(1) Provides leadership for and formulates research goals and long-range plans to accomplish the mission of the Human Genome Project, including the study of the ethical, legal, and social implications of human genome research; (2) fosters, conducts, supports, and administers research and research training programs in human genome research by means of grants, contracts, cooperative agreements, and individual and institutional research training awards; (3) provides coordination for genome research, both nationally and internationally, and serves as a focal point within NIH and the DHHS for Federal interagency coordination, collaboration with industry and academia, and international cooperation; (4) plans, supports and administers intramural, collaborative, and field research to study human genetic disease in its own laboratories, branches, and clinics; and (5) sponsors scientific meetings and symposia and collects and disseminates educational and informational materials related to human genome research to health professionals, the scientific community, industry, and the lay public.
Office of the Director - HN41

(1) Plans, directs, and coordinates the development and progress of the Institute's programs; (2) develops major policy and program decisions based on an evaluation of the status of support and accomplishments of the Institute's program areas; (3) coordinates grant review and program management activities; (4) plans and organizes conferences and workshops; and (5) communicates with the scientific community and coordinates activities with other private and government agencies.
Division of Intramural Research - HN45

(1) Plans and conducts a program of laboratory and clinical research related to the application of genome research to the understanding of human genetic disease and the development of human gene therapy; (2) acts as a focal point at NIH for genome research and maintains core facilities that serve as a resource for the entire NIH intramural research community; (3) evaluates research efforts and establishes intramural program priorities; (4) allocates funds, space, and personnel ceilings to ensure maximum utilization of available resources in the attainment of NHGRI objectives and integrates new research activities into the program structure; (5) collaborates with other NIH Institutes and Centers and external research institutions, and maintains an awareness of national and international research efforts in relevant program areas; and (6) advises the NHGRI Director and staff on intramural research programs and areas of science of interest to the Institute.
(1) Provides support and oversight for a wide variety of clinical research and training activities, including inpatient and outpatient services, education for genetic counseling studies, and training for M.D. and Ph.D. trainees engaged in the Metropolitan Washington Medical Genetics Training Program; (2) assures appropriate credentialing of NHGRI caregivers; (3) contributes to the operation of the NIH Clinical Center by providing genetic counseling consultations and participating in the Medical Executive Committee; (4) constitutes and supports the NHGRI Institutional Review Board and the Data Safety and Monitoring Board for oversight of clinical protocols; and (5) provides an infrastructure conducive to clinical research for NHGRI physicians and scientists directing investigational studies.
Medical Genetics and Genomic Medicine Training Program - HN4513

(1) Provides accredited training opportunities for Ph.D. and M.D. candidates that leads to Board Certification by the American Board of Medical Genetics; and (2) sponsors training leading to certification in Clinical Genetics, Medical Biochemical Genetics, Clinical Biochemical Genetics, Clinical Molecular Genetics, and Clinical Cytogenetics.
Bioethics Core – HN4516

(1) Provides bioethics consultation, education, and administrative infrastructure for human subjects research and (2) provides administrative infrastructure, training, and consultation for investigators related to the Institutional Review Board (IRB) process; (3) coordinates annual Responsible Conduct of Research educational sessions; (4) moderates bioethics rounds at clinical genetics case conferences; (5) participates in the Clinical Center’s ethics consultation service and Ethics Committee; (6) addresses emergent bioethics education and consultation needs; and (7) engages in a program of evaluation research to help ensure that recommendations are empirically well-grounded and responsive to the various stakeholders involved.
(1) Reviews medical records of applicants with undiagnosed disorders who seek a definitive diagnoses; (2) investigates mysterious human diseases by intensive and extensive phenotyping through week-long Clinical Center admissions, single nucleotide polymorphism array and exome sequencing analyses, and functional studies of candidate gene variants; (3) develops new methods for exome sequence analysis, such as the diploid aligner and variant filtering programs; (4) serves as an integral part of the newly established Undiagnosed Diseases Network and its Data Coordinating Center, providing experience and expertise on best practices for an undiagnosed diseases program; (5) develops algorithms for diagnostic evaluations, data submission to dbGaP, and other databases, and privacy protection for patients with undiagnosed diseases; (6) develops and conducts one-week training sessions and year-long fellowships in the analysis of Next Generation sequencing data of patients with rare and new diseases; and (7) defines and discovers new human disorders and disease mechanisms.
Cancer Genetics and Comparative Genomics Branch - HN452

(1) Defines the genetic changes involved in the susceptibility and progression of cancer. (2) Identifies both germline risk alleles and cumulative genetic changes arising throughout life that contribute to tumor formation; (3) pursues a range of experimental studies to identify the genes and variants that increase cancer susceptibility, as well as to develop an understanding of associated functional changes; (4) identifies mutations leading to susceptibility and progression of prostate, breast and endometrial tumors; (5) utilizes naturally occurring animal systems to localize cancer susceptibility genes in a background of reduced heterogeneity; and (6) develops and applies technologies, such as cDNA and SNP arrays, together with deep sequencing for genome-wide analysis aimed at identifying genetic variants, expression changes, or haplotypes that are unique to specific cancers, response to treatment, or stages of progression.
Reproductive Cancer Genetics Section - HN4522

