OMNIBUS SOLICITATION OF THE
NATIONAL INSTITUTES OF HEALTH,
CENTERS FOR DISEASE CONTROL AND PREVENTION, AND
FOOD AND DRUG ADMINISTRATION FOR

SMALL BUSINESS INNOVATION
RESEARCH (SBIR)

AND

SMALL BUSINESS TECHNOLOGY
TRANSFER (STTR)

GRANT APPLICATIONS

NIH, CDC, and FDA Program Descriptions and
Research Topics

SUBMISSION DATES

SEPTEMBER 8, 2020, JANUARY 5, 2021, AND
APRIL 5, 2021

National Institutes of Health (SBIR and STTR)
Centers for Disease Control and Prevention (SBIR)
Food and Drug Administration (SBIR)
Funding Opportunity Announcements, Application Instructions, and Appendices are contained in separate files. Follow the links below to view these documents.

FUNDING OPPORTUNITY ANNOUNCEMENTS

REMINDER: ALL APPLICATIONS MUST BE SUBMITTED IN RESPONSE TO A FUNDING OPPORTUNITY ANNOUNCEMENT THROUGH GRANTS.GOV

PHS 2020-02 OMNIBUS SOLICITATION OF THE NIH, CDC, AND FDA FOR SMALL BUSINESS INNOVATION RESEARCH GRANT APPLICATIONS (PARENT SBIR [R43/R44] CLINICAL TRIAL NOT ALLOWED)
HTTP://GRANTS.NIH.GOV/GRANTS/GUIDE/PA-FILES/PA-20-260.HTML

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ADDITIONAL SPECIAL ANNOUNCEMENTS FOR SMALL BUSINESS RESEARCH OPPORTUNITIES HTTPS://SBIR.NIH.GOV/FUNDING/INDIVIDUAL-ANNOUNCEMENTS

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PROGRAM DESCRIPTIONS AND RESEARCH GRANT TOPICS

The research topics shown in this solicitation represent program areas that may be of interest to applicant small business concerns in the development of projects that have potential for commercialization. Small business concerns are encouraged to submit SBIR/STTR grant applications in these areas. Information about the HHS SBIR/STTR programs for applicants and awardees, including resources and programs available to HHS SBIR/STTR awardees, can be found at https://sbir.nih.gov.

Applicable to NIH and CDC Only: SBIR and STTR grant applications will be accepted and considered in any area within the mission of the awarding components (i.e., Institutes and Centers (ICs)) identified in this solicitation.

Applicants are strongly encouraged to subscribe to the NIH Guide for Grants and Contracts LISTSERV (http://grants.nih.gov/grants/guide/listserv.htm) or query program administrators periodically via email to learn of new or emerging scientific interests of the NIH, CDC, and FDA awarding components.

You may also subscribe to the SBIR-STTR LISTSERV list to get timely information about the NIH SBIR/STTR Programs (https://sbir.nih.gov/engage/listserv).

Additional information on each of the awarding components (ICs) and their research interests is available electronically on the home pages shown throughout the “Research Topics” section of the solicitation.

NATIONAL INSTITUTES OF HEALTH (NIH)

NIH is the steward of medical and behavioral research for the Nation. Its mission is science in pursuit of fundamental knowledge about the nature and behavior of living systems and the application of that knowledge to enhance health, lengthen life, and reduce the burdens of illness and disability.

The goals of the agency are as follows:

1. to foster fundamental creative discoveries, innovative research strategies, and their applications as a basis for ultimately protecting and improving health;
2. to develop, maintain, and renew scientific human and physical resources that will assure the Nation’s capability to prevent disease;
3. to expand the knowledge base in medical and associated sciences in order to enhance the Nation's economic well-being and ensure a continued high return on the public investment in research; and
4. to exemplify and promote the highest level of scientific integrity, public accountability, and social responsibility in the conduct of science.

In realizing these goals, the NIH provides leadership and direction to programs designed to improve the health of the Nation by conducting and supporting research:

- in the causes, diagnosis, prevention, and cure of human diseases;
- in the processes of human growth and development;
- in the biological effects of environmental contaminants;
- in the understanding of mental, addictive and physical disorders; and
- in directing programs for the collection, dissemination, and exchange of information in medicine and health, including the development and support of medical libraries and the training of medical librarians and other health information specialists.

In addition, the NIH sponsors training of research personnel; career development of new and established scientists; construction and renovation of research facilities and provision of other research resources.
Information about the NIH SBIR/STTR programs for applicants and awardees, including resources and programs available to NIH SBIR/STTR awardees, can be found at https://sbir.nih.gov.

To carry out these responsibilities, the NIH is organized into awarding components (Institutes/Centers). Those components that have an extramural element, that is, those that provide funds for research and research training activities in organizations external to the NIH, are shown below. The NIH makes every effort to finance worthy applications, including the co-funding of such applications by one or more awarding components having relevance in the projects.

Funding levels for projects are determined through the combined interaction among peer review, grants management, program, budget, and other Institute and/or Centers (IC) staff. These levels are based on allowable costs that are consistent with the principles of sound cost management and in consideration of IC priorities, constraints on the growth of average grant costs, and the availability of funds. For specific topics, NIH may exceed the total award amounts set by the Small Business Administration (SBA) (https://www.sbir.gov). The current list of approved topics can be found in Appendix A.

Before considering and/or preparing an application to the SBIR & STTR programs, all applicants are strongly encouraged to review the agencies’ and NIH Institutes’ and Centers’ websites and to contact the SBIR-STTR program coordinators listed in the Omnibus Solicitation. The Fogarty International Center, which provides support only for conferences, postdoctoral fellowships for research in the United States and abroad, and senior scientist exchanges between the United States and other countries, does not participate in the SBIR/STTR program.

**NIH, CDC, and FDA Awarding Component Contact Information**

Questions of a general nature about the NIH SBIR/STTR program may be directed to:

NIH SBIR/STTR Program Office  
Telephone: 301-435-2688  
Email: sbir@od.nih.gov

For Agency, Institute and Center Scientific/Research (Program) and Financial/Grants Management contacts, please see here:

https://sbir.nih.gov/engage/ic-contacts
NATIONAL INSTITUTE ON AGING (NIA)

NIA’s mission is to:
- Support and conduct genetic, biological, clinical, behavioral, social, and economic research on aging.
- Foster the development of research and clinician-scientists in aging.
- Provide research resources.
- Disseminate information about aging and advances in research to the public, health care professionals, and the scientific community, among a variety of audiences.

The NIA SBIR-STTR Programs support research and product development focusing on aging and aging-related conditions and diseases, as well as other problems and needs unique to older Americans. NIA supports SBIR and STTR research and product development under four divisions: Behavioral and Social Research, Biology of Aging (Aging Biology), Geriatrics and Clinical Gerontology, and Neuroscience.

The NIA will consider any application relevant to the NIA’s mission, even if it does not directly address one of the topics below. For additional information about NIA’s SBIR and STTR programs please visit: https://www.nia.nih.gov/research/osbr

Limited Amount of Award

For budgetary, administrative, or programmatic reasons, the NIA may not fund an application or may decrease the length of an award and/or the budget recommended by a review committee. For topics listed in APPENDIX A: National Institutes of Health SBA-Approved SBIR/STTR Topics for Awards over Statutory Budget Limitations, the NIA generally will not fund Phase I applications to the Omnibus greater than $300,000 total costs or project periods greater than 2 years; or Phase II applications greater than $2,000,000 total costs or project periods greater than 3 years. For all other topics, the NIA does not generally fund Phase I applications greater than $256,580 total costs or project periods greater than 2 years; or Phase II applications greater than $1,710,531 total costs or project periods greater than 3 years. Applicants considering a requested budget greater than these limits are strongly encouraged to contact program staff before submitting an application.

NIA-Supported Funding Opportunity Announcements (FOAs)

In addition to this Omnibus program announcement, the NIA releases targeted Funding Opportunity Announcements (FOAs) throughout the year. These FOAs are listed to inform potential applicants about other funding opportunities to which they can apply; applications submitted in response to this Omnibus program announcement are not limited to research and development areas described in the following targeted FOAs. FOA’s may specify specific budget caps that are above the caps listed for Omnibus applications. Applicants are encouraged to visit the following webpage for an up to date list of NIA SBIR/STTR funding opportunities:

https://www.nia.nih.gov/research/nia-small-business-funding-opportunities

For projects that aim to address Alzheimer’s Disease and Related Dementias, applicants are encouraged to consider the following funding opportunities which allows Phase I budgets up to $500,000 and Phase II budgets up to $2.5M (for topics covered by the approved waiver from SBA):

- Advanced Research on Alzheimer's Disease (AD) and Alzheimer's-Disease-Related Dementias (ADRD) (R43/R44 Clinical Trial Optional): Accepts Phase I, Phase II, Direct-to-Phase II and fast-track applications. Details can be found here: https://grants.nih.gov/grants/guide/pa-files/PAS-19-316.html
• Advancing Research on Alzheimer’s Disease (AD) and Alzheimer’s-Disease-Related Dementias (ADRD) (R41/R42 Clinical Trial Optional): Accepts Phase I, Phase II, and fast-track applications. Details can be found here: https://grants.nih.gov/grants/guide/pa-files/PAS-19-317.html

Phase IIB Competing Renewal Awards and Commercialization Readiness Pilot (CRP)

NIA welcomes submission of Phase IIB Competing Renewal grant applications from Phase II SBIR/STTR awardees to continue the process of developing a wide range of aging-focused products, including digital-mobile/cyber-health technology, pharmaceutical compounds, and medical devices. The Phase IIB Competing Renewal award is intended to allow small businesses the opportunity to realize further progress in commercialization, including stimulating interest in and investment by third parties.

Prospective Phase IIB Competing Renewal applicants are strongly encouraged to contact NIA’s SBIR-STTR program coordinator prior to consideration and preparation of a Phase IIB application and well in advance of the SBIR-STTR submission due dates.

NIA also welcomes the submission of CRP applications to the 3 CRP FOAs:

• SBIR/STTR Commercialization Readiness Pilot (CRP) Program Technical Assistance and Late Stage Development - Clinical Trial Not Allowed (PAR-19-333)
• SBIR/STTR Commercialization Readiness Pilot (CRP) Program Technical Assistance - Clinical Trial Not Allowed (PAR-19-334)
• SBIR/STTR Commercialization Readiness Pilot (CRP) Program Technical Assistance and Late Stage Development - Clinical Trial Required (PAR-19-335)

For questions about potential NIA SBIR/STTR grant applications and NIA’s participation in the Phase IIB or CRP programs, please contact:

Michael-David (“M-D”) ARR Kerns, M.M., M.S., Ph.D.
Program Contact, NIA Small Business R & D Programs
Telephone: 301-402-7713
Email: kernsmd@mail.nih.gov
Bio: https://www.nia.nih.gov/about/staff/kerns-michael-david

Research Topics of Interest to NIA

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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<td>Early Stage Clinical Trials for the Spectrum of Alzheimer's Disease and Age-related Cognitive Decline (R01 Clinical Trial Optional) (PAR-18-877)</td>
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<td>Late Stage Clinical Trials for the Spectrum of Alzheimer's Disease and Age-related Cognitive Decline (R01 Clinical Trial Required) (PAR-18-878)</td>
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<td>Emotional Function in Normal Aging and/or MCI and AD/ADRD (R01 Clinical Trial Optional) (PAR-18-581)</td>
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The NIA will consider any application relevant to the NIA’s mission, even if it does not directly address one of the topics below. The below topics provide an overview of interest areas for both non-clinical trial and clinical trial applications.

A. Alzheimer’s Disease (AD), AD-Related Dementias (ADRD), and Age-Related Change in Brain Function. Research and development of novel interventions to ameliorate AD/ADRD;
improve AD/ADRD care; or further the understanding of the etiology of AD/ADRD, neurodegeneration, brain connectivity, neuroplasticity, or brain—behavior relationships. This includes drug and non-drug interventions for age-related cognitive decline, delirium, sleep disorders, or other central nervous system dysfunctions, including dysfunctions of the motor, emotional, sensory, and neuroimmune systems. This also includes novel biomarkers of neural stem cell functions and new technologies or imaging devices that improve or study brain connectivity; metabolism; sleep; or cognitive, motor, emotional, or sensory activity.

a. For projects addressing AD/ADRD, you may want to consider applying to PAS-19-316 (SBIR) and PAS-19-317 (STTR), which have higher budget limits.

B. Aging in Place. Research and development of social, behavioral, and environmental interventions that promote independence and aging in place by addressing the unique needs of older adults, their healthcare providers, and caregivers. This includes prosthetics, assistive devices and robotics, digital technologies and software, and technology to mitigate age-related physical and behavioral health challenges or to improve healthcare delivery, care coordination, and disease management.

C. Age-Related Diseases and Conditions. Research and development of new diagnostic tools and methods, biomarkers, therapeutics, imaging devices, and technologies to monitor, diagnose, predict, prevent, treat, and further the understanding of the molecular mechanisms of aging or age-related diseases and conditions.

D. Research Tools. Development and validation of innovative tools, resources, or methodologies that promote the efficient, cost-effective, and high-quality collection, analysis, or interpretation of aging-related quantitative or qualitative data. This includes bioinformatics tools; screening platforms; surveying, sampling, and behavioral/behavioral economics methods; and clinical instruments to enhance the study of aging, cellular resiliencies, and aging-related diseases.

Special Emphasis Areas of Interest

Areas of particular interest related to aging biology, aging-related diseases and conditions, behavioral health, and AD/ADRD include the following:

A. Companion diagnostics and other forms of personalized medicine.

B. Bioinformatics, public health informatics, or data science technologies/methods (e.g., machine learning, artificial intelligence) to better understand and predict health outcomes.

C. Novel cell and gene therapies, as well as other novel therapeutic approaches to AD/ADRD.

D. Biomarkers and diagnostic tools for the early detection of disease.

E. Prevention and therapeutics that directly target mechanisms related to aging biology.

F. Assistive technology, devices, and mobile applications for older adults and caregivers.

G. Tools, technologies, and analytic methods to address health disparities among older adults.

For more information on research topics and questions regarding a potential application, contact:

Program Contact, NIA Small Business R & D Programs:
Michael-David ("M-D") A.R.R. Kerns, M.M., M.S., Ph.D.
National Institute on Aging (NIA)
Telephone: 301-402-7713
Email: kernsmd@mail.nih.gov
Bio: https://www.nia.nih.gov/about/staff/dea/kerns-michael-david

If there are specific questions pertaining to the interests or activities of the NIA scientific divisions, contact:

**Division of Aging Biology:**
Max Guo, Ph.D.
National Institute on Aging (NIA)
Telephone: 301-402-7747
Email: max.guo@nih.gov
Bio: https://www.nia.nih.gov/about/staff/guo-qing-bin

**Division of Behavioral and Social Research:**
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Telephone: 301-496-3131
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**Division of Geriatrics and Clinical Gerontology:**
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Bio: https://www.nia.nih.gov/about/staff/joseph-lyndon

**Division of Neuroscience:**
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National Institute on Aging (NIA)
Telephone: 301-827-7130
Email: zane.martin@nih.gov
Bio: https://www.nia.nih.gov/about/staff/martin-jones-zane
NATIONAL INSTITUTE ON ALCOHOL ABUSE AND ALCOHOLISM (NIAAA)

NIAAA supports research on the causes, prevention, control, and treatment of the major health problems associated with alcohol use. Through its extramural research programs, NIAAA funds a wide range of basic and applied research to develop new and/or improved technologies and approaches for increasing the effectiveness of diagnosis, treatment, and prevention. NIAAA also is concerned with strengthening research dissemination, scientific communications, public education, and data collection activities in the areas of its research programs. NIAAA is interested in studies that examine racial, ethnic and gender minorities as well as other underserved populations that experience more negative alcohol-related consequences of illness and premature death than the general population.

Limited Amount of Award

NIAAA will make awards compliant with all statutory guidelines as outlined above. Total funding support (direct costs, indirect costs, fees) normally may not exceed $256,580 for Phase I awards and $1,710,531 for the duration of the Phase II awards. With appropriate justification from the applicant, NIAAA may consider awards that exceed these amounts by up to 50%. NIAAA has received a budgetary guideline waiver from the Small Business Administration for applications relating to the limited list of scientific topics (Appendix A). Applicants considering a requested budget greater than the standard limits are strongly encouraged to contact the NIAAA SBIR/STTR Program Coordinator before submitting an application. For budgetary, administrative, or programmatic reasons, NIAAA may decrease the length of an award and/or the budget recommended by a review committee, or not fund an application.

Phase IIB Competing Renewal Awards

NIAAA will accept SBIR/STTR Phase IIB Competing Renewal grant applications from Phase II SBIR/STTR awardees to continue the process of developing products that require approval of a Federal regulatory agency (e.g., FDA, FCC). Such products include, but are not limited to, medical implants, drugs, vaccines, biologicals, and new treatment or diagnostic tools that require FDA approval. This renewal grant should allow small businesses to get to a stage where interest and investment by third parties is more likely. To be eligible for Phase IIB consideration, the project must retain high significance in the light of current market conditions.

Prospective applicants are strongly encouraged to contact NIH staff well in advance of submitting a Phase IIB Competing Renewal application by submitting to Ms. Megan Ryan (contact information below) a letter of intent that includes the following information:

- Name, address, and telephone number of the Principal Investigator
- Names of other key personnel
- Participating institutions
- Grant number and title
- Progress of the Phase II award
- Goals and justification for the Phase IIB request

It is expected that only a portion of NIAAA SBIR/STTR Phase II awards will be eligible for a Phase IIB Competing Renewal grant.

NIAAA supports Investigational New Drug (IND)-enabling studies for therapeutic candidates under separate FOAs (https://niaaa.nih.gov/grant-funding/funding-opportunities).

Commercialization Assistance Programs

NIAAA Phase I grantees may consider applying for the I-Corps at NIH pilot program.
Research Topics of Interest to NIAAA

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NIAAA Non-Clinical Trials Topics:

Medications Development
Areas that may be of interest to small businesses include, but are not limited to:

- Therapeutics for Alcohol Use Disorder (AUD) and Alcohol-Related Complications
  - Targets that include but are not limited to agents that attenuate excessive alcohol use and other symptoms of AUD, e.g., craving, sleep problems, and negative affect as well as those that hold promise for the treatment of alcoholic hepatitis, liver fibrosis, cirrhosis, pancreatitis, cardiomyopathy, or other alcohol-induced tissue damage.
  - Development and validation of medium to high-throughput assays
  - Early therapeutic discovery activities (e.g., target ID, lead compound target validation)
  - Preclinical drug development
  - Devices, technologies or formulations to improve medication delivery, medication compliance or improved formulations of existing medications to treat AUD

For pre-clinical questions, contact:
Mark Egli, PhD (Neuroscience and Behavior)
Telephone: 301-594-6382
Email: Mark.Egli@nih.gov

Svetlana Radaeva, PhD (Organ Damage)
Telephone: 301-433-1189
Email: Svetlana.Radaeva@nih.gov

- Preclinical development of combined pharmacological approaches to synergistically regulate multiple drug targets for AUD.
  - Areas that may be of interest to small businesses include, but are not limited to:
    - Prioritization of multi-drug targets and identification of the effective drug combinations or multi-target drugs for the medication development
    - High-throughput screening of compound libraries to identify multi-target drugs.
    - Adaption of low throughput assays to high throughput screening, development of lead compounds, and identification of drug candidate(s) with proper pharmaceutical properties for medication development.

Changhai Cui, PhD
Telephone: 301-443-1678
Email: Changhai.Cui@nih.gov

**Technological Methods for the Treatment of Hazardous Drinking and Alcohol Use Disorder**

Areas that may be of interest to small businesses include, but are not limited to:

- Development, enhancement, and/or validation of daily process data collection methods (e.g., ecological momentary assessment) for capturing real-time data for use in alcohol clinical trials and treatment paradigms.
- Development and testing computerized versions of empirically supported treatments, including but not limited to in languages other than English.
- Evaluation of telehealth and mHealth technologies to improve patient outcomes and increase and improve patient engagement in treatment.
Prevention

NIAAA is interested in the development and evaluation of the following:

- Innovative prevention/intervention programs for alcohol misuse, or specific materials for integration into existing prevention programs, that utilize state-of-the-art technology and are based on currently accepted clinical and behavioral strategies.
- Innovative prevention/intervention programs for alcohol misuse targeted at the needs of groups at elevated risk for such misuse, including racial/ethnic minority populations, adolescents, women, and sexual and gender minority populations.
- Educational materials designed to intervene with the elderly around specific age-related risks for alcohol problems.
- Prevention programs tailored specifically to the needs of any of the following groups: children of individuals with alcohol use disorders, women, ethnic and minority populations, sexual and gender minority populations, persons with disabilities, and the elderly.
- School-based prevention curricula, interactive videos, multi-media programs, and training manuals for teachers and parents.
- Development of statistical analysis tools for: imputation of missing data, analysis of small sample sizes, simulations of the distribution of outcomes under varying initial conditions, and evaluation of the effect of public policy on health and injury outcomes.
- Approaches, strategies, and platforms to increase public awareness of avoidable risks and encourage behavior change that reduces those risks through publicity about actual traffic crashes, crash outcomes (e.g., deaths, injuries, family and community effects), and the role of alcohol-impaired driving and other factors that contributed to those crashes (e.g., speeding, failure to use seatbelts).

Improving the Delivery of Alcohol Treatment Services

Areas that may be of interest to small businesses include, but are not limited to:

- Assisting clinicians in selecting and delivering evidence-based treatments
- Supporting long-term recovery, by facilitating patients’ continued engagement in recovery support services as an adjunct to or after treatment
- Assisting treatment programs and service agencies in measuring clinically relevant performance indicators or improvements in quality of service provision
- Promoting engagement and mitigate burnout among counselors and others engaged in direct treatment service delivery

Laura Kwako, PhD
Telephone: 301-451-8507
Email: Laura.Kwako@nih.gov
**Fetal Alcohol Spectrum Disorders (FASD) and Prenatal Alcohol Exposure**

Areas that may be of interest to small businesses include, but are not limited to:

- Development and assessment of novel diagnostic and/or screening methods, tools, or technologies to improve identification and diagnosis of individuals affected by prenatal exposure across the lifespan and in a variety of setting.
- Validation of biomarkers that can be used to verify prenatal alcohol exposure in newborns or later in life or to predict which individuals will display neurobehavioral deficits later in life.
- Novel strategies for therapeautic, skill-building, and educational program products that enhance behavioral, neurocognitive, social, adaptive, and motor function to improve the overall well-being of individuals with FASD and their families.
- Establishment of accurate measures of the responsiveness of individuals affected by prenatal alcohol exposure to predictors of vulnerability for alcohol-drinking or other psychopathology during adolescence and adulthood.
- Development, evaluation, and implementation of novel educational and training programs to enhance the skills of non-professional caregivers in dealing with the problems associated with FASD.
- Development and implementation of innovative strategies, methods, tools or technologies to reduce alcohol misuse in women and prevent FASD in offspring.

**For biomedical research questions, contact:**

William Dunty, PhD  
Telephone: 301-443-7351  
Email: William.Dunty@nih.gov

Elizabeth Powell, PhD  
Telephone: 301-443-0786  
Email: Elizabeth.Powell3@nih.gov

**For prevention and treatment research questions, contact:**

Tatiana Balachova, PhD  
Telephone: 301-443-5726  
Email: Tatiana.Balachova@nih.gov

Deidra Roach, MD  
Telephone: 301-443-5820  
Email: Deidra.Roach@nih.gov

**Development of Clinical Biomarkers of Alcohol Exposure and Alcohol-Induced Organ Damage**

Areas that may be of interest to small businesses include, but are not limited to:

- Biomarkers that detect alcohol intake that is measurable for two to three weeks after drinking is stopped.
- Biomarkers that detect alcohol intake that is measurable for months after drinking is stopped.
- Biomarker signatures of alcohol-induced organ damage, which are likely to be organ-specific, particularly using the early stages of alcohol-induced organ damage.
- Biomarker signatures of familial risk factors for AUD. Early identification of subjects predisposed to AUD will allow for early intervention, possible prevention, and allow the subjects to make informed personal decisions.
Characteristics of useful biomarkers are:

- Sensitivity, specificity, accuracy, and reliability
- Ease of use and acceptability to patient and provider
- Affordability
- Availability in easily obtained specimens, such as serum or plasma, urine, saliva, or hair.
- Validity, reproducibility, affordability, and transportability to a variety of settings, including AUD treatment centers, hospitals, primary care offices, or the workplace.

We encourage use of high-throughput discovery approaches using genomics, epigenomics, transcriptomics, proteomics, metabolomics, lipomics, or glycomics to identify pattern-based molecular signatures which as opposed to single component biomarkers may be predicted to provide greater sensitivity, specificity, accuracy, and reliability than single component biomarkers.

Also of interest:

- Improvement of turn-around time and cost efficiency of current assays for PEth, EtS, EtG and other alcohol biomarkers.
- Design and development of point of care devices, for use in rural or remote primary care and hospital settings

**For clinical questions, contact:**
Raye Z. Litten, PhD
Telephone: 301-443-0636
Email: Raye.Litten@nih.gov

**For pre-clinical questions, contact:**
Kathy Jung, PhD
Telephone: 301-443-8744
Email: Kathy.Jung@nih.gov

**Wearable Alcohol Biosensors**

We are seeking non-invasive or minimally invasive technologies. In particular, this topic should focus on devices that detect blood alcohol, rather than alcohol exuded across the skin. Characteristics of these technologies include:

- Real-time or near real-time detection of alcohol levels
- Ability to interpret and store the data or transmit it to a smartphone or other device by wireless transmission
- Standardization verification at regular intervals; ability to indicate loss of functionality
- A power source that is dependable and rechargeable
- Data storage and transmission that is completely secure in order to protect the privacy of the individual. A form of subject identification would be an added benefit.
- Unobtrusive design, passive in action

Also of interest is the development of appropriate data analysis systems for individual level evaluation of alcohol biosensor devices already in development, as well as for assessment of trends in research populations.

Kathy Jung, PhD.
Telephone: 301-443-8744
Alcohol Use and HIV Infection, and HIV Co-infection with HCV, HBV, or TB

Areas that may be of interest to small businesses include, but are not limited to:

• Development of therapies to mitigate alcohol-associated adverse impact on the development of viral-related liver and/or lung diseases.
• Development of biomarkers for alcohol-induced damage in those patients with HIV infection or co-infection.

For basic research questions on alcohol and HIV, and HIV co-infection, contact:
H. Joe Wang, PhD
Telephone: 301-451-0747
Email: Joe.Wang1@nih.gov

For clinical or epidemiological questions on HIV, contact:
Kendall J. Bryant, PhD
Telephone: 301-402-9389
Email: Kendall.Bryant@nih.gov

Electronic Diagnostic Clinical Assessment of Frailty among HIV+ Individuals with Past and/or Current Alcohol Use Disorder: Severity and Patterns

Innovative self-report, biological, and/or common clinical measures for the identification and diagnosis of frailty related to alcohol use among alcohol-using HIV patients and those with related comorbidities are sought.

• Should be tested in the widest range of individuals, including but not limited to racial, ethnic and gender identity minorities, at various trajectories of progression of HIV disease and patterns of alcohol use. In particular, information from measures should be able to accurately identify individuals who are “sick quitters” and/or have high degree of frailty due to either past and/or current alcohol use.
• This clinical decision-making tool should be of greatest value to diagnostic assessment and interventions within clinical settings and may include the development of audio, visual, and/or training modules to support the use of appropriate diagnostic index(es) across linguistic groups.
• Support of an electronic internet site for scoring and collection of information on HIV disease characteristics, frailty and patterns of alcohol use in clinical populations, and to provide information on a range of options for assessment of alcohol use severity in HIV+ populations (e.g. brief assessment instruments, calendar methods, biological markers, etc.).
• Identification of current and emerging methods for behavioral and/or biological intervention to reduce alcohol use in the context of HIV and improve clinical outcomes.

Kendall J. Bryant, PhD
Telephone: 301-402-0332
Email: Kendall.Bryant@nih.gov

Stem Cell Research for Alcohol-Induced Disorders

Areas that may be of interest to small businesses include, but are not limited to:
• Generation and dissemination of induced pluripotent stem cells (iPS) from adult human cells to resemble diverse individual variations in alcohol metabolism and use of these genetic variant models to study AUD and pharmacotherapy development. Examples of these genetic variations include alcohol dehydrogenase (ADH), aldehyde dehydrogenase (ALDH), cytochrome P450 isozyme CYP2E1, and glutathione S-transferase (GST).
• Generation and dissemination of disease-specific iPS cell lines for studies on the biology and signaling pathways that contribute to alcohol-related disease pathology.
• Development of models derived from human iPS cells to study biological and pathological effects of alcohol and its metabolites in vivo.
• Use of CRISPR (clustered regularly interspaced short palindromic repeats) gene editing technology on iPS cells to study alcohol-related disease

Peter Gao, MD  
Telephone: 301-443-6106  
Email: Peter.Gao@nih.gov

Role of Non-Coding RNAs in the Neuroadaptation to Alcohol Use Disorder

Areas that may be of interest to small businesses include, but are not limited to novel technologies to both measure and interpret ncRNA gene expression signatures in specific cell types and regions in the brain, and/or primary neuronal cultures following alcohol exposure.

• novel technologies to both measure and interpret ncRNA gene expression signatures in specific cell types and regions in the brain, and/or primary neuronal cultures following alcohol exposure.
• These technologies could include, but are not limited to:
  o Novel methods to tag and measure various types of ncRNAs,
  o New imaging techniques to monitor changes in ncRNAs, and
  o Novel bioinformatic analytic tools and methods to interpret alcohol-induced alterations in ncRNAs, and to predict and validate their target genes.

Hemin Chin, PhD  
Telephone: 301-443-1282  
Email: Hemin.Chin@nih.gov

In vivo Detection of Neuromodulators in Behaving Animals

Areas that may be of interest to small businesses include, but are not limited to:

• Development of effective tools to detect dynamic changes of neuromodulators in real time in the brain of behaving animals
• Development of tools to visualize and map neuromodulator changes in the intact brain coupled with other measurements of neuronal activities. To understand how neuromodulators shape neuronal activities and contribute to the alcohol use disorder.
• Development of technologies to detect concurrent changes of multiple neuromodulators with high spatial and temporal precision in vivo.
• Development of tools that allow the in vivo detection of the dynamics of neuromodulators over an extended time period.

Changhai Cui, PhD  
Telephone: 301-443-1678  
Email: Changhai.Cui@nih.gov
**Ex vivo Screens to Identify Pharmacotherapies for Alcohol Use Disorder**

Areas that may be of interest to small businesses include, but are not limited to:

- Development and validation of *ex vivo* screens capable of predicting efficacy test results in preclinical behavioral models of AUD
  
  Assays should:
  
  - include array of parameters that model aspect/aspects of AUD
  - be able to discriminate positive and negative control drugs found in the AUD pharmacotherapy literature
  - be sensitive to drugs with diverse mechanisms of action
  - be relatively rapid, simple and capable of generate readily reproducible results

Qi-Ying Liu, MD, M.Sci.
Telephone: 301-443-2678
Email: liuqiy@mail.nih.gov

**Novel Tools and Technologies to Detect the Effects of Alcohol on the CNS Structure and Activities**

Areas that may be of interest to small businesses include, but are not limited to:

- Improvement of chemical or genetic sensors to detect dynamic changes in calcium, voltage, cAMP etc.
- Development of tools and sensors to monitor structure and activities of neurons and glial cells, and their interactions
- Development of tools and sensors to monitor synaptic activities
- Definition of cell types in the neurocircuits
- Development of tools and sensors to monitor synaptic activities
- Development of miniature and nanoscale apparatus and sensors, or miniaturizing and optimizing detection apparatus for the study of alcohol effects
- Development of computational methods for the acquisition and analysis of large scale data

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Telephone: 301-443-2678
Email: liuqiy@mail.nih.gov

Changhai Cui, PhD
Telephone: 301-443-1678
Email: changhai.cui@nih.gov

**Research Tools**

The NIAAA supports the development of new or improved tools to enhance the ability to conduct alcohol-related laboratory studies on humans and animals and to more effectively analyze data from large databases. Areas that may be of interest to small businesses include, but are not limited to:

- Development of novel animal models, including transgenic animals, possessing specific traits of significance for the study of alcohol use disorder (AUD), or for the study of specific pathologic disease states which arise from excessive alcohol consumption.
- Development of a hepatocyte cell line capable of maintaining viability and metabolic functions in culture systems for an indefinite period.
- Development of specialized cell culture chambers to provide controlled administration of ethanol to *in vitro* cell systems.
• Development of experimental systems that mimic organ function, including, but not limited to, co-culture and novel approaches to three-dimensional culture.
• Development of new methods of ethanol administration to animals that produce precise dose control or that closely mimic types of alcohol exposure occurring in humans, including, but not limited to, binge drinking, acute consumption, moderate consumption or chronic consumption.
• Development of ligands which will enhance the potential usefulness of PET and SPECT neuroimaging technologies for the study of the etiology of AUD and related brain pathology.
• Development of computational, statistical or bioinformatics tools to organize and manage high throughput data obtained by genomic, functional genomic or other ‘omic strategies.
• Development of databases, methods for integration of databases, or data analysis systems for alcohol research.

Kathy Jung, PhD
Telephone: 301-443-8744
Email: Kathy.Jung@nih.gov

Data Science Tools for Integrating Alcohol Research

Areas of interest include, but are not limited to:
• Development of algorithms for integrative analysis incorporating multiple current NIAAA and public ‘big data’ sets, including machine learning, deep learning, artificial intelligence, data mining and other model based and model-free approaches.
• Creation of software applications for data interfaces for aggregation, imputation, harmonization, or visualization of data from multiple sources, including current and future NIH data systems.
• Design of algorithms and/or software tools for improving data collection, i.e. smart phone apps, extraction of specific alcohol research parameters from existing large databases and established public health studies, biological sensors or wearable devices.
• Generation and validation of computational and/or systems biology models of alcohol exposure and use on cellular, organ, network, or organism scales, including multiscale models, during periods from initial alcohol exposure and extending though alcohol use disorder, treatment, recovery and relapse.

Activities and deliverables are expected to use currently available data sets and databases. The generation of new primary data is not supported by this topic.

Elizabeth Powell, PhD
Telephone: 301-443-0786
Email: Elizabeth.Powell3@nih.gov

Development of Novel Tools to Measure Potentially Toxic Alcohol Metabolites and Advanced Glycation End Products in Body Fluids & Other Organs

Areas of interest include, but are not limited to:
• Development of tools/kits for early detection and to measure AA-adducts and AGEs in serum, CSF, brain and other organs of AUDs in animal models and in pre-clinical settings and their relationship to the biomarkers of neuro-inflammation.
• Development of tools/kits to measure AA-adducts and AGEs, like other oxidative and carbonyl stress end products; advanced lipoperoxidation end products (ALE) and advanced oxidation protein products (AOPP).
- Development of tools to study interaction of AA-adducts and AGEs with RAGE (Receptor for AGEs) and measurement of structural damage to the extracellular matrix and other components.

Mohammed Akbar, PhD  
Telephone: 301-443-6009  
Email: Akbarm@mail.nih.gov

Single Cell Genomics (RNA Sequencing): Application to Develop Molecular Markers of Alcohol Use Disorder

Areas of interest include, but are not limited to:

- Development of rapid and cost-effective RNA seq methods/technologies in human blood cells  
- Characterization of the gene expression profile in human blood cells of AUD subjects.  
- Identification and validation of specific gene profile signature as biomarkers in AUD subjects.
- Development of technology/device for rapid screening different stages of alcohol use and AUD

Abbas Parsian, PhD  
Telephone: 301-443-5733  
E-mail: Parsiana@nih.gov

**NIAAA Clinical Trials Topics:**

**Medications Development**

Areas that may be of interest to small businesses include, but are not limited to:

- Clinical development of therapeutics for AUD and Alcohol-Related Complications.  
- Novel and re-purposed compounds for the treatment of AUD.  
- Compounds that target one or more domains of the addiction cycle, including reward, stress and negative affect, incentive salience, executive function, habituation, and impulsivity/compulsivity.  
- Devices, technologies or formulations to improve medication delivery, medication compliance or improved formulations of existing medications to treat AUD.

**For questions, contact:**  
Raye Z. Litten, PhD  
Telephone: 301-443-0636  
Email: Raye.Litten@nih.gov

**Technological Methods for the Treatment of Hazardous Drinking and Alcohol Use Disorder**

Areas that may be of interest to small businesses include, but are not limited to:

- Development, enhancement, and/or validation of daily process data collection methods (e.g., ecological momentary assessment) for capturing real-time data for use in alcohol clinical trials and treatment paradigms.  
- Development and testing computerized versions of empirically supported treatments, including but not limited to in languages other than English.  
- Evaluation of telehealth and mHealth technologies to improve patient outcomes and increase and improve patient engagement in treatment.
Prevention

NIAAA is interested in the development and evaluation of the following:

- Innovative prevention/intervention programs for alcohol misuse, or specific materials for integration into existing prevention programs, that utilize state-of-the-art technology and are based on currently accepted clinical and behavioral strategies.
- Innovative prevention/intervention programs for alcohol misuse targeted at the needs of groups at elevated risk for such misuse, including racial/ethnic minority populations, adolescents, women, and sexual and gender minority populations.
- Educational materials designed to intervene with the elderly around specific age-related risks for alcohol problems.
- Prevention programs tailored specifically to the needs of any of the following groups: children of individuals with alcohol use disorders, women, ethnic and minority populations, sexual and gender minority populations, persons with disabilities, and the elderly.
- School-based prevention curricula, interactive videos, multi-media programs, and training manuals for teachers and parents.
- Development of statistical analysis tools for: imputation of missing data, analysis of small sample sizes, simulations of the distribution of outcomes under varying initial conditions, and evaluation of the effect of public policy on health and injury outcomes.
- Approaches, strategies, and platforms to increase public awareness of avoidable risks and encourage behavior change that reduces those risks through publicity about actual traffic crashes, crash outcomes (e.g., deaths, injuries, family and community effects), and the role of alcohol-impaired driving and other factors that contributed to those crashes (e.g., speeding, failure to use seatbelts).

Improving the Delivery of Alcohol Treatment Services

Areas that may be of interest to small businesses include, but are not limited to:

- Assisting clinicians in selecting and delivering evidence-based treatments
- Supporting long-term recovery, by facilitating patients’ continued engagement in recovery support services as an adjunct to or after treatment
- Assisting treatment programs and service agencies in measuring clinically relevant performance indicators or improvements in quality of service provision
- Promoting engagement and mitigate burnout among counselors and others engaged in direct treatment service delivery.
Fetal Alcohol Spectrum Disorders (FASD) and Prenatal Alcohol Exposure

Areas that may be of interest to small businesses include, but are not limited to:

- Development and assessment of novel diagnostic and/or screening methods, tools, or technologies to improve identification and diagnosis of individuals affected by prenatal exposure across the lifespan and in a variety of setting.
- Validation of biomarkers that can be used to verify prenatal alcohol exposure in newborns or later in life or to predict which individuals will display neurobehavioral deficits later in life.
- Novel strategies for therapeudic, skill-building, and educational program products that enhance behavioral, neurocognitive, social, adaptive, and motor function to improve the overall well-being of individuals with FASD and their families.
- Establishment of accurate measures of the responsiveness of individuals affected by prenatal alcohol exposure to predictors of vulnerability for alcohol-drinking or other psychopathology during adolescence and adulthood.
- Development, evaluation, and implementation of novel educational and training programs to enhance the skills of non-professional caregivers in dealing with the problems associated with FASD.
- Development and implementation of innovative strategies, methods, tools or technologies to reduce alcohol misuse in women and prevent FASD in offspring.

For biomedical research questions, contact:
William Dunty, PhD
Telephone: 301-443-7351
Email: William.Dunty@nih.gov

Elizabeth Powell, PhD
Telephone: 301-443-0786
Email: Elizabeth.Powell3@nih.gov

For prevention and treatment research questions, contact:
Tatiana Balachova, PhD
Telephone: 301-443-5726
Email: Tatiana.Balachova@nih.gov

Deidra Roach, M.D.
Telephone: 301-443-5820
Email: Deidra.Roach@nih.gov

Development of Clinical Biomarkers of Alcohol Exposure and Alcohol-Induced Organ Damage

Areas that may be of interest to small businesses include, but are not limited to:

- Biomarkers that detect alcohol intake that is measurable for two to three weeks after drinking is stopped.
- Biomarkers that detect alcohol intake that is measurable for months after drinking is stopped.
- Biomarker signatures of alcohol-induced organ damage, which are likely to be organ-specific, particularly using the early stages of alcohol-induced organ damage.
- Biomarker signatures of familial risk factors for AUD. Early identification of subjects predisposed to AUD will allow for early intervention, possible prevention, and allow the subjects to make informed personal decisions.
Characteristics of useful biomarkers are:

- Sensitivity, specificity, accuracy, and reliability
- Ease of use and acceptability to patient and provider
- Affordability
- Availability in easily obtained specimens, such as serum or plasma, urine, saliva, or hair.
- Validity, reproducibility, affordability, and transportability to a variety of settings, including AUD treatment centers, hospitals, primary care offices, or the workplace.

We encourage use of high-throughput discovery approaches using genomics, epigenomics, transcriptomics, proteomics, metabolomics, lipomics, or glycomics to identify pattern-based molecular signatures which as opposed to single component biomarkers may be predicted to provide greater sensitivity, specificity, accuracy, and reliability than single component biomarkers.

Also of interest:

- Improvement of turn-around time and cost efficiency of current assays for PEth, EtS, EtG and other alcohol biomarkers.
- Design and development of point of care devices, for use in rural or remote primary care and hospital settings

For clinical questions, contact:
Raye Z. Litten, PhD
Telephone: 301-443-0636
Email: Raye.Litten@nih.gov

For pre-clinical questions, contact:
Kathy Jung, PhD
Telephone: 301-443-8744
Email: Kathy.Jung@nih.gov

Wearable Alcohol Biosensors

We are seeking non-invasive or minimally invasive technologies. In particular, this topic should focus on devices that detect blood alcohol, rather than alcohol exuded across the skin. Characteristics of these technologies include:

- Real-time or near real-time detection of alcohol levels
- Ability to interpret and store the data or transmit it to a smartphone or other device by wireless transmission
- Standardization verification at regular intervals; ability to indicate loss of functionality
- A power source that is dependable and rechargeable
- Data storage and transmission that is completely secure in order to protect the privacy of the individual. A form of subject identification would be an added benefit.
- Unobtrusive design, passive in action

Also of interest is the development of appropriate data analysis systems for individual level evaluation of alcohol biosensor devices already in development, as well as for assessment of trends in research populations.

Kathy Jung, PhD
Telephone: 301-443-8744
Email: Kathy.Jung@nih.gov
Alcohol Use and HIV Infection, and HIV Co-infection with HCV, HBV, or TB

Areas that may be of interest to small businesses include, but are not limited to:

- Development of therapies to mitigate alcohol-associated adverse impact on the development of viral-related liver and/or lung diseases.
- Development of biomarkers for alcohol-induced damage in those patients with HIV infection or co-infection.

For basic research questions on alcohol and HIV, and HIV co-infection, contact:
H. Joe Wang, PhD
Telephone: 301-451-0747
Email: Joe.Wang1@nih.gov

For clinical or epidemiological questions on HIV, contact:
Kendall J. Bryant, PhD
Telephone: 301-402-9389
Email: Kendall.Bryant@nih.gov

Electronic Diagnostic Clinical Assessment of Frailty among HIV+ Individuals with Past and/or Current Alcohol Use Disorder: Severity and Patterns

Innovative self-report, biological, and/or common clinical measures for the identification and diagnosis of frailty related to alcohol use among alcohol-using HIV patients and those with related comorbidities are sought.

- Should be tested in the widest range of individuals, including but not limited to racial, ethnic and gender identity minorities, at various trajectories of progression of HIV disease and patterns of alcohol use. In particular, information from measures should be able to accurately identify individuals who are "sick quitters" and/or have high degree of frailty due to either past and/or current alcohol use.
- This clinical decision-making tool should be of greatest value to diagnostic assessment and interventions within clinical settings and may include the development of audio, visual, and/or training modules to support the use of appropriate diagnostic index(es) across linguistic groups.
- Support of an electronic internet site for scoring and collection of information on HIV disease characteristics, frailty and patterns of alcohol use in clinical populations, and to provide information on a range of options for assessment of alcohol use severity in HIV+ populations (e.g. brief assessment instruments, calendar methods, biological markers, etc.).
- Identification of current and emerging methods for behavioral and/or biological intervention to reduce alcohol use in the context of HIV and improve clinical outcomes.

Kendall J. Bryant, PhD
Telephone: 301-402-0332
Email: Kendall.Bryant@nih.gov

Development of Novel Tools to Measure Potentially Toxic Alcohol Metabolites and Advanced Glycation End Products in Body Fluids & Other Organs

Areas of interest include, but are not limited to:
• Development of tools/kits for early detection and to measure AA-adducts and AGEs in serum, CSF, brain and other organs of AUDs in animal models and in pre-clinical settings and their relationship to the biomarkers of neuro-inflammation.
• Development of tools/kits to measure AA-adducts and AGEs, like other oxidative and carbonyl stress end products; advanced lipoperoxidation end products (ALE) and advanced oxidation protein products (AOPP).
• Development of tools to study interaction of AA-adducts and AGEs with RAGE (Receptor for AGEs) and measurement of structural damage to the extracellular matrix and other components.
• Measurement of AGE–RAGE mediated activation of nuclear factor NF-B, cytokines and growth factors genes and increased expression of neuro-inflammatory molecules.

Mohammed Akbar, PhD
Telephone: 301-443-6009
Email: Akbarm@mail.nih.gov

Single Cell Genomics (RNA Sequencing): Application to Develop Molecular Markers of Alcohol Use Disorder

Areas of interest include, but are not limited to:
• Development of rapid and cost-effective RNA seq methods/technologies in human blood cells
• Characterization of the gene expression profile in human blood cells of AUD subjects.
• Identification and validation of specific gene profile signature as biomarkers in AUD subjects.
• Development of technology/device for rapid screening different stages of alcohol use and AUD

Abbas Parsian, PhD
Telephone: 301-443-5733
E-mail: Parsiana@nih.gov

Direct your general questions about the SBIR/STTR program or scientific/research issues to:

Megan Ryan, M.B.A.
National Institute on Alcohol Abuse and Alcoholism
6700B Rockledge Dr.
Rockville, MD 20852-1705

For administrative and business management questions, contact:

Mr. Jeff Thurston
Grants Management Specialist
National Institute on Alcohol Abuse and Alcoholism
Phone: 301-443-9801, Fax: 301-443-3891
Email: jeffrey.thurston@nih.gov
NATIONAL INSTITUTE OF ALLERGY AND INFECTIOUS DISEASES (NIAID)

NIAID conducts and supports basic and applied research to better understand, treat, and ultimately prevent infectious, immunologic, and allergic diseases.

NIAID's Division of AIDS (DAIDS), Division of Allergy, Immunology, and Transplantation (DAIT), and Division of Microbiology and Infectious Diseases (DMID) encourage SBIR/STTR applications related to their mission and activities as described below. Questions regarding specific research areas may be addressed to the NIAID Program Officials listed below. General questions about the NIAID SBIR and STTR programs or administrative and business management concerns may be directed to the NIAID Small Business Program Team (https://www.niaid.nih.gov/grants-contracts/small-business-program-team).

When possible, applicants are encouraged to use email for communication.

For information about NIAID's Small Business Programs, please visit https://www.niaid.nih.gov/grants-contracts/small-businesses.

Limited Total Amounts for Phase I and Phase II Awards

Total funding support (direct costs, indirect costs, fee) normally may not exceed $256,580 for Phase I awards and $1,710,531 for Phase II awards. Budget requests at or near these hard caps should be well justified. Phase II/IIB applicants should note that NIAID will not generally allow awards (of any duration) that exceed $1,000,000 total costs per year.

NIH has received a waiver from SBA, as authorized by statute, to exceed total award amount hard caps for specific topics. The current list of approved topics can be found in Appendix A of this document or at https://sbir.nih.gov/funding#omni-sbir. Approved topics that align with NIAID’s priority research areas are listed for each Division; however, a listed topic from any IC is sufficient to consider budget requests that exceed the hard caps. Budget requests exceeding the hard caps must be very well justified in the “Budget Justification” attachment to the Research and Related Budget form and be clearly consistent with the scope of the proposal.

For proposals that address an approved topic, NIAID will allow Phase I applications with budgets of up to $300,000 total costs per year for up to 2 years; and Phase II or Phase IIB applications with budgets of up to $1,000,000 total costs per year for up to 3 years. Requests for these budget levels must be very well-justified. In all cases, applicants should propose a budget that is reasonable and appropriate for completion of the research project.

NIAID staff cannot provide prior approval to exceed hard caps. Compliance with a pre-approved topic will be confirmed at time of award by the applicant’s Grants Management Specialist and Program Officer.

NIAID will generally not make SBIR or STTR awards with budgets that exceed these guidelines. For budgetary, administrative, or programmatic reasons, NIAID may decide not to fund an application or may decrease the length of an award and/or the budget recommended by a review committee.

