To secure the promise of personalized medicine for all Americans by expanding and accelerating genomics research and initiatives to improve the accuracy of disease diagnosis, increase the safety of drugs, and identify novel treatments.

IN THE SENATE OF THE UNITED STATES
MARCH 23, 2007
Mr. Obama (for himself and Mr. Burr) introduced the following bill; which was read twice and referred to the Committee on Health, Education, Labor, and Pensions

A BILL
To secure the promise of personalized medicine for all Americans by expanding and accelerating genomics research and initiatives to improve the accuracy of disease diagnosis, increase the safety of drugs, and identify novel treatments.

1 Be it enacted by the Senate and House of Representa-
2 tives of the United States of America in Congress assembled,

3 SECTION 1. SHORT TITLE.
4 This Act may be cited as the “Genomics and Person-
5 alized Medicine Act of 2007”.

6 SEC. 2. FINDINGS.
7 Congress makes the following findings:
(1) The completion of the Human Genome Project in 2003 paved the way for a more sophisticated understanding of diseases and drug responses, which has contributed to the advent of “personalized medicine”.

(2) Personalized medicine is the application of genomic and molecular data to better target the delivery of health care, facilitate the discovery and clinical testing of new products, and help determine a person’s predisposition to a particular disease or condition.

(3) Many commonly-used drugs are typically effective in only 40 to 60 percent of the patient population.

(4) In the United States, up to 15 percent of hospitalized patients experience a serious adverse drug reaction, and more than 100,000 deaths are attributed annually to such reactions.

(5) Pharmacogenomics has the potential to dramatically increase the efficacy and safety of drugs and reduce health care costs, and is fundamental to the practice of genome-based personalized medicine.

(6) Pharmacogenomics is the study of how genetic variation affects a person’s response to drugs. This relatively new field combines pharmacology (the
science of drugs) and genomics (the study of genes and their functions) to develop safer and more effective medications and dosing regimens that will be tailored to an individual’s genetic makeup.

(7) The cancer drug Gleevec was developed based on knowledge of the chromosomal translocation that causes chronic myelogenous leukemia, which is characterized by an abnormal growth in the number of white blood cells. The mean 5-year survival for affected patients who are treated with Gleevec is 95 percent, which contrasts to a 5-year survival of 50 percent for patients treated with older therapies.

(8) The ERBB2 gene helps cells grow, divide and repair themselves. One in 4 breast cancers are characterized by extra copies of this gene, which causes uncontrolled and rapid tumor growth. Pharmacogenomics research led to both the development of the test for this type of breast cancer as well as an effective biologic, Herceptin.

(9) Warfarin, a blood thinner used to prevent the formation of life-threatening clots, significantly elevates patient risk for bleeding in the head or gastrointestinal tract, both of which are associated with increased rates of hospitalization, disability and
death. Pharmacogenomic researchers have identified and developed tests for genetic variants in the cytochrome P450 metabolizing enzyme (CYP2C9) and vitamin K epoxide reductase complex that increase risk for these adverse events. By using a companion diagnostic test for these two genes, physicians can modify the dosing regimen and decrease the likelihood of adverse events.

(10) Although the cancer drug 6-mercaptopurine (6-MP) cures 85 percent of children with acute lymphoblastic leukemia, historically, a significant number of patients would die inexplicably from the drug. Researchers later discovered that 1 in 300 individuals inherit an inactive version of the gene encoding the metabolizing enzyme thiopurine methyltransferase (TPMT) from both their mother and father and, as a result, should receive only a fraction of the standard dose of purine drugs. In addition, 1 in 10 individuals have only 1 copy of the gene with variable function, and the dosage of 6-MP must be adjusted for a subset of these patients. Physicians now are able to screen for TPMT gene variants before administering these drugs.

(11) Research into the genetics of breast cancer identified two pivotal genes, BRCA1 and BRCA2,
mutations in which correspond to a significantly increased lifetime risk of developing breast and ovarian cancer. Individuals in affected families or with specific risk factors may use genetic testing to identify whether they carry mutations in these genes and to inform their decisions about treatment options, including prophylactic mastectomy and oophorectomy.