Conducts research into the genetic basis of reproductive cancers including, but not limited to, endometrial cancer; and conducts ongoing investigations using high-throughput nucleotide sequencing, whole genome genotyping, and linkage and association studies to understand the genetic basis of endometrial cancer susceptibility, as well as disease progression and outcome.
Comparative Genetics Section - HN4523

(1) Localizes and studies genetic variants responsible for disease susceptibility and morphologic variation; (2) analyzes high-risk families by mapping or exome sequencing to find variants that increase susceptibility to prostate cancer, particularly to aggressive forms of the disease; (3) understands the role of low penetrant but frequent variants in cancer-associated genes, particularly those that increase risk or progression of disease; and (4) uses whole genome sequencing and genome wide studies of dog breeds to identify genomic variants important in disease susceptibility and morphological variation, with a focus on traits related to cancer, body size, and skull shape.
Cancer Genomics Unit - HN4527

Adopts genomic technologies to cancer susceptibility inherited disorders such as Fanconi anemia in order to understand genotype-phenotype correlations, to develop better diagnostics and to explore novel therapeutic approaches.
Comparative Genomics Analysis Unit - HN4528

Develops, evaluates, and applies genomic algorithms and software tools using high-performance computers, in an effort to understand and interpret the increasing amounts of sequence and genotyping data available both from public sources and from collaborations with other investigators.
Genomics Core - HN4529

Consults NHGRI investigators on their genotyping needs and provides guidance as to the type of service needed, generates the appropriate genotyping data using both the ABI sequencer for STRP and Illumina technologies for SNP based genotypes, and assists with the analysis of data. Provides services related to physical mapping, DNA sequencing, and access to DNA panels.
Microarray Core - HN452A

Provides intramural investigators with full service, cost-effective, and time-efficient access to comprehensive, state-of-the-art genomics, and transcriptomics technologies; and supports analysis on all commercial microarray platforms to offer a broad range of products and services, including whole genome gene expression, genotyping (SNP), epigenetics (DNA methylation), methylation, copy number variation (CNV, LOH and CGH), non-coding and microRNA analysis, proteomic microarray, custom microarray printing, hybridization, data mining and analysis, as well as consultation on experimental design.
Genetic Disease Research Branch - HN453

(1) Plans and conducts research using genomic and genetic approaches to identify and understand mechanisms regulating human genetic diseases; (2) uses functional genomic approaches to study genetic alterations involved in human disease states; (3) utilizes genetically modified model organisms and cellular models to explore signaling pathways affected in human health, development and disease; and (4) determines the functions of cloned genes involved in human genetic diseases.
Cell Signaling and Immunity Section - HN4532

(1) Plans and conducts research on signal transduction pathways focusing on signaling in the immune system and its contribution to normal cellular development and abnormal disease states; (2) uses a combination of genetic, genomic, biochemical, and cellular approaches including both transgenic and gene-targeting technology in mice to examine the function of tyrosine kinases and related signaling molecules involved in lymphocyte development, function and responses to infection and immunization; and (3) uses functional genomic, cellular and systems approaches to understand mechanisms of human immunity and immune-mediated diseases, including genetic causes of primary immunodeficiencies.
(1) Uses genomic tools and genetic manipulation of model systems to unravel genome function and to dissect gene regulatory pathways in development and disease; (2) integrates data from basic science studies with clinical information; (3) identifies pathways that regulate mammalian development; (4) understands how alterations in these pathways lead to disease states; and (5) develops paradigms for therapeutic interventions.
Embryonic Stem Cell and Transgenic Mouse Core - HN4537

(1) Provides a shared support service to generate genetically engineered mice (GEM); (2) generates GEM by utilizing embryonal stem cell (ESC) technology and conventional transgenic methods (direct microinjection); (3) consults and guides researchers in planning transgenic and embryonic stem cell projects; (4) provides support to researchers in breeding and analyzing GEM generated through transgenic and ESC technology; (5) rederives imported mice into the animal facility and cryopreserves all important mice for investigators.; (6) provides investigators with in-house breeding stock of tester strains and other useful mice that are used across various animal users' protocols.
Cytogenetic and Microscopy Core - HN4538

(1) Provides state-of-the-art molecular cytogenetic and microscopy services; (2) serves as a source for experimental planning, implementation of protocols, and conveyance of results; (3) implements new cytogenetic and microscopy techniques; (4) mentors trainees; and (5) maintains and utilizes state-of-the-art equipment and techniques.
Genetics and Molecular Biology Branch - HN454

Uses state-of-the-art genetic and genomic technologies to study genomes and disease mechanisms, to translate research findings into improved diagnosis, treatments, and prevention of human diseases; provides rigorous mentoring to the next generation of genetics and genomics investigators and involves all trainees in translational research projects.
Hematopoiesis Section - HN4544

(1) Studies hematopoiesis in mouse models and human patients; (2) identifies novel genes involved in the differentiation of red blood cells; (3) determines the regulatory signals for red cell gene expression at the genome level; and (4) performs testing of in vivo and in vitro methods to increase red cell production.
Flow Cytometry Core - HN454B

(1) Provides investigators with state-of-the-art flow cytometry services; (2) serves as a source for experimental design and planning, execution of experiments and data analysis; (3) collaborates with researchers to reach scientific goals; (4) teaches trainees in the art of flow cytometry; and (5) maintains and utilizes state-of-the-art flow cytometry equipment.
Medical Genetics Branch - HN456

