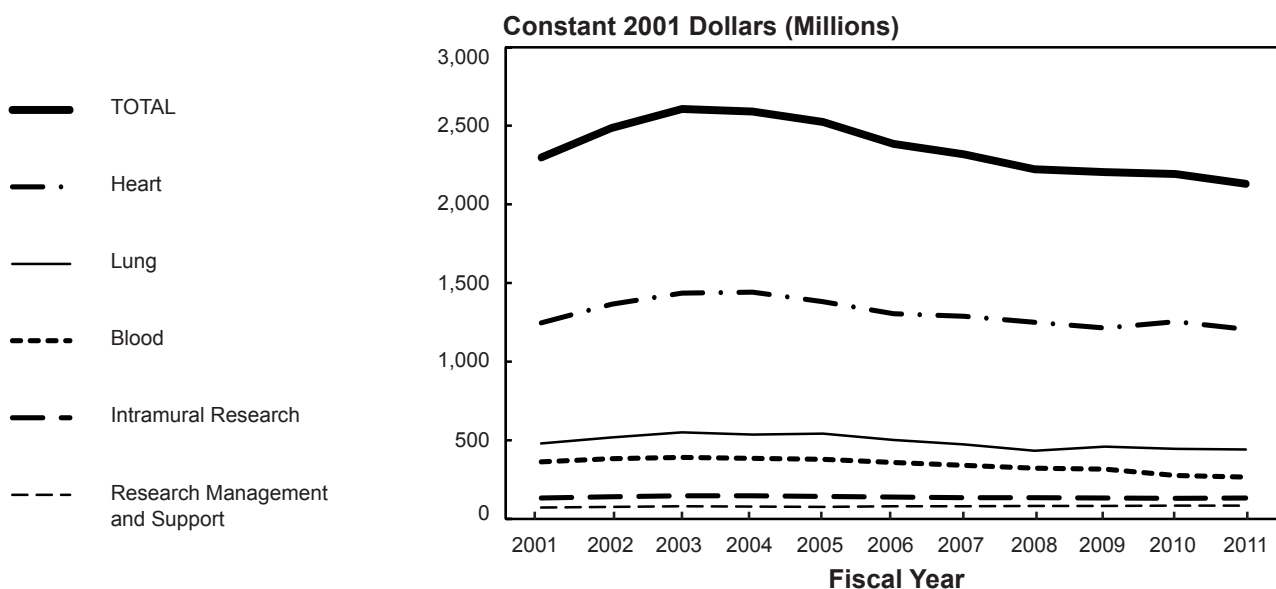
Current Dollars



Note: Beginning in 2007, the WHI funds are included in the “Heart” category and the Sleep Disorders Research funds are included in the “Lung” category. Previously they were reported separately.

NHLBI Total Obligations by Budget Category: Fiscal Years 2001–2011*

Constant 2001 Dollars



* This chart is based on the Biomedical Research & Development Price Index through 2010.

Note: Beginning in 2007, the WHI funds are included in the “Heart” category and the Sleep Disorders Research funds are included in the “Lung” category. Previously they were reported separately.

NHLBI Total Obligations by Budget Category: Fiscal Years 2001–2011

Budget Category	Current Dollars (Millions)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Extramural Research											
Heart	\$1,245.8	\$1,412.4	\$1,538.8	\$1,604.7	\$1,599.6	\$1,582.7	\$1,624.9	\$1,652.2	\$1,659.2	\$1,769.1	\$1,736.2
Lung	481.0	535.2	590.5	596.0	628.2	610.3	597.6	572.2	627.8	629.9	635.5
Blood	364.0	396.0	419.3	429.2	439.5	434.9	429.7	426.2	431.7	389.2	383.2
Intramural Research	133.7	146.7	157.8	164.2	166.3	168.3	169.5	177.5	181.7	186.2	192.1
Research Management and Support	73.5	79.4	87.3	88.5	89.0	97.2	100.6	109.2	114.1	119.1	122.6
Total	\$2,298.0	\$2,569.7	\$2,793.7	\$2,882.6	\$2,922.6	\$2,893.4	\$2,922.3	\$2,937.3	\$3,014.5	\$3,093.5	\$3,069.6

Note: From 1999 to 2006, the WHI was reported separately. In this table, it has been incorporated into the “Heart” line. The Sleep Disorders Research was reported separately from 1996 to 2006. In this table, it has been incorporated into the “Lung” line.

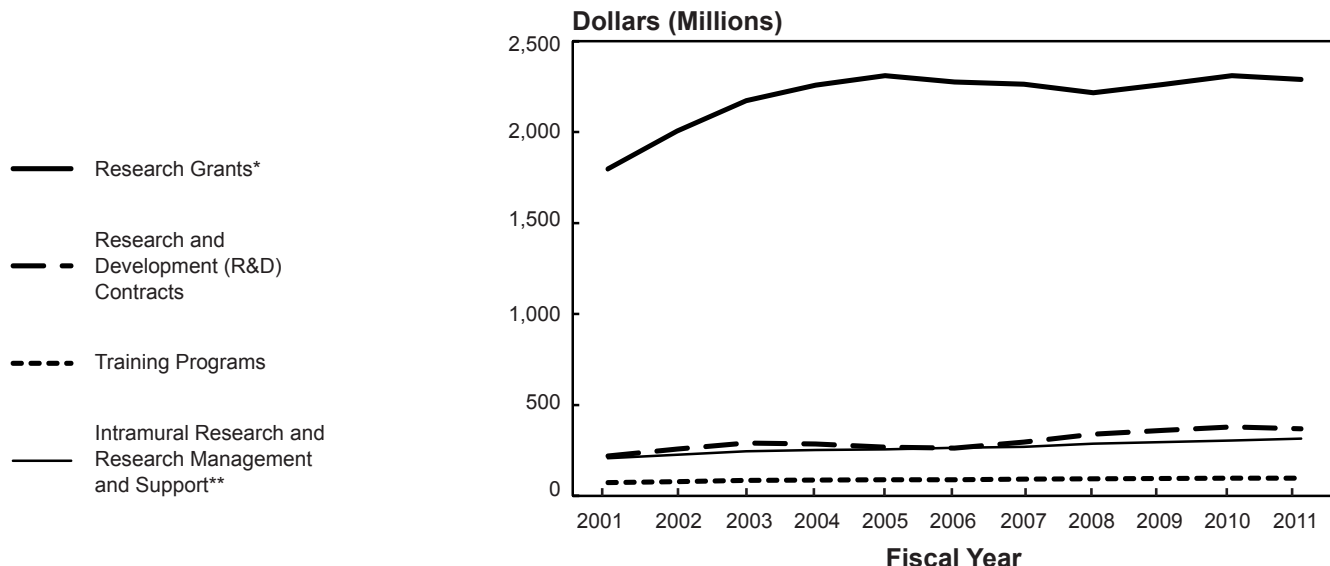
NHLBI Total Obligations by Budget Category: Fiscal Years 2001–2011

Budget Category	Constant 2001 Dollars (Millions)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Extramural Research											
Heart	\$1,245.8	\$1,366.0	\$1,435.4	\$1,441.8	\$1,381.3	\$1,304.8	\$1,289.6	\$1,250.7	\$1,213.8	\$1,253.8	\$1,205.3
Lung	481.0	517.6	550.8	535.5	542.5	503.1	474.3	433.2	459.3	446.4	441.9
Blood	364.0	383.0	391.1	385.6	379.5	358.6	341.0	322.6	315.8	275.8	265.9
Intramural Research	133.7	141.9	147.2	147.5	143.6	138.7	134.5	134.4	132.9	132.0	133.4
Research Management and Support	73.5	76.8	81.4	79.5	76.9	80.1	79.8	82.7	83.5	84.4	85.1
Total	\$2,298.0	\$2,485.2	\$2,606.1	\$2,589.9	\$2,523.8	\$2,385.4	\$2,319.3	\$2,223.5	\$2,205.2	\$2,192.4	\$2,131.7

* This chart is based on the Biomedical Research & Development Price Index through 2010.

Note: From 1999 to 2006, the WHI was reported separately. In this table, it has been incorporated into the “Heart” line. The Sleep Disorders Research was reported separately from 1996 to 2006. In this table, it has been incorporated into the “Lung” line.

NHLBI Total Obligations by Budget Mechanism: Fiscal Years 2001–2011



* Includes Research Career Programs.

** Excludes Office of the Director and DIR research contracts, which are included in R&D contracts.

NHLBI Total Obligations by Budget Mechanism: Fiscal Years 2001–2011

Funding Mechanism	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Research Grants*	\$1,796.9	\$2,006.2	\$2,172.3	\$2,257.3	\$2,310.2	\$2,275.9	\$2,263.1	\$2,216.9	\$2,261.0	\$2,310.3	\$2,288.6
Research and Development (R&D) Contracts	220.1	258.3	290.5	285.5	268.6	262.8	295.8	338.8	361.1	379.9	368.3
Training Programs	73.7	79.2	85.8	87.1	88.4	89.2	93.3	94.9	96.6	98.0	98.0
Intramural Research and Research Management and Support**	207.3	226.1	245.1	252.7	255.4	265.6	270.1	286.7	295.8	305.3	314.6
Total	\$2,298.0	\$2,569.8	\$2,793.7	\$2,882.6	\$2,922.6	\$2,893.5	\$2,922.3	\$2,937.3	\$3,014.5	\$3,093.5	\$3,069.5

* Includes Research Career Programs.

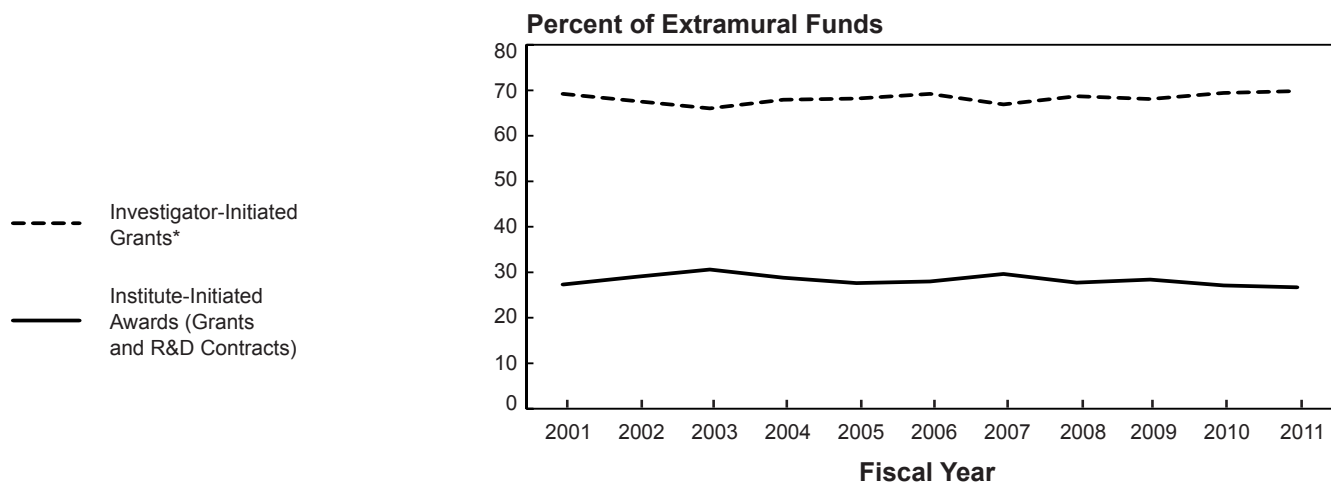
** Excludes Office of the Director and DIR research contracts, which are included in R&D contracts.

NHLBI Employment: Fiscal Years 2001–2011

Staff	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
FTEs*	868	880	880	861	796	797	814	846	856	876	876

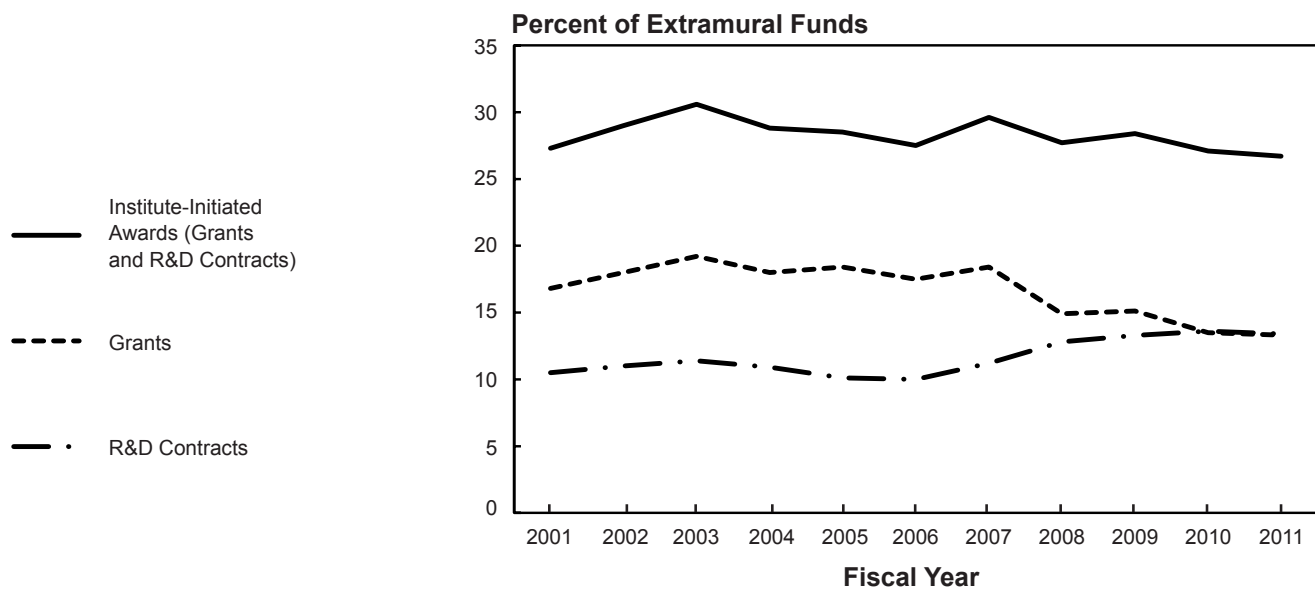
* Full-time equivalents.

NHLBI Institute-Initiated and Investigator-Initiated Awards: Fiscal Years 2001–2011



* Includes Research Career Programs.

NHLBI Grants and Research and Development Contracts as Subsets of Institute-Initiated Awards: Fiscal Years 2001–2011



NHLBI Extramural Programs: Fiscal Years 2001–2011

Funding Mechanism	Dollars (Millions)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Investigator-Initiated Awards											
Investigator-Initiated Grants*	\$1,388.8	\$1,521.4	\$1,616.1	\$1,716.8	\$1,747.2	\$1,747.0	\$1,719.3	\$1,742.1	\$1,765.5	\$1,865.9	\$1,842.8
Research Career Programs	57.5	63.5	65.8	67.8	71.0	70.4	55.4	78.7	84.6	68.0	79.1
Subtotal, Investigator-Initiated Awards	1,446.3	1,584.9	1,681.9	1,784.6	1,818.2	1,817.3	1,774.7	1,820.8	1,850.1	1,933.9	1,921.9
Institute-Initiated Awards											
Institute-Initiated Grants (RFA)	350.7	421.3	490.4	472.5	492.1	458.6	488.2	396.1	410.9	376.4	366.7
<i>Centers**</i>	127.2	128.2	138.9	140.6	151.5	141.1	141.0	107.3	90.1	72.6	55.9
R&D Contracts (RFP)	220.1	258.3	290.5	285.5	268.6	262.9	295.8	338.8	361.1	379.9	368.3
Subtotal, Institute-Initiated Awards	570.8	679.6	780.9	758.0	760.7	721.4	784.0	734.9	772.0	756.3	735.0
Training											
Individual Awards	8.9	9.5	8.6	8.8	9.7	10.0	8.2	9.0	10.3	11.7	10.6
Institutional Awards	64.8	69.7	77.2	78.4	78.7	79.1	85.1	85.8	86.2	86.3	87.4
Subtotal, Training	73.7	79.2	85.8	87.2	88.4	89.2	93.3	94.8	96.5	98.0	98.0
Total, Extramural	\$2,090.8	\$2,343.7	\$2,548.6	\$2,629.8	\$2,667.3	\$2,628.0	\$2,652.0	\$2,650.5	\$2,718.6	\$2,788.2	\$2,754.9

* Includes all R18s.

** Centers are a subset of Institute-Initiated Grants (RFAs) and are not added to the Institute-Initiated Awards subtotal as a distinct category.

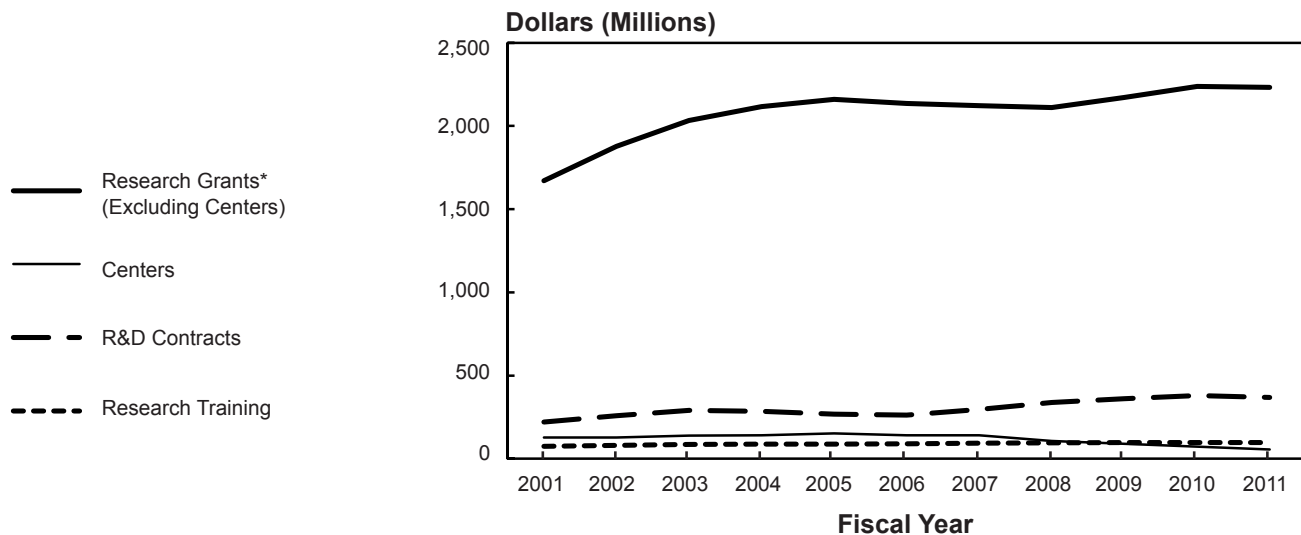
NHLBI Extramural Programs: Fiscal Years 2001–2011

Funding Mechanism	Percent of Total Extramural Budget										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Investigator-Initiated Awards											
Investigator-Initiated Grants*	66.4%	64.9%	63.4%	65.3%	65.5%	66.5%	64.8%	65.7%	64.9%	66.9%	66.9%
Research Career Programs (K04, K06)	2.8	2.7	2.6	2.6	2.7	2.7	2.1	3.0	3.1	2.4	2.9
Subtotal, Investigator-Initiated Awards	69.2	67.6	66.0	67.9	68.2	69.2	66.9	68.7	68.1	69.4	69.8
Institute-Initiated Awards											
Institute-Initiated Grants (RFA)	16.8	18.0	19.2	18.0	18.4	17.5	18.4	14.9	15.1	13.5	13.3
<i>Centers**</i>	6.1	5.5	5.5	5.3	5.7	5.4	5.3	4.0	3.3	2.6	2.0
R&D Contracts (RFP)	10.5	11.0	11.4	10.9	10.1	10.0	11.2	12.8	13.3	13.6	13.4
Subtotal, Institute-Initiated Awards	27.3	29.0	30.6	28.8	28.5	27.5	29.6	27.7	28.4	27.1	26.7
Training											
Individual Awards	0.4	0.4	0.3	0.3	0.4	0.4	0.3	0.3	0.4	0.4	0.4
Institutional Awards	3.1	3.0	3.0	3.0	3.0	3.0	3.2	3.2	3.2	3.1	3.2
Subtotal, Training	3.5	3.4	3.4	3.3	3.3	3.4	3.5	3.6	3.5	3.5	3.6
Total, Extramural	100%	100%	100%	100%	100%	100%	100%	100%	100%	100%	100%

* Includes all R18s.

** Centers are a subset of Institute-Initiated Grants (RFAs) and are not added to the Institute-Initiated Awards subtotal as a distinct category.

NHLBI Extramural Research Funding Mechanism: Fiscal Years 2001–2011



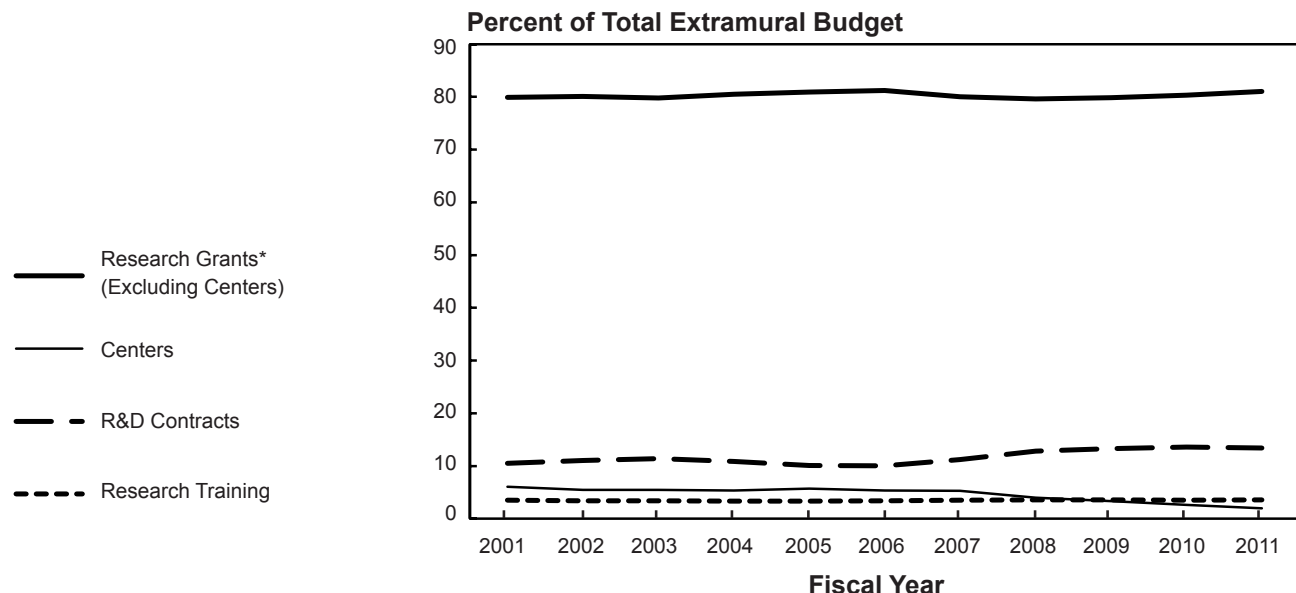
* Includes Research Career Programs; does not include Centers.

NHLBI Extramural Research Funding Mechanism: Fiscal Years 2001–2011

Funding Mechanism	Dollars (Millions)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Research Grants*	\$1,669.8	\$1,878.0	\$2,033.4	\$2,116.5	\$2,158.8	\$2,134.9	\$2,121.9	\$2,109.6	\$2,170.9	\$2,237.7	\$2,232.7
Centers	127.2	128.2	138.9	140.6	151.5	141.1	141.0	107.3	90.1	72.6	55.9
R&D Contracts	220.1	258.3	290.5	285.5	268.6	262.9	295.8	338.8	361.1	379.9	368.3
Research Training	73.7	79.2	85.8	87.2	88.4	89.2	93.3	94.8	96.5	98.0	98.0
Total, Extramural	\$2,090.8	\$2,343.7	\$2,548.6	\$2,629.8	\$2,667.3	\$2,628.0	\$2,652.0	\$2,650.5	\$2,718.6	\$2,788.2	\$2,754.9

* Includes Research Career Programs; does not include Centers.

NHLBI Extramural Research Funding Mechanism: Fiscal Years 2001–2011



* Includes Research Career Programs; does not include Centers.

NHLBI Extramural Research Funding Mechanism: Fiscal Years 2001–2011

Funding Mechanism	Percent of Total Extramural Budget										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Research Grants*	79.9%	80.1%	79.8%	80.5%	80.9%	81.2%	80.0%	79.6%	79.9%	80.3%	81.0%
Centers	6.1	5.5	5.5	5.3	5.7	5.4	5.3	4.0	3.3	2.6	2.0
R&D Contracts (RFP)	10.5	11.0	11.4	10.9	10.1	10.0	11.2	12.8	13.3	13.6	13.4
Research Training	3.5	3.4	3.4	3.3	3.3	3.4	3.5	3.6	3.5	3.5	3.6
Total, Extramural	100%	100%	100%	100%	100%	100%	100%	100%	100%	100%	100%

* Includes Research Career Programs; does not include Centers.

Note: Numbers may not add to total due to rounding.

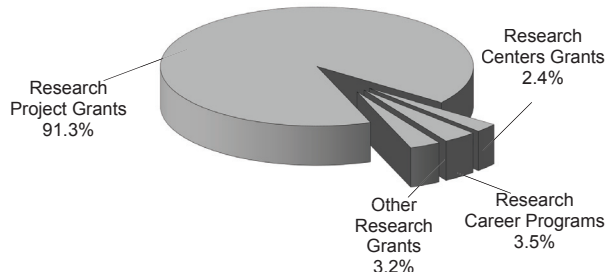


9. Research Grants

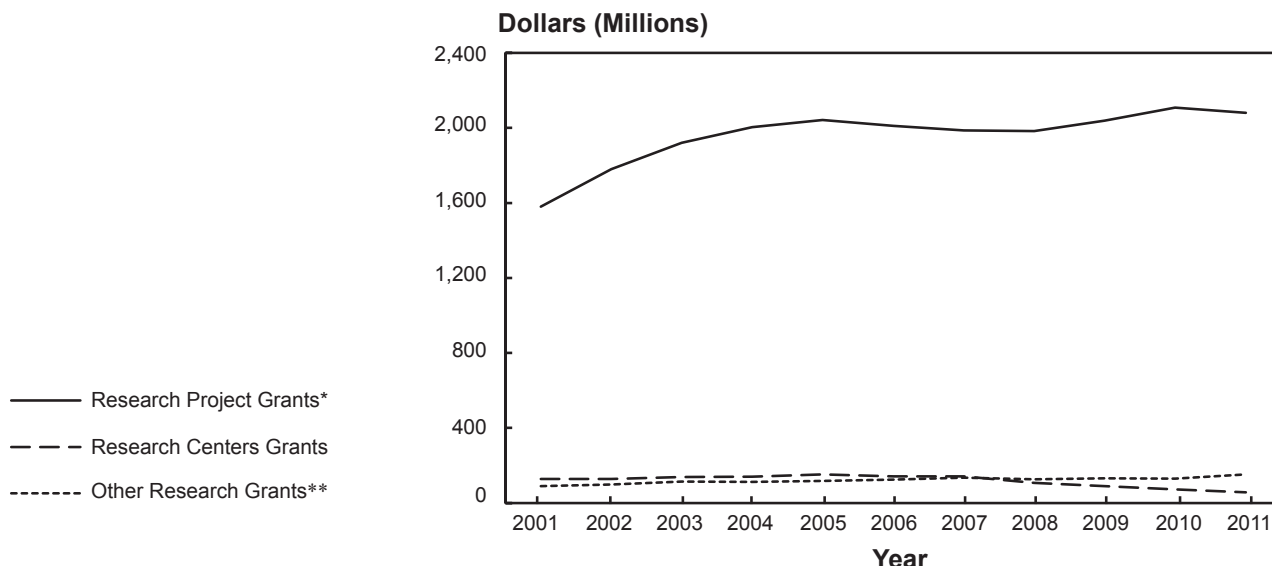
NHLBI Research Grants by Funding Mechanism: Fiscal Year 2011

	Number of Grants	Total Cost (Dollars in Thousands)	Percent of Total NHLBI Research Grant Dollars
Research Project Grants (RPGs)			
Research Project Grants (Excluding Small Business RPGs)			
Regular Research Grants (R01)	2,943	\$1,355,916	59.25%
Program Project Grants (P01)	153	321,179	14.03
Cooperative Agreements (U01)	156	180,217	7.87
Exploratory Developmental Grant (R21)	316	68,722	3.00
Method to Extend Research in Time (R37)	71	30,775	1.34
Exploratory/Developmental Grants Phase II (R33)	3	2,289	0.10
Clinical Trial Planning Grant (R34)	12	4,646	0.20
Clinical Planning Grant Cooperative Agreement (U34)	2	1,573	0.07
Academic Research Enhancement Award (R15)	9	3,547	0.15
NIH Director's New Innovator's Award (DP2)	2	4,928	0.22
Research Transition Award (R00)	75	18,606	0.81
Cooperative Agreements (U19)	3	6,513	0.28
Small Research Grants (R03)	9	693	0.03
Linked Research Grant (RL1)	—	59	0.00
NIH Director's Pioneer Award (DP1)	3	2,255	0.10
Exploratory/Developmental Cooperative Agreements Phase I (UH2)	2	988	0.04
Multi-Component Research Project Cooperative Agreements (UM1)	—	2,079	0.09
Subtotal, Research Project Grants (Excluding Small Business RPGs)	3,759	2,004,985	88.00
Small Business Research Project Grants			
Small Business Technology Transfer (STTR Phase I) (R41)	15	3,396	0.15
Small Business Technology Transfer (STTR Phase II) (R42)	11	5,608	0.25
Small Business Innovation Research (SBIR Phase I) (R43)	67	16,973	0.74
Small Business Innovation Research (SBIR Phase II) (R44)	70	48,958	2.14
Subtotal, Small Business Research Project Grants	163	74,935	3.26
Subtotal, Research Project Grants	3,922	2,079,920	91.26
Research Centers Grants			
Centers of Research Programs (P50)	30	34,366	1.50
Sickle Cell Centers (U54)	13	5,901	0.26
Centers for AIDS Research (P30)	—	3,334	0.15
Specialized Centers (Cooperative Agreements) (U54)	—	1,265	0.06
National Swine Research and Resource Center (U42)	—	1,448	0.06
Exploratory Grants (P20)	12	8,593	0.38
Center Core Grants (P30)	5	1,024	0.04
Subtotal, Research Centers Grants	60	55,931	2.44
Research Career Programs			
Mentored Research Scientist Development Award for Minority Faculty (K01)	45	5,860	0.26
Minority Institution Faculty Mentored Research Scientist Development Award (K01)	5	668	0.03
Mentored Scientist Development Award in Research Ethics (K01)	—	126	0.01
Independent Scientist Award (K02)	21	2,076	0.09
Pediatric Transfusion Medicine Academic Award (K07)	4	486	0.02
Innovators in Hemoglobinopathies Care Career Development Award (K07)	2	574	0.03
Clinical Investigator Development Award (K08)	210	27,166	1.19
Research Career Development Program in Vascular Medicine (K12)	3	2,499	0.11
Clinical Research Career Development Programs in Emergency Medicine (K12)	6	1,186	0.05
Career Enhancement Award for Stem Cell Research (K18)	5	789	0.03
Career Transition Award (K22)	3	699	0.03
Mentored Patient-Oriented Research Career Development Award (K23)	170	23,871	1.04
Midcareer Investigator Award in Patient-Oriented Research (K24)	34	5,851	0.26
Mentored Quantitative Research Career Development Award (K25)	15	2,110	0.09
Career Transition Award (K99)	47	5,129	0.22
Subtotal, Research Career Programs	570	79,090	3.46
Other Research Grants			
Cooperative Clinical Research (U10, R10)	50	44,705	1.95
Minority Biomedical Research Support (S06, SC1, SC2)	13	2,883	0.13
Other (D43, R13, R18, R24, R25, T15, U24, UH1)	115	26,094	1.14
Subtotal, Other Research Grants	178	73,682	3.22
Total, NHLBI Research Grants	4,730	\$2,288,623	100%

NHLBI Total Research Grants by Category



NHLBI Research Project Grants, Research Centers Grants, and Other Research Grant Obligations: Fiscal Years 2001–2011



* Includes R01, U01, P01, R03, R15, R21, R29, R37, R41, R42, R43, and R44; R33 beginning in 2001; DP2 and U19 beginning in 2007; DP1 and R00 beginning in 2008; R34 and U34 beginning in 2010; and UH2 and UM1 beginning in 2011.

** Includes Research Career Programs; excludes General Research Support Grants.

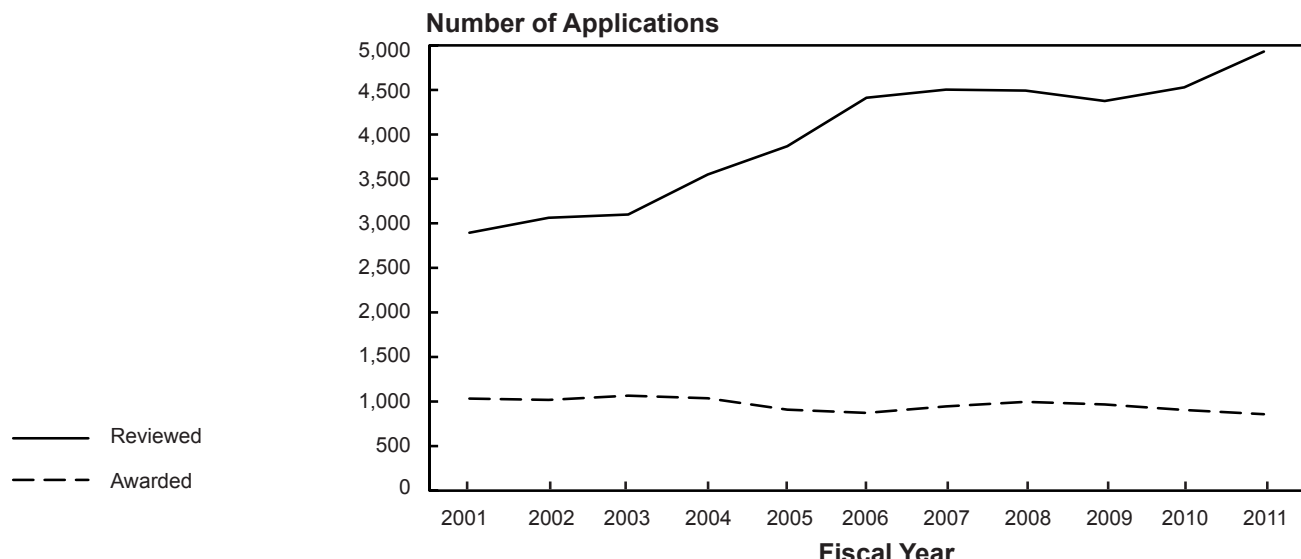
NHLBI Research Project Grants, Research Centers Grants, and Other Research Grant Obligations: Fiscal Years 2001–2011

	Dollars (Thousands)										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Research Project Grants*	\$1,580,751	\$1,779,573	\$1,920,201	\$2,003,769	\$2,042,050	\$2,011,049	\$1,986,692	\$1,983,633	\$2,039,861	\$2,108,524	\$2,079,920
Research Centers Grants	127,232	128,161	138,941	140,600	151,495	141,086	141,034	107,393	90,152	72,566	55,931
Other Research Grants**	88,958	98,460	113,172	112,785	116,713	123,802	135,284	125,942	131,001	129,245	152,772
Total	\$1,796,941	\$2,006,194	\$2,172,314	\$2,257,154	\$2,310,258	\$2,275,937	\$2,263,010	\$2,216,968	\$2,261,014	\$2,310,335	\$2,288,623

* Includes R01, U01, P01, R03, R15, R21, R29, R37, R41, R42, R43, and R44; R33 beginning in 2001; DP2 and U19 beginning in 2007; DP1 and R00 beginning in 2008; R34 and U34 beginning in 2010; and UH2 and UM1 beginning in 2011.

** Includes Research Career Programs; excludes General Research Support Grants.

NHLBI Competing Research Project Grant Applications:^{*} Fiscal Years 2001–2011 Number Reviewed and Awarded

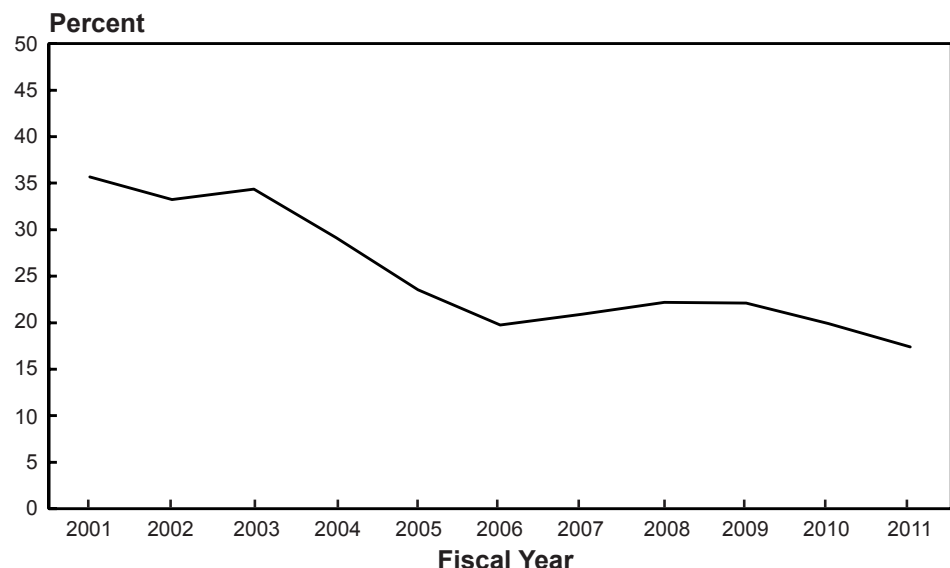


Number Reviewed and Awarded and Percent Funded

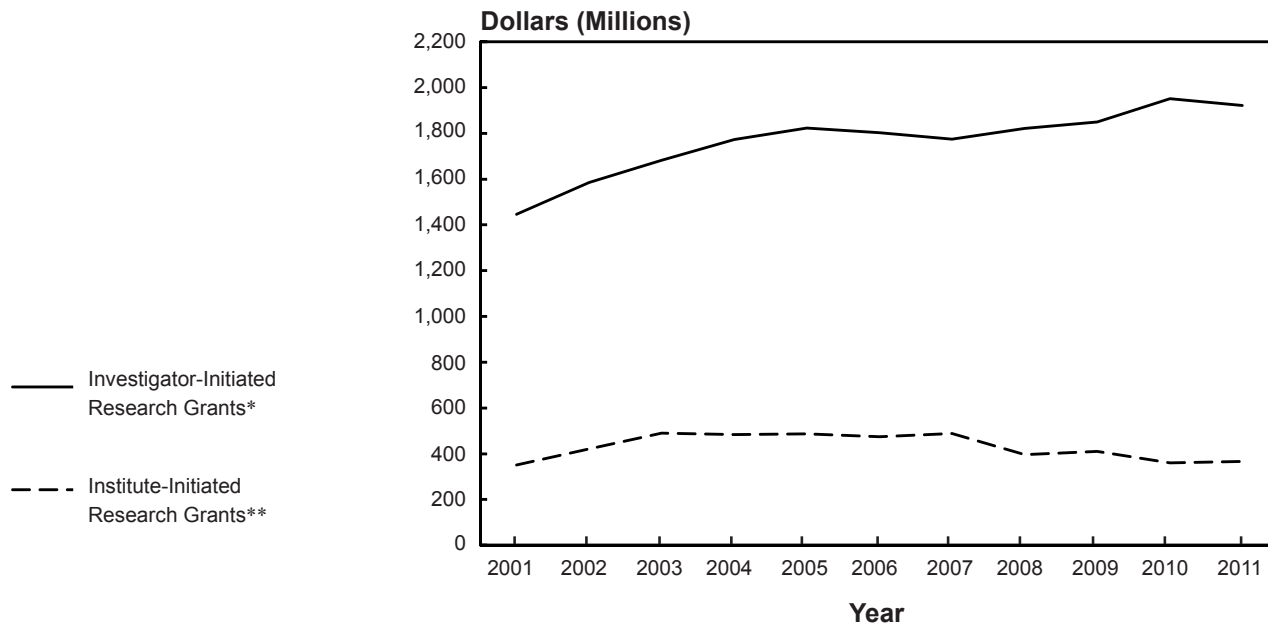
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Applications Reviewed	2,895	3,064	3,098	3,548	3,865	4,412	4,504	4,492	4,373	4,528	4,931
Applications Awarded	1,033	1,018	1,064	1,034	909	871	943	997	968	903	856
Percent Funded (Success Rate)	35.7	33.2	34.3	29.1	23.5	19.7	20.9	22.2	22.1	19.9	17.4

* Includes R01, U01, P01, R03, R15, R21, R29, and R37; R33 beginning in 2001; DP2 and U19 beginning in 2007; DP1 and R00 beginning in 2008; R34 and U34 beginning in 2010; and UH2 and UM1 beginning in 2011.

Percent of Reviewed Applications Funded (Success Rate)



NHLBI Investigator-Initiated and Institute-Initiated Grant Obligations: Fiscal Years 2001–2011



* Includes RPGs, SBIRs/STTRs, Research Career Programs, and Other Research.

** Includes RPGs, Centers Grants, Research Career Programs, Other Research, and Cooperative Agreement RFAs.

NHLBI Investigator-Initiated and Institute-Initiated Grant Obligations: Fiscal Years 2001–2011

	Dollars (Millions)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Investigator-Initiated*	\$1,446.2	\$1,584.9	\$1,681.9	\$1,773.4	\$1,822.9	\$1,802.1	\$1,774.8	\$1,820.8	\$1,850.1	\$1,950.9	\$1,921.9
Institute-Initiated**	350.7	421.3	490.4	483.8	487.3	473.8	488.2	396.1	410.9	359.5	366.7
Total	\$1,796.9	\$2,006.2	\$2,172.3	\$2,257.2	\$2,310.2	\$2,275.9	\$2,263.0	\$2,216.9	\$2,261.0	\$2,310.4	\$2,288.6

* Includes RPGs, SBIRs/STTRs, Research Career Programs, and Other Research.

** Includes RPGs, Centers Grants, Research Career Programs, Other Research, and Cooperative Agreement RFAs.

NHLBI Research Project Grants:^{*} Amount Funded by Type of Award, Fiscal Years 2001–2011

	Dollars (Millions)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Competing											
New Competing	\$ 280.0	\$ 291.2	\$ 285.5	\$ 290.5	\$ 270.0	\$ 242.9	\$ 330.9	\$ 314.2	\$ 340.2	\$ 330.5	\$ 353.1
Renewal Competing	143.9	143.9	177.2	185.5	176.1	168.3	169.4	196.9	172.6	171.6	131.6
Competing Supplements	0.4	2.3	1.0	1.3	1.7	0.4	—	1.7	0.3	0.3	—
Subtotal, Competing	424.3	437.4	463.7	477.3	447.8	411.6	500.3	512.8	513.1	501.8	484.7
Noncompeting											
Subtotal, Noncompeting	1,101.5	1,281.3	1,390.3	1,454.9	1,520.0	1,527.0	1,486.4	1,470.8	1,526.8	1,606.7	1,595.2
Total, Competing and Noncompeting	\$1,525.8	\$1,718.7	\$1,854.0	\$1,932.2	\$1,967.8	\$1,938.6	\$1,986.7	\$1,983.6	\$2,039.9	\$2,108.5	\$2,079.9

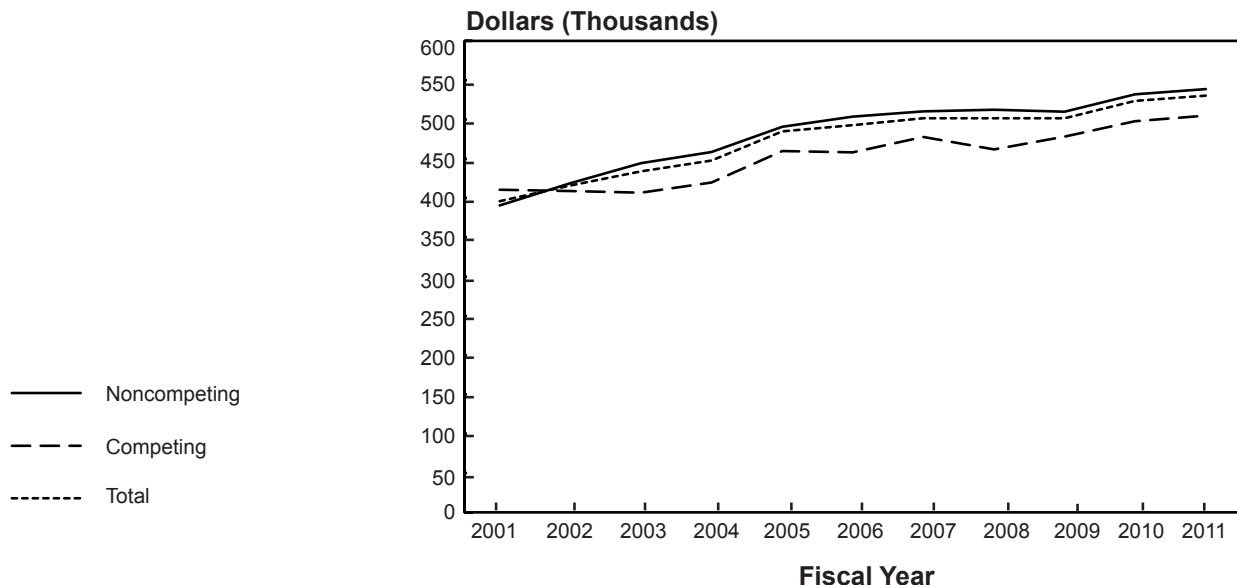
* Includes R01, U01, P01, R03, R15, R21, R29, R37, R41, R42, R43, and R44; R33 beginning in 2001; DP2 and U19 beginning in 2007; DP1 and R00 beginning in 2008; R34 and U34 beginning in 2010; and UH2 and UM1 beginning in 2011.

