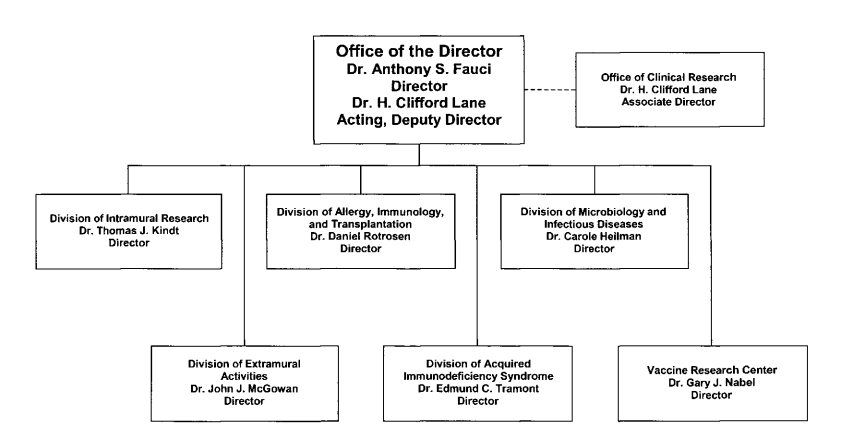
#### DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### NATIONAL INSTITUTES OF HEALTH

#### National Institute of Allergy and Infectious Diseases

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# National Institutes of Health National Institute of Allergy and Infectious Diseases Organizational Structure



#### NATIONAL INSTITUTES OF HEALTH

National Institute of Allergy and Infectious Diseases

For carrying out section 301 and title IV of the Public Health Service Act with respect to allergy and infectious diseases, [\$4,440,007,000] \$4,459,395,000: Provided, That \$100,000,000 maybe made available to International Assistance Programs, "Global Fund to Fight HIV/AIDS, Malaria, and Tuberculosis", to remain available until expended: Provided further, That up to [\$150,000,000] \$30,000,000 shall be for extramural facilities construction grants to enhance the Nation's capability to do research on biological and other agents. (Department of Health and Human Services Appropriations Act, 2005.)

## National Institutes of Health National Institute of Allergy and Infectious Diseases

**Amounts Available for Obligation 1/** 

		<del></del>	
Source of Funding	FY 2004 Actual	FY 2005 Appropriation	FY 2006 Estimate
Appropriation	\$4,335,155,000	\$4,440,007,000	\$4,459,395,000
Enacted Rescissions	(30,593,000)	(37,166,000)	0
Subtotal, Adjusted Appropriation	4,304,562,000	4,402,841,000	4,459,395,000
Real transfer to the Global Fund to fight HIV/AIDS, Malaria, and Tuberculosis	(149,115,000)	(99,200,000)	0
Real transfer under NIH Director's one-percent transfer authority to other ICs	(13,678,000)	0	0
Comparative transfer to NIBIB for Radiology Program	(321,000)	0	0
Comparative transfer to Buildings and Facilities	(1,201,000)	0	0
Comparative transfer to/from other NIH ICs for NIH Roadmap	13,678,000	0	0
Comparable transfer from the Global Fund to fight HIV/AIDS, Malaria, and Tuberculosis	149,115,000	99,200,000	0
Subtotal, adjusted budget authority	4,303,040,000	4,402,841,000	4,459,395,000
Unobligated Balance, start of year	0	0	0
Unobligated Balance, end of year	0	0	0
Subtotal, adjusted budget authority	4,303,040,000	4,402,841,000	4,459,395,000
Unobligated balance lapsing	0	0	0
Total obligations	4,303,040,000	4,402,841,000	4,459,395,000

<sup>&</sup>lt;u>1</u>/ Excludes the following amounts for reimbursable activities carried out by this account: FY 2004 - \$4,867,000 FY 2005 - \$5,969,000 FY 2006 - \$7,320,000 Excludes \$7,896,139,000 in FY 2005 and \$9,880,377,000 in FY 2006 for royalties.

#### **Justification**

#### **National Institute of Allergy and Infectious Diseases**

Authorizing Legislation: Section 301 of the Public Health Service Act, as amended.

Reauthorizing legislation will be submitted.

**Budget Authority:** 

•	FY 2004	FY	2005	FY	2006	Increas	e or
	<u>Actual</u>	<u>Approp</u>	<u>oriation</u>	<u>Est</u>	timate	Decrea	ase
<u>FTEs</u>	BA	<u>FTEs</u>	<u>BA</u>	<u>FTEs</u>	BA	FTEs	BA
.1.443	\$4.303.040.000	1.507	\$4,402.841.00	1.507	\$4.459.395,00	00 0	\$56.554.000

This document provides justification for the Fiscal Year 2006 research activities of the National Institute of Allergy and Infectious Diseases (MAID), including HIV/AIDS activities. A more detailed description of NIH-wide Fiscal Year 2006 HIV/AIDS activities can be found in the NTH section entitled "Office of AIDS Research (OAR)."

#### INTRODUCTION

The National Institute of Allergy and Infectious Diseases (NIAID) is a component of the National Institutes of Health (NIH), which is an agency of the Department of Health and Human Services (DHHS). NIAID supports basic and applied research to prevent, diagnose, and treat infectious and immune-mediated illnesses, including human immunodeficiency virus/acquired immunodeficiency syndrome (HIV/AIDS) and other sexually transmitted diseases, illness from agents with bioterrorism potential, tuberculosis, malaria, autoimmune disorders, asthma, and allergies. NIAID research has led to new therapies, vaccines, diagnostic tests, and other technologies that have improved the health of millions of people in the United States and around the world. The scope of the NIAID research portfolio has expanded considerably in recent years in response to new challenges such as bioterrorism, emerging and re-emerging infectious diseases, and the increase in asthma prevalence among children in this country. The growth of NIAID programs has also been driven by unprecedented scientific opportunities in the core NIAID scientific disciplines of microbiology, immunology, and infectious diseases. Advances in these key fields have led to a better understanding of the human immune system and the mechanisms of infectious and immune-mediated diseases.

#### BIODEFENSE: RESPONDING THROUGH RESEARCH

A terrorist attack on the United States using biological agents, once thought to be a remote possibility, occurred in the fall of 2001, when anthrax spores were sent through the mail. Recent events have raised awareness of both the possibility of a bioterrorist attack and the vulnerability of the U.S. population to such an event. In 2003 and 2004, the toxin ricin was found in an envelope at a postal facility in South Carolina, a U.S. Senate Office Building, and in several jars of baby food in California.

The threat of bioterrorism has created new challenges for medicine and public health. The nation's ability to detect and respond to acts of bioterror requires new and improved countermeasures, including diagnostics, vaccines, and therapies. Although the Department of Defense (DoD) has developed defenses for biological warfare, there are additional concerns that need to be addressed to provide an adequate civilian defense from a bioterrorist attack.

As the lead agency at NIH for infectious diseases and immunology research, NIAID has set research priorities and goals for each microorganism that might be used as an agent of bioterrorism, with particular emphasis on "Category A" agents—those considered by the Centers for Disease Control and Prevention to be the worst bioterror threats. NIAID Category B and C priority pathogens, in general, cause milder disease or fewer deaths than Category A agents and are more difficult to disseminate in populations. NIAID has developed the NIAID Strategic Plan for Biodefense Research'; the NIAID Biodefense Research Agenda for CDC Category A Agents'; and the NIAID Biodefense Research Agenda for Category B and C Priority Pathogens'. Advances in biodefense research have been rapid and significant, as outlined in the NIAID Biodefense Research Agenda for CDC Category A Agents Progress Report' and the NIAID Biodefense Research Agenda for CDC Category B and C Priority Pathogens Progress Report'.

NIAID's biodefense program includes both short- and long-term research targeted at the design, development, and evaluation of the specific public health tools or countermeasures (diagnostics, therapies, and vaccines) needed to control a bioterrorist-caused outbreak. Major biodefense research projects include: ongoing advanced development of products for protecting the public from CDC Category A agents, including better vaccines for anthrax and smallpox; large-scale genomic sequencing of agents, such as the bacteria that cause anthrax and plague, to help identify vulnerabilities and target them with drugs or vaccines; and implementation of major research resources initiated in 2003.

It is anticipated that the large investment in research on biodefense will have many positive spin-offs for other diseases. NIAID research on microbial biology and on the pathogenesis of organisms with bioterror potential will certainly lead to a better understanding of other more common and naturally occurring infectious diseases. In particular, the advancement of knowledge should have an enormous positive impact on our ability to diagnose, treat, and prevent major infectious killers, such as malaria, tuberculosis, HIV/AIDS, and a spectrum of emerging and re-emerging diseases, such as West Nile disease, dengue, influenza, and multi drug-resistant infections. Furthermore, and importantly, the NIAID biodefense research agenda promises to enhance our understanding of the molecular and cellular mechanisms of the immune system. Such knowledge will help in the search for new ways to treat and prevent a variety of immune-mediated diseases, such as type 1 diabetes and rheumatoid arthritis. New insights into the mechanisms of regulation of the human immune system will impact research on cancer, immune-mediated neurological diseases, and allergic and hypersensitivity diseases.

NIAID Strategic Plan for Biodefense Research, <a href="http://www2.niaid.nih.gov/Biodefense/Research/strategic.pdf">http://www2.niaid.nih.gov/Biodefense/Research/strategic.pdf</a>, (accessed December 6, 2004).

<sup>&</sup>lt;sup>2</sup> NIAID Biodefense Research Agenda for CDC Category A Agents, http://www2.niaid.nih.gov/Biodefense/Research/biotresearchagenda.pdf, (accessed December 6, 2004).

<sup>&</sup>lt;sup>3</sup> NIAID Biodefense Research Agenda for Category B and C Priority Pathogens, http://www2.niaid.nih.gov/Biodefense/Research/categorybandc.pdf, (accessed December 6, 2004).

<sup>&</sup>lt;sup>4</sup> NIAID Biodefense Research Agenda for CDC Category A Agents Progress Report, http://www2.niaid.nih.gov/Biodefense/Research/category A Progress Report.pdf, (accessed December 6, 2004).

<sup>&</sup>lt;sup>5</sup> NIAID Biodefense Research Agenda for Category B and C Priority Pathogens Progress Report, http://www2.niaid.nih.gov/Biodefense/Research/category\_bc\_progress\_report.pdf, (accessed December 6, 2004).

### CATEGORY A AGENTS: DEVELOPMENT OF BIOMEDICAL COUNTERMEASURES AND OTHER RESEARCH PROGRESS

#### Anthrax

NIAID is aggressively pursing the advanced development of a new anthrax vaccine suitable for civilian populations of varying age and health status. NIAID is developing a next-generation vaccine based on a recombinant form of the anthrax protective antigen (rPA). Two new contracts were awarded to support the production, testing, and evaluation of lots for consistency of rPA vaccine, including a phase II trial.

In the event of a bioterrorism incident, effective therapeutics will be needed to address the immediate health needs of the public. Through the *In Vitro and Animal Models for Emerging Infectious Diseases and Biodefense* program, NIAID is screening existing FDA-approved antimicrobials and immunomodulators for efficacy against inhalational anthrax. Five licensed antibiotics have been selected for study, with ciprofloxacin (Cipro®) as a control. NIAID also is pursuing studies to determine whether the course of antibiotic therapy can be decreased by vaccinating subjects with the rPA vaccine candidates currently under development. In addition, NIAID is supporting the development of two versions of anthrax monoclonal antibodies as potential antitoxin therapies.

NIAID also supports basic research on anthrax. In FY 2004, NIAID-funded scientists determined the three-dimensional structure of the cell-binding component of anthrax toxin, called protective antigen (PA), bound tightly to a target cell surface protein called CMG-2. This discovery offers a precise, finely detailed snapshot of a crucial step in the pathway that allows anthrax toxin to enter human cells. This work provides important new leads for the development of novel antitoxins.

A successful response to a bioterrorist threat requires diagnostics to quickly and efficiently identify the pathogen(s) involved. A team of scientists co-funded by NIAID and DoD have developed an assay to simultaneously detect, in a single sample, three Category A Priority Pathogens, *Bacillus anthracis, Yersinia pestis* and *Francisella tularensis*, and one Category B Priority Pathogen, *Burkholderia mallei*.

#### Smallpox and Other Orthopox Viruses

Smallpox is caused by the variola major virus, a member of the orthopox family of viruses. It is among the most dangerous potential biological weapons because the virus easily spreads from person-to-person, no effective treatment exists, and few people are fully immune to the virus.

The vaccine used to achieve the eradication of smallpox from the human population was based on a live, attenuated strain of vaccinia, a virus related to variola major, the virus that causes smallpox. Previous NIAID-sponsored clinical trials demonstrated that the existing stocks of the smallpox vaccine DryVax® could be diluted five-fold and still elicit a potent immune response. In 2004, another stock of vaccine that was manufactured by Aventis Pasteur in the 1950s and stored as a frozen liquid was shown to elicit a robust immune response, even when diluted tenfold. The results from these two NIAID-sponsored clinical trials, along with the recent manufacture and purchase of 225 million doses of a cell-cultured version of DryVax® secured under contract by DHHS, indicate that the smallpox vaccine stocks currently available would be sufficient to vaccinate the entire U.S. population in an emergency.

NIAID is vigorously pursing the advanced development of the next generation smallpox vaccine, modified vaccinia ankara (MVA) vaccine, an attenuated poxvirus designed to protect against variola major, the virus that causes smallpox. A 2004 study conducted by NIAID intramural researchers indicated that MVA is nearly as effective as the DryVax® vaccine in protecting monkeys against monkeypox, an animal model of human smallpox. Furthermore, NIAID researchers found that, in addition to protecting healthy mice against a lethal form of the vaccinia virus, MVA protects mice with immune deficiencies. In FY 2004, NIAID awarded two three-year contracts for the production and testing of two MVA vaccine candidates.

In an effort to identify treatments for smallpox infection, over 1,500 compounds, including most of the licensed antiviral drugs, have been evaluated for anti-poxvirus activities in cell culture. The approximately 40 compounds that were active *in vitro* were tested in animal models. The antiviral drug cidofovir was shown to have potential as a therapy for both smallpox and generalized vaccinia, a potential side-effect of smallpox vaccination. In FY 2004, a NIAID-sponsored study indicated that cidofovir applied topically may be more effective than oral or intravenous administration to control vaccinia lesions. NIAID is also evaluating immunotherapy with monoclonal antibodies against vaccinia.

#### Ebola and Other Viral Hemorrhagic Fevers

Viral hemorrhagic fevers are caused by four distinct viral families: arenaviruses, such as Lassa virus; bunyaviruses, such as hantavirus; flaviviruses, such as dengue virus; and filoviruses, such as Ebola and Marburg viruses. As a group, these diseases are characterized by hemorrhaging that begins several days after the sudden onset of high fever, muscle and abdominal pain, and extreme fatigue. NIAID, in cooperation with corporate and Federal partners, has a robust program to develop a vaccine against Ebola. As part of a cooperative program with the U.S. Army Medical Research Institute of Infectious Diseases (USAMRIID), NIAID scientists at the Vaccine Research Center (VRC) and their collaborators developed an effective vaccination strategy that protects monkeys against Ebola viral hemorrhagic fever. The first phase I trial in humans of a DNA vaccine to prevent Ebola infection has been initiated and is currently fully enrolled. The VRC is also developing an adenoviral Ebola vaccine through a Cooperative Research and Development Agreement (CRADA) with industrial partners. A single dose of this vaccine was recently shown to protect monkeys from Ebola infection. NIAID researchers have also developed candidate vaccines against each of the four dengue virus strains. No treatment for Ebola infection or other hemorrhagic fevers currently exists, although candidate drug screening is under way. NIAID also has supported the evaluation of hundreds of compounds for their in vitro activity in models for hemorrhagic fever viruses such as Yellow Fever, Pichinde virus (a surrogate for Lassa), and Punta Toro virus (a surrogate for hantavirus). Approximately 240 compounds were screened in FY 2004.

#### **Botulism**

Botulinum toxin, the cause of the disease known as botulism, is by weight the most toxic substance known—a dose of less than ten millionth of a gram is fatal to humans about 50 percent of the time. The toxin is produced by the common soil bacterium *Clostridium botulinum*. NIAID investigators recently described the A2 neurotoxin gene cluster in *C. botulinum*. Further characterization and understanding of the serotype A toxins is essential to developing countermeasures that will be broadly protective.

NIAID supports several initiatives to develop countermeasures against botulinum toxin. Through its *Food and Waterborne Diseases Integrated Research Network*, NIAID funds the development

of novel therapeutics to neutralize botulinum toxins in the blood or within neuronal cells. In addition, NIAID supports fast track development of monoclonal antibody-based therapies for botulinum neurotoxin serotype A by funding the manufacture of monoclonal antibodies that will be used in preclinical and clinical studies. Moreover, a recombinant botulinum vaccine is under development.

#### Tularemia

Tularemia is a potential bioterrorist agent because of its high level of infectivity and its ability to be aerosolized. Inhalation of as few as ten bacteria can cause disease. NIAID is collaborating with USAMRIID to develop clinical laboratory methods for working with tularemia, and with the DoD Joint Vaccine Acquisition Program to conduct planned safety and efficacy clinical trials of a newly manufactured, modernized version of the Soviet LVS tularemia vaccine. Proposals for the coordinated development of research tools needed to identify and evaluate new candidates for a safe, effective, general-use tularemia vaccine were recently solicited, and awards are scheduled to be made in FY 2005.

#### Plague

Plague is caused by the bacterium *Yersinia pestis*. Historically, plague has occurred in sporadic but severe epidemics, including the "Black Death" that occurred in Europe during 14<sup>th</sup> century. The bacterium is usually transmitted from infected animals to humans by insects, especially fleas. Inhalation of *Y. pestis* can cause a pneumonic form of the disease, which can be transmitted from person to person and is nearly always fatal.

One of the barriers to development of countermeasures against plague has been the lack of animal models that could allow studies of *Y. pestis* transmission and pathogenesis. In FY 2004, NIAID scientists developed a flea-to-mouse model of flea-borne transmission of plague. The model was used to test a recombinant plague vaccine candidate developed by USAMRIID. The vaccine, called F1-V, was found to protect mice against a flea-borne plague challenge.

In FY 2004, NIAID awarded a contract to develop recombinant plague vaccine candidates, manufacture and test the vaccines, and conduct clinical studies in healthy populations. In addition, NIAID established a cooperative program with USAMRIID to test FDA-approved antibiotics for efficacy against pneumonic plague in monkeys.