(1) Conducts clinical and basic research into the genetic contribution to human growth and development in health and disease and the application of that knowledge to genetic diagnosis, counseling and therapy; (2) provides resources and oversight for the conduct of research; (3) investigates skeletal and connective tissue biology, disorders of human brain development and the process and effectiveness of genetic education and counseling; and (4) manages a multi-disciplinary staff including M.D.'s, Ph.D.'s, nurses, genetic counselors, and trainees dedicated to achieving these goals.
(1) Investigates metabolic disorders of man at the clinical, biochemical, molecular, and cell biological levels; (2) pursues clinical endeavors including patient care, controlled therapeutic trials, investigational drug use, employment of unique diagnostic tests, and description of new disorders or new presentations of known disorders; (3) determines the primary defect in rare genetic diseases; (4) studies the pathogenesis of specific clinical manifestations; (5) investigates the mechanisms of action of cellular processes whose aberrations result in metabolic disease states; (6) describes defective processes or systems which often provide the opportunity to elucidate normal function; (7) studies disorders of vesicle formation and trafficking; (8) investigates the metabolic diseases known collectively as Hermansky-Pudlak syndrome (HPS) which provide models for the genesis of three intracellular vesicles (the melanosome, lysosome, and platelet dense body); (9) describes various aspects of HPS in different ethnic groups; (10) elucidates the many molecular bases of HPS; (9) determines the primary cellular defect responsible for the distinctive phenotype of HPS; (10) studies each type of HPS to reveal critical aspects of how vesicles form or move inside cells; (11) investigates nephropathic cystinosis (a prototypic lysosomal transport disorder), sialuria (a defect in feedback inhibition of the rate-limiting step in sialic acid synthesis), and alkaptonuria (a progressive joint disorder due to deficiency of homogentisic acid oxidase, an enzyme in the tyrosine catabolic pathway); (12) studies unique and unusually instructive diseases to gain insight into normal and abnormal intracellular processes; and (13) designs therapeutic interventions to correct specific metabolic defects from knowledge acquired from these disease states.
Human Development Section - HN4565

(1) Conducts research using molecular genetics, biological and biochemical methods on a number of projects related to human genetic disorders; (2) investigates disorders of human brain development including holoprosencephaly, craniosynostosis syndromes and ADHD; (3) manages multiple collaborative research projects involving intramural and extramural investigators; (4) studies the natural clinical history and underlying biochemical and molecular basis of a variety of rare metabolic disorders of children; (5) conducts both clinical and basic research in known metabolic diseases of carbohydrate metabolism and other disorders that lead to developmental delay in children; (6) clinically evaluates children with developmental delay and a clinical history suggestive of rare disorders of metabolism; (7) provides training to post-doctoral level PhD fellows, graduate students working towards the Ph.D. degree, and M.D. fellows to develop basic research skills; (8) provides a bioethics curriculum which includes instruction about how to conduct genetic research with regards to ethics issues related to clinical genetics and genetics policy; (9) conducts interdisciplinary research and utilizes both conceptual approaches and empirical evaluations; (10) addresses issues related to the genetic study of isolated and/or minority populations; and (11) serves as a focal point for institutions and centers representing special populations.
Molecular Neurogenetics Section - HN4568

(1) Conducts clinical and basic research into the factors contributing to the phenotypic variation observed in monogenic diseases, using Gaucher disease as a prototype disorder; (2) investigates the relationship between Gaucher disease and parkinsonism; and (3) explores new therapeutic approaches for Gaucher disease.
Cell Biology of Metabolic Disorders Unit - HN456A

Conducts and coordinates molecular and cell biological studies on a variety of human metabolic disorders, with the purpose of understanding underlying pathomechanisms and evaluating plausible therapies for these disorders. Utilizes mouse, zebrafish and cell systems of human metabolic disorders. Special interests include disorders related to intracellular formation and trafficking of lysosome-related organelles, glycobiology and sialic acid metabolism.
(1) applies genetic discoveries to improve interventions for preventing disease and promoting health and well-being; (2) applies social, behavioral, and communication theories to understand the essential elements of communicating genetic risk effectively; (3) develops and refines theory-based methods for communicating genetic risk to affected individuals, families, communities and populations; (4) understands how social factors (e.g., interpersonal, organizational, cultural and societal) influence genetic discoveries and research; and (5) investigates the ethical and public policy implications of genetic research and the use of genetics in clinical practice.
Social Network Methods Section - HN4594

(1) Develops methods to measure and model the complexities of social systems; (2) utilizes these methods to understand the social, psychological, and communicative context of families at risk for hereditary disease; (3) translates this understanding into effective network-based interventions; (4) investigates differences in the diffusion or coping processes across families from different ethnic and racial backgrounds and across different genetic diseases; (5) investigates how formal support systems, such as health care providers (i.e. genetic counselors, general practitioners), family members who are not at-risk (e.g. spouses), as well as those who are "like family" (e.g. friends) participate in the process of decision making, communication and support for at-risk family members; (6) understands the familial culture from a network perspective that provides important information for the delivery of genetic counseling services and dissemination of genetic risk information; and (7) develops network-based interventions that facilitates sharing of risk information throughout the family system.
(1) Conducts research at the intersection of neuroscience, basic behavioral science and social science to advance understanding of the etiology, clinical course and treatment of attention deficit hyperactivity disorder (ADHD); (2) defines the genetic bases of the abnormal neurodevelopmental trajectories that characterize childhood ADHD; (3) characterizes subtypes of the disorder based on neurobehavioral profiles; (4) develops tools to predict long term outcomes; and (5) explores the genetic, epigenetic, behavioral and interpersonal factors that influence clinical course and response to pharmacological and behavioral treatments.
Immersive Virtual Environment Test Unit - HN4597