Phase IIB Competing Renewal Awards

NIAID welcomes Phase IIB Competing Renewal Applications (SBIR only) for Phase II grants and contracts via the Omnibus Solicitation for SBIR Grant Applications, and as indicated by other NIAID Funding Opportunity Announcements (FOAs). Standard NIAID Phase II funding policy applies unless otherwise stated in the FOA. STTR Phase II awardees may apply but must switch programs to SBIR. Non-NIAID Phase II awardees must contact NIAID prior to submission to confirm programmatic interest.
Research Topics of Interest to NIAID

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
<th>No*</th>
<th>Yes**</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>X</td>
</tr>
</tbody>
</table>

*If No,
- This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, UNLESS the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,
- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency accept clinical trials applications under this mechanism?</th>
<th>Yes</th>
<th>No</th>
<th>Other Information:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Trials SBIR/STTR Omnibus/Parent Funding Opportunity Announcement/s</td>
<td></td>
<td>X</td>
<td>If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information.</td>
</tr>
</tbody>
</table>
NIAID Non-Clinical Trial Topics:

Division of AIDS (DAIDS)

The Division of AIDS (DAIDS) supports a global research portfolio to advance biological knowledge of HIV/AIDS, its related co-infections, and co-morbidities. With the ultimate goal of creating an “AIDS-Free Generation,” the division develops and supports the infrastructure and biomedical research needed to: 1) halt the spread of HIV through the development of an effective vaccine and biomedical prevention strategies that are safe and desirable; 2) develop novel approaches for the treatment and cure of HIV infection; 3) treat and/or prevent HIV co-infections and co-morbidities of greatest significance; and 4) partner with scientific and community stakeholders to efficiently implement effective interventions.

Director: Dr. Carl Dieffenbach
Telephone: 301-496-0545
Email: CDieffenba@niaid.nih.gov

Basic Sciences Program

Supports basic and applied research on the causes, diagnosis, treatment, and prevention of HIV and AIDS.

Director: Dr. Diana Finzi
Telephone: 301-451-2598
Email: Dfinzi@niaid.nih.gov

A. Epidemiology Branch. Population-based research, modeling, and comparative effectiveness studies (not including clinical trials) that assess the natural history, biologic, and clinical course of HIV/AIDS, and related outcomes, and could advance treatment and prevention of HIV. Specific interests include phylodynamics and other factors related to HIV transmission and associated biological and behavioral factors, basic research on immunology, virology, and antiretroviral therapy, issues surrounding care for HIV and other co-morbidities, interactions and impact on clinical outcomes. Development of novel electronic tools, including devices and computer programs to enhance behaviors, such as treatment adherence or uptake of treatment guidelines, is also of interest.

Contact: Lori Zimand
Telephone: 240-627-3212
Email: lzimand@niaid.nih.gov

B. Basic Research Branch. Innovative technologies for detection of acute HIV infection or HIV rebound following long-term suppression of viremia. Identification and validation of new targets for discovery or design of strategies to prevent HIV transmission, inhibit replication, control viremia in the absence of antiretroviral drugs, or eradicate reservoirs of HIV that persist despite long-term antiretroviral therapy. Innovative approaches for monitoring changes in the size of the persistent HIV reservoir.

Contact: Dr. Karl Salzwedel
Telephone: 301-496-5332
Email: salzwedelkd@niaid.nih.gov

C. Targeted Interventions Branch. Discovery and development of small molecule inhibitors with novel or underexplored mechanisms of action using standard and high-throughput technologies; cell-based and gene therapies; RNA-based therapeutics; next-generation biologics; novel targeting and delivery vehicles for agents active against HIV; therapeutic vaccines and monoclonal antibodies; protein chemistry-based anti-HIV approaches; assays to quantitate latent virus; animal models to facilitate evaluation of agents to treat or cure HIV infection.
Contact: Dr. Brigitte Sanders  
Telephone: 240-627-3209  
E-mail: sandersbe@niaid.nih.gov

**Vaccine Research Program**

Supports the discovery, development and clinical evaluation of an HIV/AIDS vaccine.

Director: Dr. Mary Marovich  
Telephone: 301-435-3727  
Email: mary.marovich@nih.gov

A. **Vaccine Clinical Research Branch (VCRB).** Research areas: (1) phase I, II, and III domestic and international clinical trials of candidate AIDS vaccines; (2) evaluation and characterization of immune responses in HIV-infected and uninfected immunized volunteers, using micro and macro assays; and (3) studies to identify, validate, and standardize immunologic and virologic markers for monitoring response of participants in vaccine clinical trials.

Contact: Dr. Jim Lane  
Telephone: 240-627-3033  
Email: laneji@mail.nih.gov

B. **Preclinical Research and Development Branch (PRDB).** Preclinical research and development of candidate AIDS vaccines, delivery methods, novel vaccine vectors, and adjuvants for the prevention of AIDS; promotion and evaluation of safety and efficacy of the prevention modalities, especially novel vaccine concepts identified in preclinical models including studies using non-human primates, humanized mouse, and other animal models; genetic and immunologic variation studies in relation to AIDS vaccine development; and mucosal and innate immunity in SIV, HIV, and SHIV models.

Contact: Dr. Anjali Singh  
Telephone: 240-627-3030  
Email: anjalisingh@niaid.nih.gov

C. **Vaccine Translational Research Branch (VTRB).** VTRB enables, facilitates, and operationalizes HIV vaccine translational research by advancing innovative vaccine concepts and scalable unit operations into the development of cGMP manufactured, vialled products for clinical testing. VTRB’s efforts to accelerate the development of preventive HIV-1 vaccines involves identifying, supporting and advancing: (a) cell line development to increase Env expression, production, quality, and yield; (b) evaluation of phase-appropriate upstream and downstream manufacturing processes; (c) scalable and prototype process development and purification platforms for viral vectors and HIV-1 Env proteins; (d) cGMP manufacturing of broad portfolio of vaccine products ranging from complex HIV Env protein immunogens, nanoparticle-based vaccines, viral vectors, virus-like particles (VLP), DNA and mRNA vaccines, monoclonal antibodies (neutralizing and non-neutralizing) for testing in early phase human clinical trials; (e) procuring and manufacturing new and/or alternative adjuvant analogs with similar agonist functions as those currently available for optimal immune response; (f) novel and emerging nanoparticle antigen and adjuvant delivery modalities and dosage forms, coformulation technologies and platforms for immunization; (g) antigen-adjuvant formulation development, analytics development to support product characterization, in-process operations, release, and stability testing; and (h) preclinical safety and toxicology testing.

Contact: Dr. Michael Pensiero  
Telephone: 301-435-3749  
Email: mpensiero@niaid.nih.gov
Therapeutics Research Program

Develops and oversees research and development of therapies for HIV disease, including complications, co-infections and co-morbidities, in adults.

Director: Dr. Peter Kim
Telephone: 301-451-2761
Email: peter.kim2@nih.gov

A. Drug Development and Clinical Sciences Branch. Preclinical development of experimental therapies for HIV, TB and other HIV/AIDS-related infectious diseases; including long-acting/extended release approaches; maintenance of a database of potential anti-HIV and anti-opportunistic infection compounds; management of quality assurance contracts for oversight of the quality of clinical laboratory testing in support of clinical trials; immunologic, virologic, and pharmacologic research related to the design and conduct of clinical trials; development and evaluation of practical and affordable tests to measure viral load, drug toxicities, and drug resistance to monitor populations in resource-poor settings; development of tests to detect early infection in seropositive HIV-infected adult and pediatric individuals.

Contact: Dr. Joe Fitzgibbon
Telephone: 240-627-3088
Email: jfitzgibbon@niaid.nih.gov

B. HIV Research Branch. Clinical research of treatments for acute and chronic HIV infection and approaches to achieve sustained remission or cure; strategies to augment HIV-specific immune responses, general host immunity to control or clear HIV infection, and prevention of HIV disease-associated end organ disease.

Contact: Tia Morton
Telephone: 240-627-3073
Email: frazierti@niaid.nih.gov


Contact: Dr. Chris Lambros
Telephone: 240-627-3093
Email: clambros@niaid.nih.gov

D. For evaluation of therapeutic agents or diagnostics for hepatitis B in HIV infected in adults.

Contact: Dr. Chris Lambros
Telephone: 240-627-3093
Email: clambros@niaid.nih.gov

E. Tuberculosis Clinical Research Branch. Translational and clinical research for tuberculosis, with and without HIV co-infection, to facilitate the development of biomarkers/diagnostics, therapies, and prevention/vaccines.

Contact: Dr. Richard Hafner
Telephone: 301-435-3766
Email: rhafner@niaid.nih.gov

Prevention Science Program

Supports basic research on mechanisms of HIV transmission supportive of new biomedical strategies for interrupting transmission. Supports domestic and international phase I, II, and III clinical trials to evaluate
HIV/AIDS prevention strategies, including microbicides, chemoprophylactic agents, and other biomedical and behavioral risk reduction interventions.

Director: Dr. Sheryl Zwerski
Telephone: 301-402-4032
Email: szwerski@niaid.nih.gov

A. **Preclinical Microbicides and Prevention Research Branch.** Development of non-vaccine biomedical HIV prevention products including topical microbicides, pre-exposure prophylaxis (PrEP), post-exposure prophylaxis (PEP), and multipurpose prevention technologies (MPT). Emphasis on drug delivery systems (DDS) designed to achieve systemic protection for ≥ 1 month. Development of shorter duration products, which address a compelling specific public health need. Key populations are adolescents, cisgender women, men who have sex with men (MSM), and transgender people.

Contact: Dr. James E. Cummins
Telephone: 240-292-4800
Email: cumminsje@niaid.nih.gov

B. **Clinical Microbicide Research Branch.** Clinical development of promising microbicides to prevent HIV infection with the ultimate goal to advance safe, effective, and acceptable microbicide products toward licensure.

Contact: Dr. Roberta Black
Telephone: 301-496-8199
Email: rblack@niaid.nih.gov

C. **Clinical Prevention Research Branch.** Development of safe and effective non-vaccine biomedical and integrated HIV prevention interventions to reduce the number of new HIV infections in adults and adolescents. Support the development of HIV incidence assays, biomarkers of adherence, mathematical modeling, and other tools needed to accomplish these objectives.

Contact: Dr. David Burns
Telephone: 301-435-8896
Email: burnsda@niaid.nih.gov

D. **Maternal, Adolescent and Pediatric Medicine Branch.** Therapies for cure, management, treatment and prevention of HIV and HIV-associated complications in pregnant women, infants, children, and adolescents. Strategies to reduce transmission of HIV and HIV co-infections from mother to child.

Contact: Judi Miller
Telephone: 240-292-4801
Email: jmillera@niaid.nih.gov

**Division of Allergy, Immunology, and Transplantation (DAIT)**

The Division of Allergy, Immunology, and Transplantation (DAIT) supports studies of the immune system in health and the cause, pathogenesis, diagnosis, prevention, and treatment of disease caused by immune dysfunction.

Director: Daniel Rotrosen, M.D.
Telephone: 301-496-1886
Email: drotrosen@niaid.nih.gov

A. **Allergy, Asthma and Airway Biology Branch.** Conditions of interest: asthma, food allergy, eosinophilic esophagitis and gastroenteritis in relation to food allergy, atopic dermatitis, urticaria, rhinitis, rhinosinusitis, drug allergy, sepsis. The Branch supports basic and clinical studies investigating mechanisms of disease and new approaches to diagnose, treat or...
prevent these conditions. Special interest for SBIR/STTR includes a) the development of biomarkers as diagnostic markers, markers of disease severity and predictive markers for treatment effectiveness, particularly of immunologic interventions such as allergen immunotherapy for food and respiratory allergy; b) the development of new forms of allergen immunotherapy aiming at increased tolerogenic immune responses and decreased allergenicity.

Chief: Alkis Togias, M.D.
Telephone: 301-496-8973
Email: togiasa@niaid.nih.gov

B. **Basic Immunology Branch.** The Branch supports basic and clinical research in the following areas: adjuvant discovery and development; origin, maturation, and interactions of immune cells; immune cell receptors, and ligands; cytokine biology; molecular basis of immune activation, antigen recognition, and immune tolerance; immune response regulation; hematopoiesis and stem cell biology; computational immunology; immunologic mechanisms associated with Myalgic Encephalomyelitis/Chronic Fatigue Syndrome; enhancement of vaccine effectiveness in neonates and adults, and basic immunology of vaccines and immunotherapeutics as medical countermeasures for biodefense. Special interests for SBIRs include: adjuvant discovery and/or development; bioinformatics tools for immune epitope predictions/visualization, and/or for the analysis of multi-parameter or systems immunology data; development of immunologic reagents for analysis of immunity in non-mammalian (e.g., frogs, fish, C. elegans) and under-represented mammalian (e.g., pig, ferret) models, and development of novel/improved methods to analyze human immune responses from limited amounts of human sample (tissue, cells, serum, etc.).

Chief: Dr. Alison Deckhut-Augustine
Telephone: 301-496-7551, Fax: 301-480-2381
Email: augustine@niaid.nih.gov

C. **Autoimmunity and Mucosal Immunology Branch.** Preclinical and clinical research to develop and improve therapies for the treatment of autoimmune diseases and primary immune deficiencies (not HIV); basic research of autoimmune disease mechanisms and biomarkers; immunotherapy of disease processes; disorders mediated by lymphocyte products; and mucosal immunity.

Chief: Dr. James McNamara
Telephone: 301-451-3121, Fax: 301-480-1450
Email: jmcnamara@niaid.nih.gov

D. **Transplantation Branch.** Preclinical and clinical research in organ, vascularized composite tissue and cellular transplantation: acute and chronic graft rejection, allogeneic and xenogeneic transplantation, development of immunomodulatory agents to prevent and treat graft rejection and to promote acute and long term graft acceptance and immunologic tolerance, genomics of the alloimmune response, graft versus host disease for hematopoietic stem cell transplantation, minor histocompatibility antigens, complications of immunosuppression in transplantation, and major histocompatibility complex (MHC) region genomics, technologies for MHC typing, and clinical applications of high-resolution HLA typing.

Chief: Nancy D. Bridges
Telephone: 301-496-5598
Email: nbridges@niaid.nih.gov

E. **Radiation Countermeasures Program.** The Radiation and Nuclear Countermeasures Program will consider preclinical research to support specific product development activities leading to creation of IND or IDE packages to be submitted to FDA. These IND/IDE-enabling activities could include: efficacy studies to optimize formulation, dose, and dose schedule; drug product stability studies, drug product GMP manufacturing scale-up, GLP toxicology and pharmacology safety studies, pharmacokinetic and metabolism studies, development of GLP analytical methods for efficacy studies and product characterization (including using chip technology to determine tissue-specific efficacy of...
a lead drug candidate), mechanism of action studies and completion of IND or IDE package for FDA submission. Product development efforts will advance new medical countermeasures towards Phase I clinical safety studies, GLP animal pivotal efficacy studies, and licensure/approval/clearance by the FDA.

Acting Chief: Charles Hackett, Ph.D., Associate Director
Telephone: 301-496-1886
Email: CHackett@niaid.nih.gov

Division of Microbiology and Infectious Diseases (DMID)

The Division of Microbiology and Infectious Diseases (DMID) supports research to better understand, treat, and ultimately prevent infectious diseases caused by virtually all infectious agents, except HIV. DMID supports a broad spectrum of research from basic molecular structure, microbial physiology and pathogenesis, to the development of new and improved vaccines and therapeutics. DMID also supports medical diagnostics research, which is defined as research to improve the quality of patient assessment and care that would result in the implementation of appropriate therapeutic or preventive measures. DMID does not support research directed at decontamination or the development of environmentally oriented detectors, whose primary purpose is the identification of specific agents in the environment. Note that some of the organisms and toxins listed below are considered NIAID priority pathogens or toxins for biodefense and emerging infectious disease research.

Division Small Business Representative: Dr. Barbara Mulach
Telephone: 240-627-3322
Email: bmulach@niaid.nih.gov

Division Small Business Representative: Dr. Alyssa Werner
Telephone: 301-761-7525
Email: alyssa.werner@nih.gov

A. **Bacteriology and Mycology Branch.**

The branch oversees research and product development related to:

- **Bacterial infections with emphasis on hospital-related infections**, including Acinetobacter, Klebsiella, Serratia, Legionella, Pseudomonas, Aeromonas, Enterobacter, Proteus, non-enteric E. coli, staphylococci, enterococci, actinomycetes among others;
- **Bacterial zoonoses**, including plague, anthrax, tularemia, glanders, melioidosis, Lyme disease, rickettsial diseases, anaplasmosis, ehrlichiosis, and Q fever, and leptospirosis;
- **Fungal infections** including Candida, Aspergillus, Cryptococcus, Coccidiodes, Histoplasma, Blastomyces, Pneumocystis, Microsporidia, and other pathogenic fungi.

Research is encouraged in the following general areas: (1) vaccines, adjuvants, therapeutics and diagnostics (including target identification and characterization, device or apparatus development, novel delivery, and preclinical evaluation); (2) strategies to combat antibacterial and antifungal drug resistance; (3) applied proteomics and genomics; (4) host-pathogen interactions, including pathogenesis and host response; (5) genetics, molecular, and cell biology; and (6) microbial structure and function.

Research on all of the above is welcome, but the following areas are of particular interest to the branch:

- Vaccines, therapeutics, and medical diagnostics for hospital infections
- Adjunctive therapies and non-traditional approaches to combat and treat antimicrobial resistance
• Diagnostics for invasive fungal diseases
• Novel approaches for the diagnosis of Lyme disease
• Vaccines against Coccidioidomycosis

Contact: Dr. Alec Ritchie  
Telephone: 240-627-3356  
Email: aritchie@niaid.nih.gov

B. Enteric and Sexually Transmitted Infections Branch.

Enterics Section:
Research portfolios focus on enteric bacterial pathogens such as *Campylobacter* spp., *Clostridia* spp., pathogenic *Escherichia coli*, *Helicobacter* spp., *Listeria* spp., *Vibrio* spp., *Salmonella* serovars, *Shigella* spp., enteric *Yersinia* spp., and *Bacteroides* spp. (enterotoxins only). Also within the branch portfolio are toxins such as ricin toxin, *Staphylococcus* enterotoxin B, and botulinum neurotoxins; gastrointestinal diseases associated with diarrhea, dysbiosis; and the gastrointestinal microbiota and microbiome.

Special emphasis areas include:
• Development of pediatric vaccines to prevent the major worldwide causes of diarrhea; vaccines against neurotoxins and enterotoxins, and vaccines for enteric diseases where waning immunity is an issue (i.e., *C. difficile* and *Salmonella* infection).
• Development of antimicrobials and antivirals that focus on novel targets such as host-pathogen interactions to combat the development of resistance; therapies that target toxins once they enter cells; and therapies to treat recurrent diseases.
• Development of simple, rapid point-of-care diagnostic tools for the simultaneous identification of multiple pathogens that includes their antimicrobial resistance profiles; diagnostics for use in low-resource settings, especially for typhoid; and novel diagnostics that differentiate *C. difficile* colonization from infection.

Sexually Transmitted Infections Section:
Areas of emphasis include the development of medical diagnostics including better and more rapid multiplex point of care tests, ability to rapidly determine antibiotic sensitivity, and novel technologies enabling testing in low resource settings while maintaining high sensitivity/specificity; development of new classes of antimicrobials and non-antimicrobial treatment approaches, particularly those focused on reducing the development of antibiotic resistance; novel delivery systems for multipurpose prevention technologies, vaccines and therapeutics for Sexually Transmitted Infections (STIs) and other reproductive tract syndromes such as bacterial vaginosis and pelvic inflammatory disease; understanding vaginal ecology and immunology and approaches to developing synthetic microbiota for use as biotherapeutics or as adjunct therapy to antibiotic treatment; development of epidemiologic and behavioral strategies to reduce transmission of STIs; developing and evaluating interventions and products to better serve adolescents, medically underserved populations, and minority groups who are disproportionately affected by STIs; development of multipurpose prevention technologies to prevent STIs, HIV, and unintended pregnancies; better understanding of the role of STIs in infertility, premature birth, and adverse outcomes of pregnancy and how to improve outcomes; and better understanding of the role of STIs in HIV transmission and the role of HIV in altering the natural history of STIs.

Contact: Dr. Tom Hiltke  
Telephone: 240-627-3275  
Email: thiltke@niaid.nih.gov
C. Parasitology and International Programs Branch.

Research areas: (1) protozoan infections, including amebiasis, cryptosporidiosis, cyclosporiasis, giardiasis, leishmaniasis, malaria, trypanosomiasis, toxoplasmosis; helminth infections, including cysticercosis, echinococcosis, lymphatic filariasis, schistosomiasis, onchocerciasis, others (e.g., roundworms, tapeworms, and flukes); invertebrate vectors/ectoparasites, black flies, sandflies, tsetse flies, mosquitoes, ticks, snails, and mites; (2) parasite biology (genetics, genomics, physiology, molecular biology, and biochemistry); (3) protective immunity, immunopathogenesis, and evasion of host defense; (4) clinical, epidemiological, and natural history studies of parasitic diseases; (5) research and development of vaccines, drugs, immunotherapeutics and immunoprophylaxis, and medical diagnostics; and (6) vector biology and management, mechanisms of pathogen transmissions.

Research on the above is welcome, but research on the following is of particular interest to the branch:

- New drug discovery or re-purposing of existing drugs to treat parasitic diseases
- Highly sensitive diagnostics tools for parasitic diseases
- Vaccines, monoclonal antibodies, and vaccine technologies applicable to prevention or elimination of parasitic diseases
- Technologies or approaches that address arthropod vector monitoring, management, and control, to prevent transmission of vector-borne pathogens to humans

Contact: Dr. Annie Mo
Telephone: 240-627-3320
Email: moa@niaid.nih.gov

D. Respiratory Diseases Branch.

Research areas include: (1) viral respiratory diseases caused by influenza viruses, human coronaviruses including SARS, MERS, and novel emerging coronaviruses, rhinoviruses, respiratory syncytial virus and other related paramyxoviruses; (2) mycobacterial diseases, including tuberculosis (TB) caused by bacteria of the Mycobacterium tuberculosis complex, leprosy, Buruli ulcer and non-tuberculous mycobacterial (NTM) diseases, particularly pulmonary infections in persons not afflicted with HIV/AIDS; (3) other bacterial respiratory diseases including bacterial pneumonia, pertussis, Group A and B streptococcal diseases, meningitis, upper respiratory infections, acute exacerbations of chronic obstructive pulmonary disease cystic fibrosis; and (4) mixed viral/bacterial respiratory infections.

Special emphasis areas include:

- Development of new or improved antimicrobials (especially for antimicrobial-resistant pathogens) and antivirals, including immunotherapeutics, immunomodulators, and host-directed therapies to augment anti-infectives;
- Methodologies for rapid, point-of-care evaluation of drug levels in TB patients to facilitate therapeutic drug monitoring;
- New or improved vaccines (with and without adjuvants);
- Improved delivery systems and formulations for drugs/vaccines;
- Microbial and host biomarkers and biosignatures suitable for diagnostic tests;
- Rapid multiplex diagnostic tests, including low cost point-of-care, or other tools to detect infection prior to active disease and to identify drug resistance;
- Diagnostics to distinguish viral from bacterial infections.
There is particular need for preventive and treatment countermeasures for influenza, including universal vaccine platforms and broad-spectrum antivirals; for diagnostics (including drug susceptibility tests), novel therapeutics, and vaccines (including adjuvants) against *Mycobacterium tuberculosis* (TB); for novel TB diagnostics; particularly diagnostics suitable for diagnosis in pediatric populations and rapid drug susceptibility testing; for relevant diagnostics, preventive and curative interventions against non-HIV associated pulmonary Non-tuberculous mycobacteria (NTM); and for the prevention, diagnosis, and treatment of *Bordetella pertussis*, *Group A streptococcus*, and *Streptococcus pneumoniae* infections and other antibacterial resistant infections.

Contact: Dr. Xin-Xing Gu  
Telephone: 240-627-3265  
Email: guxx@niaid.nih.gov

E. **Virology Branch.**

The Virology Branch focuses on:

a. Acute viral infections: arthropod-borne (ex.: tick-borne) and rodent-borne viral diseases (including Dengue, Zika, West Nile, Japanese encephalitis, Chikungunya, yellow fever, hantavirus, Crimean-Congo Hemorrhagic fever, severe fever with thrombocytopenia syndrome (SFTS), Heartland virus, Bourbon virus, Tick-borne Encephalitis virus (TBE), Powassan virus, LaCrosse, Rift Valley Fever virus, etc.), viral hemorrhagic fevers (Ebola, Lassa fever, etc.), Nipah, Hendra, measles, polio, coxsackie virus, enteroviruses, poxviruses, rabies, rubella, Astroviruses, Caliciviruses, Rotavirus;

b. Persistent viral infections: adenoviruses, BK virus, bornaviruses, coronaviruses, herpesviruses, human T-lymphotrophic virus, JC virus, human papillomaviruses, parvoviruses, emerging human polyomaviruses;

c. Acute infections with hepatitis viruses A, B, C, D and E (HAV, HBV, HCV, HDV, and HEV); chronic infections with hepatitis viruses, B, C, D and E;

d. Transmissible Spongiform Encephalopathies (TSE).

Areas of emphasis for SBIR/STTR applications include:

- Development of vaccines;
- Development of techniques to improve vaccine stability;
- Approaches to identify antiviral targets and agents;
- Chemical design and synthesis of novel antiviral agents;
- Development of therapeutic interventions;
- Development and validation of point of care assays for disease diagnosis and to measure response to therapy;
- Development of new preclinical animal model systems that predict clinical efficacy of vaccines, therapeutics and diagnostics.

The Virology Branch does not support applications covering environmental detection and decontamination.

Contact: Dr. Mindy Davis  
Telephone: 301-761-6689  
Email: mindy.davis@nih.gov

**NIAID Clinical Trial Topics:**

NIAID will generally consider clinical trial proposals consistent with the topics listed above. However, applicants are strongly encouraged to consult with NIAID Program Staff at least 10 weeks before the receipt date.
For further information, please consult NIAID’s Investigator-Initiated Clinical Trial Resources page:
https://www.niaid.nih.gov/grants-contracts/investigator-initiated-clinical-trial-resources

For more information on NIAID’s SBIR/STTR research topics, program policy or to identify NIAID Subject Matter Experts for a specific topic, please contact:

Dr. Natalia Kruchinin
SBIR/STTR Program Coordinator
Division of Extramural Activities
National Institute of Allergy and Infectious Diseases
Phone: 240-669-2919
Email: kruchininn@niaid.nih.gov

General Inquiries and subscription requests for NIAID’s SBIR/STTR List-Serv: NIAIDSBIR@mail.nih.gov
NATIONAL INSTITUTE OF ARTHRITIS AND MUSCULOSKELETAL AND SKIN DISEASES (NIAMS)

The mission of the National Institute of Arthritis and Musculoskeletal and Skin Diseases is to support research into the causes, treatment, and prevention of arthritis and musculoskeletal and skin diseases, the training of basic and clinical scientists to carry out this research, and the dissemination of information on research progress in these diseases.

For additional information about areas of interest to the NIAMS, please visit NIAMS Strategic Plan at [https://www.niams.nih.gov/about-niams/strategic-plan-fiscal-years-2020-2024](https://www.niams.nih.gov/about-niams/strategic-plan-fiscal-years-2020-2024).

Limited Amount of Award

For budgetary, administrative, or programmatic reasons, NIAMS may decide not to fund an application or may decrease the length of an award and/or the budget recommended by a review committee. Generally, NIAMS does not fund Phase I applications with a total cost greater than $256,580 or a project period greater than 2 years and Phase II applications with a total cost greater than $1,710,531 or a project period greater than 3 years. Applicants considering a requested budget greater than these limits are strongly encouraged to contact program staff before submitting an application. It is not the intent of NIAMS to support clinical trials through the SBIR/STTR mechanism. Applicants who wish to submit clinical trials applications to the NIAMS are encouraged to utilize one of the NIAMS FOAs listed at [https://www.niams.nih.gov/grants-funding/conducting-clinical-research/investigator-clinical-trial-policies](https://www.niams.nih.gov/grants-funding/conducting-clinical-research/investigator-clinical-trial-policies).

Research Topics of Interest to NIAMS

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
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*If No,
- This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, UNLESS the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,
- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.
NIAMS Non-Clinical Trials Topics:

The NIAMS small business program supports research and development of products and services for prevention, diagnosis and treatment of rheumatic, musculoskeletal and skin diseases. The research topics include, but are not limited to, the following:

A. **Rheumatic Diseases.** The NIAMS supports research on rheumatic and related diseases including rheumatoid arthritis (RA), juvenile idiopathic arthritis (JIA), Lyme arthritis, viral arthritis, gout, calcium pyrophosphate deposition disease (CPDD), spondyloarthropathies, and systemic autoimmune diseases such as systemic lupus erythematosus (SLE), systemic scleroderma (SSc), and autoimmune myositis.

B. **Musculoskeletal Diseases.** The musculoskeletal system is composed of the skeleton, the muscles, and connective tissues such as cartilage, tendon, and ligament. The NIAMS supports research aimed at improving the diagnosis, treatment, and prevention of diseases and injuries of the musculoskeletal system and its component tissues. The topics in this area include research on musculoskeletal diseases such as osteoporosis, osteoarthritis, muscular dystrophy, and osteogenesis imperfecta, tissue engineered products, orthopedic devices and implants, and sports medicine and fitness.

C. **Skin Diseases.** The NIAMS supports research on a wide range of skin diseases and conditions including chronic inflammatory skin diseases such as psoriasis, rosacea, acne vulgaris, and atopic dermatitis and autoimmune diseases such as pemphigus, vitiligo, and alopecia areata. The NIAMS also supports research on skin repair and regeneration in treatment of chronic wounds and reducing scar formation. Skin cancer is an area of overlap with the National Cancer Institute (NCI), with the NIAMS focus on the response of keratinocytes to UV light and early stages in the development of non-melanoma skin cancer and products for prevention of melanocyte tumorigenesis.
This is not an inclusive list of all research topics covered by the NIAMS. To learn more, please visit the NIAMS supported scientific areas at https://www.niams.nih.gov/grants-funding/funding-opportunities/supported-scientific-areas

Special Emphasis Areas of Interest to Small Businesses:

NIAMS supports all Research and Development activities within its mission. Particular areas of programmatic interest relative to small business initiatives include, but are not limited to:

A. Innovation research on rare musculoskeletal, rheumatic and skin diseases
B. Multiplex assay development for arthritis and musculoskeletal and skin diseases
C. Lab to marketplace: translation of scientific discoveries in NIAMS mission areas from labs into products on the market
D. Test and/or validation of novel, state-of-the-art candidate biomarker platforms for predicting the onset and progression of inflammatory diseases of interest to the NIAMS and for determining the pharmacodynamics, safety and/or efficacy of therapeutic agents targeting those diseases.

NIAMS Clinical Trials Topics:

NIAMS does not participate in the SBIR/STTR clinical trial funding opportunities. NIAMS NON-SBIR/STTR clinical trial funding opportunities support all research within the NIAMS mission areas.

For general SBIR/STTR program information, contact:

Dr. Xibin Wang, NIAMS SBIR/STTR Coordinator
Telephone: 301-451-3884, Fax: 301-480-1284
Email: wangx1@mail.nih.gov

For administrative and business management questions, contact:

Ms. Victoria Matthews
Telephone: 301-594-3968, Fax: 301-480-5450
Email: victoria.matthews@nih.gov
NATIONAL INSTITUTE OF BIOMEDICAL IMAGING AND BIOENGINEERING (NIBIB)

The mission of the National Institute of Biomedical Imaging and Bioengineering (NIBIB) is to improve health by leading the development and accelerating the application of biomedical technologies. The Institute is committed to integrating the physical and engineering sciences with the life sciences to advance basic research and medical care. This is achieved through: research and development of new biomedical imaging and bioengineering techniques and devices to fundamentally improve the detection, treatment, and prevention of disease; enhancing existing imaging and bioengineering modalities; supporting related research in the physical and mathematical sciences; encouraging research and development in multidisciplinary areas; supporting studies to assess the effectiveness and outcomes of new biologics, materials, processes, devices, and procedures; developing technologies for early disease detection and assessment of health status; and developing advanced imaging and engineering techniques for conducting biomedical research at multiple scales.

Research Topics of Interest to NIBIB

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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| Clinical Trials SBIR/STTR Omnibus/Parent Funding Opportunity Announcement/s | | X | If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information.
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<tr>
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NIBIB Non-Clinical Trials Topics:

A. **Artificial Intelligence, Machine Learning, and Deep Learning.** Design and development of intelligent and innovative algorithms, software, methods, and computational tools to enhance analysis of complex medical images and data. Relevant technologies include those that facilitate organization, representation, retrieval, analysis, recognition, and classification of biomedical and biological data and images. Unsupervised and semi-supervised techniques and methodologies are of particular interest.

B. **Image Processing, Visual Perception, and Display.** Design and development of algorithms for post-acquisition image processing and analysis. These algorithms include methods for image segmentation, image registration, atlas generation, morphometry measurement, and the determination of function and structure from medical images. Also supported by this program is the development of theoretical models and analysis tools to evaluate and improve the perception of medical images. This may include diagnostic-performance evaluation, assessment of computer-aided diagnosis technologies, statistical models for evaluation of observer performance, and assessment of observer variability. Finally, this program supports the development of visualization tools for improved detection.

C. **Biomedical Informatics.** Development of structures and algorithms to improve the collection, annotation, aggregation, anonymization, classification, retrieval, integration, analysis, and dissemination of quantitative and qualitative biomedical data. Examples of informatics tools and resources supported by this program are: biostatistics methods for bioinformatics, meta databases and integrative services, digital biomarkers, information-driven computer-aided diagnosis and decision support systems, digital atlases, data mining, large scale biomedical image/information databases, data fusion, and hyperspectral data analysis and -omics. This program is intended to support NIBIB’s other program areas in biomedical imaging and bioengineering researchers.

D. **Point of Care Technologies-Diagnostics.** Development of rapid in-vitro diagnostic technologies and monitoring platforms that provide real time medical evaluation and analysis of the disease status or condition at the time and place of patient care. Technology development area examples within the program include but are not limited to disposable lateral flow assays, nucleic acid testing platforms, glucose monitoring devices, etc. The program includes the delivery of healthcare that is safe, effective, timely, patient centered, efficient, and available in centralized and decentralized locations.
E. **Connected Health-Mobile Health and Telehealth.** Development of enabling technologies that emphasize the integration of wireless technologies with human and biological interfaces. This program includes the development of software and hardware for telehealth and mobile health studies. This program includes the input and delivery of healthcare information digitally for the analysis or monitoring of health or disease status. The emphasis is on developing mobile health technologies driven by clinical needs and integrating these technologies in healthcare delivery, wellness, and daily living.

F. **Bio-Electromagnetic Technologies.** Development of technologies that use static or dynamic electromagnetic fields for sensing, imaging, or therapeutic effects. The emphasis is on increasing the sensitivity, spatial/temporal resolution, efficacy, or safety of bioelectromagnetic devices through the development of novel hardware, method of operation, or pre-/post-processing techniques for single modalities or the combination of multiple modalities. This program may support the development of magnetic particle imaging, electrical impedance tomography, electroencephalography, magnetoencephalography, electromagnetic-field-induced hyperthermia/ablation, and microwave/terahertz imaging, for example.

G. **Magnetic Resonance Imaging.** Development of in vivo MR imaging and MR spectroscopy, for both animal and human research and potential clinical applications. The emphasis is on the development of MRI hardware and methodologies, including image acquisition and reconstruction techniques, that would improve the speed, spatial resolution, information content, efficiency, robustness, quality, patient experience, and safety. The emphasis should be on technological development rather than detailed applications to specific diseases or organs.

H. **Optical Imaging and Spectroscopy.** Development and application of optical imaging, microscopy, and spectroscopy techniques for improving disease prevention, diagnosis, and treatment in the medical office, at the bedside, or in the operating room. Examples of research areas include fluorescence imaging, bioluminescence imaging, OCT, SHG, IR imaging, diffuse optical tomography, optical microscopy and spectroscopy, confocal microscopy, and multiphoton microscopy. The emphasis is on development of cost effective, portable, safe, and non-invasive or minimally invasive devices, systems, and technologies.

I. **Bioanalytical Sensors.** Engineering the components and functionality of bioanalytical sensors. Detection could be based on optical, chemical, electrochemical, and/or physical (such as mechanical, gravimetric, thermal) perturbation of a sample, for example. Examples of technologies of interest include, but are not limited to, nano-textured substrates for analyte detection, DNA sensors for liquid biopsy, and small molecule detectors for diagnosing infectious diseases.

J. **Molecular Probes and Imaging Agents.** Development and biomedical application of molecular probes and imaging agents across all imaging modalities for the visualization, characterization and quantification of normal biological and pathophysiological processes and anatomy in living organisms at the molecular, cellular and organ levels. The emphasis is on engineering of targeting and responsive molecular probes of high sensitivity and specificity for PET and SPECT (radiotracers), MR (T1, T2, CEST, hyperpolarized agents), EPR, CT, optical (fluorescent and bioluminescent probes), ultrasound (microbubbles) and photoacoustic imaging.

K. **Ultrasound: Diagnostic and Interventional.** Improvement of technologies for diagnostic, interventional and therapeutic uses of ultrasound. The diagnostic ultrasound program includes, but is not limited to the design, development and construction of transducers, transducer arrays, and transducer materials, innovative image acquisition and display methods, innovative signal processing methods and devices, and optoacoustic and thermoacoustic technology. It also includes the development of image-enhancement devices and methods, such as contrast agents, image and data presentation and mapping methods, such as functional imaging and image fusion. The therapeutic ultrasound program includes, but is not limited to the design, development, and construction of transducers, transducer arrays, interventional technologies, adjunct enhancement of non-ultrasound therapy applications, high-intensity focused ultrasound (HIFU), or hyperthermia applications. It also
includes non-invasive or minimally invasive interventional surgical or therapy tools, ultrasound contrast agents for therapy, targeted drug delivery, neuromodulation, and biopsy.

L. **Image-Guided Interventions.** Development of novel image-directed technologies for guidance, navigation, tissue differentiation, and disease identification for reaching specified targets during therapeutic procedures, which may range along the continuum from non-invasive to minimally invasive to open surgical interventions. These technologies may range from molecular to macroscopic scale levels. In addition, emphasis includes technologies that expand needed procedural access for individuals otherwise excluded by disease characteristics, co-morbidities, and other parameters.

M. **Nuclear Medicine.** Research and development of technologies that create images out of the gamma-ray or positron emissions from radioactive agents that are injected, inhaled, or ingested into the body. The emphasis is on: simulation and development of new detectors, collimators, and readout methods that enhance the signal quality of detecting isotope emissions; designs of novel camera geometries; and correction methods that compensate for the radiation physics properties to improve the clinical reliability of the image. Of interest are improvements and corrections for interaction events in PET detectors and enhancement to time of flight (TOF) image generation methods (reconstructions algorithms); as well as new collimator and camera designs for SPECT.

N. **X-ray, Electron, and Ion Beam.** Simulation, design and development of new detector systems; new readout methods that enhance the signal quality for x-ray image generation; designs of novel imaging geometries; algorithms that compensate for the physical properties of the detection system to improve the clinical reliability of the image (reconstruction algorithms); and approaches to radiation dose reduction, especially in CT. Of interest are diagnostic image enhancements via photon counting, dual energy, and new applications of cone-beam tomography.

O. **Biochemical Engineering.** Development and demonstration of new approaches to control/program biology for biomedical intervention, without preference for any particular disease or application. Emphasis in this program is on engineering new biochemical materials, sensors, actuators, and other parts and modules to interface and communicate with human biology and engineered systems for biomedical intervention. These parts and modules act as biotransducers to convert chemical energy into biological action. Projects should be directed toward overcoming a technological challenge that limits biomedical adoption. This program encourages projects that use a design-build-test approach.

P. **Bioelectric Engineering.** Development and demonstration of new approaches to control/program biology for biomedical intervention, without preference for any particular disease or application. Emphasis in this program is on engineering new bioelectric materials, sensors, actuators, and other parts and modules to interface and communicate with human biology and engineered systems for biomedical intervention. These parts and modules act as biotransducers to convert electric energy to biological action. Projects should be directed toward overcoming a technological challenge that limits biomedical adoption. This program encourages projects that use a design-build-test approach.

Q. **Biomechanical Engineering.** Development and demonstration of new approaches to control/program biology for biomedical intervention, without preference for any particular disease or application. Emphasis in this program is on engineering new biomechanical materials, sensors, actuators, and other parts and modules to interface and communicate with human biology and engineered systems for biomedical intervention. These parts and modules act as biotransducers to convert mechanical energy into biological action. Projects should be directed toward overcoming a technological challenge that limits biomedical adoption. This program encourages projects that use a design-build-test approach.

R. **Bionic and Robotic Systems.** Development and demonstration of new approaches to control/program biology for biomedical intervention, without preference for any particular disease or application. Emphasis in this program is on engineering bionic and robotic systems to sense and
actuate in response to human biology for biomedical intervention. Projects should be directed toward overcoming a technological challenge that limits biomedical adoption. This program encourages projects that use a design-build-test approach.

S. **Biophotonic Engineering.** Development and demonstration of new approaches to control/program biology for biomedical intervention, without preference for any particular disease or application. Emphasis in this program is on engineering new biophotonic materials, sensors, actuators, and other parts and modules to interface and communicate with human biology and engineered systems for biomedical intervention. These parts and modules act as biotransducers to convert photonic energy to biological action. Projects should be directed toward overcoming a technological challenge that limits biomedical adoption. This program encourages projects that use a design-build-test approach.

T. **Mathematical Modeling, Simulation and Analysis.** Development of novel mathematical modeling, simulation and analysis tools that can be broadly applied across a wide spectrum of diagnostic, therapeutic, imaging, and interventional applications. Emphasis is on engineering solutions for theory-driven, physics-based, physiologically realistic, virtual representations of biomedical systems, with a particular weight on multiscale modeling. Interests include, but are not limited to: multiscale modeling, predictive modeling frameworks, non-standard methodologies, and methods to address model credibility, reproducibility, and reuse.

U. **Synthetic Biological and Biomimetic Systems.** Development and demonstration of new approaches to control/program biology for biomedical intervention, without preference for any particular disease or application. Emphasis in this program is on engineering biological and biomimetic systems to sense and actuate in response to human biology for biomedical intervention. Projects should be directed toward overcoming a technological challenge that limits biomedical adoption. This program encourages projects that use a design-build-test approach.

**NIBIB Clinical Trials Topics:**

NIBIB will accept clinical trials in any area listed above in the non-clinical trials topics.

For additional information on research topics, contact:

Mr. Todd Merchak  
National Institute of Biomedical Imaging and Bioengineering  
Telephone: 301-496-8592, Fax: 301-480-1614  
Email: NIBIB-SBIR@mail.nih.gov

Dr. Ilana Goldberg  
National Institute of Biomedical Imaging and Bioengineering  
Telephone: 301-402-3465 Fax: 301-480-1614  
Email: NIBIB-SBIR@mail.nih.gov

For administrative and business management questions, contact:

Mr. James Huff  
National Institute of Biomedical Imaging and Bioengineering  
Telephone: 301-451-4786, Fax: 301-451-5735  
Email: huffj@mail.nih.gov
**NATIONAL CANCER INSTITUTE (NCI)**

The National Cancer Institute’s SBIR Development Center program is one of the nation’s largest sources of financing for small businesses engaged in technology innovation. Its funding, mentoring and networking assistance is offered to small businesses demonstrating promising next-generation cancer cure technologies, with the ultimate goal being successful commercialization and life-changing public benefit. NCI’s SBIR/STTR Programs offer funding for therapeutic agents and devices; in vitro and in vivo diagnostics, including companion diagnostics and imaging agents; agents and technologies for cancer prevention; tools for research in cancer biology, cancer control, and epidemiology; digital health, including health information technology and bioinformatics; and many more areas of interest to the NCI.

The goal of NCI’s SBIR/STTR program is to increase small business participation and private-sector commercialization of novel technologies that can prevent, diagnose and treat cancer. The major NCI SBIR/STTR portfolio areas are listed below as a guide to general technology areas funded through the program. However, NCI will accept any applications outside these topic areas and proposing innovative cancer-related technologies with strong commercial potential is encouraged.

**Major NCI SBIR/STTR Portfolio Areas:**

- Therapeutics (e.g., Small Molecules, Biologics, Radiomodulators, and Cell-based Therapies)
- *In Vitro* and *In Vivo* Diagnostics (e.g., Companion Diagnostics and Prognostic Technologies)
- Imaging Technologies (e.g., Agents, Devices, and Image-Guided Interventions)
- Devices for Cancer Therapy (e.g., Interventional Devices, Surgical, and Radiation and Ablative Therapies, Hospital Devices)
- Agents and Technologies for Cancer Prevention
- Technologies for Cancer Control (e.g., Behavioral Health Interventions, Tools for Genetic, Epidemiologic, Behavioral, Social, and/or Surveillance Cancer Research)
- Tools for Cancer Biology Research
- Digital Health Tools and Software Platform for Cancer Related Technologies

NCI is committed to lessening the impact of cancer by propelling technological innovation. The NCI SBIR/STTR Programs are aligned with recommendations laid out by the Cancer Moonshot Blue Ribbon Panel Report: [https://www.cancer.gov/research/key-initiatives/moonshot-cancer-initiative/blue-ribbon-panel](https://www.cancer.gov/research/key-initiatives/moonshot-cancer-initiative/blue-ribbon-panel)

NCI SBIR program is technology agnostic and we welcome all innovative solutions with commercial potential that is relevant to the mission of the NCI and that reduces the burden on cancer patients, their caregivers and providers.

For up-to-date information on high priority technology areas, and to learn about programmatic initiatives and upcoming events, visit the NCI SBIR Development Center homepage: [http://sbir.cancer.gov/](http://sbir.cancer.gov/).

In addition, please see the contact list at the end of the NCI section to identify the NCI SBIR/STTR Program Director(s) that specializes in your technology area.

**Limited Amount of Award**

For budgetary, administrative, or programmatic reasons, NCI may decide not to fund an application or may decrease the length of an award and/or the budget recommended by a review committee. Generally, NCI will fund Phase I applications with budgets up to $256,580 total costs and project periods up to 2 years; Phase II applications with budgets up to $1,710,531 total costs combined over all years and with project periods up to 3 years will be considered. However, for certain research topic, the U.S. Small
Business Administration has approved an NIH SBIR/STTR Topic Waiver list for which the NCI generally will fund Phase I applications with higher budgets up to $400,000 total costs combined over all years, and project periods up to 2 years; similarly, for certain research topics NCI will consider Phase II applications with higher budgets up to $2,000,000 total costs combined over all years, and project periods up to 3 years. These NCI SBIR/STTR Waiver Topic areas can be found in the NCI Section of Appendix A below.

**Phase IIB Competing Renewal Awards**

The NCI does not accept applications for Phase IIB SBIR competing renewal awards through this Omnibus solicitation. However, the NCI offers Phase IIB opportunities in the form of the NCI SBIR Phase IIB Bridge Award, which is announced via a separate funding solicitation: [https://sbir.cancer.gov/bridge](https://sbir.cancer.gov/bridge). The NCI Phase IIB Bridge Award is designed to support the next stage of development for cancer-related technologies previously funded under SBIR or STTR Phase II awards from any Federal agency. The purpose of this award is to address the funding gap known as the "Valley of Death" between the end of the SBIR Phase II award and the subsequent round of financing needed to advance a product or service toward commercialization. To achieve this goal, the Bridge Award funding opportunity is specifically designed to incentivize partnerships between Federally-funded SBIR Phase II awardees and third-party investors and/or strategic partners. Competitive preference and funding priority will be given to applicants that demonstrate the ability to secure substantial independent third-party investor funds (i.e., third-party funds that equal or exceed the requested NCI funds).

To ensure that you will be notified upon the release of the NCI SBIR Phase IIB Bridge Award solicitation, please sign up for the NCI SBIR mailing list: [https://sbir.cancer.gov/emailsignup](https://sbir.cancer.gov/emailsignup). If you have any questions regarding the NCI SBIR Phase IIB Bridge Award, please contact your Phase II program director.

**Research Topics of Interest to NCI**

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
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*If No,
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**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,**
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<td>X</td>
<td></td>
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</tr>
<tr>
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<td>X</td>
<td></td>
<td><a href="https://www.cancer.gov/grants-training/grants-funding/funding-opportunities">R21, R01, P01, K08</a></td>
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</tbody>
</table>

**NCI Non-Clinical Trials Topics:**

NCI will accept applications in any of the NCI priority areas mentioned above or any other areas that are relevant to NCI’s mission.

**NCI Clinical Trials Topics:**

NCI will accept applications for support of clinical trials in the NCI priority area mentioned above or any other areas that are relevant to the NCI’s mission.