(12) Realizing the promise of personalized medicine will require continued Federal leadership and agency collaboration, expansion and acceleration of genomics research, a capable genomics workforce, incentives to encourage development and collection of data on the analytic and clinical validity and clinical utility of genomic tests and therapies, and improved regulation over the quality of genetic tests, direct-to-consumer advertising of genetic tests, and use of personal genomic information.

SEC. 3. DEFINITIONS.

In this Act:

(1) BIOBANK.—The term “biobank” means a shared repository of human biological specimens that may also include data associated with such specimens collected for medical or research purposes. Human biological specimens may include body fluids, tissues, blood, cells, or materials derived from
these sources, and data associated with such specimens may include health information or environmental data.

(2) **BIOMARKER.**—The term “biomarker” means an analyte found in or derived from a patient specimen that is objectively measured and evaluated as an indicator of normal biologic processes, pathogenic processes, or pharmacologic responses to a therapeutic intervention.

(3) **CLIA.**—The term “CLIA” means the Clinical Laboratory Improvement Amendments of 1988 (42 U.S.C. 263a).

(4) **ENVIRONMENT.**—The term “environment” means conditions or circumstances that are non-genetic but may have a health impact.

(5) **GENETIC TEST.**—The term “genetic test” means an analysis of human DNA, RNA, chromosomes, proteins, or metabolites, that detects genotypes, mutations, or chromosomal and biochemical changes.

(6) **LABORATORY-DEVELOPED GENETIC TEST.**—The term “laboratory-developed genetic test” means a genetic test that is designed, validated, conducted, and offered as a service by a clinical laboratory subject to CLIA using either com-
mercially available analyte specific reagents (FDA-regulated) or reagents prepared by the laboratory (not FDA-regulated), or some combination thereof.

(7) PHARMACOGENETIC TEST.—The term “pharmacogenetic test” means a genetic test intended to identify individual variations in DNA sequence related to drug absorption and disposition (pharmacokinetics) or drug action (pharmacodynamics), including polymorphic variation in the genes that encode the functions of transporters, receptors, metabolizing enzymes, and other proteins.

(8) PHARMACOGENOMIC TEST.—

(A) IN GENERAL.—The term “pharmacogenomic test” means a genetic test intended to identify individual variations in single-nucleotide polymorphisms, haplotype markers, or alterations in gene expression or inactivation, that may be correlated with pharmacological function and therapeutic response.

(B) VARIATIONS AND ALTERATIONS.—For purposes of this paragraph, the variations or alterations referred to in subparagraph (A) may be a pattern or profile of change, rather than a change in an individual marker.
(9) Secretary.—The term “Secretary” means the Secretary of Health and Human Services.

SEC. 4. GENOMICS AND PERSONALIZED MEDICINE INTER-AGENCY WORKING GROUP.

(a) In General.—Not later than 90 days after the date of enactment of this Act, the Secretary shall establish within the Department of Health and Human Services the Genomics and Personalized Medicine Interagency Working Group (referred to in this Act as the “IWG”).

(b) Duties.—The IWG shall facilitate collaboration, coordination, and integration of activities within the Department of Health and Human Services and other Federal agencies, and among such agencies and relevant public and private entities, by—

(1) reviewing current and proposed genomic initiatives, in order to identify shared interests and leverage resources;

(2) prioritizing new genomic initiatives, based on areas of need as measured by public health impact;

(3) reaching consensus on standardized genomic terminology, definitions, and data code sets for adoption and use in Federally conducted or supported programs;
(4) establishing and disseminating quality standards and guidelines for the collection, processing, archiving, storage, and dissemination of genomic samples and data for research and clinical purposes;

(5) developing and promulgating guidelines regarding procedures, protocols, and policies for the safeguarding of the privacy of biobank subjects, in accordance with the Office for Human Research Protection and Clinical Research Policy Analysis and Coordination Program at the National Institutes of Health, and other guidelines as appropriate;

(6) reviewing and making recommendations to address ownership and patient access issues with respect to genomic samples and analyses;

(7) developing and promulgating guidelines regarding procedures, protocols, and policies for access to patient data, genomic samples, and associated health information by non-governmental entities for research purposes;

(8) developing and disseminating guidelines for constructing informed consent forms that ensure patient privacy and confidentiality of associated clinical data and information, understanding of research
procedures, benefits, risks, rights, and responsibilities, and continuous voluntary participation; and

(9) providing recommendations for the establishment of a distributed database, pursuant to section 5.