Facility and Administrative (F&A) Costs of NHLBI Research Project Grants:^{*} Fiscal Years 2001–2011

Fiscal Year	Dollars (Thousands)			
	Direct Cost	F&A Cost	Total Cost	F&A Cost as a Percent of Direct Cost
2001	1,045,144	480,673	1,525,817	46.0
2002	1,182,408	536,324	1,718,732	45.4
2003	1,276,819	577,131	1,853,950	45.2
2004	1,329,106	603,133	1,932,239	45.4
2005	1,355,803	612,007	1,967,810	45.1
2006	1,334,406	604,183	1,938,589	45.3
2007	1,378,134	608,558	1,986,692	44.2
2008	1,376,276	607,357	1,983,633	44.1
2009	1,410,033	629,828	2,039,861	44.7
2010	1,459,211	649,313	2,108,524	44.5
2011	1,429,935	649,985	2,079,920	45.5

* Includes R01, U01, P01, R03, R15, R21, R29, R37, R41, R42, R43, and R44; R33 beginning in 2001; DP2 and U19 beginning in 2007; DP1 and R00 beginning in 2008; R34 and U34 beginning in 2010; and UH2 and UM1 beginning in 2011.

NHLBI Research Project Grants:^{*} Average Costs, Fiscal Years 2001–2011



* Includes R01, U01, P01, R03, R15, R21, R29, R37, R41, R42, R43, and R44; R33 beginning in 2001; DP2 and U19 beginning in 2007; DP1 and R00 beginning in 2008; R34 and U34 beginning in 2010; and UH2 and UM1 beginning in 2011.

NHLBI Research Project Grants:^{*} Average Costs, Fiscal Years 2001–2011

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Noncompeting	\$390.7	\$418.8	\$444.4	\$458.7	\$490.6	\$503.9	\$510.3	\$512.4	\$509.8	\$532.2	\$538.6
Competing	410.8	409.1	406.7	419.7	459.9	458.1	477.8	462.0	478.2	497.9	504.9
Total	\$396.1	\$416.2	\$433.8	\$447.9	\$484.8	\$492.8	\$501.7	\$501.8	\$501.4	\$523.6	\$530.3

* Includes R01, U01, P01, R03, R15, R21, R29, R37, R41, R42, R43, and R44; R33 beginning in 2001; DP2 and U19 beginning in 2007; DP1 and R00 beginning in 2008; R34 and U34 beginning in 2010; and UH2 and UM1 beginning in 2011.

NHLBI Cooperative Agreements (U01, U10, U19) Programs

Cooperative Agreements were instituted to support discrete, circumscribed projects in areas of an investigator's specific interest and competency with substantial programmatic participation by the NHLBI during performance of the activity.

	Total Obligations Prior to FY 2011	Total FY 2011 Obligations	Total Obligations to Date
Heart and Vascular Diseases			
ARIC Neurocognitive Study (ARIC-NCS)	\$4,024,211	\$5,132,469	\$9,156,680
Cardiovascular Cell Therapy Research Network	24,419,327	3,800,000	28,219,327
Cardiovascular Inflammation Reduction Trial	—	1,375,726	1,375,726
Cardiovascular Outcomes in Renal Atherosclerotic Lesions (CORAL)	25,827,224	3,029,201	28,856,425
Catheter Ablation Versus Antiarrhythmic Drug Therapy for Atrial Fibrillation (CABANA) Trial	5,985,643	3,032,875	9,018,518
Center for Cardiovascular Outcomes Research	5,203,594	4,964,931	10,168,525
Childhood Obesity Prevention and Treatment Research (COPTR)	4,058,435	3,986,427	8,044,862
Claudication Exercise Versus Endoluminal Revascularization (CLEVER)	7,647,416	1,371,307	9,018,723
Clinical Research Consortium To Improve Resuscitation Outcomes	46,447,452	9,455,378	55,902,830
Consortium of Hospitals Advancing Research on Tobacco (CHART)	3,321,649	3,505,096	6,826,745
Cross Organ Mechanism-Associated Phenotypes For Genetic Analysis of H, L, B, and S Diseases	—	4,275,465	4,275,465
Diabetes Prevention Program Outcomes Study—Phase II	2,200,000	1,100,000	3,300,000
Dynamic Evaluation of Percutaneous Coronary Intervention	8,413,711	742,170	9,155,881
Early Adult Reduction of Weight Through LifestYle Intervention (EARLY Trials*)	10,519,703	6,123,513	16,643,216
Genetics of Coronary Artery Disease in Alaskan Natives (GOCADAN)	17,504,059	51,881	17,555,940
Heart Failure Clinical Research Network	37,108,888	7,651,786	44,760,674
ISCHEMIA (International Study of Comparative Health Effectiveness with Medical Invasive Approaches) Trial	—	6,671,629	6,671,629
Look AHEAD: Action for Health in Diabetes	8,000,000	4,000,000	12,000,000
Network for Cardiothoracic Surgical Investigation in Cardiovascular Medicine	25,978,806	5,459,597	31,438,403
Next Generation Genetic Association Studies	—	5,855,804	5,855,804
NHLBI Cardiac Development Consortium	8,239,641	7,037,111	15,276,752
NHLBI Pediatric Cardiac Genomics Consortium	5,143,441	3,814,314	8,957,755
NHLBI Pediatric Translational Consortium Administrative Coordinating Center	10,453,976	—	10,453,976
NHLBI Progenitor Cell Biology Consortium Research Hubs	46,788,094	22,514,886	69,302,980
Obesity Related Behavioral Intervention Trials (ORBIT)**	10,987,451	4,297,228	15,284,679
Pediatric Heart Network	63,348,873	12,827,177	76,176,050
Pharmacogenetics Research Network	74,421,336	6,532,424	80,953,760
Ranolazine in Implantable Defibrillators (RAID) Trial	2,279,794	—	2,279,794
Rule Out Myocardial Infarction Using Computed Assisted Tomography (ROMICAT II)	4,419,188	366,525	4,785,713
Strong Heart Study	73,879,041	2,144,794	76,023,835
Surgical Treatment for Ischemic Heart Failure (STICH)	40,041,062	352,198	40,393,260
Therapeutic Hypothermia After Pediatric Cardiac Arrest (THAPCA) Trials	5,993,723	4,401,772	10,395,495
Subtotal, Heart and Vascular Diseases	582,655,738	145,873,684	728,529,422
Lung Diseases			
Asthma Networks (AsthmaNet)	23,800,000	15,500,000	39,300,000
COPD Clinical Research Network	46,330,386	2,600,000	48,930,386
Heart and Lung Failure-Pediatric Insulin Titration Trial (HALF-PINT)	—	2,685,460	2,685,460
Infant Study of Inhaled Saline in Cystic Fibrosis (ISIS)	2,244,874	628,241	2,873,115
Microbiome of the Lung and Respiratory Tract in HIV-Infected and in HIV-Uninfected Controls	10,743,005	5,324,656	16,067,661
Novel Therapies for Lung Diseases—Phase II	7,593,739	12,843,238	20,436,977

	Total Obligations Prior to FY 2011	Total FY 2011 Obligations	Total Obligations to Date
Lung Diseases (continued)			
Pharmacogenetics of Asthma Treatment	28,998,338	1,957,381	30,955,719
Prematurity and Respiratory Outcomes Program (PROP)	1,597,280	4,066,948	5,664,228
Preterm Birth in Nulliparous Women: An Understudied Population at Great Risk	500,000	3,298,000	3,798,000
Randomized Trial of Antenatal Late Preterm Steroids (ALPS)	2,133,947	2,137,414	4,271,361
Randomized Trial of Maternal Vitamin D Supplementation To Prevent Childhood Asthma	4,975,916	2,469,339	7,445,255
Sedation Management in Pediatric Patients With Acute Respiratory Failure	7,779,116	2,181,199	9,960,315
Severe Asthma Research Program	—	5,159,933	5,159,933
Study of Asthma and Nasal Steroids (STAN)	1,449,536	724,884	2,174,420
Study of Soy Isoflavones in Asthma (SOYA)	1,417,854	688,673	2,106,527
Trial of Late Surfactant (TOLSURF) To Prevent Bronchopulmonary Dysplasia (BPD)	3,765,851	1,807,148	5,572,999
Subtotal, Lung Diseases	143,329,842	64,076,514	207,406,356
Blood Diseases and Resources			
Acute Venous Thrombosis: Thrombus Removal With Adjunctive Catheter-Directed Thrombolysis (ATTRACT)**	6,272,537	243,775	6,516,312
Blood and Marrow Transplant Clinical Research Network	59,071,021	5,319,784	64,390,805
Bridging Anticoagulation on Patients Requiring Temporary Interruption of Warfarin Therapy for an Elective Invasive Procedure or Surgery (BRIDGE) Trial	15,531,903	—	15,531,903
Transfusion Medicine/Hemostasis Clinical Research Network	57,040,060	6,133,712	63,173,772
Subtotal, Blood Diseases and Resources	137,915,521	11,697,271	149,612,792
Total, NHLBI Cooperative Agreements	\$856,249,333	\$221,647,469	\$1,085,548,570

* Formerly known as Targeted Approaches to Weight Control for Young Adults.

** Formerly known as Translating Basic Behavior and Social Science Discoveries Into Interventions to Reduce Obesity

† Formerly known as Pharmacomechanical Catheter-Directed Thrombolysis for Acute DVT (ATTRACT) Trial.

Heart and Vascular Diseases Program

ARIC Neurocognitive Study (ARIC-NCS), Initiated in Fiscal Year 2010

The purpose of this study is to determine whether mid-life vascular risk factors and markers of macrovascular and microvascular disease are predictive of dementia, mild cognitive impairment, and cognitive change in a large biracial ARIC cohort.

Obligations

Funding History:

Fiscal Year 2011—\$5,132,469

Fiscal Year 2010—\$4,024,211

Total Funding to Date—\$9,156,680

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. Johns Hopkins University
Baltimore, Maryland | —HL-096812 |
| 2. University of North Carolina
Chapel Hill, North Carolina | —HL-096814 |
| 3. University of North Carolina
Chapel Hill, North Carolina | —HL-096899 |
| 4. University of Minnesota
Minneapolis, Minnesota | —HL-096902 |
| 5. University of Mississippi Medical Center
Jackson, Mississippi | —HL-096917 |

Cardiovascular Cell Therapy Research Network, Initiated in Fiscal Year 2007

See Chapter 11. Clinical Trials.

Cardiovascular Inflammation Reduction Trial, Initiated in Fiscal Year 2011

The purpose of this trial is to test the inflammatory hypothesis of atherothrombosis by evaluating whether low-dose methotrexate will reduce rates of recurrent myocardial infarction, stroke, and cardiovascular death among patients with stable CVD who are at increased risk indicated by persistent elevations of high-sensitivity C-reactive protein.

Obligations

Funding History:

Fiscal Year 2011—\$1,375,726

Total Funding to Date—\$1,375,726

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Brigham and Women's Hospital
Boston, Massachusetts | —HL-101389 |
|--|------------|

- | | |
|--|------------|
| 2. Brigham and Women's Hospital
Boston, Massachusetts | —HL-101422 |
|--|------------|

Cardiovascular Outcomes in Renal Atherosclerotic Lesions (CORAL), Initiated in Fiscal Year 2004

The purpose of this trial is to determine whether renal artery stenting adds value to optimal medical therapy in terms of cardiovascular and renal outcomes in individuals with a history of resistant hypertension and/or chronic kidney disease and stenosis (>60%) of one or both renal arteries. The trial is in an extended follow-up phase.

Obligations

Funding History:

Fiscal Year 2011—\$3,029,201

Fiscal Years 2004–2010—\$25,827,224

Total Funding to Date—\$28,856,425

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. University of Toledo Health Sciences Campus
Toledo, Ohio | —HL-071556 |
| 2. University of Minnesota, Twin Cities
Minneapolis, Minnesota | —HL-072734 |
| 3. University of Virginia
Charlottesville, Virginia | —HL-072735 |
| 4. Mid-America Heart Institute of St. Luke Hospital
Kansas City, Missouri | —HL-072736 |
| 5. Beth Israel Deaconess Medical Center
Boston, Massachusetts | —HL-072737 |

Catheter Ablation Versus Antiarrhythmic Drug Therapy for Atrial Fibrillation (CABANA), Initiated in Fiscal Year 2009

The purpose of this trial is to determine whether percutaneous left atrial catheter ablation is superior to current pharmacologic therapy for eliminating atrial fibrillation.

Obligations

Funding History:

Fiscal Year 2011—\$3,032,875

Fiscal Years 2009–2010—\$5,985,643

Total Funding to Date—\$9,018,518

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. Mayo Clinic, College of Medicine
Rochester, Minnesota | —HL-089645 |
| 2. Mayo Clinic, College of Medicine
Rochester, Minnesota | —HL-089709 |

- | | |
|--|------------|
| 3. Duke University
Durham, North Carolina | —HL-089786 |
| 4. Duke University
Durham, North Carolina | —HL-089907 |

Center for Cardiovascular Outcomes Research, Initiated in Fiscal Year 2010

The purpose of this program is to conduct cardiovascular outcomes research that focuses on patient- and clinician-relevant outcomes of health care and their determinants. The goal is to move clinical evidence into public policy and clinical practice.

Obligations

Funding History:

Fiscal Year 2011—\$4,964,931

Fiscal Year 2010—\$5,203,594

Total Funding to Date—\$10,168,525

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. University of Massachusetts Medical School
Worcester, Massachusetts | —HL-105268 |
| 2. Yale University
New Haven, Connecticut | —HL-105270 |
| 3. Boston Medical Center
Boston, Massachusetts | —HL-105342 |
| 4. Duke University
Durham, North Carolina | —HL-107023 |

Childhood Obesity Prevention and Treatment Research (COPTR), Initiated in Fiscal Year 2010

See Chapter 11. Clinical Trials.

Claudication: Exercise Versus Endoluminal Revascularization, Initiated in Fiscal Year 2005

The purpose of this study is to optimize physical functioning, increase activity levels, and reduce CVD risk in older individuals with peripheral artery disease. Investigators are determining whether a strategy of aortoiliac stenting and pharmacotherapy improves maximum walking duration better than a strategy of supervised rehabilitation, exercise, and pharmacotherapy for those with moderate to severe claudication due to aortoiliac insufficiency.

Obligations

Funding History:

Fiscal Year 2011—\$1,371,307

Fiscal Years 2005–2010—\$7,647,416

Total Funding to Date—\$9,018,723

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Rhode Island Hospital
Providence, Rhode Island | —HL-077221 |
| 2. Beth Israel Deaconess Medical Center
Boston, Massachusetts | —HL-081656 |

Clinical Research Consortium To Improve Resuscitation Outcomes, Initiated in Fiscal Year 2004

See Chapter 11. Clinical Trials.

Consortium of Hospitals Advancing Research on Tobacco (CHART), Initiated in Fiscal Year 2010

See Chapter 11. Clinical Trials.

Cross Organ Mechanism-Associated Phenotypes for Genetic Analysis of Heart, Lung, Blood, and Sleep Diseases (MAPGen for HLBS) Research Centers, Initiated in Fiscal Year 2011

The purpose of this program is to establish a MAPGen consortium of research centers. The consortium seeks to utilize evolving knowledge of cellular and molecular networks to define common mechanism-associated traits across organ systems. The goal is to redefine disease at the level of pathogenetic mechanisms and phenotype individuals based on pathobiology, rather than clinical presentation. This approach will provide the basis for the development of mechanism-based strategies for prevention, diagnosis, and treatment in individual patients.

Obligations

Funding History:

Fiscal Year 2011—\$4,275,465

Total Funding to Date—\$4,275,465

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. Brigham and Women's Hospital
Boston, Massachusetts | —HL-108630 |
| 2. University of Southern California
Los Angeles, California | —HL-108634 |
| 3. University of Pennsylvania
Philadelphia, Pennsylvania | —HL-108636 |
| 4. Yale University
New Haven, Connecticut | —HL-108638 |
| 5. University of Pittsburgh
Pittsburgh, Pennsylvania | —HL-108642 |
| 6. Stanford University
Menlo Park, California | —HL-108647 |

Diabetes Prevention Program Outcomes Study— Phase II, Initiated in Fiscal Year 2009

See Chapter 11. Clinical Trials.

Dynamic Evaluation of Percutaneous Coronary Intervention, Initiated in Fiscal Year 1997

The purpose of this program, which complements prior NHLBI percutaneous transluminal coronary angioplasty (PTCA) registries and the New Approaches to Coronary Intervention Registry, is to evaluate patterns of device usage and the immediate and follow-up outcomes in patients who are undergoing percutaneous transluminal coronary revascularization. Results will provide guidance to the cardiology community in selecting appropriate therapies and designing clinical trials to evaluate competing devices. Twenty-four percent of the patients are from various racial and ethnic minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$742,170

Fiscal Years 1997–2010—\$8,413,711

Total Funding to Date—\$9,155,881

Current Active Organization and Grant Number

1. University of Pittsburgh
Pittsburgh, Pennsylvania

—HL-033292

Early Adult Reduction of Weight Through LifestYLE Intervention (EARLY) Trials,* Initiated in Fiscal Year 2009

See Chapter 11. Clinical Trials.

Genetics of Coronary Artery Disease in Alaska Natives (GOCADAN), Initiated in Fiscal Year 2000

The purpose of this study is to document CVD and CVD risk factors in approximately 40 extended families (1,214 members from villages in Northern Alaska). Scientists are seeking to identify and characterize genes that contribute to CVD in this unique and understudied population.

Obligations

Funding History:

Fiscal Year 2011—\$51,881

Fiscal Years 2000–2010—\$17,504,059

Total Funding to Date—\$17,555,940

Current Active Organization and Grant Number

1. MedStar Research Institute
Hyattsville, Maryland

—HL-064244

Heart Failure Clinical Research Network, Initiated in Fiscal Year 2006

See Chapter 11. Clinical Trials.

ISCHEMIA (International Study of Comparative Health Effectiveness with Medical Invasive Approaches) Trial, Initiated in Fiscal Year 2011

The purpose of this trial is to define the role of an invasive approach in patients with stable ischemic heart disease and substantial ischemia. Investigators will determine whether cardiac catheterization followed by complete revascularization plus optimal medical therapy is superior to optimal medical therapy alone as the management strategy for patients with moderate-severe ischemia on stress imaging. Cost-effectiveness will also be assessed.

Obligations

Funding History:

Fiscal Year 2011—\$6,671,629

Total Funding to Date—\$6,671,629

Current Active Organizations and Grant Numbers

1. Duke University
Durham, North Carolina
2. Emory University
Atlanta, Georgia
3. Duke University
Durham, North Carolina
4. New York University School of Medicine
New York, New York

—HL-105462

—HL-105561

—HL-105565

—HL-105907

Look AHEAD: Action for Health in Diabetes, Initiated in Fiscal Year 2009

See Chapter 11. Clinical Trials.

* Formerly known as Targeted Approaches to Weight Control for Young Adults.

Network for Cardiothoracic Surgical Investigation in Cardiovascular Medicine, Initiated in Fiscal Year 2007

See Chapter 11. Clinical Trials.

Next Generation Genetic Association Studies, Initiated in Fiscal Year 2011

The purpose of this study is to investigate functional aspects of genetic variation in humans by combining cellular reprogramming strategies with molecular profiling or cellular assays, and then integrating the information with existing genotypic and clinical phenotypic data to assess how naturally occurring human genetic variation influences the activities of biological networks in cell-based models of disease. Researchers are seeking to develop the technology needed for high-throughput iPS cell line generation and differentiation and will use the technology to follow up on genomic associations with additional mechanistic information gained from cellular models of disease.

Obligations

Funding History:

Fiscal Year 2011—\$5,855,804
Total Funding to Date—\$5,855,804

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Stanford University
Stanford, California | —HL-107388 |
| 2. Stanford University
Stanford, California | —HL-107393 |
| 3. Scripps Research Institute
La Jolla, California | —HL-107436 |
| 4. Medical College of Wisconsin
Milwaukee, Wisconsin | —HL-107437 |
| 5. Massachusetts General Hospital
Boston, Massachusetts | —HL-107440 |
| 6. University of California, San Diego
La Jolla, California | —HL-107442 |
| 7. Boston University Medical Campus
Boston, Massachusetts | —HL-107443 |
| 8. Johns Hopkins University
Baltimore, Maryland | —HL-107446 |
| 9. University of Pennsylvania
Philadelphia, Pennsylvania | —HG-006398 |

NHLBI Cardiac Development Consortium, Initiated in Fiscal Year 2009

The purpose of this study is to elucidate the regulatory networks controlling cardiovascular development. A consortium of multidisciplinary research teams will

select key regulatory pathways, identify components of the pathways and targets, and disseminate data to the scientific community. Research results may lead to the development of regenerative therapies and tissue engineering approaches.

Obligations

Funding History:

Fiscal Year 2011—\$7,037,111
Fiscal Years 2009–2010—\$8,239,641
Total Funding to Date—\$15,276,752

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. University of Utah
Salt Lake City, Utah | —HL-098160 |
| 2. Harvard University Medical School
Boston, Massachusetts | —HL-098166 |
| 3. J. David Gladstone Institutes
San Francisco, California | —HL-098179 |
| 4. University of Pittsburgh
Pittsburgh, Pennsylvania | —HL-098180 |

NHLBI Pediatric Cardiac Genomics Consortium, Initiated in Fiscal Year 2009

The purpose of this study is to conduct clinical and translational research on the genetic causes of congenital heart disease and the genetic contributions to outcomes in individuals with congenital heart disease.

Obligations

Funding History:

Fiscal Year 2011—\$3,814,314
Fiscal Years 2009–2010—\$5,143,441
Total Funding to Date—\$8,957,755

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Mount Sinai School of Medicine
New York, New York | —HL-098123 |
| 2. Children's Hospital Boston
Boston, Massachusetts | —HL-098147 |
| 3. Children's Hospital of Philadelphia
Philadelphia, Pennsylvania | —HL-098153 |
| 4. Yale University
New Haven, Connecticut | —HL-098162 |
| 5. Columbia University Health Sciences
New York, New York | —HL-098163 |

NHLBI Pediatric Translational Consortium Administrative Coordinating Center, Initiated in Fiscal Year 2009

The purpose of this Coordinating Center is to provide administrative support for the Cardiovascular Development Consortium and the Pediatric Cardiac Genomics Consortium, monitor multicenter patient recruitment by the Pediatric Cardiac Genomics Consortium, and administer funds to consortium-wide cores.

Obligations

Funding History:

Fiscal Year 2011—\$0

Fiscal Years 2009–2010—\$10,453,976

Total Funding to Date—\$10,453,976

Current Active Organization and Grant Number

- | | |
|---|-----------|
| 1. New England Research Institute, Inc.
Watertown, Massachusetts | —HL-98188 |
|---|-----------|

NHLBI Progenitor Cell Biology Consortium Research Hubs, Initiated in Fiscal Year 2009

The purpose of this study is to establish virtual research hubs that focus on progenitor cell biology. Investigators are seeking to identify and characterize progenitor cell lineages, direct the differentiation of stem and progenitor cells to desired cell fates, and develop new strategies to address the unique challenges presented by the transplantation of progenitor cells.

Obligations

Funding History:

Fiscal Year 2011—\$22,514,886

Fiscal Years 2009–2010—\$46,788,094

Total Funding to Date—\$69,302,980

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Children's Hospital of Philadelphia
Philadelphia, Pennsylvania | —HL-099656 |
| 2. University of Wisconsin, Madison
Madison, Wisconsin | —HL-099773 |
| 3. Johns Hopkins University
Baltimore, Maryland | —HL-099775 |
| 4. Stanford University
Stanford, California | —HL-099776 |
| 5. Fred Hutchinson Cancer Research Center
Seattle, Washington | —HL-099993 |

6. Stanford University Stanford, California	—HL-099995
7. University of Maryland Baltimore, Maryland	—HL-099997
8. Stanford University Stanford, California	—HL-099999
9. Children's Hospital Boston Boston, Massachusetts	—HL-100001
10. Fred Hutchinson Cancer Research Center Seattle, Washington	—HL-100395
11. Stanford University Stanford, California	—HL-100397
12. Vanderbilt University Nashville, Tennessee	—HL-100398
13. University of Texas Southwestern Medical Center Dallas, Texas	—HL-100401
14. Massachusetts General Hospital Boston, Massachusetts	—HL-100402
15. University of Pennsylvania Philadelphia, Pennsylvania	—HL-100405
16. J. David Gladstone Institutes San Francisco, California	—HL-100406
17. University of Minnesota, Twin Cities Minneapolis, Minnesota	—HL-100407
18. Massachusetts General Hospital Boston, Massachusetts	—HL-100408

Obesity Related Behavioral Intervention Trials (ORBIT),* Initiated in Fiscal Year 2009

The purpose of this study is to translate findings from basic research on human behavior into more effective clinical, community, and population interventions to reduce obesity and improve obesity-related behaviors. Investigators are seeking to develop innovative obesity-reducing strategies that are effective in small-scale trials, acceptable to target populations of interest and are ready for testing in large-scale randomized clinical and community trials. Some of the projects are expected to have 50–100 percent participation from minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$4,297,228

Fiscal Years 2009–2010—\$10,987,451

Total Funding to Date—\$15,284,679

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. Claremont Graduate University
Claremont, California | —HL-097839 |
|---|------------|

* Formerly known as Translating Basic Behavioral and Social Sciences Discoveries Into Interventions To Reduce Obesity.

2. Weill Medical College of Cornell University New York, New York	—HL-097843
3. Wayne State University Detroit, Michigan	—HL-097889
4. Rush University Medical Center Chicago, Illinois	—HL-097894
5. University of California, San Francisco San Francisco, California	—HL-097973

Pediatric Heart Network, Initiated in Fiscal Year 2006

See Chapter 11. Clinical Trials.

Pharmacogenetics Research Network, Initiated in Fiscal Year 2001

The purpose of this program is to establish a network of multidisciplinary, collaborative research groups to study how genetic variation contributes to interindividual differences in responses to medications. Four studies under this initiative are investigating the pharmacogenetics of heart, lung, and blood diseases. One of the projects has 38 percent minority participation. The Pharmacogenetics Knowledgebase (PharmGKB) has been established to integrate information obtained from pharmacogenomics, phenotypes, and genotypes.

Obligations

Funding History:

Fiscal Year 2011—\$6,532,424

Fiscal Years 2001–2010—\$74,421,366

Total Funding to Date—\$80,953,760

Current Active Organizations and Grant Numbers

1. Brigham and Women's Hospital and Harvard Medical School Boston, Massachusetts	—HL-065899
2. Vanderbilt University Nashville, Tennessee	—HL-065962
3. Children's Hospital and Research Center Oakland, California	—HL-069757
4. University of Maryland Baltimore, Maryland	—HL-105198

Ranolazine in Implantable Defibrillators (RAID) Trial, Initiated in Fiscal Year 2010

The purpose of this clinical trial is to determine whether ranolazine will reduce the risk of ventricular arrhythmias and improve survival in high-risk patients who already have an implantable cardiac defibrillator. Currently,

very few options are available for treating patients at risk for ventricular arrhythmias—which often leads to death—and ranolazine may be a safe and effective treatment.

Obligations

Funding History:

Fiscal Year 2011—\$0

Fiscal Year 2010—\$2,279,794

Total Funding to Date—\$2,279,794

Current Active Organizations and Grant Numbers

1. University of Rochester Rochester, New York	—HL-096607
2. University of Rochester Rochester, New York	—HL-096610

Rule Out Myocardial Infarction Using Computed Assisted Tomography (ROMICAT II), Initiated in Fiscal Year 2009

The purpose of this study is to determine whether using cardiac computed assisted tomography early in the emergency department triage will enable immediate and safe discharge without further testing of a significant number of patients with acute chest pain.

Obligations

Funding History:

Fiscal Year 2011—\$366,525

Fiscal Years 2009–2010—\$4,419,188

Total Funding to Date—\$4,785,713

Current Active Organizations and Grant Numbers

1. Massachusetts General Hospital Boston, Massachusetts	—HL-092022
2. Massachusetts General Hospital Boston, Massachusetts	—HL-092040

Strong Heart Study, Initiated in Fiscal Year 1988

The objectives of this study are to survey CVD morbidity and mortality rates among three geographically diverse groups of American Indians and estimate their levels of CVD risk factors. Phases II and III of the cohort study extended surveillance of community mortality and assessed development of CVD and changes in CVD risk factors. In Phase III, investigators added a substudy of asthma and a pilot family study. Phase IV expanded the family study to 120 families comprising 3,600 members to investigate genetic and environmental contributors of CVD. Phase V examined the family study cohort to

assess genetic relationships to risk factor change over a 5-year period and initiated surveillance for cardiovascular morbidity and mortality.

Obligations

Funding History:

Fiscal Year 2011—\$2,144,794

Fiscal Years 1988–2010—\$73,879,041

Total Funding to Date—\$76,023,835

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. Medstar Research Institute
Hyattsville, Maryland | —HL-041642 |
| 2. Missouri Breaks Research, Inc.
Timberlake, South Dakota | —HL-041652 |
| 3. University of Oklahoma
Oklahoma City, Oklahoma | —HL-041654 |
| 4. Weill Medical College of
Cornell University
New York, New York | —HL-065521 |

Surgical Treatment for Ischemic Heart Failure (STICH), Initiated in Fiscal Year 2002

The objective of this clinical trial is to continue to follow for 5 more years, patients who were assigned to the first part of the STICH study that sought to determine whether the addition of CABG to intensive medical therapy improves long-term survival of patients with heart failure and left ventricular dysfunction. The original study found no significant difference between medical therapy alone and medical therapy plus CABG with respect to the primary endpoint of death from any cause. However, the secondary endpoint related to deaths due to cardiovascular causes showed that patients assigned to CABG had lower rates of death compared with those assigned to medical therapy alone.

The extension study (STICHES) will enable investigators to obtain long-term (10-year average) data on survivors from the original cohort. The primary endpoint is all-cause mortality. Secondary endpoints include cardiovascular mortality; cardiovascular morbidity defined by hospitalization for heart failure, myocardial infarction, or need for coronary revascularization procedure; and functional status and symptoms. Approximately 25 percent of the population will come from minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$352,198

Fiscal Years 2002–2010—\$40,041,062

Total Funding to Date—\$40,393,260

Current Active Organization and Grant Number

- | | |
|--|------------|
| 1. Duke University
Durham, North Carolina | —HL-069015 |
|--|------------|

Therapeutic Hypothermia After Pediatric Cardiac Arrest (THAPCA) Trials, Initiated in Fiscal Year 2009

The purpose of this program is to determine whether therapeutic hypothermia after pediatric cardiac arrest improves outcomes, including survival, in infants and children. Approximately 50 percent of the patients are expected to come from racial and ethnic minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$4,401,772

Fiscal Years 2009–2010—\$5,993,723

Total Funding to Date—\$10,395,495

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. University of Utah
Salt Lake City, Utah | —HL-094339 |
| 2. University of Michigan at Ann Arbor
Ann Arbor, Michigan | —HL-094345 |

Lung Diseases Program

Asthma Networks (AsthmaNet), Initiated in Fiscal Year 2009

See Chapter 11. Clinical Trials.

COPD Clinical Research Network, Initiated in Fiscal Year 2003

See Chapter 11. Clinical Trials.

Heart and Lung Failure-Pediatric Insulin Titration Trial (HALF-PINT), Initiated in Fiscal Year 2011

The purpose of this study is to determine whether safe and effective tight glycemic control can sufficiently reduce morbidity and mortality in children with heart and lung failure to justify a low risk of hypoglycemia.

Obligations

Funding History:

Fiscal Year 2011—\$2,685,460

Total Funding to Date—\$2,685,460

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Children's Hospital Boston
Boston, Massachusetts | —HL-107681 |
| 2. Children's Hospital Boston
Boston, Massachusetts | —HL-108028 |

Infant Study of Inhaled Saline in Cystic Fibrosis (ISIS), Initiated in Fiscal Year 2008

The purpose of this randomized clinical trial is to assess the efficacy and safety of 7 percent hypertonic saline inhaled twice daily for 48 weeks among infants with CF aged 4 to 15 months at enrollment. Investigators are seeking to determine whether hypertonic saline will improve hyperinflation and obstructive lung disease as measured by infant lung function testing compared with the control agent (isotonic saline).

Obligations

Funding History:

- Fiscal Year 2011—\$628,241
Fiscal Years 2008–2010—\$2,224,874
Total Funding to Date—\$2,873,115

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. Children's Hospital and Regional Medical Center
Seattle, Washington | —HL-092931 |
| 2. University of Washington
Seattle, Washington | —HL-092932 |

Microbiome of the Lung and Respiratory Tract in HIV-Infected Individuals and HIV-Uninfected Controls, Initiated in Fiscal Year 2009

The purpose of this study is to characterize the microbiome of the lung alone or in combination with the upper airways in HIV-infected individuals and matched HIV-uninfected controls. Investigators will use molecular techniques to identify bacteria, and if possible, other organisms (e.g., viruses, cell-wall deficient organisms, protozoa, and fungi). Data will be used to examine the effects of changes in the respiratory microbiome on the pathogenesis and progression of HIV disease, HIV-related respiratory complications, and anti-HIV therapies.

Obligations

Funding History:

- Fiscal Year 2011—\$5,324,656
Fiscal Years 2009–2010—\$10,743,005
Total Funding to Date—\$16,067,661

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. University of Pennsylvania
Philadelphia, Pennsylvania | —HL-098957 |
| 2. George Washington University
Washington, DC | —HL-098958 |
| 3. Indiana University-Purdue University,
Indianapolis
Indianapolis, Indiana | —HL-098960 |
| 4. University of Michigan, Ann Arbor
Ann Arbor, Michigan | —HL-098961 |
| 5. University of Pittsburgh
Pittsburgh, Pennsylvania | —HL-098962 |
| 6. University of California, San Francisco
San Francisco, California | —HL-098964 |
| 7. University of Colorado
Denver, Colorado | —HL-098966 |

Novel Therapies for Lung Diseases—Phase II, Initiated in Fiscal Year 2010

See Chapter 11. Clinical Trials.

Pharmacogenetics of Asthma Treatment, Initiated in Fiscal Year 2000

The objective of this project is to bring together research experts in asthma, epidemiology, statistics, bioinformatics, physiology, clinical trials, genetics, and genomics to focus on the pharmacogenetics of asthma treatment.

Obligations

Funding History:

- Fiscal Year 2011—\$1,957,381
Fiscal Years 2000–2010—\$28,998,338
Total Funding to Date—\$30,955,719

Current Active Organization and Grant Number

- | | |
|--|------------|
| 1. Brigham and Women's Hospital
Boston, Massachusetts | —HL-065899 |
|--|------------|

Prematurity and Respiratory Outcomes Program (PROP), Initiated in Fiscal Year 2010

The purpose of this observational clinical study is to investigate hypotheses on the molecular mechanisms that contribute to respiratory disease risk of the premature newborn with the long-term goal of improving outcomes in the first year of life.

Obligations

Funding History:

- Fiscal Year 2011—\$4,066,948
Fiscal Year 2010—\$1,597,280
Total Funding to Date—\$5,664,228

Current Active Organizations and Grant Numbers

1. Vanderbilt University Nashville, Tennessee	—HL-101456
2. Washington University St. Louis, Missouri	—HL-101465
3. University of Pennsylvania Philadelphia, Pennsylvania	—HL-101794
4. University of California, San Francisco San Francisco, California	—HL-101798
5. Children's Hospital Medical Center, Cincinnati Cincinnati, Ohio	—HL-101800
6. University of Rochester Rochester, New York	—HL-101813

Preterm Birth in Nulliparous Women: An Understudied Population at Great Risk, Initiated in Fiscal Year 2010

The purpose of this study is to create a network of clinical research sites with a Data Coordinating and Analysis Center to develop common research protocols to study the cardiovascular health of women in their first pregnancy and assess the significance of disordered breathing.

Obligations

Funding History:

Fiscal Year 2011—\$3,298,000

Fiscal Year 2010—\$500,000

Total Funding to Date—\$3,798,000

Current Active Organization and Grant Number

1. Research Triangle Institute Research Triangle, North Carolina	—HD-063036
---	------------

Randomized Trial of Antenatal Late Preterm Steroids (ALPS), Initiated in Fiscal Year 2010

The purpose of this study is to determine whether antenatal corticosteroids can potentially improve lung function and reduce respiratory morbidity in newborn infants who are born in the late preterm period (34–36 weeks). Previous studies have shown that steroids improve lung function in very premature infants. Fifty-five percent of the participants are expected to come from racial and ethnic minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$2,137,414

Fiscal Year 2010—\$2,133,947

Total Funding to Date—\$4,271,361

Current Active Organizations and Grant Numbers

1. George Washington University Washington, DC	—HL-098354
2. Columbia University Health Sciences New York, New York	—HL-098554

Randomized Trial of Maternal Vitamin D Supplementation To Prevent Childhood Asthma, Initiated in Fiscal Year 2009

The purpose of this randomized clinical trial is to determine whether supplemental vitamin D to increase the level of vitamin D in a pregnant woman will prevent asthma and allergy in her child at age 3 years. Investigators will recruit 870 pregnant women who are in the first trimester of pregnancy and randomize them to one of two treatment arms of a 4-year clinical trial: one arm being treatment with 4,000 international units of vitamin D in addition to typical prenatal vitamins and the other being treatment with typical prenatal vitamins alone. Currently, 80 percent of the participants are from racial and ethnic minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$2,469,339

Fiscal Years 2009–2010—\$4,975,916

Total Funding to Date—\$7,445,255

Current Active Organizations and Grant Numbers

1. Washington University St. Louis, Missouri	—HL-091075
2. Brigham and Women's Hospital Boston, Massachusetts	—HL-091528

Sedation Management in Pediatric Patients With Acute Respiratory Failure, Initiated in Fiscal Year 2008

The purpose of this randomized clinical trial is to test an innovative approach to sedation management that includes team education and consensus on the use of sedatives in pediatric patients supported on mechanical ventilation; team identification of each patient's trajectory of illness and daily prescription of a sedation goal; use of a nurse-implemented goal-directed comfort algorithm that guides moment-to-moment titration of opioids and benzodiazepines; and team feedback on sedation management performance. Investigators have randomized 2,754 critically ill infants and children into two study groups: sedation management intervention

and usual care. Forty-five percent of the patients are from racial and ethnic minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$2,181,199
Fiscal Years 2008–2010—\$7,779,116
Total Funding to Date—\$9,960,315

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. University of Pennsylvania
Philadelphia, Pennsylvania | —HL-086622 |
| 2. Children's Hospital Boston
Boston, Massachusetts | —HL-086649 |

Severe Asthma Research Program,* Initiated in Fiscal Year 2011

The purpose of this study is to define severe asthma at the molecular and cellular levels longitudinally to understand its evolution. Research findings will serve as a rational basis for designing mechanism-based diagnostic, prognostic, and treatment strategies for severe asthma.

Obligations

Funding History:

Fiscal Year 2011—\$5,159,933
Total Funding to Date—\$5,159,933

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Pennsylvania State University,
Hershey Medical Center
Hershey, Pennsylvania | —HL-109086 |
| 2. University of California, San Francisco
San Francisco, California | —HL-109146 |
| 3. University of Pittsburgh, Pittsburgh
Pittsburgh, Pennsylvania | —HL-109152 |
| 4. Wake Forest University Health Sciences
Winston-Salem, North Carolina | —HL-109164 |
| 5. University of Wisconsin, Madison
Madison, Wisconsin | —HL-109168 |
| 6. Brigham and Women's Hospital
Boston, Massachusetts | —HL-109172 |
| 7. University of Virginia, Charlottesville
Charlottesville, Virginia | —HL-109250 |
| 8. Washington University
St. Louis, Missouri | —HL-109257 |

Study of Asthma and Nasal Steroids (STAN), Initiated in Fiscal Year 2009

The purpose of this clinical trial is to determine whether treatment of chronic rhinitis and sinusitis with a nasal steroid improves asthma control. Investigators have randomized 380 patients with poorly controlled asthma and chronic rhinitis and sinusitis to a nasal steroid or matching placebo in addition to their regular asthma treatment. One third of participants are expected to be from minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$724,884
Fiscal Years 2009–2010—\$1,449,536
Total Funding to Date—\$2,174,420

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. University of Vermont
Burlington, Vermont | —HL-089464 |
| 2. Johns Hopkins University
Baltimore, Maryland | —HL-089510 |

Study of Soy Isoflavones in Asthma (SOYA), Initiated in Fiscal Year 2009

The purpose of this double-blind, randomized controlled trial is to determine whether genistein supplements (soy isoflavone) improves lung function in patients with poorly controlled asthma. The study includes 380 patients with low dietary soy intake, ages 12 years and older, who are taking either inhaled corticosteroids or leukotriene modifiers or both and have poorly controlled asthma. Participants are being randomly assigned to treatment with either a soy isoflavone supplement (containing genistein, daidzein, and glycinein) 100 mg daily or to placebo for 6 months. Thirty percent of participants are expected to be from minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$688,673
Fiscal Years 2009–2010—\$1,417,854
Total Funding to Date—\$2,106,527

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Northwestern University
Evanston, Illinois | —HL-087987 |
| 2. Johns Hopkins University
Baltimore, Maryland | —HL-088367 |

* The Severe Asthma Research Program began in FY 2001 and was funded under the R01 mechanism.

Trial of Late Surfactant (TOLSURF) To Prevent Bronchopulmonary Dysplasia, Initiated in Fiscal Year 2009

The purpose of this randomized controlled clinical trial is to determine whether late doses of surfactant in addition to iNO administered to extremely low gestational age neonates (< 30 weeks) who require mechanical ventilation between 7 and 14 days of age will increase survival without bronchopulmonary dysplasia.

Obligations

Funding History:

Fiscal Year 2011—\$1,807,148

Fiscal Years 2009–2010—\$3,765,851

Total Funding to Date—\$5,572,999

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. University of California, San Francisco
San Francisco, California | —HL-094338 |
| 2. University of California, San Francisco
San Francisco, California | —HL-094355 |

Blood Diseases and Resources

Acute Venous Thrombosis: Thrombus Removal With Adjunctive Catheter-Directed Thrombolysis (ATTRACT),* Initiated in Fiscal Year 2008

The purpose of this study is to determine whether adjunctive pharmacomechanical catheter-directed thrombolysis (PCDT), which includes intrathrombus administration of recombinant tissue plasminogen activator, can prevent postthrombotic syndrome in patients with symptomatic proximal deep vein thrombosis (DVT). Investigators are comparing the addition of PCDT to optimal standard DVT therapy with optimal standard DVT therapy alone.

Obligations

Funding History:

Fiscal Year 2011—\$243,775

Fiscal Years 2008–2010—\$6,272,537

Total Funding to Date—\$6,516,312

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. McMaster University
Hamilton, Ontario | —HL-088118 |
| 2. Washington University
St. Louis, Missouri | —HL-088476 |

Blood and Marrow Transplant Clinical Research Network, Initiated in Fiscal Year 2001

See Chapter 11. Clinical Trials.

Bridging Anticoagulation on Patients Requiring Temporary Interruption of Warfarin Therapy for an Elective Invasive Procedure or Surgery (BRIDGE) Trial, Initiated in Fiscal Year 2008

The purpose of this study is to determine the safety and efficacy of low molecular weight heparin (LMWH) in adults with atrial fibrillation who stop warfarin in preparation for surgery. The trial randomly allocated 3,282 patients with atrial fibrillation to either LMWH or placebo before and after surgery. Investigators hypothesize that simply withholding warfarin in a perioperative setting for patients with atrial fibrillation will not meaningfully increase the risk for arterial thromboembolism and will forestall hemorrhagic complications, compared with a strategy using LMWH before and after surgery. One-third of participants are expected to be from minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$0

Fiscal Years 2008–2010—\$15,531,903

Total Funding to Date—\$15,531,903

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. Duke University
Durham, North Carolina | —HL-086755 |
| 2. Duke University
Durham, North Carolina | —HL-087229 |

Transfusion Medicine/Hemostasis Clinical Research Network, Initiated in Fiscal Year 2002

See Chapter 11. Clinical Trials.

* Formerly known as Pharmacomechanical Catheter-Directed Thrombolysis for Acute DVT (ATTRACT) Trial.

NHLBI Centers of Research Program (P50)

The Centers of Research Program supports specialized centers that focus on multidisciplinary research and development from basic science to clinical investigation in response to announcements of the programmatic needs of the Institute. The spectrum of activities comprises a multifaceted attack on a specific disease entity or biomedical problem area.

NHLBI Centers of Research Program

Type of Center	Period of Operation	Obligations (Dollars in Thousands)		
		Prior to FY 2011	FY 2011	Total to Date
NIH Centers for Population Health and Health Disparities (CPHHD)	2010–	\$9,898	\$10,169	\$20,067
Subtotal, CPHHD		\$9,898	\$10,169	\$20,067
Specialized Centers of Clinically Oriented Research (P50)				
<i>Heart and Vascular Diseases Program</i>				
Vascular Injury, Repair, and Remodeling	2006–	69,643	1,760	71,403
Subtotal, Heart and Vascular Diseases Program		69,643	1,760	71,403
<i>Lung Diseases Program</i>				
Chronic Obstructive Pulmonary Disease	2007–	44,781	8,775	53,556
Pulmonary Vascular Disease	2007–	25,558	4,660	30,218
Subtotal, Lung Diseases Program		70,339	13,435	83,774
Subtotal, SCCOR (P50)		\$139,982	\$15,195	\$155,177
Centers for Advanced Diagnostics and Experimental Therapeutics in Lung Diseases (CADET)				
Therapeutics in Lung Diseases (CADET)	2011	—	\$9,000	\$9,000
Subtotal, CADET		—	\$9,000	\$9,000
Total, Centers of Research Program (P50)		\$149,880	\$34,364	\$184,244

NIH Centers for Population Health and Health Disparities (CPHHD) (P50)

The purpose of this program is to create centers of transdisciplinary research that will evaluate the multilevel determinants of health disparities and devise interventions to reduce them.