Applies innovative methods in order to conduct and support experimental research that anticipates and shapes the application of new genomic information for health care and public health.
Health Disparities Unit - HN4598

Conducts research that evaluates approaches to integrating new genomic knowledge into clinical settings without exacerbating inequities in healthcare delivery.
Gene and Environment Interaction Section - HN459A

(1) Carries out research aimed at understanding the interactions between genetic variation and the external environment; (2) investigates the function of gene variants in human populations as they relate to disease susceptibility; (3) uses computational, cell and animal models to understand the gene-environment interactions at the molecular level; (4) develops improved diagnostic approaches and leads to preventative therapies by understanding disease susceptibility genes and their interaction with other genes and the environment.
Social Epidemiology Research Unit - HN459B

Conducts research related to the effects of social determinants on cardiovascular disease morbidity and mortality as well as the effects of social determinants on gene expression among disproportionately affected sub-populations.
Genetic Counseling Training Program - HN45M7

Prepares students for a Master of Science degree (MSc) in Genetic Counseling from the Department of Health, Behavior and Society, Johns Hopkins Bloomberg School of Public Health; and develops, maintains, and oversees the genetic counseling coursework, student supervision, coordination of clinical rotations and American Board of Genetic Counselors accreditation.
Skeletal Genomics Unit - HN45M8

The goals of the unit are to use a translational research approach to understand the etiology, pathophysiology, diagnosis, and management of individuals with heritable disorders of skeletal biology. Specifically, to conduct clinical research to uncover the clinical spectrum and genomic basis of selected skeletal dysplasias, develop animal and cell models to understand the pathomechanism of these disorders, and develop targeted treatment approaches.
The Systems Biology and Genome Engineering Section’s goal is to systematically understand the effects of genetic differences. Researchers in this section study genetic diversity across an expansive range, such as natural variation between individuals or between species, or novel genetic changes that they create. By studying the functional effects of these variants, section researchers work to learn principles of genome function.

Section researchers use genome engineering to direct activities such as genome editing or recombination, using budding yeast as a model organism. As the space of potential genetic variants is immense, the section’s researchers study large numbers of genetic differences in parallel. They do this by using large-scale oligonucleotide synthesis to generate a diversity of instructions targeting desired modifications, in order to generate pools of cells containing thousands of unique genomic changes. In addition to existing genome engineering methods, the section is continually developing new approaches to further expand the genetic space that can be studied.
Intramural Training Office - HN45A

(1) Supports NHGRI DIR trainees’ professional needs through mentoring, training programs and outreach activities to enhance institutional experiences while enabling achievement of career goals; and (2) trains and retains a highly skilled and diverse genomic and genetics workforce, providing future leaders for the global community.
Office of Laboratory Animal Medicine - HN45C

(1) Provides veterinary medical support to research animal models; (2) trains NHGRI investigators in all aspects humane care and use of animals in biomedical and behavioral research, teaching, and testing; (3) provides research support to investigators using animal models; (4) aides in development/refinement of research biometodologies; (5) assists in Animal Study Proposal development; and (6) orders, receives, and ships research models.
Technology Transfer Office - HN45D

(1) Educates administrative and scientific personnel about patent, licensing, CRADA- and other technology transfer-related procedures and policies.; (2) facilitates the exchange of materials and information between NHGRI scientists and their collaborators and fosters productive interactions between intramural NHGRI labs and outside collaborators by effectively negotiating the legal, financial and scientific terms for these joint research projects; and 3) contributes to the successful commercialization of promising NHGRI research discoveries by facilitating interactions and providing advice and oversight to NHGRI scientists and potential collaborators and/or licensees.
NIH Intramural Sequencing Center - HN45E

(1) Performs large-scale DNA sequencing for NIH intramural investigators; (2) provides expert advice for setting up new genomic sequencing projects; and (3) serves as the focal point at NIH for genomic sequencing technologies, chemistries, and instrumentation as well as innovative sequence analysis software.
Computational and Statistical Genomics Branch - HN45H

(1) Develops and applies methods in statistics, bioinformatics, genetics, genomics, and computer science to identify genes and regulatory elements that are responsible for disease expression and trait variation in humans; (2) uses computationally intensive approaches to analyze large-scale genomic data from population-based and family-based studies of humans, as well as from studies of model organisms, to accomplish this mission; (3) employs comparative genomic approaches to understand the evolution and function of protein families and their ultimate role in human disease; (4) develops and disseminates new software tools and bioinformatic approaches; and (5) serves as a focal point for research and training in statistical genetics, genetic epidemiology, and bioinformatics.
Genometrics Section - HN45H2