For additional information about the NCI SBIR/STTR programs, please contact the NCI SBIR Development Center:

National Cancer Institute
SBIR Development Center
9609 Medical Center Drive, Suite 1W550
Rockville, MD 20850
Website: [http://sbir.cancer.gov](http://sbir.cancer.gov)
Email: NCIsbir@mail.nih.gov
Phone: 240-276-5300

For additional information on research topics, please contact a Program Officer with the relevant area of expertise:

Michael Weingarten, MA
Director, NCI SBIR Development Center
Email: weingartenm@mail.nih.gov
Greg Evans, PhD  
Program Director and Team Leader  
Email: evansgl@mail.nih.gov  
 Areas of expertise: Therapeutics (Immunotherapy, Gene Therapy), Cancer Imaging, Cancer Control, Tools for Cancer Biology Research, and Digital Health

Deepa Narayanan, MS, CCDM  
Program Director and Team Leader  
Email: narayanand@mail.nih.gov  
 Areas of expertise: Medical Devices, Clinical Trials, Cancer Imaging and Radiation Therapy

Kory Hallett, PhD  
Program Director and Team Leader  
Email: kory.hallett@nih.gov  
 Areas of expertise: Immunology, Immunotherapy/Immuno-Oncology, Monoclonal Antibodies, Hematopoietic Stem Cell Transplantation, and Cell-Based Therapies

Christie Canaria, PhD  
Program Director  
Email: Christie.canaria@nih.gov  
 Areas of Expertise: Cancer/Biological Imaging, Research Tools and Devices

Jonathan Franca-Koh PhD, MBA  
Program Director  
Email: jonathan.franca-koh@nih.gov  
 Areas of expertise: Cancer Biology, Biologics, Small Molecules, and Cell Based Therapies

Nancy Kamei, PharmD  
Program Director  
Email: nancy.kamei@nih.gov  
 Areas of Expertise: Therapeutics

Xing-Jian Lou, PhD  
Program Director  
Email: loux@mail.nih.gov  
 Areas of expertise: In Vitro Diagnostics and Therapeutics (Gene Therapy, Biologics, Small Molecules)

Monique Pond, PhD  
Program Director  
Email: monique.pond@nih.gov  
 Areas of Expertise: Biologics, Small Molecules, Therapeutic Devices, Digital Health, Regulatory Resources

Amir Rahbar, PhD, MBA  
Program Director  
Email: amir.rahbar@nih.gov  
 Areas of expertise: In Vitro Diagnostics, Proteomics, and Therapeutics (Biologics, Small Molecules)
Ashim Subedee, PhD
Program Director
Email: ashim.subedee@mail.nih.gov
Areas of Expertise: Therapeutics (Small molecules, nanotechnology-based, immunotherapy), Imaging Agents, Diagnostics and Theranostics, Digital Health, Cancer Prevention and Control, and Therapeutic Devices

Patricia Weber, DrPH
Program Director
Email: weberpa@mail.nih.gov
Areas of expertise: Digital Health and Therapeutics (Small Molecules, Biologics, Immunotherapy)

Ming Zhao, PhD
Program Director
Email: zhaoming3@mail.nih.gov
Areas of expertise: In Vitro Diagnostics, Cancer Stem Cells, Molecular Imaging, Bioinformatics, Therapeutics (Small Molecules, Biologics, Immunotherapy), and Cancer Control (Community-Based Participatory Research)

For administrative and grants management questions, please contact:

Ashley Salo
Office of Grants Administration
National Cancer Institute
9609 Medical Center Drive
West Tower, 2W502
Rockville, MD 20850
Phone: 240-276-5656
Email: ashley.salo@nih.gov

Prior to Submission

Applicants are strongly encouraged to contact SBIR/STTR staff prior to submitting any application. To schedule a meeting, please email ncisbir@mail.nih.gov with a copy of your specific aims page that includes answers to the following questions:

- What is your product?
- What would be the impact of your technology to cancer patients, providers or caregivers?
- How is your product innovative and how is it different from the current standard?
- What are your aims for the application? What will be your milestones or success criteria?
- Who is the end-user of your product? Who is the purchaser?

For NCI-related SBIR Information, visit: http://sbir.cancer.gov.
EUNICE KENNEDY SHRIVER NATIONAL INSTITUTE OF CHILD HEALTH AND HUMAN DEVELOPMENT (NICHD)

The mission of the NICHD is to lead research and training to understand human development, improve reproductive health, enhance the lives of children and adolescents, and optimize abilities for all.

The NICHD has a broad and diverse research portfolio, including biological, behavioral, and clinical research related to conception and pregnancy, normal and abnormal development in childhood, reproductive health, rehabilitation, and population dynamics across the lifespan.

For up-to-date information on priority research areas of scientific interest to the NICHD, please visit our home page at http://www.nichd.nih.gov.

Pre-submission Resources & Commercialization Assistance

There are several resources and programs available throughout the SBIR/STTR process. For more information, please visit our Commercialization Resources webpage.

Limited Amount of Award

The NICHD will accept SBIR/STTR applications up to $256K total costs for Phase I for a time period no greater than 2 years and $1.7M for Phase II for a time period no greater than 3 years. It is strongly encouraged to contact program staff prior to applying.

The NICHD received a budgetary guideline waiver from the Small Business Administration for applications relating to a limited list of scientific topics in Appendix A. For these the NICHD will accept applications up to $300K total costs for Phase I and $2M for Phase II. Applicants should propose a budget that is reasonable and appropriate for completion of the research project and requests for these budget levels must be very well justified. Applicants are strongly encouraged to contact the listed NICHD Branch Contact Program Officer for scientific-related questions about a project’s eligibility for a budgetary waiver.

For general budgetary questions, applicants are encouraged to contact NICHD’s SBIR/STTR Grants Management Coordinator. For budgetary, administrative or programmatic reasons the NICHD may decrease the budget or length of an award or decide not to fund an application recommended by scientific review.

Phase IIB Competing Renewal Awards

The NICHD accepts Phase IIB SBIR Competing Renewal applications to support additional R&D necessary for approval of a federal regulatory agency (e.g., FDA, FCC). Such products may include medical implants, pediatric devices, drugs, vaccines, and new treatment or diagnostic tools Applicants who received NICHD SBIR Phase I or Phase II support and who are currently Phase II awardees are eligible. If the project meets the criteria for a budgetary waiver (Appendix A), the Phase IIB should not exceed $3M total costs for three years. The amount of award may vary year to year depending on the research proposed. Funding priority will be given to those small business concerns that show the ability to develop innovative products and demonstrate growth towards independence from the SBIR/STTR programs.

NICHD Supported Funding Opportunity Announcements (FOAs)

In addition to the Omnibus program announcement, for up-to-date NICHD releases on targeted funding announcements and programmatic initiatives visit: https://sbir.nih.gov/funding/individual-announcements or https://www.nichd.nih.gov/grants-funding/SBIR_STTR/Pages/default.aspx.
Research Topics of Interest to NICHD

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NICHD Non-Clinical Trials Topics:

The major NICHD research priority areas for each Branch are listed below. Investigator initiated applications that have commercial potential that fall outside these topic areas but fall within the research mission of the NICHD are also considered through this Omnibus solicitation.

A. Child Development and Behavior Branch

The CDBB encourages innovative developmentally-sensitive theoretically-grounded evidence-based small business initiatives that develop technology and products addressing the psychological, social and emotional, psychobiological, language, numerical, literacy, cognitive and intellectual development and health of persons from infancy to maturity recognizing the important role others have in contributing to the healthy development of an individual. Products that target at-risk populations and/or exploit new technologies that can expand the effective reach or inclusion of underserved populations in order to encourage healthy development and/or our understanding of the influences of context and/or behavior on development are especially encouraged.

Foci of specific interest include, but are not limited to (please also see the CDBB description for research priorities):

- **Enhancing Bilingual and Biliteracy Development:** Adaptive learning technology to enhance bilingual and/or biliteracy development in English-language learning children and youth.
- **Measures of Neurodevelopment:** Develop easy to administer neurodevelopmental measures from evidence-based neurocognitive research specific to typically developing infants and toddlers that are shown to correlate with development of brain connectivity and activation.
- **Pediatric Primary Care Behavioral and Health Promotion Interventions:** Facilitate research on the impact of behavioral and health promotion interventions in pediatric primary care and related clinical settings with a focus on end result child and adolescent health outcomes.
- **Psychosocial Adjustment for Individuals in High-Risk Environments:** Develop measures to identify and tools to stimulate developmental factors and mechanisms which promote short- and long-term psychosocial adjustment for children and adolescents exposed to high-risk family and neighborhood environments.
- **School Readiness Skills in Economically and Socially Disadvantaged Children:** Develop mobile device apps and/or hand-held devices that promote the development of school readiness skills and abilities in diverse populations of children as well as measures of home, child care and preschool environments and practices that are related to child learning and development.
- **Reading, Writing, and Mathematics Struggling Learners:** Develop assistive technology to enhance learner outcomes for individuals that struggle to acquire literacy and numeracy skills.
- **Assessment and Enhancement of Reasoning Development:** Develop validated and specific assessment tools that are sensitive to contributing factors (e.g., biobehavioral, environmental, cultural, academic, and cognitive factors) to facilitate research on and the promotion of neurocognitive development of reasoning (e.g., quantitative, deductive, inductive, causal) in typically developing populations.

Kathy Mann Koepke, PhD
Phone: 301-435-6855
Email: KMK@nih.gov
B. **Contraceptive Research Branch**

The CRB supports research with an emphasis on developing new and improved methods of fertility regulation as well as research on the benefits and risks of contraceptive drugs, devices and surgical procedures.

Areas of interest include, but are not limited to:

- Development of new and improved methods of fertility regulation, for men and women, that are safe, effective, inexpensive, reversible and acceptable
- Synthesis and testing of novel chemical compounds that are potential contraceptives
- Multipurpose technologies designed to prevent sexually transmitted infections, such as HIV, as well as pregnancy

Steven C. Kaufman, MD
Phone: 301-435-6989
Email: kaufmans@exchange.nih.gov

C. **Developmental Biology and Structural Variation Branch**

The DBSVB supports biomedical research on the cellular, molecular, and genetic aspects of typical and atypical embryonic development including early embryogenesis, organogenesis, as well as topics in stem cell regenerative biology. The overall goal is to promote research on developmental biology to understand the causes of structural birth defects.

Areas of interest include but are not limited to:

- Development of new model systems (animal or other) to study developmental mechanisms and causes of structural birth defects
- Innovative technologies for *in vivo* imaging of developmental processes (cell and tissue dynamics) and gene expression
- Development of antibodies, novel ligands, and other probes to facilitate our understanding of typical and atypical embryonic development in model organisms
- Technologies for quantitative measurement of physical properties of cells/tissues *in vivo* during development
- Innovative technologies for studying metabolomics in developing vertebrate embryos
- Technologies to facilitate and advance systems biology approaches to the study of embryonic development and structural birth defects
- Technologies to facilitate and advance high throughput chemical screening (including small molecules) for advancing structural birth defects research
- Software development to facilitate the collection and analyses of data generated using medium-high throughput screening platforms in model systems (model organisms, cell-based models)
- Software development to facilitate the collection, mining and analyses of genomic and phenotypic data from children affected with structural birth defects, and cross-analysis with model organism data
- Development of user-friendly software for biomedical researchers with limited knowledge of computational biology to analyze large-scale human and other datasets associated with structural birth defects
- Technologies/methodologies to generate, and software to mine, data related to wound healing and regenerative responses across animal species
- Novel reagents for activation and mobilization of endogenous/adult stem cells to promote *in vivo* tissue regeneration
• Technologies for iPSC-based regenerative medicine in the context of structural birth defect
• Screening technologies for small molecules in human Embryonic Stem (ES) Cells or Induced Pluripotent Stem Cells (iPSCs) and disease specific iPSCs for targeted modification of regulatory networks affected in structural birth defects

Mahua Mukhopadhyay, PhD
Phone: 301-435-6886
Email: mukhopam@mail.nih.gov

D. Fertility and Infertility Branch

The FIB supports research on the reproductive processes of men and women and of animals with similar reproductive systems related to developing safer and more effective means of regulating, preserving or achieving fertility.

Areas of interest include but are not limited to:

• Development of reagents to facilitate study of reproductive and developmental processes, such as gamete and early embryo development, and reproductive track development
• Development of improved methods of growing and differentiating stem cell lines in vitro, including feeder cell-free approaches to facilitate reproductive research
• Development of novel assays, kits, and devices to monitor fertility and treat infertility and gynecological disorders
• Use of genomics and proteomics to develop novel diagnostics and treatments for reproductive diseases and disorders
• Development of high-resolution technologies to provide invasive or noninvasive assessments of reproductive and developmental competence
• Development of experimental animal models that would be useful for studying the physiology and pathophysiology of reproductive processes
• Development of techniques and identification of novel biomarkers to produce, identify, and use healthy gametes
• Development of improved and novel technologies for the preservation of human gametes
• Development of improved technologies for preimplantation genetic diagnosis
• Development of in vitro model systems that are useful for understanding human embryo implantation.
• Development of improved technologies for the reprogramming of cells, including embryonic stem cells or adult cells, into eggs and sperm
• Development of innovative technologies for point-of-care testing for infertility and reproductive diseases and disorders
• Development of new methods to alter the function of trophoblast cells so that the embryo/fetus can be protected from ill effects of maternal viral infection
• Development of genomic, epigenomic or proteomic technologies to diagnose impairments in sperm function, fertilization, ovulation, implantation, decidualization and other aspects of reproductive processes

Clara Cheng, PhD
Phone: 301-435-6992
Email: clara.cheng@nih.gov

E. Gynecologic Health and Disease Branch
The **GHDB** supports biomedical research related to gynecologic health throughout the reproductive lifespan, beginning at puberty and extending through early menopause.

Areas of interest include, but are not limited to:

- Development of new diagnostic approaches and treatments for female pelvic floor disorders, including drugs, and devices used for treatment of pelvic organ prolapse, urinary incontinence, fecal incontinence, and other female pelvic floor disorders
- Development of new diagnostic methods and novel surgical and non-surgical treatments for uterine fibroids, endometriosis, adenomyosis, and benign ovarian cysts
- Production of marketable novel or improved methods, devices, and technologies for the diagnosis, monitoring and therapy of gynecologic pain disorders including chronic pelvic pain, vulvodynia, and dysmenorrhea
- Generation of new approaches for the diagnosis, monitoring and treatment of abnormal menstrual cyclicity
- Surgical and non-surgical treatments for girls and women with reproductive tract abnormalities, including congenital structural abnormalities and complications from female genital cutting.
- Devices and/or technologies designed to address surgical challenges in gynecologic surgeries, including hysterectomy.
- Technologies designed to apply omics platforms (genomics, proteomics, metabolomics etc.) to questions of gynecologic health and disease.

Candace M. Tingen, PhD  
Phone: 301-435-6971  
Email: candace.tingen@nih.gov

**F. Intellectual and Developmental Disease Branch**

The **IDDB** sponsors research aimed at preventing, diagnosing, and ameliorating intellectual and developmental disabilities (IDD). Emphasis is on studies related IDD, including common and rare neurodevelopmental and neuromuscular disorders, such as autism spectrum, Down, Fragile X, and Rett syndromes, mitochondrial conditions, inborn errors of metabolism, autism spectrum disorders, and others.

Areas of interest include, but are not limited to:

- Innovative tools, including molecular, imaging, statistical or behavioral tools, to characterize the etiology and pathophysiology of abnormal nervous system development.
- Methods and devices to delineate genetic, genomic, and epigenetic causes of IDD and develop gene-based treatments.
- Methods or devices designed to screen for and diagnose IDD and other conditions, particularly those identified or identifiable by newborn screening.
- Assessment tools for use in the clinic or community settings to enable the accurate measurement of change in response to interventions.
- Development of early interventions leading toward the prevention, diagnosis, treatment, and management of IDD.
- Methods or devices to develop or apply smart technologies (such as wearable devices, mobile health applications to assist in remote health monitoring, point-of-care diagnostic tools, etc.) to enhance screening, diagnosis, prevention, treatment or management of IDD conditions.
- Development of assessment measures or treatments for co-morbid symptoms in those with IDD including disordered sleep, obesity, gastrointestinal dysfunction, immune
dysregulation, seizures/epilepsy, self-injurious behaviors, and ADHD and other mental health disorders.

Sujata Bardhan, PhD
Phone: 301-435-0471
Email: sujata.bardhan@nih.gov

G. Maternal and Pediatric Infection Disease Branch

The MPIDB supports domestic and international research on human immunodeficiency virus (HIV)/acquired immune deficiency syndrome (AIDS) and related infections (such as tuberculosis, hepatitis and malaria) in women of child bearing age, pregnant women, mothers, fetuses, infants, children and adolescents. Specific areas of interest include but are not limited to epidemiology, clinical manifestations, immune-pathology, pathogenesis, transmission, treatment and prevention (including immune-therapeutics like monoclonal antibodies, vaccines and other biomedical modalities) of HIV infection, Zika infection and other infectious diseases in children, adolescents and pregnant women, including prevention of mother to child transmission of HIV and other congenital infections, and HIV-related and other infectious-disease related complications in these populations.

Additional areas of interest include:

- New technologies relevant to resource-limited countries for:
  - Screening, diagnosis, and management of infectious diseases in pregnant women, infants and children, including but not limited to HIV such as congenital CMV, congenital Syphilis, and Zika virus
  - Rapid assays to monitor disease activity and response to therapy as well as immune response to vaccinations against relevant infections in infants and children (e.g. malaria, tuberculosis), which can be used at the individual level and/or as part of public health campaigns (e.g. eradication of outbreaks and prevention of spread)
  - Diagnosis and treatment of HIV-related co-morbidities (e.g., diagnosis of tuberculosis in children)
  - Diagnosis and treatment of Zika-related outcomes in mothers and infants
  - Simple and less technologically demanding point of care assays to monitor CD4 cell percentage/count, HIV viral load, or other surrogate markers of HIV disease progression in children
  - Interventions designed to promote or optimize medication adherence

- Child-friendly formulations (preferably not liquid preparations) of drugs used to treat or prevent HIV infection, complications of HIV infection, and/or other high-priority infections such as tuberculosis, hepatitis, and malaria relevant to children, particularly in resource-limited countries; Fixed-dose drug formulations and innovative methodologies for development of solid heat stable formulations capable of being administered to young children (e.g., sustained release beads, etc.) and/or improve pill or volume burden

- Innovative long-lasting drug formulations for antiretroviral and other anti-infective drugs that would allow less frequent drug administration (e.g., once daily, weekly, or monthly)

- Simple, standardized, validated tools to evaluate neurodevelopmental outcomes in children in resource-limited settings

- Innovative data collection and database development approaches to leverage and link electronic medical records and/or other health information systems to better understand HIV treatment and prevention, and related infections, among infants, children, adolescents and women of child bearing age.

- Biomedical modalities including vaccines and methods to assess efficacy of vaccines, to prevent acquisition of HIV and other infectious diseases in children, adolescents and women.
• Topical microbicide agents, wearable, implantable, or insertable devices releasing medications alone or as part of multipurpose prevention technologies (MPTs), to prevent sexual acquisition of HIV and other sexually transmitted infections in adolescents, adult women, and pregnant or postpartum women.

• New, non-invasive technologies to evaluate complications of antiretroviral drugs (e.g., mitochondrial toxicity, bone toxicity) in HIV-infected infants, children, adolescents, pregnant women, and their fetuses.

• New or improvements to existing technologies for measuring the HIV latent reservoir, including high-throughput, visualization algorithms, and improvement in assay reliability and sensitivity.

Sai Majji, PhD  
Phone: 301-661-9816  
Email: sai.majji@nih.gov

H. Obstetric and Pediatric Pharmacology and Therapeutics Branch

The OPPTB aims to assure that there are safe and effective therapeutics for children and pregnant and lactating women and that these medications are used optimally according to individual needs. The branch promotes basic, translational, and clinical research to improve the safety and efficacy of therapeutics, primarily pharmaceuticals. It is responsible for developing and supporting a comprehensive national effort to increase the knowledge base for understanding how to appropriately treat disease during pregnancy, lactation, infancy, childhood, and adolescence using evidence-based therapeutic approaches. This includes support for the development and validation devices to inform treatment decisions and enhance precision drug delivery. The goal of these efforts is to assure that medications are appropriately tested for dosing, safety, and effectiveness for individuals within their target populations.

Applications to advance the study of obstetric and pediatric therapeutics include but are not limited to:

• Development of artificial intelligence (AI) pharmacokinetic/pharmacodynamic modeling to improve therapeutic approaches during pregnancy, infancy, childhood, and adolescence.

• Development of autonomous drug delivery systems that monitor and use the status of various organ systems for real-time adjustment of therapy in neonates, especially those that use AI to learn and react to individual differences

• Development of robustly controlled autonomic specimen sampling and analysis for limited blood volumes

• Development of AI-driven approaches for computational identification of novel potential drugs or drug repurposing across multiple therapeutic areas

• Development of non-invasive devices for evaluating adherence to chronic therapy in life-threatening conditions (e.g., HIV, diabetes, asthma, and liver and kidney transplantation)

• Use of a materials science approach to overcome solubility limitations of pediatric drugs, increase bioavailability, decrease excipient exposure, and provide effective taste masking

• Development of biomarker panels and/or drug transporter panels to enhance the diagnosis and refine therapeutic treatment for pediatric and obstetric conditions

June Lee, MD, PhD  
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I. Pediatric Growth and Nutrition Branch

The PGNB supports research designed to support short and long-term health so that children can achieve their full potential through an expanded understanding of those factors that influence
metabolism, growth (body composition and linear growth) and neurodevelopment. An additional focus is on those biological (e.g., genetic, nutritional, endocrinological) factors that contribute the early life origins of non-communicable disease (e.g., obesity, diabetes, cardiovascular disease, osteoporosis). The PGNB encourages research that focuses on detecting the biological antecedents of these conditions during pregnancy, infancy, and childhood.

Areas of interest include, but are not limited to:

New research tools, improved measurement methods, and technologies that enhance our understanding of:

- **Growth:***
  - Physical growth, body composition, bone health, nutrition, and obesity
  - Determinants of normal bone mineral accretion and peak bone mass. Interactions of muscle and bone during infancy and childhood
  - Neuroendocrinology of puberty, linear growth, body composition
  - Mechanisms of hormone action during linear growth, pubertal maturation, and other aspects of physical development

- **Biological antecedents of childhood obesity and its short and long-term consequences:**
  - Genetic and molecular mechanisms of obesity, psychosocial risks of obesity, and therapeutic interventions for obesity in children and adolescents
  - Impact of early life exposures including infant feeding practices on short and long-term health and development

- **Biology of nutrition as it pertains to health and development (physical and neurological function) during pregnancy, infancy and childhood including discovery, development and deployment of biomarkers for early detection of:**
  - Mal-(over-/under) nutrition; including biomarkers of exposure, status, function and effect (i.e., impact on early life development including neurodevelopment)
  - Enhanced understanding of the role of human milk in child health and development.
  - Maternal nutrition (pre-pregnancy, pregnancy, and lactation)
  - Novel approaches to enhanced infant feeding practices in term and pre-term infants

- **Developmental origins of health and disease including:**
  - Ascertain biomarkers early in life that predict the onset of chronic diseases such as diabetes, osteoporosis, and the metabolic syndrome later in life. The PGNB emphasizes the life course model to develop primary preventive approaches to chronic diseases.
  - Develop platforms for implementation of biomarkers of disease status, nutritional status, and biological function from infancy through adolescence

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J. **Pediatric Trauma and Critical Illness Branch**

The PTCIB supports research and research training in pediatric trauma, injury prevention, and critical illness across the continuum of care. These efforts include:

Research on the prevention, treatment, and management of physical and psychological trauma and the surgical, medical, psychosocial, and systems interventions needed to improve outcomes for critically ill and injured children and youth.
Studies of the continuum of psychosocial, behavioral, biological, and physiological influences that affect child health outcomes in trauma, injury, and critical care.

Basic, clinical, and translational studies that explore short- and long-term consequences of such traumatic experiences as exposure to natural or man-made disasters, all forms of violence against children, as well as experiences of bereavement, grief and loss.

Research linking the science of pediatric emergency and critical care medicine to the epidemiology, prevention, and treatment of trauma and injury in infants, children and adolescents.

Applications of interest include, but are not limited to:

- Research and development of pediatric-specific technologies, devices and equipment used by emergency and trauma care as well as pediatric critical care personnel.
- Research and development of novel strategies or approaches in caring for injured children prior to and during transport to treatment settings.
- Development of tools and technologies for screening and determination of the nature and extent of injuries related to forms of child maltreatment.
- Research and development of devices and innovative therapeutic technologies for management of medical conditions and related problems stemming from critical illness and serious or life-threatening injuries.
- Development and testing of preventive intervention tools, materials, and technologies designed to improve clinical practice, parenting and social system support for injured or traumatized children.
- Development and testing of preventive intervention tools, materials, and technologies designed to reduce pediatric trauma exposure and the number and severity of pediatric injuries and deaths.
- Research and development of effective tools and technologies to improve the environment of pediatric intensive care including resources to promote patient safety and to enhance clinical education and training of critical care personnel.
- Development of tools and technologies that support the diagnoses and treatment of critical illness in children, including nosocomial infections and iatrogenic injury.

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K. Population Dynamics Branch

The PDB supports research and research training in demography, reproductive health, and population health. In demography, the Branch supports research on the scientific study of human populations, including fertility, mortality and morbidity, migration, population distribution, nuptiality, family demography, population growth and decline, and the causes and consequences of demographic change. In reproductive health, the Branch supports behavioral and social science research on sexually transmitted diseases, HIV/AIDS, family planning, and infertility. In population health, the Branch supports data collection and research on human health, productivity, behavior, and development at the population level, using such methods as inferential statistics, natural experiments, policy experiments, statistical modeling, and gene/environment interaction studies.

Applications are encouraged, but are not limited to these areas:

- Technological innovations or inventions to improve collection of biomarker and anthropometric data in large population-representative surveys
- Hardware or software to improve collection of accurate cause of death information or health diagnosis such as information related to maternal morbidity and mortality, in large population-representative surveys or in administrative data sets
- Methods for integrating geographical information systems (GIS), spatial network analysis, and/or simulation methods for demographic research
- Methods for improving collection, documentation, archiving, and dissemination of population representative data sets, especially data sets that are complex, multilevel or multimodal
- Methods for protecting and assuring confidentiality for human subjects when collecting, archiving, or disseminating population-representative data sets, especially data sets that are longitudinal or that include both spatial and individual-level data
- Methods for reducing cost of collecting and disseminating large-population-representative data sets
- Development of effective tools for prevention research and intervention programs related to STD/HIV, pregnancy, contraceptive use, child health, at risk youth, adolescent, young adult, and maternal mortality, and other health-related topics, and to the dissemination of such tools
- Innovative approaches to teaching population studies and other behavioral and social sciences at the undergraduate and graduate level
- Innovative approaches for research design, data collection techniques, measurement, and data analysis techniques in the social and behavioral sciences, with particular attention to methodology and measurement issues in studying diverse populations, sensitive behaviors, confidential behaviors; in issues related to the protection of research subjects; and in issues related to the archiving and disseminating complex datasets

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L. Pregnancy and Perinatology Branch

The PPB supports research in the following areas: the physiology of pregnancy and labor; high-risk pregnancies, including those with hypertensive disorders, diabetes or seizure disorders; fetal pathophysiology; premature labor and birth; diagnostic, monitoring, and therapeutic devices and instruments for newborn infants in the nursery and in Neonatal ICU setting; improving the existing products or developing new products that would improve the routine and extended care of the newborn infants; products and agents related to breastfeeding; hospital supplies specifically related to use in the care of newborn infants; nanotechnology and its application for the care of newborn infants; instruments and devices for assessing and monitoring the nursery environment (noise, lighting, and odor); disorders of the newborn; sudden infant death syndrome; and biological and behavioral antecedents of low birth weight.

The following topic areas are of high priority:

- Non-invasive (or minimally invasive) methods to assess preeclampsia; gestational diabetes; fetal well-being; spontaneous preterm birth; and stillbirth
- Methods to characterize the bioactive components of human milk
- Methods to longitudinally assess the structure and functions of the human placenta
- Devices, instruments, and tools to minimize bacterial colonization, reduce proclivity for thrombus formation, and reduce healthcare associated infection risks
- Lab-on-a-chip; specifically, non- or minimally-invasive approaches for assessing: metabolic profiles (e.g., glucose and lactate/pyruvate), ketone bodies, bilirubin (unconjugated, free, indirect, and total), and other major analytes (Na⁺, Ca⁺, Cl⁻, K⁺ etc.)
• Rapid methods for diagnosis of bacterial infections and the assessment of antibiotic sensitivity
• Improved syringes, needles, and injection set ups to help administer small doses of medications over prolonged periods (e.g., insulin for treating hyperglycemia)
• Methods to assess pain in the newborn, analgesia, and the evaluation of neonatal opioid withdrawal syndrome
• Non-invasive measures to assess brain energy utilization in the newborn, especially glucose, oxygen, lactate, ketones, and other energy substrates
• Improved devices and instruments for assisted ventilators for use in the neonatal ICU

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M. National Center for Medical Rehabilitation Research

This Center supports innovative research on the restoration, replacement, enhancement or adaptation of function for people with chronic physical disabilities. This includes rehabilitative approaches across etiologies and the lifespan, as well as the environmental and policy factors that promote full participation. We encourage studies that integrate biomedical, engineering and/or psychosocial approaches to develop practical and creative solutions to the daily functioning of people with disabilities and their families. The mission of the NCMRR is to increase the effectiveness of medical rehabilitation practices through research. Information about specific program areas within NCMRR can be found here.

Examples may include but are not limited to:

• Adaptation and Plasticity: Develop non-invasive and surrogate measures of plasticity that would be appropriate for use in a clinical setting to target rehabilitation therapies and monitor treatment effectiveness (e.g., biomarkers, imaging)
• Novel Technology: Orthotics, prosthetics, and robotics devices and interfaces; Assistive technologies; Invasive and non-invasive biological sensors, prosthetic systems or implants to improve function; New control methods and improved sensory feedback; Strategies for controlling and adapting to the environment; Advanced wheelchair designs and enhancements and other mobility devices; Biomaterials and tissue interfaces, nanotechnology, bionics
• Rehabilitation Interventions: Development and use of robotics; Gaming applications; Virtual and Augmented Reality; Simulations; M-health and other approaches to promote participation, understand and support healthy behaviors, reduce health disparities and enhance clinical compliance, especially in children with physical disabilities.
• Chronic Symptom Management: Methods to increase screening for chronic conditions or preventable secondary conditions in individuals with physical disability; Prevention and treatment strategies for mitigating symptoms associated with multiple chronic conditions in individuals with physical impairments, including persistent pain, symptoms of obesity, diabetes, cardiovascular deconditioning, fatigue, symptoms of overuse injuries, pressure ulcers, sleep disturbances, and depressive symptoms; Improving muscle capacity in chronic physical disability to include therapeutic or adaptive exercise and muscle stimulation; muscle-disuse syndromes and contractures; Rehabilitation interventions for improvement of physical disability and comorbid cognitive, sensory, or somatic consequences of impairment, disease or injury; Autonomic function in the context of injury or specific conditions.
• Rehabilitation in the Community: Strategies to build or modify community and/or environmental resources that provide effective rehabilitation and health promotion services within the individual’s own community. Development of engineering,
crowdsourcing, and social science approaches to promote, monitor, and sustain outcomes in real world settings.

Investigators proposing budgets exceeding the guidelines are encouraged to contact program staff six weeks prior to submitting the application.

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**NICHD Clinical Trials Topics:**

The NICHD will generally consider clinical trial proposals consistent with the topics listed above with the following exception:

A. **Developmental Biology and Structural Variation Branch**  
The DBSVB does not support clinical trials through the SBIR/STTR program.

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For additional SBIR/STTR program administrative information and research topics contact:

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For additional financial/business management questions contact:

Mindy Bixby  
Grants Management Specialist  
*Eunice Kennedy Shriver National Institute of Child Health and Human Development*  
Contractor  
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NATIONAL INSTITUTE ON DRUG ABUSE (NIDA)

NIDA is the lead federal agency supporting scientific research on drug use and its consequences. Our mission is to advance science on the causes and consequences of drug use and addiction and to apply that knowledge to improve individual and public health through: 1) Strategically supporting and conducting basic and clinical research on drug use (including nicotine), its consequences, and the underlying neurobiological, behavioral, and social mechanisms involved; and 2) Ensuring the effective translation, implementation, and dissemination of scientific research findings to improve the prevention and treatment of substance use disorders and enhance public awareness of addiction as a brain disorder.

SBIR and STTR programs at NIH are primarily intended to encourage private-sector commercialization of technology and to increase small business participation in federally funded research and development (R&D). The goal of NIDA SBIR/STTR program is to increase small business participation and private-sector commercialization of novel technologies that can prevent, diagnose, and treat Substance Use Disorders (SUD). For more information about NIDA’s portfolio interests, resources, funding opportunities and key contacts for a successful submission, please visit https://sbir.nih.gov/nida/index

The SBIR and STTR Programs are structured in three phases:

**Phase I.** The objective of Phase I is to establish the technical merit, feasibility, and commercial potential of the proposed R/R&D efforts and to determine the quality of performance of the small business awardee organization prior to providing further Federal support in Phase II.

**Phase II.** The objective of Phase II is to continue the R&D efforts initiated in Phase I. Funding is based on the results achieved in Phase I (or equivalent) and the scientific and technical merit and commercial potential of the project proposed in Phase II. NIDA seeks to determine that both technical feasibility and commercial feasibility are established in Phase I before making the decision about Phase II support.

**Phase III.** The objective of Phase III is for the small business concerns (SBC) to pursue commercialization objectives resulting from the Phase I/II R&D activities. The small business programs do not fund Phase III. NIDA encourages grantees to seek commitment(s) of funds and/or resources from an investor or partner organization for commercialization of the product(s) or service(s) resulting from the SBIR/STTR grant. Phase III funding may come from different sources: private investors, venture capital firms, strategic alliances, research contracts, sales of prototypes (built as part of this and/or other project), public offering, state finance programs, non SBIR-funded R&D or production commitments from a Federal agency for use by the United States government or other industrial firms. NIDA monitors SBC efforts to pursue, with non-SBIR/STTR funds, the commercialization of the Phase I and II outputs.

The NIH/NIDA dual peer review system is mandated by the statute. The first level of review is carried out by a Scientific Review Group (SRG) composed primarily of non-federal scientists who have expertise in relevant scientific disciplines and current research areas. The second level of review is performed by NIDA National Advisory Council. Only applications that are recommended for approval by both the SRG and the Advisory Council may be recommended for funding. Final funding decisions are made by the NIDA Director.

NIDA funding decisions for small business programs are based on a combination of factors:

- Programmatic priorities and current portfolio balance (for funded projects, search NIH RePORTER: https://projectreporter.nih.gov/);
- Potential for commercialization and public health impact;
- For Phase II applications: results of Phase I (or equivalent) clearly indicating that both technical feasibility and commercial feasibility were established, and the scientific/technical merit and commercial potential of the project proposed in Phase II;
- For applicants who received preceding SBIR and STTR grants: quality of prior performance and evidence of Phase III activities;
• The peer review critiques and overall impact scores;
• Availability of funds.

SPECIAL FEATURES OF NIDA SBIR and STTR PROGRAMS

Limited Amount of Award

Award budgets and project periods are listed in the Section II Award Information of the Omnibus/Parent Funding Opportunity Announcements for SBIR and STTR. Total award budgets include direct costs, indirect costs and fee, and are capped not to exceed the total award amounts listed.

NIDA also sets its own budget limit for the certain research topics which received a waiver to exceed the hard budget caps from the U.S. Small Business Administration. The current list of approved NIDA topics can be found in the “Program Descriptions and Research Topics” document, Appendix A. NIDA budget limit for Phase I on approved topic is $320,000 in total costs and project period up to 1 year. NIDA budget limit for Phase II on approved topic is $2,500,000 in total costs and project period up to 3 years.

Applicants are strongly encouraged to contact NIDA program officials prior to submitting any application in excess of the hard caps listed in the Omnibus/Parent Funding Opportunity Announcements for SBIR and STTR and early in the application planning process. In all cases, applicants should propose a budget and a project duration period that are reasonable and appropriate for completion of the research project.

Fast-Track Applications

NIDA uses Fast-Track mechanism for scientifically meritorious applications that have expressly high potential for near-term commercialization. Applicants are strongly encouraged to contact NIDA program officials prior to submitting a Fast-Track application at least 8 weeks before the application due date.

For Fast-Track consideration, NIDA encourages the following:

1) Preliminary data that clearly support the technical and commercial feasibility. If repurposing already existing drug/device for SUD diagnosis or treatment, preliminary data about existing drug/device and scientific rationale for the feasibility in SUD space;
2) Commercialization Plan that demonstrates a high probability of commercialization;
3) Clear, appropriate, meaningful and measurable goals (milestones) that should be achieved prior to initiating Phase II. A milestone is a marker in a project that signifies the point at which a major uncertainty in the project is resolved. Milestones show key events and map forward movement in the project plan. Milestones for Phase I must be written with the goal of addressing all significant questions of technical and commercial feasibility. Phase I milestones must be quantitative, not qualitative (e.g., “an affinity of x nM,” rather than “twice as potent as Y” or the “twice as potent as the best analog”). Milestones must be written utilizing the appropriate controls. All milestones must be achieved prior to initiating Phase II. Phase I milestones must be written to allow to evaluate the claims that their completion fully justifies the initiation of Phase II;
4) Letters of Phase III support/interest, additional funding commitments, and/or resources from the private sector or non-SBIR/STTR funding sources;
5) Discussion with NIH Program Staff.
Research Topics of Interest to NIDA

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
<th>No*</th>
<th>Yes**</th>
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<tbody>
<tr>
<td>X</td>
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*If No,
- This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, UNLESS the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,
- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency accept clinical trials applications under this mechanism?</th>
<th>Yes</th>
<th>No</th>
<th>Other Information: If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information.</th>
</tr>
</thead>
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<tr>
<td>X</td>
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The major NIDA SBIR/STTR portfolio areas are listed below as a guide to general technology areas funded through the program. Applications proposing innovative SUD-related technologies with strong...
commercial potential that fall outside these topic areas are also encouraged through this Omnibus solicitation.

For up-to-date information about NIDA SBIR/STTR Program, and to learn about programmatic initiatives and upcoming events, visit the homepage: https://sbir.nih.gov/nida/index

Topics of Special Interest

- Biomarker Development for Substance Use Disorders (SUD)
- Biomedical Robotic Cloud Labs
- Substance Use Disorder (SUD) Drug Discovery and Development
- Technologies for Safe and Controlled Methadone Dispensing for Use at Home
- FDA-regulated Medical Devices for Substance Use Disorder (SUD)
- Technological Approaches to Address Stigma Associated with Substance Use Disorders (SUD)
- Digital Health Technologies to Address the Social Determinants of Health in context of Substance Use Disorders (SUD)
- New technological approaches for the Investigation, Diagnosis, and Certification of Deaths Related to Drug Overdose

Biomarker Development for Substance Use Disorders (SUD)

Currently there are no biomarkers able to assess or predict treatment efficacy or to categorize SUDs into clinical subtypes. Thus, it is not possible to design treatments for effective and long-term recovery by classifying SUD patients into categories that have reproducible and predictive validity.

Biomarkers and signatures in patients with SUD can be very different from those observed in patients without SUD because long-term use of opioids and other substances alters the integrity of homeostasis, changing the endogenous opioid, endogenous cannabinoid and almost all receptor systems studied so far in the brain and peripheral immune cells. These biomarkers or potential predictive markers could serve as objective prognostic indicators to develop SUD. In addition, they could act as response predictors to SUD therapeutics in adults, or as diagnostic biomarkers for infants with neonatal abstinence syndrome (NAS). Addition of biomarker signatures that span the trajectory of substance use disorder assessment in patients with addiction disorders, and prediction of the potential to develop SUD are also interest of the program.

The proposed biomarker research should emphasize the importance of biomarker signatures that can intersect SUD and related conditions that are considered important to the mission of NIDA. Projects may include proposing biomarkers that assess the probability of SUD or allow an assessment of the treatment trajectory in patients under treatment for SUD. Grant applications solely focused on biomarkers for pain in patients with pain not associated with SUD are of limited interest.

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Biomedical Robotic Cloud Labs
The day-to-day labor required to conduct biomedical wet lab science frequently occupies a significant proportion of a scientist’s time with monotonous and repetitive labor (mixing, centrifugation, filtering, buffer preparation, weighing, pH adjusting, etc). These tasks could not just be very successfully automated, but for many, the automation would also improve standardization. Thousands of labs are doing routine genotyping, cloning, sequencing, all with slightly different protocols. This could be assisted by the use of robust IT-enabled approaches, driven by robotics and computation. When outsourcing wet lab experiments, scientists would simply ship samples, design and run the experiment, then explore and analyze the data. Essentially, a complete life science laboratory can be controlled from the individual’s computer. This capability could improve the way biomedical science is conducted in normal times, enhancing the fundamental mission of NIH. It could also be revolutionary as the biomedical research workforce adjusts to the impact of social distancing imperatives on the ability to conduct lab work.

Limited existing efforts in this area have demonstrated the advantages of this approach in making experiments cheaper and faster; more reproducible; and easy for scaling up of the successful pilots without changing the original protocol. This approach also allows for economies of scale, 24-h production, improved rigor and unlimited access.

The topic calls for the research proposals to virtualize the various procedures and operations that serve as a cornerstone of the experimental biomedical research. Virtualization of methods used in addiction research labs is of special interest.

Once proven to be amendable to virtualization through the remote use of robotics and computation, these biomedical procedures and operations will be used to equip the cloud labs, which would then contribute to expansion of the scientific workforce by providing more scientists access to certified wet labs. Scientists at smaller institutions whose research has been constrained by lack of assay capabilities would be able to pursue new research questions, and researchers with existing robust wet lab resources would be able to pivot from the more monotonous tasks covered by the robotic cloud lab toward more complicated experimental protocols and intellectual work. Importantly, the proposed approach preserves and enhances the opportunity for sophisticated intellectual pursuit by the individual scientists, while opening the door for the scientists whose access to wet labs is currently limited or nonexistent.

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Substance Use Disorder (SUD) Drug Discovery and Development

Pharmacotherapy offers a promising means for treating SUD. Currently, there are five Food and Drug Administration (FDA)-approved pharmacotherapies for the treatment of OUD and mitigation of opioid withdrawal symptoms: Methadone, Buprenorphine, Extended-release naltrexone, Naloxone and Lofexidine. Varenicline is an approved drug for the treatment of nicotine cessation. However, given the diverse nature of SUD, many patients have limited response to available medications. There are no FDA-approved medications for cocaine, methamphetamine, or cannabis use disorders. Because of this, developing and evaluating new, more efficacious medications remains a high priority. Candidate medications may include novel and re-purposed compounds. Grant applications that propose to study compounds already extensively investigated or currently being studied in patients with SUD are of limited interest. Grant applications pursuing drug discovery and development for pain should propose the research in the context of SUD.

Specific areas of interest include medications that target one or more domains of the addiction cycle, including reward, stress and negative affect, incentive salience, executive function, habituation, and
impulsivity/compulsivity. Compounds may be developed for indications such as prevention of initiation of SUDs, prevention of progression of the severity of SUDs, improvement of SUD treatment adherence, facilitation of opioid agonist discontinuation, treatment of opioid withdrawal, treatment of neonatal opioid withdrawal, reduction of lethality of opioid overdose, reduction of overdose relapse and reduction of the risk of opioid respiratory depression. Additional targets for pharmaceutical development include agents that attenuate excessive substance use and other symptoms of SUD, e.g., craving, sleep problems, and negative affect as well as those that hold promise for the treatment of from tobacco, cannabis, cocaine, methamphetamine, heroin, or prescription opiate use.

Projects proposed may include application of emerging and existing technologies and platforms for SUD medication development with the focus on products with the potential to minimize drug seeking, compulsive behavior, overdose prevention and reversal:

- Early therapeutic discovery activities ranging from target identification and validation through lead development.
- Preclinical and/or clinical drug development.
- Technologies or formulations to improve medication delivery, longer-acting formulations of existing addiction medications
- Medications that would address specific symptoms of withdrawal, such as cravings, depression, cognitive impairments, pain, and sleep problems
- Medications are those that, while not addressing addiction directly, target major risk factors for relapse, insomnia, dysphoria and depression
- Medications (neurochemicals) involved in social bonding that also modulates key processes associated with addiction, including reward and stress responses and may enhance the efficacy of psychosocial addiction treatments
- Big-data analytics and machine-learning algorithms analysis yielding insight into behavioral and biological markers of relapse risk, tools or devices to avert relapse.

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Technologies for Safe and Dose-Controlled Methadone Dispensing for Use at Home

More than 350,000 Americans are prescribed methadone maintenance treatment (MMT) to treat opioid dependence. By law, methadone can only be administered or dispensed through an opioid treatment program (OTP) that is both Substance Abuse and Mental Health Services Administration (SAMHSA)-certified and Drug Enforcement Administration (DEA)-registered. According to the SAMHSA, methadone can be administered as a pill or a liquid. To receive the medication, patients must report to OTP centers, often daily. In most OTPs, the opioid dependent patient receives a daily dose of liquid methadone, with doses adjusted and tailored to the specific patient needs, to reduce withdrawal symptoms and opioid cravings. Some of the major barriers to methadone treatment are the restricted availability of timeslots in the OTPs for daily dosing, as well as logistical constraints and associated costs (e.g., travel from/to OTP).

A “take-home” is a dose of methadone given to the patient to take unsupervised at home in place of requiring a return to the clinic the next day for observed dosing. “Take-home” doses are offered as rewards to patients with regular clinic and counseling attendance, and abstinence from illicit drug use. Even with adherence to MMT program expectations, “take-home” doses may constitute up to a maximum of 6 or 13 consecutives doses, so that eligible patients may be required to be physically present at the
OTP either every week or every other week. The physician can also provide a certification for the patient as "exception take-homes", an option that is mostly used with patients on dialysis, patients with prescribed need of additional oxygen, patients who are wheelchair-bound, patients in residential treatment facilities, or patients with rapid metabolism of methadone who require more than one daily dose.

Recently, recognizing the evolving issues surrounding COVID-19, SAMHSA expanded the previous OTP guidance, allowing states to request blanket exceptions for all stable patients in an OTP to receive 28 days of “take-home” doses of the patient’s medication for opioid use disorder. Under this guidance the state may request up to 14 days of take-home medication for those patients who are less stable but who the OTP believes can safely handle this level of take-home medication. Additionally, the new SAMHSA guidelines allow for an expanded use of telehealth solutions that allow the provider to continue to treat remotely an existing OTP patient using methadone.

The new flexibility provided by SAMHSA increases the number of take-home patients and provide the much-needed social distancing. The current take-home process is based on providing patients with a specially designed lock box, which does not afford with the tracking capability or specific safety precautions. As such, there are concerns related to treatment adherence, possible abuse and diversion, patient vulnerability to theft and violence. As such, there is an urgent need for a comprehensive technological solution to continually assess the efficacy and safety of this take-home medication strategy.

There is a need for an integrated solution specifically addressing the intricacies of the methadone clinic, which combines multiple technological innovations and provides both OTPs and patients with answers that allow for safe and effective MMT, safe access to take-home medication, while reducing the burden and stigma that accompanies the current process. This includes the technological solutions that go beyond the pill boxes/dispensers, and which can be utilized for both liquid and pill form of take-home Methadone, with features such as remote monitoring, remote dispense control, teletherapy, and deactivation of the Methadone in case of diversion or theft. NIDA emphasized the need for the development of all-inclusive, holistic approach to enable safer “take home” processes for MMT.

Specific aspects of “take home” technologies or approaches for MMT may include, but not limited to:

- Remote tracking of medication adherence
- Remote dose and time-controlled dispensing
- Personalized secure access to medication dispenser
- Tamper-proof access to medication
- Digital health solutions that allow for patient-provider interaction (e.g., reminder for patients, telehealth solution for provider)

Proposed tools and technologies should take into full considerations the unique legal, public health and community aspects of MMT.

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Areas of expertise: Therapeutics (Small Molecules, Biologics, Nanotherapeutics, Immunotherapy, Cell & Gene Based Therapies), Biomarker Development and Validation.

FDA-regulated Medical Devices for Substance Use Disorder (SUD).
Medical Devices, including Digital Therapeutics, offers a promising means to monitor, diagnose, and treat SUDs. Currently, there are only a few Food and Drug Administration (FDA)-cleared medical devices for the diagnosis or treatment of SUD. The evaluation and development of new safe and effective medical devices is a high priority. Applications in this area are expected to propose activities that will lead to the regulatory submissions for pre-market clearance/approval. Depending on the specific regulatory pathway needed, applications in this area will include some of the following interactions with the FDA: pre-submission (Q-submission), Investigational Device Exemption (IDE), and 510(k), DeNovo, or Premarket Approval (PMA) application. Prior to clinical studies, additional expected specific activities include, but are not limited to a) pre-clinical bench testing or computational modeling studies; b) safety and/or effectiveness studies in animal models; c) Good Manufacturing Practice (GMP) studies; d) Non-clinical safety studies (e.g., toxicology, biocompatibility).

Specific areas of interest include:

- Imaging devices for investigating brain function and enhancing disease diagnosis and treatment of SUD;
- Devices that directly detect and/or reduce craving;
- Devices that identify and treat neonatal opioid withdrawal syndrome (NOWS), also referred to as neonatal abstinence syndrome (NAS);
- Digital health therapeutics (e.g., Software as Medical Device, Software in Medical Device) focused on behavioral health interventions to alleviate the burden of SUD;
- Therapeutic (e.g., neuromodulation) devices and other advanced methods to improve SUD treatment outcomes and relapse prevention;
- Medical devices used to diagnose and treat opioid-induced respiratory depression;
- FDA-regulated devices for physiological monitoring, including remote detection (e.g., wearable sensors, health monitoring/emergency notification systems) — tailored to patients with SUD.

Contact:
Leonardo Angelone, PhD
Program Officer
Email: leonardo.angelone@nih.gov
Areas of expertise: FDA-regulated Medical Devices, including Digital Therapeutics.

Technological Approaches to Address Stigma Associated with SUD

Leveraging the breakthrough technologies and the latest science should allow to develop and commercialize the products and services aimed at reducing stigma around substance use disorders (SUD). Stigma is defined as an identity marked by disgrace, disapproval or shame, which often leads to discriminatory treatment by others. Stigma typically is reflective of stereotypes or negative views attributed to a person or groups of people because of behavior or other attributes that are seen being different from broad societal norms. Stigma often may be directed toward multiple identities - these may be based on conditions such as SUD, mental illness, or infectious disease; behaviors such as specific drug use practices (e.g., opioid injection); or identity statuses related to gender, sexual orientation, sexual identity, race/ethnicity, or income. Stigma may be internalized (personal endorsement of prejudice and stereotypes), enacted (experiences of discrimination from others), or anticipated (expectations of discrimination from others in the future, even if one has not experienced discrimination in the past).

Applications in this area are invited to propose the projects to demonstrate how latest technology and evidence-based science could meaningfully reduce the stigma associated with SUD. Potentially commercializable research to reduce SUD stigma in adolescent population and research on effects of providing anti-stigma training to medical professionals and non-medical providers (social workers, criminal justice, family members, and educators) is of special interest.
Examples of technological approaches could include: the use of neuromarketing tools and services to help develop and disseminate the most effective anti-stigma campaigns; digital compassion (anti-stigma) coaching for medical professionals delivering treatment to patients with SUD; digital certification program for nonprofessional care givers who provide support services for patients with SUD; virtual employee assistance programs with focus on SUD and mental health that provide employees with support for issues that affect their well-being and enhance the effectiveness of a drug-free workplace program.