(c) IWG CHAIRPERSON.—The Secretary, or his or her designee, shall serve as chairperson of the IWG.

(d) MEMBERS.—In addition to the Secretary, the IWG shall include members from the—

(1) National Institutes of Health;

(2) Centers for Disease Control and Prevention;

(3) Food and Drug Administration;

(4) Health Resources and Services Administration;

(5) Office of Minority Health;

(6) Agency for Healthcare Research and Quality;

(7) Centers for Medicare & Medicaid Services;

(8) Veterans Health Administration;

(9) Office of the National Coordinator for Health Information Technology;

(10) Department of Energy;

(11) Armed Forces Institute of Pathology;

(12) Indian Health Service; and
(13) other Federal departments and agencies as determined appropriate by the Secretary.

(c) Public Input.—The IWG shall solicit input from relevant stakeholders with respect to meeting the duties described in subsection (b).

(f) Report.—Not later than 18 months after the date of enactment of this Act, the Secretary shall prepare and submit a report to the appropriate committees of Congress and to the public on IWG deliberations, activities, and recommendations with respect to meeting the duties described in subsection (b).

(g) Termination.—The IWG shall terminate after submitting the report described in subsection (f), or later at the discretion of the Secretary.

(h) Authorization of Appropriations.—There are authorized to be appropriated to carry out this section, $1,000,000 for fiscal years 2008 and 2009.

SEC. 5. NATIONAL BIOBANKING INITIATIVE.

(a) In General.—The Secretary shall advance the field of genomics and personalized medicine through establishment of a national biobanking distributed database for the collection and integration of genomic data, and associated environmental and clinical health information, which shall facilitate synthesis and pooled analysis of such data.
(b) DATABASE.—With respect to the national biobanking distributed database, the Secretary shall—

(1) adhere to relevant guidelines, policies, and recommendations of the IWG, pursuant to section 4;

(2) establish, directly or by contract, a single point of authority to manage operations of the database;

(3) incorporate biobanking data from Federally conducted or supported genomics initiatives, as feasible;

(4) encourage voluntary submission of biobanking data obtained or analyzed with private or non-Federal funds;

(5) facilitate submission of data, including secure and efficient electronic submission;

(6) allow public use of data only—

(A) with appropriate privacy safeguards in place; and

(B) for health research purposes;

(7) determine appropriate procedures for access by nongovernmental entities to biobank data for research and development of new or improved tests and treatments, and submission of data generated from such samples to the Food and Drug Adminis-
tration as part of the approval process for drugs and devices;

(8) conduct, directly or by contract, analytical research, including clinical, epidemiological, and social research, using biobank data; and

(9) make analytic findings from biobanking initiatives supported by Federal funding publicly available within an appropriate timeframe to be determined by the Secretary.

(c) RULE OF CONSTRUCTION.—Nothing in this section shall be construed to require the submission or acceptance of biological specimens.

(d) BIOBANK INITIATIVES GRANTS.—

(1) IN GENERAL.—The Secretary shall establish a grant program for eligible entities to develop or expand biobanking initiatives to increase understanding of how genomics interacts with environmental factors to cause disease, and to accelerate the development of genomic-based tests and treatments.

(2) ELIGIBLE ENTITIES.—

(A) IN GENERAL.—For purposes of this subsection, eligible entities include academic medical centers and other entities determined appropriate by the Secretary. Eligible entities
desiring a grant under this subsection shall submit an application to the Secretary in accordance with this subsection, at such time, in such manner, and containing such additional information as the Secretary may require.