(1) Uses methods in statistical genetics to identify genetic effects (genes and regulatory elements) underlying quantitative traits and determines the statistical challenges that must be addressed in identifying those effects; (2) develops new methods of statistical genetic analysis for quantitative traits and makes the developed software freely available; (3) investigates the statistical properties of these newly developed and other statistical methods with computer simulation; and (4) applies methods and the insights gained from simulation experiments to analyze data from ongoing collaborative studies.
(1) Performs research in both theoretical and applied statistical genetics as applied to human traits and disease; (2) performs data collection; power analysis, linkage, association and interaction analysis of human data; (3) analyzes large scale sequence and other genomic datasets in humans; (4) investigates the properties of various methods of statistical genetic analysis through computer simulation; and (5) emphasizes studies on complex diseases such as human cancers, eye diseases, and malformations.
Computational Genomics Unit - HN45H5

Uses phylogenetic and comparative genomic techniques to study developmental proteins that play a fundamental role in the specification of body plan, pattern formation, and cell fate determination during metazoan development; and employs a variety of computational approaches to understand the evolution and function of these proteins and their ultimate role in human disease.
(1) Supports the research being performed by NHGRI investigators by providing expertise and assistance in bioinformatics and computational analysis; (2) facilitates access to specialized software and hardware; (3) develops generalized software solutions that can address a variety of questions in genomic research; (4) develops database solutions for the efficient archiving and retrieval of experimental and clinical data; (5) disseminates new software and database solutions to the genome community at-large; (6) collaborates with NHGRI researchers on computationally intensive projects; and (7) provides educational opportunities in bioinformatics to investigators and trainees.
Genome Informatics Section- HN45H7

(1) Designs and applies efficient algorithms for the analysis of large-scale genomic data; (2) develops fundamental software tools for the assembly, comparison, and exploration of genomes and metagenomes; (3) produces high-quality reference genomes that enable the discovery of novel genomic structures and variation; (4) actively promotes open source and open data initiatives in genomics; (5) fosters interdisciplinary collaboration between the computational, biological, and medical sciences to advance global health.
(1) Plans and conducts research into the genomic and social determinants of complex diseases with particular attention to diseases (e.g., diabetes, hypertension, heart and kidney disease, scleroderma, and podoconiosis) that are disproportionately distributed by ethnicity and geography; (2) designs and analyzes genomic data to document and interpret the non-random distribution of human genetic variation with the goal of understanding human history and how these histories inform disease distribution and etiology; and (3) facilitates the development of large genomic initiatives in populations currently under-represented in the genomic revolution.
Center for Research on Genomics and Global Health - HN45L2

Advances research in the role of culture, lifestyle, genetics and genomics in disease etiology, differential susceptibilities to disease and variable drug response at the individual and population levels; and develops genetic epidemiology models that explore the patterns and determinants of common complex diseases in populations in the US and other human populations around the world.
Inflammatory Disease Section - HN45L3

Utilizes genetic and genomic strategies to elucidate the molecular basis, pathophysiology, and treatment of human inflammatory diseases. Major areas of concentration include (1) the genetics of Mendelian inflammatory diseases, including known disorders such as familial Mediterranean fever and currently unexplained phenotypes; (2) analysis of genetically complex inflammatory disorders, such as Behçet’s disease and scleroderma; (3) studies of disease pathogenesis, using both patient samples and animal models; (4) natural history and targeted therapeutic studies in patients; and (5) integrated studies of patients at the NIH Clinical Center with laboratory investigations.
(1) Specializes in multi-disciplinary translational research, in concordance with the traditional genetic-to-phenotype imputations; (2) utilizes a combination of noninvasive cardiovascular imaging, biomedical informatics, systems biology, and genomic epidemiology to conduct very large scale data analyses; and (3) pursues understanding of the relationships and interdependencies between clinical phenotypes, human behavior, biosurveillance, molecular, genomic and social determinants on gene expression and their contribution to health disparities in cardiovascular disease.
Medical Genomics and Metabolic Genetics Branch - HN45M

(1) Plans and conducts research investigating the relationship of genetic variation to health and disease; (2) uses genetic, genomic, and metabolic approaches to understand normal and abnormal physiology; (3) develops and tests innovative approaches to the diagnosis and management of disease; and (4) facilitates the development of early career physician-scientists.
Clinical Genomics Section - HN45M2

(1) Plans and conducts clinical and laboratory-based research to understand the relationship of genotype to phenotype; (2) develops and tests novel approaches to predictive and genomic medicine; (3) understands the molecular basis of birth defect and overgrowth disorders; and (4) develops treatment approaches for inherited and somatic genetic disorders.
Molecular Genetics Section - HN45M4

(1) Conducts research to uncover the genetic basis of monogenic and polygenic diseases; (2) develops better diagnostic and therapeutic approaches for these diseases; and (3) conducts research studies on progeria and type 2 diabetes mellitus, including the use of genomic technologies to identify disease-associated variations and elucidate their functional consequences.
Metabolism, Infection and Immunity Unit - HN45M5

Designs and conducts clinical and laboratory research in order to: (1) understand metabolic perturbations brought about by immune activation; and (2) demonstrate the role of intermediary metabolism in the differentiation, activation and function of immune cells.
Organic Acid Research Section- HN45M6

(1) Plans and conducts research to diagnose and study patients with organic acidemias; (2) develops animal models of organic acidemias; (3) develops new therapies for organic acidemias; and (4) develops new assays for diagnosing and for evaluating the effectiveness of therapies for organic acidemias.
Translational and Functional Genomics Branch - HN45N