Contact:

Julia Berzhanskaya, PhD
Program Officer
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Areas of expertise: Biomedical Research Tools, Prevention and Education, Consumer Digital Products.

**Digital Health Technologies to Address the Social Determinants of Health in context of Substance Use Disorders (SUD).**

The taking of drugs of abuse is a high-risk behavior associated with immediate and long-term health consequences. The public health experts have long recognized the impact of social determinants on health outcomes. According to the World Health Organization (WHO), the social determinants of health are the conditions in which people are born, grow, live, work and age. These circumstances are shaped by the distribution of money, power and resources at global, national and local levels. Growing research is demonstrating that social determinants of health (SDH) play a far greater role in health outcomes than expected. The health of people who use drugs of abuse is inextricably bound to their social environment. Drug-taking and drug-use risk behaviors are affected by social processes, and health, in this situation, is a product of both drug-use behaviors and social determinants. Social determinants can directly shape health risk behaviors. SDH can be manifested in the living conditions and resources that indirectly exacerbate the consequences of drug use. For example, inadequate housing can increase the likelihood of infectious disease transmission, while the stable social relationships can offer protective financial and emotional resources, and more cohesive neighborhoods can have a greater likelihood of providing appropriate support and care.

A full spectrum of interventions encompassing both social determinants of health and individual-level factors should be considered in order to fully address the drug use epidemic in the Unites States. Digital technology-based solutions can offer a new path forward in addressing SDH in drug addiction, as these solutions focus on providing evidence-based, continuous and accessible experiences for individuals affected by drug use or living with SUD. The advantages of digital technology also lie in its capacity to accommodate the changing context and environments that contribute to the 21st century SDH: new communication means, mobility, cultural contexts, new consumer behaviors, family and community dynamics.

The eligible small businesses can submit the grant applications focusing on transforming family, housing, employment, justice and educational determinants of drug addiction. The proposed products should offer the most far-reaching and promising opportunities for the intended customers and end-users to meaningfully contribute to addressing the drug addiction and opioid crisis.

Illustrative topics could include, but are not limited to:

- Research, design and validate novel tools and approaches to address food and housing insecurities;
- Research and design of novel tools to enable impactful housing programs that promote health (for example, the innovative housing programs that can co-locate employment, education, and behavioral health services);
- Design and validation of curriculum for “soft skills” development for addiction treatment providers;
• Novel educational tools/novel educational delivery systems to foster compassion and eliminate stigma associated with SUD;
• Research and design of preventive systems for families to promote healthy behaviors, social skills, community opportunities, and productive social involvement;
• Novel educational tools/novel didactic delivery systems focused on social stability (community, tradition, faith, family), and self-regulation and resilience;
• Novel educational tools/novel didactic delivery systems to focus on happiness, wellbeing, belonging, positive and fruitful communal life;
• Design and validate technologies that help create and enhance productive social support networks that facilitate recovery, engagement with care, and/or access to needed services;
• Research and design tools and technologies to help facilitate continuity of care, access to services, and successful community reintegration for people re-entering communities following a period of incarceration;
• Development of technology to facilitate data sharing among organizations that serve justice-involved individuals with the goal of increasing coordination of services, enhancing service quality, and/or increasing engagement with effective services;
• Research, design and validation of novel approaches for job training (e.g. in entrepreneurship, financial literacy, IT skills), especially, delivered in recovery housing or while incarcerated;
• Develop and validate the best approaches for employer education and support to allow employers to hire, retain, and facilitate treatment for employees seeking help for SUD.

Contact:

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Program Officer
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Areas of expertise: FDA-regulated Medical Devices, including Digital Therapeutics.

New technological approaches for the Investigation, Diagnosis, and Certification of Deaths Related to Drug Overdose

Topic calls for research proposals to improve the various aspects of the practice of death investigation and autopsy, toxicological analysis, interpretation of those analyses, and death certification.

• Curation and digitizing of the jurisdiction-dependent practices and protocols;
• Improved tools to alleviate the test backlogs;
• Tools to improve communication and coordination among forensic pathologists, hospitals and lab technicians (medical examiner/coroner (ME/C));
• Improved methods to minimize the risk of infection with blood-borne pathogens (e.g., Hepatitis C or Human Immunodeficiency Virus) while performing the autopsy and toxicological analysis;
• Qualitative tests to determine a therapeutic vs toxic vs lethal doses;
• Rapid techniques and devices for field use;
• Improved immunoassays (e.g. to include urine, to decrease the number of false positives and negatives);
• Testing technologies able to rapidly to accommodate the changes in the new drug of choice.

NIDA hopes that the development of these tools will improve the detection and reporting of opioid-related deaths. Improved surveillance will reveal the magnitude of opioid-related deaths more accurately, thus clarifying attempts to decrease the number of opioid-related deaths and improving public health by monitoring the effects of these interventions.
Contact:

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Program Officer
Email: ramachandran.arudchandran@nih.gov
Areas of expertise: Therapeutics (Small Molecules, Biologics, Nanotherapeutics, Immunotherapy, Cell & Gene Based Therapies), Biomarker Development and Validation.

Prior to Submission

Applicants are strongly encouraged to request a technical assistance meeting with NIDA SBIR/STTR staff prior to submitting any application. To schedule a meeting, please email NIDASBIR@mail.nih.gov with brief answers to the following:

- What problem are you trying to solve?
- How is this problem solved now? What is the current gold standard to treat/to solve this problem?
- What is your product and how is it different from the current standard?
- Is your solution a consumer or regulated product? What is the FDA regulatory path?
- Who is the end-user of your product? Who is the purchaser?
NATIONAL INSTITUTE ON DEAFNESS AND OTHER COMMUNICATION DISORDERS (NIDCD)

The NIDCD supports research on the normal mechanisms of, as well as on diseases and disorders of hearing, balance, smell, taste, voice, speech and language. The Institute also supports research related to disease prevention and health promotion. The NIDCD addresses special biomedical and behavioral problems associated with people who have communication impairments or disorders. The NIDCD also supports efforts to create and refine devices, as well as develop cellular-based applications that may replace or substitute for lost and impaired sensory and communication functions. For more information about areas of interest to the NIDCD, please visit our home page at http://www.nidcd.nih.gov/ and the NIDCD Strategic Plan website (https://www.nidcd.nih.gov/about/strategic-plans). Potential applicants are encouraged to contact the program staff listed in the following descriptions of NIDCD program areas early in the process of preparing the application.

Limited Amount of Award

Total funding support (direct costs, indirect costs, fee) normally may not exceed $256,580 for Phase I SBIR/STTR awards and $1,710,531 for Phase II SBIR/STTR awards. Applicants considering a requested budget greater than these limits are strongly encouraged to contact program staff before submitting the application. The Small Business Administration has allowed NIDCD to make awards that exceed these amounts for the areas noted in Appendix A of this document. All applications must contain sufficient detail to justify the requested budget, and NIDCD may decrease the length of an award and/or the budget as recommended by a review committee or administrative review.

Phase IIB Competing Renewal Awards

The NIDCD will accept Phase IIB SBIR/STTR Competing Renewal grant applications to support use of the final product in:

- Clinical trials with a larger number of participants to adequately validate safety or efficacy.

Research Topics of Interest to NIDCD

The NIDCD accepts a broad range of small business applications that are significant, innovative, and relevant to its mission. Some examples of research topics within the NIDCD mission areas include topics shown below; further example can be found on the NIDCD Strategic Plan website (https://www.nidcd.nih.gov/about/strategic-plans). Priority is given to meritorious applications that are likely to develop innovative technologies, provide clear evidence of effectiveness, and bring novel products to the commercial marketplace.

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,**

- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

| Does IC/Office/Agency accept clinical trials applications under this mechanism? | Yes | No | Other Information: 
If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information. |
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<td>NIDCD accepts and supports non-SBIR/STTR clinical trial applications through specific opportunities, which can be found on the NIDCD Funding Opportunities webpage: <a href="https://www.nidcd.nih.gov/research/clinical-studies/researchers-professionals/know-what-is-available">https://www.nidcd.nih.gov/research/clinical-studies/researchers-professionals/know-what-is-available</a></td>
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**NIDCD Non-Clinical Trials Topics:**

NIDCD will accept applications for development and commercialization of novel products in any of the areas noted below.

**Hearing and Balance Program**

Development of treatment modalities to prevent or lessen the effects of hearing disorders; development of new hearing aids, cochlear implants, and other assistive devices; development of improved screening technologies to assess hearing loss, especially in neonates and infants; development of new or improved power sources for hearing aids or cochlear implants; development of technologies that provide self-fitting, self-adjusting, or other features that increase performance, accessibility, or affordability of hearing aids; development of new outcome measures for assessing the efficacy of treatments for hearing disorders; development of technologies for the study, diagnosis and treatment of tinnitus; development of technologies for the study, diagnosis and treatment of otitis media including non-invasive diagnostics to identify middle ear pathogens, novel antibacterial strategies, and prophylactic anti-microbial strategies; development of technologies for the study, diagnosis and treatment of noise-induced and age-related hearing loss.
Development of technologies for the study, diagnosis and treatment of balance disorders, particularly for the elderly; development of clinical tests and instruments to assess balance/vestibular function; development of instruments and tests measuring head stability and vestibular function during natural stimulation of the vestibular system; development of perceptual reporting techniques and psychological indices for clinical assessment of the balance-disordered patient; development of tests and new outcome measures for assessing the efficacy of physical rehabilitative regimens for balance disorders; and development of assistive devices for balance disorders, including neural prostheses for the vestibular system.

Development of new research tools to aid in the study of the auditory and/or balance systems that can provide an improved understanding of fluctuating patterns of neural circuit structure and function over time and across large assemblies of neurons; new animal models of impaired function; improved diagnostic tools for inner ear function, including DNA-based assays and biochemical markers of disease. Development of improved tests and instruments for screening and diagnosis of inner ear function; development of technologies to enable gene transfer to the inner ear, including viral vectors; development of cell type specific markers and probes to examine cell lineage in inner ear regeneration.

Roger L. Miller, Ph.D.
National Institute on Deafness and Other Communication Disorders
301-402-3458, Fax: 301-402-0390
Email: millerr@nidcd.nih.gov

Voice, Speech, and Language Programs

Development of technologies for the study, diagnosis and treatment of voice, speech, and language disorders is strongly encouraged, as are projects that focus on determining the nature, causes, treatment and prevention of communication disorders such as stuttering, Specific Language Impairment, spasmodic dysphonia, dysarthria, and aphasia. Emphasis is on research and development of diagnostic measures and intervention strategies for voice, speech, and language disorders; development of communication and other assistive devices for individuals with voice, speech, and language disorders; development of speech and language assessments and interventions for nonverbal individuals with autism; development of new systems for visual communication by individuals who are deaf or severely hearing impaired; development of new systems of communication for individuals with motor speech impairment, including a brain computer interface (BCI) communication prosthesis; development of innovative treatment delivery systems or intervention protocols; design and development of diagnostic measures or materials for early identification of voice, speech and language impairment in children; development of assessments and treatments for childhood and adult voice, speech and language impairment associated with bilingual or multi-cultural populations; development of assessment measures of sign language abilities; development of artificial intelligence computer models that simulate normal and disordered voice, speech and language.

Development of novel, low cost approaches capable of providing urgent news, events, updates in a format appropriate for users of American Sign Language. Development of technologies that assist in the access to or delivery of healthcare during a public health crisis to individuals with voice, speech and language disorders and/or those who are deaf or hard of hearing.

Judith A. Cooper, Ph.D. [Language Program]
National Institute on Deafness and Other Communication Disorders
301-496-5061, Fax: 301-402-0390
Email: cooperj@nidcd.nih.gov

Lana Shekim, Ph.D. [Voice & Speech Program]
National Institute on Deafness and Other Communication Disorders
301-496-5061, Fax: 301-402-0390
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Taste and Smell Program

Development of easily administered diagnostic tools for testing human chemosensory function throughout the lifespan; development of intervention strategies and targeted drugs for the treatment of taste and smell disorders; preventive measures to limit the deleterious effects of infections, airborne toxins, radiation, chemotherapy and other drugs on chemosensory function; novel therapies to stimulate regeneration of mature sensory neurons in damaged and/or aged tissue; development of olfactory biomarkers for neurodegenerative disease; development of tools to facilitate chemosensory research including mouse models of chemosensory dysfunction and improved neuroimaging, cell labeling, and axonal tracing techniques.

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301-402-3458, Fax: 301-402-0390
Email: millerr@nidcd.nih.gov

NIDCD Clinical Trials Topics:

NIDCD will accept applications for support of clinical trials in any of the areas noted above.

For additional information on research topics, contact:

Roger L. Miller, Ph.D.
National Institute on Deafness and Other Communication Disorders
301-402-3458, Fax: 301-402-0390
Email: millerr@nidcd.nih.gov

For administrative and business management questions, contact:

Mr. Christopher P. Myers
Grants Management Officer
National Institute on Deafness and Other Communication Disorders
301-435-0713, Fax: 301-451-5370
Email: myersc@nidcd.nih.gov
NATIONAL INSTITUTE OF DENTAL AND CRANIOFACIAL RESEARCH (NIDCR)

The NIDCR conducts and fosters research on the etiology, pathogenesis, prevention, diagnosis, and treatment of oral, craniofacial and dental diseases and conditions. For more specific information about areas of interest to the NIDCR, please visit our home page at http://www.nidcr.nih.gov.

NIDCR’s small business programs are highly focused on maximizing translational science opportunities – moving rapidly and translating basic dental and orofacial biology into useful products.

Research Topics of Interest to NIDCR

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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NIDCR Non-Clinical Trials Topics:

Developmental Biology and Mammalian Genetics and Genomics

Emphasis is on understanding the development of craniofacial complex and on the identification of the genetic and environmental contributions to craniofacial disorders. The objective of this scientific program is to elucidate the underlying causes of craniofacial disorders, thereby advancing the fields of diagnosis, treatment, and prevention. Interests in this area include but are not limited to:

A. Develop novel or improved imaging techniques and analysis methods that enable easy, cost-effective, accurate, and scalable deep phenotyping for research and/or diagnosis of dental, oral, and craniofacial conditions and disorders.

B. Develop novel or improved sample collection methods that offer superior features such as high sample quantity and quality for state-of-the-art multi-omics data generation and analyses of dental, oral, and craniofacial conditions; easiness of sampling, shipping, and storage; and overall cost effectiveness.

C. Develop early pregnancy genetic tests for genetic variants involved in inherited syndromic and non-syndromic craniofacial conditions and disorders.

D. Develop devices, including point of care units, to improve the diagnoses and treatment of inherited and acquired craniofacial conditions and disorders.

E. Develop advanced methods, assays, and reagents that allow flexible throughput to genetically engineer and functionally characterize organisms in craniofacial development and genetics studies.

Tools for Analysis of Genomic, Phenotypic, and Environmental Data

Emphasis is on development of data analysis tools that maximize the utility of existing data resources in basic and clinical research in the dental, oral, and craniofacial domains. Interests in this area include but are not limited to:

A. Develop bioinformatic and computational tools for integration and analyses of clinical, environmental, behavioral, phenotypic (including imaging and auditory), molecular, and multi-omics data that enable genetic studies, healthy lifestyles, disease prognoses, diagnoses, and treatment in dental, oral, and craniofacial domains.

B. Develop advanced analytics tools to retrieve data from diverse databases such as data repositories, literature, health records, etc. and infer hidden relations between data elements to inform basic and
clinical research in the dental, oral, and craniofacial domains. Tools that use artificial intelligence algorithms are considered highly relevant.

**Infectious Diseases and Immunity**

Research relating to the etiology, pathogenesis, prevention, diagnosis and treatment of infectious diseases of the oral cavity is supported by the NIDCR. This includes research on practical ways to effectively use the host immune system to prevent or treat oral infectious diseases and microbial-induced inflammation. Infectious diseases of the oral cavity include caries, periodontitis, candidiasis, peri-implantitis, pulpitis, and various viral, bacterial, and fungal infections of the oral mucosa. Research on the diagnosis and prevention of oral manifestations and malignancies of HIV infection and AIDS. Specific examples of technology development needs include but are not limited to:

A. Develop ways to overcome or eliminate the risk of oral infections in persons who smoke or chew tobacco, drink alcohol, or are immunosuppressed, have diabetes, are malnourished, or are psychologically stressed.

B. Explore novel methods or agents to eradicate oral biofilms (dental plaque) on teeth, oral soft tissues, and dental implants without adversely affecting the normal oral flora.

C. Isolate, synthesize or prepare new antibiotics and antimicrobial agents that can overcome bacterial and fungal resistance to current compounds. Formulate combinatorial drug regimens to attack microbes growing in oral biofilms (dental plaque).

D. Develop controlled release systems for local delivery of synthetic peptides, recombinant proteins, or other chemical or immunotherapeutic agents to prevent, control, and/or treat oral infectious diseases, or the oral manifestations of HIV infection.

E. Develop biological response modifiers or other immunological approaches to reduce or eliminate microbial-induced chronic inflammation or the tissue destruction associated with chronic inflammation in the oral cavity.

F. Develop ways to interfere with microbial colonization and growth through the use of antimicrobial agents and chemotherapy.

G. Identify and exploit the structural features of oral biofilms for increased therapeutics delivery.

H. Develop computer programs and apply systems biology approaches to model biologically active peptide regions of oral components that have anti-fungal, anti-bacterial and anti-viral activities.

I. Develop substitutes of naturally occurring chemicals (phytochemicals) known to have a role in controlling opportunistic infections induced by HIV.

J. Develop synthetic peptides and recombinant proteins of oral components with anti-fungal, anti-bacterial and anti-viral activities including those against HIV and oral opportunistic pathogens.

K. Develop oral topical formulations with combined microbicide, analgesic, and anti-inflammatory activities to enhance oral mucosal defenses and prevent and/or control oral infections and lesions in HIV-infected and/or immunosuppressed subjects.

L. Discover, test, standardize, and validate novel biomarkers present in oral biospecimens for screening and clinical diagnosis of HIV, and oral opportunistic pathogens infections and AIDS malignancies. Apply similar strategies as listed below for oral, oropharyngeal and salivary gland cancers to AIDS malignancies.

M. Develop the next generation of rapid tests and point of care devices to detect, quantify, screen, and diagnose HIV and oral opportunistic pathogens. Develop novel assays to quantify oral mucosal reservoirs for oral viruses, oral immune responses to viral prophylactic and therapeutic vaccines, and viral changes due to anti-viral treatments.
N. Develop safe and effective targeted diagnostic and therapeutic technologies in response to endemic and pandemic infections

**Oral, Oropharyngeal and Salivary Gland Cancers**

Emphasis is on molecular mechanisms of oral epithelial cell deregulation that lead to oral cancers. Research related to early detection, diagnosis, and prediction, and treatment of oral cancers is of particular interest. Examples include but are not limited to the following areas:

A. Develop imaging techniques for the early detection, diagnosis and prognosis of pre-malignant lesions.
B. Develop effective pharmacological, immunological and radiological modalities for treatment of pre-malignant and malignant lesions in preclinical models.
C. Develop novel technologies for the genetic and molecular-targeted therapy (e.g. siRNAs, peptide based therapies) in preclinical models.
D. Develop genetic animal models of oral cancer premalignancy and oral cancer progression that mimic human oral cancers, including HPV-associated oropharyngeal cancers.
E. Develop animal models to facilitate the testing of therapeutic and chemopreventive agents for oral cancers.

**Temporomandibular Disorders and Orofacial Pain**

Emphasis is on research for chronic disabling painful diseases of the oral-craniofacial-dental areas including chronic pain, neuropathies, and diseases of the temporomandibular joint. NIDCR encourages applications that include but are not limited to:

A. Develop improved methods and technologies for measuring nociceptive, chemosensory, tactile, kinesthetic, or proprioceptive function involving craniofacial structures. Such measures may be useful in screening for deficits, improving diagnosis, or for evaluating responses to orofacial treatments or interventions.
B. Develop improved biomarkers for neuropathic pain conditions affecting oral-craniofacial tissues or structures.
C. Develop assays facilitating reliable evaluations of relationships between biological and other risk factors as they relate to onset, and exacerbation of pain and for examining transition from acute pain to chronic pain conditions.
D. Identify and develop novel pharmacologic or biologic agents, including but not limited to small molecules, peptides, recombinant proteins and nucleic acids to prevent, control, and/or treat orofacial pain.
E. Develop animal models to facilitate testing of therapeutic agents for orofacial pain.

**Saliva, Salivary Diagnostics, and Salivary Gland Diseases**

Emphasis is on salivary gland physiology and pathophysiology and in the repair and restoration of the damaged gland. Examples include but are not limited to:

A. Develop viral, non-viral and gene therapy-based approaches to address compromised salivary gland function. Develop cell and tissue-based strategies and technologies for restoration of damaged or destroyed salivary gland function.
B. Develop novel compounds or materials that protect and preserve salivary glands from head and neck cancer irradiation therapy.
C. Develop non-invasive methods for the determination of efficacy and safety of artificial saliva, sialogogues, and their delivery vehicles used in addressing the diminution or lack of saliva (xerostomia) due to Sjögren’s Syndrome or head and neck cancer irradiation therapy.

D. Develop biomarker-based technologies for the identification of Sjögren’s Syndrome using blood or saliva as body fluids.

E. Identify biomarkers derived from oral fluids that are predictive of the onset, progression and recurrence of oral diseases and conditions, such as periodontal diseases, caries, and oral, oropharyngeal and salivary gland cancers.

F. Develop immunological strategies and immunotherapy-based approaches for addressing xerostomia from Sjögren’s Syndrome.

G. Improve the existing or develop new tools for early detection of salivary gland cancers.

**Biotechnology, Biomaterials, and Applications for Regeneration and Restoration of Oral, Dental and Craniofacial Tissues**

Emphasis is placed on the development of a broad range of technologies targeted at regeneration and restoration of diseased and injured hard and soft tissues of the oral and craniofacial complex and on translating these applications to the clinic. Tissues of interest include craniofacial and alveolar bone, the periodontal ligament, TMJ bone and cartilage, oral mucosa, facial skeletal muscle, vasculature and nerves. Also of interest are multi-tissue composites and organs, such as vascularized and innervated bone and muscle, salivary gland, tooth, periodontium, bone-periodontal ligament-cementum interface and osteochondral complexes. Specific topics could include but are not limited to:

A. Develop technologies for design, fabrication, and manufacturing of biomimetic and biocompatible biomaterials and scaffolds, including nanomaterials and self-assembling nano-scaffolds, for tissue engineering and regenerative medicine applications. Projects need to include assessments demonstrating the ability of biomaterials and scaffolds to support generation and regeneration of mineralized tissues that replicate the mechanical, physical and biological properties of dentin, enamel or bone.

B. Develop cell-based technologies, including stem cell-based technologies. These include, designing strategies for isolation, purification, differentiation, scaled up production, manufacturing, standardization and quality control of stem and progenitor cells and their differentiated progenies, derivation of efficient and predictable methodologies for cellular reprogramming, and advancing technologies for reconstruction of stem cell niches for augmenting tissue regeneration.

C. Develop bioreactor systems to facilitate design, fabrication, and manufacturing of soft and hard tissues of dental, oral and craniofacial complex. These bioreactors may be able to mimic biophysical forces, such as mechanical and electrical forces that normally guide tissue morphogenesis in vivo. Among other desirable features of the bioreactors are maintenance of tissue construct oxygenation and real-time tissue imaging capabilities.

D. Develop improved dental composite materials, including biomimetic and self-healing materials and adhesive sealants. These include but are not limited to materials to replace Bis-GMA resin-based systems that are suitable for restoring crowns of posterior teeth and exposed roots of the teeth. Any novel dental composite restorative components or systems must include assessments in a physiologically relevant test system that mimics microbial and physicochemical conditions found in the oral cavity.

E. Develop methods, materials, and devices for orthodontic, prosthetic, periodontic, endodontic and craniofacial applications including those that can be used for craniofacial bone distraction, reconstruction, hard and soft craniofacial tissue healing and regeneration, and scarless craniofacial tissue repair.
F. Develop miniaturized artificial tissue and organ mimics/tissue chips and organoids that can be adapted to high-throughput formats for a broad range of applications, such as analysis of biomaterial and tissue function, drug efficacy and toxicology assays, biocompatibility assays, genetic screening and elucidating mechanisms of dental, oral and craniofacial development and disease.

G. Develop mathematical, computational, and bioinformatics approaches for modeling oral and craniofacial tissues and organ function and physiology to address needs of system biology, synthetic biology, and single cell analysis.

H. Develop new approaches for utilizing novel biomolecules, including growth factors, cytokines, small molecules, siRNAs, and others for counteracting diseases and injuries of oral and craniofacial tissues and promoting their healing and regeneration.

I. Develop new approaches to study molecular or cellular interactions between hard and soft tissues such as between the nervous system and mineralized tissues. Approaches can include development of new technologies or application of existing technologies that are newly applied to the dental and craniofacial field.

J. Develop advanced viral and non-viral based biomolecule delivery approaches, including nanotechnology-based technologies that can precisely deliver and release therapeutic proteins, nucleic acids, small molecules, or combinations thereof with predictable temporal kinetics to target specific tissue sites.

K. Develop imagining diagnostics to accelerate clinical implementation of reliable, reproducible, highly specific and sensitive diagnostic instruments for various applications, including but not limited to dental caries, cracked teeth, pulp vitality, bone quality, and periodontal disease.

L. Develop safe and effective biosensors for noninvasive, dynamic real-time monitoring of physiological processes in the human body using the oral cavity as the sensing site. These biosensors will be able to assess health and disease states and receive feedback from body fluids and clinical compounds that are found in or pass through the oral cavity and in certain cases, will be able to communicate these outputs wirelessly and remotely.

M. Develop safe and effective biosensors, monitoring devices and systems, data driven and computer science tools for automated detection, diagnosis and treatment of dental, oral and craniofacial disease.

Preclinical Research

A. Preclinical research and development activities for dental and craniofacial technologies (including devices, diagnostic instruments, reconstructive materials, pharmaceuticals, therapeutics, vaccines and biologics) that require review and approval by the FDA as a regulated product before commercial distribution.

NIDCR Clinical Trials Topics:

Special statement regarding clinical trials:

Projects proposing a clinical trial within STTR/SBIR applications must provide detailed justification that funds available through these awards can adequately support the trial especially if the trial is testing a drug, device or biologic under an investigational new drug (IND) or investigational device (IDE) application. The cost and time needed to plan and deploy most Phase II and almost all Phase III clinical trials would exceed the support and project period provided under this program.

In addition to the specific clinical trial topics listed below, we also would consider clinical trials for certain program topics listed above. Products originally developed and preliminarily tested with SBIR/STTR support also can be pursued further with funding mechanisms that support clinical trials, see:
Biomedical Clinical Research

Emphasis is on development of methods, drugs and materials to diagnose or treat oral and craniofacial diseases and conditions. Areas of interest include but are not limited to projects that:

A. Develop improved methods to detect and predict progression of dental caries, periodontal disease, reversible and irreversible pulpitis.
B. Develop improved methods or materials to prevent dental, oral, and craniofacial diseases or conditions.
C. Develop new or improved methods or materials to enhance oral and craniofacial surgery. This would include both intraoral and extra-oral surgery.
D. Develop improved methods or materials to mechanically and/or biologically repair or treat tooth structure damaged by dental caries or periodontal disease.
E. Develop improved appliances to aid suckling by newborn infants with cleft palate and cleft lip.
F. Develop safe and efficacious methods to diagnose caries, pulp vitality and / or periodontal diseases utilizing non-ionizing radiation.
G. Develop technologies for local delivery of drugs to treat oral and craniofacial diseases or disorders.
H. Develop novel non-opioid pharmacological medications for management of acute dental pain.
I. Develop safe and efficacious methods or medications to manage complications of head and neck cancer treatment.
J. Develop tools for implementation of precision medicine in the oral cavity.
K. Develop methods and tools to detect soft tissue pathologies in the oral cavity.
L. Develop oral devices and materials for monitoring local and systemic conditions.

Behavioral Clinical Research

Provides support for the development of evidence-based products related to behavioral and social aspects of oral health, oral health prevention or treatment interventions, and other patient-oriented aspects of oral health. This includes support for clinical trials and patient-oriented research to establish safety and initial efficacy of products. NIDCR is especially interested in applications that significantly improve oral health by: 1) being broadly applicable to many populations, 2) contributing to meaningful oral health improvements for a specific population, 3) expediting translation of research findings into oral health improvements, and/or 4) equipping oral health care providers, educators or researchers with tools to improve public oral health. Examples of studies of interest include, but are not limited to, the following:

A. Develop and test devices or methods to improve time-sampled monitoring of behavioral adherence with preventive or therapeutic regimens specifically relevant to oral diseases/conditions. Such devices or methods could be utilized in a variety of settings, including naturalistic settings, within clinical trials, within oral health care delivery systems, etc.
B. Develop and test novel compliance and survey measures or tools to identify the underlying causes of insufficient preventive dentistry for specific underserved populations.
C. Develop, or adapt for use in a new population or setting, novel measures or methods for identifying individual, family, group, or other processes that explain oral health behavior.
D. Develop and test for safety, efficacy, and/or effectiveness of measures or materials for diagnosing, preventing, or treating oral, dental, and craniofacial conditions and disorders.
E. Develop, or adapt for use in a new population or setting, oral health interventions utilizing technology to improve efficiency of delivery (e.g., management of chronic pain related to temporomandibular joint disorders, etc.).

F. Develop, or adapt for use in a new population or setting, interventions addressing health behaviors highly associated with oral health (e.g., tobacco, alcohol, and other drug use; management of diabetes, HIV infection, or other chronic illnesses; etc.).

G. Develop technologies or modules that utilize existing web-based platforms to improve preventive oral health hygiene for children and adolescents (e.g., social marketing via web-based interaction, virtual reality "worlds", "massively multiplayer online games", etc.).

H. Develop and test innovative methods for facilitating collaborations, referrals, and/or ongoing follow-ups between oral health professionals and other health care professionals.

I. Develop and test web-based training or other innovative approaches for oral health care professionals to accelerate accurate translation of new knowledge regarding oral diseases and their effective prevention or treatment into clinical or public health practice.

J. Develop and test the effectiveness of innovative teaching tools to inform oral health professionals or the public regarding oral cancer prevention and early detection.

K. Develop and test the effectiveness of innovative teaching or educational tools or curricula to inform oral health professionals and dental students regarding the role of genetics and genomics, including the oral microbiome, in oral diseases and conditions and in oral health care.

For additional information on research topics, contact:

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NATIONAL INSTITUTE OF DIABETES AND DIGESTIVE AND KIDNEY DISEASES (NIDDK)

The mission of the National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) is to conduct and support medical research and research training and to disseminate science-based information on diabetes and other endocrine and metabolic diseases; digestive diseases, nutritional disorders, and obesity; and kidney, urologic, and hematologic diseases, to improve people’s health and quality of life. For additional information about areas of interest to the NIDDK, please visit our home page at [http://www.niddk.nih.gov](http://www.niddk.nih.gov). See our SBIR/STTR page at [https://www.niddk.nih.gov/research-funding/research-programs/small-business](https://www.niddk.nih.gov/research-funding/research-programs/small-business).

Limited Amount of Award

For budgetary, administrative, or programmatic reasons, the NIDDK may not fund an application or may decrease the length of an award and/or the budget recommended by a review committee.

Per current SBA-defined hard caps, total funding support (direct costs, indirect costs, fee) normally may not exceed $256,580 for Phase I and $1,710,531 for Phase II.

With appropriate justification from the applicant, the NIDDK may consider budgets that exceed these amounts to support research that aligns with an approved waiver topic (see APPENDIX A: National Institutes of Health SBA-Approved SBIR/STTR). Topics for Awards over Statutory Budget Limitations.

The NIDDK generally considers:

- Phase I budgets up to $300,000 total costs or project periods up to 2 years.
- Phase II applications up to $2,000,000 total costs or project periods up to 3 years (Phase II budgets generally should not exceed $1,000,000 total costs in any year).
- Phase IIB (see below) applications up to $3,000,000 total costs or project periods up to 3 years (Phase IIB budgets generally should not exceed $1,000,000 total costs in any year).

Applicants considering a requested budget greater than these limits are strongly encouraged to contact program staff before submitting an application.

The NIDDK also participates in the SBIR/STTR Commercialization Readiness Pilot (CRP) Program. Applicants should review the Award Budget section of relevant funding opportunity announcements. For Phase II awardees, especially those developing products that require clinical evaluation or approval by a Federal regulatory agency, the NIDDK strongly encourages potential applicants to apply to NIDDK’s Phase IIB program.

Phase IIB Competing Renewal Awards

NIDDK will accept Phase IIB SBIR/STTR Competing Renewal grant applications (only) from NIDDK-supported Phase II awardees that propose to continue the process of developing products that ultimately require 1) clinical evaluation, 2) approval by a Federal regulatory agency, and 3) continuing refinements to durable medical equipment (DME) designs such as cost reduction, testing for safety, durability, and reliability, and meeting or establishing standards. This renewal grant should allow small businesses to get to a stage where interest and investment by third parties is more likely. Such products include, but are not limited to biological products, devices, drugs, medical implants, etc. related to the mission of the NIDDK. These awards are intended to support completion of research needed to obtain an IND or IDE. Applicants must provide evidence that they have consulted formally with the FDA concerning the research needed for the development of a drug, biologic or medical device and that the proposed research will address these regulatory requirements. Such evidence should include FDA correspondence from a pre-IND...
meeting for an IND application or a pre-IDE meeting for an IDE application, and the status of the project in a timeline related to Federal regulatory approval processes.

Examples of research that would be considered responsive to this announcement are listed below for illustrative purposes and are not exclusive of other appropriate activities.

- Completion of studies as required by the Food and Drug Administration (FDA) for Investigational New Drug (IND) or Radioactive Drug Research Committee (RDRC) application.
- Assessment of devices with regard to performance standards related to the FDA approval process.
- Clinical studies in support of an application for clearance or approval by the FDA. See the table below and review NIDDK’s Policies for Clinical Researchers ([https://www.niddk.nih.gov/research-funding/human-subjects-research/policies-clinical-researchers](https://www.niddk.nih.gov/research-funding/human-subjects-research/policies-clinical-researchers)) when considering an application involving human subjects.

Final Progress Reports

As detailed in NOT-OD-17-085, the NIH has implemented the Final Research Performance Progress Reports (Final RPPR) for SBIR/STTR Final Progress Reports.

The NIDDK is interested in tracking the progress of the small business concerns it funds and the products they develop. Funding priority will be given to those small business concerns that show not only their ability to develop products but also their growth as a small business concern towards independence from the SBIR/STTR program.

Additional Programs and Services for NIDDK SBIR/STTR Awardees

The NIDDK encourages awardees to apply to participate in NIH Technical Assistance Programs ([https://sbir.nih.gov/tap](https://sbir.nih.gov/tap)). The NIDDK may offer additional programs throughout the year, and awardees are encouraged to keep their contact information current so that they receive announcements regarding these programs.

Research Topics of Interest to NIDDK

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
<th>No*</th>
<th>Yes**</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes**</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>

*If No,
- This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, UNLESS the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,
- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.
<table>
<thead>
<tr>
<th>Does IC/Office/Agency accept clinical trials applications under this mechanism?</th>
<th>Yes</th>
<th>No</th>
<th>Other Information:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Trials SBIR/STTR Omnibus/Parent Funding Opportunity Announcement/s</td>
<td>X</td>
<td></td>
<td>The NIDDK accepts SBIR, but not STTR applications with NIH-defined clinical trials.</td>
</tr>
<tr>
<td>Clinical Trials SBIR/STTR IC/Office/Agency - Specific Funding Opportunity Announcement/s</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply</td>
<td>X</td>
<td></td>
<td>Small businesses are eligible to apply for several non-SBIR/STTR funding opportunities. Comprehensive information on Human Subjects Research at NIDDK can be found here: <a href="https://www.niddk.nih.gov/research-funding/human-subjects-research">https://www.niddk.nih.gov/research-funding/human-subjects-research</a></td>
</tr>
</tbody>
</table>

The NIDDK has multiple R01 funding opportunities, spanning pilot & feasibility clinical trials to clinical trials with one or two research centers.

Applications for clinical trials requiring three or more research centers should be submitted to NIDDK multi-center clinical study implementation planning cooperative agreement (U34) and NIDDK multi-center clinical study cooperative agreement (U01) FOAs.

Clinical Trials Allowed and Optional funding opportunities to which NIDDK is subscribed can be found on NIDDK’s Current Funding Opportunities page: [https://www.niddk.nih.gov/research-funding/current-opportunities](https://www.niddk.nih.gov/research-funding/current-opportunities).

**NIDDK Non-Clinical Trials Topics:**

**Diabetes, Endocrinology and Metabolic Diseases**

The Division of Diabetes, Endocrinology and Metabolic Diseases supports basic and clinical research on the etiology, pathogenesis, prevention, diagnosis, and treatment of diabetes mellitus and its complications of diabetic wound healing and diabetic neuropathy; thyroid and adrenal diseases; and rare metabolic
disorders; as well as research on basic endocrine and metabolic processes. Research topics of potential interest to small businesses include, but are not limited to:

**I. SENSORS, HORMONE REPLACEMENT, AND DELIVERY DEVICES:**

A. Assessment of non-invasive, minimally invasive or implantable sensors for monitoring blood or interstitial fluid glucose for prevention of hypo- and hyperglycemia in diabetic patients. NIDDK will give priority to research that has already progressed to an *in vivo* model or to be clinically tested.

B. Integration of glucose sensor and hormone delivery systems to create an artificial pancreas.

C. Development of improved insulin and other pancreatic hormone delivery systems or devices.

D. Development of novel insulin and glucagon formulations showing improved kinetics and stability.

E. Development of novel and more accurate non-enzymatic based glucose detection technologies.

F. Development of telemedicine/remote monitoring approaches that can be incorporated as components/and or adjuvants of an artificial pancreas for better diabetes self-management.

G. Development of technologies that may promote and facilitate adherence/compliance by users of glucose monitoring and control devices.

H. Development of biomaterials that can deliver drugs or biologics to a diabetic foot ulcer to improve healing.

I. Development of sensors, algorithms and patient interfaces that can provide feedback to diabetic individuals with insensate feet to prevent or off-load diabetic foot ulcers.

**II. SCREENING TESTS, DIAGNOSTICS AND BIOLOGIC TOOLS:**

A. Development of techniques or products/biomarkers useful for predicting, preventing or delaying progression of diabetes, including tests for identifying patients at risk, and methods of monitoring disease progression.

B. High throughput - Point of care technologies (reliable, accurate, cost-effective, highly sensitive, standardized having rapid turnaround time) for autoantibody detection, T cell -subsets-auto-reactivity and other immune parameters for autoimmune diabetes diagnosis and follow-up.

C. Development of methods to measure changes in the immune status that may be used as markers to follow the immune-modulatory activity and beneficial effect (beta cell mass preservation, reduction of inflammation at the target organ, etc.,) of biologic agents tested in clinical trials for the prevention and/or treatment of T1D.

D. Development of high throughput assays based on biologic pathways likely involved in the pathogenesis of diabetes and its complications.

E. Development of methods/techniques/assays to measure adipose tissue in different depots in humans, including marrow fat.

F. Develop validated, highly sensitive, and specific assays for glucagon detection.

G. Development of reagents for assessment/manipulation of glucagon receptor activity in the pancreatic islet and other tissues.

H. Reagents to be used to improve our understanding of pancreatic alpha-cell biology particularly in response to glucagon and incretins.

I. Reagents and assays for the accurate expression and quantitation of functional incretin receptors in the endocrine/exocrine cells of the pancreas and relevant extrapancreatic tissues.
J. Development of materials and technologies for the support of microphysiological platforms used for pre-clinical testing and/or modeling of physiological and pathophysiological aspects of diabetes and metabolic disorders.

K. Development and validation of tools for use by health care providers/systems to improve diabetes care and prevention.

L. Development of techniques and tools to identify islet cell progenitors, methods to predict transplant success with recovered islet preparations, and non-invasive imaging as well as other methods for the \textit{in vivo} measurement/evaluation of pancreatic beta cell mass, function or inflammation after transplantation of pancreatic islet/beta cells.

M. Point of care low cost/portable technologies for diabetes and pre-diabetes diagnosis.

N. Development of innovative technologies to accurately measure and to predict and prevent hypoglycemia.

O. Development of clinical measures of the molecular and cellular damage from diabetes, such as oxidative stress, advanced glycation end-products and chronic inflammation that can be used as predictive or diagnostic biomarkers for diabetes complications.

P. Development and validation of biomarkers to monitor disease progression and response to therapy for diabetic neuropathy, including peripheral sensory, autonomic and painful diabetic neuropathy.

Q. Development of diagnostic and predictive biomarkers for diabetic foot ulcers that can be used to diagnose infections and biofilms, predict healing, select treatment strategies or determine risk of primary or secondary occurrence of foot ulcers.

III. INTERVENTIONS AND THERAPIES:

\textbf{Diabetes}

A. Development of immunomodulation/tolerance induction strategies to prevent or slow progression of type 1 diabetes.

B. Development of methods that protect islet grafts after transplantation with no need of immunosuppression, including alternative transplantation sites, immunoisolation strategies, and immunomodulation/tolerance induction.

C. Development of reproducible methods that improve yield/viability/function of islets and allow ex-vivo expansion of human islets/insulin producing cells while still retaining appropriate functional islet characteristics and the ability to be successfully transplanted.

D. Development of methods utilizing replenishable cell sources, especially stem cells that produce functional islet like cells/tissues that can be successfully transplanted or tested in microphysiological systems and/or \textit{in vivo} models of the disease.

E. Development of educational or psychosocial approaches that increase adherence to recommend diabetes treatment regimens.

F. Development of novel technologies that may facilitate self-management of diabetes and adherence to treatment.

G. New implantable and easy to replace technologies that may mimic the beneficial effect of gastric bypass/bariatric surgery for the treatment of diabetes without the need of a major invasive surgical procedure.

H. Development of educational approaches and new technologies that increase adherence to preventative measures for diabetic foot ulcers in high risk patients or increase adherence to off-loading and other recommended treatment regimens for diabetic foot ulcers.
I. Development of new therapies or devices to prevent and treat diabetic foot ulcers and neuropathy, including autonomic and painful diabetic neuropathy.

J. Development of new therapies to correct the underlying metabolic defects that result from diabetes, such as reactive oxygen species production and glycation of proteins.

K. Development of therapies that can prevent or mitigate episodes of hypoglycemia.

**Other Endocrine and Metabolic Disorders**

L. Identification of new ligands for previously unclassified (orphan) nuclear receptors and development of partial agonists or antagonists with therapeutic potential for obesity and endocrine diseases within the mission of NIDDK.

M. Development of Selective Receptor Modulators (SRMs) with tissue specificity and profiles that provide beneficial effects without the side effects secondary to therapies based on naturally occurring hormones.

N. Development of novel diagnostic and therapeutic strategies for autoimmune endocrine disorders.

**Digestive Diseases and Nutrition**

The Division of Digestive Diseases and Nutrition supports research in diseases and disorders of the digestive tract; esophagus, stomach, intestine, colon, anorectum, pancreas, liver, gallbladder, and biliary tract; as well as research in nutrition and obesity. Innovative investigator-initiated projects that are not mentioned below are also encouraged. Examples of areas that may be of interest to small businesses include, but are not limited to:

**I. Gastrointestinal**

A. Development of new diagnostic techniques and tests, including non-invasive tests and imaging for detecting Barrett’s esophagus, GERD, and other intestinal disorders.

B. Development of agents and techniques to measure, diagnose, stimulate regeneration of enteric neurons, and treat motility disorders.

C. Development of novel therapies to modulate/enhance GI lymphatic function for the treatment of GI pathologies.

D. Development of gut-derived biomarkers of neurodegenerative brain disease.

E. Development of techniques or modulators of neuroimmune interactions that target functional bowel disorders or inflammatory disease.

F. Development of novel proteomic or metabolomic technologies designed to study digestive diseases and their complications.

G. Development of assays and screening methods for detection of biomarkers for diagnosis, grading and staging digestive diseases.

**II. Liver**

A. Development of novel antifibrotic therapies for chronic progressive liver diseases.

B. Development of quantitative tests of hepatic “reserve” for assessment of therapeutic intervention, transplantation, or surgical risk in patients with liver disease.
C. Development of point-of-care, serologic, and rapid tests for rapid diagnosis, treatment requirements and genotyping of hepatitis.

D. Development of rapid, reliable and inexpensive tests for genetic screening and risk markers important in liver disease.

E. Development of sensitive and reliable non-invasive techniques to detect and monitor liver fibrosis and other chronic liver diseases and the associated complications.

F. Creation of bio-artificial organs for temporary hepatic support in patients with acute liver failure.

III. Pancreas


B. Development of more accurate, non-invasive approaches to the diagnosis of chronic pancreatitis by functional, radiologic, endoscopic, or pathologic/cytologic means.

IV. Nutrition/Obesity

A. Development of novel methods and tools to accurately evaluate nutritional status, physical activity, and energy expenditure.

B. Development of non- or minimally invasive technologies that allow access and/or delivery to discrete regions of the digestive tract.

C. Development of novel breath, urine, or blood tests to accurately measure dietary intake.

D. Development of technologies to detect, prevent, and treat acute gastrointestinal infections by foodborne pathogens.

Kidney, Urologic and Hematologic Diseases

The Division of Kidney, Urologic, and Hematologic Diseases provides research funding and support for basic, translational, and clinical research studies of the kidney, urinary tract, and disorders of the blood and blood-forming organs. Projects may include development of tools to improve understanding of the physiology, pathophysiology, and related diseases of the kidney, genitourinary tract, and blood and blood forming systems, or to develop rational diagnostics, treatments, and prevention strategies for these diseases. Projects may be to develop tools/technologies to support clinical care, population health and/or pragmatic research to improve health outcomes in populations with kidney diseases and/or urologic conditions. Projects to develop technologies that will enhance research in kidney, urologic and hematologic diseases are encouraged. Development of –omics, bioinformatics, and multi-scale technologies for the study of these systems, especially where these systems interact, is also encouraged. Research opportunities that may be of interest to small businesses include, but are not limited to:

I. Kidney Diseases

Areas of research include chronic kidney disease, end-stage renal disease, diabetic nephropathy, polycystic kidney disease, hypertensive nephrosclerosis, acute kidney injury, kidney donation (delayed graft function and chronic rejection), congenital kidney disorders, glomerular and tubulointerstitial diseases, IgA nephropathy, hemolytic uremic syndrome, fluid and electrolyte disorders, kidney repair and regeneration, and normal and abnormal kidney development and physiology.

Dialysis, Devices and Medical Technologies
A. Development of innovative forms of renal dialysis which improve efficiency and/or have lower associated morbidity (e.g., tissue engineered artificial kidneys, implantable or wearable dialyzers).

B. Development of pharmacological agents, devices, techniques, or diagnostics that enhance maturation and longevity of a vascular access.

C. Development of dialysis membrane technologies with enhanced biocompatibility and anti-fouling properties.

D. Development of a means to provide continuous anticoagulation to permit renal replacement therapy.

E. Development of reliable, non-invasive, online monitoring systems for real-time assessment of treatment parameters such as blood volume, access flow, and urea clearance.

F. Development of new agents for sterilizing dialysis membranes and development of agents or methods to reduce catheter-related infections in hemodialysis or peritoneal dialysis.

G. Development of hemodialysis or peritoneal dialysis catheters using improved biomaterials, which decrease the foreign body response, biofouling, and biofilm formation.

H. Development of devices or techniques to enhance the long-term success of kidney transplantation (e.g., techniques for kidney storage and preservation).

I. Development of technologies to improve kidney biopsies (i.e., to improve safety or tissue acquisition).

Health Information Technologies

J. Development of health information technologies or mobile technologies that enhance delivery of care, population health management, and/or research for patients with kidney diseases.

K. Development of applications or application programming interfaces that use health data standards (e.g., Fast Healthcare Interoperability Resources [FHIR], clinical terminologies) to improve accessibility, accuracy, and/or completeness of real-world data for research and care of individuals with kidney diseases.

Diagnostics and Imaging

L. Development of clinical assays that enable precision medicine approaches to treating kidney diseases.

M. Development of non- or minimally-invasive methods for evaluating kidney function, including in individuals with congenital genitourinary conditions.

   1. Reliable, non-invasive, non-radioactive methods of measuring glomerular filtration rate (GFR).
   2. Translation of biomarkers of acute kidney injury or chronic kidney disease with clinical utility into commercial assays.
   3. Translation of biomarkers for early detection of kidney diseases or prediction of kidney disease progression, recovery, or drug response.

N. Development of improved renal imaging techniques, differential renal function assessment, diagnostic assessment of non-malignant kidney diseases, or measurement of perinatal nephron endowment.
O. Development of technology to improve collection of real-time data (e.g., biomarkers, diet, physical activity, patient reported outcomes, vital signs, patient experience of kidney or urologic disease or its treatment, environmental factors which affect the development or progression of kidney disease), patient outcomes, and adherence for clinical studies.

P. Development of imaging or molecular analysis technologies to enhance information extraction from renal biopsies and development of antibodies or other probes for unique cell types of the kidney.

**Therapeutics Discovery and Development**

Q. Lead optimization and preclinical development of pharmacological agents that might be used to intervene in acute or chronic renal disorders and in disorders of renal hemodynamics, blood pressure, electrolyte metabolism, and extracellular volume regulation.

R. Development of drugs or biologics to stimulate productive kidney repair or regeneration.

S. Development of functional nephrons for transplantation.

T. Development of technologies to enhance the validation of kidney disease targets or to screen compounds for efficacy or toxicity (e.g., kidney organoids or tissue chips, more relevant animal models of acute kidney injury).

U. Development of data and cell banks (e.g., of diabetic kidney disease families and polycystic kidney disease families) for use by the research community.

V. Development of preventative measures for acute kidney injury (e.g., during coronary artery bypass grafting, sepsis, or treatment with nephrotoxic agents).