(B) PRIORITY.—Academic medical centers that partner with health care professionals within their communities in order to obtain diverse genomic samples shall be given priority for awards made under this subsection.

(3) REQUIREMENTS.—The Secretary shall ensure that biobanks supported by grant awards under this section—

(A) adhere to guidelines and recommendations developed pursuant to section 4;

(B) are established to complement activities related to the implementation of current Federal biobanking research initiatives, as feasible;

(C) are based on well-defined populations, including population-based registries of disease and family-based registries;

(D) collect data from participants with diverse genomic profiles, demographics, environ-
mental exposures, and presence or absence of health conditions and diseases, as appropriate;

(E) meet quality standards for the collection, processing, archiving, storage, and dissemination of data, which shall be developed by the IWG;

(F) have practical experience and demonstrated expertise in genomics and its clinical and public health applications;

(G) establish mechanisms to ensure patient privacy and protection of information from non-health applications and, as feasible, patient access to genomic samples for clinical testing purposes; and

(H) contribute genomic and associated clinical and environmental data and analyses to the national biobanking distributed database, pursuant to subsection (b).

(4) USE OF FUNDS.—An eligible entity that receives a grant under this subsection shall use the grant funds to develop or expand a biobanking initiative, which may include the following activities:

(A) Support for scientific and community advisory committees.
(B) Recruitment and education of participants.

(C) Development of consent protocols.

(D) Obtaining genetic samples and associated environmental and clinical information.

(E) Establishment and maintenance of secure storage for genetic samples and clinical information.

(F) Conduct of data analyses and evidence-based systemic reviews that allow for the following:

   (i) Identification of biomarkers and other surrogate markers to improve predictions of onset of disease, response to therapy, and clinical outcomes.

   (ii) Increased understanding of gene-environment interactions.

   (iii) Development of genetic screening, diagnostic, and therapeutic interventions.

   (iv) Genotypic characterization of tissue samples.

   (G) Other activities, as determined appropriate by the Secretary.

(5) QUALITY ASSURANCE.—The Secretary may enter into a contract with an external entity to
evaluate grantees under this subsection to ensure that quality standards are met.

(c) APPLICATION OF PRIVACY RULES.—Nothing in this Act shall be construed to supersede the requirements for the protection of patient privacy under—

(1) the Federal regulations promulgated under section 264(c) of the Health Insurance Portability and Accountability Act of 1996 (42 U.S.C. 1320d–2 note); or

(2) sections 552 and 552a of title 5, United States Code (5 U.S.C. App.).

(f) AUTHORIZATION OF APPROPRIATIONS.—There are authorized to be appropriated to carry out this section, $75,000,000 for fiscal year 2009, and such sums as may be necessary for each of fiscal years 2010 through 2014.

SEC. 6. GENOMICS WORKFORCE AND TRAINING.

(a) GENETICS AND GENOMICS TRAINING.—The Secretary, directly or through contracts or grants to eligible entities, which shall include professional genetics and genomics societies, academic institutions, and other entities as determined appropriate by the Secretary, shall improve the adequacy of genetics and genomics training for diagnosis, treatment, and counseling of adults and children for both rare and common disorders, through support of efforts to—
(1) develop and disseminate model training pro-
gram and residency curricula and teaching materials
that reflect the new knowledge and evolving practice
of genetics and genomics;

(2) assist the review of board and other certi-
ifying examinations by professional societies and ac-
creditation bodies to ensure adequate focus on the
fundamental principles of genomics; and

(3) identify and evaluate options for distance or
on-line learning for degree or continuing education
programs.

(b) INTEGRATION.—The Secretary, in collaboration
with medical professional societies and accreditation bod-
ies and associations of health professional schools, shall
support initiatives to increase the integration of genetics
and genomics into all aspects of clinical and public health
practice by promoting genetics and genomics competency
across all clinical, public health, and laboratory disciplines
through the development and dissemination of health pro-
fessional guidelines which shall—

(1) include focus on appropriate techniques for
collection and storage of genomics samples, adminis-
tration and interpretation of genetic and genomic
tests, and subsequent clinical and public health deci-
sionmaking; and
(2) specifically target health professionals without formal training or experience in the field of genomics.