NIAID Category B and C priority pathogens, in general, cause milder disease or fewer deaths than Category A agents and are more difficult to disseminate in populations. Category B agents include: inhalational bacteria, toxins, food- and water-borne pathogens, and arthropod-borne viruses. Category C agents include all emerging infectious disease threats. Significant progress has been made since the January 2003 release of the *NIAID Biodefense Research Agenda for Category B and C Priority Pathogens*. In June 2004, the *NIAID Biodefense Research Agenda for Category B and C Priority Pathogens Progress Report* was released.

Several NIAID-sponsored research programs relevant to Category B pathogens and toxins have been established and are currently under way, including work to develop a vaccine against the toxin ricin. In addition, NIAID-funded researchers have identified and cloned multiple proteins

<sup>&</sup>lt;sup>6</sup> NIAID Biodefense Research Agenda for Category B and C Priority Pathogens, <a href="http://www2.niaid.nih.gov/Biodefense/Research/categorybandc.pdf">http://www2.niaid.nih.gov/Biodefense/Research/categorybandc.pdf</a>, (accessed December 6, 2004).

NIAID Biodefense Research Agenda for Category B and C Priority Pathogens Progress Report, <a href="http://www2.niaid.nih.gov/Biodefense/Research/category-bc-progress-report.pdf">http://www2.niaid.nih.gov/Biodefense/Research/category-bc-progress-report.pdf</a>, (accessed December 6, 2004).

from the Category B pathogen *Coxiella burnetii*, the causative agent of Q fever, that react with infection-derived antibodies. These proteins are logical candidates for future vaccines against Q fever.

Food and water are potentially important routes for the dissemination of infectious agents by terrorists. NIAID is expanding its capacity to conduct translational research, including the development of diagnostics, vaccines, and therapies for food- and waterborne diseases. NIAID's programs have begun to yield results that should aid in ultimately developing medical countermeasures to combat these diseases. For example, in FY 2004, NIAID-supported research led to the determination of the entire genetic sequence of *Cryptosporidium hommis*, a common contaminate of public water systems. In addition, proteins from *Vibrio cholerae*, the causative agent of cholera, which are recognized by the human immune system when it mounts a protective response, were identified. Furthermore, transgenic mice engineered to express defensin, an antibiotic peptide found in the human intestine, were shown to be highly resistant to *Salmonella typhimurium*, a dangerous and sometimes fatal food-borne bacterium. Finally, a protocol is under development for a phase I study of a candidate *Shigella flexneri* vaccine.

#### UNDERSTANDING, ASSESSING, AND ENHANCING HOST IMMUNITY

NIAID supports research into both innate and adaptive immune responses, which may provide insights that lead to the development of new or improved interventions against agents of bioterror. Innate immune responses are nonspecific defense mechanisms that come into play soon after a pathogen enters the body (e.g., physical barriers such as the skin, chemicals in the blood, and immune system cells that attack foreign cells in the body). In contrast, adaptive immune responses are highly specific to a pathogen and are carried out by white blood cells called B and T cells. NIAID-supported scientists recently identified a mechanism through which the body alerts its immune system to an invading pathogen. They discovered that uric acid, which is released from dying cells, acts as a powerful signal to activate the earliest pathways of immunity. This discovery may lead to new immune-boosting strategies.

Findings by other researchers supported through NIAID's *Innate Immune Receptors and Adjuvant Discovery Program* indicate that the magnitude and type of immune response to vaccination can be manipulated, for example, by altering the route of immunization. These discoveries will aid in designing more effective vaccines for specific pathogens.

NIAID continues to strengthen its portfolio of research aimed at understanding the host immune system and how it responds to agents of bioterror through a broad range of initiatives. For example, NIAID supports basic, clinical, and applied research on human immune responses to all categories of agents of bioterror through its eight *Cooperative Centers for Translational Research on Human Immunology and Biodefense*. Little is known about why people respond differently to pathogens. To address this knowledge gap, NIAID recently awarded six contracts to identify associations between specific immune responses and variations in genes. Furthermore, NIAID awarded 14 contracts to support the *Large-Scale Antibody and T Cell Epitope Discovery Program*, which is aimed at identifying the regions (epitopes) of select infectious agents that elicit immune reactions.

#### RESEARCH CENTERS AND OTHER SPECIALIZED RESEARCH RESOURCES

In 2003, NIAID created a nationwide network of eight multidisciplinary *Regional Centers of Excellence (RCE) for Biodefense and Emerging Infectious Diseases Research.* These centers conduct research on NIAID Category A - C agents designed to advance the development of

diagnostics, therapeutics, and vaccines. In June 2004, NIAID announced that it intends to fund the creation of two additional RCEs; awards are anticipated for 2005.

In 2003, NIAID supported the creation of two National Biocontainment Laboratories (NBLs). These facilities, when built, will include new Biosafety Level 4 (BSL-4) laboratory space designed to safely contain the most dangerous pathogens known. NIAID also funded the construction of nine Regional Biocontainment Laboratories (RBLs). These facilities, distributed geographically throughout the country, will include new BSL-3 laboratories. In 2004, NIAID announced a new initiative to develop five to eight additional RBLs; awards are anticipated in FY 2005. In addition to the construction of extramural biocontainment facilities, NIAID has begun the construction of three intramural research facilities that will house biocontainment laboratories. NIAID has also initiated a program to fund the renovation and upgrade of existing biocontainment laboratories.

NIAID has made a significant investment in the genomic sequencing of microorganisms that are relevant to national biodefense. By the end of FY 2004, the complete genome of at least one strain of each Category A, B, and C agent has been sequenced through the combined efforts of public and private investigators. Projects are ongoing to acquire the genomic sequences of at least one additional strain of every bacterium, virus, or protozoan on the list of Category B and C pathogens.

NIAID also supports programs that supply vital research tools to biodefense researchers nationwide and provide a range of resources for preclinical testing of new therapies and vaccines in both cell culture and animal models, including nonhuman primates. In FY 2004, several new animal models for Category A-C Priority Pathogens were developed: two new animal models for viral hemorrhagic fevers, a model of flea-borne plague transmission, and two models for West Nile virus.

#### **Future Directions in Biodefense Research**

In FY 2006, NIH will continue to expand its research related to potential agents of bioterrorism as part of a broad research agenda involving other agencies within DHHS and DoD. Since 2001, NIAID has launched dozens of biodefense research initiatives, all with the overarching goal of facilitating the creation of new therapies, diagnostic tests, and vaccines that will allow the United States to mount a successful medical and public health response to a biological attack on the civilian population, should such a terrible event occur.

Continued cooperation and coordination with the pharmaceutical industry will be vital to the success of the biodefense research program as scientific advances are translated into new countermeasures that will be available in an emergency. To this end, Project BioShield legislation, which was signed into law in July 2004<sup>s</sup>, will help expedite the conduct of NIH research and development on medical countermeasures. Shortly after the signing of the Project BioShield legislation, NIAID announced several new initiatives to expand biodefense research and product development. These include *Therapeutics for CDC Category A Agents: BioShield Accelerated Countermeasure Development*, which will support projects and studies needed to obtain investigational new drug (IND) status for countermeasure candidates; two initiatives aimed at the development of countermeasures against botulism: *Neutralizing Monoclonal Antibodies for Type A Botulinum Neurotoxins*, which will support fast track development of monoclonal antibody-based therapy for botulinum toxin by funding manufacture of monoclonal

<sup>&</sup>lt;sup>8</sup> Public Law 108-276, Project BioShield Act of 2004, 42 USC 201, July 21, 2004.

antibodies for evaluation in preclinical and clinical studies, and *Recombinant Type E Botulinum Neurotoxin Vaccine*, which will support fast track development of the recombinant botulinum neurotoxin serotype E vaccine through support of the manufacturing of clinical grade lots for evaluation in preclinical and clinical studies; and *Protecting the Immune System Against Radiation: BioShield Accelerated Product Development.* In FY 2006, NIAID will continue to use its BioShield authorities to launch new biodefense initiatives will be launched in FY 2006. *Drug Development Resources for Antiinfectives* will support and accelerate development of antimicrobials by providing preclinical drug development resources to the scientific community and industry partners. Other biodefense initiatives to be unveiled in FY 2006 include *Cooperative Research Partnerships for Biodefense*, which will aim at accelerating the development of promising medical countermeasures through preclinical and early clinical stages by means of collaborative partnerships with academia and industry. In addition, other new initiatives will support development of vaccines against agents of bioterror, including *Development of Rift Valley Fever Vaccines* and *Development of a Multivalent Recombinant Botulinum Vaccine*.

In FY 2006, NIAID will continue to promote research projects through NIAID-supported genomic networks that take advantage of the availability of microbial and human genome sequence data and examine the functional analyses of gene and protein expression on a genomic scale.

NIAID will also continue to support initiatives aimed at garnishing a greater understanding of the immune response to pathogens. In FY 2006, NIAID will launch the initiative *Innate Immunity to NIAID Category B Protozoan Pathogen-Associated Molecular Patterns* and will continue to support the following FY 2005 initiatives: *Modeling Immunity for Biodefense*, which supports multi-disciplinary centers to develop novel or improved highly predictive mathematical models that simulate immune function; *Disabling Innate Immune Evasion: New Attenuated Vaccines;* and *Immune Function and Biodefense in Children, Elderly and Immunocompromised Populations*.

#### CONFRONTING INFECTIOUS DISEASES

Infectious diseases have afflicted humanity since its inception, and they will continue to be a menace as long as man and microbes coexist. For example, since AIDS was first recognized in 1981, this emerging disease has spread relentlessly throughout the world. It now threatens to surpass in total fatalities both the "Black Death" of the 14th century and the influenza pandemic of 1918-1919, two other emerging infections that each killed tens of millions of people. In the past five years alone, West Nile and monkeypox viruses emerged in the United States, while Asia experienced an unprecedented number of human infections with avian influenza viruses and the emergence of a new infectious disease, SARS.

#### MAJOR INTERNATIONAL KILLERS

#### HIV/AIDS

The human immunodeficiency virus (HIV), the virus that causes acquired immunodeficiency syndrome (AIDS), destroys a person's immune system over many years, making the infected individual highly susceptible to multiple infections and certain cancers. Despite recent progress in treatment and prevention, HIV/AIDS continues to exact an enormous toll throughout the world. Estimates on the scope of the HIV/AIDS pandemic are profoundly sobering. At the end of 2003, an estimated 40 million people worldwide were living with HIV/AIDS, five million people

worldwide were newly infected with HIV, and three million people with HIV/AIDS had died in the last year. To help turn the tide of the global HIV/AIDS pandemic, NIAID established research collaborations with international colleagues in more than 50 countries focused on HIV vaccine development and other prevention activities, therapeutics, and care for the HIV-infected person. These collaborations already have yielded important results, notably in developing methods to reduce mother-to-child-transmission of HIV.

NIAID-supported investigators have made critical discoveries about the basic biology of HIV and the immune response to HIV infection, which have led to the development of therapies that suppress the growth of the virus. Although much has been learned in recent years, questions remain about the molecular interactions involved in the regulation of HIV expression and replication, why the host immune response fails to control the infection, and how reservoirs of virus persist in the body despite highly active antiretroviral treatment (HAART). NIAID continues to search for more scientific information about how the virus attacks the body and how the body defends itself, which is critical for identifying additional targets for therapeutic interventions and vaccines.

#### Science Advances in HIV/AIDS Research

<u>HIV Patients Get Long-Term Boost with Short, Intermittent Drug Regimen.</u> NIAID scientists demonstrated that brief, widely-spaced courses of the experimental immune-boosting drug interleukin-2 (IL-2) allow people with HIV to maintain near normal levels of CD4+ T cells, a type of immune cell, for long periods. These data provide strong evidence that IL-2 therapy, which can be self-administered by patients, could be an important adjunct to a type of HIV treatment called highly active antiretroviral therapy (HAART).

Weekly Cycles of Once-Daily Antiviral Drugs Could Reduce Cost of HIV Treatment. NIAID researchers have shown that it may be feasible to treat HIV-infected patients with a simple, once-daily regimen of anti-HIV drugs given in pre-planned, 7-day-on, 7-day-off cycles. This approach, used with well-chosen drug regimens in settings where patient adherence is high, could be a powerful and cost-effective tool in treating HIV-infected individuals.

Investigational DNA Vaccines for HIV Show Promise. NIAID scientists at the Vaccine Research Center have developed a novel DNA vaccine for HIV directed at the three most globally important clades or subtypes. The vaccine incorporates genetic material from clades A, B, and C, which cause about 90 percent of all HIV infections worldwide. The genes used for the development of this vaccine were *env*, which encodes a protein on the surface of HIV, and *gag*, *pol*, and *nef*, which encode internal proteins. Initial tests of combinations of the DNA segments carrying the HIV genes showed promising immune responses in non-human primates. Such tests also suggest that a multigene, multiclade HIV DNA vaccine is feasible because the immune responses to individual genes in the vaccine are not reduced when combined with one another. This candidate vaccine is the first multigene, multiclade vaccine to enter human clinical trials.

<u>HIV Protein Vif Subverts Host Cell Antiviral Defenses.</u> NIAID-supported investigators have uncovered a novel pathway through which HIV evades the counterattack mounted against it by a host cell. An HIV protein called Virion Infectivity Factor (Vif) had previously been shown to be essential for viral replication. Vif works by suppressing the anti-HIV activity of APOBEC3G, a host protein. To better understand how Vif counteracts the antiviral function of APOBEC3G,

<sup>&</sup>lt;sup>o</sup> Joint United Nations Programme on HIV/AIDS (UNAIDS), *UNAIDS 2004 Report on the global AIDS epidemic*, Switzerland, 2004.

NIAID-supported investigators isolated and identified the host proteins with which V i f associates during infection. Their studies revealed that the interaction of Vif, APOBEC3G, and the host protein Cullin5 form a complex known as Cul5-SCF. In turn, Cul5-SCF induces the degradation of APOBEC3G. The identification of interventions that either modulate levels of APOBEC3G or block its interaction with V i f through the Cul5-SCF complex could lead to new and innovative strategies for treating HIV infection.

GB Virus C Infection Inhibits HIV Replication. HIV patients are commonly coinfected with other pathogens, such as the hepatitis C virus (HCV). Coinfection generally contributes to AIDS mortality. In contrast, men infected with both HIV and an apparently harmless virus called GB virus type C (GBV-C) for at least five years were three times less likely to die than HIV-positive men who did not have GBV-C. NIAID-supported scientists investigated the cellular mechanisms for the protective effect of GBV-C on HIV positive individuals. They found that when a type of immune cells called peripheral blood mononuclear cells (PBMCs) were infected with both HIV and GBV-C, the levels of cellular chemical messengers called chemokines were increased. In addition, they found an inverse correlation between chemokine levels and HIV replication. Moreover, they observed that the chemokine receptor CCR5, which is a co-receptor of some strains of HIV, was decreased on the surface of the HIV/GBV-C coinfected PBMCs. The elucidation of the mechanism through which GBV-C prolongs the survival of individuals infected with HIV may lead to the identification of targets for the development of novel therapeutics and vaccines to combat HIV/AIDS. In addition, this research provides clues as to why the course of HIV infection is so variable among individuals.

#### Future Directions in HIV/AIDS Research

NIAID will continue to support a broad array of domestic and international HIV/AIDS research programs that seek to increase basic knowledge of the pathogenesis, natural history, and transmission of HIV disease, and to support research that promotes progress in its detection treatment, and prevention. In addition, NIAID will continue to support basic research that seeks to increase understanding of the biology of HIV and how the immune system responds to it.

In FY 2006, NIAID will restructure all of its HIV/AIDS clinical trials networks. This reorganization, designed in response to both the changing face of the AIDS epidemic and evolving scientific challenges, will enable NIAID and its many collaborators to effectively pursue research for safe, effective, and affordable drugs and other therapeutic strategies, preventive strategies, and HIV vaccines. In addition, it will enable NIAID to more effectively respond to global research needs, particularly for people living with and most at risk for HIV/AIDS. The major scientific priorities that will be addressed with this new clinical trials matrix will be: 1) developing HIV vaccines; 2) translating research insights into therapeutic products to treat HIV disease; 3) optimizing clinical management of HIV/AIDS, including coinfections and other HIV-related conditions; 4) developing microbicides to prevent HIV acquisition and transmission; 5) preventing mother-to-child transmission of HIV; and 6) developing other methods of HIV prevention.

NIAID supports all phases of clinical trials to determine the safety, immunogenicity, and efficacy of candidate HIV vaccines. As of September 2004, NIAID had conducted or initiated, in collaboration with academic researchers and with industry co-sponsorship, over 70 vaccine trials, including 66 Phase I, 4 Phase II, and 1 Phase III trials. These studies involved over 11,000 volunteers, 51 vaccines, and 14 adjuvants. NIAID has a number of new vaccine candidates in the

preclinical pipeline, and four to eight are expected to enter Phase I studies in the next two years. These candidate vaccines will be evaluated in animals and then in early safety studies in humans.

In FY 2006, NIAID will continue to support HIV vaccine clinical trials, including a phase III clinical efficacy trial to evaluate a novel HIV vaccine strategy commonly referred to as "prime-boost." The trial, which began enrolling volunteers in 2004, is being conducted in Thailand in conjunction with the U.S. Army Medical Research and Materiel Command of the DoD. NIAID also will support several ongoing initiatives that promote the development of HIV vaccines, including the *Innovation Grants for Approaches in HIV Vaccine Research Program, HIV Vaccine Research and Design Program,* the *Integrated Preclinical/Clinical AIDS Vaccine Development Program (IPCA VD)*, and the *HIV Vaccine Design and Development Teams* (HVDDT). In FY 2006, both the IPCAVD and HVDDT programs are slated to be recompeted.

NT A ID supports the discovery and development of effective therapies for HIV/AIDS and associated complications and co-infections by facilitating and expediting research on highly promising candidate agents and novel therapeutic concepts. In FY 2006, NIAID will recompete the initiative *Novel HIV Therapies: Integrated Preclinical/Clinical Program*, which supports the discovery, development, and evaluation of innovative concepts for the treatment of HIV infection.