Obligations

Fiscal Year 2011—\$10,168,814

Current Active Organizations and Grant Numbers

- | | | | |
|---|------------|---|------------|
| 1. University of North Carolina, Chapel Hill
Chapel Hill, North Carolina | —HL-105184 | 3. Johns Hopkins University
Baltimore, Maryland | —HL-105187 |
| 2. Northeastern University
Boston, Massachusetts | —HL-105185 | 4. University of California, Los Angeles
Los Angeles, California | —HL-105188 |
| | | 5. Rush University Medical Center
Chicago, Illinois | —HL-105189 |

Specialized Centers of Clinically Oriented Research (P50)

The NHLBI initiated the Specialized Centers of Research (SCOR) program in 1971 to encourage translational research—converting basic science findings to the clinic—in high priority areas. The SCOR concept emphasized multi-disciplinary research (i.e., basic science and clinical investigations) on diseases relevant to the Institute's mission. In FY 2002, the NHLBI revised its SCOR program—primarily on recommendation from the NHLBAC—to place more emphasis on clinical research projects. The SCCOR program still requires clinical and basic scientists to work together on a unified theme, but also requires at least 50 percent of the projects to be clinical. The SCOR program ended in FY 2008.

A description of the SCCORs supported by the Institute follows.

Heart Diseases Program

Vascular Injury, Repair, and Remodeling

The purpose of this SCCOR is to foster multidisciplinary research on clinically relevant questions that will enable basic science findings to be more rapidly translated into clinical applications. Major goals of the program are to stimulate interdependent clinical and multidisciplinary basic research projects that investigate molecular and cellular mechanisms of vascular injury, repair, and remodeling; promote patient-oriented research that will improve our ability to prevent, detect, characterize, manage, and treat vascular diseases; and develop the skills and research capabilities of new clinical investigators.

Obligations

Fiscal Year 2011—\$1,759,969

Current Active Organization and Grant Number

- | | |
|--|------------|
| 1. Beth Israel Deaconess Medical Center
Boston, Massachusetts | —HL-083813 |
|--|------------|

Lung Diseases Program

Chronic Obstructive Pulmonary Disease

The purpose of this SCCOR is to foster multidisciplinary research to accelerate progress in the diagnosis, prevention, and treatment of COPD. The program includes a broad spectrum of basic and clinical research that encompasses animal models of COPD pathogenesis; human proteomic, genetic, and genomic investigations;

disease phenotypes classification; and the development of new therapeutic interventions.

Obligations

Fiscal Year 2011—\$8,775,106

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. Washington University
St. Louis, Missouri | —HL-084922 |
| 2. Weill Medical College of
Cornell University
New York, New York | —HL-084936 |
| 3. Johns Hopkins University
Baltimore, Maryland | —HL-084945 |
| 4. University of Pittsburgh
Pittsburgh, Pennsylvania | —HL-084948 |

Pulmonary Vascular Disease

The objective of this SCCOR is to facilitate multidisciplinary research that proposes original hypotheses and applies cutting-edge approaches, including genomics and proteomics, to clinical issues in pulmonary vascular disease.

Obligations

Fiscal Year 2011—\$4,660,206

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. University of Colorado, Denver
Denver, Colorado | —HL-084923 |
| 2. Johns Hopkins University
Baltimore, Maryland | —HL-084946 |

Centers for Advanced Diagnostics and Experimental Therapeutics in Lung Diseases (CADET) (P50)

The purpose of this program is to accelerate the development of novel agents for the diagnosis and treatment of lung diseases and sleep disordered breathing through the use of rational strategies based on fundamental pathobiologic processes. CADET I provides opportunities to explore potential targets for validation to determine which are amenable for development of mechanism-based modalities for direct clinical application in the prevention, diagnosis, and treatment of pulmonary diseases and sleep disordered breathing.

Obligations

Fiscal Year 2011—\$9,000,370

Current Active Organizations and Grant Numbers

1. Children's Hospital Medical Center Cincinnati Cincinnati, Ohio	—HL-107159	11. University of Alabama, Birmingham Birmingham, Alabama	—HL-107181
2. University of Chicago Chicago, Illinois	—HL-107160	12. Johns Hopkins University Baltimore, Maryland	—HL-107182
3. Brigham and Women's Hospital Boston, Massachusetts	—HL-107165	13. Washington University Saint Louis, Missouri	—HL-107183
4. Brigham and Women's Hospital Boston, Massachusetts	—HL-107166	14. Johns Hopkins University Baltimore, Maryland	—HL-107185
5. University of North Carolina, Chapel Hill Chapel Hill, North Carolina	—HL-107168	15. University of Texas Health Center at Tyler Tyler, Texas	—HL-107186
6. Johns Hopkins University Baltimore, Maryland	—HL-107169	16. University of Arizona Tucson, Arizona	—HL-107188
7. University of Chicago Chicago, Illinois	—HL-107171	17. Johns Hopkins University Baltimore, Maryland	—HL-107190
8. University of Pittsburgh Pittsburgh, Pennsylvania	—HL-107172	18. University of California, San Francisco San Francisco, California	—HL-107191
9. University of Michigan, Ann Arbor Ann Arbor, Michigan	—HL-107177	19. Brigham and Women's Hospital Boston, Massachusetts	—HL-107192
10. Duke University Durham, North Carolina	—HL-107180		

Basic and Translational Research Program (U54)

The NHLBI reconfigured the Comprehensive Sickle Cell Centers program into a Basic and Translational Research Program (BTRP). The Program emphasizes fundamental investigations and their translation into initial studies in humans, as well as community translation to promote evidence-based clinical practice. The BTRP continues to support the Sickle Cell Disease Scholars program for the career development of young investigators and the Summer-for-Sickle Cell-Science program for research training and mentoring of high-school students. These components are part of a larger effort by the NHLBI to prepare the next generation of scientists to advance the field of SCD research.

Obligation

Fiscal Year 2011—\$5,901,312

Current Active Organizations and Grant Numbers

1. Thomas Jefferson University Philadelphia, Pennsylvania	—HL-070585	4. St. Jude's Children's Research Hospital Memphis, Tennessee	—HL-070590
2. Rho Federal Systems Division, Inc. Chapel Hill, North Carolina	—HL-070587	5. Boston Medical Center Boston, Massachusetts	—HL-070819
3. University of Texas, Southwest Medical Center Dallas, Texas	—HL-070588	6. Children's Hospital Medical Center Cincinnati, Ohio	—HL-070871

7. Medical College of Wisconsin Milwaukee, Wisconsin	—HL-090503	11. Johns Hopkins University Baltimore, Maryland	—HL-090515
8. Howard University Washington, DC	—HL-090508	12. Virginia Commonwealth University Richmond, Virginia	—HL-090516
9. Children's Hospital of Los Angeles Los Angeles, California	—HL-090511	13. University of Miami, School of Medicine Coral Gables, Florida	—HL-090569
10. University of Illinois, at Chicago Chicago, Illinois	—HL-090513		

Cardiac Translational Research Implementation Program (C-TRIP) (P20)

The C-TRIP program was initiated in FY 2010 to accelerate the translation of promising new therapeutic interventions derived from fundamental research discoveries for the treatment and prevention of heart failure or arrhythmias. The program consists of two stages. Stage 1 focuses on planning and developing clinical trials to determine the safety and efficacy of interventions to be conducted during Stage 2 of the overall program. Stage 2 studies will be supported by the P50 mechanism.

Obligation

Fiscal Year 2011—\$8,592,776

Current Active Organizations and Grant Numbers

1. Mount Sinai School of Medicine New York, New York	—HL-100396	7. University of Colorado, Denver Aurora, Colorado	—HL-101435
2. Johns Hopkins University Baltimore, Maryland	—HL-101397	8. Texas Heart Institute Houston, Texas	—HL-101438
3. Brigham and Women's Hospital Boston, Massachusetts	—HL-101408	9. Mayo Clinic Rochester, Minnesota	—HL-101439
4. University of Medicine and Dentistry of New Jersey New Jersey Medical School Newark, New Jersey	—HL-101420	10. University of Miami School of Medicine Miami, Florida	—HL-101443
5. Vanderbilt University Nashville, Tennessee	—HL-101425	11. Children's Hospital of Philadelphia Philadelphia, Pennsylvania	—HL-101820
6. University of Maryland, Baltimore Baltimore, Maryland	—HL-101434	12. Brigham and Women's Hospital Boston, Massachusetts	—HL-101866

Centers for AIDS Research (P30) Program

The NHLBI, along with five other NIH Institutes, contributes to the support of six Centers for AIDS Research that were established to provide a multidisciplinary environment that promotes basic, clinical, behavioral, and translational research activities in the prevention, detection, and treatment of HIV infection and AIDS. Almost half of the patient population comes from minority groups.

Obligations

Fiscal Year 2011—\$3,333,747

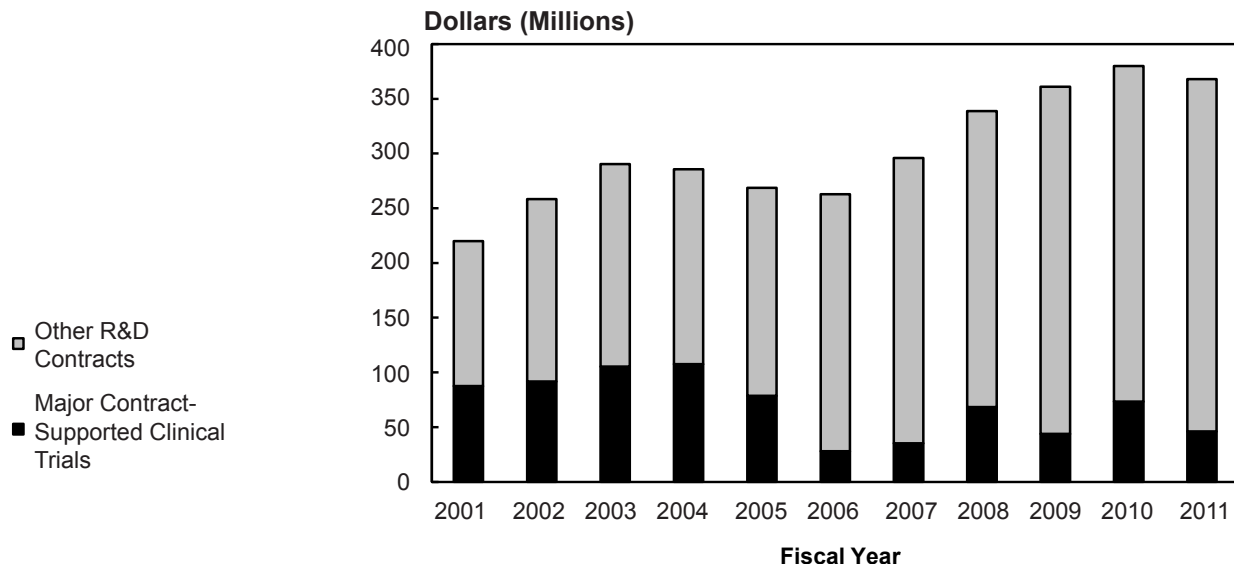
Current Active Organizations and Grant Numbers

1. University of Washington Seattle, Washington	—AI-027757	12. University of North Carolina, Chapel Hill Chapel Hill, North Carolina	—AI-050410
2. University of California, San Francisco San Francisco, California	—AI-027763	13. Yeshiva University New York, New York	—AI-051519
3. University of Alabama, Birmingham Birmingham, Alabama	—AI-027767	14. Vanderbilt University Nashville, Tennessee	—AI-054999
4. University of California, Los Angeles Los Angeles, California	—AI-028697	15. Harvard Medical School Boston, Massachusetts	—AI-060354
5. Baylor University Houston, Texas	—AI-036211	16. Duke University Durham, North Carolina	—AI-064518
6. University of California, La Jolla La Jolla, California	—AI-036214	17. University of Miami School of Medicine Coral Gables, Florida	—AI-073961
7. Case Western Reserve University Cleveland, Ohio	—AI-036219	18. University of Rochester Rochester, New York	—AI-078498
8. University of Massachusetts, Worcester Worcester, Massachusetts	—AI-042845	19. Rush University Medical Center Chicago, Illinois	—AI-082151
9. Miriam Hospital Providence, Rhode Island	—AI-042853	20. George Washington University Washington, DC	—AI-087714
10. University of Pennsylvania Philadelphia, Pennsylvania	—AI-045008		
11. Emory University Atlanta, Georgia	—AI-050409		



10. Research and Development Contracts

NHLBI Total Research and Development Contract Obligations:^{*} Fiscal Years 2001–2011



* For detailed data on contract-supported clinical trials, see Chapter 11.

NHLBI Total Research and Development Contract Obligations: Fiscal Years 2001–2011

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Heart	\$184,491	\$214,971	\$258,647	\$245,881	\$219,796	\$213,320	\$260,205	\$296,445	\$321,223	\$303,251	\$299,201
Lung	10,993	16,578	11,745	14,131	20,946	25,902	15,191	20,249	17,710	47,777	25,338
Blood	24,572	26,751	20,082	25,460	27,831	23,629	20,446	22,093	22,164	28,864	43,752
Total	\$220,056^A	\$258,300^B	\$290,474^C	\$285,472^D	\$268,573^E	\$262,851^F	\$295,842^G	\$338,787^H	\$361,097^I	\$379,892^J	\$368,291^K

A Includes Program Evaluation and IMPAC II Assessments of \$24,579,000.

B Includes Program Evaluation and IMPAC II Assessments of \$35,827,000.

C Includes Program Evaluation and IMPAC II Assessments of \$54,550,000.

D Includes Program Evaluation and IMPAC II Assessments of \$57,545,722.

E Includes Program Evaluation and IMPAC II Assessments of \$64,399,000.

F Includes Program Evaluation and IMPAC II Assessments of \$67,795,000.

G Includes Program Evaluation and IMPAC II Assessments of \$68,405,000.

H Includes Program Evaluation and IMPAC II Assessments of \$77,487,000.

I Includes Program Evaluation and IMPAC II Assessments of \$79,693,000.

J Includes Program Evaluation and IMPAC II Assessments of \$83,834,100.

K Includes Program Evaluation and IMPAC II Assessments of \$88,024,222.

Note: From 2001 to 2006 the WHI was reported separately. In this table, it has been incorporated in the “Heart” line.

Major NHLBI Research and Development Contracts by Program

	Total Obligations Prior to FY 2011	Total FY 2011 Obligations	Total Obligations to Date
Heart and Vascular Diseases			
Action to Control Cardiovascular Risk in Diabetes Follow-Up Study (ACCORDION)	—	\$13,250,149	\$13,250,149
Atherosclerosis Risk in Communities (ARIC)	157,168,999	15,012,384	172,181,383
Cardiovascular Health Study (CHS)	79,555,913	576,600	80,132,513
Coronary Artery Risk Development in Young Adults (CARDIA)	99,138,459	9,901,825	109,040,284
DNA Resequencing and Genotyping	29,209,023	9,274,939	38,483,962
Framingham Heart Study (FHS)	115,709,281	16,664,505	132,373,786
Genetically Triggered Thoracic Aortic Aneurysms and Other Cardiovascular Conditions (GenTAC): National Registry	11,115,085	3,763,038	14,878,123
Global Health Centers of Excellence	14,399,764	2,607,272	17,007,036
Hispanic Community Health Study (HCHS)	51,468,540	5,175,848	56,644,388
Interagency Registry for Mechanical Circulatory Support (INTERMACS)	7,826,196	5,299,999	13,126,195
Jackson Heart Study (JHS)	38,494,257	4,882,541	43,376,798
Multi-Ethnic Study of Atherosclerosis (MESA)	102,979,420	5,815,507	108,794,927
NHLBI Gene Therapy Resource Program (GTRP)	21,753,472	7,435,505	29,188,977
NHLBI Programs of Excellence in Nanotechnology	65,777,271	966,068	66,743,339
Proteomics Initiative	174,518,757	17,123,612	191,642,369
Pumps for Kids, Infants, and Neonates (PumpKIN)	8,388,522	10,163,382	18,551,904
Science Moving TowArds Research Translation and Therapy Program (SMARTT)	—	4,145,344	4,145,344
Lung Diseases			
Lung Tissue Research Consortium	30,000,000	7,852,716	37,852,716
Subpopulations and Intermediate Outcome Measures in COPD Study (SPIROMICS)	30,042,319	2,655,398	32,697,717
Blood Diseases and Resources			
Maintenance of NHLBI Biological Specimen Repository	19,332,708	3,209,369	22,542,077
NHLBI-CDC Registry and Surveillance System in Hemoglobinopathies (RuSH)	6,419,918	4,208,761	10,628,679
Production Assistance for Cellular Therapies (PACT)	10,826,333	11,472,212	22,298,545
Recipient Epidemiology and Donor Study III (REDS-III)*	—	19,910,527	19,910,527

* Formerly known as Retrovirus Epidemiology Donor Study.

Heart and Vascular Diseases

Action To Control Cardiovascular Risk in Diabetes Follow-Up Study (ACCORDION), Initiated in Fiscal Year 2011

The purpose of the follow-up observational study (ACCORDION) is to obtain long-term (10 year average) data on ACCORD participants. Investigators are seeking to determine whether differences in mortality, CVD events, and microvascular diseases identified during the ACCORD trial persist or change over time and whether other differences will emerge. They will monitor long-term vascular outcomes from diabetes and the effects of glucose, blood pressure lowering, and lipid treatment on those outcomes.

ACCORD was a randomized clinical trial to evaluate the ability of three treatment strategies (intensive glycemic control, intensive blood pressure control, and fibrate treatment to raise HDL-cholesterol and lower triglycerides) to prevent major CVD events in patients with type 2 diabetes who were at high risk of CVD.

After a mean 3.5 years of treatment, the intensive glycemic portion of the trial was stopped because patients in the intensive glycemic treatment group had an increased risk of all-cause mortality compared with patients in the standard treatment group even though they had a non-statistically significant 10 percent reduction in the composite primary outcome of nonfatal MI, nonfatal stroke, or CVD death. Participants in the intensive group were then transitioned to the standard treatment strategy. Initial follow-up found no overall benefit of intensive treatment (intensive blood pressure control or fibrate and statin) over standard treatment (normal blood pressure or statin).

Obligations

Funding History:

Fiscal Year 2011—\$13,250,149

Total Funding to Date—\$13,250,149*

Current Active Organizations and Contract Numbers

- | | |
|---|-----------|
| 1. Veterans Affairs Medical Center,
Albuquerque
Albuquerque, New Mexico | —HC-10100 |
| 2. Veterans Affairs Medical Center,
Memphis
Memphis, Tennessee | —HC-90350 |
| 3. Wake Forest University
Winston-Salem, North Carolina | —HC-95178 |

- | | |
|---|-----------|
| 4. McMaster University
Hamilton, Ontario | —HC-95179 |
| 5. University of Washington
Seattle, Washington | —HC-95180 |
| 6. Case Western Reserve University
Cleveland, Ohio | —HC-95181 |
| 7. Wake Forest University
Winston-Salem, North Carolina | —HC-95182 |
| 8. Minneapolis Medical Research
Foundation
Minneapolis, Minnesota | —HC-95183 |
| 9. Trustees of Columbia University of
New York
New York, New York | —HC-95184 |

Atherosclerosis Risk in Communities (ARIC), Initiated in Fiscal Year 1985

The ARIC is an epidemiology study comprising a prospective cohort component and a community surveillance component. The cohort component investigates the etiology of CHD and stroke in 15,792 participants, aged 46–64 years at baseline, by race and gender in four U.S. communities. The community surveillance component monitors trends in hospitalized myocardial infarction, fatal CHD, and heart failure (2005–2009) from the same communities.

In 2011, the study began to reexamine the cohort participants with a focus on heart failure—a major epidemic in the rapidly aging U.S. population. Three of the cohort components represent the racial mix of their respective communities, and the fourth is exclusively black.

Obligations

Funding History:

Fiscal Year 2011—\$15,012,384

Fiscal Years 1985–2010—\$157,168,999

Total Funding to Date—\$172,181,383

Current Active Organizations and Contract Numbers

- | | |
|--|-----------|
| 1. University of North Carolina at
Chapel Hill
Chapel Hill, North Carolina | —HC-55015 |
| 2. Baylor College of Medicine
Houston, Texas | —HC-55016 |
| 3. University of North Carolina at
Chapel Hill
Chapel Hill, North Carolina | —HC-55018 |
| 4. University of Minnesota, Twin Cities
Minneapolis, Minnesota | —HC-55019 |

* Total funding for ACCORD was \$142,587,546 from 1999 to 2010.

5. Johns Hopkins University Baltimore, Maryland	—HC-55020
6. Mississippi Medical Center Jackson, Mississippi	—HC-55021
7. Brigham and Women's Hospital Boston, Massachusetts	—HC-55273

Cardiovascular Health Study (CHS), Initiated in Fiscal Year 1988

The CHS is a population-based, longitudinal study of risk factors for development and progression of CHS and stroke in the elderly, 17 percent of whom are from minority populations. Extensive data and samples have been collected from nearly 6,000 participants since 1989–1990. The current CHS: Transition Phase provides partial support for an infrastructure to enable continued access to study resources and expertise, scientific collaborations, and mentorship of early-career investigators.

Obligations

Funding History:

Fiscal Year 2011—\$576,600
Fiscal Years 1988–2010—\$79,555,913
Total Funding to Date—\$80,132,513

Current Active Organization and Contract Number

1. University of Washington Seattle, Washington	—268200800007C
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Coronary Artery Risk Development in Young Adults (CARDIA), Initiated in Fiscal Year 1984

The CARDIA is a long-term study that examines the evolution of CVD risk factors and early clinical events in persons aged 18–30 years in 1985–1986. The study examines risk factors for heart and lung diseases by collecting information on body mass index, physical activity and lifestyle, genetics, cognitive functioning, serologic and metabolic components, inflammatory markers, and other subclinical measures of disease. Fifty percent of the participants are black.

Obligations

Funding History:

Fiscal Year 2011—\$9,901,825
Fiscal Years 1984–2010—\$99,138,459
Total Funding to Date—\$109,040,284

Current Active Organizations and Contract Numbers

1. Johns Hopkins University Baltimore, Maryland	—HC-45241
2. University of Alabama at Birmingham Birmingham, Alabama	—HC-48047
3. University of Minnesota, Twin Cities Minneapolis, Minnesota	—HC-48048
4. Northwestern University Chicago, Illinois	—HC-48049
5. Kaiser Permanente Division of Research Oakland, California	—HC-48050
6. University of Alabama at Birmingham Birmingham, Alabama	—HC-95095

DNA Resequencing and Genotyping Program, Initiated in Fiscal Year 2004

The purpose of this program is to provide high-quality, high-volume resequencing and genotyping of candidate genomic regions potentially important in the disease pathways of heart, lung, and blood diseases and sleep disorders. The information obtained will enable ongoing investigations to elucidate the specific genetic components involved in the causes for, variable outcomes of, and progression of the diseases and disorders.

Obligations

Funding History:

Fiscal Year 2011—\$9,274,939
Fiscal Years 2004–2010—\$29,209,023
Total Funding to Date—\$38,483,962

Current Active Organizations and Contract Numbers

1. Systems Research and Applications International, Inc. Rockville, Maryland	—268201100029C
2. University of Washington Seattle, Washington	—268201100037C

Framingham Heart Study (FHS), Initiated in Fiscal Year 1948

The original Framingham Heart Study was designed as a longitudinal investigation of constitutional and environmental factors influencing the development of CVD in individuals free of CVD symptoms at the outset. Of the original 5,209 participants, about 150 are still alive. In 1971, the Framingham Offspring Study was initiated to assess familial and genetic factors associated with CHD. More than 5,000 offspring (and their spouses) were included. In 2002, a third-generation cohort consisting of approximately 4,000 grandchildren was added

to permit examination of numerous hypotheses about the genetic contribution to CVD and CVD risk factors. Additional goals include identifying new risk factors for cardiovascular, lung, and blood diseases and developing new imaging tests that can detect very early stages of coronary atherosclerosis in otherwise healthy adults.

In 2009, the Omni Group 1 and Omni Group 2 cohorts were integrated into the NHLBI contract for the FHS. The Omni cohorts consist of minority residents of Framingham, Massachusetts (about 500 and 400 participants in Omni Group 1 and Omni Group 2, respectively), and were previously identified, recruited, and examined through NHLBI investigator-initiated grants. They were added to the FHS to reflect the growing diversity of the community.

Obligations

Funding History:

Fiscal Year 2011—\$16,664,505

Fiscal Years 1983–2010—\$115,709,281

Total Funding to Date—\$132,373,786

Current Active Organization and Contract Number

- | | |
|--|-----------|
| 1. Boston University Medical Center
Boston, Massachusetts | —HC-25195 |
|--|-----------|

Genetically Triggered Thoracic Aortic Aneurysms and Other Cardiovascular Conditions (GenTAC): National Registry, Initiated in Fiscal Year 2006

The purpose of this program is to improve the diagnosis and management of patients with thoracic aortic aneurysms and other cardiovascular complications associated with connective tissue diseases, such as Marfan, Loeys-Dietz, and Ehlers Danlos (vascular type) Syndromes. The GenTAC is establishing a medical database and biospecimens repository as a resource for qualified investigators to study the link between genes and aortic aneurysms and heart disease.

Obligations

Funding History:

Fiscal Year 2011—\$3,763,038

Fiscal Years 2006–2010—\$11,115,085

Total Funding to Date—\$14,878,123

Current Active Organization and Contract Number

- | | |
|--|-----------|
| 1. Research Triangle Institute International
Research Triangle Park, North Carolina | —HV-08238 |
|--|-----------|

Global Health Centers of Excellence, Initiated in Fiscal Year 2009

The purpose of this program is to support a worldwide network of research and training centers to prevent and control chronic cardiovascular and pulmonary diseases in developing countries. The NHLBI joined with Minneapolis-based UnitedHealth Group's Chronic Disease Initiative in establishing the UnitedHealth and NHLBI Collaborating Centers of Excellence network. Each center is led by a research institution in a developing country that is paired with at least one partner academic institution in a developed country to enhance research and training opportunities.

Obligations

Funding History

Fiscal Year 2011—\$2,607,272

Fiscal Years 2009–2010—\$14,399,764

Total Funding to Date—\$17,007,036

Current Active Organizations and Contract Numbers

- | | |
|---|-----------|
| 1. The George Institute for International Health, China
Beijing, China | —HV-98217 |
| 2. Instituto de Nutrición de Centro América y Panamá
Guatemala City, Guatemala | —HV-98218 |
| 3. University of Cape Town
Cape Town, South Africa | —HV-98220 |
| 4. Moi University School of Medicine
Eldoret, Kenya | —HV-98221 |
| 5. Universidad Peruana Cayetano Heredia
Lima, Peru | —HV-98223 |

Hispanic Community Health Study (HCHS), Initiated in Fiscal Year 2006

The purpose of this program is to determine the prevalence of and risk factors for cardiovascular and lung diseases in Hispanic populations and the role of cultural adaptation and disparities in development of these and other chronic diseases. The multicenter, 6.5-year epidemiology study comprises more than 16,400 Hispanics, aged 18–74 years, who self-identify as being of Mexican, Puerto Rican, Cuban, Dominican, or Central or South American heritage.

Obligations

Funding History:

Fiscal Year 2011—\$5,175,848

Fiscal Years 2006–2010—\$51,468,540

Total Funding to Date—\$56,644,388

Current Active Organizations and Contract Numbers

1. University of North Carolina at Chapel Hill Chapel Hill, North Carolina	—HC-65233
2. University of Miami Miami, Florida	—HC-65234
3. Albert Einstein College of Medicine New York, New York	—HC-65235
4. Northwestern University Chicago, Illinois	—HC-65236
5. San Diego State University San Diego, California	—HC-65237

Interagency Registry for Mechanical Circulatory Support (INTERMACS), Initiated in Fiscal Year 2005

The INTERMACS is a national registry for patients who are receiving mechanical circulatory support device (MCSD) therapy to treat advanced heart failure. The registry collects and analyzes clinical and laboratory data and tissue samples from patients who receive MCSDs as destination therapy for end-stage heart failure at 119 participating sites.

Obligations

Funding History:

Fiscal Year 2011—\$5,299,999
Fiscal Years 2005–2010—\$7,826,196
Total Funding to Date—\$13,126,195

Current Active Organization and Contract Number

1. University of Alabama Birmingham, Alabama	—HV-58198
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Jackson Heart Study (JHS), Initiated in Fiscal Year 1998

The JHS is an epidemiologic study of CVD in blacks in Jackson, Mississippi, similar to established studies in Framingham, Massachusetts, and Honolulu, Hawaii. The goal of the study is to identify factors related to the development and progression of CVD in blacks. The JHS conducts a variety of community education and outreach activities to promote healthy lifestyles to reduce disease burden. In addition, the JHS seeks to build research capabilities in minority institutions, address the critical shortage of minority investigators in epidemiology and prevention, and reduce barriers to dissemination and use of health information in a minority population.

Obligations

Funding History:

Fiscal Year 2011—\$4,882,541
Fiscal Years 1998–2010—\$38,494,257
Total Funding to Date—\$43,376,798

Current Active Organizations and Contract Numbers

1. Jackson State University Jackson, Mississippi	—HC-95170
2. Mississippi Medical Center Jackson, Mississippi	—HC-95171
3. Tougaloo College Tougaloo, Mississippi	—HC-95172

Multi-Ethnic Study of Atherosclerosis (MESA), Initiated in Fiscal Year 1999

The purpose of this study is to investigate the prevalence, correlates, and progression of subclinical CVD (i.e., disease detected noninvasively before it has produced clinical signs and symptoms). The cohort of 6,814 participants is 38 percent white, 28 percent black, 22 percent Hispanic, and 12 percent Asian. A fifth examination, completed in November 2011, included a repeat measurement of cardiac function with MRI to assess changes over time. Periodic monitoring of participants to identify recent hospitalizations and other clinical events will continue.

Obligations

Funding History:

Fiscal Year 2011—\$5,815,507
Fiscal Years 1999–2010—\$102,979,420
Total Funding to Date—\$108,794,927

Current Active Organizations and Contract Numbers

1. University of Washington Seattle, Washington	—HC-95159
2. Johns Hopkins University Baltimore, Maryland	—HC-95162
3. University of Vermont Colchester, Vermont	—HC-95166
4. Johns Hopkins University Baltimore, Maryland	—HC-95168

NHLBI Gene Therapy Resource Program (GTRP), Initiated in Fiscal Year 2007

The purpose of this program is to promote the translation of basic research into clinical trials. The program provides resources in the form of preclinical and clinical-grade vector production; pharmacology and toxicology testing

on animals; immunology testing; clinical trials funding assistance; and regulatory support for gene therapy research primarily in heart, lung, and blood diseases.

Obligations

Funding History:

Fiscal Year 2011—\$7,435,505

Fiscal Years 2008–2010—\$21,753,472

Total Funding to Date—\$29,188,977

Current Active Organizations and Contract Numbers

1. Social and Scientific Systems, Inc. Silver Spring, Maryland	—HV-78200
2. University of Pennsylvania Philadelphia, Pennsylvania	—HV-78202
3. Children's Hospital of Philadelphia Philadelphia, Pennsylvania	—HV-78203
4. Indiana University Indianapolis, Indiana	—HV-78204

NHLBI Programs of Excellence in Nanotechnology,* Initiated in Fiscal Year 2010

The purpose of this program is to support multi-disciplinary teams to develop nanotechnology and biomolecular engineering tools and methodologies for the diagnosis and treatment of heart, lung, and blood diseases. The focus of this phase of the program is to translate the technologies being developed toward clinical application. The program brings together bio-engineers, chemists, material scientists, biologists, and clinicians across 17 institutions for unique scientific collaborations and skills development opportunities.

Obligations

Funding History:

Fiscal Year 2011—\$966,068

Fiscal Year 2010—\$65,777,271

Total Funding to Date—\$66,743,339

Current Active Organization and Contract Number

1. Washington University St. Louis, Missouri	—268201000046C
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Proteomics Initiative, Initiated in Fiscal Year 2002

The purpose of this program is to establish highly interactive, multidisciplinary centers to enhance and develop innovative proteomic technologies directed to relevant biologic questions associated with heart, lung, blood, and sleep health and disease.

Obligations

Funding History:

Fiscal Year 2011—\$17,123,612

Fiscal Years 2002–2010—\$174,518,757

Total Funding to Date—\$191,642,369

Current Active Organizations and Contract Numbers

1. Boston University Boston, Massachusetts	—268201000031C
2. Johns Hopkins University Baltimore, Maryland	—268201000032C
3. Massachusetts General Hospital Boston, Massachusetts	—268201000033C
4. Stanford University Stanford, California	—268201000034C
5. University of California, Los Angeles Los Angeles, California	—268201000035C
6. University of Texas San Antonio, Texas	—268201000036C
7. University of Texas Galveston, Texas	—268201000037C

Pumps for Kids, Infants, and Neonates (PumpKIN), Initiated in Fiscal Year 2010

The purpose of this program is to support technologies that will expand life-saving options for infants and children who are born with congenital heart defects or those who develop heart failure. Investigators are seeking to complete animal studies and other preclinical tests for the most promising devices in order to gain approval from the FDA to begin clinical testing.

Obligations

Funding History:

Fiscal Year 2011—\$10,163,382

Fiscal Year 2010—\$8,388,522

Total Funding to Date—\$18,551,904

Current Active Organizations and Contract Numbers

1. University of Pittsburgh Pittsburgh, Pennsylvania	—268201000012C
2. Jarvik Heart, Inc. New York, New York	—268201000013C
3. University of Maryland Baltimore, Maryland	—268201000014C
4. Ension, Inc. Pittsburgh, Pennsylvania	—268201000015C

* NHLBI Programs of Excellence in Nanotechnology was a cooperative agreement from 2005 to 2009.

Science Moving TowArds Research Translation and Therapy Program (SMARTT), Initiated in Fiscal Year 2011

The purpose of this program is to support the transition of potential new therapies for heart, lung, and blood diseases from discovery in the lab to the testing needed to establish their safety and effectiveness in people. The SMARTT program provides tailored pharmacology and toxicology testing, manufacturing services, and regulatory support to investigators to expedite the transition of their discoveries to the clinic.

Obligations

Funding History:

Fiscal Year 2011—\$4,145,344
Total Funding to Date—\$4,145,344

Current Active Organizations and Contract Numbers

1. Advanced BioScience Laboratories, Inc.
Rockville, Maryland —268201100014C
2. SRI International
Menlo Park, California —268201100015C
3. Research Triangle Institute International
Research Triangle Park,
North Carolina —268201100016C
4. SRI International
Menlo Park, California —268201000017C

Lung Diseases

Lung Tissue Research Consortium, Initiated in Fiscal Year 2004

The purpose of this program is to establish a consortium for collecting lung tissues and preparing and distributing them for research. Scientists are seeking to improve management of lung diseases by increasing understanding of the pathogenetic mechanisms of lung diseases through molecular histopathological studies of tissues with and without disease. Primary emphases are on COPD and idiopathic pulmonary fibrosis.

Obligations

Funding History:

Fiscal Year 2011—\$7,852,716
Fiscal Years 2004–2010—\$30,000,000
Total Funding to Date—\$37,852,716

Current Active Organizations and Contract Numbers

1. University of Colorado Health Science Center
Denver, Colorado —HR-46159
2. University of Michigan
Ann Arbor, Michigan —HR-46207
3. Mayo Clinic College of Medicine
Rochester, New York —HR-46208
4. University of Pittsburgh
Pittsburgh, Pennsylvania —HR-46210
5. Clinical Trials and Survey Corporation
Baltimore, Maryland —HR-46211
6. Mayo Clinic College of Medicine
Rochester, New York —HR-46212

Subpopulations and Intermediate Outcome Measures in COPD Study (SPIROMICS), Initiated in Fiscal Year 2009

The purposes of this study are to define pathogenetically homogeneous subgroups of COPD subjects on the basis of biomarkers, genotypes, and computed tomography images and to identify immediate outcome measures for use in future clinical studies. Secondary aims are to clarify the natural history of COPD; develop bioinformatic resources that will enable the use and sharing of data in studies of COPD and related diseases; and create a collection of clinical, biomarker, radiographic, and genetic data that can be used by external investigators for other studies of COPD.

Obligations

Funding History:

Fiscal Year 2011—\$2,655,398
Fiscal Years 2009–2010—\$30,042,319
Total Funding to Date—\$32,697,717

Current Active Organizations and Contract Numbers

1. University of California, San Francisco
San Francisco, California —HR-96199
2. University of California, Los Angeles
Los Angeles, California —HR-96200
3. Columbia University
New York, New York —HR-96201
4. University of Michigan
Ann Arbor, Michigan —HR-96202
5. University of Utah
Salt Lake City, Utah —HR-96203
6. Wake Forest University
Winston-Salem, North Carolina —HR-96204
7. University of North Carolina, Chapel Hill
Chapel Hill, North Carolina —HR-96206

Blood Diseases and Resources

Maintenance of NHLBI Biological Specimen Repository, Initiated in Fiscal Year 1998

The purpose of this project is to establish an NHLBI Biological Specimen Repository for blood specimens from Institute-supported research. The Repository monitors storage, labeling, and testing of the specimens, and administers safe shipment of precise sample aliquots to approved investigators for future studies.

Obligations

Funding History:

Fiscal Year 2011—\$3,209,369

Fiscal Years 1998–2010—\$19,332,708

Total Funding to Date—\$22,542,077

Current Active Organization and Contract Number

- | | |
|--|-----------|
| 1. SeraCare Life Sciences, Inc.
Rockville, Maryland | —HB-87144 |
|--|-----------|

NHLBI–CDC Registry and Surveillance System in Hemoglobinopathies (RuSH), Initiated in Fiscal Year 2009

The purpose of this pilot program is to test the feasibility of developing a national data system that will enable investigators to estimate the number of people who have SCD, thalassemias, and hemoglobinopathies and to describe their sociodemographic characteristics. The Institute, along with the CDC, has created newborn screening programs with State health departments in California, Florida, Georgia, Michigan, New York, North Carolina, and Pennsylvania.

Obligations

Funding History:

Fiscal Year 2011—\$4,208,761

Fiscal Year 2010—\$6,419,918

Total Funding to Date—\$10,628,679

Current Active Organization and Contract Number

- | | |
|---|----------|
| 1. Centers for Disease Control and Prevention
Atlanta, Georgia | —HR-9045 |
|---|----------|

Production Assistance for Cellular Therapies (PACT), Initiated in Fiscal Year 2010

The purpose of this program is to facilitate the transfer of innovative cellular therapies from the bench to the bedside. The PACT offers assistance to investigators in areas ranging from translational development to production of a product for use in human clinical trials.

Obligations

Funding History:

Fiscal Year 2011—\$11,472,212

Fiscal Year 2010—\$10,826,333

Total Funding to Date—\$22,298,545

Current Active Organizations and Contract Numbers

- | | |
|---|----------------|
| 1. EMMES Corp.
Rockville, Maryland | —268201000006C |
| 2. Baylor College of Medicine
Houston, Texas | —268201000007C |
| 3. University of Minnesota
Minneapolis, Minnesota | —268201000008C |
| 4. Immune Disease Institutes
Boston, Massachusetts | —268201000009C |
| 5. University of Wisconsin
Madison, Wisconsin | —268201000010C |
| 6. Beckman Research Institutes
Duarte, California | —268201000011C |

Recipient Epidemiology and Donor Study-III (REDS-III),* Initiated in Fiscal Year 2011

The purpose of this program is to conduct research to improve transfusion practices and the safety and adequacy of the blood supply in the United States and in countries affected by the AIDS epidemic. The domestic component consists of four research hubs, and the international component consists of collaborators from blood centers in Brazil, China, and South Africa.

Building on the findings of previous REDS and REDS II programs, the REDS-III international program focuses on identifying ways to reduce and prevent the transmission of HIV/AIDS and other known and emerging infectious agents through transmission.

Obligations

Funding History:

Fiscal Year 2011—\$19,910,527

Total Funding to Date—\$1,910,527

* Formerly known as Retrovirus Epidemiology Donor Study. REDS: Total funding for FY 1989–2004, \$73,774,125. REDS II: Total funding for FY 2005–2010, \$53,016,894.

Current Active Organizations and Contract Numbers

1. Blood System Research, Inc. San Francisco, California	—268201100001	6. Yale University New Haven, Connecticut	—268201100006
2. Research Triangle Institute, Inc. Research Triangle Park, North Carolina	—268201100002	7. Blood System Research, Inc. (Brazil) San Francisco, California	—268201100007
3. Blood Center of Southeastern Wisconsin Milwaukee, Wisconsin	—268201100003	8. Johns Hopkins University (China) Baltimore, Maryland	—268201100008
4. Institute for Transfusion Medicine Pittsburgh, Pennsylvania	—268201100004	9. University of California, San Francisco (South Africa) San Francisco, California	—268201100009
5. University of California, San Francisco San Francisco, California	—268201100005		



11. Clinical Trials

A clinical trial is defined as a scientific research study undertaken with human subjects to evaluate prospectively the diagnostic, prophylactic, or therapeutic effect of a drug, device, regimen, or procedure used or intended ultimately for use in the practice of

medicine or the prevention of disease. A clinical trial is planned and conducted prospectively and includes a concurrent control group or other appropriate comparison group.

NHLBI Investigator-Initiated Clinical Trials: Fiscal Years 2001–2011

Research Grants and Cooperative Agreements (Dollars in Thousands)

	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Heart and Vascular Diseases											
Infant Heart Surgery: Central Nervous System Sequelae of Circulatory Arrest	\$ 75	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —
Women's Health Study (WHS)	—	—	—	—	889	—	868	875	919	927	—
Women's Antioxidant and Cardiovascular Study (WACS)	572	598	592	599	670	—	—	—	—	—	—
Stress Reduction and Atherosclerotic CVD in Blacks	360	376	394	—	—	—	—	—	—	—	—
Shock Trial: Should We Emergently Revascularize Occluded Coronaries for Cardiogenic Shock?	362	298	291	296	—	—	—	—	—	—	—
Women's Estrogen/Progestin Lipid Lowering Hormone Atherosclerosis Regression Trial (WELL-HART)*	32	—	—	—	—	—	—	—	—	—	—
Mode Selection Trial in Sinus Node Dysfunction (MOST)*	154	—	—	—	—	—	—	—	—	—	—
Estrogen and Graft Atherosclerosis Research Trial (EAGER)*	371	—	—	—	—	—	—	—	—	—	—
REMATCH Trial*	750	—	—	—	—	—	—	—	—	—	—
Dietary Patterns, Sodium Intake, and Blood Pressure (DASH Sodium)* **	151	387	376	395	—	—	—	—	—	—	—
Sudden Cardiac Death in Heart Failure Trial (SCD-HeFT)*	1,798	1,412	1,930	—	—	—	—	—	—	—	—
CVD Risk and Health in Post-Menopausal Phytoestrogen Users	—	304	152	—	—	—	—	—	—	—	—
Treatment of Hypertension With Two Exercise Intensities	420	—	—	—	—	—	—	—	—	—	—
Prevention of Recurrent Venous Thromboembolism (PREVENT)	543	1,272	—	—	—	—	—	—	—	—	—
PREMIER: Lifestyle Interventions for Blood Pressure Control*	2,925	1,505	—	—	—	—	—	—	—	—	—
Azithromycin and Coronary Events Study (ACES)*	720	1,254	1,137	—	—	—	—	—	—	—	—
Antiarrhythmic Effects of N-3 Fatty Acids	529	647	—	—	—	—	—	—	—	—	—
Occluded Artery Trial (OAT)*	2,604	1,724	1,963	—	—	963	1,452	1,277	1,270	1,033	—
Bypass Angioplasty Revascularization Investigation in Type 2 Diabetics (BARI 2D)*	6,515	9,342	8,189	8,265	8,304	8,592	2,647	1,971	1,130	—	—

* Paid by U01/U10.

** Previously an Institute-Initiated Clinical Trial.

NHLBI Investigator-Initiated Clinical Trials: Fiscal Years 2001–2011 (continued)

Research Grants and Cooperative Agreements (Dollars in Thousands)

	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Heart and Vascular Diseases (continued)											
Hematocrit Strategy in Infant Heart Surgery*	557	596	590	492	—	—	—	—	—	—	—
Angiotensin-II Blockade in Mitral Regurgitation	553	610	629	500	—	—	—	—	—	—	—
Heart Failure Adherence and Retention Trial (HART)	795	1,617	1,453	1,174	862	740	304	—	—	—	—
Reduction of Triglycerides in Women on HRT	708	746	555	544	721	—	625	501	—	—	—
Women's Ischemia Syndrome Evaluation (WISE)* **	1,502	1,506	1,306	1,303	996	—	—	—	—	—	—
ACE Inhibition and Novel Cardiovascular Risk Factors	—	694	656	602	—	—	—	—	—	—	—
Heart Failure: A Controlled Trial Investigating Outcomes of Exercise (HF-ACTION)*	—	7,471	9,582	7,973	4,483	4,590	2,846	652	—	—	—
Clinical Trial of Dietary Protein on Blood Pressure	—	655	610	612	504	500	—	—	—	—	—
Home Automatic External Defibrillator Trial (HAT)*	—	3,567	5,433	4,264	1,801	2,115	—	—	—	—	—
Perioperative Interventional Neuroprotection Trial (POINT)	—	553	553	562	572	378	—	—	—	—	—
Stop Atherosclerosis in Native Diabetics Study (SANDS)*	—	2,410	2,165	2,107	2,324	2,074	197	218	—	—	—
Surgical Treatment for Ischemic Heart Failure (STICH)*	—	5,709	6,542	1,613	6,082	5,583	9,396	3,639	727	1,233	352
Girls Health Enrichment Multisite Studies (GEMS)*	—	—	2,461	2,400	2,369	1,950	—	—	—	—	—
Treatment of Depression Following Bypass Surgery	—	—	964	1,132	1,181	1,193	885	—	—	—	—
Weight Loss Maintenance (WLM)*	—	—	3,687	4,368	3,099	4,015	2,151	145	150	—	—
Cardiovascular Outcomes in Renal Atherosclerotic Lesions (CORAL)*	—	—	—	4,343	5,610	4,884	3,307	3,269	—	4,414	3,029
FREEDOM Trial: Future Revascularization Evaluation in Patients With Diabetes Mellitus: Optional Management of Multivessel Disease	—	—	—	3,663	4,669	—	5,180	2,818	1,658	3,429	2,755
IMMEDIATE Trial: Immediate Myocardial Metabolic Enhancement During Initial Assessment and Treatment in Emergency Care*	—	—	—	5,170	9,514	10,966	—	—	—	—	—
AIM HIGH: Niacin Plus Statin To Prevent Vascular Events*	—	—	—	—	663	6,324	6,018	1,380	2,324	6,927	1,089
Claudication: Exercise Versus Endoluminal Revascularization (CLEVER)*	—	—	—	—	1,368	1,478	1,898	—	1,822	1,080	1,371
Interventions To Control Obesity in College	—	—	—	—	—	677	633	670	686	588	—
PACEmaker and Beta-Blocker Therapy Post-Myocardial Infarction (PACE-MI)	—	—	—	—	—	1,300	4,555	384	—	—	—
Efficacy of Smoking Quit Line in the Military	—	—	—	—	—	—	739	720	731	—	723

* Paid by U01/U10.