(1) Plans and conducts research to explore the full functional potential of the human genome; (2) investigates the genetic, epigenetic and metagenomic basis of human disorders; (3) applies genetic and genomic approaches to model organisms to understand the molecular etiology of human disorders; (4) catalyzes technology development in genetics and computational genomics; and (5) translates laboratory findings to improved diagnoses and therapeutics for human disorders.
Microbial Genomics Section - HN45N2

(1) Performs foundational studies to characterize the diversity of microbes (bacteria, fungi, other small micro-organisms) that reside on human skin; (2) characterizes changes in the human microbiome associated with skin disorders and transition through life stages; (3) examines functional host-microbial (commensal and pathogenic) interactions; (4) develops and utilizes genomic sequence technology to identify emerging hospital bacterial pathogens; and (5) tracks transmission and evolution of hospital-associated bacterial pathogens incorporating epidemiologic and genomic information.
(1) Identifies and studies genetic, genomic, and epigenetic alterations involved in the pathogenesis of leukemias and other human blood diseases; (2) investigates genetic, genomic, and epigenetic involvements, including chromosome rearrangements, point mutations, and altered expressions, in the initiation and progression of these human diseases; and (3) uses animal and cell-based models to study these disease-related genes and their encoded proteins in diverse biological processes and disease pathogenesis, and to develop novel therapeutics including small chemical and biological treatments.
Developmental Genomics Section - HN45N4

(1) Utilizes a variety of modern molecular methods to determine the functions of genes and how they relate to human genetic disease; (2) emphasizes zebrafish as a model organism and systematic mutagenesis to identify the function of new genes important in human disease; (3) focuses on isolating embryonic mutations of candidate human disease genes and mutations that disrupt tissue regeneration; (4) develops large-scale, high-throughput gene inactivation technology and analyzes gene functions on a genome-wide scale; and (5) annotates large scale functional regulatory regions of the zebrafish genome.
Genomic Functional Analysis Section - HN45N5

(1) Plans and conducts research investigating functional entities in the human genome;
(2) characterizes the relationship of the aforementioned elements to those in non-human organisms with respect to sequence alignment and phylogeny; (3) aims to analyze the relationship between mutations in functional elements and correlations with human genetic diseases; (4) integrates epigenomic and genomic data with transcriptomic outcomes; and (5) provides genomics analysis tools to the community at large for the betterment of computational genomics as a field of study.
Zebrafish Core - HN45N6

(1) Provides resources, training and services to analyze gene function and generate disease models using zebrafish for research; (2) provides microinjection services to perform transient knockdown and over-expression of desired genes; (3) performs microinjections and founder screening service to generate transgenic zebrafish lines; (4) provides services to generate genetic mutants using ZFNs, TALENs and/or CRISPR-cas9 nucleases; (5) consults with researchers in planning zebrafish experiments; (6) provides training in breeding and imaging to analyze phenotypes of mutant, morphant and transgenic zebrafish; and (7) provides chemical libraries and training to perform drug screening using zebrafish.
(1) Plans, directs, and facilitates multi-disciplinary research to understand the structure and function of genomes and their implications for biology and disease etiology, through research and training grants, research center grants, and contracts; (2) assesses the need for research and research training in genome sciences and related areas; (3) determines program priorities in genome sciences and related areas and recommends funding levels; (4) prepares reports and analyses to assist Institute staff and advisory groups in carrying out their responsibilities; (5) collaborates with the other NHGRI extramural research Divisions to establish a balance of resources, personnel, research and training budgets to achieve NHGRI goals; (6) provides expert advice to the Director, NHGRI, on various aspects of genome sciences; and (7) collaborates with the other NHGRI Divisions, other NIH ICs, and other agencies and entities, nationally and internationally, and maintains an awareness of research efforts in relevant program areas.
Division of Genomic Medicine - HN48

(1) Plans, directs, and facilitates multi-disciplinary research to identify genetic contributions to human health and disease and use them to improve diagnosis, treatment, and prevention through research and training grants, research center grants, and contracts; (2) determines program priorities in genomic medicine and related areas and recommends funding levels; (3) assesses the need for research and research training in genomic medicine and related areas; (4) prepares reports and analyses to assist Institute staff and advisory groups in carrying out their responsibilities; (5) collaborates with the other NHGRI extramural research Divisions to establish a balance of resources, personnel, research and training budgets, to achieve goals; (6) provides expert advice to the Director, NHGRI, on various aspects of genomic medicine; and (7) collaborates with the other NHGRI Divisions, other NIH ICs, and other agencies and entities nationally and internationally, and maintains an awareness of research efforts in relevant program areas.
Division of Genomics and Society - HN49

(1) Plans, directs, and facilitates, through research and training grants, research center grants, and contracts, multi-disciplinary research to understand the societal issues raised by genomics and their implications for conducting research on biology and disease etiology, prevention, treatment, and management; (2) assesses the need for research and research training in societal issues raised by genomics and related areas; (3) determines program priorities in societal issues raised by genomics and related areas and recommends funding levels; (4) prepares reports and analyses to assist Institute staff and advisory groups in carrying out their responsibilities; (5) collaborates with the other NHGRI extramural research Divisions to establish a balance of resources, personnel, research and training budgets to achieve NHGRI goals; (6) provides expert advice to the Director, NHGRI, on various aspects of the societal issues raised by genomics; and (7) collaborates with the other NHGRI Divisions, other NIH ICs, and other agencies and entities, nationally and internationally, and maintains an awareness of research efforts in relevant program areas.
Division of Extramural Operations - HN4A