#### **Tuberculosis**

The bacterium that causes tuberculosis (TB), *Mycobacterium tuberculosis* (Mtb), is estimated to infect two billion people, or about one-third of the world's population. Five to ten percent of infected people will develop active TB disease sometime in their lifetimes. Each year, approximately eight million new cases of active TB occur, and approximately two million people die of the disease<sup>10</sup>.

#### Science Advances in Tuberculosis Research

A Tuberculosis Vaccine Ready for Testing in Humans has Shown Promising Results in Two Different Animal Models. While a vaccine for tuberculosis has been in use for more than 60 years, its clinical utility is restricted to preventing pediatric complications of tuberculosis, with limited and variable impact on adolescent or adult pulmonary disease, which are the dominant modes of disease transmission. NIAID-funded scientists have reported encouraging results in animals with a new candidate TB subunit vaccine. The vaccine displayed potent immune responses in mice and guinea pigs and protected against challenge with a virulent strain of TB. Human clinical trials of this candidate vaccine began early in 2004.

#### Future Directions in Tuberculosis Research

In FY 2006, NIAID will continue to promote and support a broad range of research on TB through its intramural research program and research initiated by individual investigators, as well as through NIAID-supported research programs, such as the *Tuberculosis Research Unit*, *National Cooperative Drug Discovery Groups for Tuberculosis*, and *TB Vaccine Testing and Research Materials*. NIAID will also continue to contribute to the Global Fund to Fight AIDS, TB, and Malaria.

#### Malaria

Malaria is one of the major killers of humans worldwide, threatening the lives of more than one-third of the world's population. Caused by a single-celled parasite and transmitted by mosquitoes, malaria causes an estimated 300 million acute illnesses each year and more than one million

<sup>&</sup>lt;sup>10</sup> World Health Organization, Tuberculosis, Fact Sheet No. 104, Switzerland, 2004.

deaths". The threat posed by malaria is growing, primarily because of the spread of drugresistant strains and insecticide-resistant mosquitoes, changing weather patterns, and limitations of the medical and public health infrastructure in many endemic areas.

#### Science Advances in Malaria Research

Combining the Antimalarial Drug Sulfadoxine-Pyrimethamine with other Antimalarial Drugs Reduces Treatment Failure Rates. Resistance to antimalarial drugs, especially chloroquine, the frontline treatment for more than 50 years, has become a major problem in Africa. Affordable and effective treatment options are limited, and signs of resistance are beginning to appear with sulfadoxine-pyrimethamine (SP), a widely used second-line antimalarial agent. Combining antimalarial drugs with different mechanisms of action can improve treatment efficacy and may delay the spread of drug resistance. NIAID-supported investigators compared the effectiveness of treating patients with malaria with the antimalarial drug SP alone or combined with either artesunate (AS) or amodiaquine (AQ). The rate of relapse was higher among patients treated with SP alone than among those treated with the SP plus AQ combination. These results indicate that the SP plus AQ combination is a more effective treatment than SP alone and may both impede the spread of drug-resistant parasites, as well as prolong the therapeutic lifespan of current antimalarial drugs.

#### Future Directions in Malaria Research

In FY 2006, NIAID will continue to promote and support malaria research through research initiated by individual investigators, as well as through targeted initiatives that aim to develop new therapeutics, new and improved diagnostics, and a vaccine to prevent infection. NIAID will continue to pursue the systematic implementation of its *Malaria Vaccine Plan*<sup>22</sup>, which was designed to accelerate research leading to the development of malaria vaccines.

#### EMERGING AND RE-EMERGING INFECTIOUS DISEASES

#### Severe Acute Respiratory Syndrome

Severe Acute Respiratory Syndrome (SARS) is the first severe, newly emergent infectious disease of the 21<sup>st</sup> century. SARS is a respiratory illness caused by a newly identified virus named SARS coronavirus (SARS-CoV). The disease emerged in late 2002 and spread to several countries in early 2003. The World Health Organization reported 8,096 cases, including 774 deaths worldwide<sup>13</sup>.

#### Science Advances in SARS research

<u>SARS Animal Models Developed.</u> NIAID scientists and their collaborators developed several animal models for SARS, including mouse, hamster, and non-human primate models. They also determined that the transfer of immune sera from infected mice and hamsters, which contained antibodies against SARS-CoV, protected uninfected animals from SARS infection. These observations suggest that vaccines that induce antibodies which neutralize the SARS-CoV and immune-based therapeutic and preventive measures with anti-SARS antibodies are likely to be effective against the SARS-CoV. In other efforts to evaluate the potential of immunotherapy,

<sup>&</sup>quot; World Health Organization, *Malaria, Fact Sheet No. 94*, <a href="http://www.who.int/mediacentre/factsheets/fs094/en/">http://www.who.int/mediacentre/factsheets/fs094/en/</a> (accessed on December 4, 2004).

<sup>&</sup>lt;sup>12</sup> NIAID Research for Malaria Vaccine Development, <a href="http://www.niaid.nih.gov/dmid/malaria/malvacdv/toc.htm">http://www.niaid.nih.gov/dmid/malaria/malvacdv/toc.htm</a>, (accessed December 30, 2004).

World Health Organization, Summary of probable SARS cases with onset of illness from 1 November 2002 to 31 July 2003, <a href="http://www.who.int/csr/sars/country/table2004\_04\_21/en/">http://www.who.int/csr/sars/country/table2004\_04\_21/en/</a> (accessed December 3, 2004).

NIAID-funded researchers have developed a human monoclonal antibody that reduces viral replication in mice and protects them against SARS-CoV challenge.

Three Candidate SARS Vaccines Developed. NIAID researchers have developed three candidate SARS vaccines, a DNA vaccine and two live attenuated virus vaccines. When mice vaccinated with the DNA vaccine were exposed to SARS-CoV, viral replication was reduced one-million fold. NIAID scientists at the VRC have begun a Phase I clinical trial to study early stage safety and immune response of this DNA vaccine candidate in humans. The attenuated live virus vaccines were made by inserting the gene encoding the SARS-CoV Spike (S) protein into two existing vaccines, modified vaccinia Ankara (MVA), which was initially developed as a vaccine against smallpox and has an excellent safety record in humans, and a recombinant attenuated human parainfluenza virus 3 (BHPIV3), which is an experimental vaccine against HPIV3, a virus that can cause respiratory illness in children. The MVA and BHPIV3 vaccines act to transport the SARS-CoV S gene into the body. The MVA/S vaccine was tested in mice and the BHPIV3/S vaccine in African green monkeys. Both experimental vaccines protected the immunized animals from infection with SARS-CoV.

#### Future Directions in SARS Research

In FY 2006, NIAID will continue to support basic and clinical research on SARS through its intramural and extramural programs. Basic research is supported through *Biodefense and Emerging Infectious Diseases Research Opportunities* and *NIH Investigator-Initiated Small Research Grants*. In FY 2006, product development, including vaccine, drug and immunotherapeutic development, will be supported through *Cooperative Research Partnerships for Biodefense, the Small Business Biodefense Program* and the *NIAID Advanced Technology Program*.

#### West Nile Virus

West Nile virus (WNV) first emerged in the Western Hemisphere in 1999, in New York City. The virus, transmitted by mosquitoes, has spread rapidly throughout the Americas. By 2004, the virus had been found in birds and mosquitoes in every state except Alaska and Hawaii. People who contract WNV usually experience no symptoms or only mild symptoms—fever, headache, body aches, skin rash, and swollen lymph glands. However, if WNV enters the brain, it can cause life-threatening encephalitis or meningitis. Currently, there are no drugs to treat the virus and no vaccines available to prevent infection.

#### Science Advances in West Nile Virus research

Clinical Trial of Experimental West Nile Virus Treatment. In FY 2004, NIAID expanded a Phase I/II clinical trial of an experimental WNV treatment to about 60 sites throughout the United States and Canada. The purpose of the study is to determine whether WNV-infected individuals given antibodies to the virus are better able to fend off the severe symptoms of WNV, such as encephalitis, that contribute to the deaths of some of the individuals who become infected. The antibody preparation, Omr-IgG-am<sup>TM</sup>, contains high levels of anti-WNV antibodies that were isolated from patients who recovered from WNV disease.

#### Future Directions in West Nile Virus Research

NIAID supports basic and clinical research on WNV, including research to develop and test vaccines, drugs and other therapeutic treatments, and improved diagnostics. For example, in FY 2006, NIAID will continue to support the development and testing of experimental WNV vaccines. NIAID also will continue to support the Phase I/II clinical trial of Omr-IgG-am<sup>TM</sup>.

#### Influenza and Other Respiratory Diseases

In the United States, influenza and pneumonia are the leading infectious causes of mortality and are ranked seventh among all causes of death. Annual influenza outbreaks (epidemics) in the United States typically occur between December and March and cause approximately 36,000 deaths each year<sup>14</sup>. Influenza can also cause global outbreaks of disease in which worldwide morbidity and mortality rates significantly increase (pandemics). The potential for a pandemic exists when new strains of influenza have emerged to which the human population has little or no prior immunity, and the virus acquires the ability to quickly spread from person to person and cause severe illness and death. Information about the influenza virus and NIAID-supported influenza research is detailed on the NIAID website *Focus on the Flus*.

Science Advances in Influenza and Other Respiratory Diseases Research

[See Story of Discovery]

Production and Clinical Testing of Investigational Avian Influenza Vaccines. In early 2004, a highly virulent strain of avian influenza, H5N1, re-emerged in southeast Asia, killing 32 of 44 people infected. Although the people in this outbreak were primarily infected through direct contact with birds, experts are highly concerned that the H5N1 may mutate to become highly infectious to humans and cause pandemic influenza. In FY 2004, NIAID awarded two contracts to support the production and clinical testing of an investigational vaccine based on the H5N1 strain. If a pandemic of H5N1 avian influenza were to occur in humans, the availability and production of such a vaccine on a commercial scale could be used to protect laboratory workers, public health personnel at risk and, if needed, the general public. In addition, NIAID supported the production of an investigational vaccine based on an H9N2 strain of avian influenza virus that has infected humans and has the potential to trigger a modern-day pandemic.

Future Directions in Influenza and Other Respiratory Diseases Research
NIAID supports the majority of Federally funded influenza research, including pandemic
preparedness research outlined in the DHHS Pandemic Influenza Response and Preparedness
Plan". In FY 2006, NIAID will continue to support through its intramural and extramural
programs both basic and applied research on influenza virus biology, epidemiology,
pathogenesis, immunology, and the development of new and improved influenza diagnostics,
antiviral drugs, and vaccines. In FY 2006, NIAID plans to expand the Pandemic Preparedness in
Asia contract, which supports surveillance and characterization of avian influenza viruses with
pandemic potential in the live bird markets in Hong Kong. NIAID will also launch in FY 2006
the initiative Pandemic Preparedness: Production of H7 Inactivated Influenza Vaccines for
production of vaccine that will be used for clinical trials to evaluate its safety and
immunogenicity. In addition, NIAID, in collaboration with multiple partners, will continue to
support the Influenza Genome Sequencing Project, which conducts influenza genomic
sequencing and rapidly provides the sequence data to the public, enabling scientists to further
study how influenza flu viruses evolve, spread, and cause disease.

<sup>&</sup>lt;sup>14</sup> CDC, *Influenza: Questions and Answers: the Disease*, <a href="http://www.cdc.gov/flu/about/qa/disease.htm">http://www.cdc.gov/flu/about/qa/disease.htm</a> (accessed December 3, 2004).

<sup>&</sup>lt;sup>15</sup> NIAID, Focus on the Flu, http://www2.niaid.nih.gov/newsroom/focuson/flu04/, (accessed December 3, 2004).

World Health Organization, Situation updates - Avian influenza (accessed December 4, 2004).

<sup>&</sup>lt;sup>17</sup> DHHS, *Pandemic Influenza Response and Preparedness Plan*, <a href="http://www.hhs.gov/nvpo/pandemicplan/">http://www.hhs.gov/nvpo/pandemicplan/</a>(accessed December 7, 2004).

#### Story of Discovery — Confronting Influenza: Responding through Research

Every year in the United States, more than 200,000 people are hospitalized and about 36,000 people die from influenza, commonly referred to as the flu, and its complications. In addition to the human cost, influenza exacts a tremendous toll economically. Recent estimates put the cost of influenza epidemics to the U.S. economy at between \$71 and \$167 billion per year.

Influenza, which is caused by a virus, is a classic example of a re-emerging disease; it is not a new disease, but it continually changes. The virus also can change suddenly through the recombination of two or more influenza viruses. When this happens, it usually is a "dead end" infection that cannot readily transmit between humans. However, if a virus acquires the capability to spread efficiently from person to person, the result can be a fast-moving and deadly pandemic, such as the influenza pandemic that occurred in 1918-1919, which killed 20-50 million people worldwide<sup>2</sup>. That virus is thought to have arisen through the recombination of an avian influenza strain with a human strain.

Until recently, flu researchers believed that avian influenza A viruses could not infect humans directly. Scientists thought that an avian flu virus would first need to infect another animal, such as a pig, before being transmitted to humans. In 1997, for the first time, scientists learned that avian influenza viruses could infect humans directly when the avian influenza virus strain called H5N1 infected 18 people, six of whom died. In 2004, the H5N1 virus circulated in East Asia; as of October 25, 2004, there had been 44 reported cases of people infected with the virus and 32 people had died. Fortunately, it is not easily transmitted between people.

#### Fighting Back

NIAID has responded to the public health threat posed by influenza virus by supporting research to monitor circulating and emerging flu strains, understand how new strains evolve, and develop diagnostics, treatments, and preventive strategies.

Preparedness planning involves surveillance of influenza virus activity. NIAID-supported scientists conducting surveillance of influenza strains in Asia investigated the genetic origin and spread of a highly pathongenic avian influenza virus strain, H5N1, which caused outbreaks in poultry and humans in 2003 and 2004. Their findings indicate that a dominant form of the virus emerged and was sustained in domestic ducks in southern China and may have then been spread by wild birds to other parts of Asia. The fact that H5N1 may now be endemic in poultry in Asia indicates that it has developed an entrenched ecological niche and may be difficult to eradicate.

If this avian influenza strain or another highly pathogenic avian influenza strain were to mutate and become transmissible between humans, it would not be recognized by the human immune system and could lead to widespread infection, illness, and death. Therefore, an arsenal of treatments and preventive strategies is needed to prevent a worst case scenario from happening.

An innovative strategy developed by a team of scientists supported by NIAID may lead to the development of novel ways to treat, and perhaps prevent, influenza infection. The scientists used a technique known as RNA interference to shut down the expression of targeted influenza virus genes. The scientists designed specific short interfering RNA (siRNA) molecules to target two influenza proteins that are essential for infection. These siRNAs, when given to mice by injection or aerosol before or after viral infection, blocked virus production in the lungs and effectively prevented and treated influenza infection.

The most effective way to prevent the flu is through vaccination; however, it can take four to six months to produce vaccine in large amounts. If a flu strain with pandemic potential does evolve, scientists will need to act quickly to generate a vaccine against the new virus. Recently, NIAID-supported investigators developed a technology, plasmid-based reverse genetics, which has the potential to dramatically decrease the time it takes to produce a flu vaccine. Using this technique, the researchers were able to custom make, in only four weeks, a vaccine reference strain of the H5N1 avian virus. Previous attempts to produce a vaccine against this highly lethal virus had failed because the virus could not be grown in chicken eggs, a necessary step in the current production of flu vaccine. NIAID-supported investigators have begun clinical trials of the H5N1 vaccine strain made using reverse genetics to evaluate its safety and efficacy in humans.

- U.S. CDC, Influenza Questions and Answers: the Disease, http://www.cdc.gov/flu/about/qa/disease.htm (accessed December 4, 2004)
- WHO, Influenza, Fact Sheet No. 211, http://www.who.int/mediacentre/factsheets/fs211/en/ (accessed December 4, 2004).
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- Qing Ge et al., Inhibition of Influenza Virus Production in Virus-Infected Mice by RNA Interference. <u>Proc. Natl. Acad. Sci. U.S.A</u>. 101: 8676-8681, 2004.
- Webby et al., Responsiveness to a Pandemic Alert: Use of Reverse Genetics for Rapid Development of Influenza Vaccines. <u>Lancet</u> 363: 1099-1103, 2004

#### Transmissible Spongiform Encephalopathies

Transmissible spongiform encephalopathies (TSEs) are a family of neurodegenerative disorders that affect many mammals, including humans. These diseases include scrapie, which primarily affects sheep; bovine spongiform encephalopathy or "mad cow disease," which affects cattle; chronic wasting disease (CWD), which affects deer and elk; and Creutzfeldt-Jakob disease, which affects humans. TSEs are untreatable and invariably fatal diseases of the brain. TSEs are characterized by the presence within the brain of abnormally shaped versions of protein molecules called prion proteins, which are associated with the tissue damage observed in these diseases. Although the precise nature of TSE infectious agents remains unclear, it is known that the abnormal form of the prion protein can convert normal prion protein molecules to the abnormal form, and thus has the potential to be the infectious agent.

Science Advances in Transmissible Spongiform Encephalopathies Research

Persistent Scrapie Infection is Likely Due to Cell-Specific Factors that May Serve as Therapeutic

Targets. Normal prion protein is expressed in a wide variety of tissues, yet conversion of normal prion protein to the TSE form appears to be restricted primarily to cells of the nervous and lymphoid systems. In order to determine why some cell types are more resistant to TSE infection than others, NIAID scientists developed a tissue culture system that allows them to monitor both acute and persistent abnormal prion protein (or PrP-res) formation. They demonstrated that, while any cell type can make new PrP-res following exposure to TSE infectivity, only some cell types go on to become chronically infected and make PrP-res persistently. This suggests that there are cell-specific factors that determine the susceptibility of a cell to chronic TSE infection. These factors, once identified, could be useful in designing effective anti-TSE therapeutics.

Future Directions in Transmissible Spongiform Encephalopathies Research In FY 2006, NIAID will continue to support TSE research through its intramural and extramural research programs. The NIAID TSE research agenda is focused on four areas: understanding the infectious agents of TSEs; defining how TSEs are transmitted among animal species and across species barriers; developing diagnostic tests; and developing therapies.

#### Hepatitis

Hepatitis refers to a group of liver diseases caused by a diverse set of viruses: hepatitis A, B, C, D, and E viruses. Infection with hepatitis viruses causes liver inflammation, tissue damage and dysfunction. Both hepatitis B virus (HBV) and hepatitis C virus (HCV) cause chronic infection.