** Previously an Institute-Initiated Clinical Trial.

NHLBI Investigator-Initiated Clinical Trials: Fiscal Years 2001–2011 (continued)

Research Grants and Cooperative Agreements (Dollars in Thousands)

	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Heart and Vascular Diseases (continued)											
Vest prevention of Early Sudden Death Trial (VEST) and PREDiction of ICD Therapies Studies (PREDICTS)*	—	—	—	—	—	—	—	1,390	1,356	1,391	1,404
Randomized Trial of Interventions To Improve Warfarin Adherence	—	—	—	—	—	—	—	801	787	771	763
Planned Care for Obesity and Risk Reduction (Planned CORR)	—	—	—	—	—	—	—	784	770	769	768
Effects of Niacin on Lp(a), Oxidized LDL, and Inflammation on the AIM-HIGH Trial	—	—	—	—	—	—	—	302	312	383	—
Women's Ischemia Syndrome Evaluation (WISE) Coronary Vascular Dysfunction	—	—	—	—	—	—	—	776	742	745	761
Rule-Out Myocardial Infarction Using Computer Assisted Tomography (ROMICAT II)*	—	—	—	—	—	—	—	—	2,112	2,307	367
Collaborative Model To Improve BP Control and Minimize Racial Disparities	—	—	—	—	—	—	—	—	1,963	1,938	1,852
Multiscale Model of the Human Heart for Imaging Research	—	—	—	—	—	—	—	—	566	503	498
Catheter Ablation Versus Antiarrhythmic Drug Therapy for Atrial Fibrillation (CABANA)*	—	—	—	—	—	—	—	—	2,941	3,045	3,033
Therapeutic Hypothermia After Pediatric Cardiac Arrest (THAPCA)*	—	—	—	—	—	—	—	—	3,648	2,285	4,402
Vitamin D and Omega 3 Trial (VITAL)*	—	—	—	—	—	—	—	—	1,260	1,248	1,210
Late Sodium Blockade in High-Risk ICD Patients*	—	—	—	—	—	—	—	—	—	2,280	—
Cardiovascular Inflammation Reduction Trial	—	—	—	—	—	—	—	—	—	—	1,376
Impact of Vitamin D Supplementation on Cardiometabolic Risk in School Children	—	—	—	—	—	—	—	—	—	—	552
Lifestyle, CVD Risk, and Cognitive Impairment	—	—	—	—	—	—	—	—	—	—	785
ISCHEMIA Trial (International Study of Comparative Effectiveness with Medical Invasive Approaches)	—	—	—	—	—	—	—	—	—	—	6,672
Subtotal, Heart and Vascular Diseases	22,996	45,253	52,210	52,377	56,681	58,312	45,091	22,538	27,909	37,309	32,358
Lung Diseases											
Lung Health Study III* **	1,672	927	—	—	—	—	—	—	—	—	—
Asthma Clinical Research Network (ACRN)* **	5,705	5,863	—	—	—	—	—	—	—	—	—
Fetal Tracheal Occlusion for Severe Diaphragmatic Hernia*	181	—	—	—	—	—	—	—	—	—	—
Scleroderma Lung Study*	1,761	1,501	1,055	—	—	71	—	—	—	—	—
Inhaled Nitric Oxide for Prevention of Chronic Lung Disease*	1,803	1,764	1,442	1,245	—	—	—	—	—	—	—
Inhaled Nitric Oxide in Prevention of Chronic Lung Disease*	1,839	1,604	903	—	—	—	—	—	—	—	—
Prospective Investigation of Pulmonary Embolism Diagnosis II (PIOPED II)*	3,388	472	—	—	—	—	—	—	—	—	—

* Paid by U01/U10.

** Previously an Institute-Initiated Clinical Trial.

NHLBI Investigator-Initiated Clinical Trials: Fiscal Years 2001–2011 (continued)

	Research Grants and Cooperative Agreements (Dollars in Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Lung Diseases (continued)											
Randomized Trial To Reduce ETS in Children With Asthma	468	277	—	—	—	—	—	—	—	—	—
Apnea Positive Pressure Long-Term Efficacy Study (APPLES)*	—	3,224	3,021	3,110	3,188	—	1,532	—	—	—	—
Childhood Asthma Management Program—Continuation Study (CAMP-CS)/Phase II***	—	—	1,489	2,043	2,623	2,750	—	—	—	—	—
Acid Reflux Therapy in Asthma*	—	—	736	783	791	773	662	—	—	—	—
Impact of CPAP on Functional Outcomes in Milder Obstructive Sleep Apnea (CATNAP)	—	—	682	612	608	694	—	—	—	—	—
Outcomes of Sleep Disorders in Older Men	—	—	4,163	4,262	1,390	1,142	910	—	—	—	—
Supplemental Selenium and Vitamin E and Pulmonary Function	—	—	698	610	630	605	561	—	—	—	—
Improving Asthma Care in Minority Children in Head Start	—	—	—	721	826	1,004	779	—	—	—	—
Randomized Controlled Study of Adenotonsillectomy for Childhood Sleep Apnea	—	—	—	—	—	2,255	2,388	1,346	2,501	1,675	—
Early Insulin Therapy and Development of ARDS	—	—	—	—	—	—	489	454	464	417	386
Childhood Asthma Management Program—Continuation Study (CAMP-CS)/Phase III**	—	—	—	—	—	—	2,077	1,966	1,146	2,065	—
Infant Study of Inhaled Saline in Cystic Fibrosis (ISIS)*	—	—	—	—	—	—	—	732	737	776	628
Scleroderma Lung Study II	—	—	—	—	—	—	—	2,281	2,297	2,252	2,268
Sedation Management in Pediatric Patients With Acute Respiratory Failure*	—	—	—	—	—	—	—	568	3,885	3,327	2,181
Study of Asthma and Nasal Steroids (STAN)*	—	—	—	—	—	—	—	—	725	725	725
Outpatient Treatment of Low-Risk Patients With Pulmonary Embolism	—	—	—	—	—	—	—	—	755	349	—
Translating COPD Guidelines Into Primary Care Practice	—	—	—	—	—	—	—	—	733	719	—
Family Intervention for Pediatric Asthma Self-Management in Puerto Ricans	—	—	—	—	—	—	—	—	225	187	—
Physical Activity Self-Management in Patients With COPD	—	—	—	—	—	—	—	—	663	660	655
Study of Soy Isoflavones in Asthma (SOYA)*	—	—	—	—	—	—	—	—	775	697	689
Trial of Late Surfactant To Prevent Bronchopulmonary Dysplasia (TOLSURF)*	—	—	—	—	—	—	—	—	1,987	1,779	1,807
Randomized Trial of Maternal Vitamin D Supplementation To Prevent Childhood Asthma*	—	—	—	—	—	—	—	—	2,510	2,466	2,469

* Paid by U01/U10.

** Previously an Institute-Initiated Clinical Trial.

NHLBI Investigator-Initiated Clinical Trials: Fiscal Years 2001–2011 (continued)

Research Grants and Cooperative Agreements (Dollars in Thousands)

	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Lung Diseases (continued)											
Effects of HIV Antiretroviral Therapy on Pulmonary Function	—	—	—	—	—	—	—	—	614	516	552
Randomized Trial of Antenatal Late Preterm Steroids (ALPS)*	—	—	—	—	—	—	—	—	—	2,134	2,137
Vitamin D Supplementation in Pregnancy: Impact on Neonatal Immune Phenotype	—	—	—	—	—	—	—	—	—	323	311
Role of Beta-Catenin in Epithelial Repair in Acute Lung Injury	—	—	—	—	—	—	—	—	—	125	125
Mechanisms of Familial Pulmonary Fibrosis	—	—	—	—	—	—	—	—	—	2,330	2,252
Multicomponent Intervention To Decrease COPD-Related Hospitalizations	—	—	—	—	—	—	—	—	—	412	470
Heart and Lung Failure-Pediatric Insulin Titration (HALF-PINT)	—	—	—	—	—	—	—	—	—	—	2,685
Subtotal, Lung Diseases	17,076	18,974	15,639	14,289	10,056	9,294	9,398	7,347	20,017	23,934	20,340
Blood Diseases and Resources											
Stroke Prevention in Sickle Cell Anemia (STOP 2)*	4,200	3,166	3,168	2,366	—	—	—	—	—	—	—
Induction of Stable Chimerism for Sickle Cell Anemia*	489	525	527	551	—	—	—	—	—	—	—
Sibling Donor Cord Blood Banking and Transplantation*	1,222	1,224	1,286	1,353	—	—	—	—	—	—	—
FOCUS*	—	—	1,639	1,796	2,923	2,446	1,974	—	—	—	—
Stroke With Transfusions Changing to Hydroxyurea (SWiTCH)*	—	—	—	—	3,345	3,932	3,531	3,828	3,216	1,778	—
Bridging Anticoagulation on Patients Requiring Temporary Interruption of Warfarin Therapy for an Elective Procedure or Surgery (BRIDGE) Trial*	—	—	—	—	—	—	—	4,632	5,673	5,227	—
Acute Venous Thrombosis: Thrombus Removal With Adjunctive Catheter-Directed Thrombolysis (ATTRACT)* **	—	—	—	—	—	—	—	2,071	2,108	2,094	244
Transcranial Doppler With Transfusions Changing to Hydroxyurea (TWiTCH)	—	—	—	—	—	—	—	—	4,176	4,177	4,391
Impact of Blood Storage Duration on Physiologic Measures: RECESS Ancillary Study	—	—	—	—	—	—	—	—	—	—	389
Subtotal, Blood Diseases and Resources	4,877	4,917	5,772	6,066	6,268	6,378	5,505	10,531	15,173	13,663	5,024
Total, NHLBI	\$44,949	\$69,144	\$73,621	\$72,732	\$73,005	\$73,984	\$59,994	\$40,416	\$63,099	\$74,906	\$57,722

* Paid by U01/U10.

** Formerly known as Pharmacomechanical Catheter-Directed Thrombolysis for Acute DVT—ATTRACT Trial.

NHLBI Investigator-Initiated Clinical Trials, Fiscal Year 2011: Summary by Program

	Total Obligations Prior to 2011	FY 2011 Obligations	Total Obligation to Date
Heart and Vascular Diseases			
AIM HIGH: Niacin Plus Statin To Prevent Vascular Events*	\$23,637,416	\$1,088,603	\$24,726,019
Cardiovascular Inflammation Reduction Trial*	—	1,375,726	1,375,726
Cardiovascular Outcomes in Renal Atherosclerotic Lesions (CORAL)*	25,827,224	3,029,201	28,856,425
Catheter Ablation Versus Antiarrhythmic Drug Therapy for Atrial Fibrillation Trial (CABANA)*	5,985,643	3,032,875	9,018,518
Claudication: Exercise Versus Endoluminal Revascularization (CLEVER)*	7,647,416	1,371,307	9,018,723
Collaborative Model To Improve BP Control and Minimize Racial Disparities	3,900,897	1,852,389	5,753,286
Efficacy of Smoking Quit Line in the Military	2,189,732	723,167	2,912,899
FREEDOM Trial: Future Revascularization Evaluation in Patients With Diabetes Mellitus: Optimal Management of Multivessel Disease	21,417,459	2,754,827	24,172,286
Impact of Vitamin D Supplementation on Cardiometabolic Risk in School Children	—	552,019	552,019
Interventions To Improve Warfarin Adherence	2,358,416	763,252	3,121,668
ISCHEMIA Trial (International Study of Comparative Effectiveness with Medical Invasive Approaches)	—	6,671,629	6,671,629
Lifestyle, CVD Risk, and Cognitive Impairment	—	784,998	784,998
Multiscale Model of the Human Heart for Imaging Research	1,068,817	498,049	1,566,866
Planned Care for Obesity and Risk Reduction (Planned CORR)	2,323,037	767,849	3,090,886
Rule-Out Myocardial Infarction Using Computer Assisted Tomography (ROMICAT II)*	4,419,188	366,525	4,785,713
Surgical Treatment for Ischemic Heart Failure (STICH)*	40,041,062	352,198	40,393,260
Therapeutic Hypothermia After Pediatric Cardiac Arrest (THAPCA)*	5,933,723	4,401,772	10,335,495
Vitamin D and Omega 3 Trial (VITAL)*	2,507,778	1,210,345	3,718,123
Women's Ischemia Syndrome Evaluation (WISE) Coronary Vascular Dysfunction	2,262,918	760,707	3,023,625
Subtotal, Heart and Vascular Diseases	151,520,726	32,357,438	183,878,164
Lung Diseases			
Early Insulin Therapy and Development of ARDS	1,824,699	386,117	2,210,816
Effects of HIV Antiretroviral Therapy on Pulmonary Function	1,129,948	552,353	1,682,301
Heart and Lung Failure-Pediatric Insulin Trial (HALF-PINT)	—	2,685,460	2,685,460
Infant Study of Inhaled Saline in Cystic Fibrosis (ISIS)*	2,244,874	628,241	2,873,115
Mechanisms of Familial Pulmonary Fibrosis	2,330,347	2,251,871	4,582,218
Multicomponent Intervention To Decrease COPD-Related Hospitalizations	411,777	469,910	881,687
Physical Activity Self-Management in Patients With COPD	1,323,114	654,594	1,977,708
Randomized Trial of Antenatal Late Preterm Steroids (ALPS)*	2,133,947	2,137,414	4,271,361
Randomized Trial of Maternal Vitamin D Supplementation To Prevent Childhood Asthma*	4,975,916	2,469,339	7,445,255
Role of Beta-Catenin in Epithelial Repair in Acute Lung Injury	125,172	125,172	250,344

* Paid by U01/U10.

** Previously an Institute-Initiated Clinical Trial.

NHLBI Investigator-Initiated Clinical Trials, Fiscal Year 2011: Summary by Program (continued)

	Total Obligations Prior to 2011	FY 2011 Obligations	Total Obligation to Date
Lung Diseases (continued)			
Scleroderma Lung Study II	6,828,804	2,267,789	9,096,593
Sedation Management in Pediatric Patients With Acute Respiratory Failure*	7,779,116	2,181,199	9,960,315
Study of Asthma and Nasal Steroids (STAN)*	1,449,536	724,884	2,174,420
Study of Soy Isoflavones in Asthma (SOYA)*	1,471,854	688,673	2,160,527
Trial of Late Surfactant To Prevent Bronchopulmonary Dysplasia (TOLSURF)*	3,765,851	1,807,148	5,572,999
Vitamin D Supplementation in Pregnancy: Impact on Neonatal Immune Phenotype	323,300	311,250	634,550
Subtotal, Lung Diseases	38,118,255	20,341,414	58,459,669
Blood Diseases and Resources			
Acute Venous Thrombosis: Thrombus Removal with Adjunctive Catheter-Directed Thrombolysis (ATTRACT)* **	6,272,537	243,775	6,516,312
Impact of Blood Storage Duration on Physiologic Measures: RECESS Ancillary Study	386,523	389,405	775,928
Transcranial Doppler With Transfusions Changing to Hydroxyurea (TWiTCH)	8,353,426	4,390,891	12,744,317
Subtotal, Blood Diseases and Resources	15,012,486	5,024,071	20,036,557
TOTAL, NHLBI	\$204,651,467	\$57,722,923	\$262,374,390

* Paid by U01/U10.

** Formerly known as Pharmacomechanical Catheter-Directed Thrombolysis for Acute DVT—ATTRACT Trial.

Institute-Initiated Clinical Trials: Fiscal Years 2001–2011

Contracts

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Heart and Vascular Diseases											
Antihypertensive and Lipid-Lowering Treatment To Prevent Heart Attack Trial (ALLHAT)	7,000	3,980	2,761	3,346	—	—	—	—	—	1,235	971
Enhancing Recovery in Coronary Heart Disease Patients (ENRICHD)	596	425	70	—	—	—	—	—	—	—	—
Atrial Fibrillation Follow-Up: Investigation in Rhythm Management (AFFIRM)	2,401	802	—	—	—	—	—	—	—	—	—
Women's Angiographic Vitamin and Estrogen Trial (WAVE)	756	—	32	—	—	—	—	—	—	—	—
Women's Ischemia Syndrome Evaluation (WISE)	10	50	—	—	—	—	—	—	—	—	—
Prevention of Events With Angiotensin Converting Enzyme Inhibitor Therapy (PEACE)	—	2,849	558	—	—	—	—	—	—	—	—
Magnesium in Coronaries (MAGIC)	—	238	—	—	—	—	—	—	—	—	—
Evaluation Study of Congestive Heart Failure and Pulmonary Artery Catheterization Effectiveness (ESCAPE)	—	1,129	—	—	—	311	—	—	—	—	—
Action To Control Cardiovascular Risk in Diabetes (ACCORD)	—	1,750	18,521	33,779	26,126	—	19,484	16,343	15,461	403	—
Public Access Defibrillation (PAD) Community Trial	3,058	1,101	—	—	—	—	—	—	—	—	—
Trial of Aldosterone Antagonist Therapy in Adults With Preserved Ejection Fraction Congestive Heart Failure (TOPCAT)	—	—	—	837	5,162	5,480	2,218	7,912	4,408	898	—
Women's Health Initiative	59,200	59,010	63,222	57,483	37,826	12,124	14,873	22,609	30,615	2,409	22,766
Randomized Evaluation of VAD InterVENTion before Inotropic Therapy (REVIVE-IT)	—	—	—	—	—	—	—	—	—	—	4,953
Systolic Blood Pressure Intervention Trial (SPRINT)	—	—	—	—	—	—	—	—	3,057	29,374	—
Clarification of Optimal Anticoagulation Through Genetics (COAG)*	—	—	—	—	—	—	—	2,637	3,530	—	6,760
Subtotal, Heart and Vascular Diseases	73,021	71,334	85,164	95,445	69,114	17,915	36,575	49,501	57,071	34,319	35,450
Lung Diseases											
Pediatric Pulmonary and Cardiac Complications of HIV Infection (P2C2)	—	113	—	—	—	—	—	—	—	—	—
Childhood Asthma Management Program (CAMP)	1,330	2,786	2,287	1,475	599	—	—	—	—	—	—
Acute Respiratory Distress Syndrome Clinical Network (ARDSNet)	2,667	1,502	4,402	5,517	4,707	7,396	5,037	1,992	6,195	7,208	5,096

* Formerly known as Randomized Trial of Genotype-Guided Dosing of Warfarin Therapy.

Institute-Initiated Clinical Trials: Fiscal Years 2001–2011 (continued)

Contracts (continued)

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Lung Diseases (continued)											
National Emphysema Treatment Trial (NETT)	6,989	7,910	1,630	1,648	357	—	—	—	285	—	—
Feasibility of Retinoid Treatment in Emphysema (FORTE)	—	2,429	725	507	185	—	—	—	—	—	—
Long-Term Oxygen Treatment Trial (LOTT)	—	—	—	—	—	—	6,208	10,042	202	4,335	4,378
Subtotal, Lung Diseases	10,986	14,740	9,044	9,147	5,848	7,396	11,245	12,034	6,682	11,543	9,474
Blood Diseases and Resources											
T-Cell Depletion in Unrelated Donor Marrow Transplantation	1,144	557	774	164	—	—	—	—	—	—	—
Cord Blood Stem Cell Transplantation Study (COBLT)	1,846	2,166	588	707	822	—	—	—	—	—	—
Multicenter Study of Hydroxyurea (MSH) in Sickle Cell Anemia Adult Follow-Up	—	588	994	1,136	1,340	—	—	—	—	—	—
Pediatric Hydroxyurea Phase III Clinical Trial (BABY HUG)	405	3,100	1,112	1,964	1,526	891	3,966	5,573	1,704	—	853
Sildenafil for Sickle Cell Disease-Associated Pulmonary Hypertension (walk PHaSST)	—	—	—	—	—	1,867	2,801	3,702	963	320	212
Subtotal, Blood Diseases and Resources	3,395	6,411	3,468	3,971	3,688	2,758	6,767	9,275	2,667	320	1,065
Total, NHLBI Clinical Trials Contracts	\$87,402	\$92,485	\$97,676	\$108,563	\$78,650	\$28,069	\$54,587	\$70,810	\$66,420	\$46,182	\$45,989

Institute-Initiated Clinical Trials: Fiscal Years 2001–2011 (continued)

Cooperative Agreements

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Heart and Vascular Diseases											
Bypass Angioplasty Revascularization Investigation (BARI)	\$ 1,549	\$ 1,456	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —
Girls Health Enrichment Multisite Studies (GEMS)	2,877	2,713	—	—	—	—	—	—	—	—	—
Trial of Activity for Adolescent Girls (TAAG)	4,831	5,919	5,828	6,350	5,103	905	—	—	—	—	—
Pediatric Heart Network	3,447	4,822	5,381	4,948	3,992	6,988	6,607	12,255	7,637	7,471	12,827
Clinical Research Consortium To Improve Resuscitation Outcome	—	—	—	6,886	9,339	9,728	8,972	5,279	—	6,244	9,455
Dynamic Assessment of Patient-Reported Chronic Disease Outcomes	—	—	—	1,010	—	—	—	—	—	—	—
Heart Failure Clinical Research Network	—	—	—	—	—	5,642	7,801	7,813	7,939	7,914*	7,652
Pediatric HIV/AIDS Cohort Study (PHACS)—Data and Operations Center	—	—	—	—	—	1,000	500	490	500	769*	600
Cardiovascular Cell Therapy Research Network	—	—	—	—	—	—	4,424	7,568	6,227	6,200	3,800
Community-Responsive Interventions To Reduce Cardiovascular Risk in American Indians and Alaska Natives	—	—	—	—	—	1,419	2,314	3,151	1,999	2,071	—
Network for Cardiothoracic Surgical Investigations in Cardiovascular Medicine	—	—	—	—	—	—	6,009	8,681	3,210	8,079	5,460
EDTA Chelation Therapy for Coronary Artery Disease	—	—	—	—	—	—	—	—	2,109	—	—
Practice-Based Opportunity for Weight Reduction (POWER) Trials*	—	—	—	—	—	2,567	3,714	3,656	3,729	2,329	—
Look AHEAD: Action for Health in Diabetes	—	—	—	—	—	—	—	—	—	4,000	4,000
Diabetes Prevention Program Outcomes Study—Phase II	—	—	—	—	—	—	—	—	—	1,100	1,100
Early Adult Reduction of Weight Through LifestYle Intervention (EARLY) Trials**	—	—	—	—	—	—	—	—	4,656	5,864	6,124

* Formerly known as Weight Loss in Obese Adults With Cardiovascular Risk Factors.

**Formerly known as Targeted Approaches to Weight Control for Young Adults.

Institute-Initiated Clinical Trials: Fiscal Years 2001–2011 (continued)

Cooperative Agreements (continued)

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Heart and Vascular Diseases (continued)											
Childhood Obesity Prevention and Treatment Research (COPTR)	—	—	—	—	—	—	—	—	—	4,058	3,986
Consortium of Hospitals Advancing Research on Tobacco (CHART)*	—	—	—	—	—	—	—	—	—	3,322	3,505
Subtotal, Heart and Vascular Diseases	12,704	14,910	11,209	19,194	18,434	28,249	40,341	48,893	43,106	59,421**	58,509
Lung Diseases											
Asthma Clinical Research Network (ACRN)†	—	—	8,181	8,424	8,667	7,839	8,918	872	—	—	—
Childhood Asthma Research and Education (CARE) Network	5,314	6,005	5,610	5,292	5,704	5,735	5,916	4,887	—	—	—
COPD Clinical Research Network	—	—	6,843	6,848	8,438	7,664	6,836	3,400	3,150	3,150	2,600
Idiopathic Pulmonary Fibrosis Clinical Research Network	—	—	—	—	3,486	7,349	7,216	7,154	7,325	—	—
NICHD Cooperative Multicenter Neonatal Research Network	—	—	—	—	—	1,336	238	27	—	—	—
Asthma Network (AsthmaNet)	—	—	—	—	—	—	—	—	8,300	15,500	15,500
Novel Therapies for Lung Diseases—Phase II	—	—	—	—	—	—	—	—	—	7,594	12,843
Subtotal, Lung Diseases	5,314	6,005	20,634	20,564	26,295	29,923	29,124	16,340	18,775	26,244	30,943
Blood Diseases and Resources											
Thalassemia (Cooley's Anemia) Clinical Research Network	2,219	2,269	2,320	2,375	2,730	2,682	2,618	2,600	658	—	—
Blood and Marrow Transplant Clinical Research Network	5,360	5,899	5,950	5,972	6,460	6,845	6,709	6,952	6,351	2,507	5,319
Transfusion Medicine/Hemostasis Clinical Research Network	—	6,053	6,241	6,093	6,221	6,521	6,407	6,374	6,541	6,590	6,314
Sickle Cell Disease Clinical Research Network	—	—	—	—	—	3,761	7,498	7,173	—	—	—
Subtotal, Blood Diseases and Resources	7,579	14,221	14,511	14,440	15,411	19,809	23,232	23,099	13,550	9,097	11,453
Total, NHLBI-Initiated Clinical Trials, Cooperative Agreements	\$25,597	\$35,136	\$46,354	\$54,198	\$60,140	\$77,981	\$92,697	\$88,332	\$75,431	\$94,762**	\$100,905
Total, NHLBI-Initiated Clinical Trials	\$112,999	\$127,621	\$144,030	\$162,761	\$138,790	\$106,050	\$147,284	\$159,142	\$141,851	\$140,944**	\$146,594

* Formerly known as Effective Research on Smoking Cessation in Hospitalized Patients.

** Correction to figure that was reported in the FY 2010 Fact Book.

† Investigator-Initiated from 1998 to 2002.

Note: The line labeled "Other Clinical Trials" (\$78.8 million) that appeared in the FY 2010 Fact Book has been removed.

Institute-Initiated Clinical Trials, Fiscal Year 2011: Summary by Program

Contracts

	Total Obligations Prior to FY 2011	Total FY 2011 Obligations	Total Obligations to Date
Heart and Vascular Diseases			
Antihypertensive and Lipid-Lowering Treatment To Prevent Heart Attack Trial (ALLHAT)	\$ 84,405,430	\$ 971,242	\$ 85,376,672
Clarification of Optimal Anticoagulation Through Genetics (COAG)*	6,167,062	6,760,320	12,927,382
Randomized Evaluation of VAD InterVENTion before Inotropic Therapy (REVIVE-IT)	—	4,952,781	4,952,781
Systolic Blood Pressure Intervention Trial (SPRINT)	36,882,407	—	36,882,407
Trial of Aldosterone Antagonists Therapy in Adults With Preserved Ejection Fraction Congestive Heart Failure (TOPCAT)	26,914,315	—	26,914,315
Women's Health Initiative (WHI)**	793,070,018	22,766,227	815,836,245
Subtotal, Heart and Vascular Diseases	947,439,232	35,450,570	982,889,802
Lung Diseases			
Acute Respiratory Distress Syndrome Clinical Network (ARDSNet)	78,742,505	5,096,234	83,838,739
Long-Term Oxygen Treatment Trial (LOTT)	20,787,361	4,377,865	25,165,226
Subtotal, Lung Diseases	99,529,866	9,474,099	109,003,965
Blood Diseases and Resources			
Pediatric Hydroxyurea Phase III Clinical Trial (BABY HUG)	21,847,533	852,640	22,700,173
Sildenafil for Sickle Cell Disease-Associated Pulmonary Hypertension (walk PHaSST)	9,653,177	212,144	9,865,321
Subtotal, Blood Diseases and Resources	31,500,710	1,064,784	32,565,494
Total, NHLBI-Initiated Clinical Trials, Contracts	\$1,078,469,808	\$45,989,453	\$1,124,459,261

* Formerly known as Randomized Trial of Genotype-Guided Dosing of Warfarin Therapy.

** From 1999 to 2006, the WHI was reported separately under its own major heading. In this table, it is included in the Heart and Vascular Diseases section.

Cooperative Agreements

	Total Obligations Prior to FY 2011	Total FY 2011 Obligations	Total Obligations to Date
Heart and Vascular Diseases			
Cardiovascular Cell Therapy Research Network	\$ 24,419,327	\$ 3,800,000	\$ 28,219,327
Childhood Obesity Prevention and Treatment Research (COPTR)	4,058,435	3,986,427	8,044,862
Clinical Research Consortium To Improve Resuscitation Outcome	46,447,452	9,455,378	55,902,830
Consortium of Hospitals Advancing Research (CHART)*	3,321,649	3,505,096	6,826,745
Diabetes Prevention Program Outcomes Study—Phase II	2,200,000	1,100,000	3,300,000
Early Adult Reduction of Weight Through LifestYLE Intervention (EARLY) Trials**	10,519,703	6,123,513	16,643,216
Heart Failure Clinical Research Network	37,108,888	7,651,786	44,760,674
Look AHEAD: Action for Health in Diabetes	8,000,000	4,000,000	12,000,000
Network for Cardiothoracic Surgical Investigations in Cardiovascular Medicine	25,978,806	5,459,597	31,438,403
Pediatric Heart Network	63,348,873	12,827,177	76,176,050
Pediatric HIV/AIDS Cohort Study (PHACS): Data and Operations Center	3,259,021	600,000	3,859,021
Subtotal, Heart and Vascular Diseases	228,662,154	58,508,974	287,171,128
Lung Diseases			
Asthma Network (AsthmaNet)	23,800,000	15,500,000	39,300,000
COPD Clinical Research Network	46,330,386	2,600,000	48,930,386
Novel Therapies for Lung Diseases—Phase II	7,593,739	12,843,238	20,436,977
Subtotal, Lung Diseases	77,724,125	30,943,238	108,667,363
Blood Diseases and Resources			
Blood and Marrow Transplant Clinical Research Network	59,071,021	5,319,784	64,390,805
Transfusion Medicine/Hemostasis Clinical Research Network	57,040,060	6,133,712	63,173,772
Subtotal, Blood Diseases and Resources	116,111,081	11,453,496	127,564,577
Total, NHLBI-Initiated Clinical Trials, Cooperative Agreements	\$422,497,360	\$100,905,708	\$523,403,068
Total, NHLBI-Initiated Clinical Trials	\$1,500,967,868	\$146,595,161	\$1,647,863,029

* Formerly known as Effectiveness Research on Smoking Cessation in Hospitalized Patients.

** Formerly known as Targeted Approaches to Weight Control for Young Adults.

Heart and Vascular Diseases

Antihypertensive and Lipid-Lowering Treatment To Prevent Heart Attack Trial (ALLHAT), Initiated in Fiscal Year 1993

The purpose of this study was to compare the ability of a diuretic versus newer antihypertensive treatments (ACE inhibitor, calcium channel blocker, alpha blocker) to lower the combined incidence of fatal CHD and non-fatal MI in high-risk hypertensive patients and to determine whether lowering serum cholesterol with an HMG CoA reductase inhibitor reduced the total mortality in a subset of hypertensive patients with moderately elevated LDL cholesterol.

In February 2000, the alpha blocker arm of the study was discontinued because the CVD event rate was significantly greater among those patients compared with those in the control group. In 2002, results showed that diuretics work best to both lower blood pressure and prevent stroke and some forms of heart disease, including heart attack and heart failure.

Researchers are analyzing a post-trial follow-up (9–10 years) of participants to compare long-term effects of antihypertensive treatment with a thiazide-type diuretic, a calcium channel blocker, an ACE inhibitor, and an alpha receptor blocker when each drug was used as initial treatment, with step-up drugs added as needed, and for the lipid component, to assess long-term effects of pravastatin compared with usual care. Fifty-five percent of the participants are black.

Obligations

Funding History:

Fiscal Year 2011—\$971,242
Fiscal Years 1993–2010—\$84,405,430
Total Funding to Date—\$85,376,672

Current Active Organization and Contract Number

1. University of Texas Health Science Center Houston, Texas —26820110036C

Cardiovascular Cell Therapy Research Network, Initiated in Fiscal Year 2007

The purpose of this program is to establish a research network to evaluate innovative cell therapy strategies for individuals with CVD. The network is providing the necessary infrastructure to develop, coordinate, and

conduct multiple collaborative clinical protocols to facilitate application of emerging scientific discoveries to improve CVD outcomes.

Obligations

Funding History:

Fiscal Year 2011—\$3,800,000
Fiscal Years 2007–2010—\$24,419,327
Total Funding to Date—\$28,219,327

Current Active Organizations and Grant Numbers

1. Case Western Reserve University Cleveland, Ohio —HL-087314
2. University of Texas Health Science Center Houston, Texas —HL-087318
3. Texas Heart Institute Houston, Texas —HL-087365
4. University of Florida Gainesville, Florida —HL-087366
5. University of Minnesota, Twin Cities Minneapolis, Minnesota —HL-087394
6. Vanderbilt University Nashville, Tennessee —HL-087403

Childhood Obesity Prevention and Treatment Research (COPTR), Initiated in Fiscal Year 2010

The purpose of this research consortium is to test interventions to prevent excess weight gain in children and to reduce weight in obese children. Two obesity prevention trials are developing and testing approaches that target home, community, and primary care settings for preschool children living in low-income and ethnically diverse neighborhoods. Two obesity treatment trials are examining the therapies on overweight and obese children, 7- to 15-year olds, in school and home settings in collaboration with local youth organizations. More than 50 percent of the participants are expected to be from racial and ethnic minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$3,986,427
Fiscal Year 2010—\$4,058,435
Total Funding to Date—\$8,044,862

Current Active Organizations and Grant Numbers

1. University of North Carolina at Chapel Hill Chapel Hill, North Carolina —HL-103561

2. Vanderbilt University School of Medicine Nashville, Tennessee	—HL-103620
3. Case Western Reserve University Cleveland, Ohio	—HL-103622
4. Stanford University Palo Alto, California	—HL-103629

Clarification of Optimal Anticoagulation Through Genetics (COAG),* Initiated in Fiscal Year 2008

The purpose of this randomized, multicenter clinical trial is to compare two approaches to the initiation of warfarin therapy for optimal anticoagulation. One approach uses clinical information and an individual's genotype based on the genes known to influence warfarin response ("genotype-guided dosing"), and the other uses only clinical information ("clinical-guided dosing"). Approximately 1,200 participants are being randomized over a 3-year period at 18 medical centers throughout the United States. The primary endpoint is anticoagulation control during the first month of therapy. The trial is assessing anticoagulation control, bleeding problems and other complications, quality of life, and cost of therapy up to 6 months after initiation of therapy.

Obligations

Funding History:

Fiscal Year 2011—\$6,760,320
Fiscal Years 2008–2010—\$6,167,062
Total Funding to Date—\$12,927,382

Current Active Organization and Contract Number

1. University of Pennsylvania Philadelphia, Pennsylvania	—26800800003C
---	---------------

Clinical Research Consortium To Improve Resuscitation Outcomes, Initiated in Fiscal Year 2004

The purpose of this program is to conduct research in cardiopulmonary arrest and severe traumatic injury to facilitate the rapid translation of promising scientific and clinical advances to improve resuscitation outcomes. The Consortium conducts multiple, collaborative clinical trials and studies that target primarily out-of-hospital clinical life compromising events, such as cardiac arrest and life threatening trauma.

Obligations

Funding History:

Fiscal Year 2011—\$9,455,378
Fiscal Years 2004–2010—\$46,447,452
Total Funding to Date—\$55,902,830

Current Active Organizations and Grant Numbers

1. University of Washington Seattle, Washington	—HL-077863
2. Medical College of Wisconsin Milwaukee, Wisconsin	—HL-077866
3. University of Washington Seattle, Washington	—HL-077867
4. University of Pittsburgh Pittsburgh, Pennsylvania	—HL-077871
5. St. Michael's Hospital Toronto, Ontario	—HL-077872
6. Oregon Health and Science University Portland, Oregon	—HL-077873
7. University of Alabama at Birmingham Birmingham, Alabama	—HL-077881
8. Ottawa Health Research Institute Ottawa, Ontario	—HL-077885
9. University of Texas Southwestern Medical Center Dallas, Texas	—HL-077887
10. University of California, San Diego La Jolla, California	—HL-077908

Consortium of Hospitals Advancing Research on Tobacco (CHART), Initiated in Fiscal Year 2010**

The purpose of this study is to evaluate the effectiveness of smoking cessation interventions in hospitalized patients. A network of six projects are assessing the effectiveness and cost effectiveness of smoking cessation interventions that are initiated during hospitalization and continued post-discharge in more than 10,000 hospitalized smokers across a geographically diverse group of nearly 20 private, public, academic, and community hospitals. Participation from minority populations is strong in some of the projects.

Obligations

Funding History:

Fiscal Year 2011—\$3,505,096
Fiscal Year 2010—\$3,321,649
Total Funding to Date—\$6,826,745

* Formerly known as Randomized Trial of Genotype-Guided Dosing of Warfarin Therapy.

**Formerly known as Effective Research on Smoking Cessation in Hospitalized Patients.

Current Active Organizations and Grant Numbers

1. University of Michigan at Ann Arbor Ann Arbor, Michigan	—HL-105218	6. St. Luke's Roosevelt Institute for Health Sciences New York, New York	—DK-048404
2. New York University School of Medicine New York, New York	—HL-105229	7. Indiana University-Purdue University at Indianapolis Indianapolis, Indiana	—DK-048406
3. Kaiser Foundation Research Institute Oakland, California	—HL-105231	8. University of New Mexico Albuquerque, New Mexico	—DK-048407
4. University of Kansas Medical Center Kansas City, Kansas	—HL-105232	9. University of Tennessee Health Science Center Memphis, Tennessee	—DK-048411
5. Kaiser Foundation Research Institute Oakland, California	—HL-105233	10. Seattle Institute for Biomedical and Clinical Research Seattle, Washington	—DK-048413
6. University of California, San Diego San Diego, California	—CA-159533	11. University of California, Los Angeles Los Angeles, California	—DK-048443
7. University of Alabama at Birmingham Birmingham, Alabama	—DA-031515	12. Thomas Jefferson University Philadelphia, Pennsylvania	—DK-048468

Diabetes Prevention Program Outcomes Study—Phase II, Initiated in Fiscal Year 2009

The purpose of this multicenter clinical trial is to determine the efficacy of treatments to prevent or delay the development of type 2 diabetes in a population at high risk due to the presence of impaired glucose tolerance. The Phase II trial will continue to follow the original cohort to determine the long-term effects of the interventions (metformin versus lifestyle) on further diabetes development, microvascular outcomes, and CVD and CVD risk factors. Forty-five percent of participants are from diverse minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$1,100,000
Fiscal Years 2009–2010—\$2,200,000
Total Funding to Date—\$3,300,000

Current Active Organizations and Grant Numbers

1. University of Colorado Aurora, Colorado	—DK-048375
2. Louisiana State University Pennington Biomedical Research Center Baton Rouge, Louisiana	—DK-048377
3. Northwestern University Chicago, Illinois	—DK-048380
4. University of Chicago Chicago, Illinois	—DK-048381
5. MEDSTAR Research Institute Hyattsville, Maryland	—DK-048387

6. St. Luke's Roosevelt Institute for Health Sciences New York, New York	—DK-048404
7. Indiana University-Purdue University at Indianapolis Indianapolis, Indiana	—DK-048406
8. University of New Mexico Albuquerque, New Mexico	—DK-048407
9. University of Tennessee Health Science Center Memphis, Tennessee	—DK-048411
10. Seattle Institute for Biomedical and Clinical Research Seattle, Washington	—DK-048413
11. University of California, Los Angeles Los Angeles, California	—DK-048443
12. Thomas Jefferson University Philadelphia, Pennsylvania	—DK-048468
13. Johns Hopkins University Baltimore, Maryland	—DK-048485
14. George Washington University Washington, DC	—DK-048489
15. University of Texas Health Science Center San Antonio, Texas	—DK-048514

Early Adult Reduction of Weight Through LifestYle Intervention (EARLY) Trials,* Initiated in Fiscal Year 2009

The purpose of this program is to conduct two-phase clinical research to refine and test innovative behavioral approaches for weight control—using mobile phones, social networks, and Web-based curricula—in young adults, ages 18–35 years, who are at high risk for weight gain. Phase I involves refining proposed intervention, recruitment, retention, and adherence strategies. Phase II deals with testing the efficacy of the interventions that address weight loss, prevention of weight gain, or prevention of excessive weight gain during pregnancy. Targeted populations include pregnant and postpartum women, community college and university students, and young adults who are trying to quit smoking.

Obligations

Funding History:

Fiscal Year 2011—\$6,123,513
Fiscal Years 2009–2010—\$10,519,703
Total Funding to Date—\$16,643,216

* Formerly known as Targeted Approaches to Weight Control for Young Adults.

Current Active Organizations and Grant Numbers

1. University of Tennessee Health Science Center Memphis, Tennessee	—HL-096628
2. University of California, San Diego La Jolla, California	—HL-096715
3. Duke University Durham, North Carolina	—HL-096720
4. Cornell University Ithaca, New York	—HL-096760
5. University of Minnesota, Twin Cities Minneapolis, Minnesota	—HL-096767
6. University of Pittsburgh Pittsburgh, Pennsylvania	—HL-096770

Heart Failure Clinical Research Network, Initiated in Fiscal Year 2006

The purpose of this network is to accelerate research in the diagnosis and management of heart failure to improve outcomes through optimal application of existing therapies and evaluation of innovative therapies. The network provides the necessary infrastructure to develop, coordinate, and conduct multiple collaborative clinical protocols to facilitate application of emerging basic science discoveries into clinical investigations.

Obligations

Funding History:

Fiscal Year 2011—\$7,651,786
Fiscal Years 2006–2010—\$37,108,888
Total Funding to Date—\$44,760,674

Current Active Organizations and Grant Numbers

1. Minneapolis Medical Research Foundation, Inc. Minneapolis, Minnesota	—HL-084861
2. Duke University Durham, North Carolina	—HL-084875
3. Brigham and Women's Hospital Boston, Massachusetts	—HL-084877
4. University of Utah Salt Lake City, Utah	—HL-084889
5. Baylor College of Medicine Houston, Texas	—HL-084890
6. Morehouse School of Medicine Atlanta, Georgia	—HL-084891
7. University of Vermont and State Agriculture College Burlington, Vermont	—HL-084899
8. Duke University Durham, North Carolina	—HL-084904

9. Mayo Clinic College of Medicine Rochester, Minnesota	—HL-084907
10. Montreal Heart Institute Montreal, Quebec, Canada	—HL-084931

Look AHEAD: Action for Health in Diabetes, Initiated in Fiscal Year 2009

The purpose of this multicenter randomized clinical trial is to determine the effect of a lifestyle intervention—designed to achieve and maintain weight loss over the long term through decreased caloric intake and exercise—in obese individuals with type 2 diabetes. More than 5,000 participants have been randomly assigned to one of two interventions—either Lifestyle Intervention or Diabetes Support and Education—and will be followed for up to 11.5 years. The trial is monitoring major cardiovascular events, such as heart attack, stroke, and cardiovascular-related death. Investigators are also examining the impact of the interventions on other cardiovascular disease-related outcomes, cardiovascular risk factors, all-cause mortality, diabetes control and complications, fitness, general health, and health-related quality of life and psychological outcomes. One of the 16 clinical centers is targeting American Indians.

Obligations

Funding History:

Fiscal Year 2011—\$4,000,000
Fiscal Years 2009–2010—\$8,000,000
Total Funding to Date—\$12,000,000

Current Active Organizations and Grant Numbers

1. University of Alabama at Birmingham Birmingham, Alabama	—DK-057008
2. University of Tennessee Health Science Center Memphis, Tennessee	—DK-057078
3. University of Pennsylvania Philadelphia, Pennsylvania	—DK-057135
4. University of Colorado Denver, Colorado	—DK-057151

Network for Cardiothoracic Surgical Investigations in Cardiovascular Medicine, Initiated in Fiscal Year 2007

The purpose of this program is to establish a network to evaluate newer surgical techniques, technologies, devices, and innovative pharmaceutical and bioengineered products directed at CVD. The Network conducts randomized clinical trials and clinical studies that provide

a strong evidence base to inform surgical practice and disseminates its findings to the broader scientific community. The Network also serves as a clinical trials training ground for fellows and junior faculty.