(c) Authorization of Appropriations.—There are authorized to be appropriated to carry out this section $5,000,000 for fiscal year 2008 and such sums as may be necessary for each of fiscal years 2009 through 2013.

SEC. 7. REALIZING THE POTENTIAL OF PERSONALIZED MEDICINE.

(a) National Academy of Sciences Study.—Not later than 180 days after the date of enactment of this Act, the Secretary shall enter into a contract with the National Research Council of the National Academy of Sciences to study and recommend appropriate incentives to encourage—

(1) codevelopment of companion diagnostic testing by a drug sponsor;

(2) development of companion diagnostic testing for already-approved drugs by the drug sponsor;

(3) companion diagnostic test development by device companies that are not affiliated with the drug sponsor; and

(4) action on other issues determined appropriate by the Secretary.

(b) Genetic Test Quality.—
(1) IN GENERAL.—The Secretary shall improve the availability of information on, and safety and efficacy of, genetic tests, including pharmacogenetic and pharmacogenomic tests.

(2) INSTITUTE OF MEDICINE STUDY.—Not later than 30 days after the date of enactment of this Act, the Secretary shall enter into a contract with the Institute of Medicine to conduct a study and prepare a report that includes recommendations to improve Federal oversight and regulation of genetic tests, with specific recommendations on the implementation of the decision matrix under paragraph (3). Such study shall take into consideration relevant reports by the Secretary’s Advisory Committee on Genetic Testing and other groups and shall be completed not later than 1 year after the date on which the Secretary entered into such contract.

(3) DECISION MATRIX.—

(A) IN GENERAL.—Not later than 18 months after the date of enactment of this Act, the Secretary, taking into consideration the recommendations of the Institute of Medicine report under paragraph (2), shall implement a decision matrix (referred to in this section as the “matrix”) to improve the oversight and regula-
tion of genetic tests, including pharmacogenomic and pharmacogenetic tests by determining—

(i) the classification of all genetic tests;

(ii) which categories of tests, including laboratory-developed tests, require review and the level of review needed for such categories of tests;

(iii) which agency shall have oversight over the review process of such categories of tests that are determined to require review; and

(iv) to the extent practicable, which requirements the agency shall apply to the types of tests identified in clause (ii).

(B) LEVEL OF REVIEW.—In determining the level of review needed by a genetic test, the Secretary shall take into consideration—

(i) performance characteristics of the test and its target disease or condition;

(ii) intended use of the test;

(iii) potential for improved medical conditions and patient harms; and

(iv) social consequences of the test.
(C) **COMPARATIVE ANALYSIS.**—To inform implementation of the matrix, the Secretary shall undertake a comparative analysis of laboratory review requirements under CLIA and those of the Food and Drug Administration to—

(i) assess and reduce unnecessary differences in such requirements;

(ii) eliminate redundancies and decrease burden of review, as practicable; and

(iii) specify which elements of the test constitute a device that may be regulated by the Food and Drug Administration and which elements comprise a service that may be regulated under CLIA.

(D) **REGULATIONS.**—The Secretary shall promulgate regulations to implement the matrix not later than the date specified under subparagraph (A).

(E) **TRANSITION PERIOD.**—The Secretary may not require a laboratory to submit a report under section 510(k) or an application under section 515 of the Federal Food, Drug and Cosmetic Act (21 U.S.C. 301 et seq.) until 180
days after the regulations promulgated under
subparagraph (D) take effect.

(4) ADVERSE EVENTS.—The Secretary, acting
through the Commissioner of Food and Drugs and
the Administrator of the Centers for Medicare &
Medicaid Services, shall—

(A) develop or expand adverse event re-
porting systems to encompass reports of ad-
verse events resulting from genetic testing;

(B) respond appropriately to any adverse
events resulting from such testing; and

(C) facilitate the use of genetic and
genomic approaches, as feasible, to assess risk
for, and reduce incidence of, adverse drug reac-
tions.