In the United States There are approximately 1.2 million HBV carriers and approximately 2.7 million HCV carriers<sup>19</sup>.

Science Advances in Hepatitis Research

Researchers Identify Better Hepatitis C Treatment for People with HIV. NIAID-supported scientists determined that that the preferred treatment for HCV, peginterferon and ribavirin, is safe for people who are also infected with HIV. Moreover, this treatment proved superior for the treatment of HCV in HIV-coinfected persons when compared with the previously accepted treatment, standard interferon and ribavirin.

<sup>19</sup> U.S. Centers for Disease Control and Prevention, *Hepatitis C Fact Sheet*, http://www.cdc.gOv/ncidod/diseases/hepatitis/c/cfact.pdf (accessed December 3, 2004).

<sup>&</sup>lt;sup>18</sup> U.S. Centers for Disease Control and Prevention, *Frequently Asked Questions About Hepatitis B*, <a href="http://www.cdc.gOv/ncidod/diseases/hepatitis/b/faqb.htm">http://www.cdc.gOv/ncidod/diseases/hepatitis/b/faqb.htm</a> (accessed December 3, 2004).

#### Future Directions in Hepatitis Research

In FY 2006, NIAID will continue to conduct and support basic and clinical research on the hepatitis viruses. The new initiative *Non-Biodefense Partnerships: Vaccines for Hepatitis C* will be launched to stimulate industry participation in the development of vaccines for hepatitis C. In addition, NIAID will continue to support several ongoing projects, including the *Hepatitis C Cooperative Research Centers*, which conduct multidisciplinary H C V research, and the *Animal Models for the Prevention and Treatment of Hepatitis B and C* initiative.

#### Antimicrobial Resistant Microbes

Antimicrobials have transformed the treatment of many infectious diseases that were killers only a few decades ago. Over time, however, many pathogens have developed resistance to these powerful drugs. All major groups of pathogens—viruses, fungi, parasites, and bacteria—can become resistant to antimicrobials. Due to the emergence and spread of antimicrobial resistance, several bacterial infections such as methicillin-resistant *Staphylococcus aureus* (MRSA), vancomycin-resistant *S. aureus*, vancomycin-resistant *Enterococcus*, multi-drug-resistant *M. tuberculosis*, and penicillin-resistant *Streptococcus pneumoniae* are difficult to treat and have negative clinical outcomes and increased treatment costs.

#### Science Advances in Antimicrobial Resistance Research

<u>Understanding How Resistance Develops.</u> Originally appearing only in hospital settings, methicillin-resistant *S. aureus* (MRSA) has now emerged as a community-acquired infection. To determine the origins of MRSA strains circulating within the community, NIAID-supported investigators isolated bacteria from a high-risk population and conducted genetic analysis. The researchers found 58 percent of MRSA infections were caused by strains traceable to hospitals or long-term care facilities. Infections linked to injection drug use, however, were not linked to health care facilities. Therefore, while hospitals appear to remain the main source of MRSA in the community, the presence of other genetically diverse strains indicates that some MRSA strains now originate from the community.

#### Future Directions in Antimicrobial Resistance Research

In FY 2006, NIAID will support a new initiative called *Drug Development Resources for Antiinfectives*. NIAID also plans to continue research through an initiative launched in FY 2005, *Sepsis and Community Acquired Pneumonia: Partnerships for Diagnostics Development* to support development of diagnostic technologies for early detection for pathogens that cause hospital-acquired diseases, including drug-resistant respiratory and systemic infections. NIAID will also continue support for several antimicrobial resistance-related research networks, including the *Network on Antimicrobial Resistance in S. aureus*. Furthermore, NIAID will continue collaborations with industry for the development of novel products to address resistant bacterial infections in healthcare settings.

#### OTHER INFECTIOUS DISEASES

#### Sexually Transmitted Infections

Sexually transmitted infections (STIs) can be caused by a variety of viruses, bacteria, and parasites. More than 25 STIs have been identified, and each year they affect more than 15 million people in the United States<sup>20</sup>. STIs can lead to infertility, complications in pregnancy, cervical cancer, low birth weight, congenital/perinatal infections, other chronic conditions such as neurosyphilis, and increased risk of HIV infection. Treatment and prevention of STIs have

<sup>20</sup>U.S. Centers for Disease Control and Prevention, *Tracing the Hidden Epidemics: Trends in STDs in the United States 2000*, <a href="http://www.cdc.gov/nchstp/dstd/Stats">http://www.cdc.gov/nchstp/dstd/Stats</a> Trends/Trends2000.pdf(accessed December 3, 2004).

become critical global and national health priorities because of their devastating impact on women and infants and their inter-relationship with HrV/AIDS.

Science Advances in Sexually Transmitted Infections Research

Once-Daily Antiviral Treatment with Valacyclovir Reduces the Risk of Transmission of Genital Herpes to Uninfected Partner. A randomized controlled clinical trial conducted by NIAID-supported scientists demonstrated that once-daily antiviral suppressive therapy with valacyclovir significantly reduced the risk of transmission of genital herpes from an infected partner to an uninfected heterosexual partner. This was the first time an antiviral had been shown to reduce the risk of transmission of a sexually transmitted infection.

<u>Universal Chlamydia Screening of Adolescent Males on Entry to a Detention System Is a Cost-Effective Prevention Strategy.</u> NIAID intramural researchers and their collaborators determined that universal chlamydia screening of young males on entry to a detention system was the most cost-effective strategy to prevent pelvic inflammatory disease in their recent and future partners. Screening detained male youth using a urine-based test provides a public health opportunity to significantly reduce chlamydia infections in youth at risk for STIs.

Future Research Directions in Sexually Transmitted Infections Research
In FY 2006, NIAID will continue to support basic and clinical research studies on mechanisms of pathogenesis of STIs and prevention strategies for the control of these infections through the NIAID-funded Sexually Transmitted Diseases Cooperative Research Centers and the Sexually Transmitted Infections Clinical Trials Group, as well as several other related initiatives.

#### Enteric Diseases

Bacterial and viral infections of the gastrointestinal tract can lead to diarrheal disease as well as chronic conditions such as ulcers and stomach cancer. In the United States, diarrhea is the second most common infectious illness, accounting for one of every six (16 percent) of all infectious diseases<sup>21</sup>. Data compiled by the World Health Organization indicate that diarrheal diseases account for 15 to 34 percent of all deaths in some countries and worldwide cause more than two million deaths per year<sup>22</sup>.

Science Advances in Enteric Diseases Research

Mechanisms by which *Helicobacter pylori* Evades the Host's Immune Response. Infection with *H. pylori* is a major risk factor for developing peptic ulcer disease, stomach cancer, and primary gastric B cell lymphoma. NIAID researchers have identified mechanisms that may aid in the ability of the bacteria to colonize the stomach and persist for decades. Most *H. pylori* strains secrete a vacuolating toxin (VacA), which has been implicated as an important virulence factor in the pathogenesis of peptic ulceration and gastric cancer. New evidence demonstrates that VacA suppresses the increase in numbers of *H. pylori-specific* immune cells. These effects may contribute to the capacity of *H. pylori* to evade the adaptive immune response and establish persistent infection.

Future Directions in Enteric Diseases Research

In FY 2006, NIAID will continue to support research aimed at understanding, preventing, and treating enteric diseases through a wide array of initiatives, including investigator-initiated

<sup>&</sup>lt;sup>a</sup> Mead PS et al., Food-Related Illness and Death in the United States. Emerg Infect Pis 5: 607-625, 1999.

<sup>&</sup>lt;sup>22</sup> World Health Organization. *Communicable Diseases 2002: Global Defence Against the Infectious Disease Threat, Geneva,* Switzerland, 2003.

efforts and targeted programs, such as the *Food and Waterborne Diseases Integrated Research Network* and the *Cooperative Research Partnerships for Biodefense*.

#### Lyme and Other Insect-Borne Disease

Ticks, mosquitoes, lice, and fleas often spread viral, bacterial, and parasitic diseases. Lyme disease, an infection caused by the bacterium *Borrelia burgdorferi*, remains the most prevalent tick-borne infectious disease in the United States. The U.S. incidence of reported cases of Lyme disease declined 10 percent from 2002 to 2003, from 23,763 cases in 2002 to 21,273 in 2003<sup>23</sup>.

Science Advances in Lyme and Other Insect-Borne Diseases Research

Key to Survival of the Bacterium that Causes Lyme Disease. NIAID-supported investigators discovered that two bacterial proteins, outer surface protein A (OspA) and B (OspB), are essential for the colonization and survival of *B. burgdorferi* in ticks, thereby playing a crucial role in the transmission of Lyme disease to humans. Other studies have shown that if ticks are fed on mice immunized against OspA or OspB, or if ticks are permitted to feed on mice that have been treated with antibodies specific for OspB, the colonization of *B. burgdorferi* in the ticks is significantly impaired. This suggests that the development of vaccines that stimulate production of antibodies specific for OspA or OspB in wildlife populations may be an effective strategy for preventing the spread of Lyme disease.

Future Directions in Lyme Disease Research and Other Insect-Borne Diseases Research In FY 2006, NIAID will continue its long-standing commitment to research on Lyme disease, ehrlichiosis and Rocky Mountain spotted fever through both intramural and extramural research. Studies will include research on vectors and reservoirs, pathogenesis of disease; immune mechanisms in the host; animal models for basic and applied research; and improved diagnosis, treatment and prevention of disease.

#### Fungal Diseases

Severe, sometimes life-threatening, systemic infections caused by fungi have long been recognized in all age groups in all parts of the world. Treatment of fungal infections requires prolonged administration of relatively toxic drugs, which are sometimes ineffective, even in otherwise healthy patients. Fungal infections are recognized as a major cause of morbidity and mortality in patients with an impaired immune system. In addition, advances in modern medicine involving immunosuppression have contributed to the increase in fungal diseases in hospital settings.

#### Science Advances in Fungal Diseases Research

Monoclonal Antibodies Protect Against the Fungus *Histoplasma capsulatum*. The fungus *H. capsulatum* is a significant cause of potentially fatal infection in persons with weakened immune systems, such as individuals with HIV/AIDS. NIAID-funded investigators have discovered an antibody that protects mice from infection with *H. capsulatum*. This monoclonal antibody sticks to a protein on the fungal surface and helps the host's immune cells to destroy the fungus. This is the first demonstration of the use of an antibody for protection against infection with this fungus.

#### Future Directions in Fungal Diseases Research

In FY 2006, NIAID will support basic and applied mycology research. Research areas include: molecular biology, immunobiology, pathogenesis, therapy, and genomics and proteomics. The *Mycology Research Units (MRUs)* bring together teams of investigators to develop and improve <sup>23</sup> U.S. Centers for Disease Control and Prevention, Final 2003 Reports of Notifiable Diseases. MMWR 53: 688-696, 2004.

methods for the diagnosis, prevention, and treatment of fungal infections; three new MRUs were awarded in FY 2004. The *Bacteriology and Mycology Biostatistical and Operations Unit* and the *Bacteriology and Mycology Study Group* will continue to support clinical trials on fungal and resistant bacterial infections.

#### CONFRONTING IMMUNE-MEDIATED DISEASES

The immune system is a collection of cells and proteins that works to protect the body from potentially harmful, infectious microorganisms such as bacteria, viruses and fungi and plays an important role in the control of cancer and other diseases. It also is the culprit in allergies; hypersensitivity; immunologic diseases; and the rejection of transplanted organs, cells, tissues, and medical implants. The past two decades of intensive and highly productive research on the immune system have resulted in a wealth of new information and extraordinary growth in its conceptual understanding.

#### Immune Tolerance

The successful induction of immune tolerance is a major therapeutic goal for the treatment of many immune-mediated diseases, including asthma and allergic diseases; autoimmune disorders, such as rheumatoid arthritis, type 1 diabetes, and multiple sclerosis; and rejection of transplanted organs, tissues, and cells. Tolerance induction strategies aim to selectively block or prevent deleterious immune responses, while leaving protective immunity intact. Advances in tolerance induction will provide valuable new therapeutic strategies that do not require life-long, global immunosuppressive therapy with its associated deleterious side effects, and the ability to modulate tolerance will also be important for enhancing protective immunity in response to vaccines for tumors and infectious diseases.

#### Science Advances in Immune Tolerance

In Vitro Expanded Regulatory T Cells Suppress Autoimmune Diabetes. Research in animal models suggests that successful tolerance induction requires the elimination of tissue-damaging cells and activation of a type of immune cell called regulatory T (T<sub>reg</sub>) cells which modulate pathogenic immune responses. Studies further suggest that the T<sub>res</sub> cell population is diminished or functionally impaired in patients and animals with autoimmune disease. It has been difficult to assess the therapeutic potential of the T<sub>ses</sub> cell population because it is difficult to isolate a sufficient number for study. NIAID-funded investigators recently developed an in vitro method for robust expansion of the T<sub>reg</sub> cell population. They confirmed that these cells retain their immunomodulatory activity, as evidenced by their ability to reverse new onset and chronic diabetes in a mouse model of type 1 diabetes, and to prevent islet graft rejection in the diabetic mice. Most significantly, only very small numbers of T<sub>reg</sub> cells specifically reactive to a self protein in autoimmune diabetes are needed to accomplish this reversal of disease. Increased numbers of T<sub>1.12</sub> cells will allow further study of their characteristics, mechanisms of action, and application to other models of immune-mediated diseases, such as graft rejection and graftversus-host disease. Ultimately, in vitro T<sub>reg</sub> cell expansion and reinfusion into patients may be used to treat a variety of immune-mediated diseases.

#### Future Directions in Immune Tolerance Research

NIAID will continue to support research on the induction, maintenance, and loss of tolerance. These efforts will focus on: understanding the basic mechanisms of immune tolerance; manipulating the immune response for tolerance induction; and evaluating tolerance induction regimens in transplantation and autoimmune disease models. NIAID, in conjunction with the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) and the Juvenile

Diabetes Research Foundation International (JDRF), supports the *Immune Tolerance Network* (*ITN*), an international consortium of over 80 investigators dedicated to the clinical evaluation of novel, tolerance-inducing therapies; the ITN will be recompeted in FY 2006. NIAID also plans to recompete the *Innovative Grants on Immune Tolerance* program in FY 2006.

#### Autoimmune Diseases

Autoimmune diseases result from a dysfunction of the immune system in which the body attacks its own organs, tissues, and cells. The body has safeguards to prevent the immune system from attacking its own tissues, but when these safeguards are breached, an autoimmune disease can result. Medical science has identified more than 80 clinically distinct autoimmune diseases, including systemic lupus erythematosus, type 1 diabetes, severe lupus nephritis, Sjögren's syndrome, Crohn's disease, and multiple sclerosis. Collectively, autoimmune diseases afflict an estimated five to eight percent of the U.S. population and disproportionately affect women<sup>24</sup>.

#### Scientific Advances in Autoimmune Diseases

Early Autoantibodies Serve as Warning for Lupus. Systemic lupus erythematosus (SLE) is a potentially life-threatening autoimmune disease that can harm the kidneys, lungs, central nervous system, and heart. Because autoantibodies are central to this damage, scientists believe that their development coincides with or precedes clinical disease. NIAID-supported researchers recently discovered that clinical manifestations of SLE are preceded by autoimmune changes that are underway, and continue to progress, for many years before diagnosis. This study indicates that the presence of autoantibodies could be used to diagnose lupus before symptoms appear, thereby enabling clinicians to offer treatment earlier in the disease course and improve treatment outcome.

#### Future Directions in Autoimmune Diseases Research

NIAID supports a broad range of basic and clinical research programs in autoimmunity, including several multicenter research programs. For example, NIAID has established nine *Autoimmunity Centers of Excellence* (ACEs) that conduct collaborative basic and clinical research on autoimmune diseases. The ACEs are co-sponsored by the NIDDK and the NIH. Office of Research on Women's Health (ORWH). In FY 2006, NIAID will recompete the *Autoimmune Diseases Prevention Centers*, which are cosponsored by NIDDK, the National Institute of Child Health and Human Development (NICHD), ORWH, and JDRF. The Prevention Centers will focus on prevention of autoimmune disease before clinical onset by mechanisms other than global immunosuppression. In addition, in FY 2006, NIAID plans to recompete the *Multiple Autoimmune Diseases Genetics Consortium* and the *Stem Cell Transplantation for the Treatment of Autoimmune Diseases* program.

#### Asthma and Allergic Diseases

Allergic diseases and asthma are major causes of illness and disability in the United States; more than 50 million Americans suffer from allergies, asthma, or both, and the cost to the health care system is more than \$18 billion annually<sup>25</sup>. An allergy is a specific reaction of the body's immune system to a normally harmless substance, one that does not bother most people. Allergic antibodies initiate allergic inflammation, rhinitis, and asthma.

<sup>25</sup> NIAID, *Allergy Statistics*, <a href="http://www.niaid.nih.gov/factsheets/allergystat.htm">http://www.niaid.nih.gov/factsheets/allergystat.htm</a>, (accessed December 4, 2004).

<sup>&</sup>lt;sup>24</sup> U.S. Department of Health and Human Services, National Institutes of Health, *Autoimmune Diseases Coordinating Committee, Autoimmune Disease Research Plan. Bethesda*, MD, 2002.

Scientific Advances in Asthma and Allergic Diseases Research

Chronic Sinusitis Sufferers Have Enhanced Immune Responses to Fungi. Nearly 30 million people were diagnosed with sinusitis in 2002, and direct costs of the illness exceed \$5.6 billion per year. Scientists supported by NIAID discovered that people with chronic sinus inflammation, many of whom also suffer from allergies and asthma, have an exaggerated immune response to common airborne fungi. By comparing blood samples and nasal secretions taken from people diagnosed with chronic sinusitis with samples from healthy volunteers, the investigators observed that the levels of fungal proteins in nasal secretions were similar in both groups, thus indicating that the mere presence of fungi in the airways is not enough to cause sinusitis. The investigators also looked for evidence that immune cells from people with sinusitis respond abnormally to harmless fungi. They found that the immune cells of chronic sinusitis sufferers released significantly greater amounts of three immune-modulating chemicals called cytokines than healthy volunteers. This study is the first to show a possible immunologic basis for chronic sinusitis.