II. UROLOGIC DISEASES

Areas of research include benign prostatic hyperplasia, lower urinary tract symptoms (LUTS) including urinary incontinence, urinary tract infections, urinary stone disease, erectile dysfunction, urologic chronic pelvic pain syndromes (including interstitial cystitis and chronic prostatitis), congenital urologic disorders, repair and regeneration of lower urinary tract organs, and normal and abnormal lower urinary tract development, and genitourinary physiology.

**Diagnostics and Imaging**

A. Translation of blood or urine biomarkers in the lower urinary tract or other urologic disorders into commercial assays with clinical utility.

B. Development of non-invasive or minimally-invasive methods to diagnose bladder inflammation or changes in the urothelium that are not of a cancerous origin.

C. Development of new technologies or methods with reduced radiation dose for evaluating vesico-ureteral reflux in children and infants.

D. Development of diagnostic modes to clinically and non-invasively or minimal-invasively measure bladder outlet obstruction before and after surgical or pharmaceutical intervention.

E. Development of objective diagnostic devices or methods for the assessment of urinary storage and voiding disorders, including stress, urge, and mixed incontinence, in both adults and children.
F. Development of wireless and non-invasive or minimally-invasive measurement technologies for real-time assessment of lower urinary tract function, which can include neuro-pharmacological/neuro-physiological urodynamics.

G. Development of radiation-free and accurate imaging technologies for urinary stone disease.

**Drug and Device (Therapeutic) Interventions**

H. Lead optimization and preclinical development of pharmacological agents for treatment or prevention of urinary stone disease, urological chronic pelvic pain syndromes, urinary tract infections, or other benign urologic diseases or conditions.

I. Development of novel neuromodulation devices, which restore function or mitigate pain conditions of the lower urinary tract.

J. Development of urinary catheters which reduce the incidence of infection in the urinary tract and decrease urethral and bladder inflammation.

K. Development of technologies for treatment of bladder outlet obstruction.

L. Development of health information technologies or mobile/wireless technologies that enhance delivery of care for patients with benign urologic diseases or conditions, including transition in lifelong care of congenital genitourinary conditions.

M. Development of bioengineered materials or structures, including cell-laden structures, for the repair or regeneration of genitourinary organs.

**Research Tools**

N. Development of tools for elucidating the role of urinary or gut microbiome in urinary stone disease or other benign urologic diseases or conditions.

O. Development of novel models of benign prostatic hyperplasia.

P. Development of technology to improve collection of real-time data (e.g., biomarkers, diet, physical activity, vital signs, psychological parameters, and environmental factors), patient-reported outcomes, and adherence for clinical studies (e.g., studies of gene-environment interactions in the manifestation of urologic diseases).

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**III. HEMATOLOGIC DISEASES**

The NIDDK hematology research program focuses on understanding basic cellular and molecular mechanisms that underlie the production and function of blood cells in health and disease. The program emphasizes translational applications of new insights and knowledge gained from basic research in these areas toward the development of novel or improved approaches for the diagnosis, stratification, and treatment of hematologic diseases. This includes the development of disease biomarkers, gene targeted therapies, hematopoietic stem cell transplantation in heritable blood diseases (e.g., sickle cell disease, thalassemia, hemochromatosis, hemoglobinopathies, iron overload, anemia, and cytopenia), and the measurement and chelation of tissue iron in iron overload disorders. The NIDDK hematology research program provides resources for basic and preclinical development efforts leading up to IND or IDE submissions but does not fund clinical trials. The program has a particular focus on myeloid lineage and hematopoietic stem cells, including the effects of aging on hematopoiesis.

**Drug Discovery and Development**
A. Establishment of robust in vitro or animal models of benign hematologic diseases for drug discovery or development.

B. Development of therapeutics that target elements of hematopoietic stem cell niches (e.g., stromal cells, osteoblasts, endothelium, macrophages, pericytes, nerve cells).

C. Development of therapeutics that modulate blood cell production from hematopoietic stem cells and progenitors based upon understanding of physical and chemical regulatory pathways.

D. Development of therapeutics that modulate metabolism, storage, and transport of iron.

**Cell Therapies**

E. Development of equipment, chemically-defined reagents, and methods for high volume ex vivo expansion, isolation, and/or differentiation of highly purified human hematopoietic stem and progenitor cells.

F. Development of therapeutics that induce fetal hemoglobin synthesis by chemical means, genome editing, or other means.

G. Development of therapeutics that target blood cell membrane structure.

**Diagnostics and Imaging, Medical Technologies, and Research Tools**

H. Development and validation of sensitive, specific, reproducible, quantitative, and clinically applicable assays for measuring levels or expression of iron regulatory molecules (e.g., hepcidin).

I. Development of technologies to track, purify, monitor or assay single-cells in vivo or in vitro.

J. Development of non-invasive systems for monitoring circulating blood cells, blood chemistry or blood cell production.

K. Development of imaging technology for the non-invasive measurement of bone marrow cellularity and function.

L. Development of imaging technology for the non-invasive measurement of tissue iron loading and distribution.

M. Development of technologies to understand the roles of mitochondria in benign hematologic diseases.

**NIDDK Clinical Trials Topics:**

NIDDK will accept clinical trials in most of the areas listed above in the non-clinical trials topics. NIDDK does not support clinical trials in hematologic diseases.

For additional information on research topics, contact:

**DIABETIC TECHNOLOGY, TYPE 1 DIABETES AND ENDOCRINE DISEASES**

Dr. Guillermo Arreaza-Rubín
National Institute of Diabetes and Digestive and Kidney Diseases
301-594-4724
Email: ga96b@nih.gov

**DIABETIC WOUND HEALING AND NEUROPATHY AND TYPE 2 DIABETES**

Dr. Teresa L. Z. Jones  
National Institute of Diabetes and Digestive and Kidney Diseases  
301-435-2996  
Email: teresa.jones@nih.gov

**DIGESTIVE DISEASES AND NUTRITION**

Ms. Christine Densmore  
National Institute of Diabetes and Digestive and Kidney Diseases  
301-402-8714  
Email: densmorec@mail.nih.gov

**KIDNEY, UROLOGIC AND HEMATOLOGIC DISEASES**

Dr. Daniel R. Gossett  
National Institute of Diabetes and Digestive and Kidney Diseases  
301-594-7723  
Email: daniel.gossett@nih.gov

For administrative and business management questions, contact:

Ms. Pamela Love  
National Institute of Diabetes and Digestive and Kidney Diseases  
301-435-6198  
Email: pl48m@nih.gov
NATIONAL INSTITUTE OF ENVIRONMENTAL HEALTH SCIENCES (NIEHS)

The mission of the National Institute of Environmental Health Sciences [www.niehs.nih.gov](http://www.niehs.nih.gov) is to discover how the environment affects people in order to promote healthier lives, with a vision of providing global leadership for innovative research that improves public health by preventing disease and disability. NIEHS achieves its mission and vision through a multidisciplinary biomedical research program, prevention and intervention efforts, and a communication strategy that encompasses training, education, technology transfer and community outreach.

For additional information about NIEHS's Small Business Programs, please visit [https://www.niehs.nih.gov/funding/grants/mechanisms/sbir/](https://www.niehs.nih.gov/funding/grants/mechanisms/sbir/). In addition to this omnibus program announcement, the NIEHS releases targeted SBIR/STTR Funding Opportunity Announcements (FOAs); signup for the listserv ([https://list.nih.gov/cgi-bin/wa.exe?SUBED1=sbir-niehs&A=1](https://list.nih.gov/cgi-bin/wa.exe?SUBED1=sbir-niehs&A=1)) to be notified of FOAs.

Limited Amount of Award

For all NIEHS research interest topic areas other than Hazardous Substances Remediation and Site Characterization SBIR Program included in this PHS 2020 Omnibus SBIR/STTR Solicitation, NIEHS will accept SBIR/STTR applications up to $256,580 total costs for Phase I and $1,710,531 for Phase II. In all cases, applicants should propose a budget that is reasonable and appropriate for completion of the research project and the budget request must be well justified. The Hazardous Substances Remediation and Site Characterization SBIR Program has different limits on budget requests for both Phase I and Phase II; check under that topic below for the details. NIEHS will generally not fund applications at budget levels exceeding these budget guidelines. For budgetary, administrative, or programmatic reasons, NIEHS may decide not to fund an application or may decrease the length of an award and/or the budget.

Final Progress Reports

As detailed in NOT-OD-17-085, the NIH has implemented the Final Research Performance Progress Reports (Final RPPR) for SBIR/STTR Final Progress Reports.

The NIEHS is interested in tracking the progress of the small business concerns it funds and the products they develop. It is expected that small businesses who have received previous SBIR/STTR grants have had success in commercializing their previously supported technologies. Small businesses that are primarily interested in research and development (and not commercialization) should consider other grant mechanisms at NIH, rather than the SBIR/STTR program. Funding priority will be given to those small business concerns that demonstrate their ability to develop and commercialize products.

Phase IIB Competing Renewal Awards

NIEHS will accept Phase IIB SBIR Competing Renewal grant applications only in response to specific RFAs focused on the validation of environmental exposure assessment sensor technologies that were previously developed with Phase II SBIR or STTR funding from NIEHS or federal agencies. The proposed work must align with the NIEHS mission.

Research Topics of Interest to NIEHS

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.
<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
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<th>Yes**</th>
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<tr>
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</table>

*If No,

- This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, **UNLESS** the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,**

- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

<table>
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<tr>
<th>Does IC/Office/Agency accept clinical trials applications under this mechanism?</th>
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<th>No</th>
<th>Other Information: If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information.</th>
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<tr>
<td>Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply</td>
<td>X</td>
<td>ES 20-003 Environmental Health Sciences Core Centers (EHSCC) (P30 Clinical Trial Optional) ES 18-007 Virtual Consortium for Translational/Transdisciplinary Environmental Research (VICTER) (R01 Clinical Trial Optional)</td>
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**NIEHS Non-Clinical Trials Topics:**

**Exposure Assessment Tools**

The NIEHS Exposure Biology and the Exposome Program encompasses the totality of the exposures that a person experiences from conception to death along with the associated biological responses to those exposures. Validated tools are needed to measure, analyze, and predict a wide range of internal and external exposures and health outcomes across diverse geographic populations. These tools should be designed fit-for-purpose in collaboration with the stakeholders (e.g., community outreach programs,
citizen scientists, disaster response personnel, epidemiologists, or clinical researchers). Examples include:

**Sensors and other exposure assessment tools**

- Technologies to assess personal exposure in population studies, including networks of stationary and wearable monitors
- Devices for collecting exposure measurements across multiple stressors and scales (*i.e.*, recording time and location of exposures), with an emphasis on high sensitivity and specificity and low-cost devices, when feasible. High-priority analytes include emerging contaminants (*e.g.*, perfluorinated compounds and herbicides) as well as ultrafine particulates, PAHs, and pesticide exposures
- Technologies to collect environmental samples for subsequent targeted and untargeted laboratory analysis
- Sensor technologies that can be integrated into existing smart devices for sensing personal environment
- Personal sensors that are easily worn and durable that can be rapidly deployed after a disaster by researchers to emergency response workers and individuals in the community to help understand dermal and/or airborne exposure levels, locations, and times.

**Computational and Informatics-based Tools and Methods**

- Informatics tools and platforms to organize, store, retrieve, extract, and integrate information on exposures and health effects data
- Application of machine learning methods and natural language processing for extracting and integrating diverse data types and for generating causal networks from experimental data and public knowledgebases.
- Computational and statistical approaches to integrate exposure data from different sources, including publicly available databases and information from monitoring approaches (*e.g.*, sensors, remote sensing, and biomonitoring), to provide quantitative exposure estimates
- Adapting or developing new methods and tools for automating literature and systematic reviews, including article selection and prioritization, data extraction, study quality evaluation, and summarization of articles. This may include new methods for scanning and compiling information from gray literature and web content for environmental health impacts
- Informatic tools that can be used by the research community to rapidly build environmental health disaster research protocols similar to the NIEHS RAPIDD Protocol [https://dr2.nlm.nih.gov/](https://dr2.nlm.nih.gov/) from existing information, tools, and platforms (*e.g.*, PhenX, PROMIS, and Disaster Research Response DR2 Repository) to support rapid research response efforts in the U.S. and globally.
- Informatic and data management tools for disaster response that enable rapid collation and integration of data from stationary sources and personal exposure monitors and survey information collected from individuals using mobile platforms.
- Mobile devices and Apps for collecting information on environmental exposures from study participants involved in disaster research responses.

Information on the NIEHS Exposure Biology and the Exposome Program can be found at [http://www.niehs.nih.gov/research/supported/exposure/bio/](http://www.niehs.nih.gov/research/supported/exposure/bio/)

**Nano Environmental Health and Safety**

The NIEHS Nano Environmental Health and Safety (Nano EHS) program is interested in the detection of engineered nanomaterials (ENMs) in the environment, in consumer products, and in biological samples, and is interested in technologies or methods that can predict toxicity potential of ENMs.
High priority engineered nanomaterials of interest are those with a potential for human exposure.

Examples include:

- Sensors that can detect engineered nanomaterials in air, water, and consumer products, and provide a contextual assessment of the toxicological potential
- Mid- to high-throughput and high-content assays using in vitro or tissue chip technologies to screen and rank toxicity of emerging engineered nanomaterials for cytotoxicity, genotoxicity, and metabolic toxicity.
- Methods and tools to assess leaching of engineered nanomaterials from nanotechnology-based water filtration systems
- Technologies to assess the life cycle of nanomaterials from nano-enabled products in the market
- Development of tools and technology platforms for the isolation, characterization, and quantitation of various forms of nanoplastics from diverse aqueous sources and food samples

Information on the Nano EHS program can be found at http://www.niehs.nih.gov/research/supported/exposure/nanohealth/index.cfm

Vaping and Electronic Nicotine Delivery Systems (ENDS)

NIEHS is interested in technologies to assess exposure to aerosols from e-cigarettes and other vaping devices, including analyses of the chemical constituents in these aerosols. In addition, approaches to test the toxicity and biological responses to ENDS aerosol constituents are of interest.

Toxicity Screening, Testing, and Modeling

NIEHS supports research to identify the hazards, as well as the mechanistic understanding, of the effects of environmental stressors on biological systems that can lead to adverse human health outcomes. To increase the ability to characterize or predict the toxicity and hazard of environmental stressors, the National Toxicology Program (NTP) http://ntp.niehs.nih.gov/ at NIEHS is interested in technologies to support the goals and initiatives of the Tox21 Program http://ntp.niehs.nih.gov/results/tox21/index.html.

Technologies that support Tox21 and other NTP goals may include the development of in vitro physiologically-relevant cell-based systems that effectively model responses in humans or animals and may be used to reduce or replace in vivo animal use. High priority areas are the development of metabolically competent in vitro screening models and assay systems for various tissue types (e.g., cardiac, neurological, liver, GI tract, kidney, mammary gland, lung, and immune function).

Examples include:

- Improved human organotypic culture models (OCM) and microphysiological systems (MPS) that more accurately predict in vivo function for characterizing toxicity and/or disease processes. Priority areas are improved capability for generating more mature cells from embryonic stem (ES) or induced pluripotent (iPS) cells for organotypic models and the ability to conduct in vitro pathology studies using OCM, MPS or 3D culture models.
- Organotypic models using isolated primary cells from rat or mouse models or other experimental animal models, which can enable comparisons between in vivo and in vitro toxicity endpoints
- Approaches to characterize and integrate key molecular and cellular changes related to effects of toxicant exposures in carcinogenicity, developmental neurotoxicity, or cardiotoxicity
- Data-rich in vitro approaches that incorporate medium-throughput ‘omics and/or high-content imaging for toxicity screening
• *In vitro* toxicology screening models to predict 'idiosyncratic' chemical-induced effects in humans (e.g., drug-induced liver injury or cytokine storm)
• *In vitro* model systems that incorporate barrier functionality and transport functions into tissue models (e.g., kidney, placenta, or blood-brain barrier)
• Enhanced lower organism models (e.g., zebrafish or *C. elegans*) for toxicity screening
• Stem cell models and assays for evaluating the effects of toxicants on cell differentiation with multiple functional endpoints
• Screening systems that incorporate genetic diversity into toxicology testing (e.g., panels of human IPS cells or rodent stem cells)
• *In vitro* assays to model chronic inflammatory responses to xenobiotics
• Short-term tests, assays, or systems designed specifically to reduce or replace existing regulatory animal studies for acute toxicity (oral or inhalation), reproductive or developmental toxicity, carcinogenicity, or ocular toxicity

**Computational approaches for predictive toxicology**

• New computational systems and tools for integrating toxicity data, including *in vivo* and *in vitro* data, to analyze and visualize data across different screening systems
• Computational tools to integrate and visualize transcriptomic and metabolomic data into affected signaling and biochemical pathways.
• Improved experimental and computational tools for *in vitro* to *in vivo* extrapolation of xenobiotic exposures across a range of assay types
• Computational tools for quantitatively modeling metabolic transformation of xenobiotics
• Computational tools or systems for rapidly assessing results of literature searches, short-term tests, assays, or other relevant testing to help provide screening level risk characterization of complex mixed chemical exposures in response to disasters

**Other technologies for enhanced toxicology testing**

• Alternative or improved methods for fixing and preserving tissues that maintain cellular structure for histopathology while minimizing degradation of nucleic acids (RNA, miRNA, DNA, methylated DNA), proteins or metabolites, so that archival tissue blocks can be better used for molecular analysis
• Liquid biopsy methods for isolation and novel assays of circulating nucleic acids that reflect environmental chemical exposures or toxicity. These could include exosome-packaged or cell-free nucleic acids.
• Approaches to improve efficiency of laboratory-based toxicology studies, including more rapid, cost-effective methods for RNA isolation from tissue lysate, and novel methods for removing necrotic cells in complex cell culture models
• Alternative or improved methods for extracting high quality RNA, miRNA, DNA, methylated DNA, proteins, or metabolites from existing archived tissues

**Biomarkers of Exposure and Response**

To better understand the risks to human health from environmental agents, NIEHS supports the development and validation of biomarkers of exposure, including improved measures of internal dose, DNA adduct identification, and untargeted analysis for metabolite identification, and biomarkers of response, including assays that can distinguish reversible from irreversible changes in target organs or surrogate tissues. Examples include:

**Biomonitoring technology**

• Personal or point-of-care monitoring technologies for rapid detection of multiple exposures in biospecimens using non- or minimally-invasive approaches
• Improved methods to detect DNA or protein adducts resulting from exogenous exposures

**Biological response markers**

• Markers of oxidative stress, inflammation, DNA damage response, immune function, mitochondrial dysfunction, or altered epigenetic regulation

• High priority human biomarkers include, but are not limited to: inflammation biomarkers, plasma- or serum-based markers that reflect altered RNA, protein expression, or metabolite profiles, markers developed in exhaled breath, buccal cells, or other easily accessible, non-invasive biological samples, miRNA or other exosome biomarkers, and epigenetic markers in surrogate tissue reflecting modifications in target tissues

**Intervention Technologies**

NIEHS supports efforts to prevent or reduce exposures to environmental stressors that affect human health. Technologies to reduce exposure may include:

• Technologies for removing contaminants from drinking water for home use

• Approaches for use in the home, workplace, and school settings for reducing volatile compounds and other inhaled toxicants. Examples may include improved air filtration systems as well as technologies to monitor the efficacy of filtration systems

• Technologies and applications that can provide real-time alert about relevant environmental exposures in sensitive populations (such as asthmatic populations)

**Education/Outreach**

As part of its Partnerships for Environmental Public Health (PEPH) Program, NIEHS is interested in developing tools that build capacity, improve environmental health literacy, and support citizen science endeavors. These approaches or resources should be fit for purpose to meet the needs of the following audiences: community members, health care and public health professionals, educators, and students of all ages. Approaches may include:

• Mobile applications that provide environmental health information about exposures of concern in food, air, water, or consumer products. These may include
  o Apps that provide the context for the exposures such as single or multiple, interacting exposures, level of exposure, frequency and proximity to source
  o Apps that can be adapted for various age groups (e.g., children or the elderly), races, ethnicities and/or languages
  o Apps that visualize exposure risks with respect to levels of exposure, sources and health risks

• Devices for collecting and reporting information on exposures in environmental samples for educational purposes in schools or communities

• Systems that can utilize public and voluntary population data from sensors, activity trackers, GIS enabled devices, social communications, and surveillance cameras; for example, to assist disaster response and communication

• Educational resources related to environmental health in school settings or community education programs (e.g., Photovoice projects or GIS mapping)

• Training materials for wider dissemination of risk information (e.g., resources for high school students to train younger students; or community leaders to build capacity of other community residents)

• Continuing medical education classes, on-line courses, or on-line tools to build the environmental health literacy of health care professionals
• Tools to disseminate current information on exposure risks or hazards. This may include signage and other materials to communicate hazards, including risks from fish consumption, marine toxins, or air quality alerts in multiple languages for affected communities

Information on the PEPH program can be found at https://www.niehs.nih.gov/research/supported/translational/peph/index.cfm

Hazardous Substances Remediation and Site Characterization SBIR Program

The NIEHS Superfund Research Program (SRP) “Hazardous Substances Detection and Remediation Program” supports Small Business Innovation Research Grants (SBIR R43, R44) to foster the commercialization of novel technologies, products, and devices for remediation and detection of hazardous substances in the environment. The SRP is specifically interested in proposals applying new engineering, materials science, and biotechnology approaches to develop sustainable strategies to characterize and remediate hazardous substances at contaminated sites.

Topics of interest include, but are not limited to:

**Remediation**

• Novel technologies for *in situ* remediation of contaminated sediments, soils, and groundwater
• Innovative bioremediation and phytoremediation technologies including development and culturing/propagation of plants, bacterial strains, or fungal species for implementing bioremediation
• Remediation strategies designed to emulate/enhance biological processes or functions (e.g. biomimicry) by applying novel replacement materials/technologies to source areas or downgradient plumes
• Technologies to remediate chemical mixtures in environmental media
• New strategies for delivery of reagents/amendments for groundwater remediation and/or recovery/extraction of contaminants in groundwater
• New amendments to stabilize contaminants and/or to stabilize caps for soil and sediment remediation
• Technologies and strategies to cleanup large complex sites with multiple sources

**Site Characterization**

• Computational, geographical information system-based, or modeling products for predicting fate and transport of contaminants, rates of remediation, bioavailability, or for identifying contamination sources
• Real-time, field deployable, on-site analysis: soil, surface water, groundwater, subsurface, sediments, air (such as volatile releases from sites), etc.
  • monitoring and screening of contaminants
  • bioavailability (i.e. biologically-accessible portion of contamination)
  • miniaturized toxicity-screening kits
• Products that allow for rapid sample clean-up/preparation for analysis of environmental samples
• Non-targeted or multi-analyte field sampling devices or kits, including sample collection products that can sequester a suite of analytes for later analysis
• Novel techniques, sensors, and field analytical methods and real-time mapping/data visualization for development of subsurface conceptual site models

**Examples of remediation and site characterization needs:**
- Devices to detect and measure vapor intrusion and solutions for mitigation, including tools to determine when vapor mitigation is complete
- Devices to detect and measure non-aqueous phase liquids (NAPLs) and dense non-aqueous phase liquids (DNAPLs) in the subsurface
- Site characterization techniques and strategies for complex geology (fractured, karst and heterogeneous layered deposits) including understanding the fate of contaminants within rock matrices and properties that affect back diffusion
- Technologies for rapid extraction or processing of soil for incremental sampling methodologies (ISM)
- Technologies for rapid analysis of total and/or biologically-accessible (e.g. bioavailable/bioaccumulating) metal fractions in soils and sediments (including easy-to-use kits for soil/sediment screening)
- In-well real-time and/or continuous monitoring tools to assess the efficacy of remediation; presence/absence of key factors required for remediation (e.g. biological, geological, chemical); and/or to identify rebound events
- Technologies for automated elongated mineral fiber counting (e.g. for asbestos samples)
- Active or passive remediation technologies for mining influenced water; technologies to mitigate effects from acidic drainage; portable neutralization treatment systems; and strategies to target remediation of sources such as mining waste piles
- Soil, sediment, and groundwater remediation technologies for mixtures and degradation byproducts of poly- and perfluorinated alkyl substances (PFAS), including sustainable solutions with low energy input and/or minimal secondary waste generation
- Performance enhancements to existing remedial technologies (permeable reactive barriers, funnel and gate systems, capping to limit infiltration, etc.)
- Novel detection technologies and remediation approaches that: improve energy efficiency and reduce waste generation; and/or are capable of improving resilience such as: withstanding fire, flooding, land use changes, and other catastrophic events
- Technologies for measuring/treating environmental contamination as part of a disaster response effort

Applicants must demonstrate that the proposed technologies are relevant to Superfund. Per program mandates described in the Superfund Amendment Reauthorization Act (SARA), SRP does not accept applications targeting oil or gas site characterization/remediation. Applicants are strongly encouraged to stay within the statutory budget guidelines whereby total funding support (direct costs, indirect costs, fees) does not exceed $168,087 for Phase I awards and $1,120,586 for Phase II awards. Applicants are encouraged to contact NIH program officials prior to submitting any award budget for the “Hazardous Substances Remediation and Site Characterization Small Business Innovation Research Program” in excess of these amounts. Please note: the NIEHS Superfund Research Program (SRP) "Hazardous Substances Remediation and Site Characterization Small Business Innovation Research Program" no longer accepts Small Business Technology Transfer Grant (STTR: R41, R42) applications.

https://www.niehs.nih.gov/research/supported/centers/srp/funding/hwaerp/index.cfm

Worker Training Program

The NIEHS Worker Training Program (WTP) is interested in the development of e-Learning health and safety Advanced Technology Training (ATT) products from a variety of delivery methods to assist both students and instructors in the training and education process. These ATT products are for the health and safety training of hazardous materials (HAZMAT) workers; waste treatment personnel; skilled support personnel associated with an emergency/disaster; emergency responders in biosafety response, infectious disease training and cleanup; emergency responders in disasters; and resiliency training. ATT
as defined by the Worker Training Program (WTP) includes, but is not limited to, online training, virtual
reality, and serious gaming, which complement all aspects of training from development to evaluation
including advance technologies that enhance, supplement, improve, and provide health and safety
training for hazardous materials workers. WTP accepts solicitations via requests for applications
(RFA). Please contact Kathy Ahlmark ahlmark@niehs.nih.gov for information on the next solicitation
date, which differs from the standard receipt dates of this NIH omnibus.

Information on the WTP program can be found at https://www.niehs.nih.gov/careers/hazmat/about_wetp/

NIEHS DOES NOT Fund

• Technologies for the detection and remediation of pathogens in the environment.

NIEHS Clinical Trials Topics:

NIEHS will accept SBIR/STTR applications that propose clinical trials related to:

• Development and testing of sensor technology, biomarkers, or biomanitoring technologies,
including field testing of new technologies for exposure assessment and biological responses
to environmental exposures

• Evaluation of tools or approaches for education and dissemination of information on
environmental hazards, including evaluation of changes in behavior

For additional information on research topics, contact:

Dr. Daniel Shaughnessy
National Institute of Environmental Health Sciences
Division of Extramural Research and Training
POB 12233 (K3-12)
Research Triangle Park, NC 27709
(984) 287-3321
Email: shaughn1@niehs.nih.gov

Dr. Lingamanaidu Ravichandran
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Division of Extramural Research and Training
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(984)-287-3309
Email: lingamanaidu.ravichandran@nih.gov

For information on the NIEHS Superfund Research Program - Hazardous Substances Remediation and
Site Characterization SBIR Program, contact:

Dr. Heather Henry
National Institute of Environmental Health Sciences
Division of Extramural Research and Training
POB 12233 (K3-12)
Research Triangle Park, NC 27709
(984) 287-3268
Email: henryh@niehs.nih.gov
For administrative and business management questions contact:

Mr. Aaron Nicholas
National Institute of Environmental Health Sciences
Division of Extramural Research and Training
Grants Management Branch
(984) 287-3297
Email: nicholaa@nih.gov
NATIONAL EYE INSTITUTE (NEI)

The mission of the NEI is to conduct and support research, training, health information dissemination, and other programs with respect to blinding eye diseases, visual disorders, mechanisms of visual function, preservation of sight, and the special health problems and requirements of the blind.

Limited Amount of Award

For budgetary, administrative, or programmatic reasons, NEI may decrease the length of an award and/or the budget recommended by a review committee, or not fund an application. For topics listed in APPENDIX A: National Institutes of Health SBA-Approved SBIR/STTR Topics for Awards over Statutory Budget Limitations, NEI does not generally fund Phase I applications greater than $300,000 total costs or project periods greater than 2 years; or Phase II applications greater than $1,800,000 total costs or project periods greater than 3 years. Applicants are strongly encouraged to contact program officials prior to submitting any application in excess of the hard caps listed above and early in the application planning process.

Phase IIB Competing Renewal Awards

The NEI will only accept SBIR Phase IIB Competing Renewal grant applications from Phase II SBIR awardees to continue the process of developing technologies that ultimately require federal regulatory approval or require extraordinary time and effort in the Research and Development phase. Such technologies include, but are not limited to, pharmacologic agents, biological products, and devices. These technologies should be clearly related to the mission of the NEI. This renewal grant should allow small businesses to reach a stage in the project where interest and investment by third parties is more likely. The NEI expects that the Phase IIB grant will accelerate the transition of SBIR Phase II projects to the commercialization stage. The NEI encourages applicants to establish business relationships with third-party investors and/or strategic partners who can provide substantial financing to help accelerate the commercialization of promising new products and technologies that were initiated with SBIR funding. The Competing Renewal application must be a logical extension of a previously completed Phase II (R44) SBIR grant. NEI grantees seeking SBIR Phase IIB Competing Renewal funding must submit an application within a period no later than the first six receipt dates following expiration of the previous Phase II budget period. Cumulative budgets should not to exceed $1,8000,000 total costs, or time periods beyond three (3) years.

Although matching funds are not required, the NEI strongly encourages that applicants obtain significant private investment. Competitive preference and funding priority will be given to applicants that demonstrate the ability to secure substantial independent third-party investor funds.

Applicants are strongly encouraged to contact the NEI Program Officer, Dr. Paekgyu Lee (contact information provided below) prior to submitting any application in excess of the hard caps listed above and early in the application planning process.

Research Topics of Interest to NEI

The following topics are meant for illustrative purposes only and are not exclusive of other appropriate activities.

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.
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<td>Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply</td>
<td>X</td>
<td>R01 PA-18-351 R24 PAR-18-707 UG1 PAR-18-521, PAR-18-522, PAR-18-523</td>
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**NEI Non-Clinical Trials Topics:**

A. **General Research and Development Topics:** NEI is interested in providing support for the development of new technologies, strategies, research tools, reagents and methods that can be applied to basic and translational research which will benefit vision health. This encompasses research and development of innovative enabling technologies in areas of genomics, proteomics and nanotechnology. More specific topics include drug and high throughput assays; drug delivery systems; gene therapy, cell-based therapy and regenerative medicine; development of in vitro and in vivo disease models; surgical devices and materials; telemedicine, mobile health, and
health education; and design/fabrication of new or improved ophthalmic instruments for diagnosis
and treatment of eye disorders.

B. **Retinal Diseases:** New therapeutic approaches for inflammatory and degenerative diseases and
for inhibition of abnormal angiogenesis in the retina and choroid; Better methods of diagnosing
and treating diabetic retinopathy and other vascular diseases; Non-invasive techniques for early
diagnosis of macular degeneration and other retinal degenerative diseases; Instruments and
procedures for improved surgical management of retinal detachments; Retinal prostheses to help
restore visual function; Gene therapy/optogenetic methods for light sensitivity restoration in the
retina; Better methods for cell or tissue transplantation; New animal models/systems that better
mimic human retinal disease.

C. **Corneal Diseases:** New diagnostic tools, therapeutic agents and drug delivery methods for the
treatment of corneal injury, infection, dry eye, ocular pain, and other ocular surface disorders;
New biomaterials for corneal prostheses and corneal transplants; Instruments and procedures for
correcting the refractive power of the cornea and/or measuring the cornea's optical properties or
other physiological properties.

D. **Lens and Cataract:** New approaches in the post-operative management of cataract surgery;
New surgical instruments for cataract extraction and new biomaterials for replacement of the
natural lens; Design/fabrication of aspheric, toric, multifocal and accommodating intraocular
lenses.

E. **Glaucoma and Optic Neuropathies:** New therapeutic agents, instruments, and procedures for
the diagnosis and treatment of glaucoma; Non-invasive methods to measure changes in the optic
nerve head and retinal fiber layer.

F. **Strabismus, Amblyopia, and Refractive Error:** New approaches to detect and treat strabismus,
amblyopia, and myopia; New tools and techniques for vision screening; New or improved
methods and materials for correcting the refractive power of the eye and/or measuring the eye’s
optical properties or other physiological properties; New materials and manufacturing processes
for eyeglasses and contact lenses; prosthetic devices (both cortical and subcortical) for vision
restoration.

G. **Visual Impairment and Blindness:** Instruments and methods to better specify, measure, and
categorize residual visual function; New or improved devices, systems, or programs that meet the
rehabilitative, adaptive, and everyday living needs of visually-impaired/blind people.

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**Additional Information**

The NEI's programs are described in more extensive detail in documents which are available from the Institute. For additional information about the research programs of the NEI, please visit our home page at [http://www.nei.nih.gov](http://www.nei.nih.gov).

For more information on research topics, contact:

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Email: kyr@nei.nih.gov
NATIONAL INSTITUTE OF GENERAL MEDICAL SCIENCES (NIGMS)

The NIGMS supports research and research training in the basic biomedical sciences and in specific clinical areas (i.e., clinical pharmacology, trauma and burn injury, sepsis, wound healing, and anesthesiology). NIGMS supports research of potential interest to small businesses and their collaborators through the:

- Division of Biophysics, Biomedical Technology, and Computational Biosciences
- Division of Pharmacology, Physiology, and Biological Chemistry
- Division of Training, Workforce Development, and Diversity
- Division for Research Capacity Building

For additional information about areas of interest to the NIGMS, please visit our home page at http://www.nigms.nih.gov. This site includes staff contact information by program area (http://www.nigms.nih.gov/about/pages/contactbyarea.aspx).

Limited Amount of Award

According to statutory guidelines, total funding support (direct costs, indirect costs, fee) normally may not exceed a hard cap ($256,580 for Phase I and $1,710,531 for Phase II and Phase IIB). NIGMS will not accept applications with budget requests exceeding this hard cap with the exception of projects fitting within the list of SBA-approved topics for awards over the statutory budget limitations; the entire list for NIH (including NIGMS) may be found in Appendix A of this document.

If considering a project with a budget exceeding the hard cap, applicants are strongly encouraged to contact NIGMS program officials prior to submission, and preferably earlier during application preparation. In all cases, applicants should propose a budget that is reasonable and appropriate for completion of the research project.

Phase IIB Competing Renewal Awards

NIGMS will accept Phase IIB SBIR-only Competing Renewal grant applications to continue the process of developing products that ultimately require 1) clinical evaluation, 2) approval by a Federal regulatory agency, or 3) continuing refinements that include but are not limited to cost reduction, testing for performance, safety, reliability and/or durability, and meeting or establishing standards, particularly for basic or clinical research instrumentation or durable medical equipment (DME) designs. This renewal grant should enhance the likelihood that small business will attract interest and investment by third parties. Such products include, but are not limited to research equipment, biological products, devices, drugs, medical implants, etc. within the mission of the NIGMS. Budgets for this Phase IIB Competing Renewal opportunity must follow the guidelines for Phase II applications (described above). For awards that are intended to support completion of research needed to obtain an Investigational New Drug application (IND) or Investigational Device Exemption (IDE), applicants must provide evidence that they have consulted formally with the FDA concerning the research needed for the development of a drug, biologic or medical device and that the proposed research will address these regulatory requirements. Such evidence should include FDA correspondence from a pre-IND meeting for an IND application or a pre-IDE meeting for an IDE application, and the status of the project in a timeline related to Federal regulatory approval processes.

Prospective applicants considering a Phase IIB Competing Renewal application are strongly encouraged to contact either the Program person of record for the Phase II award or NIH staff listed at the end of this NIGMS topics announcement.
To assist NIGMS in planning for Phase IIB applications, it is helpful for prospective applicants to submit to the NIGMS SBIR/STTR Coordinator (listed below) a letter of intent that includes the following information:

- Descriptive title of the proposed research
- Phase II grant number
- Name, address, and telephone number of the Principal Investigator
- Names of other key personnel
- Participating institutions
- Funding Opportunity Announcement Number

The letter is non-binding and does not enter into the review process. It is anticipated that only a small number of NIGMS SBIR Phase II awards would be eligible for a Phase IIB Competing Renewal award.

**Research Topics of Interest to NIGMS**

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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https://www.nigms.nih.gov/Research/mechanisms/Pages/SBIR.aspx

https://www.nigms.nih.gov/Research/mechanisms/Pages/STTR.aspx

**NIGMS Non-Clinical Trials Topics:**

**Division of Biophysics, Biomedical Technology, and Computational Biosciences**

This Division facilitates advances in basic biomedical research by supporting the development of biophysical and computational methods and tools for understanding basic biological questions; physical and theoretical methodologies, bioinformatics tools, and sophisticated quantitative approaches to lay a foundation for advances in disease diagnosis, treatment, and prevention in health and disease; and the creation of innovative tools and new technologies for the study of macromolecular, cellular, and organelle processes and function. Research on membrane synthesis, structure, and function; membrane models; membrane transport; cell division; cell organization; cell motility; and biophysics of proteins, nucleic acids, and biological assemblies, including viral entry, packaging, maturation, and release, as well as the development of instrumentation, components, and methods for the analysis of cellular components and macromolecules. Research on developing a better understanding of fundamental processes and mechanisms of development and inheritance in health and disease, and population genetics.

Example areas that may be of interest to small businesses include, but are not limited to:

1. Development of instrumentation, devices, and methods for detecting, analyzing, and separating biologically important compounds, macromolecules, and their interactions.
2. Development of new methods and materials directed toward the solution of biological macromolecule structures, assemblies and complexes by, but not limited to, x-ray diffraction, electron diffraction, NMR, cryo-EM and mass spectroscopy.
3. Development of new or improved instruments, devices, and related methodologies to facilitate biomedical research. Instrumentation includes, but is not limited to mass spectrometry, nuclear magnetic resonance, optical or laser spectroscopies, X-ray absorption/diffraction/scattering,
detectors, electron or confocal microscopies, electrophoresis and other separation techniques, bioreactors, centrifugation, and flow cytometry.
4. Technology for microscopy and imaging: development of new or improved microscopic techniques, instruments, reagents, and supporting software that measures the location and dynamics or molecules in situ, organelles, cells, or tissues on the nano- and micro-scale.
5. Imaging probes and sensors, other reagents and methods, instrumentation, software for microscopy, spectroscopy, and single molecule analysis of molecules, cells, tissues, embryos and small model organisms. Technologies for applications of microscopy, spectroscopy and single molecule analysis in basic biomedical research, including but not limited to light, electron, X-ray and scanning probe microscopy and fluorescence, magnetic and electron paramagnetic spectroscopy. NOT included are small animal and preclinical imaging.
6. Development of tools including but not limited to detectors, cameras, light sources, optics, and automated data collection and analysis systems, for studying the structures of biomolecules and biospecimens in the size range of peptides to cells, using diffraction, microscopy and/or spectroscopy techniques.
7. Development of technologies for investigating and manipulating cells: development of tools and methods that manipulate or investigate the properties of cells and their environment. Development of tools for cell engineering, molecular transport and partitioning, and assays for cellular phenotype.
8. Development of novel technologies for proteomics, glycomics, metabolomics, and other aspects of systems biology for discovery and clinical applications, (e.g., sample handling, separations, mass spectrometry, and computational tools for protein identification, data curation and mining, and for integrating genome variation, pathways and networks with biological function).
9. Development and improvement of methods for the expression, solubilization, and purification of regulatory, cellular, and membrane associated proteins, as well as for the preparation of specifically labeled macromolecules and the recovery of proteins from inclusion bodies.
10. Development of novel ligands, inhibitors, and other probes for spectroscopic and microscopic analysis of cellular assemblies and viral structures, macromolecules and components, their localization and function in vivo and at a single molecule level.
11. Improvement of methodology (technology) for genetic analysis (e.g., gene expression, probes), including procedures for the separation and analysis of nucleic acids and proteins as these relate to genetic problems.
12. Improvement in procedures (statistical, computational, laboratory) for the analysis of gene flow and gene dynamics in human populations.
15. Development of tools and technologies to detect and monitor complex human phenotypes or traits.
16. Development of technology to derive and expand pluripotent cell populations from non-embryonic sources, for example, induced pluripotent stem cells (iPS), to scale up the growth of induced pluripotent stem cells in culture and to regulate their differentiation state.
17. Development of markers, reagents and tools to characterize the unique properties of iPS cell lines and to distinguish them from adult stem cells and more differentiated cells.
18. Development of existing human embryonic stem cell lines and new or existing iPS cells as a model system for drug discovery.
19. Development or improvement of methodology for generation of antibodies or other affinity reagents for proteins and other small molecules in non-mammalian genetic model systems.
20. Improvement in procedures (statistical, computational, laboratory) for the high- and medium-throughput analysis of gene expression patterns and regulatory networks.
21. Development or improvement of methods for high throughput detection of epigenomic changes.
22. Development or improvement of methods for characterizing the metabolic interactions of complex communities of microorganisms particularly those involved in host-microbe interactions.
23. Development of improved or novel methodology for structure/function analysis of very large macromolecular complexes involved in transmission or expression of genetic material. Development of new or innovative tools and methods in bioinformatics and computational biology.
24. Development of information and communication technology in support of biomedical research, that apply best practices and proven methods for software design, construction and implementation to promote adoption by a broad biomedical research community.
25. Computational methods for analysis, prediction, and improving methods for determination of macromolecular structures and structure-function relationships.
26. Development of computerized tools that might be used in the presentation of the concepts of cell and structural biology to audiences at a variety of levels.
27. Development of computer software for the analysis of the primary and secondary structures of nucleic acids as these relate to genetic problems.
28. Development of tools and methods for the modeling, simulation or analysis of complex biological systems.
29. Development of collaborative environments and technologies to translate Big Data to knowledge, including but not limited to development of knowledge environments, data integration, data and metadata curation methods, and tools that address data security and privacy issues.
30. Development of tools and methods to collect, interpret, analyze and visualize scientific data through integration and interoperability of different data types.
31. Design and development of software and hardware for improving the effectiveness of computational approaches in biomedical research.
32. Development of computational biology software packages for integrative analysis of biomedical data.
33. Development and enhancement of databases and data formats for biomedical research activities.
34. Development of tools and technologies for a biomedical data science ecosystem for biomedical research. Technologies for findability, interconnectivity, and interoperability of biomedical data sets and resources, integration of existing data management tools and development of new ones, universalization of innovative algorithms and tools.

**Division of Pharmacology, Physiology, and Biological Chemistry**

The Division’s research interests include: an improved understanding of drug action and of anesthesia; mechanisms underlying responses to drugs; new methods and targets for drug discovery; advances in natural products synthesis; carbohydrate structure and glycan biological function; an enhanced understanding of biological catalysis; knowledge of metabolic regulation and fundamental physiological processes; drug metabolism and drug delivery strategies; critical illness and injury; sepsis.

Examples include, but are not limited to:

**A. Biochemistry and Bio-related Chemistry**

2. Development of synthetic methodology to improve the efficiency of discovery, development, and production of bio-medically relevant compounds.
3. Development of enzymes, catalytic antibodies, ribozymes, artificial enzymes, and host molecules as drugs, diagnostic reagents or synthetic tools.

5. Development and application of methods and materials for the elucidation of membrane protein structures and multimeric complexes at or near atomic resolution.

6. Development of high-throughput methods for sequencing and re-sequencing of mitochondrial genes and relevant nuclear genes and for proteomic and/or functional profiling of mitochondria heteroplasmy.

7. Development of new metal ion chelators and other tools to probe and/or alter the localization and concentration of metal ions in cells and in whole organisms. Research to exploit metal metabolism and metal-regulated cellular control and cell-cell signaling processes to probe and/or alter cell function. Research to develop investigational and therapeutic applications of metal-complexes and to understand the factors governing their pharmacology and toxicology.

8. Development of high-throughput methods and strategies to characterize the function of G protein-coupled receptors and other membrane proteins, ion channels, transporters, and/or enzymes. Research to define the functional interrelationships between proteins and enzymes.

9. Development of tools to characterize oxidative stress and oxidative stress related molecules (e.g., NO, peroxynitrite, hydrogen peroxide, lipoxidation products, modified proteins, DNA modifications, etc.) including the extent and/or localization (by organ/tissue/cell/organelle) of oxidative stress.

10. Development of technologies and methodologies to measure enzymatic activities in native environments such as cells and organelles and to measure metabolic flux of transient multienzyme complexes.

B. Pharmacological and Physiological Sciences

11. Isolation, characterization, and development of factors and strategies, methods, or treatments involved in critical illness and injury including tissue repair, wound healing, sepsis and associated pain management. Research may involve emergency, peri-operative, and/or critical care conditions.

12. Improved systems for collection, processing, and analysis of real time physiological data from injured or critically ill patients.

13. Development of tools, software, algorithms, etc. needed to combine different types of data (such as clinical, demographic, physiologic, genomic, proteomic) obtained from injured or critically ill patients particularly in the context of sepsis.

14. Assays and tools to enable molecular based (-omic) analyses of critically ill patients.

15. Development of strategies, methods, or new technologies to improve the delivery, monitoring, safety and efficacy of anesthesia.

16. Research to improve drug design and delivery.

17. Research to improve drug bioavailability by improved understanding of factors that influence absorption, metabolism, transport, or clearance of therapeutics and underlying mechanisms. Application of pharmacokinetic and pharmaceutical principles to the study of large biomolecules, such as proteins, polypeptides, and oligonucleotides.
18. Development of novel in vivo and in vitro methods to predict the safety and toxicities of pharmacologic agents.

19. Development of bioinformatic, mathematical, and/or computational approaches/resources and/or pharmacokinetic modeling programs which utilize ADME parameters of drugs and information from individual patients or patient populations, to reduce adverse drug reactions.

20. Alternative and replacement models for animal testing of sepsis therapeutics.


22. Biomarker panels to enable rapid diagnosis and/or optimize treatment of critically ill patients.

23. Predictive clinical algorithms, electronic health record tools, and point of care diagnostics particularly those that will enable bedside use of molecular/omic information in critically ill patients.

24. Clinical decision support technologies that address early recognition of sepsis, patient trajectories, and resolution of sepsis.

25. Microfluidic technologies for use in sepsis research, diagnosis, and/or treatment.

26. Diagnostic tests for early detection of sepsis.

**Division of Training, Workforce Development, and Diversity**

This division supports the development technologies and tools to enhance the research skills of post-high school individuals in the biomedical research workforce pathway, or to increase the efficiencies of NIGMS research training programs. The technologies may be new products or adaptation of existing products designed to be more efficient, cost-effective, culturally appropriate, and/or user-friendly in promoting the development of the biomedical research workforce. Examples for skills development projects include but are not limited to web-based resources, instructional software, interactive media, research-focused curriculum materials, and active learning toolkits. Projects aimed at enhancing NIGMS training programs include but are not limited to technologies to track career outcomes of students and trainees and/or assist in the evaluation of workforce development programs (e.g., survey instruments and/or training activity tracking systems). Projects that will develop skills of individuals from underrepresented groups (see the NIH interest in Diversity) or increase the efficiencies of diversity enhancing research training programs are encouraged.

**Division for Research Capacity Building**

Research and development of products, tools, databases, or services in the mission of Division for Research Capacity Building (DRCB), which supports research, research training, faculty development and research infrastructure improvements in states that historically have not received significant levels of research funding from NIH. It also supports faculty research development at institutions that have a historical mission focused on serving students from underrepresented groups, and conducts a science education program designed to improve life science literacy. Topics of interest include, but are not limited to, research capacity building, faculty and professional development, mentoring, and development of educational software and course materials targeting undergraduate and community college students, on topics that range from basic molecular and cellular biology to human diseases, including areas of health disparities, that disproportionately affect rural, tribal and hard-to-reach populations. To meet the information needs of the target audiences, health education materials must be culturally appropriate.

DRCB is composed of four programs: Institutional Development Award (IDeA), Support of Competitive Research (SCORE) and Native American Research Centers for Health (NARCH), and Science Education Partnership Awards (SEPA).
A. **Science Education Partnership Awards:** Funding opportunities are available for the development of discovery-oriented educational software, Serious STEM Interactive Digital Media (IDM) and the application of educational technology and tools for health science topics that target pre-kindergarten to grade 12 (P-12) students, teachers and families, and the general public, particularly those from underserved communities. Development of software, IDM technology, or other educational tools may be focused on new products or adaptation of existing products designed to be more efficient, cost-effective, and user-friendly in promoting problem solving, interactive learning, dissemination, and promotion of health science. These resources are intended to create user-friendly, culturally appropriate and effective educational resources.

Examples of responsive applications may include but are not limited to:
1. Web-based, stand-alone computational tools, instructional software, or other interactive media for dissemination of science education;
2. Big Data and bioinformatics tools, software and apps for students, teachers and the business community.
3. Curriculum materials, interactive teaching aids, models for classroom instruction, and teacher education workshops;
4. Serious Science, Technology, Engineering and Mathematics (STEM) IDM resources;
5. Development of health promotion and disease prevention/intervention products that are culturally appropriate for the target populations and communities.

Projects that target the following constituencies are strongly encouraged:
1. P-12 students, teachers, and parents;
2. Students of community colleges, tribal colleges, undergraduate colleges, and minority-serving institutions;
3. Patients and families with health conditions that disproportionately affect minorities and other medically underserved populations, including members of disadvantaged urban and rural communities.

**NIGMS Clinical Trials Topics:**

NIGMS will accept clinical trials in any area listed above in the non-clinical trials topics.