(1) Provides leadership and advice to the Director on implementing and coordinating extramural research grant, contract, and training program operations and policies; (2) provides a full range of activities that empower the scientific divisions in implementing and coordinating cross-cutting, multi-disciplinary activities in the mission areas of NHGRI; (3) represents NHGRI on overall NIH extramural and collaborative program policy committees and coordinates such policies within the institute; (4) manages the activities of the National Advisory Council for Human Genome Research; (5) develops and provides extramural staff training and enhances communication across the Institute regarding standardized approaches policies, methods, and procedures; (6) manages institute–wide extramural services and activities; (7) provides oversight and direction for the initial scientific merit review of research, resource, and training applications and proposals assigned to the Institute; (8) provides oversight and direction for grants management activities, ensuring that all awards are made in accordance with applicable statutes, regulations, and policies; (9) provides data, reports and analyses to assist NHGRI staff and advisory groups in carrying out their responsibilities; and (10) maintains effective relationships with other NIH institutes and divisions, the research community, other federal agencies, and professional societies.
Scientific Review Branch - HN4A2

(1) Plans and administers scientific review activities, including the organization and management of Initial Review Groups (IRGs), constituted to review NHGRI's extramural grants, cooperative agreements, and contract proposals; (2) performs initial scientific and administrative review of research, research center, program project, conference, and research training grant applications, cooperative agreements, and contracts; (3) plans and directs site-visits and IRG meetings; (4) establishes review criteria and standards for the IRGs and provides technical leadership to the review process; and (5) prepares summary statements and provides pertinent information concerning the review of applications to the Council.
Grants Administration Branch - HN4A3

(1) Develops and implements Institute policies on the business management aspects of grant and fellowship programs; (2) develops and implements guidelines, procedures and internal controls to ensure proper and continuing implementation of NHGRI, NIH, PHS, and DHHS policies; (3) administers the Institute's grant programs including negotiation of awards, post award approvals, audits, and closeout; (4) participates in awarding functions, authorizing awards and changes to active awards; and (5) prepares and distributes grant-related materials required by the NHGRI Advisory Council.
Division of Management - HN4B

(1) Plans and directs administrative management functions including administrative management services, management analysis and evaluation, financial management, information technology and human resources; (2) advises the senior leadership on developments in administrative management and their implications and effects on program management; and (3) coordinates administrative management activities in support of their programs.
Administrative Services Branch - HN4B2

1) Negotiates for and secures the resources and services needed to run the Office of the Director and the Division of Extramural Research; (2) manages the Institute's RMS Budget, monitoring and tracking obligations and expenditures to assure the availability of funds and prevent budget deficiencies; (3) manages the personnel, procurement, property, space, travel, and other administrative functions as necessary to ensure the efficient and effective implementation and operation of programs; (4) advises NHGRI staff of administrative policies and procedures; (5) performs analytical studies related to the administrative organization, processes and procedures of the Institute and establishes effective administrative controls; (6) represents the Institute on NIH committees or task groups; (7) and serves as liaison and contact point for the NHGRI with the central services areas of NIH, other agencies, and outside businesses.
Financial Management Branch - HN4B3

(1) Plans, directs, coordinates the financial management activities of the Institute; (2) serves as the principal advisor to the Institute Director, Deputy Director, and Associate Director for Management on the management of the financial and personnel resources of the Institute; (3) manages the Institute budget process which includes providing guidance to Institute staff on budget preparation and management; coordinating the preparation and submission of the Institute's budget to NIH, OMB, and the President; explaining and defending the budget; and monitoring budget execution; (4) administers the Institute's budgetary, accounting, and funds control systems assuring that they are compatible with NIH and Departmental accounting systems and provide Institute management and staff with reliable accounting, financial management, and personnel resource utilization data; (5) performs analysis and provides information on the historic, current, and prospective financial status of the Institute for briefing and reporting purposes; and (6) participates in budget hearings by preparing background, briefing, and source material for the Institute Director and others appearing before committees or examiners, by attending budget hearings as part of the Institute contingent on an as-needed basis, and by coordinating responses to questions concerning the financial management of the Institute from the Congress and others.
Information Technology Branch - HN4B4

(1) Provides technical leadership and advice to all levels of Institute management in order to obtain maximum utilization of current ADP resources and advancements in the field of information systems technology and communications; (2) determines requirements, designs, and implements and coordinates the Institute's management information systems which collect, maintain, and report various types of administrative information; (3) advises the NHGRI Director, Deputy Director, Associate Director for Management, and other Institute staff on the technological and policy impact and implications of developments in information systems and related fields within and outside the government; (4) coordinates staff activities with those of contractors, other components of NIH, and other Federal and non-Federal data processing agencies; (5) provides user support, including training, in LAN/information systems capabilities, programs, and procedures.
Intramural Administrative Management Branch - HN4B5

Plans, directs, coordinates, and provides comprehensive administrative and management support services for the Division of Intramural Research; (2) provides technical and advisory services in financial management, human resources, procurement, facility management, travel services, property management and other administrative functions, as necessary, to ensure the efficient and effective implementation and operation of programs; and (3) develops policies, guidelines, and procedures on matters relating to administrative management and disseminates to relevant staff.
Acquisition and Logistics Section - HN4B52