Obligations

Funding History:

Fiscal Year 2011—\$5,459,597

Fiscal Years 2007–2010—\$25,978,806

Total Funding to Date—\$31,438,403

Current Active Organizations and Grant Numbers

- | | |
|--|------------|
| 1. University of Virginia, Charlottesville
Charlottesville, Virginia | —HL-088925 |
| 2. Emory University
Atlanta, Georgia | —HL-088928 |
| 3. Albert Einstein College of Medicine
of Yeshiva University
Bronx, New York | —HL-088939 |
| 4. Columbia University Health Sciences
New York, New York | —HL-088942 |
| 5. Mount Sinai School of Medicine
New York, New York | —HL-088951 |
| 6. Duke University
Durham, North Carolina | —HL-088953 |
| 7. Case Western Reserve University
Cleveland, Ohio | —HL-088955 |
| 8. University of Pennsylvania
Philadelphia, Pennsylvania | —HL-088957 |
| 9. Montreal Heart Institute
Montreal, Quebec, Canada | —HL-088963 |

Pediatric Heart Network, Initiated in Fiscal Year 2001

The objective of this study is to establish a clinical network to evaluate innovative treatment and management strategies for children with structural congenital heart disease, inflammatory heart disease, heart muscle disease, or arrhythmias.

Obligations

Funding History:

Fiscal Year 2011—\$12,827,177

Fiscal Years 2001–2009—\$63,348,873

Total Funding to Date—\$76,176,050

Current Active Organizations and Grant Numbers

- | | |
|---|------------|
| 1. New England Research Institute, Inc.
Watertown, Massachusetts | —HL-068270 |
| 2. Children's Hospital Medical Center
Cincinnati, Ohio | —HL-109673 |

- | | |
|---|------------|
| 3. University of Michigan at Ann Arbor
Ann Arbor, Michigan | —HL-109737 |
| 4. Baylor College of Medicine
Houston, Texas | —HL-109741 |
| 5. University of Utah
Salt Lake City, Utah | —HL-109743 |
| 6. Hospital for Sick Children
Toronto, Ontario | —HL-109777 |
| 7. Medical University of South Carolina
Charleston, South Carolina | —HL-109778 |
| 8. Emory University
Atlanta, Georgia | —HL-109781 |
| 9. Children's Hospital
Boston, Massachusetts | —HL-109816 |
| 10. Children's Hospital of Philadelphia
Philadelphia, Pennsylvania | —HL-109818 |

Pediatric HIV/AIDS Cohort Study (PHACS)—Data and Operations Center, Initiated in Fiscal Year 2006

The purpose of this study is to create a body of data that will enable researchers to understand more fully the effects of HIV on sexual maturation, pubertal development, and socialization of perinatally HIV-infected preadolescents and adolescents and to acquire more definitive information about the long-term safety of antiretroviral agents when used during pregnancy and in newborns.

Obligations

Funding History:

Fiscal Year 2011—\$600,000

Fiscal Years 2006–2010—\$3,259,021

Total Funding to Date—\$3,859,021

Current Active Organization and Contract Number

- | | |
|--|------------|
| 1. Harvard University
Boston, Massachusetts | —HD-052102 |
|--|------------|

Randomized Evaluation of VAD InterVENTion before Inotropic Therapy (REVIVE-IT), Initiated in Fiscal Year 2011

The purpose of this clinical feasibility study is to explore the potential benefit of mechanical circulatory support therapy using ventricular assist devices (VADs) in functionally-impaired advanced heart failure patients who have not yet developed serious consequences from their disease. The study will serve to inform a pivotal trial directed at a large and growing patient population for whom VADs could offer substantial benefit beyond current medical therapies.

Obligations

Funding History:

Fiscal Year 2011—\$4,952,781

Total Funding to Date—\$4,952,781

Current Active Organization and Contract Number:

- | | |
|---|----------------|
| 1. University of Michigan at Ann Arbor
Ann Arbor, Michigan | —268201100026C |
|---|----------------|

Systolic Blood Pressure Intervention Trial (SPRINT), Initiated in Fiscal Year 2009

The purpose of this study is to determine whether intensive lowering of systolic blood pressure below the currently recommended standard reduces the risk of cardiovascular and kidney diseases or age-related cognitive decline. About 7,500 participants with systolic blood pressure ≥ 130 mm Hg are being recruited at more than 80 clinics in 5 clinical center networks during a 2-year period and will be followed for 4–6 years. Among the participants, approximately 4,300 will have chronic kidney disease and 3,250 will be aged 75 years and older. The primary endpoints are nonfatal MI, acute coronary syndrome, stroke, heart failure, or CVD mortality. Secondary endpoints include decline in renal function or development of end stage renal disease. The MIND substudy is focusing on the effectiveness of lowering systolic blood pressure on reducing the decline in cognitive function.

Obligations

Funding History:

Fiscal Year 2011—\$0

Fiscal Years 2009–2010—\$36,882,407

Total Funding to Date—\$36,882,407

Current Active Organizations and Contract Numbers

- | | |
|---|------------|
| 1. Wake Forest University Health
Science Center
Winston-Salem, North Carolina | —HC-95240 |
| 2. University of Utah
Salt Lake City, Utah | —HC-95255 |
| 3. University of Birmingham
Birmingham, Alabama | —HC-95256 |
| 4. Wake Forest University Health
Science Center
Winston-Salem, North Carolina | — HC-95257 |
| 5. Case Western Reserve University
Cleveland, Ohio | —HC-95258 |
| 6. Department of Veterans Affairs, Memphis
Memphis, Tennessee | —HV-0514 |

Trial of Aldosterone Antagonists Therapy in Adults With Preserved Ejection Fraction Congestive Heart Failure (TOPCAT), Initiated in Fiscal Year 2004

The purpose of this international randomized trial is to evaluate the effectiveness of spironolactone, a generic and inexpensive drug, to reduce cardiovascular mortality and heart failure hospitalization in patients who have heart failure with preserved systolic function (left ventricular ejection fraction ≥ 45 percent). The trial is recruiting up to 3,500 patients in Argentina, Brazil, Canada, Republic of Georgia, Russia, and the United States.

Obligations

Funding History:

Fiscal Year 2011—\$0

Fiscal Years 2004–2010—\$26,914,315

Total Funding to Date—\$26,914,315

Current Active Organizations and Contract Numbers

- | | |
|--|-----------|
| 1. HHS Program Support Center, Supply
Service Center
Perry Point, Maryland | —HC-4071 |
| 2. New England Research Institutes, Inc.
Watertown, Massachusetts | —HC-45207 |

Women's Health Initiative (WHI), Initiated in Fiscal Year 1992

The WHI was established to elucidate the etiology and prevention of CVD, cancers, and osteoporosis in women aged 50–79 years. The program consists of three primary components: randomized controlled clinical trials of hormone therapy, dietary modification, and calcium/vitamin D supplementation; an observational study to identify predictors of disease; and a study of community approaches to developing healthful behaviors. Total participation includes 161,808 women, for whom 17 percent are from minority populations.

Currently, investigators are determining the long-term effects of prior hormone therapy on the cohort that participated in the clinical trials of hormone therapy. Minority participants are being followed to improve statistical power for genetic association. Self-reported outcome data are being collected and will be available for ancillary studies and a new generation of clinical trials. The rich resources of data and specimens are available to the scientific community and for training young investigators.

Obligations

Funding History:

Fiscal Year 2011—\$22,766,227

Fiscal Years 1992–2010*—\$793,070,018

Total Funding to Date—\$815,836,245

Current Active Organizations and Contract Numbers

1. State University of New York, Buffalo Buffalo, New York	—268201100001
2. Ohio State University Research Foundation Columbus, Ohio	—268201100002
3. Stanford University Stanford, California	—268201100003
4. Wake Forest University Health Sciences Winston-Salem, North Carolina	—268201100004
5. Fred Hutchinson Cancer Research Center Seattle, Washington	—268201100046
6. Wake Forest University Health Sciences Winston-Salem, North Carolina	—268200464221

Lung Diseases

Acute Respiratory Distress Syndrome Clinical Network (ARDSNet), Initiated in Fiscal Year 1994

The purpose of this network is to develop and conduct randomized controlled clinical trials to prevent and treat acute lung injury, ARDS, and other related critical illnesses and improve the outcome of patients with them.

Obligations

Funding History:

Fiscal Year 2011—\$5,096,234

Fiscal Years 1994–2010—\$78,742,505

Total Funding to Date—\$83,838,739

Current Active Organizations and Contract Numbers

1. Baystate Medical Center Springfield, Massachusetts	—HR-56165
2. University of California, San Francisco San Francisco, California	—HR-56166
3. University of Colorado Health Sciences Center Denver, Colorado	—HR-56167

4. Cleveland Clinic Lerner College of Medicine-Case Western Reserve University Cleveland, Ohio	—HR-56168
5. Duke University Medical Center Durham, North Carolina	—HR-56169
6. Johns Hopkins University Baltimore, Maryland	—HR-56170
7. LDS Hospital Salt Lake City, Utah	—HR-56171
8. Louisiana State University New Orleans, Louisiana	—HR-56172
9. University of Washington Seattle, Washington	—HR-56173
10. Vanderbilt University Medical Center Nashville, Tennessee	—HR-56174
11. Wake Forest University Health Sciences Winston-Salem, North Carolina	—HR-56175
12. Mayo Clinic College of Medicine Rochester, Minnesota	—HR-56176
13. Massachusetts General Hospital Boston, Massachusetts	—HR-56179

Asthma Network (AsthmaNet), Initiated in Fiscal Year 2009

The purpose of this network is to develop and conduct multiple clinical trials of asthma management in pediatric and adult populations. Investigators are seeking to identify optimal therapies for a variety of asthma phenotypes, genotypes, and racial and ethnic backgrounds. They are also conducting a limited number of proof-of-concept studies to advance the development of innovative therapies and perform studies to investigate the mechanistic bases for the interventions. Approximately 30 percent of the participants will be from diverse minority populations.

Obligations

Funding History:

Fiscal Year 2011—\$15,500,000

Fiscal Years 2009–2010—\$23,800,000

Total Funding to Date—\$39,300,000

Current Active Organizations and Grant Numbers

1. National Jewish Health Denver, Colorado	—HL-098075
2. University of Wisconsin, Madison Madison, Wisconsin	—HL-098090

* This figure reflects funding for the clinical trials and observational studies only. From 1992 to 1998, major support was provided through the Office of the Director, NIH. The Community Prevention Study receives funding through an inter-Agency agreement with the CDC: \$4,000,000 in FY 1999 and \$12,000,000 from FY 1996–1998.

3. Northwestern University Chicago, Illinois	—HL-098096
4. Washington University St. Louis, Missouri	—HL-098098
5. Brigham and Women's Hospital Boston, Massachusetts	—HL-098102
6. Wake Forest University Health Science Center Winston-Salem, North Carolina	—HL-098103
7. University of California, San Francisco San Francisco, California	—HL-098107
8. University of Arizona Tucson, Arizona	—HL-098112
9. Pennsylvania State University Hershey, Pennsylvania	—HL-098115
10. University of Pittsburgh Pittsburgh, Pennsylvania	—HL-098177

COPD Clinical Research Network, Initiated in Fiscal Year 2003

The purpose of this network is to investigate disease management approaches for patients with moderate-to-severe COPD and to ensure that the findings are rapidly disseminated to the medical community.

Obligations

Funding History:

Fiscal Year 2011—\$2,600,000
Fiscal Years 2003–2010—\$46,330,386
Total Funding to Date—\$48,930,386

Current Active Organizations and Grant Numbers

1. Harbor-UCLA Research and Education Institute Torrance, California	—HL-074407
2. Temple University Philadelphia, Pennsylvania	—HL-074408
3. Denver Health and Hospital Authority Denver, Colorado	—HL-074409
4. Minnesota Veterans Research Institute Minneapolis, Minnesota	—HL-074416
5. University of Alabama at Birmingham Birmingham, Alabama	—HL-074418
6. University of Michigan at Ann Arbor Ann Arbor, Michigan	—HL-074422
7. University of Minnesota, Twin Cities Minneapolis, Minnesota	—HL-074424
8. Brigham and Women's Hospital Boston, Massachusetts	—HL-074428
9. University of California, San Francisco San Francisco, California	—HL-074431

10. University of Pittsburgh Pittsburgh, Pennsylvania	—HL-074439
11. University of Maryland Baltimore Professional School Baltimore, Maryland	—HL-074441

Long-Term Oxygen Treatment Trial (LOTT), Initiated in Fiscal Year 2007

The purpose of this program is to determine the effectiveness and safety of long-term oxygen therapy in patients with COPD. Approximately 1,100 patients with moderate COPD are being enrolled to determine whether supplemental oxygen can improve their quality of life and extend their lifespan. Research findings will help the Centers for Medicare and Medicaid Services decide whether to extend Medicare coverage for home oxygen treatment for patients with moderately severe disease.

Obligations

Funding History:

Fiscal Year 2011—\$4,377,865
Fiscal Years 2007–2010—\$20,787,361
Total Funding to Date—\$25,165,226

Current Active Organizations and Contract Numbers

1. Brigham and Women's Hospital Boston, Massachusetts	—HR-76183
2. Cleveland Clinic Foundation Cleveland, Ohio	—HR-76184
3. Denver Health and Hospital Authority Denver, Colorado	—HR-76185
4. Duke University Medical Center Durham, North Carolina	—HR-76186
5. Kaiser Foundation Hospitals Portland, Oregon	—HR-76187
6. Los Angeles Biomedical Institute/Harbor-UCLA Los Angeles, California	—HR-76188
7. Ohio State University Columbus, Ohio	—HR-76189
8. Temple University Philadelphia, Pennsylvania	—HR-76190
9. University of Alabama at Birmingham Birmingham, Alabama	—HR-76191
10. University of Michigan at Ann Arbor Ann Arbor, Michigan	—HR-76192
11. University of Pittsburgh Pittsburgh, Pennsylvania	—HR-76193
12. University of Utah Salt Lake City, Utah	—HR-76194
13. University of Washington Seattle, Washington	—HR-76195

14. Washington University St. Louis, Missouri	—HR-76196
15. Johns Hopkins University Baltimore, Maryland	—HR-76197

Novel Therapies for Lung Diseases—Phase II, Initiated in Fiscal Year 2010

The purpose of this study is to conduct proof-of-concept Phase II clinical trials of innovative interventions for a sleep-associated lung disease or a cardiopulmonary disorder. Investigators are seeking to identify interventions that will have the potential to improve clinical management.

Obligations

Funding History:

Fiscal Year 2011—\$12,843,238
Fiscal Year 2010—\$7,593,739
Total Funding to Date—\$20,436,977

Current Active Organizations and Grant Numbers

1. Brigham and Women's Hospital Boston, Massachusetts	—HL-102225
2. University of Colorado Denver Aurora, Colorado	—HL-102235
3. University of Iowa Iowa City, Iowa	—HL-102288
4. Fred Hutchinson Cancer Research Center Seattle, Washington	—HL-102547
5. University of Wisconsin Madison, Wisconsin	—HL-105365
6. Brigham and Women's Hospital Boston, Massachusetts	—HL-105371
7. Johns Hopkins University Baltimore, Maryland	—HL-105569
8. Mayo Clinic Rochester, Minnesota	—HL-108712
9. University of California, San Francisco San Francisco, California	—HL-108713
10. Johns Hopkins University Baltimore, Maryland	—HL-108730

Blood Diseases and Resources

Blood and Marrow Transplant Clinical Research Network, Initiated in Fiscal Year 2001

The purpose of this network is to compare innovative treatment methods and management strategies of potential benefit for patients undergoing blood or marrow transplantation.

Obligations

Funding History:

Fiscal Year 2011—\$5,319,784
Fiscal Years 2001–2010—\$59,071,021
Total Funding to Date—\$64,390,805

Current Active Organizations and Grant Numbers

1. University of Nebraska Medical Center Omaha, Nebraska	—HL-069233
2. Fred Hutchinson Cancer Research Center Seattle, Washington	—HL-069246
3. Dana Farber Cancer Institute Boston, Massachusetts	—HL-069249
4. National Childhood Cancer Foundation Arcadia, California	—HL-069254
5. Duke University Durham, North Carolina	—HL-069274
6. City of Hope Medical Center Duarte, California	—HL-069278
7. University of Pennsylvania Philadelphia, Pennsylvania	—HL-069286
8. University of Minnesota, Twin Cities Minneapolis, Minnesota	—HL-069290
9. Stanford University Stanford, California	—HL-069291
10. Medical College of Wisconsin Milwaukee, Wisconsin	—HL-069294
11. University of Florida Gainesville, Florida	—HL-069301
12. Johns Hopkins University Baltimore, Maryland	—HL-069310
13. Sloan Kettering Institute for Cancer Research New York, New York	—HL-069315
14. University of Michigan at Ann Arbor Ann Arbor, Michigan	—HL-069330
15. University of Texas MD Anderson Cancer Center Houston, Texas	—HL-069334
16. Case Western Reserve University Cleveland, Ohio	—HL-069348
17. Baylor College of Medicine Houston, Texas	—HL-108945
18. H. Lee Moffitt Cancer Center Tampa, Florida	—HL-108987
19. Washington University St. Louis, Missouri	—HL-109137
20. Ohio State University Columbus, Ohio	—HL-109322
21. Northside Hospital Atlanta Atlanta, Georgia	—HL-109526

Pediatric Hydroxyurea Phase III Clinical Trial (BABY HUG), Initiated in Fiscal Year 2000

The objective of this clinical trial is to determine if hydroxyurea therapy is effective in preventing chronic end organ damage in pediatric patients with SCD.

Obligations

Funding History:

Fiscal Year 2011—\$852,640

Fiscal Years 2000–2010—\$21,847,533

Total Funding to Date—\$22,700,173

Current Active Organizations and Contract Numbers

1. Children's Research Institute Washington, DC	—HB-07150
2. Duke University Medical Center Durham, North Carolina	—HB-07151
3. Howard University Washington, DC	—HB-07152
4. Johns Hopkins University Baltimore, Maryland	—HB-07153
5. Medical University of South Carolina Charleston, South Carolina	—HB-07154
6. St. Jude Children's Research Hospital Memphis, Tennessee	—HB-07155
7. The Research Foundation of SUNY New York, New York	—HB-07156
8. University of Miami Miami, Florida	—HB-07157
9. University of Mississippi Medical Center Jackson, Mississippi	—HB-07158
10. University of Texas Southwestern Medical Center Dallas, Texas	—HB-07159
11. Clinical Trials and Surveys Corporation Baltimore, Maryland	—HB-07160

Sildenafil for Sickle Cell Disease-Associated Pulmonary Hypertension (walk PHaSST), Initiated in Fiscal Year 2006

The purpose of this clinical trial is to evaluate the safety and efficacy of 16 weeks of therapy with sildenafil, a nitric oxide potentiator, in adult patients with SCD and pulmonary hypertension. Exercise endurance and pulmonary artery pressure were measured. Pulmonary hypertension occurs in up to 30 percent of SCD cases and is strongly associated with mortality in adults with SCD. All of the participants are black.

The trial was stopped nearly 1 year early because the treatment group experienced either increased

hospitalizations for sickle cell crises or severe pain compared with the control group. The follow-up study is using datasets and biorepository specimens from the closed trial to assess the effect of genetic factors on hematologic, cardiopulmonary, and pain phenotypes in individuals with hemoglobin SS disease and to explore phenotypic variations in individuals with hemoglobin SC disease.

Obligations

Funding History:

Fiscal Year 2011—\$212,144

Fiscal Years 2006–2010—\$9,653,177

Total Funding to Date—\$9,865,321

Current Active Organizations and Contract Numbers

1. Rho Federal Systems Division, Inc. Chapel Hill, North Carolina	—HB-67182
2. Imperial College of London London, England	—HB-67183
3. Children's Hospital of Pittsburgh Pittsburgh, Pennsylvania	—HB-67184
4. University of Colorado Denver, Colorado	—HB-67185
5. Children's Hospital and Research Center at Oakland Oakland, California	—HB-67186
6. University of Illinois at Chicago Chicago, Illinois	—HB-67187
7. Johns Hopkins University Baltimore, Maryland	—HB-67188
8. Howard University Washington, DC	—HB-67189
9. Albert Einstein College of Medicine of Yeshiva University New York, New York	—HB-67190

Transfusion Medicine/Hemostasis Clinical Research Network, Initiated in Fiscal Year 2002

The purpose of this network is to compare new management strategies for individuals with hemostatic disorders, such as idiopathic thrombocytopenia and thrombotic thrombocytopenic purpura, and evaluate new and existing blood products and cytokines for treatment of hematologic disorders.

Obligations

Funding History:

Fiscal Year 2011—\$6,133,712

Fiscal Years 2002–2010—\$57,040,060

Total Funding to Date—\$63,173,772

Current Active Organizations and Grant Numbers

1. University of Iowa Iowa City, Iowa	—HL-072028	10. Duke University Durham, North Carolina	—HL-072289
2. Case Western Reserve University Cleveland, Ohio	—HL-072033	11. Blood Center of Southeastern Wisconsin Milwaukee, Wisconsin	—HL-072290
3. University of Minnesota, Twin Cities Minneapolis, Minnesota	—HL-072072	12. Children's Hospital Boston, Massachusetts	—HL-072291
4. Johns Hopkins University Baltimore, Maryland	—HL-072191	13. Massachusetts General Hospital Boston, Massachusetts	—HL-072299
5. Weill Medical College of Cornell University New York, New York	—HL-072196	14. Puget Sound Blood Center Seattle, Washington	—HL-072305
6. Emory University Atlanta, Georgia	—HL-072248	15. University of Pittsburgh Pittsburgh, Pennsylvania	—HL-072331
7. New England Research Institutes, Inc. Watertown, Massachusetts	—HL-072268	16. University of Pennsylvania Philadelphia, Pennsylvania	—HL-072346
8. Tulane University of Louisiana New Orleans, Louisiana	—HL-072274	17. University of North Carolina at Chapel Hill Chapel Hill, North Carolina	—HL-072355
9. University of Oklahoma Health Sciences Center Oklahoma City, Oklahoma	—HL-072283	18. University of Maryland Baltimore Professional School Baltimore, Maryland	—HL-072359



12. Activities To Promote Diversity and Address Health Disparities

Throughout its history, the NHLBI has been a leader in conducting and supporting research and programs to eliminate health disparities that exist between various segments of the U.S. population. The Institute has not only initiated research projects with significant racial and ethnic minority participation to compare health status between various populations, but has also given high priority to programs that focus exclusively on minority health issues.

Since FY 1991, the Institute has had procedures in place to ensure full compliance with the NIH Policy on Inclusion of Women and Minorities as Subjects in Clinical Research. As a result, all NHLBI-supported research that involves human subjects includes minorities, with the exception of a very few projects for which a compelling justification for limited diversity in the study population exists. Thus, all segments of the population, minority and nonminority, women, and children, stand to benefit from the Institute's research programs.

It has long been a goal of the NHLBI to increase the number of individuals from underrepresented groups in biomedical and behavioral research. Selected FY 2011 activities addressing this goal include the following:

- NHLBI Research Centers at Minority Serving Institutions: Supports establishment of research centers at minority serving institutions to augment and strengthen research capabilities and resources related to heart, lung, and blood diseases and disorders, with the goal of enabling the institutions and their investigators to become fully competitive
- Sickle Cell Scholars Program: Supports career development of young or new investigators in SCD research as part of the Basic and Translational Research Program (BTRP) (see Chapter 9)
- Summer-for-Sickle Cell-Science Program: Supports career development of young or new investigators in SCD research as part of the BTRP program

- Clinical Research Education and Career Development in Minority Institutions: Encourages the development and implementation of curriculum-dependent programs in minority institutions to train selected doctoral and postdoctoral candidates in clinical research leading to a Master of Science in Clinical Research or Master of Public Health in a clinically relevant area
- Short-Term Research Education Program To Increase Diversity in Health-Related Research: Promotes diversity in undergraduate and health professional student populations by offering short-term education support to stimulate career development in cardiovascular, lung, and blood diseases and sleep disorders research
- Program To Increase Diversity Among Individuals Engaged in Health-Related Research (PRIDE): Encourages scientists and research-oriented faculty from diverse backgrounds to expand their research skills and gain experience in advanced methods and experimental approaches in basic and applied sciences in heart, lung, and blood diseases and sleep disorders so that they can compete for external funding for research in the biomedical and behavioral sciences
- Mentored Career Development Award To Promote Faculty Diversity/Re-Entry in Biomedical Research: Promotes an increase in the number of highly trained investigators—from diverse backgrounds (i.e., faculty members who are from underrepresented racial and ethnic groups or who have disabilities or who are from disadvantaged backgrounds) or those who have experienced an interruption in their research careers—whose basic and clinical research interests are grounded in the advanced methods and experimental approaches needed to solve problems related to cardiovascular, lung, and blood diseases and sleep disorders
- Mentored Career Award for Faculty at Minority Serving Institutions: Encourages eligible faculty members at institutions that promote diversity to undertake special studies and supervised research under a mentor who is an accomplished

investigator in the research area proposed and who is experienced in developing independent investigators

- Minority Institutional Research Training Program: Supports the training of pre- and postdoctoral students and certain health professional students at minority serving institutions in order to provide additional opportunities to develop research skills and to encourage the pursuit of research careers.
- Support of Competitive Research (SCORE) Program: Fosters the development of faculty at minority serving institutions to increase their research competitiveness in the areas of heart, lung, and blood diseases and sleep disorders and to promote their transition to non-SCORE external sources of funding; and supports pilot awards for individuals at the beginning stages of a research career who are interested in testing a new idea or generating preliminary data, and for more experienced investigators who are interested in switching to a different field of research

The Office of Research Training and Minority Health (ORTMH) within the Office of the Director provides oversight for, and coordinates, supports, and evaluates Institute programs related to minority health outcomes, including research, research training and career development, public outreach, and translation of research findings. The ORTMH also coordinates activities to foster greater participation of underrepresented minorities, individuals from disadvantaged backgrounds, and individuals with disabilities in NHLBI research and research training and career development programs. Selected FY 2011 activities include the following:

- Issuing four training and career development RFAs to increase the number of highly trained individuals from diverse backgrounds, including individuals from underrepresented racial and ethnic groups, individuals from disadvantaged backgrounds, and individuals with disabilities
- Participating in HHS-Endorsed Minority Organization Internship Programs by supporting positions in NHLBI extramural and intramural divisions for students from the National Association for Equal Opportunity in Higher Education, the Hispanic Association of Colleges and Universities, the Washington Internships for Native Students programs, and the Directors of Health Promotion and Education Internship Program/CDC

- Sponsoring Out of the Box, a project for the Cherokee and Smokey Mountain Elementary Schools that is designed to create awareness and interest in the importance of science, medicine, and health; eliminate gaps in quality of health among diverse groups by encouraging an interest in health-related careers; and empower children to take responsibility for their health
- Supporting the African American, Hispanic, and Native American Youth Initiatives to bring minority students to the NIH campus for scientific presentations, an introduction to NHLBI research training and career development programs, and a tour of NHLBI laboratories
- Providing undergraduate students from the Tougaloo College Scholars program with an opportunity to visit the NIH for 3 days to learn about biomedical research and research training opportunities at the NHLBI/NIH
- Increasing recruitment of individuals for the NHLBI intramural and extramural training programs by representing the Institute at four diversity-focused research meetings to raise awareness of research and research training and career development opportunities supported by the NHLBI
- Coordinating the Biomedical Research Training Program for Individuals From Underrepresented Groups, which offers opportunities for underrepresented health professional degree students and post-baccalaureate individuals to receive training in fundamental biomedical sciences and clinical research as they relate to the etiology and treatment of heart, blood vessel, lung, and blood diseases
- Serving as the NHLBI contact for guidance to candidates applying for the NIH Pathway to Independence Award and the NHLBI Career Transition Award for extramural programmatic issues

In addition, the DCVS Office of Research Training and Career Development sponsored the 18th annual Diversity Supplement Trainee Meeting prior to the American Heart Association Annual Scientific Meetings. The meeting brought together awardees, faculty mentors, and NHLBI program staff for scientific and poster presentations, discussions, and opportunities to network with colleagues.

See Chapter 13 for additional NHLBI-supported research training and career development programs for individuals from diverse backgrounds.

The following text describes selected current projects that focus on minority populations and reflect the Institute's research portfolio related to minority health. Additional information can be found in Chapters 9–11.

Heart and Vascular Diseases

Epidemiology

Long-term epidemiologic studies are critical to uncovering risk factors that lead to disease. The Institute has initiated several major studies of heart disease focused significantly or completely on minority populations:

- CARDIA (see Chapter 10): To determine the evolution of CHD risk factors and lifestyle characteristics in young adults that may influence development of risk factors prior to middle age. Fifty percent of participants are black.
- ARIC (see Chapter 10): To investigate the association between CHD risk factors and the development of atherosclerosis and CVD in an adult population. Approximately 30 percent of participants are black.
- Strong Heart Study (see Chapter 9): To compare risk factor levels and morbidity and mortality from CVD among American Indians from three different geographic locations. In 2000, the study began focusing on large, extended families on which linkage analyses and gene discovery studies could be conducted.
- JHS (see Chapter 10): To identify environmental and genetic factors influencing evolution and progression of CVD in blacks.
- MESA (see Chapter 10): To examine the characteristics of subclinical CVD that predict progression to clinically overt CVD and related risk factors that predict subclinical disease in blacks, whites, Hispanics, and Asians. Sixty-two percent of participants are from minority populations.
- HCHS (see Chapter 10): To identify risk factors for cardiovascular and lung disease in Hispanic populations in the United States and determine the role of acculturation in their prevalence and development.

- GOCADAN (see Chapter 9): To study the relative contributions of genetic and environmental factors to CVD in Eskimos in the Norton Sound region of Alaska.

The Institute supports components of the NHANES that track the prevalence of disease and risk factors for cardiovascular and lung diseases by race and ethnicity in the U.S. population and the National Longitudinal Mortality Study that analyzes socioeconomic, demographic, occupational, and racial differentials in mortality in the United States.

The NHLBI also supports a variety of investigator-initiated research activities across a range of racial and ethnic groups on risk factors, genetic contributors, and health disparities in heart, lung, and blood diseases and sleep disorders. Many of them are ancillary studies to NHLBI-initiated cohort studies.

Risk Factors

Investigator-initiated studies on cardiovascular risk factors in underrepresented racial and ethnic groups range in focus from biological to environmental, psychosocial, and cultural. One study among blacks in the JHS is characterizing the relationships between vascular function and CVD risk factors and indicators of subclinical CVD, and detecting gene variants that influence vascular function. A second study is investigating the effects of low vitamin D levels in blacks in the biethnic ARIC cohort. A third study involves echocardiographic examination of a subsample of adult participants in the HCHS to determine the prevalence of abnormal cardiac function, the degree of heterogeneity in cardiac function between Hispanic subgroups, and the relationship between cardiac structure and function and such factors as diabetes biomarkers, psychosocial measures, and measures of socioeconomic status in Hispanics.

Additional studies are determining geographic and ethnic variations in the prevalence of CHD risk factors; investigating risk factors linked to atherosclerosis and disease progression in South Asians that live in America; and examining the effects of arsenic exposure on diabetes and CVD in American Indians.

Genetic Epidemiology

Genetic epidemiology is concerned with the role of genetic factors in the etiology of disease within groups of relatives and the interplay of genetic factors with environmental factors. NHLBI-supported studies include those focusing on gene discovery through both linkage studies in family-based samples and genome-wide association studies (GWASs) in population-based samples; the effects of gene–environment interactions on risk factors and health; and genotypic characterization in relationship to intermediate phenotypes, such as biomarkers.

Genetic epidemiologic research is also beginning to transition to predicting and assessing genetic risk and reporting genetic results to participants of research studies. The activities offer unique insights into specific populations and general observations that are relevant to all populations. Examples are studies of genetics of hypertension in populations of West African origin; the role of stress in gene–environment interaction in a multi-ethnic population; the contribution of genetic variation to obstructive sleep apnea, impaired endothelial function, and central blood pressure in Mexican Americans; salt sensitivity and blood pressure in Chinese populations; genetics of high serum triglycerides and metabolic traits in Mexican Americans; and genetic variation that underlies obesity and obesity-related phenotypes among Samoan adults.

A new study is investigating the molecular mechanisms that underlie the genetic basis of left ventricular hypertrophy. This study is leveraging epidemiological and genetic work of the NHLBI-supported Hypertension Genetic Epidemiology Network Echo Study. Fifty-five percent of participants are black.

Health Disparities

The NHLBI is committed to supporting research that will contribute to reducing health disparities among racial and ethnic minorities.

- CPHD (see Chapter 9): To promote transdisciplinary research in health disparities in order to improve health outcomes and quality of life for populations at high risk for CVD. The population will consist primarily of blacks and Hispanics.
- NHLBI Centers for Cardiovascular Outcomes Research (see Chapter 9): To conduct research that focuses on measuring, evaluating, and improving

outcomes of cardiovascular care. Approximately 30 percent of participants will be from racial and ethnic minority populations.

A project within the Mentored Career Development Award To Promote Faculty Diversity/Re-Entry in Biomedical Research program is conducting research on racial disparities between blacks and whites who live in the inner city by focusing on issues related to physical fitness and cardiovascular risk reduction.

Education

The NHLBI hosts *Children and Clinical Studies* (see <http://www.ChildrenAndClinicalStudies.nhlbi.nih.gov>), an educational Web site, for children and their families and health care providers, to improve understanding of pediatric research. The site offers information in English and Spanish about the importance of research on children, safety measures in clinical trials, and potential effects on a family when a child is enrolled in a study. The site also has a section about minority interests and concerns.

The NHLBI, through the DARD, translates research findings into practice by facilitating the development of clinical practice guidelines; communicating research advances; and disseminating health information to physicians, health care professionals, patients, and the public on ways to prevent or treat diseases within the Institute's mandate.

The Institute supports the following activities to improve cardiovascular health in racial and cultural/ethnic groups:

- The Community Health Worker Initiative: To improve cardiovascular health among black, Hispanic, American Indian, Alaska Native, and Filipino-American communities using the community health worker model to promote evidence-based, heart health knowledge, attitudes, and behaviors.
- NHLBI-U.S. Department of Housing and Urban Development HOPE VI: To address cardiovascular health disparities in public housing settings. With the aid of the NHLBI evidence-based, heart health curriculum, residents of the black community are trained to become health educators of other public housing residents.

- Salud para su Corazón: To develop networks and partnerships to disseminate information and strategies and to train lay educators about evidence-based CVD prevention and control, in order to promote heart healthy knowledge, attitudes, and behaviors in Hispanic communities.
- Honoring the Gift of Heart Health: To train community health workers to deliver culturally appropriate, evidence-based curricula to prevent and control CVD risk factors in American Indian tribal communities.
- Healthy Heart, Healthy Family: To train community health workers to deliver culturally and linguistically appropriate curricula and to conduct outreach activities that increase community awareness of heart disease and its associated risk factors and that promote heart healthy lifestyles among the growing population of Filipino heritage in the United States.
- The Heart Truth® Campaign: To raise awareness of heart disease in women through community-based interventions and social marketing media. Special populations are especially targeted through the Heart Truth's Women of Color Initiative, a partnership with national black and Hispanic organizations.
- **We Can!**® (Ways to Enhance Children's Activity & Nutrition): To help children ages 8–13 years maintain a healthy weight by providing curricula, tools, tips, and other resources to parents, caregivers, communities, and other organizations. Special attention is directed to black, Hispanic, and American Indian/Alaska Native populations.

In addition to the activities mentioned above, the Institute has prepared and distributed publications on CVD prevention for minority populations. They include the following:

- *With Every Heartbeat Is Life: A Community Health Worker's Manual for African Americans* (and a series of picture cards)
- *On the Move to Better Heart Health for African Americans* (easy-to-read booklet on heart healthy living)
- *Your Choice for Change: Honoring the Gift of Heart Health for American Indians* (easy-to-read booklet on heart healthy living)

- *Healthy Heart, Healthy Family: A Community Health Worker's Manual for the Filipino Community* in English and Tagalog (and a series of picture cards)
- *Healthy Heart, Healthy Family Series* (six easy-to-read English and Tagalog booklets on heart healthy living for the Filipino community)
- *Your Heart Is Golden: Heart Health Promotion Activities for Vietnamese Communities*
- *Your Heart, Your Life: A Health Educator's Manual for the Latino Community* in English and Spanish (and a series of bilingual picture cards and a DVD)
- *Healthy Hearts, Healthy Homes Series* (six easy-to-read English and Spanish booklets on heart healthy living)
- *The Heart Truth for African American Women: An Action Plan*
- *The Heart Truth for Latinas: An Action Plan*

The educational materials listed throughout this chapter can be obtained from the NHLBI public Web site or through the NHLBI online catalog.

Arrhythmias

Arrhythmia is a disorder of the normal rhythms of the heart, whereby it beats too fast (tachycardia), too slow (bradycardia), or irregularly. The NHLBI is supporting basic and genetic research on the mechanisms that underlie cardiac arrhythmias to improve their diagnosis, treatment, and prevention in all racial and ethnic groups in the United States. One study is evaluating variants of the gene *NOS1AP* that affect the QT interval in various racial and ethnic populations. Another study is evaluating the distributions of electrocardiographic predictors of atrial fibrillation in racial and ethnic groups to determine whether minority populations have a higher risk for this arrhythmia than whites. A third study is investigating the role of sodium channel polymorphisms in cardiac arrhythmias in families with Long QT syndrome and Brugada Syndrome; approximately 66 percent of participants will be from racial and ethnic minority populations.

Scientists have identified an association between variations in certain receptors that are activated during sympathetic nervous system stimulation and an increased risk of sudden cardiac death, most often due to ventricular arrhythmia. Although significant differences in

associated risk of sudden cardiac death were not found between blacks and whites, continued research is expected to advance the understanding of differences in genetic predisposition for cardiac arrhythmias among racial and ethnic groups and ultimately lead to improved therapy.

Investigators identified two mutations in a gene encoding the major intracellular calcium release channel in two cases of SIDS. One mutation was identified in 1 of 50 black infants and the other in 1 of 83 white infants. Researchers reported a pathogenic mechanism for SIDS, whereby SIDS-linked mutant ion channels become “leaky” during stress and thus potentially trigger fatal cardiac arrhythmias. Importantly, researchers have found further evidence that cardiac arrhythmias of genetic origin contribute to SIDS, a concept that offers a chance to prevent some avoidable tragedies.

Heart Failure

Heart failure (heart muscle dysfunction) affects approximately 5 million Americans and is a growing public health concern. It is frequently the end result of other conditions, such as hypertension, diabetes, and prior heart attacks.

Findings from the CARDIA study showed that heart failure before age 50 is substantially more common in blacks than in whites. Heart failure in black participants in the study was often preceded by such risk factors as high blood pressure, obesity, and chronic kidney disease 10 to 20 years prior to the heart failure.

The Institute has initiated a study to facilitate the translation of basic science discoveries into clinical applications:

- Heart Failure Clinical Research Network (see Chapter 9): To develop, coordinate, and conduct multiple collaborative proof-of-concept clinical protocols to improve heart failure outcomes. The Network includes a historically black medical center with minority investigators and access to a high-risk, underserved population.

Investigator-initiated research is targeting heart failure among minority populations. A study is examining the role of impaired ATP synthesis and delivery in contractile dysfunction in heart failure patients; approximately 40 percent of participants are from racial minority

populations. Another study is examining the underlying genetic variations that result in familial dilated cardiomyopathy, an inherited form of heart dysfunction. A study is focusing on angioedema—a severe allergic reaction and a life-threatening side effect of ACE-inhibitor drugs that is more common in blacks than in whites. Investigators are determining the mechanisms that cause this side effect and are studying the genetic profile of affected individuals and their families to determine who should avoid taking the drugs. Another study is focusing on drugs commonly used to treat heart failure to determine whether beta blockers are as effective in black as in white heart failure patients; researchers will study genetic and other factors that affect the difference.

Individuals with heart failure typically have a poor prognosis and a diminished quality of life. Inadequate health literacy skills, depression, and emotional stress are common and are often associated with poor health outcomes. One study is determining whether cognitive behavioral intervention is superior to supportive clinical management for depressed outpatients with heart failure; approximately 30 percent of participants are expected to be black. Another study is examining the role of emotional stress on heart failure exacerbations; a large proportion of the population is black. A third study is investigating whether an exercise intervention, such as tai chi, can increase physical function, cardiac functional capacity, and quality of life in patients with heart failure; nearly 50 percent of participants are expected to come from minority populations.

High Blood Pressure

Etiology

High blood pressure is a serious health problem that is especially prevalent and severe among minorities.

The NHLBI supports a number of investigator-initiated studies to identify genes linked to hypertension in blacks, Mexican Americans, Asians, and whites to determine if part of the disparity in prevalence can be attributed to genetic differences among the groups. Genes under investigation include those associated with the renin-angiotensin system, the autonomic nervous system, and sodium transport.

The role of dietary factors, particularly macronutrients, in the etiology of high blood pressure is another area of investigation. Scientists are conducting

epidemiologic studies among participants with diverse ethnicity, SES, and dietary habits in four countries to determine the effect of selected dietary components (proteins, lipids, carbohydrates, amino acids, calcium, magnesium, sodium, potassium, antioxidants, fiber, and caffeine) on blood pressure.

Researchers are also seeking to understand the role of obesity in the development of high blood pressure. A new study will investigate the effect of natriuretic peptides (hormones produced by the heart) in blood pressure regulation and dietary salt sensitivity, and relate findings to previous findings that obese individuals have decreased numbers of circulating natriuretic peptides. Approximately 40 percent of study participants are expected to be from minority subgroups.

Treatment and Prevention

Identifying effective treatment strategies for various populations requires large-scale studies with representative populations in sufficient numbers.

- SPRINT (see Chapter 11): To determine whether intensive lowering of systolic blood pressure below the currently recommended standard will reduce the risk of cardiovascular and kidney diseases or age-related cognitive decline. Approximately 25 percent of participants will be from racial and ethnic populations.

The Institute also supports a number of investigator-initiated studies to prevent hypertension and improve blood pressure control in racial and ethnic minorities. Interventions target both lay and medical communities. Strategies being tested include communication skill enhancement, organizational change, educational programs, lifestyle and nutritional counseling, use of technology, case management, pharmacy-based interventions, and provision of care by community health workers and other nontraditional providers. One study is testing a church-based lifestyle intervention to reduce blood pressure using group classes and motivational interviews to help participants make and maintain therapeutic lifestyle changes; approximately 400 blacks with high blood pressure are expected to participate.

Education

The NHLBI has developed a number of outreach activities to inform minority populations of the importance of blood pressure control. Several publications and Web-based products have been developed and distributed for health professionals, patients, and the public. Some examples are:

- *Presión Arterial Alta: NHLBI Diseases and Conditions Index*
- *Keep the Beat: Control Your High Blood Pressure* in English and Spanish
- *Help Your Heart: Control Your High Blood Pressure* in Tagalog and English
- *Keep Your Heart in Check—Know Your Blood Pressure Number* in Vietnamese and English
- *Your Choice for Change: Honoring the Gift of Heart Health for American Indians*

High Serum Cholesterol

Etiology

The Institute supports investigator-initiated studies to identify genes that influence the lipoprotein profile in various racial and ethnic groups. Research findings could offer an explanation for differences in susceptibility to CHD found among the groups. A project involving extended families of Mexican Americans in the San Antonio Family Heart Study has detected and mapped many quantitative trait loci (QTLs) for CVD risk factors, including some that influence HDL and LDL levels. Scientists are seeking to identify genes for QTLs that are related to lipoproteins.

Treatment

Research has shown that patients with elevated LDL levels who have been advised to make lifestyle changes and to take statins often do not comply with the prescribed regimens. An investigator-initiated study is seeking to develop and evaluate an interactive virtual environment system to increase the initiation and maintenance of medication adherence and therapeutic lifestyle change in patients who are at risk for CHD. Patients will be able to access the system to seek advice from a virtual health care provider and get assistance in developing an effective care plan that is based on clinical guidelines.

Education

The Institute has prepared the following publications on blood cholesterol for minority audiences:

- *Healthy Hearts, Healthy Homes—Do You Know Your Cholesterol Levels?* in English and Spanish
- *Heart-Healthy Home Cooking African American Style*
- *Delicious Heart-Healthy Latino Recipes* in English and Spanish
- *Healthy Heart, Healthy Family—Be Heart Smart: Keep Your Cholesterol in Check* in Tagalog and English
- *Serve Up a Healthy Life—Give the Gift of Good Nutrition* in Vietnamese and English

Obesity

Etiology

Obesity is a major health concern that affects children and adults. Minorities—including American Indians, blacks, and Mexican Americans—are especially at risk. Data from the 2007–2008 NHANES show that 34 percent of adults and 17 percent of children aged 2–19 years are obese. Understanding the causes of obesity could lead to effective strategies to combat it. A long-term investigator-initiated study is examining parental and extended family influences on the development of childhood obesity in Mexican American children.

An investigator-initiated study seeks to elucidate interconnected biological and social pathways associated with adolescent obesity and risk for later development of type 2 diabetes and CVD in Latin American youth. The goal of the study is to identify modifiable conditions in order to prevent obesity and related diseases.

The NHLBI funds several studies that focus on genetic risk factors for obesity in one or more minority populations. Evidence for obesity genes has been identified on chromosome 4 in American Indians, on chromosome 9 in Mexican Americans, and on chromosomes 5 and 6 in blacks. In some cases, the results confirm those found in European Americans, and in other cases, the results represent novel findings.

Researchers have found that black and Hispanic children are especially likely to develop sleep apnea. An investigator-initiated study will assess the role of obesity

in the development of abnormalities that increase the likelihood of developing sleep apnea and whether the problem can be corrected with weight loss. Blacks comprise approximately 80 percent of participants. Another study with multi-ethnic participation is determining whether obese children with sleep apnea are at increased risk of cognitive impairment and vascular disease.

Treatment and Prevention

The NHLBI has initiated programs to test approaches for treating or preventing obesity:

- EARLY Trials (see Chapter 11): To develop and evaluate innovative approaches for weight control in young adults from ethnically and socioeconomically diverse populations who are at high risk for weight gain.
- ORBIT (see Chapter 9): To translate findings from basic research on human behavior into more effective clinical, community, and population interventions to reduce obesity and improve obesity-related behaviors. Some of the studies are expected to have 50- to 100-percent participation from minority populations.
- COPTR Consortium (see Chapter 11): To test interventions to prevent excess weight gain in non-overweight and overweight youth and to reduce weight in obese and severely obese youth. More than 50 percent of participants are expected to be from racial or ethnic minority populations.