(5) AUTHORIZATION OF APPROPRIATIONS.—
There are authorized to be appropriated to carry out
this subsection, $6,000,000 for fiscal year 2008, and
such sums as may be necessary for each of fiscal
years 2009 through 2013.

(c) FOOD AND DRUG ADMINISTRATION.—

(1) IN GENERAL.—

(A) SUMMARY INFORMATION.—If a genetic
test that is determined to be within the jurisdic-
tion of the Food and Drug Administration but
that does not require review as determined
under the matrix, the sponsor of such test shall
provide the Secretary with summary informa-
tion on how such test was validated and its per-
formance characteristics. Such information shall
be in a standardized format and with standard-
ized content as specified by the Food and Drug
Administration, and shall be made easily acces-
sible to the public.

(B) SOURCE OF INFORMATION.—The in-
formation described under subparagraph (A)
may be obtained from the labeling submitted
for CLIA complexity categorization.

(2) ENCOURAGEMENT OF COMPANION DIAG-
NOSTIC TESTING.—The Secretary may encourage
the sponsor of a drug or biological product—

(A) to codevelop a companion diagnostic
test, after filing an investigational new drug ap-
plication or a new drug application to address
significant safety concerns of the drug or bio-
logical product;

(B) to develop a companion diagnostic test
if phase IV data demonstrate significant safety
or effectiveness concerns with use of the drug
or biological product; and
(C) to relabel the drug or biological product to require validated companion diagnostic testing when evidence of improved outcomes has been established in practice or if data demonstrate significant safety concerns with use of such drug or biological product.

(3) Pharmacogenomic Data Submission.—

The Secretary shall encourage and facilitate voluntary pharmacogenomic data submission from drug sponsors, which may include—

(A) the development and dissemination of guidance on relevant policies, procedure and practice regarding such submission;

(B) the provision of technical assistance;

(C) the establishment of a mechanism to store, maintain and analyze such data, in collaboration with the National Institutes of Health and the Centers for Disease Control and Prevention;

(D) determining when such data may be used to support an investigational new drug or a new drug application;

(E) the conduct of a study of the use of genomic approaches to understand and reduce adverse drug reactions; and
(F) other activities determined appropriate by the Commissioner.

(4) **TERMINATION OF CERTAIN ADVERTISING CAMPAIGNS.**—The Food and Drug Administration shall collaborate with the Federal Trade Commission to identify and terminate, pursuant to section 5 of the Federal Trade Commission Act (15 U.S.C. 45), advertising campaigns that make false, misleading, deceptive, or unfair claims about the benefits or risks of genetic tests.

(d) **CENTERS FOR MEDICARE & MEDICAID SERVICES.**—

(1) **IN GENERAL.**—If a genetic test that is determined to be within the jurisdiction of the Centers for Medicare & Medicaid Services but that does not require review as determined under the matrix, the sponsor of such test shall provide the Administrator of the Centers for Medicare & Medicaid Services with summary information on how the test was validated and its performance characteristics. Such information shall be in a standardized format and with standardized content as specified by the Centers for Medicare & Medicaid Services, and shall be made easily accessible to the public.
(2) **SPECIALTY AREA.**—To ensure the accuracy, validity, and reliability of clinical genetic tests that do not require premarket approval by or notification to the Food and Drug Administration, and to improve oversight of genetic test laboratories, the Director of the Division of Laboratory Services of the Survey and Certification Group of the Center for Medicaid and State Operations of the Centers for Medicare & Medicaid Services, in collaboration with the Clinical Laboratory Improvement Advisory Committee at the Centers for Disease Control and Prevention, shall establish a specialty area for molecular and biochemical genetic tests, in order to—

(A) develop criteria for establishing analytic and clinical validity for genetic tests that are determined to require review under the matrix;

(B) specify requirements for proficiency testing for laboratories;

(C) provide guidance regarding the scope of duty for laboratory directors;

(D) make information easily accessible to the public about—

(i) laboratory certification; and
(ii) analytic and clinical validity for genetic tests that are determined to require high level review under the matrix; and

(E) conduct other activities at the discretion of the Administrator of the Centers for Medicare & Medicaid Services.