(1) Plans, directs and coordinates acquisition activities for the intramural staff using efficient, timely and appropriate methods; and (2) plans, directs, and coordinates logistical support for the DIR including management of DIR property activities.
Management Analysis and Workforce Development Branch - HN4B6

(1) Coordinates NHGRI-wide administrative/management initiatives including organizational analysis, workflow management, delegations of authority, policy and procedure analysis, and privacy and records management; (2) develops, executes and maintains an NHGRI-wide risk management program; (3) coordinates/directs cross-functional teams and study groups in finance, IT, and general administration; and (4) coordinates workforce management and professional development activities.
(1) Administers a comprehensive NHGRI ethics program that reflects statutory responsibilities and integrity in service to the public; (2) develops and recommends policies and procedures related to employee standards of conduct, financial interests and disclosure, outside activities, gifts administration, official duty activities, sponsored travel, and procurement integrity; (3) administers the annual public and confidential financial disclosure process including reviewing and certifying financial disclosure reports and reports of holdings in substantially affected organizations, and develops new employee ethics agreements; (4) reviews and approves requests for outside activities, official duty requests, and sponsored travel or related gifts for conformance with regulations and policies; (5) provides advice and assistance to employees regarding the application of the ethics laws, regulations, and policies; (6) develops and provides NHGRI ethics training (7) provides liaison to the HHS Office of the General Counsel, the Office of Government Ethics, the NIH Ethics Office, other agencies, and outside organizations as needed; (8) provides advice to the Office of the Director regarding conflict of interest of individuals involved in the conduct of biomedical research, including Government employees, advisory committee members, and non-Government employees such as peer reviewers, Data Safety Monitoring Board (DSMB) members, and members of working groups; (9) reviews: (a) procurements over one million dollars involving justification of other than full and open competition; (b) gifts acceptance under NHGRI statutory authorities; (c) memoranda of understanding of public private partnership proposals and co-sponsorships with non-federal entities; and (d) conflicts concerning prior employment ties to academic institutions and private entities; (10) identifies management issues requiring action by the Office of General Counsel such as copyright, intellectual property, contract, or personnel authorities; and (11) reviews clinical protocols, conducts conflict of interest (COI) analysis to confirm no COI exists between investigators' official duties on the protocol and their personal or imputed financial interests.
Division of Policy, Communication and Education - HN4C

(1) Advises on a comprehensive range of issues affecting the NHGRI research portfolio and programs, as well as the broader genomics and biomedical research enterprise; (2) provides internal and external leadership and coordination for policy and program activities as they relate to the mission, representing the Institute and serving as a resource to internal and external stakeholders; (3) collaborates with leadership and scientific program staff to evaluate on-going programs in order to plan for future research and policy needs and provide analysis to the Director for use in the management of Institute resources, including all Institute tracking and reporting functions and programmatic activities related to budget development; (4) provides primary staff support to the Director for presentations and meetings discussing Institute programs and relevant policy matters; (5) develops and disseminates education and public engagement activities using a robust array of outreach and communication strategies and platforms, including the production of general and targeted information to promote understanding of genomics research and its applications to health and society; and (6) acts as a focal point for the synthesis of trans-Institute expertise on research, education, and policy topics relevant to the implications of genomics research and technology developments, including sponsoring consensus conferences and public workshops to consider research or policy directions, or to engage health professionals and the general public in discussions addressing genomics research and its applications.
Communications and Public Liaison Branch - HN4C2

(1) Advises the Institute Director and senior staff on a broad range of issues regarding the dissemination of information about the agency's programs; (2) distributes information about goals, programs, and advances in genomics, NHGRI's human genetics and ELSI research; (3) helps prepare reports, pamphlets, articles, news releases and other written, audiovisual or web-based materials about NHGRI research accomplishments, program, policies and goals; (4) communicates this information to audiences including the lay public, news media, Congress, other federal agencies, universities, and professional and consumer organizations; (5) coordinates the institute's public and private web sites; and (6) assists the Institute Director and other senior staff in developing and delivering talks to public and professional audiences.
Policy and Program Analysis Branch - HN4C3

(1) Advises the Institute Director and senior staff on a broad range of policy and program matters; (2) develops, implements, and reports analyses of Institute programs; (3) analyzes and tracks legislation relevant to the mission of the Institute and makes recommendations for legislative proposals; and (4) conducts and coordinates policy analysis related to the ethical, legal and social implications of human genomic and genetic research and develops programs to address these issues.
Education and Community Involvement Branch - HN4C4

(1) Advises the Institute Director and senior staff on a broad range of issues regarding public education and community involvement matters; (2) develops, implements, and evaluates public education programs; (3) conducts and coordinates public education initiatives; (4) develops community involvement programs to engage a broad range of the public in understanding genomics and accompanying ELSI issues; (5) conducts, coordinates, and evaluates community involvement initiatives; and (6) acts as the liaison between the Institute and community-based organizations.
Genomic Healthcare Branch - HN4C5

(1) Advises the Institute Director and senior staff on a wide range of topics related to genomics in health care; (2) develops, implements, and evaluates educational tools as well as other types of tools that enhance the use of genomic technologies in healthcare; (3) facilitates intra- and extra-governmental partnerships that promote the effective use of genomic technologies in healthcare; (4) develops programs that promote healthcare provider and healthcare organization involvement at all levels of the process of translating genomic discoveries to effective health interventions; and (5) acts as a liaison between the Institute and healthcare organizations.