The Institute supports a number of investigator-initiated studies on the effectiveness of obesity prevention and control interventions among diverse populations. They include testing an integrated school- and community-based intervention that involves physical activity and diet to reduce the prevalence of obesity among Hispanics, people of Asian heritage, and whites; evaluating, in a primarily minority population, the effect of adding environmental approaches to a standard family-based intervention to reduce overeating in obese children; exploring whether naturally occurring social support networks can help parents manage their children's weight in a population where 40 percent of the children are expected to be from racial and ethnic minority populations; and evaluating an intervention that supports primary care treatment of obesity in adults with at least one other cardiovascular risk factor; one project has strong Hispanic participation.

Many obese adults have difficulty breathing upon physical exertion and therefore are unable to exercise sufficiently. It is unclear whether this is because the individuals are in poor physical condition and could be helped with endurance exercise training, or whether obesity-related respiratory changes have occurred that necessitate weight loss before exercise training can be effective. Researchers are planning to investigate this question in obese individuals (approximately 40 percent of whom will be from minority populations) by assessing the effect of endurance exercise training (without weight loss) versus weight loss (without exercise training) on breathing difficulties after physical exertion.

Education

The NHLBI has prepared and distributed health information for minorities on losing excess weight:

- *Healthy Hearts, Healthy Homes—Do You Need To Lose Weight?* in English and Spanish
- *¿En Qué Consiste el Sobrepeso y la Obesidad? (What are Overweight and Obesity?)* in the NHLBI Diseases and Conditions Index
- **We Can!**[®]: Many bilingual (English and Spanish) publications on energy balance are available on the Web site at <http://wecan.nhlbi.nih.gov>

Physical Inactivity

Despite substantial research about the benefits of physical activity on CVD and its risk factors, physical inactivity is highly prevalent, especially among minority populations. Researchers have observed an age-related decline in physical activity or aerobic capacity in the biracial cohorts of two Institute-initiated longitudinal cohort studies.

The transition from elementary school to middle school marks a critical stage in the development of young people and the period during which physical activity tends to decline dramatically. An investigator-initiated study is investigating factors that contribute to the decline and the potential moderating effects of gender, race, socioeconomic status, and neighborhood environment on factors that influence changes in physical activity. A majority of participants will be from racial and ethnic minority populations. Another study will

collect data on aspects of neighborhood environments that are most often associated with physical activity in adolescents and determine whether this information can be used, via interventions offered in the offices of pediatricians, to help children increase their physical activity. A third study will develop and evaluate policies to increase physical activity and improve nutrition during summer and afterschool programs sponsored by the YMCA; approximately 50 percent of the children in the programs being evaluated are expected to be black.

Regular physical activity is important for cardiovascular health throughout life. Investigators are designing and evaluating a long-term, multilevel physical activity intervention for sedentary residents who are living in retirement communities. The intervention uses a variety of elements, including self-monitoring with a pedometer, group sessions and peer mentoring, and such environmental components as tailored walking maps. Approximately 25 percent of participants are expected to be from racial and ethnic minority populations.

Several investigator-initiated studies are evaluating culturally appropriate interventions to increase physical activity. Projects include those that use faith-based approaches involving church leaders and congregations to increase activity levels in blacks and those that test culturally targeted interventions in schools or among pregnant women and parents with young children.

Several projects are using mobile phone technology to increase physical activity and decrease sedentary behaviors. These studies capitalize on recent advances in communication technologies, such as “smart phones,” that offer a new way to deliver convenient and sustainable adherence strategies. In one study, women are receiving prompts, video clips, and individualized feedback via their cell phones to help them increase their physical activity levels; approximately 60 percent of the women are expected to be from racial and ethnic minority populations. In another study with large minority participation, researchers are assessing a program that is designed to improve diet and activity levels in sedentary people with poor quality diets; all participants use “smart phones” to monitor themselves and transmit information to a personal coach.

Education

The Institute has prepared and distributed the following publications for minorities on the importance of physical activity and ways to become more physically active:

- *On the Move to Better Health for African Americans*
- *American Indian and Alaska Native People: Be Active for Your Heart!*
- *Are You at Risk for Heart Disease?* in Tagalog and English
- *Be Active for a Healthier Heart* in Vietnamese and English
- **We Can!®**: Many bilingual (English and Spanish) publications on physical activity and energy balance are available on the Web site at <http://wecan.nhlbi.nih.gov>

The Institute also has developed a Web-based application on physical activity for lay health educators in English and Spanish, which can be found at <http://hin.nhlbi.nih.gov/salud/pa/index.htm>.

Smoking

Smoking is a major risk factor for CHD, stroke, COPD, and other cardiovascular and respiratory conditions and is the leading cause of preventable death. Although considerable progress has been made in reducing smoking rates and providing effective prevention and cessation interventions, additional research is needed to extend these efforts and improve the maintenance of behavior change.

The Institute has initiated smoking intervention programs in specialized groups:

- CHART (see Chapter 11): To evaluate the translation of efficacious smoking cessation strategies initiated during hospitalization and continued post-discharge into effective programs that can be widely implemented in routine clinical practice and assess the cost-effectiveness of these interventions. One of the projects will have approximately 75 percent participation from Asian, Hispanic, and black populations.
- Longitudinal Studies of HIV-Associated Lung Infections and Complications: To develop and evaluate a specialized smoking cessation intervention for the

treatment of nicotine dependence in HIV-seropositive smokers who are at high risk of developing accelerated emphysema; 40 percent of participants are black.

The NHLBI supports a number of investigator-initiated studies of smoking cessation in underserved populations. One study among predominately black women who live in public housing neighborhoods is evaluating smoking cessation interventions that use a combination of strategies—including contact with community health workers, small-group behavioral counseling, and neighborhood support groups. Another study is assessing the efficacy of telemedicine as a way to provide smoking cessation counseling to primary care patients who live in rural areas. Two studies among military personnel are testing interventions that focus on smoking cessation and subsequent abstinence; approximately 25–40 percent of participants are expected to come from racial and ethnic minority populations.

Education

The Institute has prepared the following publications on smoking cessation for minorities:

- *Enjoy Living Smoke Free* in English and Spanish
- *Be Heart Healthy: Enjoy Living Smoke Free* in Tagalog and English
- *Don't Burn Your Life Away—Be Good to Your Heart* in Tagalog and English and in Vietnamese and English

Psychosocial Factors

Etiology

A large and consistent body of evidence has demonstrated that psychosocial factors—such as depression, stress, and low social support—are associated with elevated risk for CVD and major adverse cardiac events in heart disease patients. Additionally, race and ethnicity, gender, and social class are important factors that can influence these associations in important ways.

The NHLBI is funding research to identify more precisely the nature of the relationship between depression and adverse cardiac outcomes and the conditions associated with successful treatment of depression. Research results will guide clinical care and inform future trials of depression treatment in heart patients and minority

patient populations. One study is seeking to increase understanding of the mechanisms that lead to death in depressed patients with heart disease by assessing the effects of stress on heart and brain function in heart disease patients with and without depression; approximately 45 percent of participants are expected to be from minority groups. Another study is examining the potential epigenetic mechanisms that link depression and cardiovascular disease. More than 50 percent of participants are expected to be black.

The Institute supports investigator-initiated research on the interactions of psychosocial factors with race and ethnicity, environmental factors, and low SES in the development of CHD. Scientists are investigating the contribution of biobehavioral factors (hostility, anxiety, and heightened cardiovascular reactivity to stress) in the etiology, pathogenesis, and course of CHD. Racial differences in stress-induced physiologic responses are also being examined.

The nature of the relationship between acute and chronic forms of stress and cardiac morbidity and mortality is particularly relevant to minority populations, because stress induced by environmental, social, and discriminatory influences can be significant. One study is investigating whether the effect of acute and chronic exposure to established risk factors (depressive symptoms, major life events, and lack of social support) for CVD during a 5-year period is related to a 2-year increase of subclinical CVD in a sample of women undergoing menopause. Fifty-four percent of participants will be black. Another study is examining whether racial disparity in AMI-recovery outcomes in older individuals is partially due to stress related to age and racial stigmas. Fifty percent of participants will be black. Other studies are seeking to clarify the physiologic pathways through which stress affects cardiovascular outcomes—including hemodynamic, sympathetic nervous system and neuroendocrine mechanisms, and inflammatory processes. The goal is to determine whether stress management interventions can alter these physiological indicators and improve cardiovascular outcomes.

Treatment

The Institute supports several investigator-initiated studies to develop and evaluate interventions to improve cardiovascular outcomes. One study is seeking to determine the efficacy of a mindfulness-based personalized health plan intervention on reducing inflammation, a

significant predictor of future CVD, via reductions in traditional risk factors, selected psychosocial attributes, and stress-reactivity among at-risk adults.

Approximately 30 percent of participants will be from minority populations. Another study is evaluating the effectiveness of stress management training combined with exercise-based cardiac rehabilitation as a way to reduce stress in vulnerable cardiac patients. Approximately 25 percent of participants are black. A third study is developing and evaluating an intervention among racial minorities that reduces the impact of bias by reducing stereotypic perceptions that render patients and providers less able to communicate effectively and impair the effect of the visit to improve patient self-management behavior. Additional studies are investigating whether stress management in a high-risk population of blacks with hypertension can influence cardiovascular risk factors.

Diabetes

Etiology

Diabetes mellitus is a strong risk factor for CVD. Individuals with type 2 diabetes are 2 to 4 times more likely to be at risk for CHD than the general population. Using GWAS, investigators have identified several genetic variants for CHD in the general population. An investigator-initiated study is seeking to identify new genetic variants for excessive risk of CHD in diabetic patients, assess the genetic effects on intermediate biochemical changes, and examine gene–environment interactions. One of the data sources will come from the Costa Rican Diabetes–CHD case-control study.

Treatment and Prevention

The NHLBI supports clinical trials to determine the benefits of various strategies to reduce CVD among patients with diabetes or treat patients with coronary artery disease and diabetes:

- ACCORDION (see Chapter 10): To obtain long-term (10 years on average) data on ACCORD participants. More than 33 percent of participants are from minority populations.
- Diabetes Prevention Program Outcomes Study—Phase II (see Chapter 11): To determine the efficacy of treatments to prevent or delay the development of type 2 diabetes in a population at high risk due to the presence of impaired glucose tolerance. Forty-five percent of participants will be from minority populations.

- Look AHEAD (see Chapter 11): To test the effectiveness of a lifestyle intervention in obese participants with type 2 diabetes over a long-term period. One of the clinical centers will direct its interventions toward American Indians.

Education

The Institute has prepared the following publications on diabetes for minorities:

- *Healthy Hearts, Healthy Homes—Protect Your Heart Against Diabetes* in English and Spanish
- *Healthy Heart, Healthy Family—Protect Your Heart: Prevent and Control Diabetes* in Tagalog and English

HIV-Related Cardiovascular Diseases

Use of multidrug antiretroviral therapy has improved life expectancy of HIV-infected individuals to the point that HIV/AIDS is now a chronic condition for many patients. As a result, CVD is now causing an increasing proportion of the deaths of HIV-infected individuals.

The NHLBI initiated research on the development of CVD in HIV-infected patients and potential management strategies:

- Mechanisms and Management of Cardiovascular and Metabolic Complications of HIV/AIDS: To elucidate the underlying mechanisms of metabolic and anthropometric abnormalities found in HIV-infected patients and relate the mechanisms to CVD risk; evaluate biomarkers and imaging methods for assessing coronary artery disease and risk in HIV patients; and identify strategies and approaches that reduce cardiovascular risk and optimize the medical management of HIV infection. Approximately 30–70 percent of participants are expected to come from racial and ethnic minority populations.

Lung Diseases

The NHLBI supports research on several lung diseases—such as asthma, sarcoidosis, TB, and HIV-related lung diseases—that disproportionately affect minorities. The following section provides examples of research to address health disparities in lung diseases; selected sleep disorders are also included.

Asthma

Asthma is a chronic lung disease that inflames and narrows the airways. It affects people of all ages and most often starts in childhood. In the United States, more than 24 million people are known to have asthma, and more than 7 million of them are children. Prevalence rates are especially high in blacks and Puerto Ricans.

Etiology

The NHLBI has initiated additional studies to improve understanding of the etiology and pathophysiology of asthma:

- Severe Asthma Research Program (See Chapter 9): To define severe asthma at the molecular and cellular levels over time in order to gain an understanding of the pathogenesis of the disease and provide a basis for design of mechanism-based diagnostic, prognostic, and treatment strategies for children and adults with severe asthma. Several of the projects have strong participation from minority populations.
- Airway Smooth Muscle Function and Targeted Therapeutics in Human Asthma: To investigate the complex role that airway smooth muscle plays in the development of asthma and to identify innovative therapeutic targets. Two projects expect 30–50 percent of participants to be from minority populations.

The Institute also supports investigator-initiated projects on the etiology and pathophysiology of asthma. Several projects focus on the role of genetics in the development of asthma. One study will identify genetic, biologic, and immunologic characteristics and environmental exposures that interact in children who experience severe bronchiolitis caused by the respiratory syncytial virus early in life and subsequently determine their role in the development of asthma, airway hyperactivity, and allergy. Forty percent of participants will be black. Another study will improve understanding of the etiology of asthma and the response to asthma drugs by performing GWAS to determine genetic factors that are associated with asthma, asthma severity, and bronchodilator response in two Latino subgroups. Additional studies will identify the genes and structural genetic variations that contribute to childhood asthma in the major racial and ethnic groups in the United States and in Hispanic populations outside the United States.

Investigators are also interested in the role of stress in the development of asthma. One study is using a systems biology approach to determine multiple biologic pathways by which stress can contribute to asthma. Scientists are investigating whether maternal stress immediately before or after the birth of a child can adversely affect the child's risk for wheezing and impair lung function later in childhood. Scientists hypothesize that multiple stressors that are prevalent in disadvantaged populations can cumulatively influence immune system development and airway inflammation in early life, thus making the populations more susceptible to other environmental factors and genetic risk factors explaining, in part, observed asthma disparities associated with SES and race and ethnicity.

A group of scientists has speculated that vitamin D deficiency in pregnant mothers may lead to faulty immune system development in the neonate, predisposing the neonate to asthma and allergy. To test the hypothesis, scientists will determine whether supplemental vitamin D intake to increase the level of vitamin D in pregnant women will prevent asthma and allergy in their children at age 3 years. More than 50 percent of participants are from racial and ethnic minority populations.

Research findings suggest that obesity and asthma are complex disorders that may have shared genetic determinants. An investigator-initiated study is seeking to identify single nucleotide polymorphisms (SNPs) that are jointly associated with asthma and obesity, using data from the CAMP study, and subsequently validate the SNPs in three independent and diverse (Hispanic, black, and white) cohorts.

Treatment and Control

The Institute has initiated research to identify optimal strategies for treatment and management of asthma. Because the burden of asthma disproportionately affects minority children, it is important for them to be well represented in clinical trials.

- AsthmaNet (see Chapter 11): To develop and conduct multiple clinical trials to identify optimal therapies for a variety of asthma phenotypes, genotypes, and racial and ethnic backgrounds in pediatric and adult populations. Approximately 30 percent of participants will be from minority populations.

The Institute also supports two investigator-initiated clinical trials to evaluate treatment strategies in asthma patients. STAN (see Chapter 9) is seeking to determine whether treatment of chronic rhinitis and sinusitis with nasal steroids will improve the control of asthma. Thirty-three percent of participants are expected to be from minority populations. SOYA (see Chapter 9) expects to determine whether supplementation with soy isoflavones among persons with poorly controlled asthma improves both lung function and markers of airway inflammation. Thirty-three percent of participants are expected to be from minority populations.

Quality of life (QOL) measures can assist health care providers in the treatment of asthma. Based on recommendations by the Asthma Related Quality of Life Subcommittee of the Asthma Outcomes Workshop held in 2010, new instruments are being developed and tested to measure more comprehensively the effects of asthma on QOL. Approximately 30 percent of participants will be from racial and ethnic minority populations.

One way to reduce asthma health disparities and reduce harm is to begin treatment in early childhood. Studies have shown that asthma education programs can improve overall management of asthma in preschool children. A study in Baltimore, Maryland, is partnering with Head Start to compare the efficacy of early intervention plus asthma education versus asthma education alone in reducing asthma morbidity. Nearly all participants will be black.

The Institute is supporting several investigator-initiated studies focusing on finding effective treatment for various populations. One study in whites, blacks, and Hispanics is creating an asthma self-management skills training program for children and their parents. The program is being created for Web, CD-ROM, and DVD formats and focuses on improving understanding of asthma, preventing asthma attacks, and creating an asthma management plan. Another study is seeking to improve health among urban black adolescents with asthma by using peer support—enhanced by a culturally sensitive, technology-based MP3 player platform—to increase adherence to daily controller medications. A third study is evaluating a Web-based intervention program that addresses asthma management and avoidance of tobacco use in rural, black adolescent boys. A fourth study is

assessing the effects of heart rate variability biofeedback on airway reactivity and inflammation to determine whether biofeedback can be useful for treating asthma; approximately 35 percent of participants will be from minority groups.

Many individuals with asthma have poor disease management. An investigator-initiated study is determining whether an intervention designed to stimulate communication between caregivers and clinicians and to contain individualized guideline-based recommendations for care, administered in urban primary care offices, reduces morbidity among urban children with asthma; 50 percent of participants are black. Another study is addressing asthma disparities that persist among high-risk children who live in rural, medically underserved areas by testing a school-based telemedicine approach that will deliver asthma education to rural children with asthma, their caregivers, and school nurses. The approach also prompts the children's primary care physicians with treatment recommendations. Approximately 75 percent of participants will be black.

A study in high-risk black adolescents with moderate to severe asthma is testing the effectiveness of an intensive home- and community-based psychotherapy intervention to improve asthma management and reduce the number of hospitalizations and visits to emergency departments. Black women will be the target of a study to improve asthma management by using a highly tailored telephone counseling approach to foster a partnership between women and a clinician. A study of obese adults, for which approximately 40 percent will be from minority populations, will evaluate the efficacy of an evidence-based lifestyle weight loss intervention to control asthma.

Investigators are also interested in evaluating whether cultural competency training for primary care physicians who primarily serve black or Hispanic communities will improve the asthma outcomes of their patients.

Symptoms of depression are commonly found in patients with asthma. A pilot study using an antidepressant to treat outpatients with major depression and asthma showed that patients who received treatment experience greater sustained remission of depressive symptoms and require significantly less oral corticosteroids for asthma management than the group who

received placebo. Based on these findings, scientists are implementing a definitive antidepressant study in patients with asthma and major depression. Most participants will be black or Hispanic.

Education

The Institute is supporting several education activities through the National Asthma Control Initiative (NACI), which was developed by the National Asthma Education and Prevention Program (NAEPP) and is coordinated by the NHLBI. The NACI seeks to accelerate the adoption of six priority action messages from the latest asthma guidelines to improve clinical practices and outcomes. NACI Demonstration Projects, Strategic Partner Projects, and Clinical Champions Projects are implementing strategies in racial and ethnic minority communities to address asthma health disparities in diverse populations.

The Institute has developed materials on asthma treatment and control:

- *So You Have Asthma*
- *My Asthma Wallet Card*
- *Asthma Action Plan*
- *Asthma and Physical Activity in the School*

The Institute also disseminates clinical practice guidelines for the treatment and monitoring of asthma, patient education materials, and information on environmental control of allergens in the United States and throughout the world.

Chronic Obstructive Pulmonary Disease

COPD—a disease in which the lungs are damaged, making breathing difficult—is the third leading cause of death in the United States. It is responsible for more than 500,000 hospitalizations and more than 130,000 deaths in the United States each year.

Etiology

The NHLBI is supporting a large, investigator-initiated study of genetic factors that determine the risk of developing COPD or that influence the type and extent of damage done to the body by the disease. The COPDGene™ study has enrolled more than 3,000 blacks with a substantial history of cigarette smoking, obtained extensive baseline clinical and phenotypic data regarding the individuals, and compared the severity and character

of COPD in the subjects to analyses of their DNA. Genome-wide genetic assays will be performed on a substantial fraction of this cohort.

Although COPD is less common among blacks than among whites, it is nevertheless the seventh leading cause of death among blacks. Any disparity, whether higher or lower in the minority group, may reflect racial differences in the biology of the disease that would require use of different treatments or drugs for optimal disease management. If the genes found to be determinants of COPD risk differ in blacks and whites, this will provide clues to how the roles of specific pathogenetic pathways of COPD differ among races.

Treatment and Control

The Institute has established a research network to determine effective disease management approaches for individuals with moderate-to-severe COPD:

- COPD Clinical Research Network (see Chapter 11): To perform collaborative, therapeutic interventional trials of medications, devices, and disease management strategies in individuals with moderate-to-severe COPD. In addition to evaluating treatment efficacy, network studies include examinations of genetic factors, biomarkers, or genomic/proteomic profiles that may identify patients who are more or less likely to benefit from various treatments.
- Twenty-two percent of participants are minorities.

In addition, in collaboration with the Centers for Medicare and Medicaid Services, the Institute is sponsoring a clinical trial of supplemental oxygen treatment in COPD patients:

- LOTT (see Chapter 11): To test whether long-term oxygen therapy can reduce disability and prolong life in COPD patients who have moderate resting hypoxemia or severe hypoxemia during exercise.

Education

The NHLBI has developed a number of outreach activities associated with COPD. Several publications and Web-based products have been developed and distributed for health professionals, patients, and the public. Some examples include:

- COPD education Web site: <http://www.nhlbi.nih.gov/health/public/lung/copd/index.htm>
- COPD Learn More Breathe Better Campaign

Sarcoidosis

Sarcoidosis is an inflammatory disease of unknown etiology characterized by persistent granulomas with damage to surrounding tissue. The Institute supports research into the basic mechanisms of sarcoidosis and new and improved treatments for it.

Sarcoidosis occurs more frequently and with more severity in blacks than in whites, suggesting the presence of genetic determinants to disease predisposition. To increase understanding of the disease, researchers are seeking to identify genes of African ancestry that play a significant role in its etiology and pathogenesis. Other possible causes of sarcoidosis are also being investigated.

A mentored research project conducted within the Black Women's Health Study is investigating potential risk factors for sarcoidosis.

Diagnostic tools and treatment approaches for sarcoidosis are lacking. A new study supported by the Institute-initiated CADET Program (see Chapter 9) is seeking to develop a skin test for diagnosing sarcoidosis and to establish an approach for treating sarcoidosis with oral vaccines.

Sleep Disorders

Etiology

Sleep-disordered breathing (SDB), a condition characterized by repetitive interruption in breathing, is a common disorder that disproportionately affects blacks. It is associated with an increased risk of CVD, including hypertension and stroke, and is particularly prevalent in patients with heart failure. Ongoing programs are assessing the interrelationship between sleep disorders and heart failure and the mechanisms leading to cardiovascular stress when the two intersect.

The Institute also supports investigator-initiated projects to elucidate cardiovascular and other health consequences of SDB, sleep deprivation, and shift work in various community settings. One study is testing associations between indices of sleep apnea and the quality, duration, and timing of sleep with indices of cardiovascular risk and disease to address the role of sleep disorders in CVD development in minority populations.

Approximately 60 percent of participants are expected to be from racial and ethnic minority groups. Another study

is measuring the association between sleep duration and nervous system, cardiovascular, and metabolic health and dysfunction and overall body inflammation; approximately 70 percent of participants are expected to be from minority populations. A third study is investigating sleep apnea in stroke patients, collecting and analyzing data on prevalence, relationship to ethnicity, and the association between stroke type and outcomes; approximately 65 percent of participants are expected to be from racial and ethnic minority groups.

In an ancillary study to the Hispanic Community Health Study, investigators are collecting measurements of sleep patterns from several Hispanic subgroups. Investigators will analyze the prevalence of altered sleep patterns and define the importance of psychosocial factors (e.g., home- and work-related factors and mood, as well as cultural factors) in predicting abnormal sleep patterns. They will also evaluate associations between poor sleep and cardiovascular health outcomes, including obesity, hypertension, diabetes, and heart disease.

Treatment and Control

The Institute initiated a pilot study to guide design of phase III clinical trials to test whether treatment of obstructive sleep apnea (OSA) with positive airway pressure (PAP) reduces CVD risk:

- Reducing Cardiovascular Disease Risk Through Treatment of Obstructive Sleep Apnea: To evaluate the treatment of OSA using PAP. The study will provide information about the feasibility of long-term (12 to 18 months) PAP treatment of OSA in patients who are at risk of CVD. The study will also provide data about the effects of PAP on surrogate markers of cardiovascular risk. Approximately 40 percent of participants will be from minority populations.

A new study supported by the Institute-initiated CADET Program is seeking to develop reliable urinary biomarker measures that can be used to screen and diagnose OSA in children, thereby facilitating timely treatment and prevention of OSA-associated morbidities. Approximately 80 percent of participants are expected to be from minority populations.

Investigator-initiated research will also assess treatment strategies in minorities. One study is developing in-home personalized sleep plans to improve nightly

sleep duration, neurocognitive function, and behavioral disorders in lower income minority children (ages 5 and 6 years). Another study will measure effects of continuous compliance to PAP therapy on neurocognitive function (including academic achievement, attention, working memory, decision making, and mental flexibility) in children. Parental reports of child behaviors and sleep patterns will also be assessed. Approximately 60 percent of participants will be from minority populations. A third study will advance understanding of endothelial activation processes in OSA patients and may enable early identification of OSA patients who are at risk for vascular diseases. Results may also provide the basis for developing new therapeutic strategies for preventing or reversing vascular risk in OSA patients. Approximately 90 percent of participants are expected to be from minority populations.

HIV-Related Lung Diseases

HIV infection disproportionately affects minority populations in the United States and due to multidrug antiretroviral therapy, has become a chronic condition for many patients. Among them, HIV-associated lung complications are frequent causes of illness and death. But the long-term consequences of HIV infection and HIV-associated lung infections and complications are unknown. Little is known about drug-resistant *Pneumocystis*, the prevalence and pathogenesis of HIV-associated COPD, HIV-associated pulmonary hypertension, and immune reconstitution syndromes. In developing countries where millions of people are infected with HIV, many have serious or fatal lung complications, including TB and bacterial pneumonias, that have never been characterized well. Lung and cardiovascular diseases, usually more prevalent in older populations, are having an increasing effect on HIV-infected populations in the United States and other industrialized nations where the average age of HIV-infected individuals has increased due to effective antiretroviral therapy. The effect is exacerbated because lung and cardiovascular complications are occurring in relatively young age groups in HIV-infected populations.

Etiology

In addition to supporting investigator-initiated research on the etiology and pathogenesis of HIV-associated lung diseases, the Institute has initiated research to understand their causes and impact and to

identify potential therapeutic targets and preventive strategies:

- Longitudinal Studies of HIV-Associated Lung Infections and Complications: To accelerate research on lung complications associated with HIV-infection by characterizing lung infections, other HIV-associated lung complications, and their consequences in longitudinal studies in existing HIV-infected cohorts and other established groups of patients who are HIV-infected. Depending on the center, participation from minority populations ranges from approximately 40 to 100 percent.
- Microbiome of the Lung and Respiratory Tract in HIV-Infected Individuals and HIV-Uninfected Controls (see Chapter 9): To characterize the microbiome of the lung alone or in combination with the upper airways in HIV-infected individuals and matched HIV-uninfected controls using molecular techniques to identify bacteria and if possible other organisms (e.g., viruses, cell-wall deficient organisms, protozoa, and fungi). Enrollment from minority populations is expected to range from 35 to 76 percent.

Tuberculosis

TB is a common and often deadly infectious disease caused by *Mycobacterium tuberculosis*. In the United States, an estimated 10–15 million people are infected with the TB bacterium. Although the majority of those infected remain healthy (latent TB infection), a small percentage develops active tuberculosis. The rates of both TB infection and active TB among minorities and in the foreign-born remain high. In 2008, according to the CDC, more than 58 percent of all active TB cases in the United States were among the foreign-born. Racial disparity in TB rates was greatest for U.S.-born blacks, whose rate was 7-times higher than the rate for U.S.-born whites.

Etiology

The immune response to TB infection is complex and involves the formation of granulomas in the lungs of infected individuals. In 2010, the Institute began a systems biology approach to identify the mechanisms of TB latency and reactivation.

- Systems Biology Approach to the Mechanisms of TB Latency and Reactivation: To investigate mechanisms of latency and reactivation of TB in the host

using integrated systems biology approaches. A collaborative program consisting of five Tuberculosis Systems Biology Centers and a Data Coordinating Center will integrate data from humans and animal models with computational and mathematical models in a comprehensive systems biology approach to increase understanding of latent TB and the factors that lead to its reactivation. Depending on the center, minority enrollment in the U.S. populations under study is expected to range from approximately 30 to 66 percent. Several of the grants in this program will study international populations in Africa and Asia.

The Institute also supports investigator-initiated research that characterizes genes associated with TB susceptibility; investigates host lung defenses, including immune responses to infection; and examines the impact of TB on HIV disease. A genetics study will seek to fine-map chromosomal regions that have been linked to resistance to TB. The study also plans to analyze innate immune responses and model genetic predictors of resistance using data from a long-term household contact study conducted in Uganda.

Treatment and Control

The NHLBI supports a number of investigator-initiated studies on understanding the relationship between the immune system and TB. Most of the studies are being conducted among patients from minority populations. One study is seeking new approaches to diagnosing and treating active TB in sub-Saharan Africa and other parts of the world. The study may also help to identify new markers that can predict response to TB therapy. Another study is determining whether different strains of *Mycobacterium tuberculosis* cause different responses in individuals from various racial and ethnic groups. Results of this study may increase understanding about tailoring vaccines to specific populations. A third study is testing the efficacy of daily vitamin D supplementation, added to multidrug therapy, to improve antimicrobial immune response to TB infection in residents of Mongolia.

Blood Diseases

The NHLBI supports basic, translational, and clinical research on SCD and thalassemia (Cooley's anemia) with the goal of curing the disorders and improving patient care. The Institute also supports a deep vein thrombosis and venous disease program and programs in transfusion

medicine, blood banking and blood products safety, and such cellular therapies as bone marrow transplantation.

Sickle Cell Disease

Basic and Translational Research

SCD, the most common inherited blood disorder in the United States, affects an estimated 70,000 to 100,000 Americans, most of whom are black. SCD occurs in about 1 in 500 blacks and 1 in 36,000 Hispanics. The disease is characterized by anemia, severe infections, acute and chronic pain, and organ damage. SCD, the first molecular disease described, was shown to be due to a single amino acid substitution on the beta chain of hemoglobin.

Since 1972, the NHLBI has supported an extensive research program to improve understanding of the pathophysiology of SCD, identify better approaches for its diagnosis and treatment, and prevent complications.

Basic and translational research currently focuses on genetic influences on disease manifestations, regulation of hemoglobin synthesis, discovery of drugs to increase fetal hemoglobin production, transplantation of blood-forming stem cells, gene therapy, pain research, and development of animal models for preclinical studies. Institute-initiated programs include:

- BTRP (see Chapter 9): To conduct comprehensive research, training, and education efforts related to SCD. The BTRP seeks to improve understanding of SCD pathophysiology and develop cures or improved medical management of the disease.
- Ancillary Studies in Clinical Trials: To conduct time-sensitive ancillary studies in conjunction with ongoing Phase II-III clinical trials or network clinical trials related to heart, lung, and blood diseases and sleep disorders. One study is using proteomic approaches to identify biomarkers of early cerebral ischemia in children with SCD. Identifying such circulating biomarkers may allow earlier therapeutic intervention in these children.
- Exploratory Studies in the Neurobiology of Pain in Sickle Cell Disease: To conduct basic and translational research on the neurobiology of pain in SCD and to develop effective pharmacologic treatments.

Investigator-initiated studies are exploring new methods to reactivate fetal hemoglobin (HbF) expression for

the treatment of SCD and thalassemia; elucidating the mechanism of hypercoagulation in SCD; examining the relationships between analgesia, biomarkers, and pain levels in adults with SCD; and investigating the mechanisms underlying pain in SCD in order to develop more effective pain therapies.

Gene therapy is another area of focus. One project is exploring lentiviral gene therapy for SCD, and another is studying the application of induced pluripotent stem cell technology to replace the defective sickle beta-globin gene with a normal gene in a SCD mouse model. Researchers are seeking to translate their results to human cells that will become the foundation for future clinical trials.

Through the SBIR and the STTR programs, the NHLBI supports translational projects to develop therapeutics and tools that can be used to treat patients with SCD and other hemoglobinopathies. Recent projects are investigating a safer and more affordable method to correct autologous hematopoietic stem cells in SCD and other genetic disorders; identifying antisickling compounds in a mixture of botanicals that can provide leads for developing new synthetic antisickling compounds; evaluating agents to increase hemoglobin F expression; testing a humanized monoclonal antibody drug for treatment of vaso-occlusive crisis; and advancing use of small molecule organic catalysts to create a new class of drugs to prevent sickle cell hemoglobin polymerization. The NHLBI also supports a project to develop a Web interface tool that can be used to improve the quality of care received by SCD patients during hospital emergency room visits.

Clinical Research

The NHLBI is committed to finding improved treatments and ultimately a cure for SCD and other hemoglobinopathies. Institute-initiated studies have begun to yield therapies that will alleviate the symptoms of sickle cell anemia and procedures that should ultimately cure the disease.

- BABY HUG (see Chapter 11): To assess the effectiveness of hydroxyurea during a 2-year treatment period in preventing onset of chronic organ damage in young black children who have sickle cell anemia. Although the primary goals of preserving spleen and renal function were not achieved, data show markedly reduced numbers of vaso-occlusive

events and improved hematologic counts. An observational study is following the cohort to learn more about long-term effects of hydroxyurea.

- Sildenafil for Sickle Cell Disease-Associated Pulmonary Hypertension (walkPHaSST) Follow-Up (see Chapter 11): To use datasets and biorepository specimens from the closed trial to assess the effects of genetic factors on hematologic, cardiopulmonary, and pain phenotypes in individuals with hemoglobin SS disease and to explore phenotypic variations in individuals with hemoglobin SC disease.
- Planning Grants for Clinical Trials in Hemoglobinopathies: To support pilot studies to obtain data that are critical for the design of robust clinical trials, especially Phase II and III clinical trials in the major hemoglobinopathies, SCD, and the thalassemias.
- Ancillary Studies in Clinical Trials: To conduct time-sensitive ancillary studies in conjunction with ongoing clinical trials and other large clinical studies that are related to heart, lung, and blood diseases and sleep disorders. One study is assessing the treatment response of children who are experiencing acute sickle cell pain episodes to determine the effectiveness of magnesium therapy (parent trial).

The Institute supports research to assist hematologists in their ability to assess clinical outcomes:

- Adult Sickle Cell Quality of Life Measurement Information System (see Chapter 10): To develop, validate, and disseminate a psychometrically sound and clinically useful instrument to assess health-related quality of life among adults with SCD. This system is fully integrated into an NIH Roadmap Patient-Reported Outcomes Measurement Information System.

The Institute is supporting two investigator-initiated clinical trials for children with SCD. One trial in children with abnormal transcranial Doppler (TCD) velocities is comparing standard therapy (transfusions) with alternative therapy (hydroxyurea) for maintenance of TCD velocities. The other is seeking to determine the effect of hydroxyurea treatment on the cumulative incidence of conversion from conditional to abnormal TCD velocities.

The Institute is also supporting a follow-up study on cohorts of children from the Stroke Prevention in Sickle Cell Anemia (STOP) and STOP II trials. Although the trials had major implications for managing SCD in children, the real-world effects of the tested approach have

yet to be determined. The current study uses a retrospective chart review of databases that contain trial and post-trial data to determine the effectiveness of chronic transfusion therapy and TCD screening in actual clinical practice rather than in just a randomized clinical trial. The study will gather information to support the development of treatment guidelines and improve the care of individuals with SCD.

Education

The NHLBI has developed the Sickle Cell Disease Information Center (<http://www.nhlbi.nih.gov/new/sicklecell.htm>), a Web site that contains information for the public and health professionals.

Thalassemia

Thalassemia is an inherited disorder in which red blood cells with abnormal forms of hemoglobin are produced. The disorder, which results in excessive destruction of red blood cells and anemia, affects primarily people of African, Asiatic Indian, Chinese, Mediterranean, and Southeast Asian origin.

Institute-initiated activities include:

- RuSH (See Chapter 10): To test, in a pilot project, the feasibility of developing a national data system that will enable investigators to estimate the number of people who have SCD, thalassemias, and hemoglobinopathies and to describe their sociodemographic characteristics.
- Innovators in Hemoglobinopathies Academic Career Development Award: To advance the development of clinician scientists who have implemented innovative programs for patients with SCD or thalassemia but who have not been on a research track.

An investigator-initiated study is also examining hematopoietic transplantation and gene therapy approaches to cure thalassemia.

Deep Vein Thrombosis and Venous Disease

Deep vein thrombosis (DVT) is a serious condition that can cause significant disability and death if not promptly diagnosed and effectively treated.

Approximately 2–3 million individuals in the United States develop venous thromboembolism each year, and of them, 60,000 die primarily of pulmonary embolism. The Institute initiated a program to improve understanding of deep vein thrombosis and venous disease.

- Deep Vein Thrombosis and Venous Disease: To improve diagnosis, therapy, and prevention of venous thrombotic diseases in order to enhance patient health and well-being. Several of the projects have substantial participation from minority populations.

Warfarin sodium, an anticoagulant drug, is prescribed to millions of patients each year to prevent thromboembolism. However, warfarin has an unusually narrow therapeutic range and is difficult to dose properly. To address this problem, the Institute is supporting several investigator-initiated studies to determine environmental and genetic factors that influence patient response to warfarin. Fifty to 60 percent of participants are from racial and ethnic minority populations.

Transfusion Medicine and Cellular Therapeutics

The NHLBI supports research on the use, safety, and availability of blood and blood components for transfusion and cellular therapies that are seeking to reach minority populations:

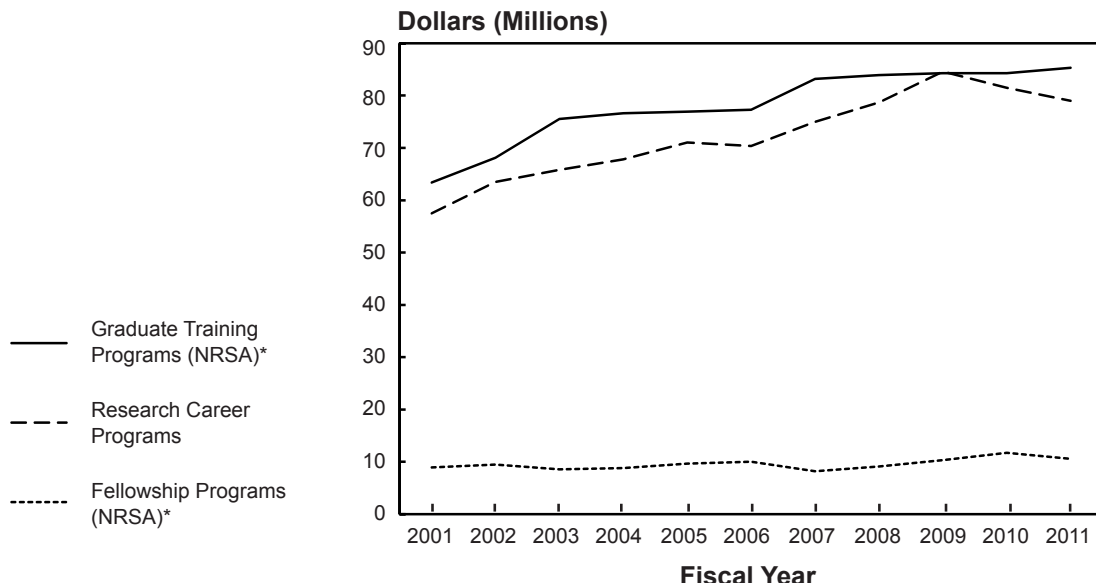
- Blood and Marrow Transplant Clinical Research Network (BMT CRN) (see Chapter 11): In collaboration with the NCI, to perform clinical trials that advance hematopoietic stem cell transplantation.

To reach minority populations, the BMT CRN supports bilingual transplant center personnel and provides public Web pages and educational materials. In addition, the Network is working with the National Marrow Donor Program to develop strategies and implement procedures to enhance enrollment of patients from minority groups.

- The Sickle Cell Unrelated Transplant Trial: To assess unrelated donor marrow and umbilical cord blood transplantation for severe SCD. The trial, supported by the BMT CRN and the Sickle Cell Disease Clinical Research Network, is the first Phase II study to assess the promise of this therapy as a curative option for patients who are severely affected by SCD.
- PACT (see Chapter 10): To manufacture clinical grade mesenchymal stem cells (MSC) for a clinical trial that will examine the safety of infusing MSC during the transplantation process. Data from other studies demonstrate that MSC can facilitate the acceptance of the foreign cells being infused into a patient and help to repair blood vessels that have been damaged during the circulation of “sickled” red blood cells.

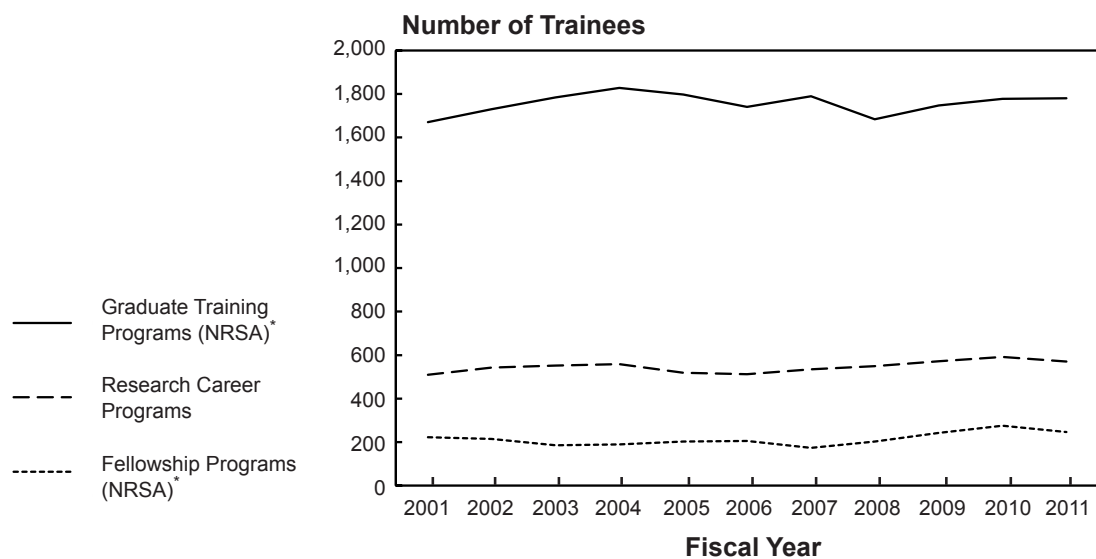
13. Research Training and Career Development Programs

NHLBI Research Training and Career Development Obligations: Fiscal Years 2001–2011



* National Research Service Awards (NRSA).

NHLBI Full-Time Training Positions: Fiscal Years 2001–2011



* National Research Service Awards (NRSA).

Note: Numbers of awards and trainees may not agree with other tables due to the method of counting supplements.

Training Awards, Full-Time Training Positions, and Obligations by Activity: Fiscal Year 2011

	Number of Awards Obligated	Trainees (Full-time Training Positions)	Direct Cost	Indirect Cost	Total Cost	Percent of Total NHLBI Training Program Dollars
Fellowship Programs						
Individual Predoctoral NRSA for M.D./Ph.D. (F30)	88	88	\$ 3,297,797	\$ —	\$ 3,297,797	3.4%
Predoctoral Individual NRSA (F31)	46	46	1,561,809	—	1,561,809	1.6
Postdoctoral Individual NRSA (F32)	112	112	5,766,781	—	5,766,781	6.0
Senior Fellowships NRSA (F33)	—	—	—	—	—	—
Subtotal, Fellowships	246	246	10,626,387	—	10,626,387	7.7
Graduate Training Programs						
Institutional NRSA (T32)	237	1,667	76,108,416	6,428,051	82,536,467*	87.1
Minority Institutional NRSA (T32)	4	27	885,279	63,229	948,508	1.0
Off-Quarter Professional Student Training NRSA (T34, T35)	16	86	1,746,722	157,501	1,904,223	2.0
Short-Term Training for Minority Students (T35M)	—	—	—	—	—	—
Subtotal, Graduate Training Programs	257	1,780	78,740,417	6,648,781	85,389,198	90.0
Total, Training Programs	503	2,026	\$89,366,804	\$6,648,781	\$96,015,585	100.0%

* Excludes assessment of \$1,982,000.

History of Training Obligations by Activity: Fiscal Years 2001–2011

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Fellowship Programs											
Individual Predoctoral NRSA for M.D./Ph.D. (F30)	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ 641	\$ 2,191	\$ 3,024	\$ 3,297
Predoctoral Individual NRSA (F31)	264	478	563	549	794	1,202	1,509	1,888	2,009	2,094	1,562
Postdoctoral Individual NRSA (F32)	8,515	8,887	7,868	8,128	8,813	8,790	6,684	6,487	6,012	6,559	5,767
Senior Fellowships NRSA (F33)	147	84	112	144	58	53	—	59	118	—	—
Subtotal, Fellowships	8,926	9,449	8,543	8,821	9,665	10,045	8,193	9,075	10,330	11,677	10,626
Graduate Training Programs											
Institutional NRSA (T32)	58,516 ^A	62,999 ^B	69,951 ^C	71,229 ^D	70,524 ^E	71,831 ^F	78,343 ^G	80,373 ^H	81,453 ^I	81,319 ^J	82,536 ^K
Minority Institutional NRSA (T32)	996	1,092	1,006	734	1,184	743	780	688	349	1,050	949
Off-Quarter Professional Student Training NRSA (T34, T35)	1,974	1,987	1,975	1,993	2,233	2,215	2,411	2,021	2,202	1,941	1,904
MARC (T36)	5	—	—	—	—	—	—	—	—	—	—
Short-Term Training for Minority Students (T35M)	1,877	2,057	2,594	2,671	2,976	2,527	1,673	804	283	—	—
Subtotal, Training Grants	63,368	68,135	75,526	76,627	76,917	77,316	83,207	83,886	84,287	84,310	85,389
Total, Training Programs	\$72,294^A	\$77,584^B	\$84,069^C	\$85,448^D	\$86,582^E	\$87,361^F	\$91,400^G	\$92,961^H	\$94,617^I	\$95,987^J	\$96,015^K

A Excludes Assessment of \$1,424,000.