For scientific questions about NIGMS-funded SBIR/STTR research, contact:

**DIVISION OF PHARMACOLOGY, PHYSIOLOGY, AND BIOLOGICAL CHEMISTRY**

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**DIVISION OF BIOPHYSICS, BIOMEDICAL TECHNOLOGY, AND COMPUTATIONAL BIOSCIENCES**

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Research and Development in Science Education
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**ADMINISTRATIVE AND BUSINESS MANAGEMENT:**

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Email: ilene.Glassman@nih.gov

Ms. Julie Chang
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Email: julie.chang@nih.gov

For additional information on NIGMS research topics and the SBIR/STTR application process, contact:

**NIGMS SBIR/STTR COORDINATOR**

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301-435-0752, Fax: 301-480-0884
Email: dmitriy.krepkiy@nih.gov
NATIONAL HEART, LUNG, AND BLOOD INSTITUTE (NHLBI)

For the most up-to-date information, please visit the NHLBI SBIR/STTR website (https://sbir.nih.gov/nhlbi) and subscribe to our listserv (http://bit.ly/NHLBI-SBIR-Updates). You can also follow us on Twitter @NHLBI_SBIR. NHLBI encourages potential applicants to contact us at http://bit.ly/ContactNHLBIsbir.

NHLBI plans, conducts, and supports research, clinical trials, and demonstration and education projects related to the causes, prevention, diagnosis, and treatment of heart, blood vessel, lung, and blood diseases, and sleep disorders. It also supports research on the clinical use of blood and all aspects of the management and safety of blood resources. The NHLBI SBIR/STTR programs foster basic, applied, and clinical research on all product and service development related to the mission of the NHLBI. The NHLBI has four extramural program divisions, described below.

Phase II Applications

The NHLBI strongly encourages applicants to include a robust regulatory strategy with corresponding milestones in Phase II applications. Applicants are also encouraged to include letters of support or other evidence documenting their regulatory strategy. The NHLBI will consider the strength of the regulatory plan when making funding decisions. For assistance regarding regulatory strategy, explore the “Small Biz Hangout” series on the NHLBI YouTube channel and submit specific regulatory questions to Chris Sasiela at http://bit.ly/ContactNHLBIsbir.

For assistance regarding the commercialization plan, watch the “Small Biz Hangout” for advice on Writing Your Phase II Commercialization Plan (http://bit.ly/Ph2CommPlanHangout) and contact Mike Pieck (nhlbi_sbir@mail.nih.gov) with specific questions.

NHLBI-Supported Funding Opportunity Announcements (FOAs)

In addition to this Omnibus program announcement, the NHLBI releases targeted Funding Opportunity Announcements (FOAs) throughout the year. Sign up for the listserv (http://bit.ly/NHLBI-SBIR-Updates) to be notified of new FOAs.

These FOAs are listed to inform potential applicants about other funding opportunities to which they can apply; applications submitted in response to this Omnibus program announcement are not limited to research and development areas described in the following targeted FOAs. The NHLBI also encourages mission-aligned applications for innovative technologies outside these targeted areas.

(Funding Opportunity Announcements can be released or expire at any time throughout the year; please refer to the NHLBI SBIR/STTR website for active announcements supported by NHLBI.)


Phase IIB Competing Renewal Awards

The NHLBI does not accept applications for Phase IIB competing renewal awards through this Omnibus solicitation; however, the NHLBI offers SBIR Phase IIB opportunities through the NHLBI Phase IIB Bridge
Awards and the NHLBI Phase IIB Small Market Awards using separate funding opportunity announcements (Bridge Award: RFA-HL-19-017; Small Market Award: RFA-HL-19-018)

The purpose of the NHLBI Phase IIB program is to accelerate the transition of SBIR/STTR Phase II projects to the commercialization stage by promoting partnerships between SBIR/STTR Phase II awardees and third-party investors and/or strategic partners. NHLBI SBIR Phase IIB program encourages business relationships between applicant small business concerns and third-party investors/strategic partners who can provide substantial financing to help accelerate the commercialization of promising new products and technologies that were initiated with SBIR/STTR funding. In particular, applicants are expected to leverage their previous SBIR/STTR support, as well as the opportunity to compete for additional funding through the NHLBI Phase IIB program, to attract and negotiate third-party financing needed to advance a product or technology toward commercialization. Development efforts may include preclinical R&D needed for regulatory filings (e.g., IND or IDE) and/or clinical trials. The Phase IIB Small Market Award focuses on supporting technologies addressing rare diseases or pediatric populations.

The Phase IIB Bridge or Small Market application must represent a continuation of the research and development efforts performed under a previously funded SBIR or STTR Phase II award. The NHLBI welcomes applicants previously funded by any NIH Institute or Center or any other Federal agency, as long as the proposed work applies to the NHLBI mission. Applications may be predicated on a previously funded SBIR or STTR Phase II grant or contract award. Applicants with Phase II contracts or awards from another Federal agency must contact the NHLBI to ensure their application can be received.

Applicants are strongly encouraged to contact Mike Pieck at nhlbi_sbir@mail.nih.gov for additional information.

Limited Amount of Award

For budgetary, administrative, or programmatic reasons, the NHLBI may not fund an application or may decrease the length of an award and/or the budget recommended by a review committee. NIH has received a waiver from SBA, as authorized by statute, to exceed the statutory budget limitations of $256,580 for Phase I and $1,710,531 for Phase II for specific topics relevant to the NHLBI that can be found below. Generally, the NHLBI does not fund Phase I applications greater than $300,000 total costs or project periods greater than 2 years. In addition, the NHLBI does not generally fund Phase II applications greater than $2,000,000 total costs or project periods greater than 3 years. Applicants with budget questions or considering requesting a budget greater than these amounts are strongly encouraged to contact Mike Pieck at http://bit.ly/ContactNHLBIsbir before submitting an application.

NHLBI Topics for Awards over Statutory Budget Limitations

A. Biomedical technologies (medical devices, instruments, pharmaceuticals, drugs, therapeutics, vaccines, diagnostics and biologics) for heart, lung, blood, and sleep related diseases and disorders requiring Federal regulatory approval (FDA) or clearance to be commercialized.

B. Small and large animal testing of products of tissue engineering and regenerative medicine, drugs, medical devices, therapeutics, and biologics and studies involving in vivo animal experiments for heart, lung, blood, and sleep related diseases and disorders.

C. Clinical trials and other experiments involving human subjects for heart, lung, blood, and sleep related diseases and disorders.

D. Therapeutics (drugs, devices, or biologics) development for heart, lung, blood, and sleep related diseases and disorders.

E. Device development for heart, lung, blood, and sleep related diseases and disorders.
F. Diagnostic development for heart, lung, blood, and sleep related diseases and disorders.

G. Investigation of biomarkers and biosignatures of heart, lung, blood, and sleep related diseases and disorders.

H. Technologies to enhance clinical research for heart, lung, blood, and sleep related diseases and disorders.

I. Advanced instrumentation and high throughput tools for biomedical research in heart, lung, blood, and sleep related diseases and disorders.

J. Tools and platforms to improve the dissemination and implementation of evidence-based interventions for heart, lung, blood, and sleep related diseases and disorders.

**Final Progress Reports**

As detailed in NOT-OD-17-085, the NIH has implemented the Final Research Performance Progress Reports (Final RPPR) for SBIR/STTR Final Progress Reports.

The NHLBI is interested in tracking the progress of the small business concerns it funds and the products they develop. Funding priority will be given to those small business concerns that show not only their ability to develop products but also their growth as a small business concern towards independence from the SBIR/STTR program.

**Programs and Services for NHLBI Small Business Awardees**

The NHLBI offers free assistance to applicants and awardees regarding intellectual property, commercialization, and business plan development. Visit https://sbir.nih.gov/nhlbi/pdss to request services.

The NHLBI hosts “Small Biz Hangouts” - a free educational series covering the basics of biomedical technology development. Previous Hangouts are archived on the NHLBI YouTube channel: http://bit.ly/SmallBizHangouts-YouTube

Sign up for the NHLBI listserv (http://bit.ly/NHLBI-SBIR-Updates) to learn about upcoming live events. Learn more about available resources at http://www.nhlbi.nih.gov/about/org/dera/otac/resources.

**Research Topics of Interest to NHLBI**

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,**

- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

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<tr>
<td>Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply</td>
<td>X</td>
<td></td>
<td>For information on non-SBIR/STTR clinical trials funding mechanisms for which small businesses are eligible, please visit the NHLBI clinical trials website</td>
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**NHLBI Non-Clinical Trials Topics:**

**Cardiovascular Sciences**

The Division of Cardiovascular Sciences (DCVS) supports basic, clinical, population, and health services research on the causes, prevention, and treatment of cardiovascular diseases. The research programs of the Division encompass investigator-initiated research, Institute-initiated research in targeted areas of research need and scientific opportunity, specialized centers of research focused on selected research topics, and clinical trials. Research supported by the Division is concerned with the etiology, pathogenesis, prevention, diagnosis, and treatment of coronary artery disease and atherothrombosis; structural heart disease; heart failure and arrhythmias; and hypertension and vascular diseases. A broad array of epidemiological studies is supported by the DCVS to describe disease and risk factor patterns in populations and to identify risk factors for disease. Also supported are clinical trials of interventions to prevent and treat disease; studies of genetic, behavioral, sociocultural, and environmental influences on disease risk and outcomes; and studies of the application of prevention and treatment strategies to determine how to improve clinical care and public health.

**Lung Diseases**
The Division of Lung Diseases (DLD) supports research on the causes, diagnosis, prevention, and treatment of lung diseases and sleep disorders. Research is funded through investigator-initiated and Institute-initiated grant and contract programs in areas including asthma, bronchopulmonary dysplasia, chronic obstructive pulmonary disease, cystic fibrosis, respiratory neurobiology, sleep-disordered breathing, critical care and acute lung injury, developmental biology and pediatric pulmonary diseases, immunologic and fibrotic pulmonary disease, rare lung disorders, pulmonary vascular disease, and pulmonary complications of AIDS and tuberculosis.

Blood Diseases and Resources

The Division of Blood Diseases and Resources (DBDR) supports research on the causes, prevention, and treatment of nonmalignant blood diseases, including anemias, sickle cell disease, and thalassemia; premalignant processes such as myelodysplasia and myeloproliferative disorders; hemophilia and other abnormalities of hemostasis and thrombosis; and immune dysfunction. Research supported by the Division encompasses a broad spectrum of topics ranging from basic biology to medical management of blood diseases. The Division has a major responsibility for research to improve the adequacy and safety of the nation’s blood supply. It also plays a leading role in transfusion medicine and blood banking, including research to evaluate blood donation screening, manufacturing, and processing technologies. The Division also has a major responsibility supporting research in hematopoiesis and stem cell biology and disease. It also supports hematopoietic stem cell transplantation research, and the application of stem cell biology findings to the development of new cell-based therapies to repair and regenerate human tissues and organs.

Center for Translation Research and Implementation Science

The Center for Translation Research and Implementation Science (CTRIS) plans, fosters, and supports an integrated and coordinated program of research to understand the multi-level processes and factors that are associated with successful integration of evidence-based interventions within specific clinical and public health settings such as worksites, communities, and schools; identifies and makes readily available to implementation and dissemination practitioners emergent knowledge about the late phases of translation research, especially the “T4” phase, for rapid and sustained adoption of effective interventions in real world settings; leads the NHLBI effort in the rigorous, systematic evidentiary reviews and subsequent NHLBI participation in the collaborative model for clinical practice guidelines development; supports training and career development of personnel in “T4” translation research and health inequities relating to heart, lung, and blood diseases; provides a focal point for advice and guidance on matters pertaining to minority health, health inequities and minority participation in research; represents the NHLBI to other governments, other Federal Departments and agencies, international organizations, and the private sector on global health issues; and provides data analytics and portfolio analysis to evaluate and inform future directions of implementation research programs.

The NHLBI encourages applications through this Omnibus solicitation proposing innovative technologies related to any area within the NHLBI mission.

The NHLBI maintains a list of notices of special interest (NOSI) (https://www.nhlbi.nih.gov/grants-and-training/funding-opportunities-and-contacts/all-funding-opportunity-announcements) to the Institute. Instructions for submitting applications in response to these topics are posted on the web page. The list is revised throughout the year, so please check regularly for updates. For more information, contact the NHLBI Small Business team at http://bit.ly/ContactNHLBIsbir or the Division contact associated with your technology area listed at the end of the NHLBI section.

NHLBI Clinical Trials Topics:

Cardiovascular Sciences
The Division of Cardiovascular Sciences (DCVS) supports basic, clinical, population, and health services research on the causes, prevention, and treatment of cardiovascular diseases. The research programs of the Division encompass investigator-initiated research, Institute-initiated research in targeted areas of research need and scientific opportunity, specialized centers of research focused on selected research topics, and clinical trials. Research supported by the Division is concerned with the etiology, pathogenesis, prevention, diagnosis, and treatment of coronary artery disease and atherothrombosis; structural heart disease; heart failure and arrhythmias; and hypertension and vascular diseases. A broad array of epidemiological studies is supported by the DCVS to describe disease and risk factor patterns in populations and to identify risk factors for disease. Also supported are clinical trials of interventions to prevent and treat disease; studies of genetic, behavioral, sociocultural, and environmental influences on disease risk and outcomes; and studies of the application of prevention and treatment strategies to determine how to improve clinical care and public health.

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For additional information on research areas, please contact:

**CARDIOVASCULAR SCIENCES**

Albert Lee  
Division of Cardiovascular Sciences  
Advanced Technologies and Surgery Branch  
301-435-0567  
Email: albert.lee3@nih.gov

**LUNG DISEASES and SLEEP DISORDERS**

Lei Xaio  
Division of Lung Diseases  
301-827-7852  
Email: lei.xiao@nih.gov

**BLOOD DISEASES and RESOURCES**

Ron Warren  
Division of Blood Diseases and Resources  
Translational Blood Science and Resources Branch  
301-827-8288  
Email: ronald.warren@nih.gov

**CENTER FOR TRANSLATION RESEARCH and IMPLEMENTATION SCIENCE**

Cheryl Boyce  
Center for Translation Research and Implementation Science  
Translation Research Branch  
301-496-1051  
Email: Cheryl.Boyce@nih.gov
NATIONAL HUMAN GENOME RESEARCH INSTITUTE (NHGRI)

The National Human Genome Research Institute (NHGRI) has been guided, since the inception of the Human Genome Project in 1990, by a sequential series of plans, each of which has been developed with considerable input from the scientific community. These plans have always laid out ambitious goals and measurable objectives to gauge progress. NHGRI initiated a planning process in 2008 and concluded with the publication in February 2011 of the strategic plan, “Charting a Course for Genomic Medicine from Base Pairs to Bedside,” (Nature, 10 February 2011; Volume 470). The phenomenal advances that have marked genomics and have allowed genomic applications to transform many important fields have made it an opportune time for the Institute to take a new look at genomics and its future. A new strategic planning process is underway for publication in October 2020.

Research Topics of Interest to NHGRI

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### NHGRI Non-Clinical Trials Topics:

#### A. Technology and Methods Development

Technology development in DNA sequencing and genotyping are examples of activities that have changed the nature of what scientific research questions are practical to address, have enabled new approaches, and have facilitated the development of new community resource data sets. Many areas of critical importance to the realization of the genomics-based vision for biomedical research require continued technological and methodological developments before pilots and then large-scale approaches can be attempted. Accordingly, the NHGRI will continue to support the development of new, fundamental technologies in all areas of genomics. Important areas in which technology development applications would be responsive to this Program Announcement include (but are not limited to) experimental technologies and computational methods to analyze gene expression and other molecular phenotypes; discovery and characterization of genetic variation; identification of the genetic contributions to health, disease, and drug response; statistical analytic methods for understanding human genomic variation and its relationship to health and disease; and chemical genomics. There is also continued need to support technology development for the comprehensive discovery of functional elements in the human and model organism genomes, and new DNA sequencing technology. Many of these assays would benefit from the ability to work with very small amounts of starting material, to the limit of single cells, along with minimally-invasive human specimens that are easy to collect, handle, and store. As these technologies mature, emphasis should be on high throughput, cost-effective methods that consistently produce very high-quality data.

The Institute also places high priority on contributing selectively to the development of new and needed technology in related areas, such as proteomics and systems biology research, when NHGRI funding can be used to further a truly unique development that will have a significant impact on the field.

Further information on opportunities related to technology and methods development is available at this website: [https://www.genome.gov/10000368/](https://www.genome.gov/10000368/).

#### B. Bioinformatics, Computational Genomics and Data Science

The ongoing development of new sequencing technologies has dramatically increased the amount of data produced for genomics in basic science and translation to medicine. NHGRI encourages new computational approaches for the analysis, visualization and integration of genomic information in basic and clinical research and in applications to improve its utility in healthcare. These approaches may include the development of methods for processing, annotating, interpreting, analyzing and sharing of sequencing data with associated phenotypes.
and other large-scale genomic data sets such as haplotype maps, genetic variants, transcriptome measurements, functional elements, and in some cases protein interactions. NHGRI also encourages the development of better computational solutions for storage, access, compression, secure sharing, privacy and transfer of large genomic datasets by biomedical researchers.

Some genomic data analysis and display tools have been developed that already are used in the community but would benefit from additional work to support broader dissemination, for example making them efficient, reliable, robust, well-documented, and well-supported, or for deploying them in containers or at scale in a cloud-based platform. NHGRI will support projects to extend the support for these informatics tools to make them more easily adopted by any biomedical research laboratory that wishes to use genomic technologies to address biomedical questions.

Genomic databases are essential resources for the biological and biomedical research communities and relevant regulatory agencies. The creation and maintenance of effective and sustainable databases that take advantage of technology improvements are an important component of NHGRI research funding strategy.

Where possible, existing or emerging community data standards, models, and methods for data representation and exchange should be used in the development of these new methods and tools as well as other approaches to enhance reproducibility. Standards-based approaches are also encouraged to integrate and share genomics and phenotype data for data mining with other sources including for clinical application. Projects that will make genomic digital objects Findable, Accessible, Interoperable, Reusable (FAIR) in the broader community are highly recommended.

Further information on programs related to NHGRI supported research in computational genomics and data science is available at this web site: http://www.genome.gov/10001735.

C. Population Genomics and Genomic Medicine

Population genomics applies genomic technologies, such as genome-wide association testing and sequencing, to population studies to identify genes or variants that affect common etiologically complex conditions and predict individual risk. Genomic medicine investigates the value of applying genomic methods in clinical care for the diagnosis, treatment, and prevention of diseases. The research scope of Population Genomics and Genomic Medicine at NHGRI includes: developing resources and statistical methods for observational studies and clinical trials incorporating advanced genomic technologies; conducting proof-of-principle studies that apply genomic technologies to epidemiologic and clinical research; developing research methods and infrastructure needed for future epidemiologic and clinical studies of genetic and environmental contribution to disease; investigations of whether and how clinical genome sequencing impacts disease diagnosis and treatment; studies of approaches to improve the identification and interpretation of genomic variants for dissemination in clinical settings; assessing phenotypic manifestations of genetic variants through electronic medical records (EMRs); integrating genomic results and clinical decision support into EMRs; studies that address current barriers to the implementation of clinical genome sequencing, and assessing the impact of genetic information on clinical utility, health outcomes and delivery of care. For additional information about Genomic Medicine NHGRI, please visit this web site: http://www.genome.gov/27550079.

D. Ethical, Legal and Social Implications

NHGRI, through the ELSI Research Program, supports research studies that examine issues and, where appropriate, develop policy options regarding the ethical, legal and social implications of genomics. These studies may focus on issues associated with genomic research, genomic medicine or broader societal effects of genomic information and technologies. More detailed
information on specific ELSI research priorities within each of these broad areas is available on the ELSI Research priorities web site: http://www.genome.gov/27543732.

**NHGRI Clinical Trials Topics:**

The National Human Genome Research Institute (NHGRI) will accept applications designated as clinical trials for all program areas supported by the Institute as outlined above for non-clinical trials SBIRs/STTRs. The broadened definition of clinical trials as defined in NOT-OD-15-015 and https://grants.nih.gov/policy/clinical-trials/definition.htm is not intended to expand the scope of applications accepted by NHGRI beyond studies that have a major genomic or Ethical, Legal and Social Implications (ELSI) component and relate clearly to NHGRI’s mission. Information on areas of research interest is available on the NHGRI Research Funding Divisions homepage: https://www.genome.gov/27552836/nhgri-research-funding-divisions/ and the ELSI Research Domains website: https://www.genome.gov/27543732/elsi-research-domains/. Additionally, applicants are strongly encouraged to discuss their research plans with NHGRI Program Staff prior to submitting their application.

For more information on research topics, contact:

Michael W. Smith, Ph.D.
Technology Development Program &
Small Business Lead
Coordinator SBIR Grants
301-496-7531
Email: smithmw@mail.nih.gov

Heidi Sofia, Ph.D.
Informatics & Genomic Medicine
Coordinator STTR Grants
301-496-7531
Email: Heidi.Sofia@nih.gov
NATIONAL INSTITUTE OF MENTAL HEALTH (NIMH)

The mission of the National Institute of Mental Health (NIMH) is to transform the understanding and treatment of mental illnesses through basic and clinical research, paving the way for prevention, recovery, and cure. Mental disorders constitute an immense burden on the U.S. population, with suicide as one of the leading causes of death in the US, major depression, the leading cause of disability in the U.S., and schizophrenia, bipolar disorder, and obsessive-compulsive disorder ranked among the ten leading causes of disability. NIMH also takes a leading role in understanding the impact of behavior on HIV transmission and pathogenesis, and in developing effective behavioral preventive interventions. The NIMH conducts a wide range of research, research training, research capacity development, as well as public information outreach and dissemination to fulfill its mission.

For the Institute to continue fulfilling this vital public health mission, it must foster innovative thinking and ensure that a full array of novel scientific perspectives are used to further discovery in the evolving science of brain, behavior, and experience. In this way, breakthroughs in science can become breakthroughs for all people with mental illnesses.

The NIMH SBIR/STTR programs support small businesses to develop technologies that can advance the mission of the Institute, including in basic neuroscience research relevant to mental disorders, translational and clinical research of mental disorders, clinical diagnosis or treatment of mental disorders, and dissemination of evidence-based mental health care. The NIMH Strategic Plan (http://www.nimh.nih.gov/about/strategic-planning-reports/index.shtml) and the National Advisory Mental Health Council’s workgroup report “From Discovery to Cure” http://www.nimh.nih.gov/about/advisory-boards-and-groups/namhc/reports/fromdiscoverytocure_103739.pdf present key scientific priorities across these domains, and describe the need for technologies to realize these priorities. Research priorities for the NIMH further include aspects of HIV/AIDS prevention, treatment, and care, in accordance with the Trans-NIH Plan for HIV-Related Research (http://www.oar.nih.gov/strategicplan/).

For additional information about areas of interest to the NIMH, please visit our home page at http://www.nimh.nih.gov. For additional information on NIMH requirements for clinical trials, please visit: https://www.nimh.nih.gov/funding/opportunities-announcements/clinical-trials-foas/index.shtml.

Also visit the NIMH SBIR/STTR home page: http://www.nimh.nih.gov/research-funding/small-business/index.shtml.

Important notes:

1. Potential SBIR/STTR applicants should contact NIMH prior to submitting an application to ensure the application is of priority/interest to NIMH. Please see the contacts section.

2. An additional criterion that the federal government considers in supporting a small business with SBIR/STTR funds, is past commercialization performance. It is expected that small businesses who have received previous SBIR/STTR grants, have had success in commercializing their previously supported technologies. Small businesses that are mostly interested in research and development (and not commercialization) should consider other grant mechanisms at NIH, rather than the SBIR/STTR program. Program staff at NIMH can help identify the most appropriate grant mechanism to use.

3. The NIH has received a waiver from the SBA, regarding the funding cap. The technology areas that are included in this waiver can be found in the topic list located in Appendix A of this document. The technologies listed in the Appendix A (under NIMH) are of priority to this institute.

Phase IIB Competing Renewal Awards

The NIMH will accept Phase IIB SBIR/STTR Competing Renewal grant applications in two categories: 1) to continue research and development of technologies that ultimately require federal regulatory approval,
and 2) to continue research and development of complex instrumentation, clinical research tools, or behavioral/digital health interventions and treatments.

Technologies in the former category (those that ultimately require federal regulatory approval) include, but are not limited to: pharmacologic agents and drugs, biological products, medical devices, vaccines, etc. related to the mission of the NIMH. Phase IIB SBIR/STTR Competing Renewal grants for such technologies should allow small businesses to move research and development to a stage where interest and investment by third parties is more likely.

Companies that are developing technologies that do not focus on drug development, but that require federal regulatory approval prior to commercialization, may be eligible to submit a Phase IIB Competing Renewal application.

For both technology areas, Phase IIB applications may be submitted through the Omnibus SBIR/STTR funding opportunity announcement. Generally, for this opportunity, budget limits of $3 million total costs and time periods up to 3 years may be requested. These budget allowances have been approved by the SBA through a waiver.

The following examples would make appropriate topics for proposed NIMH SBIR/STTR Phase IIB Competing Renewal projects. These are meant for illustrative purposes only and are not exclusive of other appropriate activities:

- Preclinical studies, including pharmacology and toxicology, beyond those conducted under the Phase I (R43/R41) and initial Phase II (R44/R42) grants. Some in vivo or in vitro studies would be expected to have been carried out in Phase I or the initial Phase II grant.
- Completion of studies as required by the Food and Drug Administration (FDA) for Investigational New Drug (IND) or Radioactive Drug Research Committee (RDRC) application.
- Studies in normal healthy volunteers to determine a drug’s safety profile, metabolism, etc.
- Assessment of devices with regard to performance standards related to the FDA approval process.
- Safety and effectiveness studies of novel medical devices.
- Evaluation of novel imaging approaches for diagnostic purposes.
- Clinical studies in support of Pre-Market Approval for biomarkers/medical devices by the FDA.

Although technologies in the latter category listed above (complex instrumentation, clinical research tools, or behavioral interventions/treatments) may not require federal regulatory approval, extraordinary time and effort is needed for their research and development. Therefore, NIMH supports Phase IIB Competing Renewal awards of existing Phase II grants for such technologies. The Phase IIB Competing Renewal award for these would provide up to an additional three years of support at total cost funding levels of up to generally $3 million for the project. These budget allowances have been approved by the SBA through a waiver.

Please contact the Program Director in the appropriate Division or Dr. Margaret Grabb (listed below) before beginning the process of putting an application together. In addition, prospective applicants are encouraged to submit to the program contact a letter of intent that includes the following information:

- Descriptive title of the proposed research
- Name, address, and telephone number of the Principal Investigator
- Names of other key personnel
- Participating institutions
- Funding Opportunity Announcement (e.g. PA-19-273).

Although a letter of intent is not required, is not binding, and does not enter into the review of a subsequent application, the information that it contains allows NIH staff to estimate the potential review
workload and plan the review. It is expected that only a portion of NIMH SBIR Phase II awards will be eligible for a Phase IIB Competing Renewal grant.

Research Topics of Interest to NIMH

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
<th>No*</th>
<th>Yes**</th>
</tr>
</thead>
<tbody>
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<td></td>
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<td>X</td>
</tr>
</tbody>
</table>

*If No,
- This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, UNLESS the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,
- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency accept clinical trials applications under this mechanism?</th>
<th>Yes</th>
<th>No</th>
<th>Other Information: If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Trials SBIR/STTR Omnibus/Parent Funding Opportunity Announcement/s</td>
<td>X</td>
<td>NIMH will prioritize funding for SBIR/STTR applications with a clinical trial focus that are consistent with the stated research goals and priorities relevant to clinical trials as outlined in the clinical trials FOAs. For more information see: [<a href="https://www.nimh.nih.gov/funding/opportunities-announcements/c">https://www.nimh.nih.gov/funding/opportunities-announcements/c</a> clinical-trials-foas/support-for-clinical-trials-at-nimh.shtml](<a href="https://www.nimh.nih.gov/funding/opportunities-announcements/c">https://www.nimh.nih.gov/funding/opportunities-announcements/c</a> clinical-trials-foas/support-for-clinical-trials-at-nimh.shtml)</td>
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<tr>
<td>Clinical Trials SBIR/STTR IC/Office/Agency - Specific Funding Opportunity Announcement/s</td>
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<td>NIMH will prioritize funding for SBIR/STTR applications with a clinical trial focus that are consistent with the stated research goals and priorities relevant to clinical trials as outlined in the clinical trials FOAs. For more information see: [<a href="https://www.nimh.nih.gov/funding/opportunities-announcements/c">https://www.nimh.nih.gov/funding/opportunities-announcements/c</a> clinical-trials-foas/support-for-clinical-trials-at-nimh.shtml](<a href="https://www.nimh.nih.gov/funding/opportunities-announcements/c">https://www.nimh.nih.gov/funding/opportunities-announcements/c</a> clinical-trials-foas/support-for-clinical-trials-at-nimh.shtml)</td>
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</tr>
<tr>
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<tr>
<td>Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply</td>
<td>X</td>
<td></td>
<td>NIMH will prioritize funding for SBIR/STTR applications with a clinical trial focus that are consistent with the stated research goals and priorities relevant to clinical trials as outlined in the clinical trials FOAs. For more information see: <a href="https://www.nimh.nih.gov/funding/opportunities-announcements/clinical-trials-foas/support-for-clinical-trials-at-nimh.shtml">https://www.nimh.nih.gov/funding/opportunities-announcements/clinical-trials-foas/support-for-clinical-trials-at-nimh.shtml</a> and: • First in Human and Early Stage Clinical Trials of Novel Investigational Drugs or Devices for Psychiatric Disorders (U01 Clinical Trial Required) <a href="#">PAR-18-427</a> • Early Stage Testing of Pharmacologic or Device-based Interventions for the Treatment of Mental Disorders (R61/R33- Clinical Trial Required) <a href="#">RFA-MH-18-702</a> • Early Stage Testing of Pharmacologic or Device-based Interventions for the Treatment of Mental Disorders (R33- Clinical Trial Required) <a href="#">RFA-MH-18-703</a> • Development of Psychosocial Therapeutic and Preventive Interventions for Mental Disorders (R61/R33- Clinical Trial Required) <a href="#">RFA-MH-18-704</a> • Development of Psychosocial Therapeutic and Preventive Interventions for Mental Disorders (R33 Clinical Trial Required) <a href="#">RFA-MH-18-705</a></td>
</tr>
</tbody>
</table>
Does IC/Office/Agency accept clinical trials applications under this mechanism?  Yes  No  Other Information: If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information.

Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply

- Confirmatory Efficacy Clinical Trials of Non-Pharmacological Interventions for Mental Disorders (R01 Clinical Trial Required)  RFA-MH-18-707
- Pilot Effectiveness Trials for Treatment, Preventive and Services Interventions (R34- Clinical Trial Required)  RFA-MH-18-706
- Clinical Trials to Test the Effectiveness of Treatment, Preventive, and Services Interventions (R01 Clinical Trial Required)  RFA-MH-18-701
- Clinical Trials to Test the Effectiveness of Treatment, Preventive, and Services Interventions (Collaborative R01 - Clinical Trial Required)  RFA-MH-18-700

NIMH Non-Clinical Trials Topics:

**Division of Neuroscience and Basic Behavioral Science (DNBBS)**

The Division of Neuroscience and Basic Behavioral Science provides support for research programs in the areas of basic neuroscience, genetics, basic behavioral science, research training, resource development, technology development, drug discovery, and research dissemination. The Division has the responsibility, in cooperation with other components of the Institute and the research community, for ensuring that relevant basic science knowledge is generated and then harvested to create improved diagnosis, treatment, and prevention of mental and behavioral disorders.

In this Division, the SBIR and STTR programs support research and the development of tools related to basic brain and behavioral science, genetics, and drug discovery and development relevant to the mission of the NIMH. Such tools include: software (such as informatics tools and resources and tools for analyzing data); hardware (such as the development of instrumentation or devices); wetware (such as the use of iRNAs or other bioactive agents as research tools or molecular imaging agents or genetic approaches to label neural circuits or modify circuit functions); and drug discovery related technologies such as high throughput screening (HTS) or computational pharmacology approaches.

**AREAS OF EMPHASIS**

- Novel imaging probes to study brain structure and function at all levels, from the molecular level to the whole organ, using any imaging modality (PET, fMRI, optical, etc.).

- Drug discovery/drug development of novel compounds which act on molecular pathways (receptors, enzymes, second messengers, etc.) that are not typically targeted with currently available psychiatric drugs, and that have a strong biological justification as a novel mechanism for treatment of psychiatric disorders.

- Novel screening assays for high throughput acquisition and analysis of data about behavior and the brain, from the level of genes to behavior.
• Novel technologies that would enable researchers to study how populations of neural cells work together within and between brain regions, in order to understand how changes in neural activity contributes to mental disorders.

• Develop informatics tools to facilitate the analysis and sharing of data between laboratories about behavior and the brain. This could include common data element efforts, but is not limited to that area.

• Technologies consistent with the goals of the BRAIN Initiative: [http://www.braininitiative.nih.gov/](http://www.braininitiative.nih.gov/).

Prospective applicants are strongly encouraged to contact Dr. Margaret Grabb (listed below) with questions about the relevance of their interests to the mission of this division.

**Division of Translational Research (DTR)**

The DTR directs, plans, and supports programs of research and research training that translate knowledge from basic science to discover the etiology, pathophysiology, and trajectory of mental disorders and develops effective interventions for children and adults. DTR supports integrative, multidisciplinary research on the following areas: the phenotypic characterization and risk factors for psychiatric disorders; neurobehavioral mechanisms of psychopathology; trajectories of risk and resilience based on the interactive influences of genetics, brain development, environment, and experience; and design and testing of innovative psychosocial, psychopharmacologic, and somatic treatment interventions.

In this Division, the SBIR and STTR Programs support research aimed at facilitating the validation and commercialization of new methods of assessing psychopathology and measuring treatment response to therapeutic agents. In addition, the SBIR and STTR Programs support the clinical development of interventions, including novel pharmacologic agents or brain stimulation devices as well as technology development used to deliver novel psychosocial approaches to the treatment of mental illness in adults, pediatrics and geriatrics. For more information on NIMH supported clinical trials and requirements, see: [https://www.nimh.nih.gov/funding/opportunities-announcements/clinical-trials-foas/index.shtml](https://www.nimh.nih.gov/funding/opportunities-announcements/clinical-trials-foas/index.shtml)

**AREAS OF EMPHASIS**

• Develop, test, and validate biological markers (e.g., genetic, proteomic, imaging) for diagnosing or detecting risk/vulnerability, onset, progression, and/or severity of mental disorders, or to evaluate treatment response.

• Develop valid measures of the various constructs in the Research Domain Criteria (RDoC) matrix (see [https://www.nimh.nih.gov/research/research-funded-by-nimh/rdoc/index.shtml](https://www.nimh.nih.gov/research/research-funded-by-nimh/rdoc/index.shtml)), e.g., behavioral tasks, psychometrically sophisticated self-report measures, and measures of physiological and neural activity, into a commercial product.

• Web-based tools and biosensors to enhance prevention, early identification and treatment of pediatric mental disorders by various educational and health professionals.

• Development of hardware and software tools to support operations of multi-site clinical trials.

• Development of novel methods to enhance efficiency of early phase clinical trials.

• Novel technologies and data analytic tools to enable quantification of behavioral data that is relevant to research or clinical trials in mental disorders and/or autism.

• Development of imaging technologies that can reveal specific pathologies in major mental disorders.
Prospective applicants are strongly encouraged to contact Dr. Margaret Grabb (listed below) with questions about the relevance of their interests to the mission of this division.

**Division of AIDS Research (DAR)**

The NIMH DAR supports scientific research to understand and alleviate the consequences of HIV infection on the central nervous system, and research to strengthen the provision and outcomes of HIV/AIDS prevention and treatment. Examples of high-priority research areas for SBIR/STTR applications are described below.

- Develop and test novel, non-invasive diagnostic approaches (instrumentation, imaging, biomarkers, central nervous system [CNS] cell-based *in vitro* models) to detect HIV-1 associated CNS dysfunction and innovative technologies to study the mechanisms involved in HIV-1 associated neuropathogenesis and persistence of HIV-1 in the CNS.

- Design and test novel therapeutic interventions aimed at amelioration of HIV-1 associated CNS dysfunction, and/or eradication of HIV-1 from CNS reservoirs, and/or strategies to prevent viral resurgence in the CNS upon cessation of anti-retroviral therapy.

- Tools to assess neurotoxicity profiles of antiretroviral medications and pharmacological strategies to reduce adverse effects of anti-retroviral drugs (neuropsychiatric side effects and drug-drug interactions).

- Develop new tools/techniques to aid in deciphering the complex neuro-immune interactions at a molecular and cellular level in the context of HIV.

- Develop or adapt neurological/neuropsychological/neurobehavioral assessments to evaluate HIV-1 associated abnormalities in adults or children in resource limited environments that are adaptable to different cultures and languages. Designing neurobehavioral assessments could occur retrospectively (not a clinical trial – using EMR).

- Build and optimize informatics tools to aid in analyzing and characterizing the phenotype of CNS disease modalities associated with HIV by using machine learning, big data and systems biology-based approaches.

- Develop technologies, instruments and tools to aid in improving uptake, adherence, and persistence to biomedical HIV prevention and treatment regimens; Increasing regular HIV testing among those most at risk of acquiring HIV and translating findings from basic behavioral and social science research into processes to improve engagement in HIV care.

- Develop innovative approaches to improve the scientific assessment of HIV exposure due to sexual behavior or the scientific assessment of HIV medication adherence through wireless technologies, remote sensing devices, biomarkers, assays, or other novel methods. Designing approaches for assessment could occur retrospectively (not a clinical trial – using EMR).

- Develop and test tools, curricula, and strategies that seek to reduce documented racial/ethnic, gender, and age-related disparities in HIV infection, HIV testing, HIV care engagement, or in HIV treatment adherence and treatment outcomes. Designing strategies for reducing disparities could occur retrospectively (not a clinical trial – using EMR).

- Develop novel tools and approaches to identify, recruit, enroll, and/or retain those most vulnerable to HIV/AIDS (e.g., African-American MSM, adolescents) in HIV prevention research and/or services. Developing tools for recruitment and retention of key populations could occur retrospectively (not a clinical trial – using EMR).
• Develop and test tools, curricula, and strategies to identify the potent, modifiable mechanisms and processes linking mental health and HIV and developing interventions to address the high prevalence of mental health disorders among those living with HIV and those at substantial risk for HIV infection.

Prospective applicants are strongly encouraged to contact Dr. Vasudev R Rao (listed below) with questions about the relevance of their interests to the mission of this division.

Division of Services and Intervention Research (DSIR)

The Division of Services and Intervention Research (DSIR) SBIR/STTR supports two critical areas of research:

• Intervention research to evaluate the effectiveness of pharmacologic, psychosocial, somatic, rehabilitative and combination interventions on mental and behavior disorders—including acute and longer-term therapeutic effects on functioning across domains for children, adolescents, and adults.

• Mental health services research to improve the access, continuity, cost, quality and outcomes of mental health care, as well as to improve the dissemination and implementation of effective interventions in clinical and community settings, to strengthen the public health impact of NIMH research.

The intervention research program addresses the effectiveness of treatment and preventive interventions in usual practice and community settings with the purpose of informing clinicians, patients, families, and health policy makers on evidence-based practices. In funding decisions, special emphasis is placed on the potential clinical impact of the research activities and on the implications of the research findings for improving community practice and health outcomes. Types of interventions include the full range of behavioral, psychotherapeutic, pharmacologic, and non-pharmacologic somatic or alternative interventions, as well as rehabilitation or other adjunctive services, e.g., integrated approaches to chronic mental illness. Examples of areas of interest are:

• Analyses of naturalistic databases to evaluate the effectiveness of preventive and treatment interventions.

• Identifying moderators and mediators of intervention effects as a step to design and test personalized interventions. **Moderator/mediator identification could occur retrospectively (not a clinical trial – using EMR).**

• Evaluating the combined or sequential use of interventions. **Evaluation of combined/sequential interventions could occur retrospectively (not a clinical trial – using EMR).**

• Determining the optimal duration, frequency and intensity of an intervention to optimize improvements in symptoms and functioning, establishing the utility of continuation or maintenance treatment (that is, for prevention of relapse or recurrence). **Evaluation of the optimal length of an intervention could occur retrospectively (not a clinical trial – using EMR).**

• Evaluating the long-term impact of therapeutic interventions on symptoms, functioning, and quality of life. **Evaluation of the optimal length of an intervention could occur retrospectively (not a clinical trial) – using EMR or survey data.**

Services research covers all mental health services across the lifespan for all mental health disorders and aligns with NIMH strategic research priority 4 ([https://www.nimh.nih.gov/about/strategic-planning-reports/strategic-research-priorities/srp-objective-4/index.shtml](https://www.nimh.nih.gov/about/strategic-planning-reports/strategic-research-priorities/srp-objective-4/index.shtml)), which includes but is not limited to:
• Service settings at the patient, provider, health system, and cross system levels to include primary care, specialty mental health, integrated care, general health, and other delivery settings (such as employment, educational, veteran, military, and criminal justice settings).

• Enhanced capacity for conducting services research by developing and utilizing innovative and established methodologies, including health economics, to inform decisions about service delivery and financing of care.

• The clinical epidemiology of mental disorders to include development and use of data sets from health surveillance activities, decision support tools, administrative claims, mobile apps and similar technologies, electronic health record, disease registries, and other databases where epidemiological data (to include big data) reside.

For both interventions and services research, DSIR supports the development and testing of digital health tools. These tools include technology-assisted approaches to assessment (e.g., technology-assisted screening and diagnosis) and intervention (e.g., m-health and other technology platforms to support the delivery of preventive, therapeutic, and services interventions). DSIR encourages efforts to employ technology-assisted approaches to expand the reach, efficiency, continuity, quality, and/or boost the therapeutic benefit of research-informed strategies, rather than mere translation of research-supported strategies onto new or emerging technology platforms. Collaboration with NIMH supported researchers for the development of software for new analytic techniques and/or decision-making algorithms is encouraged. Also supported is research and the development or adaptation of tools and technologies to be used to enhance the training and development of new generations of researchers and practitioners and to keep established researchers and practitioners up-to-date on the findings, implementation, and methods of interventions and services research.

Prospective applicants are strongly encouraged to contact Dr. Adam Haim (listed below) with questions about the relevance of their interests to the mission of this division.

NIMH Clinical Trials Topics:

Division of Neuroscience and Basic Behavioral Science (DNBBS)

AREAS OF EMPHASIS

• Novel imaging probes to study brain structure and function at all levels, from the molecular level to the whole organ, using any imaging modality (PET, fMRI, optical, etc.), in human studies.

• First in human drug trials.

• Novel screening assays for high throughput acquisition and analysis of data about behavior and the brain, from the level of genes to behavior.

• Novel technologies that would enable researchers to study how populations of neural cells work together within and between brain regions, in order to understand how changes in neural activity contributes to mental disorders, when applied to humans.

• Human/clinical-based technologies consistent with the goals of the BRAIN Initiative: http://www.braininitiative.nih.gov/.

Prospective applicants are strongly encouraged to contact Dr. Margaret Grabb (listed below) with questions about the relevance of their interests to the mission of this division.

Division of Translational Research (DTR)
AREAS OF EMPHASIS

- Develop valid measures of the various constructs in the Research Domain Criteria (RDoC) matrix (see http://www.nimh.nih.gov/research-funding/rdoc/index.shtml), e.g., behavioral tasks, psychometrically sophisticated self-report measures, and measures of physiological and neural activity, into a commercial product.

- Conduct early stage, proof of concept clinical trials to advance the development of novel therapeutics. The clinical trials are expected to include biological/behavioral data to assess target engagement and to help determine potential success or failure of the compound before moving on to larger clinical trials (see NOT-MH-11-015 http://grants.nih.gov/grants/guide/notice-files/NOT-MH-11-015.html).

- Develop, test and validate reliable and stable biomarkers that can identify at-risk individuals prior to disease onset, improve diagnosis and classification, predict treatment response, or to measure disease progression. Biomarkers are also needed in clinical trials to identify dose ranges, to identify a specific subpopulation of subjects to enroll in a treatment trial, or to measure efficacy or toxicity/side effects.

- Development of novel diagnostic tools and innovative measures of treatment response and disease progression, preclinical or clinical efficacy testing, or toxicity measures for drug development.

- Development of hardware and software tools to enable refined physiological and behavioral assessment of normal and atypical neurodevelopment focused on pediatrics, adult and geriatric age ranges.

- Web-based tools and biosensors to enhance prevention, early identification and treatment of pediatric mental disorders by various educational and health professionals.

- Development of hardware and software tools to support operations of multi-site clinical trials.

- Development of novel methods to enhance efficiency of early phase clinical trials.

- Novel technologies and data analytic tools to enable quantification of behavioral data that is relevant to research or clinical trials in mental disorders and/or autism.

- Development of imaging technologies that can reveal specific pathologies in major mental disorders.

Prospective applicants are strongly encouraged to contact Dr. Margaret Grabb (listed below) with questions about the relevance of their interests to the mission of this division.

Division of AIDS Research (DAR)

The NIMH DAR supports scientific research to understand and alleviate the consequences of HIV infection on the central nervous system, and research to strengthen the provision and outcomes of HIV/AIDS prevention and treatment. Examples of high-priority research areas for SBIR/STTR applications are described below.

- Develop and test novel, non-invasive diagnostic approaches (instrumentation, imaging, biomarkers, central nervous system [CNS] cell-based in vitro models) to detect HIV-1 associated CNS dysfunction and innovative technologies to study the mechanisms involved in HIV-1 associated neuropathogenesis and persistence of HIV-1 in the CNS.
• Design and test novel therapeutic strategies aimed at amelioration HIV-1 associated CNS dysfunction, and/or eradication of HIV-1 from CNS reservoirs, and/or strategies to prevent viral resurgence in the CNS upon cessation of anti-retroviral therapy

• Tools to assess neurotoxicity profiles of antiretroviral medications and pharmacological strategies to reduce adverse effects of anti-retroviral drugs (neuropsychiatric side effects and drug-drug interactions).

• Develop or adapt domain based neurological/neuropsychological/neurobehavioral assessments to evaluate HIV-1 associated abnormalities in adults or children in resource limited environments that are adaptable to different cultures and languages.

• Build and optimize informatics tools to aid in analyzing and characterizing the phenotype of CNS disease modalities associated with HIV by using machine learning, big data and systems biology-based approaches

• Develop technologies, instruments and tools to aid in improving uptake, adherence, and persistence to biomedical HIV prevention and treatment regimens; Increasing regular HIV testing among those most at risk of acquiring HIV and translating findings from basic behavioral and social science research into processes to improve engagement in HIV care

• Develop innovative approaches to improve the scientific assessment of HIV exposure due to sexual behavior or the scientific assessment of HIV medication adherence through wireless technologies, remote sensing devices, biomarkers, assays, or other novel methods.

• Develop and test tools, curricula, and strategies that seek to reduce documented racial/ethnic, gender, and age-related disparities in HIV infection, HIV testing, HIV care engagement, or in HIV treatment adherence and treatment outcomes.

• Develop novel tools and approaches to identify, recruit, enroll, and/or retain those most vulnerable to HIV/AIDS (e.g., African-American MSM, adolescents) in HIV prevention research and/or services.

• Develop and test tools, curricula, and strategies to identify the potent, modifiable mechanisms and processes linking mental health and HIV and developing interventions to address the high prevalence of mental health disorders among those living with HIV and those at substantial risk for HIV infection

Prospective applicants are strongly encouraged to contact Dr. Vasudev R. Rao (listed below) with questions about the relevance of their interests to the mission of this division.

**Division of Services and Intervention Research (DSIR)**

The Division of Services and Intervention Research (DSIR) supports two critical areas of research:

• Intervention research to evaluate the effectiveness of pharmacologic, psychosocial, somatic, rehabilitative and combination interventions on mental and behavior disorders-including acute and longer-term therapeutic effects on functioning across domains for children, adolescents, and adults.

• Mental health services research to improve the access, continuity, cost, quality and outcomes of mental health care, as well as to improve the dissemination and implementation of effective interventions in clinical and community settings, to strengthen the public health impact of NIMH research.
The intervention research program addresses the effectiveness of treatment and preventive interventions in usual practice and community settings with the purpose of informing clinicians, patients, families, and health policy makers on evidence-based practices. In funding decisions, special emphasis is placed on the potential clinical impact of the research activities and on the implications of the research findings for improving community practice and health outcomes. Types of interventions include the full range of behavioral, psychotherapeutic, pharmacologic, and non-pharmacologic somatic or alternative interventions, as well as rehabilitation or other adjunctive services, e.g., integrated approaches to chronic mental illness. Examples of areas of interest are:

- Randomized clinical trials evaluating the effectiveness of preventive and treatment interventions.
- Identifying moderators and mediators of intervention effects as a step to design and test personalized interventions. Moderator/mediator identification could occur prospectively (within the context of a clinical trial).
- Evaluating the combined or sequential use of interventions. Evaluation of combined/sequential interventions could occur prospectively (within the context of a clinical trial).
- Determining the optimal duration, frequency and intensity of an intervention to optimize improvements in symptoms and functioning, establishing the utility of continuation or maintenance treatment (that is, for prevention of relapse or recurrence). Evaluation of the optimal length of an intervention could occur prospectively (within the context of a clinical trial).
- Evaluating the long-term impact of therapeutic interventions on symptoms, functioning, and quality of life. Evaluation of the optimal length of an intervention could occur prospectively (within the context of a clinical trial).

Services research covers all mental health services across the lifespan for all mental health disorders and aligns with NIMH strategic research priority 4 (https://www.nimh.nih.gov/about/strategic-planning-reports/strategic-research-priorities/srp-objective-4/index.shtml), which includes but is not limited to:

- Interventions and other research to improve access, continuity, engagement, quality, uptake, equity, efficiency, and cost of care.
- The dissemination and implementation of evidence-based interventions, programs, support tools, or other practices or technologies into service settings.