<u>Customized Program Reduces Asthma Symptoms in Inner-City Children</u>. Results from the Inner-City Asthma Study, which was co-funded by NIAID and the National Institute of Environmental Health Sciences, demonstrated that an environmental intervention that targets allergens and tobacco smoke in the home resulted in fewer asthma symptoms in children participating in the program than in those who did not. The environmental intervention, which was specifically tailored to each child's needs, substantially lowered levels of cockroach and house dust mite allergens, and investigators observed a direct relationship between reduction in allergen levels and a decline in asthma symptoms.

Future Directions in Asthma and Allergy Research

NIAID has been at the forefront of many advances and discoveries leading to the characterization of asthma and allergic diseases as immunological disorders. In FY 2006, NIAID will continue to support several ongoing research programs. For example, the network of 13 *Asthma and Allergic Diseases Research Centers* (AADRCs), which will be recompeted in FY 2006, conducts basic and clinical research on the mechanisms, diagnosis, treatment, and prevention of asthma and allergic diseases. The *Inner-City Asthma Consortium: Immunologic Approaches to Reduce Asthma Severity,* which was initiated in FY 2002, is a network of basic scientists and clinical investigators who are evaluating the efficacy of immune-based therapies for reducing asthma severity and preventing onset in inner-city children. NIAID plans to establish the *Food Allergy Research Consortium* in FY 2005 to provide critical information on the pathophysiology and natural history of the disease and develop effective preventive interventions.

#### **Transplantation**

According to the Organ Procurement and Transplantation Network, more than 25,000 solid organ transplants were performed in the United States in 2003<sup>27</sup>. In addition, as of August 2004, more than 86,000 people had their names on waiting lists for organs such as livers, kidneys, hearts, lungs, and intestines <sup>28</sup>. Although organ replacement prolongs survival for people suffering from end-stage organ failure, it rarely restores normal life expectancy and can

28 Ibid.

<sup>&</sup>lt;sup>26</sup> U.S Centers for Disease Control and Prevention, *Chronic Sinusitis*, <a href="http://www.cdc.gov/nchs/fastats/sinuses.htm">http://www.cdc.gov/nchs/fastats/sinuses.htm</a> (accessed December 4, 2004)

The U.S. Organ Procurement and Transplantation Network and the Scientific Registry of Transplant Recipients, 2003 OPTN/SRTR Annual Report, <a href="http://www.optn.org/AR2003/default.htm">http://www.optn.org/AR2003/default.htm</a>, (accessed December 4, 2004).

sometimes lead to health problems associated with long-term use of immunosuppressive drugs, which reduce the risk of transplant rejection but also weaken the immune system against disease. Hematopoietic cell transplantation (HCT), the transplantation of stem cells found in bone marrow and cord blood, is used to treat many forms of hematologic malignancies, non-malignant diseases, and inborn errors of immunity and metabolism. Over 17,000 patients received HCT procedures in 2003 <sup>29</sup>. However, HCT carries the risk of potentially fatal complications, such as graft failure, opportunistic infections, and graft-versus-host disease.

#### Scientific Advances in Transplantation

A Protective Gene for Graft-Versus-Host Disease following Bone Marrow Transplantation. Hematopoietic cell transplantation is an effective therapy for a number of life-threatening diseases, including leukemias. The primary complication of HCT is graft-versus-host disease (GVHD), in which donor immune cells called T cells that accompany the transplanted blood cells produce an immune response against host organs and tissues, often with fatal consequences. In a study of nearly 1,000 recipients of HCT from unrelated donors, NIAID-funded researchers discovered that a genetic variant of the gene encoding the immune system molecule interleukin-10 decreases the risk of acute GVHD and death after HCT by close to 50 percent.

#### Future Directions in Transplantation Research

In FY 2006, NIAID will continue to support transplantation research aimed at: understanding the mechanisms whereby the immune system recognizes and either accepts or rejects transplanted organs, tissues, and cells; evaluating promising therapies to improve graft survival and function and to prevent and treat graft rejection; and exploring the challenges to xenotransplantation, the transplantation of an organ, tissue, or cells between two different species. The *Clinical Trials in Organ Transplantation* program, co-sponsored by NIAID, NIDDK, and NHLBI, supports a clinical consortium dedicated to improving the success of organ transplantation. In FY 2005, NIAID will launch the *HLA Region Genetics in Immune-Mediated Diseases* initiative to continue research on defining associations between genes or markers in the HLA region and immune-mediated diseases. The genes that encode the HLA help regulate several aspects of the immune response and code for "self markers on all body cells.

#### Primary Immunodeficiency Diseases

Primary immunodeficiency diseases are caused by inherited defects in the immune system that increase susceptibility to infections. Unlike secondary or acquired immune deficiency diseases, which are caused by infectious, chemical, or radiological agents, the estimated 80 primary immunodeficiency diseases are inherited conditions in which specific cells of the immune system do not function properly. Approximately 25,000 to 50,000 Americans are severely affected by primary immunodeficiency diseases, and there are believed to be another 500,000 persons who remain undiagnosed<sup>30</sup>.

#### Scientific Advances in Primary Immunodeficiency Diseases

<u>Thymus Tissue Transplantation Restores Immune System Function.</u> DiGeorge syndrome is a congenital primary immunodeficiency disorder in which the thymus gland, heart, and parathyroid glands fail to develop normally. The thymus gland is required for the normal development and maturation of T lymphocytes, a type of immune cell. NIAID-funded researchers previously demonstrated that thymus transplantation is an effective therapy for

<sup>29</sup> Ibid.

<sup>&</sup>lt;sup>30</sup>NIAID, *Primary Immune Deficiency*, <a href="http://www.niaid.nih.gOv/factsheets/pid.htm#more">http://www.niaid.nih.gOv/factsheets/pid.htm#more</a>, (accessed December 3, 2004).

patients with complete DiGeorge syndrome, a disorder characterized by the lack of a thymus. These researchers have extended their previous studies to include infants with partial DiGeorge syndrome, in which thymus development and function are impaired, but a small number of T cells are present. Because there was no tissue matching between the thymus donor and the infant and because the small number of T cells that remain in partial DiGeorge syndrome infants increase the risk of graft rejection, the infants were given T cell depleting agents prior to transplantation. One year post transplant, T cell function showed normal or greatly improved results as compared with the pre-transplant period, demonstrating that host T cells can proliferate and mature in a non-matched donor thymus, even after the use of T cell depleting agents. These results may also have important implications for patients requiring T cell depleting therapies for a variety of diseases.

Future Directions in Primary Immunodeficiency Diseases Research

The major goals of NIAID-supported research in primary immunodeficiency diseases are: to understand the causes and immune mechanisms leading to their development, including identifying gene mutations and other contributing factors; to expand the genetics knowledge base to improve diagnosis, facilitate genetic counseling and decision-making for affected individuals; and to provide protective and curative treatments, including gene therapy. In FY 2006, NIAID, in conjunction with NICHD, will continue to support the *Primary Immunodeficiency Research Consortium*, a coalition of the world's most prominent researchers in the field of primary immunodeficiency diseases who are working to prioritize and coordinate future research directions and develop new resources for the study of these comparatively rare disorders.

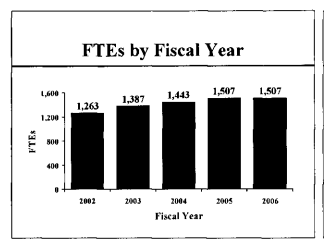
#### NIH ROADMAP: ACCELERATING MEDICAL RESEARCH PROGRESS

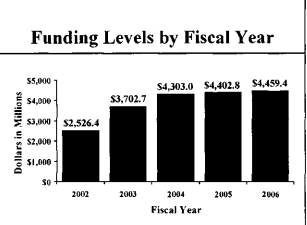
NIAID will explore novel approaches for diagnosis, treatment and prevention of disease using innovative research technologies such as nanomedicine, high throughput molecular screening, metabolomics, and bioinformatics through the NIH Roadmap. The Roadmap will enable NIAID to accelerate the translation of basic research discoveries into clinical practice by fostering interdisciplinary research and training and promoting public-private partnerships. The *Director's* Pioneer Award will support individual scientists with highly innovative ideas and approaches to contemporary challenges in biomedical research. Through this award, NIAID will support the study of novel approaches to HIV/AIDS vaccine development. Nanomedicine initiatives will support highly specific medical intervention at the molecular level. The Roadmap will support studies on nanomedicine immunomodulation, interdisciplinary research on antimicrobial resistance, vector-borne disease control in urban environments, and research centers - the Exploratory Centers for Interdisciplinary Research and an Exploratory Center for Vaccinology Research. The Berkeley Nanomedicine Center in Membrane Signaling may uncover mechanisms of immune regulation. High Throughput Molecular Screening Assay Development will encourage the development and automation of biological assays that enable high throughput screening (HTS) of thousands of distinct chemical entities to better understand biological processes and identify promising therapeutics. Examples of Roadmap funded HTS activities that support NIAID's mission include identification of host factors important for Listeria infection, cellular regulators of hepatitis C virus infection, and potential inhibitors for both smallpox and malaria.

#### **Budget Policy**

The Fiscal Year 2006 budget request for the NIAID is \$4,459,395,000, an increase of \$56,554,000 and 1.3 percent over the FY 2005 Appropriation. Also included in the FY 2006 request, is NIAID's support for the trans-NIH Roadmap initiatives, estimated at 0.89% of the FY 2006 budget request. This Roadmap funding is distributed through the mechanisms of support, consistent with the anticipated funding for the Roadmap initiatives. A full description of this trans-NIH program may be found in the NIH Overview.

A five year history of FTEs and Funding Levels for NIAID are shown in the graphs below.





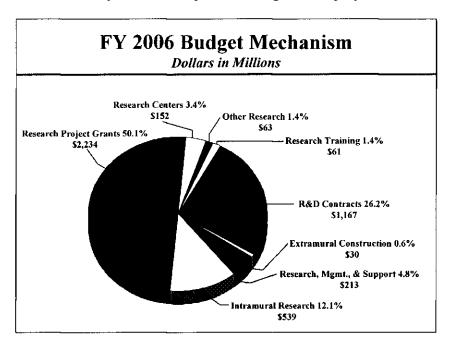
NIH's highest priority is the funding of medical research through research project grants (RPGs). Support for RPGs allows NIH to sustain the scientific momentum of investigator-initiated research while pursuing new research opportunities. The average cost of competing RPGs will be held at the FY2005 level. However, the NIAID average cost for competing RPGs is skewed by the large average cost of HIV/AIDS Clinical Trials Networks. Excluding these large grants, the average cost of comparable competing RPGs will be held at the FY 2005 level. There will be no inflationary increases for direct, recurring costs in Noncompeting continuation RPGs.

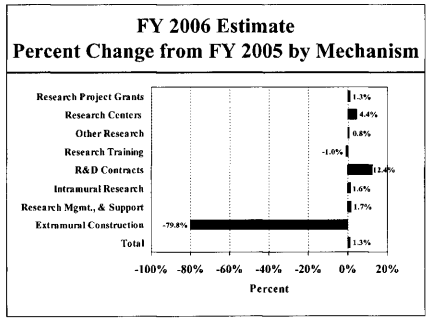
Advancement in medical research is dependent on attracting the best and the brightest to pursue careers in biomedical research. In the Fiscal Year 2006 request, stipend levels for post-doctoral recipients supported through the Ruth L. Kirschstein National Research Service Awards will increase by 4.0% for those with 1-2 years of experience, with all other stipends remaining at the FY2005 levels. NIH will also provide an increase of \$500 per post-doctoral fellow, for increased health insurance costs. This increase in stipends and health benefits is financed within the FY2006 request by reducing the number of Full-Time Training Positions by -52. NIAID will support 1,292 pre- and postdoctoral trainees in full-time training positions.

The Fiscal Year 2006 request includes funding for 44 research centers, 450 other research grants, including 362 research career awards, and 271 R & D contracts. Intramural Research and Research Management and Support receive increases of 1.6 percent and 1.7 percent, respectively.

Additionally, the FY 2006 budget request includes \$34.0 million to support the expansion of the extramural HIV Vaccine Research Center (VRC). At the G-8 Summit in June 2004, President Bush endorsed the establishment of a Global HIV Vaccine Enterprise, a virtual consortium to accelerate HIV vaccine development, and urged his G-8 counterparts to increase their commitment to the development of an HIV vaccine. Conceptually, the extramural VRC will function similarly to the NIAID Vaccine Research Center, but will be established within the extramural community and be virtual in nature (i.e., we will not build a discrete building to house the center as we did with the NIAID VRC).

The mechanism distribution by dollars and percent change are displayed below:





Budget	Mechanism	-	Total

<del></del>		Mechanism		г	
	FY 2004 FY 2005		FY 2006		
	Actual	Aŗ	Appropriation		Estimate
No.	Amount	No.	Amount	No.	Amount
				I	
2,956	\$1,423,484,000	3,086	\$1,611,918,000	2,954	\$1,471,494,000
(119)	62,821,000	(70)	17,250,000	(67)	16,499,000
1		1		1	
304	125,693,000	284	113,290,000	246	157,054,000
863	431,539,000	796	365,151,000	708	487,598,000
1	126,000	1	126,000	11	126,000
1,168	557,358,000	1,081	478,567,000	955	644,778,000
4,124	2,043,663,000	4,167	2,107,735,000	3,909	2,132,771,000
258	97,162,000	303	97,914,000	312	101,263,000
4,382	2,140,825,000	4,470	2,205,649,000	4,221	2,234,034,000
32	113,132,000	36	138,543,000	38	143,923,000
0	0	0	0	0	0
3	3,234,000	3	4,282,000	6	5,346,000
0	0	0	0	0	0
0	2,388,000	0	2,440,000	0	2,440,000
35	118,754,000	39	145,265,000	44	151,709,000
T					
324	38,294,000	352	43,170,000	362	44,893,000
0	0	0	0	0	0
0	0	0	0	0	0
1	77,000	1	101,000	1	117,000
0	1,030,000	0	1,052,000	0	1,052,000
77	16,608,000	87	18,075,000	87	16,836,000
402	56,009,000	440	62,398,000	450	62,898,000
4,819	2,315,588,000	4,949	2,413,312,000	4,715	2,448,641,000
T				<del></del>	
FTTPs		<u>FTTPs</u>		FTTPs	
173	7,100,000	166	6,973,000	160	6,973,000
1,113	51,706,000	1,178	54,985,000	1,132	54,395,000
1,286	58,806,000	1,344	61,958,000	1,292	61,368,000
296	1,241,494,000	253	1,039,022,000	271	1,167,456,000
(2)	(192,000)	(0)	(0)	(0)	(0)
FTEs		FTEs		FTEs	
763	490,639,000	763	530,671,000	763	539,207,000
1			, ,		212,723,000
0	0	0	0	0	0
				-	30,000,000
	0		0		0
1,443		1,507		1,507	4,459,395,000
1 ,					(38,983,000)
<del>                                     </del>					(730,893,000)
	No.  2,956 (119)  304 863 1 1,168 4,124 258 4,382 32 0 3 0 0 35  324 0 0 1 0 77 402 4,819  FTTPs 173 1,113 1,286 296 (2) FTEs 763 680	FY 2004	FY 2004         Actual         April No.           No.         Amount         No.           2,956         \$1,423,484,000         3,086           (119)         62,821,000         (70)           304         125,693,000         284           863         431,539,000         796           1         126,000         1           1,168         557,358,000         1,081           4,124         2,043,663,000         4,167           258         97,162,000         303           4,382         2,140,825,000         4,470           32         113,132,000         36           0         0         0           3         3,234,000         3           0         0         0           3         3,234,000         39           35         118,754,000         39           324         38,294,000         352           0         0         0           0         0         0           1         77,000         1           0         1,030,000         0           77         16,608,000         87           402         56,009	FY 2004   Actual   Appropriation   No.   Amount   No.   Amount   Amount   2,956   \$1,423,484,000   (119)   62,821,000   (70)   17,250,000   304   125,693,000   284   113,290,000   1   126,000   1   126,000   1   126,000   1   126,000   1   126,000   1   126,000   1,168   557,358,000   1,081   478,567,000   4,124   2,043,663,000   4,167   2,107,735,000   258   97,162,000   303   97,914,000   4,382   2,140,825,000   4,470   2,205,649,000   33   3,234,000   36   138,543,000   0   0   0   0   0   0   0   0   0	FY 2004   Actual   Appropriation   No.   Amount   No.   Amount   No.   Amount   No.   Amount   No.