(3) REIMBURSEMENT.—

(A) CODING.—To foster adoption of genetic screening tools, the Administrator of the Centers for Medicare & Medicaid Services shall—

(i) assess and update current procedure terminology codes to encourage the rapid review and coverage of novel tests through the creation of new HCPCS codes and adoption of new CPT codes and without undue reliance on national coverage determinations; and

(ii) determine and implement fair and reasonable coverage policies and reimbursement rates for medically necessary genetic and genomic treatments and services, including laboratory testing.

(B) BUDGET NEUTRALITY.—Before enhancing payment for a year pursuant to this
paragraph, the Secretary shall, if necessary, provide for an adjustment to payments made under part B of title XVIII of the Social Security Act (42 U.S.C. 1395j et seq.) in that year to ensure that such payments shall be equal to aggregate payments that would have been made under such part in that year if this paragraph had not been enacted.

(e) CENTERS FOR DISEASE CONTROL AND PREVENTION.—

(1) DIRECT-TO-CONSUMER MARKETING.—Not later than 2 years after the date of enactment of this Act, the Director of the Centers for Disease Control and Prevention, with respect to genetic tests for which consumers have direct access, shall—

(A) conduct an analysis of the public health impact of direct-to-consumer marketing to the extent possible from available data sources;

(B) analyze the validity of claims made in direct-to-consumer marketing to determine whether such claims are substantiated by competent and reliable scientific evidence; and

(C) make recommendations to the Secretary regarding necessary interventions to pro-
tect the public from potential harms of direct-
to-consumer marketing and access to genetic
tests.

(2) Public Awareness.—The Director shall
expand efforts to educate and increase awareness of
the general public about genomics and its applica-
tions to improve health, prevent disease and elimi-
nate health disparities. Such efforts shall include
the—

(A) ongoing collection of data on the
awareness, knowledge and use of genetic tests
through public health surveillance systems, and
analysis of the impact of such tests on popu-
lation health; and

(B) integration of the use of validated ge-
netic and genomic tests in public health pro-
grams as appropriate.

(3) Authorization of Appropriations.—
There are authorized to be appropriated to carry out
this subsection, $10,000,000 for fiscal year 2008,
and such sums as may be necessary for each of fis-
cal years 2009 through 2013.

(f) Agency for Healthcare Research and
Quality.—The Director of the Agency for Healthcare
Research and Quality, after consultation with the IWG
and other public and private organizations based in the United States and abroad, as appropriate, shall support the assessment of the clinical utility and cost-effectiveness of companion diagnostic tests that guide prescribing decisions, through research that—

(1) develops standardized tools and methodologies to assess the clinical utility and cost-effectiveness of such tests, as well as criteria for use;

(2) establishes and validates drug dosing algorithms for which such tests can improve outcomes, taking into consideration—

(A) a reduction in toxicity, adverse events, and mortality;

(B) improved clinical outcomes and quality of life, including decreased requirements for monitoring and laboratory testing; and

(C) the impact on the direct and indirect costs of health care, which may include costs due to length of hospital stay, length of time to identify safe and effective dosing for patients, toxicity and adverse events, and other measures of health care utilization and outcomes;

(3) supports and expedites the development of clinical decision tools for clinical use of genetic tests, as warranted; and
(4) prioritizes the development of such tests for diseases and health conditions that have a significant public health impact because of prevalence, risk of complications from treatment, and other factors determined appropriate by the Director.

(g) AUTHORIZATION OF APPROPRIATIONS.—There are authorized to be appropriated to carry out this section, $10,000,000 for fiscal year 2008, and such sums as may be necessary for each of fiscal years 2009 through 2013.