For both interventions and services research, DSIR supports the development and testing of digital health tools. These tools include technology-assisted approaches to assessment (e.g., technology-assisted screening and diagnosis) and intervention (e.g., m-health and other technology platforms to support the delivery of preventive, therapeutic, and services interventions). DSIR encourages efforts to employ technology-assisted approaches to expand the reach, efficiency, continuity, quality, and/or boost the therapeutic benefit of research-informed strategies, rather than mere translation of research-supported strategies onto new or emerging technology platforms. Collaboration with NIMH supported researchers for the development of software for new analytic techniques and/or decision-making algorithms is encouraged. Also supported is research and the development or adaptation of tools and technologies to be used to enhance the training and development of new generations of researchers and practitioners and to keep established researchers and practitioners up-to-date on the findings, implementation, and methods of interventions and services research. Prospective applicants are strongly encouraged to contact Dr. Adam Haim (listed below) with questions about the relevance of their interests to the mission of this division.
Program Contacts

Margaret Grabb, Ph.D. (general questions about the NIMH SBIR program, Phase IIB program, DNBBS, DTR divisional interests)
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NATIONAL INSTITUTE ON MINORITY HEALTH AND HEALTH DISPARITIES (NIMHD)

The mission of the National Institute on Minority Health and Health Disparities (NIMHD) is to promote minority health and to lead, coordinate, support, and assess the National Institutes of Health (NIH) effort to reduce and ultimately eliminate health disparities. In this effort, the NIMHD conducts and supports basic, clinical, social and behavioral research; facilitates the development of research infrastructure and training; fosters emerging programs; and reaches out to racial/ethnic minority and other health disparity communities.

The Small Business Innovation Research (SBIR) Program and the Small Business Technology Transfer (STTR) Program enable the Nation’s small businesses to apply their unique research and development capabilities toward accomplishing NIMHD’s mission. NIMHD has developed the NIMHD Research Framework (https://ajph.aphapublications.org/doi/pdf/10.2105/AJPH.2018.304883) and small businesses are encouraged to consider the factors operating within and across the frameworks’ multiple ecosocial levels and domains before initiating the design of products for potential research and development by NIMHD. The framework, initially develop for researchers, can also inform small businesses research and development of new technologies and products for improving, sustaining or enhancing minority health and extending longevity. It can also inform the research and development of technologies and products for reducing or eliminating health disparities. The factors identified in the framework are known to contribute to the creation, recreation, and perpetuation and replication of poor minority health and health disparities over time and place. Entrepreneurs are encouraged to consider these and other factors when conceptualizing, designing, and prototyping novel products seeking NIMHD SBIR and STTR funding. Minority health and health disparity academic researchers are encouraged to consider partnering with small businesses to assist in translating NIMHD or NIH funded research findings into potentially commercializable products for improving minority health or eliminating health disparities.

Through small business Phase I, Phase II, and Fast-track awards, NIMHD supports multi- and trans-disciplinary research and development leading to novel and or improved products capable of contributing to NIMHD’s mission. Research and development informed by the NIMHD Research Framework or other framework may proceed or be initiated at the molecular, cellular, individual, community or population level. Funding support for focus groups, phase I/II clinical trials, and other studies involving human participants needed to develop and test the proposed product may be requested. Additionally, NIMHD seeks innovative strategies for improving minority health, eliminating health disparities, and enhancing health and well-being via novel partnership between small businesses and health disparity communities where the community assists in product design activities from conception, application submission, and through completion of NIMHD funding periods and beyond. The NIMHD Research Framework acknowledges the value of small businesses partnering with community-based or -located organizations or small businesses, and with health care providers and health care-organizations. Applications developing innovative technologies or services for enhancing minority health and well-being through partnerships with community-based small businesses, such as beauty salons, barbershops, pharmacies, etc., that engage with racial and ethnic minority or health disparity populations on a regular basis, can also be requested. Technology that leverages indigenous community advisors and supporters in health promotion or prevention efforts may contribute to overall community health improvement and well-being through the processes of community empowerment and increased community cohesion.

An overarching objective of NIMHD’s investments in SBIR/STTR programs is to ensure that racial and ethnic minorities and health disparity populations benefit equally from innovations in health promotion, educational and medical curricula, prevention interventions, biotechnology, imaging technologies, technologies for computational biology and informatics, including, e.g., systems and structural biology; and technologies designed to advance personalized medicine and health, electronic health records, etc. New or improved instruments, devices, and related methodologies to facilitate biomedical or behavioral research and efforts that seek to simplify via redesign or design new instruments, devices, and methods likely to increase access, reduce costs, and improve quality of care and outcomes are of special interest.
Research Topics of Interest to NIMHD

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NIMHD Non-Clinical and Clinical Trials Topics:

Applicants are encouraged to engage in research and development that results in a product, process or service that will improve minority health and eliminate health disparities and that by design targets or involves any of the topics listed in the NIMHD waiver list or otherwise will contribute to the NIMHD mission.

For additional information about the areas of interest to the NIMHD, please visit our home page at http://www.nimhd.nih.gov/.

For additional information on research topics, contact:

Derrick C. Tabor, PhD
National Institute on Minority Health and Health Disparities, NIH
6707 Democracy Blvd.
Suite 800, MSC 5465
Bethesda, MD 20892-5465
301-402-1366, Fax: 301-480-4049
Email: derrick.tabor@nih.gov

For administrative and business management questions, contact:

Ms. Priscilla Grant, J.D., C.R.A.
Grants Management Officer
National Institute on Minority Health and Health Disparities, NIH
6707 Democracy Blvd.
Suite 800, MSC 5465
Bethesda, MD 20892-5465
301-594-8412, Fax: 301-480-4049
Email: pg38h@nih.gov
NATIONAL INSTITUTE OF NEUROLOGICAL DISORDERS AND STROKE (NINDS)

The mission of NINDS is to reduce the burden of neurological disease—a burden borne by every age group, by every segment of society, by people all over the world (https://www.ninds.nih.gov/About-NINDS/Who-We-Are/Mission). To this end, the Institute supports and conducts research on the healthy and diseased brain, spinal cord, and peripheral nerves. The NINDS SBIR/STTR (https://www.ninds.nih.gov/Funding/Small-Business-Grants) program funds small business concerns to conduct innovative neuroscience research and/or development (R/R&D) that has both the potential for commercialization and public benefit.

Limited Amount of Award and Budget Waivers

NINDS will allow awards to exceed the statutory guideline amounts by up to 50% (up to $256,580 for Phase I and $1,710,531 for Phase II, total funding support) with appropriate justification from the applicant. In addition, NINDS will allow Phase I project periods of up to 2 years and Phase II project periods of up to 3 years with appropriate justification. For budgetary, administrative, or programmatic reasons, NINDS may decrease the length of an award and/or the budget recommended by a review committee, or not fund an application.

NIH has received a waiver from SBA to exceed the hard cap for specific topics that can be found in APPENDIX A: National Institutes of Health SBA-Approved SBIR/STTR Topics for Awards over Statutory Budget Limitations and only these specific topics can apply and receive awards over the hard cap. Generally, NINDS does not fund Phase I applications greater than $700,000 total funding support, with no more than $500,000 total cost in any year or project periods greater than 2 years. In addition, the NINDS does not generally fund Phase II applications greater than $3,000,000 total funding support, with no more than $1,500,000 total cost in any year, or project periods greater than 3 years. Information about the NINDS budget guidelines can be found on the NINDS SBIR webpage: https://www.ninds.nih.gov/Funding/Small-Business-Grants/Budget-Information. Applicants considering a requested budget greater than $256,580 for Phase I or $1,710,531 for Phase II are strongly encouraged to contact program staff before submitting an application.

For all other funding opportunities, applications should follow the guidelines in the Award Budget section of those announcements carefully. Additional information can be found on the NINDS SBIR/STTR webpage at: https://www.ninds.nih.gov/Funding/Small-Business-Grants/Budget-Information.

Phase IIB Competing Renewal Awards

NINDS only accepts Phase IIB SBIR/STTR Competing Renewal applications through specific opportunities that focus on the commercialization of SBIR and STTR developed technologies. These opportunities can be found on the NINDS Funding Opportunities webpage: https://www.ninds.nih.gov/Funding/Find-Funding-Opportunities. Contact Emily Caporello at emily.caporello@nih.gov for additional information.

Research Topics of Interest to NINDS

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**NINDS Non-Clinical Trials Topics:**

**General Areas of Interest**

The NINDS accepts a broad range of small business applications that are significant, innovative, and relevant to its mission. Examples of research topics within the mission of NINDS are shown below. This list is not all inclusive and some research areas fall into multiple categories.

1. Therapeutics and Diagnostics Development for Neurological Disorders, including biomarker and diagnostic assays, therapeutics (drugs, biologics, and/or devices) for treatment of neurological disorders, and technologies/methodologies to deliver therapeutics to the nervous system.
2. Clinical and Rehabilitation Tools, including intraoperative technologies for neurosurgeons, rehabilitation devices and programs for neurological disorders, and brain monitoring systems
3. Technology and Tools, including technologies to image the nervous system, neural interfaces technologies, and tools for neuroscience research and drug development.
In addition to the research topics listed, NINDS also encourages applications in specific program areas. For additional information about NINDS funding opportunities, please visit the NINDS Funding Opportunities webpage at: https://www.ninds.nih.gov/Funding/Find-Funding-Opportunities.

NINDS Priorities

NINDS priorities are given to meritorious research proposals with the greatest potential to advance the NINDS mission (see https://www.ninds.nih.gov/About-NINDS/Who-We-Are/Mission). NINDS is especially interested in:

1. Novel and innovative technologies that are new to the SBIR or STTR programs.
2. Technologies coming to the SBIR or STTR programs for their first indication or market opportunity.
3. Companies and applicants that are new to the SBIR and STTR programs.
4. NINDS Cooperative Agreement (U44) Translational Programs. NINDS has specific translational programs that utilize the SBIR cooperative agreement mechanism (U44) as noted below. If eligible, companies are encouraged to apply through these programs.

NINDS SBIR and STTR funding decisions are based on a combination of factors:

1. potential for high impact on advancing the NINDS mission and the other programmatic priorities described in NOT-NS-18-002 (https://grants.nih.gov/grants/guide/notice-files/NOT-NS-18-002.html);
2. potential for commercialization;
3. portfolio balance (to determine whether similar projects have already been funded, search NIH Reporter http://projectreporter.nih.gov/reporter.cfm);
4. the quality of the previous performance of the applicant and/or company in the SBIR and/or STTR program, including evidence of Phase III activities;
5. for Phase II applicants, the results of the Phase I;
6. the peer review scores and critiques; and
7. availability of funds.

Translational Research

The NINDS offers a variety of specific funding opportunities and programs to accelerate the preclinical discovery and development of new therapeutic interventions for neurological disorders. These programs have specific funding opportunities and allow for budgets over the hard cap. All three programs utilize the cooperative agreement (U44) mechanism, which is milestone-driven and involves NIH program staff participation in developing the project plan, monitoring research progress, and appropriate go/no-go decision-making. SBIR applicants considering projects involving translational research are strongly encouraged to contact program staff well in advance of submission.

- **Cooperative Research to Enable and Advance Translational Enterprises for Biotechnology Products and Biologics (CREATE Bio)** is dedicated to biotechnology product- and biologics-based therapies, which broadly include modalities such as peptides, proteins, oligonucleotides, gene therapies, and cell therapies. The program supports lead optimization, IND-enabling studies for the candidate, and early-phase clinical trials. Contact: Ann-Marie Broome (ann-marie.broome@nih.gov)

- **Translational Neural Devices Program** provides support for projects that focus on pre-clinical and pilot clinical studies for therapeutic devices. Activities supported in this program include implementation of clinical prototype devices, preclinical safety and efficacy testing, design verification and validation activities, pursuit of regulatory approval for the clinical study, and a clinical study. Contact: Nick Langhals (nick.langhals@nih.gov)
- **Blueprint Neurotherapeutics Network (BPN)** provides both funding and non-dilutive support for small molecule drug discovery and development, from hit-to-lead chemistry through phase I clinical testing. The program offers funding, access to NIH-funded contract research organizations (CROs), and access to consultants with expertise in various aspects of drug discovery and development. Contact: Charles Cywin (cywincl@mail.nih.gov)

- **NIH Countermeasures Against Chemical Threats (CounterACT)** supports research and development on new and improved therapeutics to prevent or mitigate the toxic effects from exposure to chemical threats, defined as toxic chemical agents that could be used in a terrorist attack against civilians, or those that could be released at toxic levels by accident or natural disaster. NINDS supports partnerships between small business and not-for-profit laboratories engaged in research related to the CounterACT program that falls within the NINDS mission, including devices that could be used to reduce morbidity and mortality during a chemical emergency involving mass casualties, as well as some research on therapeutics. Contact: David Jett (david.jett@nih.gov)

- **Neuroscience Biomarker Program** is focused on improving the quality and efficiency of neurotherapeutic clinical research toward Phase II and beyond by supporting rigorous biomarker validation. To achieve this goal, the program will: 1) promote rigorous biomarker identification and validation through milestone-driven funding opportunities, 2) centralize information about existing NINDS and NIH biomarker sample and data repository resources and 3) facilitate the development of future resources focused on bridging the gaps in the biomarker development pipeline. Contact: Mary Ann Pelleymounter (mary.pelleymounter@nih.gov).

Information about these and other programs can be found at [https://www.ninds.nih.gov/Current-Research/Research-Funded-NINDS/Translational-Research](https://www.ninds.nih.gov/Current-Research/Research-Funded-NINDS/Translational-Research).

**Brain Research through Advancing Innovative Neurotechnologies (BRAIN) Initiative**

The Brain Research through Advancing Innovative Neurotechnologies (BRAIN) Initiative is a Presidential project aimed at revolutionizing our understanding of the human brain. NIH is one of several federal agencies involved in the BRAIN Initiative. Planning for the NIH component of the BRAIN Initiative is guided by the long-term scientific plan, “BRAIN 2025: A Scientific Vision,” which details seven high-priority research areas. This report can be found at [https://braininitiative.nih.gov/](https://braininitiative.nih.gov/).

NIH has several specific funding opportunity announcements through the BRAIN Initiative that are targeted to small business concerns. These funding opportunities can be found at [https://www.braininitiative.nih.gov/funding/](https://www.braininitiative.nih.gov/funding/). Applicants are encouraged to consider if these funding opportunities may be appropriate to their research. Contact Emily Caporello at emily.caporello@nih.gov for additional information.

**Helping to End Addiction Long-term (HEAL) Initiative**

The Helping to End Addiction Long-term (HEAL) Initiative is an aggressive, trans-agency effort to speed scientific solutions to stem the national opioid public health crisis. Further information on the HEAL Initiative can be found at [https://heal.nih.gov/](https://heal.nih.gov/).

NIH has several specific funding opportunity announcements through the HEAL Initiative that are targeted to small business concerns. These funding opportunities can be found at [https://heal.nih.gov/funding](https://heal.nih.gov/funding). Applicants are encouraged to consider if these funding opportunities may be appropriate to their research. Contact Emily Caporello at emily.caporello@nih.gov for additional information.
NINDS Clinical Trials Topics:

NINDS accepts clinical trials in any of the “General Areas of Interest” topics listed above in the “NINDS Non-Clinical Trials Topics” section.

NINDS is committed to identifying effective treatments for neurological disorders by supporting well-executed clinical trials. NINDS accepts and supports SBIR and STTR clinical trial applications within the NINDS mission through specific opportunities, which can be found on the NINDS SBIR webpage: [https://www.ninds.nih.gov/Funding/Small-Business-Grants/Areas-Interest#CT](https://www.ninds.nih.gov/Funding/Small-Business-Grants/Areas-Interest#CT). Other human subjects research can be submitted through the SBIR and STTR Parent (Clinical Trials Not Allowed) solicitation. However, NINDS may decline funding of any application that includes human subjects research for programmatic or administrative reasons. SBIR applicants considering projects involving human subjects research are strongly encouraged to contact program staff in advance of submission.

For additional information on the NINDS SBIR/STTR program, contact:
Emily Caporello, Ph.D.
Scientific Project Manager
301-496-1779
Email: emily.caporello@nih.gov

For financial and grants management questions, contact:
Chief Grants Management Officer
National Institute of Neurological Disorders and Stroke (NINDS)
Email: ChiefGrantsManagementOfficer@ninds.nih.gov
NATIONAL INSTITUTE OF NURSING RESEARCH (NINR)

The National Institute of Nursing Research (NINR) supports research focused on biological and behavioral aspects of critical health problems that confront the Nation. Rapid advances in technology and genomic science, as well as significant changes in demographics and health care policies and practice, have inspired the field of nursing science to identify innovative approaches and interventions that improve health outcomes. Emphasis is on seeking ways to reduce the burden of illness and disability by understanding and easing the effects of acute and chronic illness and associated symptoms, improving health-related quality of life by preventing or delaying the onset of disease or slowing its progression, establishing better approaches to promote health and prevent disease, and improving clinical environments by testing interventions that influence patient health outcomes and reduce costs and demand for care.

For additional information about areas of interest to the NINR, please visit our home page at http://www.ninr.nih.gov/, and also at http://www.ninr.nih.gov/researchandfunding.

NINR does not accept Phase IIB applications.

Research Topics of Interest to NINR

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General Topic Areas
NINR is interested in the development of technologies, products or services related to:
1. Self-management of acute and chronic conditions to improve quality of life,
2. Symptom management and personalized health strategies,
3. Promotion of wellness and disease/disability prevention, and
4. Tools that enhance end-of-life care and palliative care across the lifespan.

Priority Topics
Digital health tools offer significant new opportunities to improve medical outcomes and enhance the efficiency of care via the convergence of devices, data interoperability and/or artificial intelligence. Individuals can take advantage of these tools to track and better manage their own health while healthcare providers can use them to improve their ability to accurately diagnose, effectively treat disease and enhance patient treatment plans. Accordingly, NINR is interested in the translational development and implementation of a broad array of digital health technologies, products or services. The list of priority topics below is not all inclusive and some research areas fall into multiple categories.

NINR Non-Clinical Trials Topics:
A. Digital Health Technology, including but not limited to:
   - **Wearable or remote sensing devices** such as the development of technologies that passively monitor biology, behavior and context of use to maintain/promote health and/or minimize the physical and psychological burdens on caregivers while caring for individuals with acute or chronic conditions
   - **Point-of-care (POC) diagnostic/intervention devices**, especially as they relate to pain, sleep and the recovery from infection
   - **mHealth technologies** such as the development of evidence-based apps for mobile and wireless devices (smartphones, tablets, etc.), especially for use in combination with wearable or remote health-related sensors
• **Smart speaker / smart home technologies** for remote monitoring, telehealth, self-management and patient assistance, coordinated family/provider care, communication, end-of-life palliative care, etc.

• **Telehealth / telemedicine** such as the development of technologies to facilitate the diagnosis, consultation, education/communication, clinical care and self-management of an individual’s health and wellness, etc.

• **Health information/integration technologies (hIT)** such as the development of an integrative technology that leverages ‘big data’ from wearables, mHealth, EHR, biotypes, biomarkers, and/or social determinants of health, etc. to holistically inform personalized health strategies, shared decision making and reduced health disparities – especially technologies that use the FHIR® standard ([NOT-OD-19-122](#))

• **Artificial Intelligence (AI)-based tools** such as ML/DL-based algorithms/software within a telehealth platform that leverages hIT ‘big data’ to predict disease risk/progression and improve clinical decision making, especially related to end-of-life and palliative care

B. **Medical Robot Systems** including, but not limited to:

• **Rehabilitative robots** such as systems that enhance motor learning and functional recovery after injury or illness

• **Physically-assistive robots** such as systems that assist individuals with activities of daily living

• **Psychosocially-assistive robots** such as interactive systems to enhance mood, mitigate the effects of loneliness, and enhance social connection and communication

• **Telepresence robots** such as systems that allow for access to specialty services in a rural health clinic

C. **Development of medical devices**, including AI/ML-based software as a medical device

D. **Interventions for decision support** such as the development and implementation of interventions that enhance communication between individuals, caregivers and health care providers or tools that promote caregiver support

**NINR Clinical Trials Topics:**

NINR accepts clinical trials in any of the topic areas above.

For additional information please contact:

Dr. Kristopher Bough  
Director, Small Business Innovation Research Program  
National Institute of Nursing Research (NINR)  
Office of Extramural Programs (OEP)  
National Institutes of Health (NIH)  
6701 Democracy Blvd, Room 727  
Bethesda, MD 20892-4870  
Office: (301) 496-2604  
Email: kristopher.bough@nih.gov
NCATS is transforming translational science to improve human health; it relies on the power of data, new technologies and teamwork to develop, demonstrate and disseminate innovations that reduce, remove or bypass costly and time-consuming bottlenecks in translational research.

NCATS small business funding is designed specifically to transform the translational science process so that new treatments and cures for diseases can be delivered to patients more quickly. The Center supports the development of technologies, assays, drugs, devices, instruments, and methodologies that may have broad application to any stage of the translational process from preclinical development to clinical research and to implementation science in patient care and public health. For additional information, please visit http://www.ncats.nih.gov.

NCATS is committed to supporting small business Phase I, Phase II, Fast-track and Phase IIB Competing Renewal awards through the Small Business Innovation Research (SBIR) and Small Business Technology Transfer Programs (STTR). For additional information, please visit http://ncats.nih.gov/smallbusiness.

Limited Amount of Award

For budgetary, administrative or programmatic reasons, NCATS may decide not to fund an application or may decrease the length of an award and/or the budget recommended by a review committee. Generally, NCATS will not fund:

- Phase I applications greater than $256,580 total costs or project periods greater than 2 years
- Phase II applications greater than $1,710,531 total costs or project periods greater than 3 years

For certain topical areas (Appendix A), the Small Business Administration has approved an NIH SBIR/STTR Topic Waiver list for which the NCATS generally will not fund:

- Phase I applications greater than $325,000 total costs or project periods greater than 2 years
- Phase II applications greater than $2,000,000 total costs or project periods greater than 3 years

Applicants considering a requested budget greater than these limits are strongly encouraged to contact program staff before submitting an application.

Phase IIB Competing Renewal Awards

Occasionally, NCATS may accept Phase IIB SBIR Competing Renewal grant applications of NCATS supported Phase II awards to continue research and development of products that have a potential to address bottlenecks in the translational process, and where additional time and effort is needed to reach a stage where interest and investment by third parties would be likely. Such products are expected to have broad applicability and be consistent with the mission of NCATS. Applicants are strongly encouraged to speak to NCATS Program staff prior to submitting their Phase IIB application. Budgets for Phase IIB grant applications must be approved by NCATS Program staff prior to submission.

Research Topics of Interest to NCATS

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<td>Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply</td>
<td>X</td>
<td>U01 – however the SBC can only participate if repurposing an existing drug or biologic (therapeutics) that have already completed at least a Phase I trial for a different indication by the time an award is made. These pharma drugs and biologics are listed in PAR-18-332. <a href="https://grants.nih.gov/grants/guide/pa-files/PAR-18-332.html">https://grants.nih.gov/grants/guide/pa-files/PAR-18-332.html</a></td>
<td></td>
</tr>
</tbody>
</table>

**NCATS Non-Clinical Trials Topics:**

Preclinical Drug Discovery and Development
- Innovative platforms for identification and prioritization of targets for therapeutic intervention with clear clinical impact; such as those that are: implicated for disease, have genetic variations that have been identified in functional regions of receptor targets, and/or have high potential for biased signaling that would promote the beneficial effects of receptor signaling and reduce the unwanted effects
- Tools and technologies to enable high throughput screening of compound activity on currently “non-druggable” targets
- Assays for high-throughput screening of rare-diseases-related targets
- Co-crystallization high-throughput screening techniques
- Fluorescence probes to replace antibodies for determination of cellular protein translocation
- Phenotypic assay development, including stem cell technology platforms for human “disease-in-a-dish” applications and the evaluation of toxicity
- Interventions that target molecular pathways or mechanisms common to multiple diseases
- Platforms for non-antibody biologics, cell-based therapies and gene therapy discovery
- Small molecule and biologics analytical characterization
- Accelerated bioengineering approaches to the development and clinical application of biomedical materials, devices, therapeutics and/or diagnostics
- Development of novel technologies for enzyme replacement therapies (e.g., new cell culture/expression system) to solve major bottlenecks in rare diseases research
- Innovative methods to determine alternative uses for existing therapeutic interventions for high priority areas, such as rare diseases and pain.
- Tools and technologies that increase the predictivity or efficiency of medicinal chemistry, biologic or other intervention optimization
- Technologies to deliver nucleic acid therapeutics to tissues other than the liver
- Methodologies and technologies to increase efficiencies of manufacturing therapeutics
- Development of novel high-throughput technologies that focus on making translational research more efficient
- GMP production of exosome/extracellular vesicles
- Generation of producer lines for large scale production of exosomes/extracellular vesicles
- Extracellular RNA-based biomarkers and therapeutics of human diseases
- Approaches to targeting the human microbiome for therapeutic or diagnostic purposes
- Scale up, manufacturing and characterization of IPS cells
- 3D printing technologies
- Technologies to substantially improve the efficiency and reduce the cost of clinical grade gene therapy vector manufacturing
- Development of in vitro human tissue models (organs, 3D printing)
- Technologies to allow therapeutic proteins other than lysosomal enzymes to be secreted and taken up by other cells via cross-correction
- Novel strategies to prevent deleterious immune responses to gene therapy, genome editing and/or enzyme replacement therapy
- Establishing more robust phenotypic screens that may help prioritize candidate compounds for further testing
- Innovative technology for non-small molecule delivery
- High-throughput epigenetics screening/characterization tools and technologies
- Microphysiological systems (MPS)/Tissue Chips, including MPS/Tissue Chips that incorporate known functional variants, e.g. ACMG 59 or CPIC A alleles, for study comparison using the same derived genetic background across a set of tissue chips with the functional variant
• Searchable access to information about research resources, facilities, methods, cells, genetic tests, molecules, biologic reagents, animals, assays, and/or technologies with evidence about their use in research studies
• Cloud-based tools and methods for meaningful sharing, re-use and integration of research data
• Novel platforms, technologies and tools for: (1) enabling clinical and translational research, particularly those with mechanisms for inclusion of patient-reported data and (2) integration of patient data collected from multiple devices and multiple/diverse clinical studies
• Development of personalized phenotypic profiling (as well as personalized intervention) based on patient-centered integration of data from multiple data sources, including social media
• Development of predictive models for translational science
• Digital applications and tools (including telemedicine platforms) that facilitate/enhance translational research and medicine in rural populations
• Generic Disease Registry template platforms that can be reused for multiple diseases.
• Mobile device validation tools to ensure data from different brands or versions have compatible results.
• Tools to assess with algorithms developed with artificial intelligence, machine learning.
• Tools that allow for persistent identifier and attribution for data contributors that give credit to the data producers while ensuring that shared data has not been altered
• Patient Mobile Tool Platforms that facilitate tool developers to build "apps" that integrate into their medical records.
• Tools and environments that enable an easy interrogation of publicly available data

Clinical, Dissemination and Implementation Research

• Tools and technologies that increase the efficiency of human subjects research, that facilitate the rapid diagnosis and/or clinical trial recruitment and subject tracking, institutional review board evaluation, and/or regulatory processes
• Increased efficiency of clinical research conduct, including but not limited to regulatory decision support, patient eligibility analysis and recruitment and retention tracking
• Tools, technologies, and other strategies to evaluate and improve the process of informed consent
• Educational tools for clinical and translational science
• Computational or Web-based health research methods, including:
  o Platforms for generally applicable and scalable multi-disease registries and natural history studies
  o Clinical trial designs and analyses (e.g., for pragmatic clinical trials)
• Approaches, tools, platforms and environments to integrate data in novel ways for development of new biomarkers that can be tested in translational research paradigms for which there are barriers or bottlenecks
• Strategies to enhance the quality and accelerate the conduct of dissemination and implementation research
• Sustainable solutions for effective tools and environments in translational research
• Development and validation of patient reported outcomes, clinician reported outcomes, and biomarkers for rare diseases that are not already supported by a disease-specific NIH ICs
• Tools, technologies and other strategies that address medication adherence in clinical settings
• Tools, technologies and other strategies that address and improve community engagement
• Tools and technologies that address the rapid diagnosis, clinical management of rare diseases
• Patient empowerment tools/apps that allow users to compare their treatment, outcomes to normative populations existing treatment guidelines
• Telemedicine or digital health applications that focus on research in rural populations
• Tools and technologies that help characterize human disease states and assist in assessing the impact of interventions

**NCATS Clinical Trials Topics:**

NCATS will not accept SBIR and STTR applications that propose clinical trials, and all of the topics listed must be for projects that do not propose clinical trials.

For additional information on research topics, please contact:

Lili M. Portilla, MPA
NCATS SBIR/STTR Program Director
National Center for Advancing Translational Sciences, NIH
Phone: 301-827-7170
Email: [NCATS-SBIRSTTR@mail.nih.gov](mailto:NCATS-SBIRSTTR@mail.nih.gov)

For Administrative, business management and grants policy questions, please contact:

Ms. Betsy Snell
Grants Management Specialist, SBIR/STTR Project Liaison
Phone: 301-827-8454, Fax: 301-480-3777
Email: [betsy.snell@nih.gov](mailto:betsy.snell@nih.gov)
NATIONAL CENTER FOR COMPLEMENTARY AND INTEGRATIVE HEALTH (NCCIH)

The mission of NCCIH is to define, through rigorous scientific investigation, the usefulness and safety of complementary and integrative health interventions and their roles in improving health and health care.

The following narrative indicates the scope of projects suitable for the SBIR/STTR program that fit within the mission of NCCIH. For additional information about areas of interest to NCCIH and a listing of NCCIH’s currently funded applications, please visit http://www.nccih.nih.gov/research. Business concerns interested in exploring SBIR/STTR grant opportunities with NCCIH are encouraged to contact NCCIH program directors prior to submitting an application.

Limited Amount of Award

Generally, NCCIH will not fund:

- Phase I applications greater than $256,580 total costs for the duration of the project or project periods greater than 2 years
- Phase II applications greater than $1,710,531 total costs for the duration of the project or project periods greater than 3 years.

These total award amounts will be adjusted according to NIH guidelines.

The Small Business Administration (SBA) has approved an NIH SBIR/STTR Topic Waiver list for “National Institutes of Health SBA-Approved SBIR/STTR Topics for Awards Over Statutory Budget Limitations,” for which NCCIH generally will not fund:

- Phase I applications greater than $325,000 total costs per year or project periods greater than 2 years.
- Phase II applications greater than $2,000,000 total costs for the duration of the project or project periods greater than 3 years.

Applicants considering a requested budget greater than these limits are strongly encouraged to contact program staff before submitting an application.

Research Topics of Interest to NCCIH

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,
See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

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| Clinical Trials SBIR/STTR Omnibus/Parent Funding Opportunity Announcement/s | X | | For applications involving clinical studies that fall within the NIH definition of a clinical trial, NCCIH will not support clinical trials aiming to test efficacy/effectiveness (meaning the study is powered on a primary outcome that is a clinical assessment used in clinical diagnosis of disease or monitoring of disease severity) of an intervention as a part of an SBIR/STTR Phase I application. Applicants seeking to conduct efficacy or effectiveness clinical trials should pursue funding via other Funding Opportunity Announcements (FOAs) such as the Omnibus SBIR/STTR Phase II and Fast-Track.

NCCIH recognizes a difference between "clinical trials" that are designed to answer specific questions about the clinical effect of interventions and mechanistic studies that have the primary goal of understanding how an intervention works.

A clinical outcome study has the objective of determining the clinical safety, tolerability, feasibility, efficacy, and/or effectiveness of pharmacologic, nonpharmacologic, behavioral, biologic, surgical, or device (invasive or noninvasive) interventions.

A mechanistic study has the objective to understand the mechanism(s) of action of an intervention, a biological or behavioral process, or the pathophysiology of a disease/condition.

NCCIH continues to accept clinical trials of all types on Omnibus SBIR/STTR Phase II and Fast-Track applications.

See NOT-AT-19-012 for “NCCIH Policy for SBIR and STTR Phase I Applications Proposing Clinical Trials to the Omnibus Solicitations”
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| Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply | X |  | Mechanisms Underlying the Contribution of Sleep Disturbances to Pain (R01 Clinical Trial Optional)  
Mechanisms Underlying the Contribution of Sleep Disturbances to Pain (R21 Clinical Trial Optional)  
Feasibility Clinical Trials of Mind and Body Interventions for NCCIH High Priority Research Topics (R34 Clinical Trial Required)  
Center of Excellence for Research on Complementary and Integrative Health (P01 Clinical Trial Optional)  
Clinical Coordinating Center for NCCIH Multi-Site Investigator-Initiated Clinical Trials of Mind and Body Interventions (Collaborative UG3/UH3 Clinical Trial Required)  
Fundamental Science Research on Mind and Body Approaches (R21 Clinical Trial Optional)  
Investigator Initiated Clinical Trials of Complementary and Integrative Interventions Delivered Remotely or via mHealth (R01 Clinical Trial Required)  
Exploring the Mechanisms Underlying Modulation of Glymphatic-Lymphatic Systems by Complementary and Integrative Health Approaches (R01 Clinical Trial Optional)  
Exploring the Mechanisms Underlying Modulation of Glymphatic-Lymphatic Systems by Complementary and Integrative Health Approaches (R21 Clinical Trial Optional)  
Notice of Special Interest: Exploring the Mechanisms Underlying Analgesic Properties of Minor Cannabinoids and Terpenes |
NCCIH Non-Clinical Trials Topics:

**Natural Products** (including botanicals, herbs, probiotics, prebiotics, dietary supplements, special medicinal diets, and microbiome-/microbial-based therapeutics):

- Development and validation of technologies for standardization and characterization of biologically active ingredients in natural products
- Development and validation of technologies for taxonomic identification of botanical raw materials or detection of adulterants
- Development and validation of technologies for the identification and characterization of bioactive metabolites derived from oral consumption of natural products
- Development and validation of methods for the sustainable production of low-yield natural products of commercial interest
- Development of novel analytical tools and technologies to study the microbiome, including its composition, genetics, and bioactivity, that can help clarify associations between the human microbiome and brain function and health
- Development of gut microbiome monitoring assays for validating safety and functional analysis of genomic and microbiota interactions
- Development of complementary and integrative therapeutic approaches to modify and balance the gut microbiota in healthy populations and individuals with disrupted microbiota and related diseases

**Mind and Body Approaches** (including meditation, mindfulness, hypnosis, yoga, tai chi, acupuncture, manual therapies, and music/art therapies):

- Development, testing, and validation of appropriate objective and/or quantitative measures and instruments to assess or monitor mind and body approaches in different contexts (e.g., classrooms, families, child welfare, juvenile justice systems)
- Development, testing, and validation of measures and tools to assess training or fidelity of implementation of mind and body approaches in different settings (e.g., health care, community, families, schools, child welfare, juvenile justice systems)
- Development and testing of technologies for the implementation of mind and body approaches in group or individual settings and/or self-care strategies. Examples may include but are not limited to the use of mobile health technologies such as smartphone apps, sensors, online delivery, or phone-based delivery
- Development and validation of methods for standardization and characterization of the active components of mind and body approaches
- Development and validation of methods for standardization of multimodal interventions to study whole person health
- Development and validation of imaging tools or instruments for studying manual therapies, including but not limited to massage, acupuncture, or spinal manipulation
- Development and testing of innovative technologies for multisensory delivery of mind and body approaches
- Development, testing, and validation of innovative technologies to enhance light- or olfaction-based therapies

**General Tool/Technology Development:**

- Development and validation of biomarkers that correlate with efficacy of complementary and integrative health approaches
- Development and validation of standardized, reliable, and cost-effective tools that correlate with brain imaging in response to mind and body interventions
• Development and validation of tools, technologies, and instruments, including gaming and virtual reality technologies, for the accurate assessment of adherence and/or fidelity to the use of mind and body practices and interventions
• Development and validation of tools to improve patient-reported outcome measures of importance in clinical studies of complementary and integrative health approaches
• Development, pilot testing, and validation of wireless technologies for real-time data collection and monitoring of brain activity or other physiological signals for mind and body approaches
• Development or adaptation of biochemical or epigenetic monitoring devices for complementary and integrative health approaches
• Development and validation of tools to improve biological and physiological outcome measures for use in clinical studies of complementary or integrative health approaches
• Development or adaptation of technologies for objective assessment of pain with relevance to complementary and integrative health approaches
• Development of sleep monitoring technologies or biomarker panels to assess sleep deprivation, sleep deficiency, circadian rhythm dysregulation, and connection of sleep disturbances with health risks
• Development and testing of in vivo labeling technology of tissues or cells responsible for generating signals in response to different internal senses (e.g., mechanical force, temperature, osmolarity, oxygen levels)
• Development and testing of technology or methods quantifying biomechanical forces applied to internal tissues or cells
• Development and testing of mobile health technology or nonmobile technology and methods to monitor or quantify physical and/or emotional well-being, breathing, or sleep

**NCCIH Clinical Trials Topics:**

**Natural Products** (including botanicals, herbs, probiotics, prebiotics, dietary supplements, or special medicinal diets):

• Development and validation of technologies for standardization and characterization of biologically active ingredients in natural products
• Clinical testing of natural products for the management of hard-to-treat symptoms such as pain, sleep disorders, or mild-to-moderate anxiety and depression to allow development of an evidence base that would accelerate FDA approval of a drug indication for the natural product
• Development and validation of technologies for the identification and characterization of bioactive metabolites derived from oral consumption of natural products
• Development of novel analytical tools and technologies to study the microbiome, including its composition, genetics, and bioactivity, that can help clarify associations between the human microbiome and brain function and health
• Development of gut microbiome monitoring assays for validating safety and functional analysis of genomic and microbiota interactions

**Mind and Body Approaches** (including meditation, mindfulness, hypnosis, yoga, tai chi, acupuncture, manual therapies, and music/art therapies):

• Development, testing, and validation of appropriate objective and/or quantitative measures and instruments to assess or monitor mind and body approaches
• Development, testing, and validation of measures and tools to assess training or fidelity of implementation of mind and body approaches in different settings (e.g., health care, community, families, schools, child welfare, juvenile justice systems)
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For additional information on research topics, please contact:

**Merav Sabri, PhD**
Program Director, Division of Extramural Research
National Center for Complementary and Integrative Health (NCCIH)
6707 Democracy Boulevard, Suite 401
Telephone: 301-496-2583
Email: merav.sabri@nih.gov

**Anastasia Solis**
Program Analyst, Division of Extramural Research
6707 Democracy Boulevard, Suite 401
Bethesda, MD 20892-5475
Telephone: 301-594-8018
Email: anastasia.solis@nih.gov
NATIONAL LIBRARY OF MEDICINE (NLM)

The National Library of Medicine (NLM) offers support for research and development projects in biomedical informatics and data science. Biomedical informatics and data science research applies computer and information sciences to improve the access, storage, retrieval, management, dissemination and use of biomedical information. Grants are made to U.S. small businesses that seek to undertake informatics research and development leading to commercialization. Critical research areas include: representation of medical knowledge in computers; organization and retrieval issues for image databases; enhancement of human intellectual capacities through virtual reality, dynamic modeling, artificial intelligence, and machine learning; medical decision-making; linguistic analyses of medical languages and nomenclatures; investigations of topics relevant to health information or library science; biotechnology informatics issues; and informatics for disaster management. For additional information about areas of interest to NLM and a listing of NLM funded applications, please visit http://www.nlm.nih.gov/ep. Business concerns interested in exploring SBIR/STTR grant opportunities with NLM are encouraged to contact the NLM representatives prior to submitting an application.

Research Topics of Interest to NLM

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

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---|---|---|---
Clinical Trials SBIR/STTR IC/Office/Agency - Specific Funding Opportunity Announcement/s | X | |  
Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply | X | |  

NLM’s SBIR/STTR grant programs are focused on areas of particular interest from small business. The following narrative indicates the scope of projects suitable for the SBIR/STTR program that fit within the mission of NLM. They are not listed in priority order.

**NLM Non-Clinical Trials Topics:**
- Visualization approaches to complex biomedical data at multiple levels
- Approaches to help consumers and patients visualize their health data in the context of social and behavioral determinants
- Tools for modeling the impact of behavior and social determinants of health on physiological systems
- Tools and approaches for data integration of large disparate data resources
- Technological approaches to protecting confidentiality while storing, sharing and using biomedical data
- AI techniques for annotating, curating and managing biomedical data resources
- Self-managing, self-documenting datasets
- Tools to enhance security of biomedical data, including personal health data, for multi-party use (meaning, two or more users using the data at once)
- New Technologies that facilitate real-time decisions in clinical practice and public health, using personal health data in electronic records or clinical trials
- Use of formal and informal data sources to track disease outbreaks, pandemics
- Tools for understanding and predicting climate and environmental effects on human health
- AI techniques for characterizing and minimizing the impact of errors or incompleteness in a health data set

**NLM Clinical Trials Topics:**
NLM will not accept SBIR applications that propose clinical trials, and all of the topics listed must be for projects that do not propose clinical trials.
For additional information on research topics, contact:

Dr. Jane Ye  
Program Officer  
Division of Extramural Programs  
National Library of Medicine  
301-594-4882, Fax: 301-402-2952  
Email: yej@mail.nih.gov

For administrative and business management questions, contact:

Ms. Samantha Tempchin  
Grants Management Officer  
Extramural Programs Division  
National Library of Medicine  
301-496-4222, Fax: 301-402-0421  
Email: samantha.tempchin@nih.gov
DIVISION OF PROGRAM COORDINATION, PLANNING, AND STRATEGIC INITIATIVES (DPCPSI), OFFICE OF RESEARCH INFRASTRUCTURE PROGRAMS (ORIP)

ORIP supports high-quality, disease-free animal models and specialized animal research facilities to help meet the needs of biomedical researchers to understand, detect, treat, and prevent a wide range of human diseases. This support enables discoveries at molecular, cellular, and organ levels that lead to animal-based studies which then are translated to patient-oriented clinical research, aiming to find treatments to cure or ameliorate common and rare diseases. Through the small business Phase I, Phase II, Fast-track, and Phase IIB Competing Renewal awards, ORIP is especially interested in funding research to develop preclinical methods and technologies that improve animal models of human diseases, and the care, use, and management of laboratory animals. ORIP also encourage the development and implementation of technologies to directly benefit the welfare of research animals and to directly improve animal facilities that support biomedical and behavioral research.

A list of some potential ORIP program topics follows the description of our Phase IIB Competing Renewal Awards. For additional information about ORIP, please visit our home page at https://orip.nih.gov/.

Phase IIB Competing Renewal Awards

ORIP will only accept Phase IIB SBIR Competing Renewal grant applications of ORIP-supported Phase II awards to continue research and development of methods, technologies, tools and devices for basic or translational research where extraordinary time and effort is needed for completion of these projects. The Phase IIB Competing Renewal award is intended to allow small businesses the opportunity to reach a stage where interest and investment by third parties would be more likely. Such products are expected to have broad applicability, consistent with the mission of ORIP. Budgets that do not exceed $1 M per year in total costs (for up to 2 years), may be requested for this Phase IIB Competing Renewal opportunity; however it is expected that, in most cases, the requested budget would not exceed the final year budget of the applicant’s previous Phase II award. This opportunity is available for the SBIR program only.

Please contact your Program Officer before beginning the process of preparing a Phase IIB Competing Renewal application. In addition, prospective applicants are strongly encouraged to submit to the Program Contact (listed after each section), a letter of intent that includes the following information:

- Descriptive title of the proposed research
- Name, address, and telephone number of the Principal Investigator
- Names of other Key Personnel
- Participating organizations
- Funding Opportunity Announcement Number (e.g., PA-18-XXX)

A letter of intent is not required, is not binding, and does not enter into the review of a subsequent application. It is expected that only a few of ORIP SBIR Phase II awards will be eligible for a Phase IIB Competing Renewal grant.

Research Topics of Interest to ORIP

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NIH, CDC, and FDA Program Descriptions and Research Topics  DPCPSI/ORIP 171
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**ORIP Non-Clinical Trials Topics:**

**Division of Comparative Medicine**

A. Development of improved reagents and cost-effective methods and technologies to accurately screen and diagnose selected diseases of laboratory animal, and to perform overall assessments of animal quality and health status. An urgent need currently exists for the development of improved methods for detection of active tuberculosis in nonhuman primates.

B. Development of technology to identify molecular phenotype(s) of a single stem cell or induced pluripotent stem cell from laboratory animals.

C. Development of improved reagents, techniques, and equipment for genomic and transcriptomic analysis and data mining from tissue or cells of laboratory animals and animal models of human diseases.
D. Development of new technologies for rapid characterization and deep phenotyping of large numbers of animals.

E. Development of technologies to identify biomarkers for clinical diagnostics in well validated disease models.

F. Development of vaccines and new therapeutic agents to prevent and/or control selected laboratory animal diseases. One high priority need is to develop methods to control and prevent Herpes B virus in nonhuman primates.

G. Identification, development, and characterization of spontaneous and engineered vertebrate animal models for studies of various human diseases, excluding most random mutagenesis projects performed on rodents.

H. Development of technologies and robust tools for the effective preservation of biomedical models.

I. Development and refinement of high throughput technologies and devices for the cryopreservation, and long-term maintenance and revival of cells, tissues, and laboratory animal embryos, and gametes, especially for Drosophila, zebrafish and other aquatic stocks.

J. Development of technologies and devices for the effective monitoring of frozen and cryopreserved cells, biological materials/tissues and laboratory animal embryos, and gametes.

K. Development of technologies for improved embryo transfer within a single animal species or of intraspecific embryo transfer to allow preservation of rare, or unique, animal species that may have unique value in biomedical research as animal models for human disease.

L. Development of improved reagents, artificial intelligence/machine learning technologies, devices, and high throughput technology to perform, analyze, capture and process data gathered in "omics" studies (genomics, transcriptomics, phenomics, proteomics, glycomics, epigenomics, metabolomics, among others) in normal, disease and intervention conditions in animal/biological models to support/validate pre-clinical analyses.

M. Development of biological tools and reagents for reconstruction, remodeling, repair and regeneration of tissues damaged by injury or disease. Development of the technologies and procedures to test efficacy and safety of these experiments in animal models. Approaches to detect and track the implanted cells and tissues in vivo.

N. Development of in vitro animal cell culture techniques and computational (in silico) methods to reduce the number of animals used in studies and replace certain tests conducted in animal models with new complementary methods.

O. Development of acellular biomaterials, biosensors and reagents to promote, detect and track reconstruction, remodeling, repair and regeneration of tissues and organs damaged by injury or disease.

P. Development of reagents (including antibodies), assays, and technologies that will facilitate research using aquatic biomedical models such as zebrafish or Xenopus for understanding basic aspects of development, physiology, or genetics.

Q. Development of reagents (including antibodies), assays, and technologies that will facilitate research using nonhuman primates (NHPs) for understanding basic aspects of development, physiology, or genetics. A high priority need are reagents for other NHP species besides rhesus macaque.

R. Development of rapid and sensitive technology for the accurate detection and diagnosis of polymicrobial infections in biomedical laboratory animal models, including those agents involved in vertical transmission of diseases into embryos and larvae.

S. Technologies for improved sex determination of embryos, neonatal, and juvenile stages of animals, with one high priority need being nonmammalian species.
T. Development of rapid and sensitive technology for the detection of emerging human pathogens in animal models.

U. Development of non-invasive, micro sensing technology (embedded devices, microchips) for the characterization of a wider range of NHP and other live animal models for collecting data such as neuroimaging, behavioral and cognitive assessments, microbiome and metabolism. Of special interest are sensitive and selective probes/sensors for detecting physiological fluctuations in living animals, able to monitor at deep tissues level.

V. Development of technologies for cell-based therapies that could be used as implantable biocomputers in animal models of human disease, to perform complex logic computations that integrated signals from multiple metabolites. This include remote-controlled switches and natural, nontoxic, highly soluble, and potentially beneficial to health trigger molecules.

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Office of the Director
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Division of Construction and Instruments

The Division of Construction and Instruments supports the development and implementation of technologies to directly benefit the welfare of research animals and to directly improve animal facilities that support biomedical and behavioral research. In particular, the areas being supported include research on tools and equipment, their use to improve and ease care, and to facilitate monitoring of healthy animals. Another area of interest encompasses research to improve laboratory equipment to maintain the environmental conditions and to upkeep the infrastructure of animal facilities. Of special importance is the employment of green technologies. Examples of topics of special interest include (but are not limited to) research leading to the development of better, more reliable, and more efficient:

A. Equipment such as vacuum cleaners, air filters, hoods, snorkels, autoclaves for animal research facilities, for barrier facilities, and other facilities with special needs and requirements;

B. Equipment to distribute water and food, and monitor their intake by research animals;

C. Equipment to increase the quality of life and prevent injuries of research animals and research staff and investigators;

D. Equipment to monitor and protect the well-being of animals, including IT-supported tools;

E. Equipment and its use for maintenance of disease-free colonies and healthy animals;

F. Equipment to disinfect devices, furnishings, and other apparatus in animal facilities such as aquaria, cages, tunnels, and racks;

G. Cost-effective husbandry and colony management techniques, equipment, and/or new approaches to improve laboratory animal welfare and assure efficient and appropriate research use;

H. Specialized equipment and caging for laboratory animals to permit optimal environmental control and operational efficiency, including improvements in caging, identification/tagging of animals and remote monitoring in animal facilities.
ORIP Clinical Trials Topics:

ORIP will not accept SBIR applications that propose clinical trials, and all of the topics listed must be for projects that do not propose clinical trials.

For additional information on research topics, contact:

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For administrative and business management questions, contact:

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Grants Management Specialist,
National Heart, Lung, and Blood Institute (NHLBI),
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CENTERS FOR DISEASE CONTROL AND PREVENTION (CDC)

CDC will accept and consider SBIR grant applications in any area within the mission of the agency and awarding components (i.e., Institutes and Centers (ICs)) identified in this solicitation on September 8, 2020; January 5, 2021; and April 5, 2021 submission dates.