B Excludes Assessment of \$1,584,000.

C Excludes Assessment of \$1,716,000.

D Excludes Assessment of \$1,744,000.

E Excludes Assessment of \$1,764,000.

F Excludes Assessment of \$1,818,000.

G Excludes Assessment of \$1,916,000.

H Excludes Assessment of \$1,912,000.

I Excludes Assessment of \$1,960,000.

J Excludes Assessment of \$1,976,000.

K Excludes Assessment of \$1,982,000.

Full-Time Training Positions by Activity: Fiscal Years 2001–2011

	Number of Positions										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Fellowship Programs											
Individual Predoctoral NRSA for M.D./Ph.D. (F30)	—	—	—	—	—	—	—	20	63	85	88
Predoctoral Individual NRSA (F31)	12	18	19	18	25	32	44	56	59	62	46
Postdoctoral Individual NRSA (F32)	208	194	164	168	176	171	130	125	118	128	112
Senior Fellowships NRSA (F33)	3	2	2	3	1	2	—	1	2	—	—
Subtotal, Fellowships	223	214	185	189	202	205	174	202	242	275	246
Graduate Training Programs											
Institutional NRSA (T32)	1,425	1,482	1,542	1,578	1,540	1,512	1,585	1,525	1,602	1,660	1,667
Minority Institutional NRSAs (T32)	43	39	42	32	35	26	23	18	19	26	27
Off-Quarter Professional Student Training NRSA (T34, T35)	109	179	93	99	95	104	105	93	102	91	86
Short-Term Training for Minority Students (T35M)	93	30	107	119	128	99	77	48	24	—	—
Subtotal, Training Grants	1,670	1,730	1,784	1,828	1,798	1,741	1,790	1,684	1,747	1,777	1,780
Total, Training Positions	1,893	1,944	1,969	2,017	2,000	1,946	1,964	1,886	1,989	2,052	2,026

NHLBI Research Career Programs: Fiscal Years 2001–2011

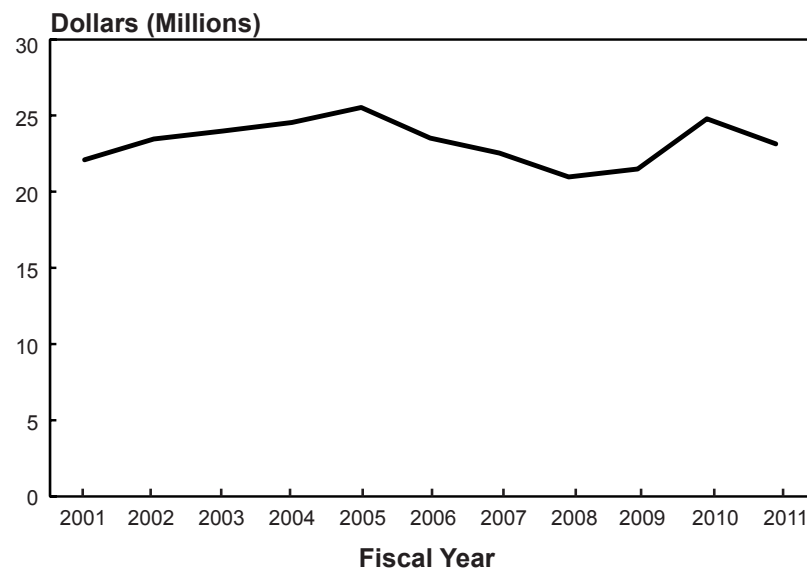
	Number of Awards										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Mentored Research Scientist Development Award for Minority Faculty (K01)	44	54	47	46	45	40	35	35	37	47	45
Minority Institution Faculty Mentored Research Scientist Development Award (K01)	9	2	7	6	4	4	5	7	5	6	5
Mentored Scientist Development Award in Research Ethics (K01)	—	—	2	2	3	3	3	1	1	—	—
Independent Scientist Award (K02)	34	33	32	31	32	24	25	22	19	19	21
Research Career Award (K06)	2	2	2	1	1	1	—	—	—	—	—
Tuberculosis Academic Award (K07)	5	—	—	—	—	—	—	—	—	—	—
Sleep Academic Award (K07)	12	8	—	—	—	—	—	—	—	—	—
Nutrition Academic Award (K07)	19	19	9	9	—	—	—	—	—	—	—
Pediatric Transfusion Medicine Academic Award (K07)	—	—	—	—	—	—	4	4	4	4	4
Cultural Competence and Health Disparities Academic Award (K07)	—	—	—	8	14	18	18	18	9	4	—
Innovators in Hemoglobinopathies Care Career Development Award (K07)	—	—	—	—	—	—	—	—	—	—	2
Clinical Investigator Development Award (K08)	241	236	240	229	239	226	214	210	232	218	210
Research Career Development Program in Vascular Medicine (K12)	—	—	—	—	—	2	7	7	7	—	3
Clinical Hematology Research Career Development Program (K12)	—	—	—	—	—	6	6	6	6	6	—
Genetics and Genomics of Lung Diseases Career Development Program (K12)	—	—	—	—	—	—	8	8	8	8	—
Clinical Research Career Development Programs in Emergency Medicine (K12)	—	—	—	—	—	—	—	—	—	—	6
Minority School Faculty Development Award (K14)	1	—	—	—	—	—	—	—	—	—	—
Career Enhancement Award for Stem Cell Research (K18)	—	—	1	5	3	2	4	6	3	4	5
NHLBI Career Transition Award (K22)	—	—	—	1	2	1	1	1	1	1	3
Mentored Patient-Oriented Research Career Development Award (K23)	58	90	110	122	127	122	120	133	149	160	170
Midcareer Investigator Award in Patient-Oriented Research (K24)	27	37	38	32	32	33	29	29	34	35	34
Mentored Quantitative Research Career Development Award (K25)	2	7	9	12	17	16	15	15	15	15	15
Clinical Research Curriculum Award (K30)	55	55	55	55	—*	14	16	—	—	—	—
Career Transition Award (K99)	—	—	—	—	—	—	24	47	42	64	47
Total, Research Career Programs	509	543	552	559	519	512	534	549	572	591	570

* In FY 2005, NHLBI relinquished management of the K30 program and as a result did not receive the grant count.

NHLBI Research Career Program Obligations: Fiscal Years 2001–2011

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Mentored Research Scientist Development Award for Minority Faculty (K01)	\$ 5,556	\$ 5,711	\$ 6,156	\$ 6,150	\$ 6,088	\$ 5,453	\$ 4,718	\$ 4,574	\$ 4,745	\$ 6,089	\$ 5,860
Minority Institution Faculty Mentored Research Scientist Development Award (K01)	1,143	1,703	991	867	588	567	698	949	663	804	668
Mentored Scientist Development Award in Research Ethics (K01)	—	—	255	253	355	358	357	102	164	62	126
Independent Scientist Award (K02)	3,202	3,130	3,099	3,079	3,218	2,421	2,511	2,184	1,880	1,847	2,076
Research Career Award (K06)	70	69	69	34	34	34	—	—	—	—	—
Tuberculosis Academic Award (K07)	396	—	—	—	—	—	—	—	—	—	—
Sleep Academic Award (K07)	1,081	722	—	—	—	—	—	—	—	—	—
Nutrition Academic Award (K07)	2,869	2,906	1,472	1,516	—	—	—	—	—	—	—
Pediatrics Transfusion Medicine Academic Award (K07)	—	—	—	—	—	—	486	486	486	486	486
Cultural Competence and Health Disparities Academic Award (K07)	—	—	—	925	1,620	2,109	2,232	2,197	1,138	562	—
Innovators in Hemoglobinopathies Care Career Development Award (K07)	—	—	—	—	—	—	—	—	—	—	574
Clinical Investigator Development Award (K08)	29,263	29,295	30,288	29,037	30,429	28,973	27,286	27,005	29,706	28,165	27,166
Research Career Development Program in Vascular Medicine (K12)	—	—	—	—	—	772	3,206	5,499	7,325	—	2,499
Clinical Hematology Research Career Development Program (K12)	—	—	—	—	—	2,360	2,367	2,364	2,375	2,371	—
Genetics and Genomics of Lung Diseases Career Development Program (K12)	—	—	—	—	—	—	3,154	3,190	3,190	3,194	—
Clinical Research Career Development Programs in Emergency Medicine (K12)	—	—	—	—	—	—	—	—	—	—	1,186
Minority School Faculty Development Award (K14)	98	—	—	—	—	—	—	—	—	—	—
Career Enhancement Award for Stem Cell Research (K18)	—	—	243	980	512	213	652	1,014	477	706	789
NHLBI Career Transition Award (K22)	—	—	—	185	364	178	160	162	162	162	699
Mentored Patient-Oriented Research Career Development Award (K23)	7,570	11,909	14,571	16,216	17,086	16,720	16,419	18,556	20,831	22,368	23,871
Midcareer Investigator Award in Patient-Oriented Research (K24)	2,877	4,058	4,368	3,815	3,929	4,315	4,037	4,161	5,078	5,942	5,851
Mentored Quantitative Research Career Development Award (K25)	272	921	1,195	1,622	2,206	2,184	2,077	2,082	1,996	2,134	2,110
Clinical Research Curriculum Award (K30)	3,073	3,090	3,110	3,115	4,589	3,708	2,520	—	—	—	—
Career Transition Award (K99)	—	—	—	—	—	—	2,074	4,190	4,430	6,652	5,129
Total, Research Career Program Obligations	\$57,470	\$63,514	\$65,817	\$67,794	\$71,018	\$70,365	\$74,954	\$78,715	\$84,646	\$81,544	\$79,090

NHLBI Minority Biomedical Research Training, Career Development, and Research Supplements Program Obligations: Fiscal Years 2001–2011



NHLBI Minority Biomedical Research Training, Career Development, and Research Supplements Program Obligations: Fiscal Years 2001–2011

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
MARC Summer Research Training Program	\$ 20	\$ 15	\$ 4	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —	\$ —
Mentored Research Scientist Development Award for Minority Faculty	5,556	5,711	6,156	6,150	6,088	5,453	4,718	4,574	4,745	6,089	5,860
MARC	5	—	—	—	—	—	—	—	—	—	—
Minority Biomedical Research Support (MBRS)	3,165	2,793	3,600	2,806	2,846	2,403	2,475	1,527	2,167	2,540	3,228
Minority Institution Faculty Mentored Research Scientist Development Award	1,143	1,703	991	867	588	567	698	949	663	804	668
Minority Institution Research Training Program	996	1,092	1,006	734	1,184	743	780	688	349	1,050	949
Minority Predoctoral Fellowship	264	278	308	374	545	1,012	1,115	1,728	1,979	2,064	1,562
Diversity Research Supplements Program	8,587	9,822	9,323	10,938	11,214	10,680	10,834	10,303	10,412	11,198	10,260
Minority School Faculty Development Award	98	—	—	—	—	—	—	—	—	—	—
Reentry Supplements	384	—	—	—	96	132	245	401	887	1,050	621
Short-Term Training for Minority Students	1,876	2,057	2,594	2,671	2,976	2,526	1,673	804	283	—	—
Total, Minority Programs	\$22,094	\$23,471	\$23,982	\$24,540	\$25,537	\$23,516	\$22,538	\$20,974	\$21,485	\$24,795	\$23,148

NHLBI Research Supplements Program by Award Type on Grants: Fiscal Years 2001–2011

	Number of Awards										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Diversity Supplements											
Investigator	33	46	47	35	29	27	31	25	22	17	10
Postdoctoral	41	33	38	37	52	49	43	42	45	50	49
Graduate	43	45	57	61	80	74	73	69	71	71	66
Undergraduate	12	17	18	17	12	11	16	17	18	13	9
High School	3	3	4	3	7	3	3	3	2	7	10
Post-Master/Post-Baccalaureate	—	2	8	17	16	11	4	9	21	21	33
Reentry Supplements	3	—	—	3	2	1	1	3	9	8	5
Disability Supplements	4	5	4	3	2	2	4	1	—	2	2
Total, Research Supplements Program	139	151	176	176	200	178	175	169	188	189	184

NHLBI Research Supplements Program by Award Type on Contracts: Fiscal Years 2001–2011

	Number of Awards										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Diversity Supplements											
Investigator	—	—	—	1	2	1	2	2	4	3	3
Postdoctoral	—	—	2	1	2	1	1	0	1	4	4
Graduate	—	—	2	7	7	2	5	2	5	5	6
Undergraduate	—	—	—	—	—	1	1	1	1	1	—
High School	—	—	1	—	—	—	—	—	—	—	—
Post-Master/Post-Baccalaureate	—	—	—	1	—	—	—	—	—	—	—
Total, Research Supplements Program	—	—	5	10	11	5	9	5	11	13	13

NHLBI Research Supplements Program Obligations by Award Type on Grants: Fiscal Years 2001–2011

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Diversity Supplements											
Investigator	\$3,430	\$ 5,046	\$3,844	\$ 4,256	\$ 3,552	\$ 3,343	\$ 3,719	\$ 3,285	\$ 2,679	\$ 2,183	\$ 1,601
Postdoctoral	3,086	2,554	2,655	2,713	3,432	3,542	3,284	3,074	3,284	3,928	3,595
Graduate	1,818	1,864	2,181	2,439	3,208	3,114	3,021	3,029	3,212	3,533	3,389
Undergraduate	235	260	301	282	179	178	350	424	386	240	151
High School	18	33	33	13	30	18	16	26	28	61	75
Post-Master/Post-Baccalaureate	—	65	309	597	618	352	156	367	823	1,076	1,284
Reentry Supplements	384	—	—	495	96	132	245	401	887	1,050	621
Disability Supplements	187	474	360	143	99	133	288	98	—	177	165
Total, Research Supplements Program	\$9,158	\$10,296	\$9,683	\$10,938	\$11,214	\$10,812	\$11,079	\$10,704	\$11,299	\$12,248	\$10,881

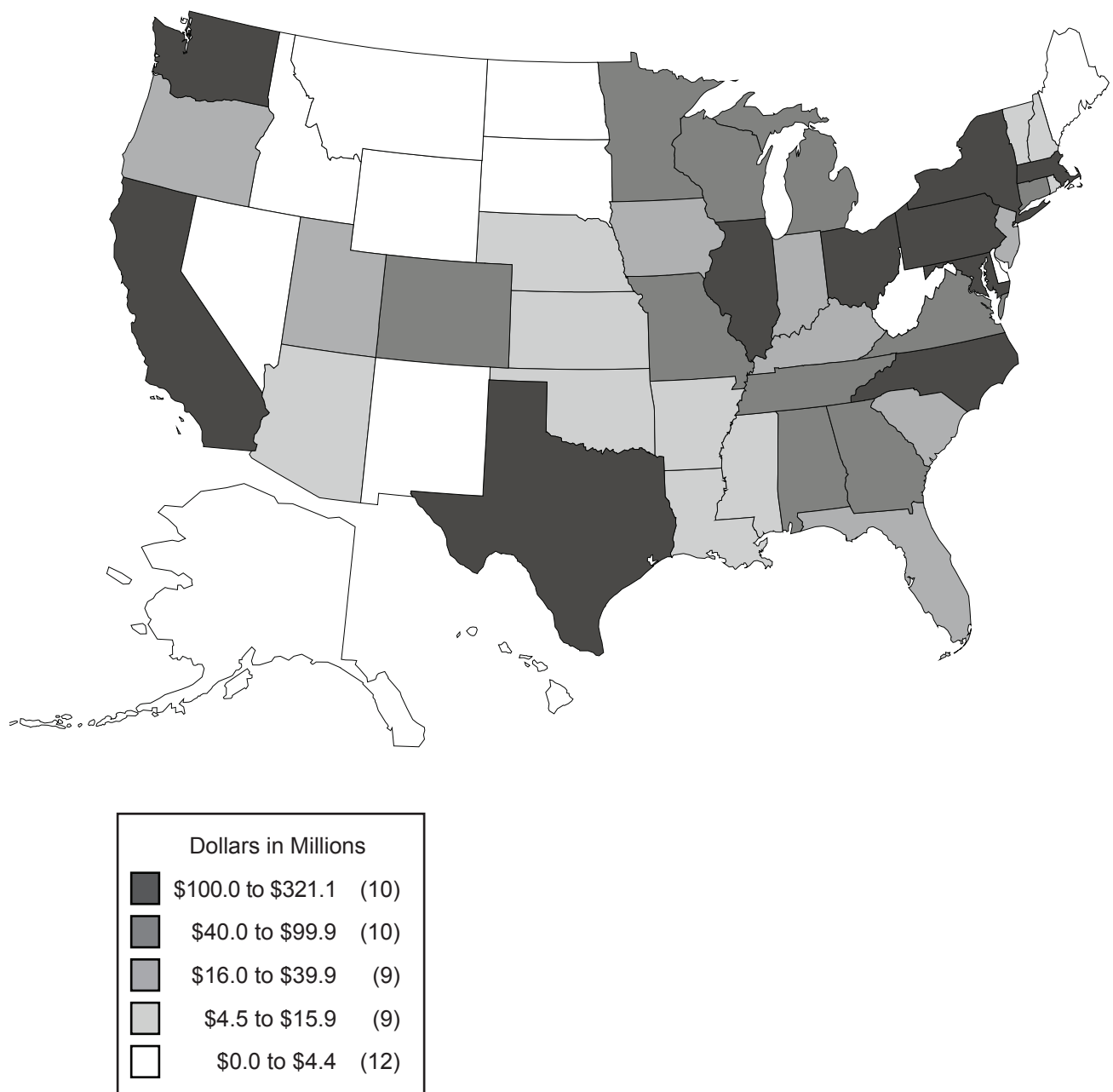
NHLBI Research Supplements Program Obligations by Award Type on Contracts: Fiscal Years 2001–2011

	Dollars (Thousands)										
	Fiscal Year										
	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011
Diversity Supplements											
Investigator	\$—	\$—	\$—	\$142	\$296	\$148	\$261	\$271	\$541	\$376	\$420
Postdoctoral	—	—	246	71	137	62	62	—	155	391	245
Graduate	—	—	108	323	229	101	294	79	155	143	215
Undergraduate	—	—	—	—	—	26	13	20	16	8	—
High School	—	—	7	—	—	—	—	—	—	—	—
Post-Master/Post-Baccalaureate	—	—	—	51	—	—	—	—	—	—	—
Total, Research Supplements Program	\$—	\$—	\$361	\$587	\$662	\$337	\$630	\$370	\$867	\$918	\$880



14. Geographic Distribution of Awards: Fiscal Year 2011

Geographic Distribution of Awards by State: Fiscal Year 2011



Geographic Distribution of Awards by State or Country: Fiscal Year 2011

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
Alabama								
Alabama University	2	5,783,466	—	—	—	—	2	5,783,466
Auburn University	2	910,233	2	910,233	—	—	—	—
Elgavish Paramagnetics, Inc.	1	590,498	1	590,498	—	—	—	—
University of Alabama at Birmingham	66	28,669,503	57	22,797,713	7	1,638,523	2	4,233,267
University of South Alabama	18	4,639,369	16	4,412,365	2	227,004	—	—
Total Alabama	89	40,593,069	76	28,710,809	9	1,865,527	4	10,016,733
Arizona								
Arizona State University, Tempe	3	1,269,263	3	1,269,263	—	—	—	—
Carl T. Hayden Medical Research Foundation	1	637,832	1	637,832	—	—	—	—
Genomics USA, Inc.	1	115,762	1	115,762	—	—	—	—
Mayo Clinic Arizona	4	1,259,028	4	1,259,028	—	—	—	—
Translational Genomics Research Institute	1	513,725	1	513,725	—	—	—	—
University of Arizona	29	10,274,328	26	9,540,410	3	733,918	—	—
Western Research Company, Inc.	1	471,157	1	471,157	—	—	—	—
Total Arizona	40	14,541,095	37	13,807,177	3	733,918	—	—
Arkansas								
Arkansas Children's Hospital Research Institute	4	1,263,368	4	1,263,368	—	—	—	—
University of Arkansas Medical Sciences, Little Rock	14	4,255,092	12	4,186,489	2	68,603	—	—
Total Arkansas	18	5,518,460	16	5,449,857	2	68,603	—	—
California								
Ablation Numerics, Inc.	1	361,816	1	361,816	—	—	—	—
Allosteros Therapeutics, Inc.	2	462,087	2	462,087	—	—	—	—
American Stem Cell, Inc.	2	638,488	2	638,488	—	—	—	—
Beckman Research Institute	8	3,982,145	7	2,325,860	—	—	1	1,656,285
BioCardia, Inc.	1	136,593	1	136,593	—	—	—	—
Biodata Innovation Systems, LLC	1	222,600	1	222,600	—	—	—	—
Biological Dynamics, Inc.	1	313,955	1	313,955	—	—	—	—
Blood Systems Research Institute	10	5,848,582	6	1,775,654	—	—	4	4,072,928
Board of Trustees of the Leland Stanford Junior University	2	4,057,384	—	—	—	—	2	4,057,384
Califia Bio, Inc.	1	218,300	1	218,300	—	—	—	—
California Institute of Technology	3	183,600	1	90,000	2	93,600	—	—
California Pacific Medical Center Research Institute	1	2,038,854	1	2,038,854	—	—	—	—
California Polytechnic State University	1	727,368	1	727,368	—	—	—	—
California State University, Northridge	1	310,346	1	310,346	—	—	—	—

Institution	Research Training and Career Development							
	Totals		Grants		Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
Capricor, Inc.	1	397,217	1	397,217	—	—	—	—
Cedars-Sinai Medical Center	15	5,606,155	14	5,545,193	1	60,962	—	—
ChemRegen, Inc.	2	344,595	2	344,595	—	—	—	—
Children's Hospital and Research Center at Oakland	8	5,226,990	7	5,132,511	1	94,479	—	—
Children's Hospital Los Angeles	6	2,518,381	6	2,518,381	—	—	—	—
Children's Hospital of Orange County	1	425,000	1	425,000	—	—	—	—
Claremont Graduate University	1	925,215	1	925,215	—	—	—	—
Cytobank, Inc.	1	153,813	—	—	—	—	1	153,813
Cytograft Tissue Engineering, Inc.	1	768,475	1	768,475	—	—	—	—
Diagnostics for the Real World, Ltd.	1	836,222	1	836,222	—	—	—	—
DNAMicroarray, Inc.	1	252,503	1	252,503	—	—	—	—
Heartflow, Inc.	1	338,268	1	338,268	—	—	—	—
HeartVista, Inc.	2	1,011,717	2	1,011,717	—	—	—	—
IntelaMetrix, Inc.	1	136,663	1	136,663	—	—	—	—
J. David Gladstone Institutes	7	7,978,880	7	7,978,880	—	—	—	—
Kaiser Foundation Research Institute	6	7,217,650	5	5,386,799	—	—	1	1,830,851
LA BioMedical Research Institute at Harbor-UCLA Medical Center	2	538,718	2	538,718	—	—	—	—
La Jolla Bioengineering Institute	2	1,002,546	2	1,002,546	—	—	—	—
La Jolla Institute for Allergy & Immunology	6	3,264,524	5	3,231,724	1	32,800	—	—
Loma Linda University	4	1,466,878	4	1,466,878	—	—	—	—
NanoVasc, Inc.	1	413,404	1	413,404	—	—	—	—
National Childhood Cancer Foundation	1	133,158	1	133,158	—	—	—	—
Northern California Institute Research and Education	6	2,581,570	6	2,581,570	—	—	—	—
Open Source Medical Software	1	997,860	—	—	—	—	1	997,860
Orthopaedic Hospital	1	414,302	1	414,302	—	—	—	—
Palo Alto Institute for Research and Education, Inc.	5	1,779,090	5	1,779,090	—	—	—	—
Palo Alto Medical Foundation Research Institute	1	774,705	1	774,705	—	—	—	—
Panorama Research, Inc.	1	289,018	1	289,018	—	—	—	—
Rand Corporation	5	2,622,743	5	2,622,743	—	—	—	—
Regents of the University of California	4	2,892,757	—	—	—	—	4	2,892,757
Salk Institute for Biological Studies	3	1,320,421	3	1,320,421	—	—	—	—
San Diego State University	15	6,064,016	14	5,216,494	—	—	1	847,522
Sanford-Burnham Medical Research Institute	12	5,944,549	12	5,944,549	—	—	—	—
Scripps Research Institute	25	11,957,476	23	11,632,812	2	324,664	—	—
Singulex, Inc.	1	99,406	1	99,406	—	—	—	—
SRI International	3	2,448,252	1	600,598	—	—	2	1,847,654
Stanford University	64	29,279,511	55	27,735,112	9	1,544,399	—	—
Targeson, Inc.	1	259,721	1	259,721	—	—	—	—
The Regents of the University of California	1	2,877,806	—	—	—	—	1	2,877,806

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
	Tristan Technologies, Inc.	1	419,555	1	419,555	—	—	—
University of California, Berkeley	7	2,157,607	5	2,097,719	2	59,888	—	—
University of California, Davis	35	14,623,437	33	14,008,033	2	615,404	—	—
University of California, Irvine	19	6,489,811	17	6,397,011	2	92,800	—	—
University of California, Lawrence Berkeley Laboratory	3	1,903,760	3	1,903,760	—	—	—	—
University of California, Los Angeles	72	37,194,412	61	35,687,242	10	1,343,473	1	163,697
University of California, Merced	3	896,317	3	896,317	—	—	—	—
University of California, Riverside	3	779,271	2	748,831	1	30,440	—	—
University of California, San Diego	92	48,969,756	81	46,866,544	11	2,103,212	—	—
University of California, San Francisco	105	41,203,157	91	38,744,284	13	1,915,221	1	543,652
University of Southern California	14	6,865,063	14	6,865,063	—	—	—	—
Vala Sciences, Inc.	1	880,128	—	—	—	—	1	880,128
Ventrix, Inc.	1	496,930	1	496,930	—	—	—	—
Veterans Medical Research Foundation, San Diego	6	1,982,532	6	1,982,532	—	—	—	—
Virogenics, Inc.	1	1,064,156	1	1,064,156	—	—	—	—
Total California	618	298,988,185	540	267,854,506	57	8,311,342	21	22,822,337
Colorado								
Colorado State University, Fort Collins	6	1,596,272	5	1,566,639	1	29,633	—	—
Kestrel Labs, Inc.	1	165,117	1	165,117	—	—	—	—
Keystone Symposia	8	114,900	8	114,900	—	—	—	—
National Jewish Health	28	14,476,111	25	13,518,731	2	104,368	1	853,012
Quest Product Development Corporation	1	685,252	1	685,252	—	—	—	—
Taiga Biotechnologies, Inc.	1	759,221	1	759,221	—	—	—	—
TDA Research, Inc.	1	149,996	1	149,996	—	—	—	—
University of Colorado, Boulder	8	2,810,419	7	2,565,539	1	244,880	—	—
University of Colorado, Denver	62	21,975,891	53	20,018,451	7	1,526,556	2	430,884
Total Colorado	116	42,733,179	102	39,543,846	11	1,905,437	3	1,283,896
Connecticut								
Connecticut Children's Medical Center	1	121,230	1	121,230	—	—	—	—
Hartford Hospital	1	429,312	1	429,312	—	—	—	—
John B. Pierce Laboratory, Inc.	1	238,669	1	238,669	—	—	—	—
RxGen, Inc.	1	718,304	1	718,304	—	—	—	—
University of Connecticut Health Center	5	3,190,784	5	3,190,784	—	—	—	—
University of Connecticut, Storrs	5	1,201,175	4	1,169,655	1	31,520	—	—
Yale University	84	36,406,535	70	32,338,012	12	1,910,970	2	2,157,553
Total Connecticut	98	42,306,009	83	38,205,966	13	1,942,490	2	2,157,553

Institution	No.	Dollar	Research Training and Career Development					
			Totals		Grants		Development	
			No.	Dollar	No.	Dollar	No.	Dollar
Delaware								
University of Delaware	5	1,316,545	4	1,261,811	1	54,734	—	—
Total Delaware	5	1,316,545	4	1,261,811	1	54,734	0	0
District of Columbia								
Academy for Education Development	1	2,370,712	—	—	—	—	1	2,370,712
American Institute for Research	2	5,438,370	—	—	—	—	2	5,438,370
American Society of Hematology	1	20,000	1	20,000	—	—	—	—
Children's Research Institute	7	2,811,132	7	2,811,132	—	—	—	—
The George Washington University	7	4,651,793	7	4,651,793	—	—	—	—
Georgetown University	5	3,343,966	5	3,343,966	—	—	—	—
Howard University	6	1,256,637	4	1,149,782	1	15,000	1	91,855
National Academy of Science	2	100,000	—	—	—	—	2	100,000
Ogilvy Public Relations	3	4,014,080	—	—	—	—	3	4,014,080
Porter Novelli Public Service	1	1,000,324	—	—	—	—	1	1,000,324
U.S. Bureau of Census	1	535,000	—	—	—	—	1	535,000
Total District of Columbia	36	25,542,013	24	11,976,673	1	15,000	11	13,550,340
Florida								
Altor Bioscience Corporation	1	1,062,358	1	1,062,358	—	—	—	—
ArchieMD, Inc.	1	490,980	1	490,980	—	—	—	—
GeNO, LLC	1	294,031	1	294,031	—	—	—	—
H. Lee Moffitt Cancer Center and Research Institute	1	165,330	1	165,330	—	—	—	—
HeartWare, Inc.	1	385,813	1	385,813	—	—	—	—
Nemours Children's Clinic	1	109,300	1	109,300	—	—	—	—
Nova Southeastern University	1	339,708	1	339,708	—	—	—	—
Self-Determined Health, Inc.	1	430,557	1	430,557	—	—	—	—
Sharklet Technologies, LLC	1	215,835	1	215,835	—	—	—	—
Torrey Pines Institute for Molecular Studies	2	844,081	2	844,081	—	—	—	—
University of Central Florida	3	1,175,702	3	1,175,702	—	—	—	—
University of Florida	38	14,853,744	34	14,455,836	4	397,908	—	—
University of Miami School of Medicine	30	13,724,577	27	12,597,929	2	642,471	1	484,177
University of South Florida	5	1,352,854	4	1,089,750	1	263,104	—	—
Total Florida	87	35,444,870	79	33,657,210	7	1,303,483	1	484,177
Georgia								
Emory University	58	24,299,843	52	23,101,636	6	1,198,207	—	—
Expression Therapeutics	1	885,011	1	885,011	—	—	—	—
Georgia Health Sciences University	31	17,542,157	30	17,421,999	1	120,158	—	—
Georgia Institute of Technology	9	3,371,472	9	3,371,472	—	—	—	—
MedShape Solutions, Inc.	1	407,885	1	407,885	—	—	—	—
Morehouse School of Medicine	9	2,016,631	8	1,756,375	1	260,256	—	—

Institution	Research Training and Career Development							
	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
Northside Hospital Atlanta	1	97,532	1	97,532	—	—	—	—
Syntermed, Inc,	1	216,081	1	216,081	—	—	—	—
University of Georgia	4	1,426,250	4	1,426,250	—	—	—	—
U.S. Centers for Disease Control and Prevention	5	7,301,761	—	—	—	—	5	7,301,761
Total Georgia	120	57,564,623	107	48,684,241	8	1,578,621	5	7,301,761
Hawaii								
University of Hawaii, Manoa	4	1,660,486	4	1,660,486	—	—	—	—
Total Hawaii	4	1,660,486	4	1,660,486	—	—	—	—
Illinois								
Advanced Diamond Technologies, Inc.	1	149,817	1	149,817	—	—	—	—
Aqualung Therapeutics Corp.	1	320,967	1	320,967	—	—	—	—
Biomedical Acoustics Research Company	1	339,058	1	339,058	—	—	—	—
Children's Memorial Hospital	7	1,036,514	6	1,007,932	1	28,582	—	—
Coramed Technologies	1	999,186	1	999,186	—	—	—	—
Illinois Institute of Technology	2	556,750	2	556,750	—	—	—	—
Loyola University, Chicago	10	3,484,003	9	3,437,203	1	46,800	—	—
Northshore University HealthSystem	1	429,730	1	429,730	—	—	—	—
Northwestern University	82	35,559,950	67	30,791,540	13	1,409,691	2	3,358,719
Rosalind Franklin University of Medicine and Science	1	368,800	1	368,800	—	—	—	—
Rush University Medical Center	11	5,328,276	11	5,328,276	—	—	—	—
SonoGene, LLC	1	361,937	1	361,937	—	—	—	—
University of Chicago	47	16,712,464	40	14,880,798	7	1,831,666	—	—
University of Illinois, Chicago	61	31,775,801	56	29,798,108	5	1,977,693	—	—
University of Illinois, Urbana-Champaign	13	3,343,354	8	3,119,257	5	224,097	—	—
Total Illinois	240	100,766,607	206	91,889,359	32	5,518,529	2	3,358,719
Indiana								
Clarian Health Partners, Inc.	1	168,750	1	168,750	—	—	—	—
Indiana University-Purdue University at Indianapolis	46	18,502,398	39	16,689,483	6	880,625	1	932,290
International Aldosterone Conference	1	15,000	1	15,000	—	—	—	—
Purdue University, West Lafayette	3	920,976	3	920,976	—	—	—	—
SonarMed, Inc.	1	224,593	1	224,593	—	—	—	—
University of Notre Dame	2	749,251	2	749,251	—	—	—	—
Total Indiana	54	20,580,968	47	18,768,053	6	880,625	1	932,290

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
Iowa								
Exemplar Genetics, LLC	1	639,811	1	639,811	—	—	—	—
Iowa State University	1	322,821	1	322,821	—	—	—	—
University of Iowa	61	31,788,154	52	28,982,851	9	2,805,303	—	—
Vida Diagnostics, Inc.	1	506,107	1	506,107	—	—	—	—
Total Iowa	64	33,256,893	55	30,451,590	9	2,805,303	—	—
Kansas								
Kansas State University	2	392,880	1	365,546	1	27,334	—	—
Pinnacle Technology, Inc.	1	463,406	1	463,406	—	—	—	—
University of Kansas Medical Center	11	3,896,691	11	3,896,691	—	—	—	—
Total Kansas	14	4,752,977	13	4,725,643	1	27,334	—	—
Kentucky								
Eastern Kentucky University	1	389,445	1	389,445	—	—	—	—
Pharmacogenetics Diagnostic Laboratories	1	1,070,149	1	1,070,149	—	—	—	—
University of Kentucky	20	7,318,522	15	6,693,538	5	624,984	—	—
University of Louisville	21	11,192,859	20	10,661,829	1	531,030	—	—
W-Z Biotech, LLC	1	175,322	1	175,322	—	—	—	—
Total Kentucky	44	20,146,297	38	18,990,283	6	1,156,014	—	—
Louisiana								
Children's Hospital (New Orleans)	1	305,584	1	305,584	—	—	—	—
Louisiana State University and A&M College, Baton Rouge	2	692,604	2	692,604	—	—	—	—
Louisiana State University Health Sciences Center, New Orleans	14	5,090,295	11	4,884,783	2	56,813	1	148,699
Louisiana State University Health Sciences Center, Shreveport	3	1,086,250	3	1,086,250	—	—	—	—
Louisiana State University Pennington Biomedical Research Center	2	840,272	2	840,272	—	—	—	—
NuPotential, Inc.	1	230,011	1	230,011	—	—	—	—
Tulane University of Louisiana	13	5,421,997	12	5,240,810	1	181,187	—	—
Total Louisiana	36	13,667,013	32	13,280,314	3	238,000	1	148,699
Maine								
Jackson Laboratory	4	2,356,850	3	2,319,250	1	37,600	—	—
Maine Medical Center	4	1,473,900	4	1,473,900	—	—	—	—
University of Maine, Orono	1	593,233	1	593,233	—	—	—	—
Total Maine	9	4,423,983	8	4,386,383	1	37,600	—	—
Maryland								
Advance Bioscience Laboratories, Inc.	1	1,271,757	—	—	—	—	1	1,271,757
American Institutes for Research	1	1,687,214	—	—	—	—	1	1,687,214

Institution	Research Training and Career Development							
	Totals		Grants		Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
CDM Group, Inc.	1	479,000	—	—	—	—	1	479,000
Cerecor, Inc.	1	292,476	1	292,476	—	—	—	—
Clinical Trials & Surveys Corporation	3	5,676,215	1	1,099,694	—	—	2	4,576,521
Danya International	1	50,000	—	—	—	—	1	50,000
Emmes Corporation	1	1,255,602	—	—	—	—	1	1,255,602
Engineering and Scientific Research Association	1	257,949	1	257,949	—	—	—	—
Federation of American Societies for Experimental Biology	1	10,000	1	10,000	—	—	—	—
Henry M. Jackson Foundation for the Advancement of Military Medicine	1	382,500	1	382,500	—	—	—	—
Health Resources and Services Administration, Maternal and Child Health Bureau	1	150,000	—	—	—	—	1	150,000
Innoscion, LLC	1	133,446	1	133,446	—	—	—	—
Johns Hopkins University	185	104,932,095	156	90,683,398	20	4,057,308	9	10,191,389
Medical Decision Logic, Inc.	2	550,955	1	349,070	—	—	1	201,885
MedStar Research Institute	1	916,209	1	916,209	—	—	—	—
National Human Genome Research Institute	1	2,527,000	—	—	—	—	1	2,527,000
National Institutes of Health	5	2,490,079	—	—	—	—	5	2,490,079
National Library of Medicine	2	1,017,015	—	—	—	—	2	1,017,015
North American Vascular Biology Organization	2	18,401	2	18,401	—	—	—	—
RetroTherapy, LLC	1	521,305	1	521,305	—	—	—	—
Science Applications International Corporation	1	2,999,960	—	—	—	—	1	2,999,960
SeraCare Life Sciences	2	3,209,369	—	—	—	—	2	3,209,369
Social and Scientific Systems, Inc	1	3,891,835	—	—	—	—	1	3,891,835
University of Maryland, Baltimore	42	23,855,324	38	21,200,528	3	764,006	1	1,890,790
University of Maryland, College Park	3	643,939	3	643,939	—	—	—	—
U.S. Department of Agriculture	1	115,000	—	—	—	—	1	115,000
U.S. Department of Health and Human Services, Indian Health Service	1	739,459	—	—	—	—	1	739,459
U.S. Food and Drug Administration	2	232,000	—	—	—	—	2	232,000
Veracity Biotechnology, LLC	1	164,310	1	164,310	—	—	—	—
Westat	1	1,523,238	—	—	—	—	1	1,523,238
Total Maryland	268	161,993,652	209	116,673,225	23	4,821,314	36	40,499,113

Massachusetts

Abiomed, Inc.	1	663,647	1	663,647	—	—	—	—
Aphios Corporation	1	149,440	—	—	—	—	1	149,440
Baystate Medical Center	2	739,913	1	436,788	—	—	1	303,125
Beth Israel Deaconess Medical Center	51	28,629,562	46	26,771,903	5	1,857,659	—	—
BioSurfaces	1	379,528	1	379,528	—	—	—	—
Boston Biomedical Research Institute	3	1,762,058	3	1,762,058	—	—	—	—

Institution	No.	Dollar	Research Training and Career Development					
			Totals		Grants		Development	
			No.	Dollar	No.	Dollar	No.	Dollar
Boston Medical Center	12	7,023,812	12	7,023,812	—	—	—	—
Boston University	7	6,505,343	6	3,700,080	—	—	1	2,805,263
Boston University Medical Campus	48	39,411,313	41	20,148,392	6	2,598,416	1	16,664,505
Brigham and Women's Hospital	172	97,064,725	155	91,528,877	16	4,322,381	1	1,213,467
Cardiovascular Engineering, Inc.	1	640,161	1	640,161	—	—	—	—
Charles Stark Draper Laboratory	2	447,366	2	447,366	—	—	—	—
Children's Hospital Boston	49	27,649,874	46	26,559,727	3	1,090,147	—	—
Dana-Farber Cancer Institute	10	3,824,181	10	3,824,181	—	—	—	—
DecImmune Therapeutics, Inc.	1	1,044,768	1	1,044,768	—	—	—	—
E-TROLZ, Inc.	1	191,624	1	191,624	—	—	—	—
Franklin W. Olin College of Engineering	1	273,000	1	273,000	—	—	—	—
Harvard Pilgrim Health Care, Inc.	3	1,084,619	3	1,084,619	—	—	—	—
Harvard University	4	2,003,375	3	1,776,229	1	227,146	—	—
Harvard University Medical School	10	4,893,069	8	4,827,063	2	66,006	—	—
Harvard University School of Public Health	14	7,428,814	12	6,624,280	2	804,534	—	—
Heartlander Surgical, Inc.	1	98,519	1	98,519	—	—	—	—
Immune Disease Institute, Inc.	6	7,404,959	5	4,574,207	—	—	1	2,830,752
Immunetics, Inc.	1	299,989	1	299,989	—	—	—	—
Infoscitex Corporation	2	1,798,818	2	1,798,818	—	—	—	—
Joslin Diabetes Center	3	2,391,224	3	2,391,224	—	—	—	—
Kala Pharmaceuticals, Inc.	1	180,415	1	180,415	—	—	—	—
Levitronix, LLC	1	897,979	1	897,979	—	—	—	—
Massachusetts General Hospital	72	37,681,979	64	32,336,768	6	1,410,946	2	3,934,265
Massachusetts Institute of Technology	4	1,519,498	3	1,468,172	1	51,326	—	—
Medi-Solve Coatings, LLC	1	194,164	1	194,164	—	—	—	—
MedMinder Systems, Inc.	1	193,573	1	193,573	—	—	—	—
New England Research Institutes, Inc.	1	8,923,478	1	8,923,478	—	—	—	—
Northeastern University	2	2,246,822	2	2,246,822	—	—	—	—
Oxus Medical, Inc.	1	188,628	1	188,628	—	—	—	—
Phoenicia Biosciences, Inc.	1	304,344	1	304,344	—	—	—	—
Radiation Monitoring Devices, Inc.	2	828,730	2	828,730	—	—	—	—
Radikal Therapeutics, Inc.	3	694,397	3	694,397	—	—	—	—
Spaulding Rehabilitation Hospital	1	359,003	1	359,003	—	—	—	—
Springfield College	—	101,592	—	101,592	—	—	—	—
Tufts Medical Center	18	5,316,152	17	4,817,973	1	498,179	—	—
Tufts University, Boston	11	4,172,734	9	3,990,436	2	182,298	—	—
Tufts University, Medford	1	249,000	1	249,000	—	—	—	—
University of Massachusetts Medical School	29	12,523,145	28	12,470,103	1	53,042	—	—
University of Massachusetts, Amherst	1	438,295	1	438,295	—	—	—	—
VasoTech, Inc.	1	534,636	1	534,636	—	—	—	—
Total Massachusetts	559	321,352,265	505	280,289,368	46	13,162,080	8	27,900,817

Institution	No.	Dollar	Research Training and Career Development					
			Totals		Grants		Contracts	
			No.	Dollar	No.	Dollar	No.	Dollar
Michigan								
Accord Biomaterials, Inc.	1	873,517	1	873,517	—	—	—	—
AlphaCore Pharma, LLC	1	231,470	1	231,470	—	—	—	—
Altarum Institute	1	110,431	1	110,431	—	—	—	—
Arbor Ultrasound Technologies, LLC	1	101,323	1	101,323	—	—	—	—
Central Michigan University	1	406,580	1	406,580	—	—	—	—
Epsilon Imaging, Inc.	1	883,531	1	883,531	—	—	—	—
Henry Ford Health System	7	6,401,254	6	6,342,572	1	58,682	—	—
Integrated Sensing Systems, Inc.	1	172,911	1	172,911	—	—	—	—
L-VAD Technology, Inc.	1	308,409	1	308,409	—	—	—	—
MC3, Inc.	4	1,835,734	4	1,835,734	—	—	—	—
MedArray, Inc.	1	999,718	1	999,718	—	—	—	—
Michigan State University	9	3,319,713	8	3,280,584	1	39,129	—	—
Michigan Technological University	2	382,757	2	382,757	—	—	—	—
Oakland University	1	240,514	1	240,514	—	—	—	—
University of Michigan, Ann Arbor	113	52,487,986	99	43,067,353	10	2,239,368	4	7,181,265
Wayne State University	15	5,962,405	14	5,928,714	1	33,691	—	—
Total Michigan	160	74,718,253	143	65,166,118	13	2,370,870	4	7,181,265
Minnesota								
Advanced Circulatory Systems, Inc.	1	210,878	1	210,878	—	—	—	—
Discovery Genomics, Inc.	1	716,704	1	716,704	—	—	—	—
Gel-Del Technologies, Inc.	1	1,395,176	1	1,395,176	—	—	—	—
HealthPartners Research Foundation	5	2,610,060	5	2,610,060	—	—	—	—
Koronis Biomedical Technologies Corporation	1	158,785	1	158,785	—	—	—	—
Mayo Clinic, Rochester	53	25,217,879	47	23,590,876	3	314,363	3	1,312,640
Minneapolis Medical Research Foundation, Inc.	1	304,616	1	304,616	—	—	—	—
Minnetronix, Inc.	2	367,020	2	367,020	—	—	—	—
National Marrow Donor Program, Inc.	1	10,000	1	10,000	—	—	—	—
Recombinetics, Inc.	1	318,202	1	318,202	—	—	—	—
ResQSystems, Inc.	1	256,286	1	256,286	—	—	—	—
University of Minnesota, Duluth	1	445,666	1	445,666	—	—	—	—
University of Minnesota, Twin Cities	78	43,522,179	66	36,016,972	9	1,534,739	3	5,970,468
Total Minnesota	147	75,533,451	129	66,401,241	12	1,849,102	6	7,283,108
Mississippi								
Jackson State University	2	469,108	2	307,702	—	—	—	161,406
Tougaloo College	—	35,371	—	—	—	—	—	35,371
University of Mississippi Medical Center	21	15,914,661	18	8,949,247	1	380,849	2	6,584,565
Total Mississippi	23	16,419,140	20	9,256,949	1	380,849	2	6,781,342