#### **Budget Authority by Activity**

		(dolla	rs in the	ousands)				
	FY 2004 Actual		FY 2005 Appropriation			Y 2006 Estimate	(	Change
ACTIVITY	FTEs	Amount	FTEs	Amount	FTEs	Amount	FTEs	Amount
Extramural Research: Allergy, Immunology, and								
Infectious Diseases		\$3,615,888		\$3,663,092		\$3,707,465		\$44,373
	:							
Subtotal, Extramural research		3,615,888		3,663,092		3,707,465		44,373
Intramural research	763	490,639	763	530,671	763	539,207	0	8,536
Research management & support	680	196,513	744	209,078	744	212,723	0	3,645
Cancer Control & Prevention	0	0	0	0	0	0	0	0
Total	1,443	4,303,040	1,507	4,402,841	1,507	4,459,395	0	56,554

<u>Summary</u>	of Change	<u> </u>		·-·
FY 2005 Appropriation				\$4,402,841,000
FY 2006 Estimated Budget Authority		<u> </u>		4,459,395,000
Net change				56,554,000
		FY 2005		
	Apj	propriation	Chang	ge from Base
		Budget		Budget
CHANGES	FTEs	Authority	FTEs	Authority
A. Built-in:				
1. Intramural research:				
a. Within grade increase	1	\$95,300,000		\$1,347,000
b. Animalization of January				
2005 pay increase		95,300,000		894,000
c. January 2006 pay increase	1	95,300,000		1,667,000
d. One less day of pay		95,300,000		(381,000)
e. Payment for centrally furnished services	ľ	74,027,000		370,000
f. Increased cost of laboratory supplies,				
materials, and other expenses		361,344,000		6,846,000
Subtotal				10,743,000
2. Research Management and Support:				
a. Within grade increase		82,392,000		1,408,000
b. Annualization of January				
2005 pay increase		82,392,000		775,000
c. January 2006 pay increase		82,392,000		1,446,000
d. One less day ofpay		82,392,000		(331,000)
e. Payment for centrally furnished services		27,013,000		135,000
f. Increased cost of laboratory supplies,				
materials, and other expenses		99,673,000		1,829,000
Subtotal				5,262,000
Subtotal, Built-in				16,005,000

#### **Summary of Changes—continued**

	T	FY 2005		
	A	ppropriaton	Char	ige from Base
CHANGES	No.	Amount	No.	Amount
B. Program:				
1. Research project grants:	İ			
a. Noncompeting	3,086	\$1,629,168,000	(132)	(\$141,175,000)
b. Competing	1,081	478,567,000	(126)	166,211,000
c. SBIR/STTR	303	97,914,000	9	3,349,000
Total	4,470	2,205,649,000	(249)	28,385,000
2. Research centers	39	145,265,000	5	6,444,000
3. Other research	440	62,398,000	10	500,000
4. Research training	1,344	61,958,000	(52)	(590,000)
5. Research and development contracts	253	1,039,022,000	271	128,434,000
Subtotal, extramural				163,173,000
	<u>FTEs</u>		<u>FTEs</u>	
6. Intramural research	763	530,671,000	0	(2,207,000)
7. Research management and support	744	209,078,000	0	(1,617,000)
8. Cancer control and prevention	0	0	0	0
9. Construction		148,800,000		(118,800,000)
10. Building and Facilities		0		0
Subtotal, program		4,402,841,000		40,549,000
Total changes	1,507		0	56,554,000

**Budget Authority by Object** 

Buuget Auti	hority by Object	r	<del></del>
			<u> </u>
	FY 2005	FY 2006	Increase or
· · · · · · · · · · · · · · · · · · ·	Appropriation	Estimate	Decrease
Total compensable workyears:			
Full-time employment	1,507	1,507	0
Full-time equivalent of overtime & holiday hours	5	5	0
Average ES salary	\$0	\$0	\$0
Average GM/GS grade	11.4	11.4	0.0
Average GW/ GB grade	11	11	0.0
Average GM/GS salary	\$73,995	\$75,697	\$1,702
Average salary, grade established by act of			
July 1, 1944(42 U.S.C. 207)	\$79,356	\$81,181	\$1,825
Average salary of ungraded positions	101,817	104,159	2,342
	FY 2005	FY 2006	Increase or
OBJECT CLASSES	Appropriation	Estimate	Decrease
Personnel Compensation:			
11.1 Full-Time Permanent	76,483,000	79,548,000	3,065,000
11.3 Other than Full-Time Permanent	43,425,000	45,125,000	1,700,000
11.5 Other Personnel Compensation	4,253,000	4,423,000	170,000
11.7 Military Personnel	3,345,000	3,476,000	131,000
11.8 Special Personnel Services Payments	15,476,000	16,066,000	590,000
Total, Personnel Compensation	142,982,000	148,638,000	5,656,000
12.0 Personnel Benefits	33,151,000	34,464,000	1,313,000
12.1 Military Personnel Benefits	1,488,000	1,546,000	58,000
13.0 Benefits for Former Personnel	70,000	73,000	3,000
Subtotal, Pay Costs	177,691,000	184,721,000	7,030,000
21.0 Travel & Transportation of Persons	8,145,000	8,212,000	67,000
22.0 Transportation of Things	826,000 3,695,000	827,000	1,000
<ul><li>23.1 Rental Payments to GSA</li><li>23.2 Rental Payments to Odiers</li></ul>	1,764,000	3,714,000	19,000
23.3 Communications, Utilities &	1,704,000	1,772,000	8,000
Miscellaneous Charges	5,958,000	6,153,000	195,000
24.0 Printing & Reproduction	837,000	837,000	1,53,000
25.1 Consulting Services	4,894,000	4,938,000	44,000
25.2 Other Services	136,041,000	137,862,000	1,821,000
25.3 Purchase of Goods & Services from		,,	1,021,000
Government Accounts	386,328,000	398,929,000	12,601,000
25.4 Operation & Maintenance of Facilities	28,170,000	28,505,000	335,000
25.5 Research & Development Contracts	924,783,000	1,051,173,000	126,390,000
25.6 Medical Care	3,661,000	3,706,000	45,000
25.7 Operation & Maintenance of Equipment	9,727,000	9,820,000	93,000
25.8 Subsistence & Support of Persons	-	<u>-</u>	-
25.0 Subtotal, Other Contractual Services	1,493,604,000	1,634,933,000	141,329,000
26.0 Supplies & Materials	39,647,000	39,945,000	298,000
31.0 Equipment	37,861,000	38,219,000	358,000
32.0 Land and Structures	33,000	33,000	-
33.0 Investments & Loans	-	-	-
41.0 Grants, Subsidies & Contributions	2,632,760,000	2,540,009,000	(92,751,000)
42.0 Insurance Claims & Indemnities	<u>-</u>	-	-
43.0 Interest & Dividends	20,000	20,000	-
44.0 Refunds	4 225 150 000	4 274 674 000	40.524.000
Subtotal, Non-Pay Costs	4,225,150,000	4,274,674,000	49,524,000
Total Budget Authority by Object	4,402,841,000	4,459,395,000	56,554,000

#### **Salaries and Expenses**

	FY 2005	FY 2006	Increase or
OBJECT CLASSES	Appropriation	Estimate	Decrease
Personnel Compensation:			
Full-Time Permanent (11.1)	\$76,483,000	\$79,548,000	\$3,065,000
Other Than Full-Time Permanent (11.3)	\$43,425,000	\$45,125,000	\$1,700,000
Other Personnel Compensation (11.5)	\$4,253,000	\$4,423,000	\$170,000
Military Personnel (11.7)	\$3,345,000	\$3,476,000	\$131,000
Special Personnel Services Payments (11.8)	\$15,476,000	\$16,066,000	\$590,000
Total Personnel Compensation (11.9)	\$142,982,000	\$148,638,000	\$5,656,000
Civilian Personnel Benefits (12.1)	\$33,151,000	\$34,464,000	\$1,313,000
Military Personnel Benefits (12.2)	\$1,488,000	\$1,546,000	
Benefits to Former Personnel (13.0)	\$70,000	\$73,000	\$3,000
Subtotal, Pay Costs	\$177,691,000	\$184,721,000	\$7,030,000
Travel (21.0)	\$8,145,000	\$8,212,000	\$67,000
Transportation of Things (22.0)	\$826,000	\$827,000	\$1,000
Rental Payments to Others (23.2)	\$1,764,000	\$1,772,000	\$8,000
Communications, Utilities and			
Miscellaneous Charges (23.3)	\$5,958,000	\$6,153,000	\$195,000
Printing and Reproduction (24.0)	\$837,000	\$837,000	\$0
Other Contractual Services:			
Advisory and Assistance Services (25.1)	\$4,894,000	\$4,938,000	\$44,000
Other Services (25.2)	\$136,041,000	\$137,862,000	\$1,821,000
Purchases from Govt. Accounts (25.3)	\$386,328,000	\$398,929,000	\$12,601,000
Operation & Maintenance of Facilities (25.4)	\$28,170,000	\$28,505,000	\$335,000
Operation & Maintenance of Equipment (25.7)	\$9,727,000	\$9,820,000	\$93,000
Subsistence & Support of Persons (25.8)	\$0	\$0	\$0
Subtotal Other Contractual Services	\$565,160,000	\$580,054,000	\$14,894,000
Supplies and Materials (26.0)	\$39,647,000	\$39,945,000	\$298,000
Subtotal, Non-Pay Costs	\$622,337,000	\$637,800,000	\$15,463,000
Total, Administrative Costs	\$800,028,000	\$822,521,000	\$22,493,000

#### NATIONAL INSTITUTES OF HEALTH

# **National Institute for Allergy and Infectious Disease**

# SIGNIFICANT ITEMS IN HOUSE AND SENATE APPROPRIATIONS COMMITTEE REPORTS

## FY 2005 House Appropriations Committee Report Language (H. Rpt. 108-636)

# Item

**Asthma** - The Committee is pleased with NIAID's leadership regarding asthma research and management. The Committee encourages NIAID to continue to improve its focus and effort on asthma management, especially as it relates to children. The Committee also encourages NIAID to collaborate more aggressively with voluntary health organizations to support asthma prevention, treatment, and research activities. Additionally, recent studies suggest that a variety of viral and bacterial agents may play a role in the development of asthma. The Committee suggests the Institute expand research into the role that infections and vaccines may play in the development of asthma, (p. 82)

#### Action to be taken

Recognizing the impact treatment for asthma attacks has on healthcare delivery systems, specifically emergency department visits, and asthma morbidity, the National Institute of Allergy and Infectious Diseases (NIAID) continues to pursue research toward the prevention, treatment, and management of asthma. An example of NIAID's research efforts is the Inter-City Asthma Study (ICAS) and the Inner-City Asthma Consortium (ICAC) cosponsored by the National Institute for Environmental Health Sciences (NIEHS).

The ICAS demonstrated that a physician feedback intervention reduced unscheduled asthma visits for children with moderate to severe asthma and that an environmental intervention reduced unscheduled asthma visits and missed school days and led to approximately three weeks of additional symptom-free days per year. NIAID sponsored a workshop at the May 2004 annual meeting of the American Thoracic Society to present new results of the ICAS.

In 2004, the ICAC, comprised of community health organizations and health care providers nationwide, launched a cockroach allergen standardization protocol, a study to evaluate the usefulness of measurements of exhaled nitric oxide in the clinical management of asthma in children, and a birth cohort to investigate the allergic and environmental factors that contribute to the development of asthma in inner-city children. The ICAC continues to evaluate the safety and efficacy of promising immune-based therapies; investigate the mechanisms of action of immune-based therapies; develop and validate biomarkers of disease stage, progression, and therapeutic effect; and investigate the immunopathogenesis of asthma in inner-city children.

Advancing the understanding of the immune mechanisms underlying the development of asthma and the causes of asthma exacerbations, including the role of viral and bacterial infections,

remains a high priority for NIAID. For example, in FY2004, NIAID co-sponsored with the National Heart, Lung and Blood Institute (NHLBI) the initiative, "Immune System Development and the Genesis of Asthma," to support research on the early life changes in immune function that lead to the development of asthma and the cellular and molecular processes involved in the onset of asthma, including the effects of bacteria and viruses on the function of the developing immune system.

Additionally, NIAID has participated with other NIH Institutes and Centers, Federal agencies, and non-federal investigators in the Asthma Workgroup of the National Children's Study. Led by the National Institute of Child Health and Human Development, the National Children's Study plans to follow a cohort of 100,000 children from before birth to at least age 25 to evaluate the effects of environmental exposures on the natural history of diseases, including asthma.

NIAID will continue to support research to understand the mechanisms of asthma; in FY 2005, the Institute will co-sponsor with NHLBI the initiative, "Asthma Exacerbations: Biology and Disease Progression," to increase the understanding of the mechanisms involved in acute exacerbations of asthma, including the cellular and molecular processes that cause some viral infections to trigger asthma attacks.

# <u>Item</u>

Atopic dermatitis - Atopic dermatitis (AD) is one of the most common skin disorders experienced in infants and children. Over 90% of cases are diagnosed before the age of five. Patients with AD suffer with chronic skin inflammation and itching that disrupt sleep and reduce quality of life. An estimated 17 percent of children in the United States have atopic dermatitis, a dramatic increase above the pre-1960s level of approximately 3 percent. The reason for this increase is unknown, but mirrors the increased rates of asthma and requires greater study. Of additional concern, individuals who have active or dormant AD are at high risk for serious adverse reaction to the smallpox vaccine. The Committee encourages NIAID to work with NIAMS to spearhead a multidisciplinary, multi-institute initiative to encourage investigator-initiated research on AD as it relates to smallpox vaccination as well as the progression to asthma and other allergic diseases, (p. 83)

# Action taken or to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) and other components of the National Institutes of Health are committed to supporting research efforts aimed at reducing the incidence and severity of atopic dermatitis and its complications. For example, through the Immune Tolerance Network, NIAID-supported scientists are currently developing a clinical trial of children with atopic dermatitis and food allergy. This trial will help determine whether oral administration of allergens will prevent the development of asthma and other allergic diseases in children with atopic dermatitis.

In addition, NIAID remains committed to reducing adverse reactions to the smallpox vaccine. Because vaccinia, the virus used to vaccinate against smallpox, can be transmitted by close contact, individuals with atopic dermatitis are at risk not only if they are immunized with

vaccinia, but also if their families or other close contacts have been immunized. Protection against this complication is a critical element in biodefense programs. The Institute supports contracts to two pharmaceutical manufacturers for the advanced development of modified vaccinia Ankara (MVA), a live, non-replicating form of vaccinia virus, vaccine to prevent smallpox. These contracts require studies of the safety and efficacy of the MVA vaccine in immunocompromised individuals and those with atopic disorders. The goal of MVA-based vaccine research is to develop a vaccine that can be used safely in all populations who are at risk for complications from the current vaccine made from a live form of vaccine virus.

In FY 2004, NIAID, with expert advice and support from the National Institute of Arthritis and Musculoskeletal and Skin Diseases, established the Atopic Dermatitis and Vaccinia Immunization Network (ADVN). The ADVN will develop short- and long-term approaches to reduce the incidence and severity of eczema vaccinatum and protect individuals with atopic dermatitis from the adverse consequences of vaccinia exposure; several human and animal protocols are currently in development. Components of the ADVN include the Clinical Studies Consortium for investigating the immune system of atopic dermatitis patients, as well as their immune responses to cutaneous viruses; the Animal Studies Consortium for developing animal models of atopic dermatitis; and the Statistical and Data Coordinating Center for data analysis, clinical coordination, regulatory activities and patient registry development.

# <u>Item</u>

**Condom Effectiveness** - The Committee recognizes the interest in the June 2000 NIH-STD Condom Report and the attention that this report calls to the epidemic of sexually transmitted diseases (STDs) that the nation is experiencing. The Committee encourages NIH to continue its practice of making advances in STD research available to the public and to health practitioners through web sites and other publications, (p. 83-84)

#### Action to be taken

The National Institute of Allergy and Infectious Diseases (NIALD) supports a diverse portfolio of research toward more effective prevention and treatment approaches to control sexually transmitted infections (STIs). This research includes basic research on pathogenesis, immunity, molecular and structural biology of sexually transmitted pathogens, and the impact of STIs in various populations; research for safe and effective vaccines, topical microbicides, therapeutics, and strategies for prevention and treatment of STIs and their sequelae; and the development of rapid and more effective diagnostic tools for STIs.

In December 2002, NIAID participated in an NICHD-sponsored conference entitled "Critical Issues in Study Design of Research on Condoms and Prevention of Sexually Transmitted Infections," which followed up on those scientific gaps identified during the NIAID 2000 condom effectiveness workshop. The goal of this second workshop was to focus on the development of study design guidance for less experienced researchers to use when designing new research projects on condoms and STIs.

While the U.S. Centers for Disease Control and Prevention is the lead Federal agency for the publication and dissemination of educational material on public health, NIAID maintains, and will continue to maintain, a comprehensive web site on STI information for the general public and health care practitioners.

#### Item

**Genomics** - The Committee is pleased that NIAID has focused on research efforts associated with multiple categories of pathogens. The Committee understands that microarray technology has enhanced the progress of pathogen-related research. The Committee encourages NIAID to continue to use this technology to further support an aggressive agenda of pathogen research activities, (p. 83)

#### Action to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) maintains a robust extramural research portfolio in the area of microbial genomics and continues to encourage scientists to take full advantage of emerging genomic technologies, such as microarrays, to study key questions in microbiology and infectious diseases. For example, NIAID continues to support the Pathogen Functional Genomics Resource Center (PFGRC) at The Institute for Genomic Research, which was established in FY 2001. The PFGRC provides and distributes to the broader research community a wide range of genomic and related resources and technologies for the functional analysis of microbial pathogens, including those that cause emerging and re-emerging infectious diseases and those that are considered to be potential agents of bioterrorism.

The number of organism-specific microarrays produced and distributed through the PFGRC to the scientific community increased to twenty in FY 2004 and includes microarrays for the causative agents for infectious diseases such as SARS, anthrax, tuberculosis, and plague. NIAID will continue to support the PFGRC in FY 2005 when the PFGRC will make available to the scientific community additional genomic resources, including protein expression clone sets and DNA microarrays. In addition, the PFGRC will undertake the genomic analysis of additional human pathogens as well as a comparative genetic analysis to identify genetic variations and relatedness within and between species for forensic strain identification and for the development of therapeutics, vaccines, and diagnostics. The expansion will also include the development and validation of protein arrays for future distribution to the scientific community.

In addition to the PFGRC, NIAID supports two Microbial Genome Sequencing Centers, which allow for the rapid, high quality, and cost-efficient sequencing of microbial genomes. These Centers have the capacity to respond to national needs and Federal agencies' priorities for genome sequencing and providing genomic sequencing data for multiple usages, including basic research on the biology of microbes, forensic identification of strains, and the identification of targets for therapeutics, vaccines, and diagnostics to combat infectious diseases. Two additional initiatives launched in FY 2004, the Bioinformatic Resource Centers and the Biodefense Proteomics Research Programs, will provide comprehensive genomic, bioinformatics, functional genomics, and proteomic research resources to the basic and applied infectious diseases research community.

In addition to the extramural program in microbial genomics, NIAID also supports a Microarray Research Facility (MRF) to provide its intramural investigators with the expertise and resources for all phases of microarray-based research projects. NIAID scientists have used microarrays produced by the MRF to elucidate the genes involved in the host immune responses to viral hepatitis, to understand the molecular mechanisms underlying resistance to malarial drugs, and to examine the human immune response to *Borrelia burgdorferi*, the causative agent of Lyme disease.

Recognizing the significance of microbial genomics to infectious disease research, NIAID will continue to expand its genomics-related activities in this area.

# <u>Item</u>

**Hemophilia** - The Committee appreciates NIAID's efforts to improve HIV and hepatitis C virus (HCV) treatment for persons with hemophilia or other bleeding disorders and encourages NIAID to work with private non-profit groups to strengthen support for research on liver disease progression, (p. 82)

#### Action to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) remains committed to supporting research targeted to the hemophiliac population. Transmission of HIV to people with hemophilia has been virtually eliminated by the use of treated blood products. As the population of HIV-infected hemophiliacs declines, the focus of hemophilia research has shifted to the impact of hepatitis C virus. To meet the research needs in this area, the Institute is supporting a University of Cincinnati study of liver disease progression and HCV genomic variability in HIV-infected hemophiliacs. NIAID will continue to support this study in FY 2005.

NIAID and the National Hemophilia Foundation continue to encourage the participation of people with hemophilia in its extra- and intramural research studies, including clinical trials for HIV infection and its sequelae.

#### Item

**Hepatitis C vaccine** - The Committee is encouraged to learn that a small hepatitis C vaccine human trial has been awarded and urges the consideration of other creative approaches and new paradigms, including the development of DNA vaccines, (p. 82)

# Action taken or to be taken

Research to develop vaccines against hepatitis C virus (HCV) remains a high priority for the National Institute of Allergy and Infectious Diseases (NIAID). NIAID supports an HCV research agenda that includes basic research as well as the development of vaccine candidates, including preclinical evaluation and clinical trials of these candidate vaccines.

The Institute also supports the establishment of research resources to facilitate research towards HCV vaccines. For example, NIAID continues to foster the development of an *in vitro* culture system for HCV as well as new animal models for basic research and for adequately testing vaccine candidates and antiviral drugs. To this end, in FY 2005, NIAID plans to renew the Hepatitis Animal Model Network, which will focus on the development of animal models to screen therapies and vaccines for HCV as well as hepatitis B.

NIAID scientists continue their efforts to develop a DNA vaccine against hepatitis C using various forms of the hepatitis C envelope proteins El and E2. In 2004, the Institute launched a Phase I trial using Chiron Corporation's prototype E1/E2 hepatitis C vaccine. This study will evaluate the safety, tolerability and immunogenicity of the vaccine in healthy, uninfected human subjects. In addition, NIAID-supported investigators at the New York Blood Center found that recombinant vaccinia viruses carrying hepatitis C virus DNA provided strong cell-mediated immune responses. Primate studies to assess the immune responses to this recombinant virus are currently in progress.

NIAID is planning to hold a workshop on HCV vaccines in early 2005 to discuss and evaluate the current status of efforts towards development of HCV vaccines with the goal of spurring their development and testing.

#### <u>Item</u>

Inflammatory Bowel Disease - The Committee continues to note with interest a scientific research agenda for Crohn's disease and ulcerative colitis (collectively known as inflammatory bowel disease) entitled "Challenges to Inflammatory Bowel Disease (IBD)." This report identifies strong linkages between the functions of the immune system and IBD. The Committee encourages the Institute to enhance its support of research focused on the immunology of IBD, as well as the interaction of genetics and environmental factors in the development of the disease, (p. 82)

# Action to be taken

The National Institute of Allergy and Infectious Diseases (NIAID), through its extra- and intramural programs, maintains a strong commitment to basic and clinical research to improve the diagnosis, treatment, and prevention of inflammatory bowel disease (IBD). For example, NIAID scientists have completed a Phase II trial of a novel immune-based therapy for Crohn's disease, monoclonal anti-interleukin-12 antibody, and have demonstrated that it is an effective treatment for active Crohn's disease. This finding is paving the way for a Phase III trial in a large patient cohort.

The Institute, in cooperation with the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK), the National Institute of Arthritis and Musculoskeletal and Skin Diseases (NIAMS), and the National Institutes of Health (NIH) Office of Research on Women's Health (ORWH), supports the Autoimmunity Centers of Excellence (ACEs), which conduct collaborative basic and clinical research on multiple autoimmune diseases, including clinical trials and mechanistic studies of immunomodulatory therapies. A Phase II clinical trial of anti-

CD3 (Visilizumab) therapy for ulcerative colitis is currently under development to be conducted through the ACEs.

NIAID intramural researchers have expanded their investigations of IBD, including basic studies of the immunology of IBD. NIAID scientists recently published an important finding which explains the mechanism by which certain mutations (CARD15/NOD2) cause Crohn's diseases. The study reveals a new mechanism by which immune responses to bacterial products are regulated; abnormalities in this regulation lead to inflammation and Crohn's disease. Intramural researchers are also conducting other studies to evaluate the safety and efficacy of therapies such as G-CSF in Crohn's diseases and interferon-beta in ulcerative colitis. Lastly, NIAID scientists are conducting a longitudinal study of IBD that combines immune monitoring and conventional therapy; data from this study will allow investigators to dissect the interaction of genetics and environmental factors in the development of disease.

The support of research to understand how genetics and environmental factors play a role in the development of IBD remains a priority for the Institute. For example, NIAID supports the Multiple Autoimmune Diseases Genetics Consortium, which collects clinical information and DNA and cell samples from families who have members with more than one autoimmune disease, including IBD. The information and samples collected from the more than 260 families who have enrolled to date will be used for genetic studies to understand the genetic factors that play a role in the development of autoimmune diseases.

NIAID also supports research to study the role of infectious agents in the development of LBD. For example, the Institute continues to fund several research projects focused on IBD through the initiative, "Microbial Etiology of Chronic Diseases," including research that is studying the possible role of *Mycobacterium paratuberculosis* in Crohn's disease. In addition, NIAID is planning to hold a workshop in the summer of 2005 to discuss the most recent research on IBD and to identify scientific opportunities in this area.

#### Item

*Meningococcal disease* - Although meningococcal disease is vaccine-preventable in most cases, approximately 30 percent of the deaths and disabilities from this bacterial infection are attributed to serogroup B, which is not vaccine-preventable. The Committee encourages NIAID to increase research efforts to develop an effective, low-cost vaccine against serogroup B that will help protect infants and adolescents, (p. 83)

# Action to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) is committed to supporting and conducting research to prevent meningococcal disease. In October 2004, NIAID and the National Vaccine Program Office co-sponsored a workshop titled "Carbohydrate Moieties as Vaccine Candidates." This workshop brought together research scientists, clinicians, and representatives from industry to identify research needs and scientific gap areas in an effort to promote vaccine development for meningococcal disease. The workshop examined the mechanisms involved in generating an appropriate immune response to selected antigens,

highlighted recent advances and discussed how this information could be used in the development of effective vaccines. Additional focus was given to discussing the obstacles involved with developing a vaccine against serogroup B meningococcus. The fruits of this discussion are still under evaluation for potential use in generating a serogroup B vaccine.

The Institute continues to support pre-clinical and clinical studies to control selected human respiratory pathogens through its Respiratory Pathogen Research Units (RPRUs). Respiratory pathogens studied at the RPRUs include meningococci, pneumococci, group A streptococcus, pseudomonas, *Chlamydia pneumoniae* and non-typable *Haemophilus influenzae*.

NIAID will continue to pursue research in support of development and licensing of vaccines and therapeutic agents for respiratory pathogens. Among the Institute's goals are to further understand the etiology and long-term health impact of acute respiratory infections, and to stimulate basic research to provide additional information on the pathogenesis, immunity, and functional components of respiratory pathogens.

#### Item

Nasal aerosol and spray vaccine delivery systems - Recent developments exploring new routes of immunization such as delivery of measles vaccine via aerosol and nasal spray may generate significant savings and result in fewer side effects than immunization by injection. The Committee encourages NIAID to support research in developing and testing these new approaches, building upon the testing already completed in older children, to investigate this delivery method in younger children, (p. 83)

# Action to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) intra- and extramural research programs have a long history of research and development of vaccines against childhood diseases and of research on the nasal delivery of vaccines. For example, NIAID both conducted and supported the research and development of intranasally administered live, attenuated influenza vaccines. This work was critical to the development of FluMist, the licensed nasal spray influenza vaccine.

NIAID researchers are developing several candidate vaccines that are specifically designed for nasal delivery. For example, an experimental intranasal combination vaccine against respiratory syncytial virus (RSV), the most important pediatric respiratory pathogen worldwide, and parainfluenza virus type 3 (PTV3) has been tested for safety and immunogenicity in children 6-18 months of age; further studies in larger numbers of young children are planned. NIAID researchers are also developing nasally administered vaccine candidates against human metapneumovirus (HMPV), a virus first identified in 2001 that is a significant cause of respiratory tract disease around the world.

Scientists at NIAID have also developed a live, attenuated vaccine for severe acute respiratory syndrome (SARS) that is administered directly to the respiratory tract. In preclinical studies, a

single immunization with this vaccine induced a high level of protective immunity. This experimental vaccine would be appropriate for further evaluation in infants and young children, for whom the vaccine might be more effective than for adults.

In addition to intramural research, the Institute also supports extramural research on alternative vaccine delivery systems, including nasal vaccine delivery. In December 2003, NIAID cosponsored, with the Department of Health and Human Services, the Centers for Disease Control and Prevention, and the Food and Drug Administration, a conference on innovative administration systems for vaccines, including jet injectors, transdermal administration (i.e., a patch), and transmucosal administration (i.e., oral or nasal administration). The conference was a forum for academic, clinical, and industry communities as well as Federal agencies to discuss the state of the science and current status of the development of the various vaccine delivery techniques.

NIAID will continue to conduct and support the research and development of novel approaches and technologies for vaccine candidates that are suitable for nasal delivery in young children and to test promising candidates in clinical trials.

# <u>Item</u>

**Scleroderma** - The Committee encourages NIAID to conduct research to study the cause and treatment of scleroderma, a chronic progressive disease that predominantly strikes women. Scleroderma is disfiguring and can be life-threatening, affecting multiple systems. More research is needed in order to develop safe, effective treatments and to identify the causes of scleroderma and its complications. NIAID is encouraged to consider including scleroderma as one of the diseases in the Autoimmune Centers of Excellence (ACE) program in order to address these important questions, (p. 83)

# Action to be taken

Although we have gained considerable understanding of the mechanisms that mediate tissue injury in autoimmune diseases, significant gaps remain. The National Institute for Allergy and Infectious Diseases (NIAID) remains committed to understanding the cause and improving the treatment of scleroderma, an autoimmune disease. For example, the Autoimmunity Centers of Excellence (ACEs) conduct collaborative basic and clinical research on autoimmune diseases, including scleroderma.

The ACEs support close interaction between clinicians and basic researchers to facilitate the identification of effective tolerance induction and immune modulation strategies to treat or prevent disease, and accelerate the translation of scientific advances to the clinic. NIAID expanded the number of ACEs from four to nine in FY 2004. This expansion will enable a wider range of autoimmune diseases to be studied as it facilitates collaboration and draws on the expertise of a larger network of scientists and physicians. The ACEs are cosponsored by NIAID, the National Institute for Diabetes and Digestive and Kidney Diseases (NIDDK) and National Institutes of Health (NIH) Office of Research on Women's Health (ORWH).

In addition to the research supported through the ACEs, NIAID will continue to pursue research to further understanding of the mechanisms of and treatment for autoimmune diseases including scleroderma. For example, the Stem Cell Transplantation for Autoimmune Diseases Consortium is developing a clinical trial to assess the efficacy of hematopoietic stem cell transplantation to treat scleroderma. The consortium will also conduct studies of the immune mechanisms underlying scleroderma.

Finally, in December 2004, the NIH Autoimmune Diseases Coordinating Committee plans to submit its second report to Congress. This report will summarize NTH funding for autoimmune diseases research, and accomplishments and activities, including ongoing research projects and future initiatives.

#### Item

**Transplantation research** - The Committee urges NIAID to convene an expert conference during fiscal year 2005 to develop a Transplantation Research Action Plan identifying the most urgently needed research to facilitate an increase in the success of organ transplantation. The Committee requests a report on the results of this conference including a breakdown of resources committed to this category of research, (p. 83)

#### Action to be taken

Although one-year organ transplantation survival has improved markedly over the last fifteen years, there has been little success in reversing the decline in long-term graft and patient survival. The National Institute of Allergy and Infectious Diseases (NIAID), through its extra-and intramural programs, supports an extensive portfolio of research aimed at understanding the mechanisms of immune-mediated graft rejection in solid organ, tissue and cell transplantation. Understanding these mechanisms will lead to the development of immunosuppressive therapies with fewer side effects, reductions in the numbers of re-transplants, and improvement in long-term graft survival without the need for life-long, global immunosuppressive therapy.

An example of NIAID support for clinical research on solid organ, tissue, and cell transplantation is the Cooperative Clinical Trials in Pediatric Transplantation (CCTPT) program, established in 1994, to conduct multicenter clinical trials of novel approaches to prevent acute and chronic graft rejection in pediatric kidney transplant recipients. Also, in FY 2004, NIAID launched the Clinical Trials in Organ Transplantation consortium, a multicenter initiative to evaluate novel therapies for preventing graft rejection and prolonging transplant graft survival in kidney, liver, and heart transplantation. This consortium, cosponsored by the National Institute of Diabetes and Digestive and Kidney Diseases and the National Heart, Lung, and Blood Institute, has been charged with defining a five-year scientific agenda for clinical research in organ transplantation and with implementing collaborative multicenter clinical trials in organ transplantation with associated mechanistic studies.

In FY 2005, the Institute will continue to support research to understand the mechanisms whereby the immune system recognizes and either accepts or rejects transplanted organs, tissues or cells. For example, NIAID will support the initiative, "HLA Region Genetics in Immune-

Mediated Diseases," which will support research to identify the variations in immune response genes that may account for the increased susceptibility of certain individuals to immunemediated diseases, such as graft rejection.

NIAID will lead NIH-wide efforts to convene an expert panel on transplantation research in FY 2005. This panel will assess basic and clinical research programs and make recommendations for future efforts. These recommendations will form the basis of the Transplantation Research Action Plan, which will be provided to Congress.

#### Item

**Tuberculosis** - The World Health Organization estimates that nearly one billion people will become infected with tuberculosis (TB), 200 million will become sick, and 70 million will die worldwide between now and 2020 of this disease. The Committee is pleased with NIAID's efforts to develop an effective TB vaccine. The Committee encourages the Institute to continue its TB vaccine development work and to expand efforts to develop new drugs to treat TB. (p. 82)

# Action taken or to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) Tuberculosis (TB) Program supports research to gain in-depth knowledge about *Mycobacterium tuberculosis* (Mtb) and how the body responds to this invasive pathogen, and to translate this knowledge into improved health care interventions. The focus of the program is to develop drugs that will shorten TB therapy and make it easier for patients to complete therapy, as well as to develop vaccines that will lead to long-lasting protection against TB.

This year, as a result of NIAID's longstanding support for TB vaccine development, the first new TB vaccine in 60 years underwent human clinical testing in the United States. The trial is being conducted by Corixa and GlaxoSmithKline Biologicals. NIAID funding was instrumental in the discovery, development and animal testing of this vaccine candidate. A second TB vaccine developed with NIAID grant support also entered human clinical trials recently; this candidate vaccine is a derivative of the currently available TB vaccine, BCG.

A key component of NIAID's successful contribution to TB vaccine development is the recently renewed TB Research Materials and Vaccine Testing contract. This contract provides exploratory and preclinical evaluation of promising new TB vaccine candidates in animal models. In addition, TB researchers around the world can request a variety of Mtb-derived research reagents from the TB Research Materials and Vaccine Testing program, which enables them to study the disease without handling contagious and technically demanding mycobacterial pathogens.

Many promising new anti-TB drug candidates, developed by NIAID-supported scientists, are now nearing initial evaluations in humans. The most promising candidate, SQ109, which is being developed as a part of NIAID's Challenge Grant program, is nearing completion of preclinical development and may enter human trials in 2005. Another drug, PA-824, is currently

in preclinical development by the Global Alliance for TB Drug Development, a public-private partnership involving NIAID.

NIAID also maintains a robust intramural research program on TB. For example, NIAID scientists are collaborating with colleagues at GlaxoSmithKline and St. Jude Children's Research Hospital to develop an improved drug to treat TB based on the drug thiolactomycin. In addition, researchers at the Institute are synthesizing analogs of PA-824 to elucidate the drug's mechanism of action and to improve the characteristics of this class of compounds as drug candidates. This work has been enhanced by NIAID scientists' development of a novel DNA microarray tool for determining the molecular mechanisms of anti-TB drug action. This tool will allow scientists studying new drugs against TB to immediately understand their mechanism of action, and will greatly facilitate the drug discovery process.

FY 2006 Senate Appropriations Committee Report Language (S. Rpt. 108-345)

#### Item

**Arthritis** - The Committee encourages the NIAID to coordinate its research efforts with other NIH institutes to find a cure for arthritis and related diseases, (p. 125)

# Action taken or to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) collaborates with other National Institutes of Health (NIH) Institutes and Centers (ICs) as well as with private research groups to conduct research toward a cure for rheumatoid arthritis, which afflicts more than 2 million Americans. For example, NIALD, along with the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK), and the NIH Office for Research on Women's Health (ORWH), supports nine Autoimmunity Centers of Excellence (ACEs), which conduct collaborative basic and clinical research on autoimmune diseases, including rheumatoid arthritis. Phase II studies of Anti-CD20 and Lovastatin for the treatment of rheumatoid arthritis are currently in development to be conducted through the ACEs.

The Institute also sponsors with NICHD, NIDDK, ORWH, and JDRF, the Cooperative Study Group on Autoimmune Disease Prevention. This group currently supports three core projects and three pilot projects to investigate the prevention of rheumatoid arthritis.

Another NIAID-supported program, the International Histocompatibility Working Group (IHWG), which is cosponsored by the National Cancer Institute, NIDDK, the National Human Genome Research Institute, the National Center for Biotechnology Information at the National Library of Medicine, the Centers for Disease Control and Prevention, and the Juvenile Diabetes Research Foundation, aims to identify the variations in immune response genes that may account for the increased susceptibility of certain individuals to immune-mediated diseases including rheumatoid arthritis. This program will be renewed in FY 2005 as the initiative, "HLA Region Genetics in Immune-Mediated Diseases."

NIAID further collaborates with other NIH ICs in supporting the "Hyperaccelerated Award/Mechanisms in Immunomodulation Trials". The National Institute of Arthritis and Musculoskeletal and Skin Diseases (NIAMS), NIDDK, and the National Institute of Neurological Disorders and Stroke also support this initiative which conducts mechanistic studies to investigate at the molecular level how immunomodulatory interventions can impact immune-mediated diseases, such as rheumatoid arthritis. NIAID also cosponsors with NIAMS and the Arthritis Foundation the North American Rheumatoid Arthritis Consortium, a registry and repository of information on families with rheumatoid arthritis.

Lastly, the NIH Autoimmune Diseases Coordinating Committee (ADCC), chaired by NIAID, was established in FY 1998 at the request of Congress to increase collaboration and facilitate coordination of research among NIH ICs, other federal agencies, and private organizations and patient advocacy groups with an interest in autoimmune diseases, including rheumatoid arthritis. In December 2004, the ADCC will submit its second report to Congress. This report will summarize NIH funding for autoimmune diseases research, and accomplishments and activities, including ongoing research projects and future initiatives that address components of the ADCC Autoimmune Diseases Plan.