Institution	Research Training and Career Development							
	Totals		Grants		Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
Missouri								
CardiaLen, Inc.	1	216,027	1	216,027	—	—	—	—
Children's Mercy Hospital, Kansas City	1	253,466	1	253,466	—	—	—	—
Nanova, Inc.	1	220,243	1	220,243	—	—	—	—
Saint Louis University	8	2,631,596	8	2,631,596	—	—	—	—
Saint Luke's Hospital	1	363,663	1	363,663	—	—	—	—
University of Missouri, Columbia	24	10,099,513	22	9,999,789	2	99,724	—	—
University of Missouri, Kansas City	1	187,500	1	187,500	—	—	—	—
Washington University	112	46,571,996	93	41,413,852	17	3,913,788	2	1,244,356
Total Missouri	149	60,544,004	128	55,286,136	19	4,013,512	2	1,244,356
Montana								
Montana State University, Bozeman	3	1,068,750	3	1,068,750	—	—	—	—
Total Montana	3	1,068,750	3	1,068,750	—	—	—	—
Nebraska								
Creighton University	3	1,306,514	3	1,306,514	—	—	—	—
University of Nebraska, Lincoln	1	304,405	1	304,405	—	—	—	—
University of Nebraska Medical Center	7	3,803,563	7	3,803,563	—	—	—	—
Total Nebraska	11	5,414,482	11	5,414,482	—	—	—	—
Nevada								
University of Nevada, Reno	8	2,453,962	7	2,425,908	1	28,054	—	—
Total Nevada	8	2,453,962	7	2,425,908	1	28,054	—	—
New Hampshire								
Celdara Medical, LLC	1	747,897	1	747,897	—	—	—	—
Dartmouth College	9	2,856,719	9	2,856,719	—	—	—	—
Xemed, LLC	2	1,374,918	2	1,374,918	—	—	—	—
Total New Hampshire	12	4,979,534	12	4,979,534	—	—	—	—
New Jersey								
Artann Laboratories, Inc.	1	223,219	1	223,219	—	—	—	—
CircuLite, Inc.	1	1,621,110	1	1,621,110	—	—	—	—
DVX, LLC	2	939,116	2	939,116	—	—	—	—
Hackensack University Medical Center	1	220,125	1	220,125	—	—	—	—
Menssana Research, Inc.	1	992,862	1	992,862	—	—	—	—
Newark Beth Israel Medical Center	1	151,881	1	151,881	—	—	—	—
NovaFlux Technologies, Inc.	1	299,989	1	299,989	—	—	—	—
Princeton University	1	241,500	1	241,500	—	—	—	—
Rutgers the State University of New Jersey, New Brunswick	3	1,160,059	3	1,160,059	—	—	—	—

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
University of Medicine & Dentistry of New Jersey, New Jersey Medical School	25	9,825,798	22	9,601,165	3	224,633	—	—
University of Medicine & Dentistry of New Jersey, Robert Wood Johnson Medical School	5	2,443,491	5	2,443,491	—	—	—	—
Vasade Biosciences, Inc.	1	556,030	1	556,030	—	—	—	—
Total New Jersey	43	18,675,180	40	18,450,547	3	224,633	—	—
New Mexico								
Lovelace Biomedical and Environmental Research	2	972,000	2	972,000	—	—	—	—
Southwest Sciences, Inc.	—	98,792	—	98,792	—	—	—	—
University of New Mexico	7	2,332,321	6	2,063,039	1	269,282	—	—
University of New Mexico Health Sciences Center	3	724,702	3	724,702	—	—	—	—
Total New Mexico	12	4,127,815	11	3,858,533	1	269,282	—	—
New York								
Albany College of Pharmacy	1	347,369	1	347,369	—	—	—	—
Albany Medical College	5	1,681,315	4	1,638,265	1	43,050	—	—
Albert Einstein College of Medicine, Yeshiva University	30	12,618,203	24	10,265,670	4	361,867	2	1,990,666
Angion Biomedica Corporation	1	1,484,711	1	1,484,711	—	—	—	—
Circulatory Technology, Inc.	1	634,135	1	634,135	—	—	—	—
City College of New York	4	1,711,084	4	1,711,084	—	—	—	—
Columbia University	73	30,944,636	64	29,122,382	8	1,639,676	1	182,578
Cornell University, Ithaca	8	3,542,830	8	3,542,830	—	—	—	—
Feinstein Institute for Medical Research	3	1,240,661	3	1,240,661	—	—	—	—
General Electric Global Research Center	1	832,322	1	832,322	—	—	—	—
Herbert H. Lehman College	1	285,250	1	285,250	—	—	—	—
Hospital for Special Surgery	2	559,908	2	559,908	—	—	—	—
Ithaca College	1	117,121	1	117,121	—	—	—	—
Jarvik Heart, Inc.	1	2,531,000	—	—	—	—	1	2,531,000
Masonic Medical Research Laboratory, Inc.	1	437,500	1	437,500	—	—	—	—
Micro Photo Acoustics, Inc.	1	100,000	1	100,000	—	—	—	—
Mount Sinai School of Medicine	38	20,821,370	34	20,164,939	4	656,431	—	—
Nanometrics, LLC	1	243,469	1	243,469	—	—	—	—
New York Blood Center	5	1,752,557	5	1,752,557	—	—	—	—
New York Institute of Technology	1	356,235	1	356,235	—	—	—	—
New York Medical College	14	7,595,823	11	7,473,815	3	122,008	—	—
New York University	2	424,278	1	383,372	1	40,906	—	—
New York University School of Medicine	38	24,189,884	34	23,408,592	4	781,292	—	—
Pulmokine, Inc.	1	72,500	1	72,500	—	—	—	—
Queens College	1	193,750	1	193,750	—	—	—	—
Rensselaer Polytechnic Institute	2	982,732	2	982,732	—	—	—	—

Institution	No.	Dollar	Research Training and Career Development					
			Totals		Grants		Development	
			No.	Dollar	No.	Dollar	No.	Dollar
Rockefeller University	3	1,539,672	3	1,539,672	—	—	—	—
Roswell Park Cancer Institute Corp.	2	2,136,723	2	2,136,723	—	—	—	—
Sloan-Kettering Institute for Cancer Research	4	1,659,045	4	1,659,045	—	—	—	—
St. Luke's-Roosevelt Institute for Health Sciences	1	652,615	1	652,615	—	—	—	—
State University New York, Stony Brook	6	2,266,870	6	2,266,870	—	—	—	—
State University of New York, Buffalo	14	5,615,925	14	5,615,925	—	—	—	—
State University of New York, Downstate Medical Center	5	1,957,571	5	1,957,571	—	—	—	—
TheraSource, LLC	1	571,120	1	571,120	—	—	—	—
Transonic Systems, Inc.	2	1,225,000	2	1,225,000	—	—	—	—
University of Buffalo	1	1,527,870	—	—	—	—	1	1,527,870
Union College	1	300,000	1	300,000	—	—	—	—
University of Rochester	60	23,801,150	53	22,428,024	7	1,373,126	—	—
Upstate Medical University	8	2,657,638	8	2,657,638	—	—	—	—
Visiting Nurse Service of New York	—	229,932	—	229,932	—	—	—	—
Wadsworth Center	1	385,921	1	385,921	—	—	—	—
Weill Medical College of Cornell University	32	19,258,955	29	18,823,010	3	435,945	—	—
Winifred Masterson Burke Medical Research Institute	1	470,206	1	470,206	—	—	—	—
Total New York	379	181,958,856	339	170,270,441	35	5,454,301	5	6,232,114

North Carolina

Duke University	95	47,195,222	86	44,974,806	8	1,821,658	1	398,758
East Carolina University	4	1,060,669	4	1,060,669	—	—	—	—
Gramercy Research Group, LLC	1	821,249	1	821,249	—	—	—	—
Heart Imaging Technologies, LLC	1	249,214	1	249,214	—	—	—	—
LifeSciTech, LLC	1	407,336	1	407,336	—	—	—	—
North Carolina Central University	1	280,000	1	280,000	—	—	—	—
North Carolina State University, Raleigh	5	1,565,785	3	1,135,388	2	430,397	—	—
Parion Sciences, Inc.	1	244,797	1	244,797	—	—	—	—
Research Triangle Institute	7	15,663,444	1	3,559,385	—	—	6	12,104,059
Rho Federal Systems Division, Inc.	1	1	1	1	—	—	—	—
University of North Carolina, Chapel Hill	87	40,977,208	71	33,120,732	11	1,286,104	5	6,570,372
Wake Forest University Health Sciences	36	33,204,306	26	14,819,217	4	1,055,533	6	17,329,556
Wake Forest University School of Medicine	1	345,112	1	345,112	—	—	—	—
Total North Carolina	241	142,014,343	198	101,017,906	25	4,593,692	18	36,402,745

Ohio

Arteriocyte, Inc.	1	586,833	1	586,833	—	—	—	—
Biofunc Research	1	159,108	1	159,108	—	—	—	—
Case Western Reserve University	57	20,516,005	51	19,018,137	4	1,004,806	2	493,062
Cincinnati Children's Hospital Medical Center	71	24,668,022	63	23,961,703	8	706,319	—	—

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
	46	29,980,105	43	29,594,765	3	385,340	—	—
Cleveland Clinic-Lerner College of Medicine	3	1,171,915	3	1,171,915	—	—	—	—
Cleveland Medical Devices, Inc.	2	666,010	2	666,010	—	—	—	—
Cleveland State University	1	247,272	1	247,272	—	—	—	—
H-Cubed, Inc.	2	987,657	2	987,657	—	—	—	—
Kent State University, Kent	7	1,585,730	5	1,267,129	2	318,601	—	—
Nationwide Children's Hospital	2	840,865	2	840,865	—	—	—	—
Northeast Ohio Medical University	2	1,223,006	2	1,223,006	—	—	—	—
NovelMed Therapeutics, Inc.	41	13,281,300	38	12,565,793	2	94,092	1	621,415
Ohio State University	1	248,847	1	248,847	—	—	—	—
University of Akron	30	10,894,061	27	10,450,715	3	443,346	—	—
University of Cincinnati	1	1,404,344	—	—	—	—	1	1,404,344
University of Toledo Health Sciences Campus	5	5,159,187	5	5,159,187	—	—	—	—
Walsh University	6	356,556	1	356,556	—	—	—	—
Wright State University	280	115,819,754	254	110,348,429	22	2,952,504	4	2,518,821
Oklahoma								
Oklahoma Medical Research Foundation	5	3,640,696	5	3,640,696	—	—	—	—
Oklahoma State University, Stillwater	11	365,470	1	365,470	—	—	—	—
University of Oklahoma Health Sciences Center	17	5,271,212	11	5,271,212	—	—	—	—
Total Oklahoma	17	9,277,378	17	9,277,378	—	—	—	—
Oregon								
Aronora, LLC	2	1,782,390	2	1,782,390	—	—	—	—
Fanconi Anemia Research Fund, Inc.	1	10,000	1	10,000	—	—	—	—
Gamma Therapeutics, Inc.	2	520,322	1	520,322	—	—	—	—
Oregon Center for Applied Science, Inc.	36	968,255	30	968,255	—	—	—	—
Oregon Health and Science University	1	15,736,615	30	14,373,546	6	1,363,069	—	—
Oregon State University	1	359,770	1	359,770	—	—	—	—
Portland State University	1	365,000	1	365,000	—	—	—	—
University of Oregon	47	609,300	2	573,123	1	36,177	—	—
Total Oregon	47	20,351,652	40	18,952,406	7	1,399,246	—	—
Pennsylvania								
Carmell Therapeutics Corporation	3	51,482	3	51,482	—	—	—	—
Carnegie Mellon University	1	915,093	1	915,093	—	—	—	—
Cereve, Inc.	1	1,156,255	—	—	—	—	—	—
Children's Hospital of Pennsylvania	1	1,311,380	—	—	—	—	1	1,311,380
Children's Hospital of Philadelphia	40	19,309,524	37	18,480,070	3	829,454	—	—
Children's Hospital of Pittsburgh	1	106,487	—	—	—	—	1	106,487
Convance Research Product	1	84,738	—	—	—	—	1	84,738

Institution	Research Training and Career Development							
	Totals		Grants		Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
Discovery Laboratories, Inc.	1	581,839	1	581,839	—	—	—	—
Drexel University	1	386,250	1	386,250	—	—	—	—
Ension, Inc.	1	2,164,774	—	—	—	—	1	2,164,774
In Silico Molecular, LLC	1	236,765	1	236,765	—	—	—	—
Institute of Transfusion Medicine	2	1,514,117	—	—	—	—	2	1,514,117
Lehigh University	1	458,143	1	458,143	—	—	—	—
Magee-Women's Research Institute and Foundation	1	325,091	1	325,091	—	—	—	—
NanoDynamics Life Sciences, Inc.	1	556,284	1	556,284	—	—	—	—
National Disease Research Interchange	—	151,286	—	151,286	—	—	—	—
University of Pennsylvania	1	6,760,320	—	—	—	—	1	6,760,320
Pennsylvania State University, Hershey Medical Center	18	16,150,357	17	16,101,959	1	48,398	—	—
Pennsylvania State University, University Park	4	946,717	3	910,435	1	36,282	—	—
Philadelphia College of Osteopathic Medicine	1	396,250	1	396,250	—	—	—	—
Progenra, Inc.	2	764,300	2	764,300	—	—	—	—
Salus University	1	350,144	1	350,144	—	—	—	—
Shifa Biomedical	1	655,327	1	655,327	—	—	—	—
Strategic Polymer Sciences, Inc.	1	374,904	1	374,904	—	—	—	—
Temple University	41	13,515,863	33	11,254,720	6	642,863	2	1,618,280
Thomas Jefferson University	23	9,548,847	21	9,453,649	2	95,198	—	—
University of Pennsylvania	134	70,521,644	116	64,249,473	17	4,972,171	1	1,300,000
University of Pittsburgh	127	56,121,300	110	49,175,899	14	2,654,871	3	4,290,530
Wistar Institute	3	2,597,866	3	2,597,866	—	—	—	—
Total Pennsylvania	413	208,013,347	356	179,583,484	44	9,279,237	13	19,150,626

Rhode Island

Brown University	5	1,147,164	3	1,070,534	2	76,630	—	—
Butler Hospital, Providence	1	489,901	1	489,901	—	—	—	—
Gordon Research Conferences	9	157,000	9	157,000	—	—	—	—
Memorial Hospital of Rhode Island	1	662,903	1	662,903	—	—	—	—
Miriam Hospital	5	2,624,780	4	2,389,203	1	235,577	—	—
Pro-Change Behavior Systems, Inc.	1	205,150	1	205,150	—	—	—	—
Rhode Island Hospital	12	5,333,767	10	4,697,846	2	635,921	—	—
Roger Williams Hospital	1	423,169	1	423,169	—	—	—	—
University of Rhode Island	1	260,590	1	260,590	—	—	—	—
Total Rhode Island	36	11,304,424	31	10,356,296	5	948,128	—	—

South Carolina

Cell and Tissue Systems, Inc.	2	502,990	2	502,990	—	—	—	—
Clemson University	4	1,109,773	4	1,109,773	—	—	—	—
Medical University of South Carolina	31	10,029,206	27	9,052,085	4	977,121	—	—

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
	1	372,579	1	372,579	—	—	—	—
MicroVide, LLC	13	4,611,639	13	4,611,639	—	—	—	—
University of South Carolina, Columbia	51	16,626,187	47	15,649,066	4	977,121	—	—
Total South Carolina								
South Dakota								
Missouri Breaks Research, Inc.	—	300,000	—	300,000	—	—	—	—
South Dakota State University	1	360,970	1	360,970	—	—	—	—
University of South Dakota	3	522,460	2	516,958	1	5,502	—	—
Total South Dakota	4	1,183,430	3	1,177,928	1	5,502	—	—
Tennessee								
East Tennessee State University	4	1,358,500	4	1,358,500	—	—	—	—
Meharry Medical College	5	998,096	2	527,800	3	470,296	—	—
St. Jude Children's Research Hospital	8	5,543,986	7	5,490,944	1	53,042	—	—
University of Tennessee Health Science Center	25	10,778,783	25	10,778,783	—	—	—	—
University of Tennessee, Knoxville	1	372,500	1	372,500	—	—	—	—
Vanderbilt University	105	45,713,230	92	42,920,823	12	2,553,423	1	238,984
Total Tennessee	148	64,765,095	131	61,449,350	16	3,076,761	1	238,984
Texas								
Baylor College of Medicine	61	26,259,630	48	20,408,696	11	2,180,376	2	3,670,558
Baylor Research Institute	1	818,840	1	818,840	—	—	—	—
BioTex, Inc.	1	150,000	1	150,000	—	—	—	—
Kardia Therapeutics, Inc.	1	659,436	1	659,436	—	—	—	—
Methodist Hospital Research Institute	4	837,770	4	837,770	—	—	—	—
Rice University	5	1,287,685	5	1,287,685	—	—	—	—
Southern Methodist University	1	334,511	1	334,511	—	—	—	—
Southwest Research Institute	1	223,109	1	223,109	—	—	—	—
Telehealth Holdings, LLC	1	199,965	1	199,965	—	—	—	—
Texas A&M University Health Science Center	13	3,163,100	13	3,163,100	—	—	—	—
Texas Biomedical Research Institute	5	8,542,776	5	8,542,776	—	—	—	—
Texas Engineering Experiment Station	1	695,356	1	695,356	—	—	—	—
Texas Heart Institute	2	1,613,322	2	1,613,322	—	—	—	—
Texas Southern University	1	204,359	1	204,359	—	—	—	—
Texas Tech University	1	216,694	1	216,694	—	—	—	—
University of Houston	2	795,778	2	795,778	—	—	—	—
University of North Texas Health Science Center	7	3,680,253	7	3,680,253	—	—	—	—
University of Texas, Austin	4	3,042,451	4	3,042,451	—	—	—	—
University of Texas, El Paso	1	149,500	1	149,500	—	—	—	—
University of Texas Health Science Center, Houston	28	13,164,415	28	13,164,415	—	—	—	—

Institution	No.	Research Training and Career Development							
		Totals		Grants		Development		Contracts	
		Dollar	No.	Dollar	No.	Dollar	No.	Dollar	No.
University of Texas Health Science Center, San Antonio	15	7,361,342	10	3,639,924	3	409,331	2	3,312,087	
University of Texas Health Science Center, Tyler	6	1,992,247	6	1,992,247	—	—	—	—	
University of Texas, MD Anderson Cancer Center	4	1,367,888	4	1,367,888	—	—	—	—	
University of Texas Medical Branch, Galveston	7	4,544,311	6	2,403,388	—	—	1	2,140,923	
University of Texas, Southwestern Medical Center	46	21,444,549	39	20,105,727	6	1,157,822	1	181,000	
University of the Incarnate Word	1	124,149	1	124,149	—	—	—	—	
Total Texas	220	102,873,436	194	89,821,339	20	3,747,529	6	9,304,568	
Utah									
IHC Health Services, Inc.	1	239,228	—	—	—	—	1	239,228	
LDS Hospital	—	23,132	—	—	—	—	—	23,132	
Navigen, Inc.	1	567,237	1	567,237	—	—	—	—	
NuView Life Sciences, Inc.	1	182,972	1	182,972	—	—	—	—	
University of Utah	42	16,651,936	36	15,420,747	4	868,690	2	362,499	
World Heart Corporation	1	554,504	1	554,504	—	—	—	—	
Total Utah	46	18,219,009	39	16,725,460	4	868,690	3	624,859	
Vermont									
University of Vermont and State Agriculture College	28	14,042,740	25	13,082,322	3	960,418	—	—	
Vermont University	1	529,650	—	—	—	—	1	529,650	
Total Vermont	29	14,572,390	25	13,082,322	3	960,418	1	529,650	
Virginia									
American Psychosomatic Society	1	10,000	1	10,000	—	—	—	—	
Barron Associates, Inc.	1	198,426	1	198,426	—	—	—	—	
College of William and Mary	1	214,157	1	214,157	—	—	—	—	
Eastern Virginia Medical School	1	358,750	1	358,750	—	—	—	—	
HemoShear, LLC	1	2,963,069	1	2,963,069	—	—	—	—	
Paragon Technology Group	1	6,952,458	—	—	—	—	1	6,952,458	
System Research and Application	1	2,945,412	—	—	—	—	1	2,945,412	
University of Virginia, Charlottesville	55	19,848,301	50	18,980,644	5	867,657	—	—	
Virginia Commonwealth University	27	10,811,737	23	10,456,201	4	355,536	—	—	
Virginia Polytechnic Institute and State University	3	1,197,117	3	1,197,117	—	—	—	—	
Total Virginia	92	45,499,427	81	34,378,364	9	1,223,193	2	9,897,870	
Washington									
Adaptive TCR Corporation	1	148,999	1	148,999	—	—	—	—	
Battelle Pacific Northwest Laboratories	1	1,671,409	1	1,671,409	—	—	—	—	

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
	3	3,245,235	3	3,245,235	—	—	—	—
Benaroya Research Institute at Virginia Mason	1	374,865	1	374,865	—	—	—	—
DRVision Technologies, LLC	1	175,000	1	175,000	—	—	—	—
EKOS Corporation	23	30,771,355	22	16,091,478	—	—	1	14,679,877
Fred Hutchinson Cancer Research Center	2	632,456	2	632,456	—	—	—	—
Group Health Cooperative	1	400,000	1	400,000	—	—	—	—
Insilicos	1	22,043	1	22,043	—	—	—	—
Institute for Systems Biology	1	342,981	1	342,981	—	—	—	—
Matrexia, LLC	2	766,200	2	766,200	—	—	—	—
Puget Sound Blood Center	—	117,123	—	117,123	—	—	—	—
Seattle Biomedical Research Institute	10	5,292,930	10	5,292,930	—	—	—	—
Seattle Children's Hospital	1	218,735	1	218,735	—	—	—	—
Seattle Institute for Biomedical and Clinical Research	114	68,795,437	98	54,982,796	12	3,288,773	4	10,523,868
Syntrix Biosystems, Inc.	1	226,288	—	—	—	—	1	226,288
Talaria, Inc.	1	811,424	1	811,424	—	—	—	—
University of Washington	1	306,380	1	306,380	—	—	—	—
VA Puget Sound Health Care System, Seattle	4	1,577,223	4	1,577,223	—	—	—	—
VisionGate, Inc.	1	202,356	1	202,356	—	—	—	—
VPDiagnostics, Inc.	1	257,337	1	257,337	—	—	—	—
Washington State University	171	116,355,776	153	87,636,970	12	3,288,773	6	25,430,033
Total Washington								
West Virginia								
West Virginia University	8	2,949,496	7	2,630,478	1	319,018	—	—
Total West Virginia	8	2,949,496	7	2,630,478	1	319,018	—	—
Wisconsin								
Aurora Health Care, Inc.	—	593,121	—	593,121	—	—	—	—
BloodCenter of Wisconsin, Inc.	12	8,821,118	9	5,817,719	1	106,855	2	2,896,544
Board of Regents of the University of Wisconsin System	1	1,734,092	—	—	—	—	1	1,734,092
Medical College of Wisconsin	58	28,381,021	53	27,526,190	5	854,831	—	—
Morgridge Institute for Research, Inc.	1	1,084,148	1	1,084,148	—	—	—	—
University of Wisconsin, Madison	50	25,767,758	42	24,166,598	8	1,601,160	—	—
Total Wisconsin	122	66,381,258	105	59,187,776	14	2,562,846	3	4,630,636
Puerto Rico								
Universidad Central Del Caribe	2	393,073	2	393,073	—	—	—	—
University of Puerto Rico Medical Sciences	2	329,931	2	329,931	—	—	—	—
Total Puerto Rico	4	723,004	4	723,004	—	—	—	—
Total U.S.	5,395	2,649,970,022	4,713	2,284,943,395	511	97,219,215	171	267,907,412

Institution	No.	Dollar	Research Training and Career Development					
			Totals		Grants		Development	
			No.	Dollar	No.	Dollar	No.	Dollar
Canada								
Hospital for Sick Children (Toronto)	1	323,911	1	323,911	—	—	—	—
McMaster University	1	243,775	1	243,775	—	—	—	—
Montreal Heart Institute	1	125,000	1	125,000	—	—	—	—
Ottawa Health Research Institute	1	288,360	1	288,360	—	—	—	—
St. Michael's Hospital	1	180,360	1	180,360	—	—	—	—
University Health Network	1	326,176	1	326,176	—	—	—	—
Total Canada	6	1,487,582	6	1,487,582	—	—	—	—
China								
The George Institute of China	2	600,499	—	—	—	—	2	600,499
Total China	2	600,499	—	—	—	—	2	600,499
Colombia								
Malaria Vaccine Development Center	1	260,787	1	260,787	—	—	—	—
Total Colombia	1	260,787	1	260,787	—	—	—	—
Guatemala								
Instituto de Nutrición de Centro América y Panamá	1	498,819	—	—	—	—	1	498,819
Total Guatemala	1	498,819	—	—	—	—	1	498,819
Hungary								
Institute of Enzymology, Biological Research Center	—	26,730	—	26,730	—	—	—	—
Total Hungary	—	26,730	—	26,730	—	—	—	—
Kenya								
Moi University School of Medicine	1	476,711	—	—	—	—	1	476,711
Total Kenya	1	476,711	—	—	—	—	1	476,711
Peru								
Universidad Peruana Cayetano Heredia	1	537,459	—	—	—	—	1	537,459
Total Peru	1	537,459	—	—	—	—	1	537,459
South Africa								
University of Cape Town	1	493,784	—	—	—	—	1	493,784
Total South Africa	1	493,784	—	—	—	—	1	493,784
Uganda								
Makerere University	—	100,000	—	100,000	—	—	—	—
Total Uganda	—	100,000	—	100,000	—	—	—	—

Institution	Totals		Grants		Research Training and Career Development		Contracts	
	No.	Dollar	No.	Dollar	No.	Dollar	No.	Dollar
United Kingdom								
University College London	1	338,308	1	338,308	—	—	—	—
Total United Kingdom	1	338,308	1	338,308	—	—	—	—
Zimbabwe								
University of Zimbabwe	—	100,000	—	100,000	—	—	—	—
Total Zimbabwe	—	100,000	—	100,000	—	—	—	—
Total, Other	14	4,920,679	8	2,313,407	—	—	6	2,607,272
Grand Total	5,409	\$2,654,890,701	4,721	\$2,287,156,802	511	\$97,219,215	177	\$270,514,684



Appendices

Types of Research Activity

List of Abbreviations and Acronyms

Index



Types of Research Activity

Research Projects

Research Project Grants (R01): To support discrete and specific projects to be performed by one or several investigators in areas of the investigator's particular interests and competencies.

Research Projects (Cooperative Agreements) (U01): To support discrete, circumscribed projects in areas of an investigator's specific interest and competency involving substantial programmatic participation by the NHLBI during performance of the activity.

Research Program (Cooperative Agreement) (U19): To support a research program of multiple projects, requiring a broadly-based, multidisciplinary and often long-term approach, directed toward a specific major objective, common theme, or program goal relevant to the Institute's mission. The award involves substantial programmatic involvement by NHLBI staff to assist investigators during performance of the research activities.

Research Program Projects (P01): To support broadly based, multidisciplinary, often long-term research projects that have specific major objectives or basic themes directed toward a well-defined research program goal. Usually, a relatively large, organized group of researchers conducts individual subprojects, the results of which help achieve objectives of the program project.

Small Research Grants (R03): To provide limited support for extended analyses of research data generated by clinical trials, population research, and demonstration and education studies.

Academic Research Enhancement Awards

(AREA) (R15): To support small-scale research projects conducted by faculty in primarily baccalaureate degree-granting domestic institutions. Awards are for up to \$75,000 for direct costs (plus applicable indirect costs) for periods not to exceed 36 months.

Exploratory/Developmental Grants (R21): To encourage the development of new research activities in heart, lung, and blood diseases and sleep disorders program areas.

Exploratory/Developmental Grant (R33): To provide phase II support for innovative exploratory and developmental research activities initiated under the R21 mechanism.

Clinical Trial Planning Grant (R34): To support the initial development of a clinical trial, including establishment of the research team; development of tools for data management and oversight of the research; and development of a trial design and other essential elements of the study, such as the protocol, recruitment strategies, and procedure manuals.

Method To Extend Research in Time (MERIT) Award (R37): To provide long-term research grant support to investigators whose research competency and productivity are distinctly superior and thus are likely to continue to perform in an outstanding manner. Investigators may not apply for a MERIT award; instead, they are selected by the NHLBI on the basis of their current grant applications and their present and past grant support.

Clinical Planning Grant Cooperative Agreement (U34): To support the initial development of a clinical trial, including establishment of the research team; development of tools for data management and oversight of the research; and development of a trial design and other essential elements of the study, such as the protocol, recruitment strategies, and procedure manuals. The award involves substantial programmatic involvement by NHLBI staff to assist investigators during performance of the research activities.

NIH Director's Pioneer Award (DP1): To support individual scientists of exceptional creativity who propose pioneering approaches to major contemporary challenges in biomedical research.

NIH Director's New Innovator Award (DP2): To support exceptionally creative new investigators who propose highly innovative approaches that have the potential to produce an unusually high impact. The New Innovator Award will emphasize the importance and potential impact of the scientific problem, the novelty and innovativeness of the approach, and the applicant's potential for creative and innovative research.

Linked Research Project Grant (RL1): To support a discrete, specified, circumscribed project that is administratively linked to another project or projects.

Exploratory/Developmental Cooperative Agreements Phase I (UH2): To support the development of new research activities in program areas that are relevant to the Institute's mission. Support is generally restricted in level of support and in time. The award requires substantial programmatic involvement by NHLBI staff to assist investigators during the performance of research activities.

Multi-Component Research Project Cooperative Agreements (UM1): To support large-scale cooperative agreements that involve complex clinical trials with multiple components (e.g., clinical networks that are relevant to the Institute's mission). The components represent a variety of supporting functions and are not independent of the research projects. The award requires substantial programmatic involvement by NHLBI staff to assist investigators during the performance of research activities.

Small Business Technology Transfer (STTR) Grants—Phase I (R41): To support cooperative R&D projects between small business concerns and research institutions, limited in time and amount, to establish the technical merit and feasibility of ideas that have potential for commercialization. Awards are made to small business concerns only.

Small Business Technology Transfer (STTR) Grants—Phase II (R42): To support in-depth development of cooperative R&D projects between small business concerns and research institutions, limited in time and amount, whose feasibility has been established in phase I and that have potential for commercialization. Awards are made to small business concerns only.

Small Business Innovation Research (SBIR) Grants, Phase I (R43): To support projects, limited in time and amount, to establish the technical merit and feasibility of research and development ideas that may ultimately lead to commercial products or services.

Small Business Innovation Research (SBIR) Grants, Phase II (R44): To support research project ideas that have been shown to be feasible in phase I and that are likely to result in commercially marketable products or services.

Research Centers

Exploratory Grants (P20): To support planning for new programs, expansion or modification of existing resources, and feasibility studies to explore various approaches to the development of interdisciplinary programs that offer potential solutions to problems of special significance to the mission of the NHLBI.

Center Core Grants (P30): To support shared resources and facilities for basic, clinical, behavioral, and translational research in the prevention, detection, and treatment of HIV infection and AIDS.

Specialized Centers of Clinically Oriented Research (SCCOR) Grants (P50): To foster multidisciplinary research on clinically relevant questions enabling basic science findings to be applied more rapidly to clinical problems. Research focuses on clinical and basic scientific issues related to diseases and disorders that are relevant to the mission of the NHLBI. The SCCOR program places more emphasis on clinical research than the SCOR program and requires at least 50 percent of the funded projects to be clinical.

National Swine Research and Resource Center (U42): To support a National Swine Research and Resource Center that will serve as a resource for depositing, maintaining, preserving, and distributing swine models for studies of human diseases, as well as cryopreservation, storage, and reconstitution of embryos and germplasm.

Comprehensive Specialized Research Center Grants (U54): To support a large, interrelated biomedical research program focused on a disorder within the Institute's mandate; to initiate and expand community education, screening, and counseling programs; and to

educate medical and allied health professionals concerning problems of diagnosis and treatment of specific diseases such as sickle cell anemia.

Research Career Programs

Mentored Research Scientist Development Award for Minority Faculty (K01): To support underrepresented minority faculty members with varying levels of research experience to prepare them for research careers as independent investigators.

Mentored Scientist Development Award in Research Ethics (K01): To provide support for training in research ethics for health professionals working at academic and other health-related institutions in biomedical, behavioral, or public health research, particularly research involving human participants.

Minority Institution Faculty Mentored Research Scientist Development Award (K01): To support faculty members at minority institutions who have the interest and potential to conduct state-of-the-art research in cardiovascular, pulmonary, or hematologic disease or in sleep disorders.

Independent Scientist Award (K02): To enhance the research capability of promising individuals in the formative stages of their careers of independent research in the sciences related to heart, lung, and blood diseases; blood resources; and sleep disorders.

Academic Award (K07): To support an individual with an academic appointment to introduce or improve a disease curriculum that will enhance the academic or research environment of the applicant institution as well as further the individual's own career. This award series included the Tuberculosis Academic Award, the Sleep Academic Award, and the Nutrition Academic Award. Currently, the Pediatric Transfusion Medicine Academic Award and the Innovators in Hemoglobinopathies Care Career Development Award programs are being supported.

Clinical Investigator Development Award (K08): To provide an opportunity for clinically trained physicians to develop research skills and gain experience in advanced research methods and experimental approaches in basic and applied sciences relevant to cardiovascular, pulmonary, and hematological diseases.

Research Career Development Program in Vascular Medicine (K12): To promote comprehensive clinical research training for physicians wanting to specialize in vascular medicine. The goal is to prepare clinicians for academic roles in mentoring and leadership in clinical research in vascular medicine.

Research Career Development Program in Clinical Hematology (K12): To develop and evaluate multidisciplinary career development programs in clinical hematology research that will equip new academic researchers with the knowledge and skills to address complex problems in blood diseases, transfusion medicine, and cellular therapies.

Research Career Development Program in the Genetics and Genomics of Lung Diseases (K12): To develop multidisciplinary career development programs in genetics and genomics of lung diseases that will equip new investigators with the knowledge and skills to elucidate the etiology and pathogenesis of such diseases.

Clinical Research Career Development Programs in Emergency Medicine (K12): To promote multidisciplinary clinical research training programs in emergency medicine that prepare clinician-scientists for independent research careers and academic leadership roles in emergency medicine. Programs will provide comprehensive research training to evaluate innovative approaches in the diagnosis and clinical management of patients with acute, life-threatening manifestations of cardiovascular, pulmonary, and hematologic diseases and severe trauma in emergency department settings.

Career Enhancement Award for Stem Cell Research (K18): To enable established investigators to acquire new research capabilities in the use of human or animal embryonic, adult, or cord blood stem cells. All candidates must have a sponsor, either within their own or at another institution, who is a well-qualified stem cell expert to serve as a mentor.

NHLBI Career Transition Award (K22): To support the postdoctoral research training of an outstanding individual in an NHLBI intramural laboratory for up to 3 years and subsequently, to support the individual's successful transition from postdoctoral research to an extramural environment as an independent researcher.

Mentored Patient-Oriented Research Career

Development Award (K23): To provide support for career development to investigators who have made a commitment to focus their research endeavors on patient-oriented research.

Midcareer Investigator Award in Patient-Oriented

Research (K24): To provide support for clinicians to allow them “protected time” to devote to patient-oriented research and to act as mentors for beginning clinical investigators.

Mentored Quantitative Research Career

Development Award (K25): To provide support to investigators with quantitative science or engineering backgrounds who have made a commitment to focus their research on basic or clinical biomedicine, bioengineering, bioimaging, or behavioral sciences.

Pathway to Independence (K99/R00): To provide up to 5 years support in two phases to highly promising postdoctoral scientists to pursue research relevant to the Institute. The K99 phase (Career Transition Award) consists of 1 or 2 years of mentored support, followed by the R00 phase (Research Transition Award) of up to 3 years of independent support, which is contingent on securing an independent research position. Award recipients will be expected to compete successfully for independent research grant support from the NIH or other Institutions during the independence phase to ensure continued support and a smooth transition to independence.

Other Research Grants

Cooperative Clinical Research (R10) (U10): To support studies and evaluations of relevant clinical problems. These grants usually involve collaborative efforts among several institutions and principal investigators and are conducted under a formal protocol.

Conference Grants (R13): To support national and international scientific meetings, conferences, or workshops at which research is discussed.

Research Demonstration and Education Projects

(R18): To provide support designed to develop, test, and evaluate health-related activities and to foster application of existing knowledge to the control of heart, lung, and blood diseases and sleep disorders.

Resource-Related Research Projects (R24): To support research projects that will enhance the capability of resources to serve biomedical research in areas related to cardiovascular, lung, and blood health and diseases; blood resources; and sleep disorders.

Education Projects (R25): To provide support for the development and implementation of a program as it relates to a category in one or more of the areas of education, information, training, technical assistance, coordination, or evaluation.

Minority Biomedical Research Support Grants

(S06): To strengthen the biomedical research and research training capability of minority institutions and to assist in increasing the involvement of minority faculty and students in biomedical research.

Research Enhancement Award (SC1): To support individual investigator-initiated research projects aimed at developing researchers at minority-serving institutions to a stage where they can transition successfully to other extramural support.

Pilot Project Award (SC2): To support underrepresented minorities who are at the beginning stages of a research career and interested in testing a new idea or generating preliminary data, or who are more experienced investigators and interested in switching to a different field of research.

Continuing Education Training Grant (T15): To assist professional schools and other public and nonprofit institutions to establish, expand, or improve programs of continuing professional education, especially for programs dealing with new scientific developments.

Resource-Related Research Projects (U24): To support research projects contributing to improvement of the capability of resources to serve biomedical research.

Historical Black College and University Scientist

Award (UH1): To strengthen and augment the human resources at historically black colleges and universities (HBCU) by recruiting an established research scientist into their biomedical or behavioral sciences department; to enhance the career of the recruited research scientist; and to strengthen other HBCU resources for the conduct of biomedical or behavioral research in areas related to cardiovascular, lung, and blood health and disease; transfusion medicine; and sleep disorders.

Individual National Research Service Awards (NRSA)

Individual Predoctoral M.D./Ph.D. NRSA (F30): To provide predoctoral individuals with supervised research training in areas related to heart, lung, and blood diseases; blood resources; sleep disorders leading toward a combined M.D./Ph.D. degree. Training under this award is designed to provide a foundation for a career as a physician-scientist in the areas of interest to the NHLBI.

Predoctoral Individual NRSA (F31): To provide predoctoral individuals with supervised research training in areas related to heart, lung, and blood diseases; blood resources; and sleep disorders leading toward the research degree (e.g., Ph.D.).

Postdoctoral Individual NRSA (F32): To provide postdoctoral research training to individuals to broaden their scientific background and extend their potential for research in areas related to heart, lung, and blood diseases and blood resources.

NRSA for Senior Fellows (F33): To provide experienced scientists with an opportunity to make major changes in the direction of their research careers, to broaden their scientific background, to acquire new research capabilities, to enlarge their command of an allied research field, or to take time from regular professional responsibilities for the purpose of broadening their research capabilities.

Institutional National Research Service Awards (NRSA)

Institutional NRSA (T32): To enable institutions to make awards to individuals selected by them for predoctoral and postdoctoral research training in areas related to heart, lung, and blood diseases; blood resources; and sleep disorders.

Minority Institutional Research Training Program (T32M): To support full-time research training for investigative careers at minority schools in areas of cardiovascular, pulmonary, and hematologic diseases and sleep disorders. Graduate students, postdoctoral students, or health professions students may be supported under this program.

MARC Undergraduate NRSA Institutional Grants

(T34): To support institutional training grants for underrepresented minority undergraduates to obtain research training and improve their preparation for graduate training in the biomedical and behavioral sciences.

NRSA Short-Term Research Training (T35 and T35M): To provide individuals with research training during off-quarters or summer periods to encourage research careers or to encourage research in areas of national need. This program includes the Short-Term Training for Minority Students Program and short-term training for students in health professional schools.

MARC Visiting Professors for Minority Institutions

(T36): To increase the number of well-trained minority scientists in biomedical disciplines and to strengthen the research and teaching capabilities of minority institutions.

Other Support

Research and Development Contracts (N01): To develop or apply new knowledge or test, screen, or evaluate a product, material, device, or component for use by the scientific community.

Small Business Innovation Research (N43): To support projects, limited in time and amount, to establish the technical merit and feasibility of R&D ideas that may ultimately lead to a commercial product(s) or service(s).

NIH Inter-Agency Agreements (Y01): To provide a source of funds to another Federal Agency to acquire specific products, services, or studies.

NIH Intra-Agency Agreements (Y02): To provide a source of funds to another NIH component to acquire specific products, services, or studies.

Minority Research Supplements Programs: To provide supplemental funds to active NHLBI grants to support the research of minority high school, undergraduate, and graduate students; postdoctoral trainees; and investigators.

List of Abbreviations and Acronyms

ACCORDION	Action To Control Cardiovascular Risk in Diabetes Follow-Up	CLRD	chronic lower respiratory diseases
ACE	angiotensin-converting enzyme	COPD	chronic obstructive pulmonary disease
AHEAD	Action for Health in Diabetes	COPTR	Childhood Obesity Prevention and Treatment Research
AIDS	acquired immunodeficiency syndrome	CPHHD	Centers for Population Health and Health Disparities
ALLHAT	Antihypertensive and Lipid-Lowering Treatment To Prevent Heart Attack Trial	CVD	cardiovascular diseases
AMI	acute myocardial infarction	DARD	Division for the Application of Research Discoveries
ARDSNet	Acute Respiratory Distress Syndrome Clinical Network	DASH	Dietary Approaches To Stop Hypertension
ARIC	Atherosclerosis Risk in Communities	DBDR	Division of Blood Diseases and Resources
BABY HUG	Pediatric Hydroxyurea Phase III Clinical Trial	DCVS	Division of Cardiovascular Sciences
BEE	Board of Extramural Experts	DERA	Division of Extramural Research Affairs
BTRP	Basic and Translational Research Program	DIR	Division of Intramural Research
CABG	coronary artery bypass graft	DLD	Division of Lung Diseases
CADET	Centers for Advanced Diagnostics and Experimental Therapeutics in Lung Diseases	EARLY	Early Adult Reduction of Weight Through LifestYle Intervention
CARDIA	Coronary Artery Risk Development in Young Adults	EDTA	ethylene diamine tetra-acetic acid
CDC	Centers for Disease Control and Prevention	FY	fiscal year
CF	cystic fibrosis	GOCADAN	Genetics of Coronary Artery Disease in Alaska Natives
CHART	Consortium of Hospitals Advancing Research on Tobacco	GWAS	genome-wide association study
CHD	coronary heart disease	HCHS	Hispanic Community Health Study
		HEW	Department of Health, Education, and Welfare (now HHS)

HHS	Health and Human Services (formerly HEW)	ORBIT	Obesity Related Behavior Intervention Trials
HIV	human immunodeficiency virus	OSA	obstructive sleep apnea
ICD	International Classification of Diseases	PA	Program Announcement
JHS	Jackson Heart Study	PACT	Production Assistance for Cellular Therapies
LOTT	Long-Term Organ Treatment Trial	PAR	Program Announcement with special receipt, referral, or review
MARC	Minority Access to Research Careers	PHS	Public Health Service
MESA	Multi-Ethnic Study of Atherosclerosis	RFA	Request for Applications
NAEPP	National Asthma Education and Prevention Program	RFP	Request for Proposals
NCEP	National Cholesterol Education Program	RPG	research project grant
NCHS	National Center for Health Statistics	RuSH	Registry and Surveillance System in Hemoglobinopathies
NCSDR	National Center on Sleep Disorders Research	SBIR	Small Business Innovation Research
NHANES	National Health and Nutrition Examination Survey	SCD	sickle cell disease
NHBPEP	National High Blood Pressure Education Program	SCCOR	Specialized Center of Clinically Oriented Research
NHI	National Heart Institute	SCOR	Specialized Center of Research
NHLBAC	National Heart, Lung, and Blood Advisory Council	SDB	sleep disordered breathing
NHLBI	National Heart, Lung, and Blood Institute (formerly NHI and NHLI)	SES	socioeconomic status
NHLI	National Heart and Lung Institute	SIDS	sudden infant death syndrome
NICHHD	National Institute of Child Health and Human Development	SOYA	Study of Soy Isoflavones in Asthma
NIH	National Institutes of Health	SPRINT	Systolic Blood Pressure Intervention Trial
NINDS	National Institute of Neurological Disorders and Stroke	STAN	Study of Asthma and Nasal Steroids
NRSA	National Research Service Award	STTR	Small Business Technology Transfer
		TB	tuberculosis
		WHI	Women's Health Initiative

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*Discrimination Prohibited:
Under provisions of applicable
public laws enacted by
Congress since 1964, no person
in the United States shall, on
the grounds of race, color,
national origin, handicap, or
age, be excluded from participa-
tion in, be denied the bene-
fits of, or be subjected to dis-
crimination under any program
or activity (or, on the basis of
sex, with respect to any educa-
tion program or activity)
receiving Federal financial
assistance. In addition,
Executive Order 11141 pro-
hibits discrimination on the
basis of age by contractors and
subcontractors in the perform-
ance of Federal contracts,
and Executive Order 11246
states that no federally funded
contractor may discriminate
against any employee or appli-
cant for employment because
of race, color, religion, sex, or
national origin. Therefore,
the Heart, Lung, and Blood
Institute must be operated in
compliance with these laws
and Executive Orders.*



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