#### Item

**Asthma** - The Committee is pleased with NIAID's leadership regarding asthma research and management. The Committee urges NIAID to continue to improve its focus and effort on asthma management, especially as it relates to children. The Committee also urges the NIAID to collaborate more aggressively with voluntary health organizations to support asthma prevention, treatment, and research activities. Additionally, recent studies suggest that a variety of viral and bacterial agents, including agents used for immunization may play a role in the development of asthma. The Committee urges the Institute to expand research into the role that infections and vaccines play in the development of asthma, (p. 125)

#### Action to be taken

Please refer to page 37 of this document for NIAID's response to this significant item regarding asthma.

#### Item

Atopic dermatitis - Atopic dermatitis (AD) is one of the most common skin disorders experienced by infants and children. Patients with AD suffer with chronic skin inflammation and itching that disrupt sleep and reduce quality of life. Of additional concern, individuals who have active or dormant AD are at high risk for serious adverse reaction to the smallpox vaccine. The Committee encourages NIAID to work with NIAMS to spearhead a multidisciplinary, multi-institute initiative to encourage investigator-initiated research on AD as it relates to smallpox vaccination as well as the progression to asthma and other allergic diseases, (p. 125)

# Action taken or to be taken

Please refer to page 38 of this document for NIAID's response to this significant item regarding atopic dermatitis.

#### **Item**

*Genomics* - The Committee is pleased that the NIAID has focused its attentions on research efforts associated with multiple categories of pathogens. The Committee understands that commercially available research tools, such as microarray technology, can enhance the progress of pathogen-related research. The Committee urges the NIAID to utilize this technology to further support an aggressive agenda of pathogen research activities, (p. 125)

#### Action to be taken

Please refer to page 40 of this document for NIAID's response to this significant item on genomics.

# <u>Item</u>

*Hemophilia* - The Committee appreciates NIAID's efforts to improve HIV and hepatitis C virus (HCV) treatment for persons with hemophilia or other bleeding disorders. The Committee encourages NIAID to work with the National Hemophilia Foundation in strengthening its support for research on liver disease progression to improve HIV and HCV treatment among persons with bleeding disorders, (p. 125)

# Action to be taken

Please refer to page 41 for NIAID's response to this significant item of hemophilia.

# <u>Item</u>

**Hepatitis C vaccine** - The Committee is encouraged to learn that a small hepatitis C vaccine human trial has been awarded and urges the consideration of other creative approaches and new paradigms, including the development of DNA vaccines, (p. 125)

# Action taken or to be taken

Please refer to page 41 of this document for NIAID's response to this significant item regarding hepatitis C vaccine.

#### Item

*Hepatitis* - The Committee continues to be concerned about the prevalence of hepatitis and urges NIAID to work with public health organizations to promote liver wellness, education, and prevention of hepatitis, (p. 125)

# Action taken or to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) supports biomedical research, spanning basic through clinical research, toward the prevention and treatment of viral hepatitis. Although the U. S. Centers for Disease Control and Prevention is the lead agency for outreach and education activities related to public health, NIAID does support some activities to promote liver wellness and education on hepatitis. For example, the NIAID web site includes several publications that focus on hepatitis disease management. In addition, the NIAID fact sheet on hepatitis C includes links to web sites for public health organizations that promote liver wellness and hepatitis prevention.

NIAID also educates clinicians and researchers about advances in hepatitis research and about resources available through the Institute. In December 2003, NIAID staff presented a talk at the "Frontiers in Drug Development for Viral Hepatitis" to promote NIAID resources and reagents available to the hepatitis research community, such as repository contents and available therapeutic screening options. In November 2004, NIAID participated in the Princeton Hepatitis B Virus Workshop, an annual workshop sponsored by the Hepatitis B Foundation to discuss critical issues in basic hepatitis research and disease management. This workshop is attended by investigators from academia, industry, and the government. The Hepatitis B Foundation supports hepatitis research, promotes disease awareness, supports immunization and treatment initiatives and serves as a source of information for patients and their families, the medical and scientific communities, and the general public.

NIAID will continue to support a robust research agenda in viral hepatitis and to maintain a comprehensive web site on hepatitis information for the general public and health care practitioners.

# <u>Item</u>

Inflammatory Bowel Disease - Recent research identifies strong linkages between Crohn's disease and ulcerative colitis (collectively known as inflammatory bowel disease) and the functions of the immune system. The Committee encourages the Institute to expand its research partnerships with the IBD community, and to increase funding for research focused on the immunology of IBD and the interaction of genetics and environmental factors in the development of the disease, (p. 125-126)

#### Action to be taken

Please refer to page 42 of this document for NIAID's response to this significant item regarding inflammatory bowel disease.

## <u>Item</u>

*Juvenile Diabetes* - The Committee is encouraged by the Institute's efforts regarding the Autoimmune Prevention Centers, and it encourages the Institute to develop pilot projects,

initiated with the support of the Prevention Centers of Autoimmunity, into full research proposals with special emphasis on pre-clinical studies related to juvenile diabetes (p. 126)

#### Action to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) remains deeply committed to supporting research to improve the diagnosis, treatment, and prevention of many immune-mediated diseases, including type 1 diabetes. For example, in FY 2001, NIAID, the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK), the National Institute of Child Health and Human Development (NICHD), the National Institutes of Health (NIH) Office of Research on Women's Health (ORWH), and the Juvenile Diabetes Research Foundation International (JDRF) established the Centers for the Prevention of Autoimmune Diseases (Autoimmunity Prevention Centers) to conduct basic research on the development of new targets and approaches to prevent autoimmune diseases, with a special emphasis on type 1 diabetes, and to evaluate these novel approaches in pilot and clinical trials.

Since their inception, the Autoimmunity Prevention Centers have supported 25 pilot projects, eight of which are directly related to type 1 diabetes. For example, the Autoimmunity Prevention Centers recently funded a collaborative pilot project which seeks to describe the spectrum of immune responses and gene and protein expression in a mouse model of type 1 diabetes. This project will generate a publicly available dataset that may serve as a starting point for numerous hypothesis-driven projects related to the prevention of type 1 diabetes. Of the eight pilot projects on type 1 diabetes that are supported by the Autoimmunity Prevention Centers, two have been developed into full investigator-initiated research grants.

The objective of the Centers for the Prevention of Autoimmune Diseases is to continue to identify promising interventions, including vaccines, which may prevent the onset of type 1 diabetes. These candidate interventions will then be evaluated through NIAID clinical research programs such as the Autoimmunity Centers of Excellence, which is cosponsored by NIDDK, the National Institute of Arthritis and Musculoskeletal and Skin Diseases, and NIH ORWH; the Immune Tolerance Network, which is cosponsored by NIDDK and JDRF; and the Type 1 Diabetes TrialNet, which is cosponsored by NIDDK and NICHD. NIAID plans to renew the Centers for the Prevention of Autoimmune Diseases in FY 2006.

#### Item

**Nontuberculous Mycobacteria (NTM)** - Mycobacteria are environmental organisms found in both water and soil that can cause significant respiratory damage. The Committee is aware of the increasing incidence of nontuberculous mycobacteria (NTM) pulmonary infections in women, particularly involving rapidly growing mycobacteria, an inherently resistant subspecies. The Committee encourages NIAID to advance diagnostic and treatment protocols for patients suffering from NTM diseases, (p. 126)

# Action taken or to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) supports a broad range of research on respiratory pathogens, including non-tuberculosis mycobacteria (NTM). For example, NIAID scientists are conducting a clinical study to examine the symptoms, course of disease, and treatment of NTM infections, as well as the genetics involved in these infections. The goals of this study are to identify the critical mechanisms involved in mycobacterial resistance, identify and develop therapies based on these mechanisms of resistance, and develop novel therapies for the treatment of severe mycobacterial infections. These researchers have identified immune system abnormalities that may aid in diagnosis of NTM infection and screening of those at higher risk of NTM infection.

Many of NIAID's NTM research activities focus on NTM as a cause of opportunistic infections in HIV-infected individuals. For example, prior to the introduction of highly active antiretroviral therapy (HAART) for the treatment of HIV/AIDS, disseminated infection with *Mycobacterium avium* complex was a common, life threatening infection in HIV-positive patients. The standard of care was lifelong treatment with antibiotics. NIAID-supported scientists determined that this maintenance therapy can be safely discontinued in patients on HAART regimens. This change contributes significantly to an improved standard of care for **HrV**-infected persons.

The Institute awarded a seven year contract to California Pacific Medical Center Research Institute, San Francisco, in FY 2002 to test new drug candidates against *Mycobacterium avium* complex in cell culture and animal models. Additionally, N T M are used as model organisms in NIAID-supported research on tuberculosis. It is likely that such research may lead to further understanding of these mycobacteria.

NIAID will continue to support both investigator-initiated research and its intramural research program on NTM, including both basic research to understand the biology of NTM and research towards therapeutics to treat NTM infections.

#### Item

Primary Immunodeficiency Diseases — The Committee notes that more than 150 primary immune deficiency diseases have been identified to date. These diseases, which impair the body's immune system, strike more severely in children, many of whom do not survive beyond their teens or early twenties. Primary immune deficiencies afflict more than 50,000 Americans, regardless of age, race, or gender. The Committee believes that NIAID should play a significant role in addressing this seriously under-diagnosed class of diseases. Research is being funded at many institutions, including several of the twelve Jeffrey Modell Diagnostic and Research Centers, as part of a consortium created to expand and enhance the research in this group of diseases. However, that research only helps people if physicians know to look for the disease; the public is aware of it; and, patients are diagnosed early and accurately. For this reason, the Committee encourages NIAID to increase its support for the public outreach campaign principally funded by the CDC, while maintaining its research portfolio. The Committee commends NIAID for the establishment of its primary immunodeficiency disease research consortium (USIDNet) in partnership with the Immune Deficiency Foundation and encourages

continued support for this program, (p. 126)

#### Action to be taken

The National Institute of Allergy and Infectious Diseases (NIAID) is deeply committed to supporting research efforts aimed at understanding the causes and immune mechanisms leading to the development of primary immunodeficiency diseases and to working with other agencies within the Department of Health and Human Services to increase public awareness of primary immunodeficiency diseases.

The recently initiated U.S. Immunodeficiency Network (USIDNet) is sponsoring training sessions for new and young investigators as well as providing support for exploratory research. USIDNet provides leadership and mentoring; facilitates research collaborations; enhances the coordination of primary immunodeficiency diseases research efforts; and solicits, reviews, and makes awards for pilot and small research projects. In addition, USIDNet maintains a primary immunodeficiency diseases registry to provide data to the research community about the clinical characteristics and prevalence of these diseases and a repository of specimens from subjects with primary immunodeficiency diseases. The Consortium has funded nine research proposals and continues to review new research proposals three times a year, with the goal of funding 6-9 new research proposals each of the next several years.

NIAID continues to work with the Centers for Disease Control and Prevention and other organizations such as the Immune Deficiency Foundation and Jeffrey Modell Foundation in public outreach on primary immunodeficiency diseases. NIAID scientists are active members of the Medical Advisory Board of the Immune Deficiency Foundation. NIAID scientists also participate as expert speakers at patient advocate group meetings where they update patients and families about current and future therapies. In addition, NIAID researchers actively seek patients for enrollment in clinical studies of primary immunodeficiency disease and have developed information booklets about these diseases for distribution to patients and their families as well as to physicians in the community who may care for these patients. NIAID clinicians also provide lectures to the clinical community about inherited primary immunodeficiency diseases and participate in scientific meetings and conferences which disseminate important information about research and treatments these diseases.

In addition to public outreach activities, the Institute also supports research to improve the diagnosis of primary immunodeficiencies, such as the development of a computer algorithm to identify potentially immunodeficient hospital patients, including underdiagnosed minority patients, and the development of a testing system to allow the identification of some primary immunodeficiency diseases as part of newborn screening.

#### Item

**Scleroderma** - The Committee encourages NIAID to undertake research to study the cause and treatment of scleroderma, a chronic progressive disease that predominantly strikes women. Scleroderma is disfiguring and can be life-threatening, affecting multiple systems. More research is critically needed in order to develop safe, effective treatments and to identify the cause or

causes of scleroderma and its complications. Therefore, the Committee urges NIAID to include scleroderma research in the portfolio of the Autoimmune Centers of Excellence, (p. 126)

# Action taken or to be taken

Please refer to page 45 of this document for NIAID's response to this significant item regarding scleroderma.

#### Item

**Transplantation research** - The Committee urges NIAID to convene an expert conference during fiscal year 2005 to develop a Transplantation Research Action Plan identifying the most urgently needed research to facilitate an increase in the success of organ transplantation. The Committee requests a report on the results of this conference including a breakdown of resources committed to this category of research, (p. 127)

# Action to be taken

Please refer to page 46 of this document for NIAID's response to this significant item on transplantation research.

#### <u>Item</u>

**Tuberculosis** - The World Health Organization estimates that nearly 1 billion people will become infected with tuberculosis [TB], 200 million will become sick, and 70 million will die worldwide between now and 2020. The Committee is pleased with NIAID's efforts to develop an effective TB vaccine and encourages the Institute to continue its TB vaccine development work and to expand efforts to develop new drugs to treat TB. (p. 127)

#### Action taken or to be taken

Please refer to page 47 of this document for NIAID's response to this significant item regarding tuberculosis.

**Authorizing Legislation** 

PHS Act/ Other Citation	U.S.Code Citation	2005 Amount Authorized	FY 2005 Appropriation	2006 Amount Authorized	2006 Budget Estimate
Section 301	42§241	Indefinite		Indefinite	
Section 446	42§285f	Indefinite _J	\$4,340,883,000	Indefinite \(	\$4,398,027,000
Section 487(d)	42§288	a/	61,958,000	<b>b</b> /	61,368,000
			4 402 941 000		4,459,395,000
	Other Citation  Section 301  Section 446	Other Citation Citation  Section 301 42§241  Section 446 42§285f	Other Citation Citation Authorized  Section 301 42§241 Indefinite  Section 446 42§285f Indefinite _J	Other Citation Citation Authorized Appropriation  Section 301 42§241 Indefinite  Section 446 42§285f Indefinite _J  \$4,340,883,000	Other Citation Citation Authorized Appropriation Authorized  Section 301 42§241 Indefinite  Section 446 42§285f Indefinite

a/ Amounts authorized by Section 301 and Title IV of the Public Health Act.

 $<sup>\</sup>underline{b}$ / Reauthorizing legislation will be submitted.

**Appropriations History** 

Fiscal	Budget Estimate	House	Senate	
Year	to Congress	Allowance	Allowance	Appropriation <u>1/</u>
1997	\$584,362,000 <u>2/</u>	\$1,256,149,000	\$595,016,000 <u>3/</u>	\$1,257,794,000 <u>4/</u>
1998	634,272,000 21	1,339,459,000	1,359,688,000	1,351,655,000
1999	703,723,000 <u>2/5/</u>	1,470,460,000	1,540,102,000	1,570,102,000
Rescission	0	0	0	(1,039,000)
2000	789,156,000 <i>2[</i>	1,714,705,000	1,786,718,000	1,803,063,000
Rescission				(5,025,000)
2001	935,166,000 2[	2,062,126,000	2,066,526,000	2,069,388,000
Rescission				(1,084,000)
2002	2,355,325,000	2,337,204,000	2,375,836,000	2,535,778,000
Rescission				(1,239,000)
2003	3,983,693,000	2,674,213,000	3,727,473,000	3,730,973,000
Rescission				(24,251,000)
2004	4,335,255,000	4,335,255,000	4,335,255,000	4,335,155,000
Rescission	0	0	0	(30,593,000)
2005	4,440,007,000	4,440,007,000	4,456,300,000	4,440,007,000
Rescission	0			(37,166,000)
2006	4,459,395,000			

<sup>21</sup> Excludes funds for HIV Research Activities consolidated in the NIH Office of AIDS Research.

<sup>3/</sup> Excludes enacted administrative reductions of \$569,000,000.

<sup>4/</sup> Excludes enacted administrative reductions of \$575,000,000.

<sup>5/</sup> Reflects an increase of \$1,683,000 for the budget amendment for bioterrorism.

Detail of Full-Time Equivalent Employment (FTEs)

FY 2004 Actual	FY 2005	FY 2006
		FY 2006
Actual		
1	Appropriation	Estimate
256	271	271
67:	67	67
134	134	134
98	147	147
125	125	125
763	763	763
1,443	1,507	1,507
(6)	(6)	(6)
11.4 11.4 11.4		
11.4		
	134 98 125 763 1,443	67 67 134 134 134 98 147 125 125 763 763 763 763 1,443 1,507 (6) (6) Average GM/GS Gravel 11.4 11.4 11.4 11.4 11.4

T	•	T
Detail	ΛŤ	<b>Positions</b>

Detail of Positions					
GRADE	FY 2004 Actual	FY 2005 Appropriation	FY 2006 Estimate		
Total - ES Positions	0	0	0		
Total - ES Salary	\$0	\$0	\$0		
GM/GS-15	66	70	70		
GM/GS-14	209	221	221		
GM/GS-13	209	221	221		
GS-12	175	185	185		
GS-11	168	178	178		
GS-10	4	4	4		
GS-9	111	117	117		
GS-8	47	50	50		
GS-7	67	71	71		
GS-6	24	25	25		
GS-5	9	10	10		
GS-4	14	15	15		
GS.3	5	5	5		
GS-2	6	6	6		
GS-1	4	4	4		
Subtotal	1,118	1,182	1,182		
Grades established by Act of	1,110		,,,,,,,		
July 1, 1944 (42U.S.C. 207):					
(120.8.8. 207).					
Assistant Surgeon General	1	1	1		
Director Grade	18	18	18		
Senior Grade	15	15	15		
Full Grade	7	7	7		
Senior Assistant Grade	0	0	0		
Assistant Grade	1	1	1		
Subtotal	42	42	42		
Ungraded	429	429	429		
Total permanent positions	1,138	1,184	1,184		
Total positions, end of year	1,589	1,653	1,653		
Total full-time equivalent (FTE)					
employment,end of year	1,443	1,507	1,507		
Average ES salary	\$0	\$0	\$0		
Average GM/GS grade	11.4	11.4	11.4		
Average GM/GS salary	\$71,355	\$73,995	\$75,697		