**CDC’s Mission:** CDC works 24/7 to protect America from health, safety and security threats, both foreign and in the U.S. Whether diseases start at home or abroad, are chronic or acute, curable or preventable, human error or deliberate attack, CDC fights disease and supports communities and citizens to do the same.

CDC increases the health security of our nation. As the nation’s health protection agency, CDC saves lives and protects people from health threats. To accomplish our mission, CDC conducts critical science and provides health information that protects our nation against expensive and dangerous health threats and responds when these arise.

**CDC Role:**
- Detecting and responding to new and emerging health threats
- Tackling the biggest health problems causing death and disability for Americans
- Putting science and advanced technology into action to prevent disease
- Promoting healthy and safe behaviors, communities and environment
- Developing leaders and training the public health workforce, including disease detectives
- Taking the health pulse of our nation

Those functions are the backbone of CDC’s mission. Each of CDC’s component organizations undertakes these activities in conducting its specific programs. The steps needed to accomplish this mission are also based on scientific excellence, requiring well-trained scientists and public health practitioners dedicated to high standards of quality and ethical practice.

CDC encourages investigator-initiated applications that focus on support for prevention, detection and response to emerging health threats. CDC is particularly interested in applications that address the current SARS-CoV-2 / COVID-19 pandemic response.

Before considering and/or preparing an application to the CDC SBIR program, all applicants are strongly encouraged to review CDC websites (agency, Center, Institute) and to contact the overall SBIR Program Manager or the research program coordinators listed in the Omnibus Solicitation.

**Research Topics of Interest to CDC**

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
<th>No*</th>
<th>Yes**</th>
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*If No,
This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, **UNLESS** the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,**

- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.

| Does IC/Office/Agency accept clinical trials applications under this mechanism? | Yes | No | Other Information:
If funding an SBC for clinical trials under a NON-SBIR/STTR award, please identify activity code(s) and other relevant information. |
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<td>Clinical Trials SBIR/STTR Omnibus/Parent Funding Opportunity Announcement/s</td>
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<tr>
<td>Clinical Trials SBIR/STTR IC/Office/Agency - Specific Funding Opportunity Announcement/s</td>
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<tr>
<td>Clinical Trials NON-SBIR/STTR Funding Opportunity Announcement/s where Small Businesses are eligible to apply</td>
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CDC will accept SBIR applications that both propose clinical trials and do not propose clinical trials, and all of the topics listed below cover both. CDC does not accept STTR applications.

**Limited Amount of Award**

CDC will not fund:

- Phase I applications greater than $243,500 total costs.
- Phase II applications greater than $1,000,000 total costs

For additional information about CDC, please visit our home page at [http://www.cdc.gov](http://www.cdc.gov).
Questions of a general nature about the CDC SBIR program should be directed to:

Sean David Griffiths, MPH  
Small Business Innovation Research Program (SBIR) Manager  
Office of Science, Office of Technology and Innovation  
Centers for Disease Control and Prevention (CDC)  
1600 Clifton Road NE, Mailstop H21-8  
Atlanta, GA 30329  
Phone: 404-639-4641; Fax: 404-639-4903  
Email: SBIR@cdc.gov

Or

Miriam Kelly, PhD  
Innovation Team Lead  
Office of Science, Office of Technology and Innovation  
Centers for Disease Control and Prevention  
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CENTER FOR GLOBAL HEALTH (CGH)

The Center for Global Health (CGH) leads the execution of the CDC’s global strategy; works in partnership to assist Ministries of Health to plan, manage effectively, and evaluate health programs; achieves U.S. Government program and international organization goals to improve health, including disease eradication and elimination targets; expands CDC’s global health programs that focus on the leading causes of mortality, morbidity and disability, especially chronic disease and injuries; generates and applies new knowledge to achieve health goals; and strengthens health systems and their impact.

Please visit their web site at: http://www.cdc.gov/globalhealth/index.html

(1) Development of Immersive Virtual or Augmented Reality Laboratory (AR/VR) Training Modules

**Background:** Laboratory science plays a vital role in the early detection, monitoring and response to a variety of diseases and health events as well as in surveillance to monitor the impact of vaccination programs. Efficient laboratory systems enable countries to better detect, respond, control and prevent the spread of disease. CDC global health programs support the improvement of global laboratory systems through training programs focused on various aspects of laboratory operations and management. Specific training includes workforce development, core laboratory competencies and management, facility and equipment maintenance, quality management system implementation, biosafety and biosecurity, as well as specific disease diagnostics (e.g. Zika, Ebola). Although the traditional hands-on practical approach is the mainstay of laboratory trainings, new and emerging technology-based approaches, such as electronic and mobile learning (E-Learning and M-learning); virtual learning (virtual classrooms), and immersive virtual reality (tactical and strategic immersions) techniques have been increasingly applied. CDC is actively promoting the development and utilization of cost-effective public health laboratory competency training tools using these innovative technologies.

**Specific Research Areas of Interest:** The goals for the proposed research are to address the limited availability of innovative virtual and augmented reality (VR/AR) training tools to improve public health laboratory capacity globally.

Examples of specific research areas of interest include, but are not limited to:
1. Development and field testing of immersive virtual and/or augmented reality laboratory training tools covering modules on guided instruction of standard operating procedures (SOPs) followed by immersive simulation to perform the SOPs.

2. Development and field testing of immersive virtual reality laboratory training tools covering visualization and interactive operation guides for complex, priority laboratory equipment (e.g. equipment used for molecular techniques, serology, immunology and culture).

3. Development and field testing of immersive virtual reality laboratory training tools covering step-by-step processes to mitigate laboratory biohazards, such as proper use and maintenance of engineering controls (e.g. biosafety cabinet) and personal protective equipment (PPE) (e.g. respirator, gowing & gloving) to optimize performance and maximize safety of laboratory operations.

**Impact and Commercialization Potential:** This research will result in increased availability of innovative and cost-effective immersive VR/AR laboratory training tools for global public health laboratory competency training programs that will complement the resource-intensive, hands-on, "wet" laboratory trainings that utilize live clinical specimens. The commercialization potential of these training products is high, as VR/AR training tools for other highly technical subjects have been successfully commercialized by different companies (e.g. Virtual Surgical Training, Flight Simulators, and Undergraduate Science Training etc.).

Visit the CGH homepage for more information on CGH's research program areas [http://www.cdc.gov/globalhealth/index.html](http://www.cdc.gov/globalhealth/index.html)

For CGH programmatic information, contact:

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NATIONAL CENTER ON BIRTH DEFECTS AND DEVELOPMENTAL DISABILITIES (NCBDDD)

The mission of CDC’s National Center on Birth Defects and Developmental Disabilities (NCBDDD) is to promote the health of babies, children and adults and to enhance the potential for full, productive living. To achieve its mission, the Center works to identify the causes of birth defects and developmental disabilities, helps children to develop and reach their full potential, and promotes health and well-being among people of all ages with disabilities, including blood disorders. NCBDDD seeks to accomplish these goals through research, partnerships, and prevention and education programs.

Please visit their web site at: http://www.cdc.gov/ncbddd/index.html

(2) Improving Newborn Screening of Coarctation of the Aorta

Background: There are approximately 6000 infants born each year in the United States with nonsyndromic critical congenital heart disease, a condition that typically requires surgical intervention during infancy in order to survive. Before the advent of newborn screening for critical congenital heart disease, approximately 30% of these children were discharged home after birth without a diagnosis having been made. In 2011, critical congenital heart disease was added to the United States Recommended Uniform Screening Panel, with all states now including this condition as part of their newborn screening panel as of 2018. Newborn screening for critical congenital heart disease has been shown to lead to earlier diagnosis, earlier treatment, and, as a result, improved survival of children with this condition. However, the sensitivity of current screening practices remains poor at only 50%. A key driver of this low overall sensitivity is the poor sensitivity of current pulse oximetry screening for coarctation of the aorta.

The current non-invasive method of newborn screening for critical congenital heart disease includes using pulse oximetry on the right hand and either foot of an infant to determine the oxygen saturation of blood. Low levels of oxygen saturation or notable differences in saturation between the right hand and either foot suggest that a child might have critical congenital heart disease. Subsequent testing is then performed to either confirm or rule out critical congenital heart disease. This method works well for most of the 12 main types of critical congenital heart disease. However, for the most common type, coarctation of the aorta, screening with pulse oximetry has a sensitivity of only 36%.

Coarctation of the aorta is a narrowing of the aorta such that the lower half of a child’s body does not receive adequate blood flow. Left untreated, this condition may lead to cardiogenic shock and death. Sometimes a patent ductus arteriosus, a blood vessel bypassing the narrowing in the aorta, provides blood flow with a lower oxygen saturation to the bottom half of the body. Current screening practices may detect coarctation if a patent ductus arteriosus is present, but this is often not the case. Attempts have been made to improve detection by studying the perfusion index, a measure of the pulsatility of blood flow in the distal circulation. However, these efforts have not been successful thus far. Therefore, there is an unmet need for a reliable method to screen for coarctation of the aorta in the newborn.

Specific Research Areas of Interest: The goal of this project is to design, develop, and test the feasibility of a new and more sensitive method of screening for coarctation of the aorta in a newborn infant at approximately 24 hours of age. This new method may include new equipment, new software that can be added to existing equipment (e.g. pulse oximeters), or a new application of existing technology. This new method must balance the screening ideals of high sensitivity, high specificity, affordability, and ease of use. Ideally any new method would add minimal burden and cost to the existing method of screening using pulse oximetry.
Impact and Commercialization Potential: Reliable, sensitive screening of coarctation of the aorta among all U.S. newborns would lead to earlier diagnosis and treatment for up to 560 additional newborns each year in the United States; this, in turn, would lead to improved survival for infants with coarctation of the aorta. Furthermore, with a highly sensitive screening method for coarctation of the aorta, the overall sensitivity of newborn screening for critical congenital heart disease would increase from 50% to as high as 82% if all cases of coarctation of the aorta are detected.

Successful technology can be implemented for screening of all newborns (approximately 3.8 million newborns in the United States each year). This can be either new stand-alone technology that is added to current newborn screening practices, or adjunct technology that is incorporated into current pulse oximetry practices.

For NCBDDD programmatic information, contact:

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**National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP)**

The CDC’s National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP) carries out a variety of activities that improve the nation's health by preventing a range of chronic diseases such as arthritis, cancer, diabetes, heart disease, obesity and stroke, while promoting health and wellness in the areas of reproductive health, oral health, nutrition and physical activity. The Center’s activities include supporting states’ implementation of public health programs; public health surveillance; translation research; and developing tools and resources for stakeholders at the national, state, and community levels.

Please visit their web site at: [http://www.cdc.gov/chronicdisease/index.htm](http://www.cdc.gov/chronicdisease/index.htm)

**(3) Using Innovative Technologies to Track Policies and Enhance Evidence-Based Policymaking**

**Background:** Evidence-based policies are approaches that have the greatest potential for impact on health because they reach entire populations of people at once. For example, seat belts reduce serious crash-related injuries and deaths by about half and saved almost 15,000 lives in 2016. Similarly, residents of cities with comprehensive smoking bans have been shown to be 21.1% less likely to currently smoke. Recent changes in Federal law prompted guidance for the consistent application of accepted evaluation standards and the implementation of best practices for Federal evaluation. The Foundations for Evidence-Based Policymaking Act of 2018 along with the recent expansion of legal epidemiology (defined as the scientific study and deployment of law as a factor in the cause, distribution, and prevention of disease and injury in a population) have stimulated the opportunity to create robust, systematic public health policy datasets. To date, development of policy datasets appropriate for use in rigorous evaluations have been costly to create and resource intensive to maintain. Because of this, public health researchers often rely on cross-sectional (single point in time) policy datasets and simple pre-post assessment methods to examine public health policies. Strengthening these designs with available longitudinal (repeated measures over time) data will enhance the ability to produce actionable information and draw causal inferences essential to meet emerging evaluation needs.

To date, the systematic analysis of law at state and local levels requires skilled policy analysts to search, capture and code complex information from multiple sources. However, once an initial dataset is created, there is an opportunity to use technology such as artificial intelligence (AI) to automate data processing when identifying changes in law over time. Additionally, the use of machine learning algorithms could be used to further improve the time, cost, and accuracy of coding longitudinal datasets to dramatically enhance their feasibility and availability for evaluation purposes. This novel application of AI and related machine learning techniques could immediately benefit local, state and federal government agencies, academic and private researchers, non-governmental organizations, and public health practitioners in analyzing the distribution and impacts of law on health in a population.

**Specific Research Areas of Interest:** Recognizing that longitudinal datasets are crucial to understanding the ongoing impact of law and policy on health outcomes at a population level, the goal of this SBIR is to pilot the use of machine learning techniques to support the development of longitudinal public health policy datasets for tracking and assessment purposes.

The initial project test case will be used to inform the accuracy, efficiency, and performance of applied machine learning algorithms to populate and code policy variables across time and jurisdiction. Specifically, the pilot will address the following key questions:

- What AI technology resources are needed for initial testing and implementation? Do these needs change over time? If so, how?
- What are the labor resources needed for development and maintenance of AI assisted longitudinal datasets over time?
- What modifications are needed to existing policy datasets or processes to facilitate implementation of machine learning algorithms to enhance coding accuracy and efficiency?

- Are results sufficient in terms of both specificity and sensitivity?

- How could future use be expanded to track new and emerging issues rather than relying solely on topic areas for which an existing policy dataset already exists (i.e., develop and create new legal datasets on new topics)?

- What are the limitations and challenges related to use for state versus local policy tracking?

- Can the approach be adapted to work with a variety of source material, i.e., if the initial test case scans systems capturing enacted legislation, can future applications scan systems that track proposed legislation at both the state and local levels?

The proposed activity will require an applicant with capacity to:

- Conduct a preliminary assessment to determine the existing processes for searching, capturing, and coding policy information. Using this information, identify several ‘gold standard, trained datasets to serve as baselines for which future machine learning algorithms can be assessed against for accuracy, clarity, and nuances in coding legal data. The information collected will be used to examine the utility of existing datasets for testing machine learning algorithms.

- Consider the functionality and feasibility of existing commercially available and open source AI/machine learning platforms and products (e.g., IBM Watson, Tensor Flow, Amazon Lex, Python) given the unique needs and objectives of the pilot.

- Implement the machine learning test case against the previously identified ‘gold standard,’ trained datasets. Work to include inputting and parsing text from selected policy datasets; building a neural network based on identified search parameters and capture criteria; determining potential challenges in capturing information from primary data sources including Westlaw; training, testing and refining the model using the ‘gold standard’ policy datasets.

- Produce 2-3 chronic disease-related policy datasets with accompanying assessment metrics (e.g., F1 score which is the weighted average of precision and recall) to determine the utility for future work.

- Summarize test case findings in a summary report outlining the feasibility of multiple approaches, successes, failures, areas for further testing, and unresolved issues.

References:


**Impact and Commercialization Potential:** This system could have wide reaching implications for policy research. The resulting machine learning approach to create longitudinal public health policy datasets developed as part of this project will enable public health researchers and practitioners to more quickly and easily assess the public health impacts of policies. The approach, if successful, will dramatically reduce the time, cost, and expense of developing policy datasets. Additionally, it will enhance the quality of policy assessments by providing ready access to longitudinal information and transform the field of legal epidemiology. Skilled policy analysts and public health attorneys will be able to focus on issues of meaning and impact that address causation (analyzing linked legal epidemiology datasets and health/economic outcomes) rather than repetitive and time intensive data capture and coding tasks. State and local public health practitioners will be better able to identify the strengths and challenges of different policy approaches, identify existing gaps, and determine health impacts of policy on disparate populations across jurisdictions.

The commercialization potential is multi-faceted. (1) The resulting algorithms developed as part of this research could be replicated and offered as a Platform as a Service (PaaS) to organizations that currently develop policy datasets. The platform could operate on a subscription-based revenue model or a product licensing model. (2) Opportunities could be developed to offer a subscription or fee-based approach to provide direct access to a databank of developed longitudinal policy datasets. (3) The algorithm developed as part of this study could be sold to an institution or other entity. End users include local, state and federal government agencies, academic and private researchers, non-governmental organizations, public health practitioners, as well as a range of public and private businesses that routinely track change in public health law.

(4) **Implementation Resources for Prescribing of Hormonal Contraception by Pharmacists**

**Background:** Almost half of U.S. pregnancies are unintended. Unintended pregnancy is associated with greater risk of adverse outcomes for both mothers and infants as well as increased health care costs. To reduce unintended pregnancy and related adverse health outcomes, and to promote optimal birth spacing for improved maternal and child health, strategies to increase access to and use of effective contraceptive methods are critical. However, many women face substantial barriers to accessing contraception, including availability and accessibility of healthcare providers, transportation and logistics challenges, and cost.

Pharmacists play a key role in assisting patients to choose and use contraception successfully. Currently, 9 states authorize pharmacists to prescribe certain hormonal contraceptive methods (e.g., oral contraceptives, patches, rings, and injections) directly to patients, and at least 17 more states are considering such legislation. Pharmacist-prescribed contraception services can improve contraceptive access to reproductive-aged women by decreasing some of the access and cost issues associated with a clinician visit. Most states allow pharmacist-prescribed contraception for all ages; in these states, adolescents may particularly benefit from contraceptive access through pharmacists, as they be more likely than adult women to face barriers to clinical services. Expanding access to contraception through pharmacists takes on added importance during national public health emergencies, such as the current COVID-19 response, when access to family planning and primary care providers may be limited.

While there have been early successes in providing pharmacist-prescribed contraception services in some locations, documented barriers to implementation include training, logistics, and reimbursement issues. While all states that allow pharmacist-prescribed contraception services have minimum training requirements, pharmacists report on-going training needs. Point-of-care tools and job aids on contraceptive counseling, medical eligibility criteria, and contraception initiation and management practices would give pharmacists on-going access to resources and continuing education. Logistics
around confidentiality, private space for counseling, and workflow in the pharmacy need to be addressed. Finally, a major obstacle to implementation has been billing and reimbursement issues. Tools to establish reimbursement infrastructure, including billing insurance, are necessary for successful and sustainable implementation of these services.

**Specific Research Areas of Interest:** The goal of the project is to develop an implementation package for pharmacists, pharmacy managers, and pharmacies who want to implement or improve implementation of pharmacist-prescribed contraception services. The implementation package could include resources for: a) patient-centered contraceptive counseling and provision, b) logistics of pharmacist-prescribed contraception services, c) billing and reimbursement strategies, and d) monitoring and evaluation of implementation activities (e.g., at pharmacy, region, chain, or state levels). This package could serve as the blueprint for pharmacies to successfully implement, monitor, and improve pharmacist-prescribed contraception services in a sustainable model within the pharmacy.

The applicant should have the capacity to:

1. develop a package of implementation tools for pharmacist-prescribed contraception services;
2. partner effectively with multiple stakeholders, including pharmacists and pharmacies (including independent pharmacies, health care system-based pharmacies, and large chain pharmacies), as well as professional organizations such as the American Pharmacists Association (APhA);
3. create a commercialization plan for the implementation package that will appeal to relevant stakeholders.

Development activities should be clearly centered within a compatible implementation sciences framework (e.g., Consolidated Framework for Implementation Research [CFIR; https://cfirguide.org/]). Strong consideration should also be given to whether any of these tools could be embedded into existing pharmacist tools or resources, rather than as a stand-alone resource for contraception services.

**Phase 1 activities may include, but are not limited to:**

1. Conduct a needs assessment around resources for pharmacist-prescribed contraception services, including a review of any available tools or training resources, and any validated evaluation or performance measures. Some existing resources include CDC’s evidence-based guidance for contraception and family planning services (US Medical Eligibility Criteria for Contraceptive Use, US Selected Practice Recommendations for Contraception Use, and Providing Quality Family Planning Services; https://www.cdc.gov/reproductivehealth/contraception/contraception_guidance.htm), and training resources already developed for pharmacist-prescribed contraception services by individual states and pharmacy boards, as well as from APhA (https://www.pharmacist.com/).

2. Develop a prototype of an implementation package, for example, building out a select number of components for testing and evaluation purposes.
3. Pilot test the prototype with key end-users.
4. Develop plans to refine and complete the implementation package to ready the package for private sector commercialization, based on lessons learned.

**Impact and Commercialization Potential:** Pharmacist-prescribed contraception services have the potential to substantially increase access to contraception by removing many barriers faced by patients, including access and cost issues of visiting a family planning provider or clinic. While surveys of pharmacists and patients show broad interest and support for expanding contraceptive access in pharmacies, direct provision of contraception by pharmacists is relatively new and pharmacists face several barriers to implementing these services, such as training needs, logistics, and billing and
reimbursement strategies. However, pharmacists have overcome similar barriers in other direct patient care services (e.g., vaccination, chronic disease management). Therefore, providing pharmacies and pharmacists with the implementation tools needed now, as these services start to expand, will give them the best chance for success. The larger impact of this implementation package will be to increase opportunities for contraception provision in a setting that is widely accessible to patients in need of initiating contraception, managing problems with contraception, or switching to a more suitable contraceptive method. Overall, implementation of these services could increase the number of women screened for the need and desire for contraceptive services, increase contraception provision to those in need, and potentially decrease unintended pregnancies. In addition, pharmacists have indicated that participating in pharmacist-prescribed contraception and other direct patient care services would increase job satisfaction. Effective implementation tools for sustainable pharmacist-prescribed contraception services may also be attractive to pharmacies seeking new revenue opportunities.

The pharmacist-prescribed contraception services implementation package could be marketed to independent pharmacies, large chain pharmacies, large health systems, states who have recently passed legislation and want to jump-start implementation, and other stakeholders interested in effective and sustainable pharmacist-prescribed contraceptive services. As pharmacist-prescribed contraception services continue to grow, the implementation package could be continuously updated as new implementation strategies are developed and evaluated. In addition, these tools, especially around logistics and reimbursement, could be adapted and expanded to address similar issues with other pharmacy services.

(5) Electronic Transmission of Patient-reported Pregnancy Outcomes after Assisted Reproductive Technology

**Background:** Infertility is a disease of the reproductive system and an important public health issue that affects millions of families in the United States and globally. In the U.S., one in six couples in which the woman is older than 35 years have fertility problems. Since 1978, in vitro fertilization (IVF), the most common type of Assisted Reproductive Technology (ART) that involves handling of eggs or embryos for the purpose of establishing pregnancy, has been successfully used to help millions of infertile couples. In addition, many clinics offer fertility treatments that do not involve ART, such as those performing ovulation induction and intrauterine insemination (non-IVF fertility treatments). The unprecedented growth of the field of fertility treatments during the last four decades is expected to continue because of the growing demand resulting from delayed childbearing in developed nations and increasing availability of these treatments. Although most countries have national registries to monitor the use and effectiveness of ART, collecting data on pregnancy outcomes is challenging because fertility treatment and obstetric care are usually provided by different specialists. In addition, ART registries are typically not designed to collect data beyond delivery. As a result, our knowledge of the long-term effect of ART on maternal and child health is limited.

In the United States, the Fertility Clinic Success Rate and Certification Act (FCSRCA) of 1992 mandates that clinics performing ART annually provide pregnancy success rates data to the Centers for Disease Control and Prevention (CDC). Fertility clinics are required to report ART cycle-level data, including details of each ART procedure performed, outcomes of the ART procedure (e.g. pregnant versus not pregnant), and outcomes of any resultant pregnancy (e.g. live birth versus no live birth). Since fertility clinics provide care to patients undergoing ART treatment, detailed information about the treatment procedure is collected directly by the fertility clinic. However, if pregnancy is achieved, the patient typically receives subsequent perinatal and obstetric care from another provider (obstetrician/gynecologist). Reporting ART data to CDC is a burden for fertility clinics that often need to hire additional staff for this purpose. ART clinics may face obstacles in collecting pregnancy outcome information effectively and efficiently from ART patients that have achieved pregnancy or from their subsequent obstetric providers. They rely on traditional mechanisms, including telephone, mail, and email, to obtain these data and they often have to make multiple attempts to get the required information. This process can be even more challenging if patients come from outside of the state or outside of the U.S. to receive ART treatment and
then return home to care for the pregnancy. Some fertility clinics may lack a user-friendly, convenient, secure, and verified communication system for patients to either: (1) personally report pregnancy outcomes to the fertility clinic in a safe and secure manner and/or (2) provide information on their obstetric provider and consent to the release of pregnancy outcome information to the fertility clinic.

Pregnancy outcome information is a vital component of ART surveillance and research. The accuracy of this information in the ART surveillance system is critical for reporting ART success rates and understanding maternal and treatment characteristics associated with treatment success. CDC uses the ART surveillance data, including the pregnancy outcome data, for several purposes, including the publishing of annual clinic-specific success rates reports and other annual reports which describe the characteristics of ART treatment procedures and outcomes at a national and state level. Accurate pregnancy outcome data are also very important for ensuring data linkages between ART surveillance data, vital records, and other disease registries. These data linkages are used to generate more robust surveillance and research datasets to allow additional research on short- and long-term maternal and child health outcomes such as complications of labor and delivery, neonatal complications, and other conditions identified after birth. This information will facilitate research on effectiveness and safety of ART and may help to improve maternal and child health outcomes of fertility treatments.

**Specific Research Areas of Interest:** The primary goal of this project is development of an effective technology solution, such as software or phone application, to facilitate electronic transmission of pregnancy outcome data from the patient and/or obstetric provider to the fertility clinic. The technology solution should be user-friendly, available on multiple platforms (such as iOS, Android, and Windows), and convenient for both the ART patients, clinicians, and the fertility clinic. It should be secure, as it may involve transmission of personally identifiable information.

A secondary goal is to use the technology solution as a mechanism for infertility patients and fertility treatment providers to communicate necessary health information during the treatment process via a secure channel. For example, patients could track the timing of medication administration, report side effects, and ask questions, while providers could track adherence to the treatment plan, adjust dosage, and schedule appointments.

In addition, after receiving patient consent, the technology solution can serve as a platform for collecting information directly from patients to enable patient-centered outcome research on the long-term health of patients and their children.

**Impact and Commercialization Potential:** The development of an effective technology solution that facilitates electronic transmission of information between infertility patients and fertility clinics will benefit: (1) infertility patients undergoing ART, (2) fertility clinics who collect and/or report this information, (3) organizations performing ART surveillance, such as the CDC, and (4) reproductive health researchers interested in collecting longitudinal information on maternal and child health post treatment. Infertility patients will benefit from having a seamless and secure communication channel with their fertility treatment provider, which can save them time and allow reporting of pregnancy outcome data at their convenience. It may also provide a more sensitive approach for collecting the information in the event that a pregnancy does not result in a live birth. At the clinic level, having a convenient and secure information delivery mechanism will help fertility clinics save resources typically spent on sending patient reminders or contacting patients. In addition, this technology could be used by providers to assess regimen adherence and satisfaction with services among their patients. Organizations performing ART surveillance may benefit in receiving more complete and accurate pregnancy outcome data, particularly for outcomes such as miscarriage or fetal death. Improved completion and accuracy of pregnancy outcome data will ultimately improve ART surveillance and the quality of research aimed at improving perinatal outcomes of fertility treatments. Reproductive health researchers will receive previously unavailable opportunity to develop registries for patient-centered outcome research using this platform to collect longitudinal data on infertility patients and their children to better understand long-term outcomes of fertility treatments.
Overall, this project has the potential to strengthen collaboration between small technology-promoting businesses, healthcare, science, and public health in the area of infertility.

The field of infertility treatments is not only a successful field of medicine, but also a multi-billion-dollar industry. This project facilitates the creation of an effective technology solution that facilitates secure electronic transmission of ART outcome information between patients and/or their obstetric providers and fertility clinics. This should provide a tangible product such as a software or phone application, which could be sold to fertility clinics or reproductive health researchers to establish secure line of communication with infertility patients. The market for the technology solution will not be limited to the U.S. alone – almost every country in the world has national ART surveillance which faces similar challenges of collecting information from infertility patients after treatment, and therefore could benefit from the development of this product. Nearly 500 fertility clinics in the U.S. perform ART and are required to report ART surveillance data per the FCSRCA. In addition, clinics that offer non-IVF fertility treatments may be interested in utilizing the solution because many clinics track their success rates to understand the effectiveness of the treatment they offer even if reporting to a national surveillance system is not required. The technology solution can also be marketed to reproductive health researchers interested in studying ART outcomes.

Visit the NCCDPHP homepage for more information on NCCDPHP’s research program areas [http://www.cdc.gov/chronicdisease/index.htm](http://www.cdc.gov/chronicdisease/index.htm)

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NATIONAL CENTER FOR EMERGING AND ZOONOTIC INFECTION Diseases (NCEZID)

The National Center for Emerging and Zoonotic Infectious Diseases (NCEZID) aims to prevent disease, disability, and death caused by a wide range of infectious diseases. NCEZID focuses on diseases that have been around for many years, emerging diseases (those that are new or just recently identified), and zoonotic diseases (those spread from animals to people). Work is guided in part by a holistic "One Health" strategy, which recognizes the vital interconnectedness of microbes and the environment. Through a comprehensive approach involving many scientific disciplines, better health for humans and animals and an improved environment can be attained.

Please visit their web site at: http://www.cdc.gov/ncezid

(6) Antibiotic Resistant Healthcare-Associate Infections

**Background:** Healthcare-associated infections (HAI) are a threat to patient safety. CDC provides national leadership in surveillance, outbreak investigations, laboratory research, and prevention of healthcare-associated infections. CDC uses knowledge gained through these activities to detect infections and develop new strategies to prevent healthcare-associated infections. Healthcare-associated infections (HAIs) can be found to effect 1 in 25 hospitalized patients on any given day in the United States, leading to an annual burden of 722,000 infections and 75,000 deaths. Meanwhile, among 18 antibiotic resistant (AR) organisms identified by CDC in 2013 as urgent, serious, and concerning threats, nearly half are primarily healthcare-associated. Whereas 1 in 7 HAIs in hospitals overall are caused by AR-threat bacteria, in some types of hospitals, AR-threat bacteria cause 1 in 4 infections. In all cases, HAIs caused by AR-threats are more difficult to treat and some are now untreatable. There are three broad, current strategies that clinicians and nurses need to employ to prevent these AR HAIs: following recommendations for preventing invasive device and surgical procedure-related infections, preventing cross-transmission of AR HAI pathogens, and practicing optimal antibiotic stewardship. In addition, there is a great need for innovation and commercial development in the following three priority areas.

**Specific Research Areas of Interest:**
The goals for the proposed research are to address antibiotic resistant healthcare-associated infections.

Examples of specific research areas of interest include, but are not limited to:

1) Development of novel diagnostics that either: A) offer a more rapid and definitive diagnosis of whether a patient does or does not require an antibiotic (alternatively whether it is safe to stop an antibiotic), or B) better detect (i.e., earlier, more rapidly, and more accurately) whether a patient is infected or colonized (and thereby may transmit) with an AR HAI pathogen.

2) Novel therapeutics and preventatives based upon preservation or restoration of the human microbiome

**Impact and Commercialization Potential:** This research will lead to the development of practical and innovative solutions to address the matrix of complex problems caused by antibiotic resistant healthcare-associated infections. Successful and novel innovation that will reduce disease, disability, and death will have huge commercial potential across many markets.


**Background:** Overuse of antimicrobial drugs in agriculture, medicine, and industry has resulted in continual pressure for pathogenic organisms to evolve mechanisms by which to evade these drugs. The National Antimicrobial Resistance Monitoring System (NARMS) is a collaborative effort of state public health departments, FDA, CDC, and USDA to monitor trends in antimicrobial resistance over time using a ‘farm to fork’ approach. Although the advent of advanced molecular detection techniques has increased
our ability to detect bacterial resistance patterns, there are knowledge gaps that remain to be addressed. Detection of resistance to clinically relevant drugs requires a laboratory setting and takes days, if not longer, and more research is needed to link data generated by molecular detection to clinical outcome. Finally, the laboratory community has realized that the exciting potential of culture-independent tests may also have an undesired outcome; the loss of important organic material for future study.

**Specific Research Areas of Interest:** The goals for the proposed research are to detect, transmit, and prevent antimicrobial resistance in enteric bacteria.

Examples of specific research areas of interest include, but are not limited to:
1. Rapid, portable, point of care diagnostic and field assays that simultaneously identify bacterial agents and clinically relevant resistance markers
   - Lateral flow technology to detect biomarkers
   - High throughput molecular tests
2. Development of an in vitro system to simulate myriad physiological conditions (human or ruminant gut, for example) in which enteric bacteria develop drug resistance
   - 3D polymer scaffold or 3D-printed substrate ‘organ’ for growth of bacteria in the presence of secretory immune factors to which antimicrobials may be applied or dosed
3. A matrix for archiving bacterial cultures that does not require a cold-chain or frozen storage
   - Preserves the integrity of the organisms
   - Storage matrix requires a tiny footprint, similar to filter paper.

**Impact and Commercialization Potential:** Using an effective in vitro “microbiome” system to study development, rate of transmission and ecology of antimicrobial resistance would require less time and human capitol than the large clinical studies that are required to evaluate current and new antimicrobial pharmaceuticals. A system such as this could help to narrow one of the major knowledge gaps in understanding antimicrobial resistance; the correlation between laboratory-determined antimicrobial breakpoints and clinical outcomes. A simple rapid method of preserving important bacteria and organic material related to bacterial resistance would be embraced by the reference and research community. Existing rapid tests could be modified to add detection of clinically relevant resistance markers, thereby dramatically decreasing time to treatment decision. Reduction of footprint and ambient storage would reduce operating and shipping costs that are currently associated with these materials.

(8) Vector Borne Diseases: Detection, Prevention, Diagnosis and Response

**Background:** Bacterial and viral vector borne diseases are transmitted to humans primarily through vectors such as an infected mosquito, tick or flea. Some of these diseases have long been present in the United States while others have recently emerged.

Vector-borne diseases are a major public health concern. Lyme disease causes over 300,000 estimated human illnesses annually in the U.S. Tick-borne rickettsial diseases, such as Rocky Mountain spotted fever, ehrlichiosis, and anaplasmosis, are responsible for over 4,000 U.S. cases each year, including some that result in death. Dengue fever causes millions of cases worldwide, including thousands of cases in Puerto Rico each year. Outbreaks of arboviral diseases such as West Nile encephalitis and Chikungunya fever have been reported in recent years. In May 2015, the Pan American Health Organization (PAHO) issued an alert regarding the first confirmed Zika virus infection in Brazil and on February 1, 2016, the World Health Organization (WHO) declared Zika virus a public health emergency of international concern. Local transmission has been reported in many other countries and territories. Less common, but often deadly threats such as Yersinia pestis causes the ancient disease plague. Local plague outbreaks occur in the southwestern U.S., and it is a significant health threat in Africa and Asia.

**Specific Research Areas of Interest:** The goal of this project is to encourage research that will enhance prevention, detection, diagnosis and response capabilities to vector borne diseases through funding innovative solutions that address the following:
1. Mitigate the spread and impact of vector borne diseases
2. Improve our ability to prevent, detect and respond to outbreaks of vector borne diseases
3. Develop diagnostic tests to differentiate among vector borne diseases
4. Develop vaccines effective against vector borne diseases

Examples of specific research areas of interest include, but are not limited to:

- Development of improved laboratory tests to diagnose vector borne diseases in the field or in healthcare settings (e.g., new diagnostics to detect and differentiate among vector borne diseases after infection, etc.)
- Development of tools to improve monitoring and reporting cases of vector borne disease infection and sequelae
- Development of tools to improve surveillance for vector borne diseases in the US and elsewhere (e.g., better surveillance applications, improved clinical, laboratory, and epidemiological data linkage, interchange, analysis, and visualization, etc.)
- Development of tools to improve linkage to and monitoring of services for vector borne disease-affected families
- Development of tools to improve mosquito, tick and flea control in and around individual houses and in the exterior environment

**Impact and Commercialization Potential:** Vector-borne diseases continue to cause morbidity and mortality in endemic areas where the threat from these diseases is recurrent. In addition, these diseases can emerge rapidly and unpredictably causing wide-spread outbreaks. Similar symptoms to other diseases can make diagnosis based on symptoms alone difficult and current diagnostic tests can cross-react among different causative agents (e.g., dengue and Zika virus infections). Effective vaccines are not available and environmental control is in need of improvement. Given the large number of individuals affected by these diseases, and the challenges to public health for their containment, improved detection through better diagnostic tests and improved prevention through vaccination would have a great impact on the health of the nation. The proposed research should lead to the development of practical solutions for the detection, prevention and diagnosis of vector-borne diseases. The products and innovations developed through this process will have significant commercial potential and will improve public health and the healthcare system’s response to vector-borne diseases.

Visit the NCEZID homepage for more information on NCEZID’s research program areas [http://www.cdc.gov/ncezid](http://www.cdc.gov/ncezid)

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Technology to Develop a Monitor to Detect Iodine Deficiency

**Background:** Iodine deficiency (IdD) is linked to many neurodevelopmental disorders and is the world’s leading cause of intellectual deficits. While IdD is well known in Africa, India, and central Asia, IdD also exists in the US. Although the overall U.S. population is not iodine deficient, CDC National Health and Nutrition Examination Survey (NHANES) data suggests that a significant fraction of the U.S. population may be iodine-deficient. Lack of quality iodine nutrition in pregnant and lactating women places the fetus/infant at risk of impaired neurodevelopment.

Although IdD spot checks in pregnant and lactating women can indicate a problem in a population, epidemiological investigations suggest that 24h urinary iodine anion samples corrected for creatinine are better for case identification, since much of the iodine is quickly excreted in the urine (along with potassium and sodium). Although randomized sampling and spectrometry also show promise in accurately detecting IdD, analysis is best determined by catalytic reduction of cerium ammonium sulfate in the presence of arsnic acid after the urine is acid digested under mild conditions. Such methodology is complicated, time consuming and is not readily available to physicians to aid in diagnosis.

While rapid strip tests based on the iodide-catalyzed oxidation of 3,3’,5,5’-tetramethylbenzidine by peracetic acid/H2O2 yield a colorimetric output to determine uncorrected iodine ranges, these strip tests are not sensitive and lack the specificity of the laboratory methods in that they provide only aggregate data and cannot probe with specificity for iodine levels below 100 ug/L. An affordable, reliable, sensitive method for IdD detection (at least to 33 ug/L) is needed. Ideally, the method should be capable of being carried out in a physician's office as part of a routine checkup.

**Specific Research Areas of Interest:** The goal of this research is to develop technology that will provide a simple sensitive IdD detection method for OBGYNs and pediatricians to use in their offices to identify urinary iodine at <33 ug/L. The ideal device would screen multiple samples with clearly quantified measurements. Phase I (6 months): The awardee should conduct a feasibility study to develop a prototype IdD monitor capable of detecting urinary iodine at <33 ug/L. The awardee should provide research data regarding the sensitivity and specificity of the IdD monitor.
**Impact and Commercialization Potential:** While there are several methods to identify IDd in pregnant and lactating women, high tech laboratory methods are expensive and time-consuming, and strip tests, while inexpensive, are too insensitive to measure levels of iodine that are expected to be harmful. Successful development and subsequent use of an IDd monitor capable of detecting iodine at levels <33 ug/L will improve identification of IDd in expectant and lactating mothers, and in infants. IDd monitoring technology developed from research funded under this topic has the potential to prevent new cases neurological disorders resulting from IDd. There is an approximate need for 15,000 devices in the US and multiples more worldwide as extreme IDd is found in 54 countries. This product has commercial appeal to hospitals, clinics, public health departments, and physician practices in which IDd monitoring is conducted.

Links to associated patents:


Strip method detection is patented here: [https://patents.google.com/patent/US6689618](https://patents.google.com/patent/US6689618)

Colorimetric reaction is described here: [https://academic.oup.com/jcem/article/83/3/1007/2865523](https://academic.oup.com/jcem/article/83/3/1007/2865523)

The ICP-MS method is discussed here: [https://file.scirp.org/pdf/AJAC_2017041415342563.pdf](https://file.scirp.org/pdf/AJAC_2017041415342563.pdf)

Visit the NCEH homepage for more information on NCEH’s research program areas [http://www.cdc.gov/nceh/](http://www.cdc.gov/nceh/)

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NATIONAL CENTER FOR HIV/AIDS, VIRAL HEPATITIS, STD, AND TB PREVENTION (NCHHSTP)

The National Center is committed to our vision of a future free of HIV/AIDS, viral hepatitis, STDs, and TB. NCHHSTP is responsible for public health surveillance, prevention research, and programs to prevent and control HIV and AIDS, other STDs, viral hepatitis, and TB. CDC’s National Center for HIV, Viral Hepatitis, STD, and TB Prevention’s (NCHHSTP) Strategic Plan Through 2020 articulates a vision, guiding principle, and overarching goals and strategies through 2020 to influence and enhance our programs. The three overarching goals highlighted in this plan are to decrease:

- Incidence of infection,
- Morbidity and mortality, and
- Health disparities.

Every year, millions of Americans are infected with HIV, viral hepatitis, STDs, or TB and tens of thousands die from their infection. Most of these infections share commonalities, from modes of transmission to demographic, social, and economic conditions that increase risk. As a prevention leader, NCHHSTP focuses on high impact prevention and control efforts to reduce incidence, morbidity, mortality, and health disparities due to these infections.

Please visit their web site at: http://www.cdc.gov/nchhstp/

(10) Improved Diagnostic Tests for HIV, STDs, Hepatitis and TB

**Background:** It is estimated that just over 1.2 million people in the United States are living with HIV infection, and almost 1 in 8 (12.8%) are unaware of their infection. Because there are several treatment and prevention options for HIV, a major goal of CDC, other public health agencies and our public and private partners is to further improve the percentage of people that know their HIV status. For individuals that are at risk and uninfected, it allows them to focus on prevention. For those that are infected, there is growing evidence that the sooner a person knows they are infected and can start treatment, the better their overall health can be maintained. There is also emerging evidence that early diagnosis leads to preventing further spread of the virus due to changes in behavior by those who know their status.

Whereas there are specific benefits for HIV testing and treatment, testing and treatment for comorbid pathogens such as TB, hepatitis (B and C), gonorrhea and syphilis are also of great benefit in populations at risk for HIV. These diseases (STDs, TB and hepatitis) can all lead to worse health outcomes for HIV infected individuals. Furthermore, having diseases such as syphilis and gonorrhea can increase the chances of someone acquiring HIV. Because there is an effective vaccine for hepatitis B and effective therapy for hepatitis C, syphilis, gonorrhea and TB, improving tests and testing for these pathogens can lead to a further decrease in HIV transmission or morbidity.

Whereas there is tremendous value in testing and diagnosis for each of the described diseases, some at risk individuals are never tested or do not receive their tests results and often times are only tested for one of the diseases when testing for a combination of the diseases would be more beneficial both for the individual and for public health.

Prognostic tests (e.g. viral load, drug resistance monitoring) also play an important role in improving health outcomes for individuals infected with HIV and the ability to predict recent or long term HIV infection can be used for public health action.

**Specific Research Areas of Interest:** The major goal of the project is development of diagnostic reagents, tests or testing platforms, that will further improve diagnosis or monitoring of HIV or comorbid pathogens such as hepatitis (B and C), syphilis, gonorrhea or TB. The specific area of interest is innovative approaches or novel technology that would allow for diagnosis of HIV and other comorbid pathogens such as hepatitis (B and C), syphilis, gonorrhea and TB alone or in any combination using a
single test device or platform. The preferred reagents, test format or technology would facilitate testing that allows for rapid results (preferably less than one hour), is affordable (comparable to currently available tests) and can be performed at the point of care or in a laboratory capable of performing moderately complex tests.

Consideration will also be given to innovative technology that provides prognostic (monitoring) results such as viral load (HIV, HCV, HBV), drug resistance detection (HIV, syphilis, gonorrhea, TB), or disease staging (i.e., acute/recent, longstanding or latent infection).

**Impact and Commercialization Potential:** It is known that early diagnosis and treatment of HIV infection as well as diagnosis and treatment of comorbid pathogens can improve health outcomes for individuals infected with HIV. Furthermore, such testing has the potential for decreasing transmission of HIV and better health outcomes and optimal treatment for the comorbid pathogens. Estimations show that 1.2 million people living in the United States are living with HIV infections, and out of those, 1 in 8 are unaware of their infection. Faster turnaround times, lower cost, and more efficient detection would be highly impactful for these individuals, their partners and the community. Diagnostic reagents, tests or testing platforms, that will further improve diagnosis or monitoring of HIV or comorbid pathogens such as hepatitis (B and C), syphilis, gonorrhea or TB, would be in great demand by the health-care and public health systems as well as other sectors engaged in using diagnostics to treat this patient population.

Visit the NCHHSTP homepage for more information on NCHHSTP’s research program areas [http://www.cdc.gov/nchhstp/](http://www.cdc.gov/nchhstp/)

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NATIONAL CENTER FOR IMMUNIZATION AND RESPIRATORY DISEASES (NCIRD)

The mission of the National Center for Immunization and Respiratory Diseases (NCIRD) is the prevention of disease, disability, and death through immunization and by control of respiratory and related diseases. NCIRD balances its efforts in the domestic and global arenas as well as accommodates the specific needs of all populations at risk of vaccine preventable diseases from children to older adults.

Please visit their web site at: http://www.cdc.gov/ncird/

(11) Prevention and Diagnosis of Acute Respiratory Infections in the US and Globally

**Background:** Acute respiratory infections kill an estimated 3.9 million people annually and in developing countries are the leading cause of mortality in children under 5 years of age. Specific respiratory virus infections such as influenza and respiratory syncytial virus, are major contributors to this burden of disease, as are other respiratory bacterial and viral pathogens. Respiratory virus infections are frequent events in all age groups and impose a substantial burden on social and healthcare delivery systems.

**Specific Research Areas of Interest:** The goal of this research includes, but is not limited to activities that support the development and evaluation of tools for: 1) the prevention of acute respiratory infections such as pneumonia, influenza, and Legionnaire’s disease; 2) rapid recognition and containment of outbreaks; and 3) advanced diagnostic technologies including point-of-care testing, advanced molecular detection and whole genome sequencing.

**Impact and Commercialization Potential:** This research will lead to the development of practical solutions for the prevention and diagnosis of vaccine preventable diseases that have a substantial impact on the economy, health and wellbeing of society. The goal of the research supported through this mechanism is expected to begin shifting viral and bacterial infections from common occurrences to rare events. The innovative technologies and solutions developed through this process will make it possible to improve the public health and healthcare system’s response in a variety of settings, thus making the commercialization potential unlimited.

Visit the NCIRD homepage for more information on NCIRD’s research program areas http://www.cdc.gov/ncird/

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NATIONAL CENTER FOR INJURY PREVENTION AND CONTROL (NCIPC)

For more than 20 years, CDC's National Center for Injury Prevention and Control (the Injury Center) has helped protect Americans from injuries and violence. We are the nation’s leading authority on injury and violence. We study violence and injuries and research the best ways to prevent them, applying science and creating real-world solutions to keep people safe, healthy, and productive. NCIPC will prioritize funding meritorious applications that address the NCIPC program topics listed in this program announcement. NCIPC may also consider meritorious applications that address current NCIPC research priorities. To learn more about NCIPC research priorities, please visit our web site at: https://www.cdc.gov/injury/researchpriorities/index.html

Please visit their web site at: http://www.cdc.gov/injury/index.html

(12) Technological Innovations to Reduce Deaths and Injuries from Motor Vehicle Crashes

Background: Motor vehicle crashes are a leading cause of death among those aged 1-54 years in the United States, killing over 37,000 people every year and injuring over 3 million more. The cost of medical care and productivity losses associated with motor vehicle-related injuries in the United States can exceed $63 billion annually. In 2017, 10,874 people died in alcohol-impaired driving crashes. Among passenger vehicle occupants killed in 2017, almost half (47%) were known to be unrestrained (i.e., not buckled up) at the time of the crash. Speed is a known risk factor for injury and death in a crash. In 2017, there were 9,717 speeding-related deaths. In 2017, there were 3,166 deaths in distraction-affected crashes (8.5% of the total fatalities).

Motor vehicle crashes can result from a single or combination of environmental, human behavioral, and vehicle-related risk factors including hazardous road conditions, driving too fast for the environment, driver perception deficits, non-compliance with vehicle safety devices, impaired driving, lack of seat belt use, lack of helmet use, distracted or drowsy driving, and sub-optimal vehicle performance. Reducing any of these risk factors can lower the likelihood of a crash and increase the chance of survival in the event of a crash.

Adaptive technologies can generate feedback loops about the road and environment, driver fitness, and vehicle performance, and applications of these adaptive technologies in both private and commercial vehicles can reduce risks associated with motor vehicle crashes. Currently, there are a limited number of adaptive technologies to warn drivers of potential dangers associated with driving; most of these technologies focus on vehicle related performance (e.g., collision warning, electronic stability control, lane departure warning, adaptive headlights, etc.). Innovative adaptive technologies that can assist in alerting drivers to risks associated with the road or environment and vehicle performance, and that can facilitate drivers to modify personal risk behavior, including impaired or drowsy driving, are sought. These adaptive technologies can result in the development of tools or systems that will reduce the likelihood and severity of motor vehicle crashes and assist drivers in making potentially life-saving decisions more quickly and more intuitively.
**Specific Research Area of Interest:** The goal of this project is the development of improved technologies that have the potential to further reduce motor vehicle crashes and resulting injuries. These technologies will address risks such as distracted driving, impaired driving, non-compliance with use of vehicle safety equipment (e.g., seat belts), environmental conditions (including road quality), vehicle performance and other factors that may impact driving. A successful awardee would be one that would be able to develop and pilot test new technology designed to reduce motor vehicle crashes and resulting injuries.

**Impact and Commercialization Potential:** The availability of technologies to reduce motor vehicle crashes and resulting injuries can reduce the related public health burden and save lives. Development of these technologies has commercial viability. Commercial applications of this technology might be of interest to drivers, motor vehicle manufacturers, insurance companies, patients, clinicians, and health systems.

(13) Innovative Technology or Media to Prevent Violence Affecting Children/youth

**Background:** Violence is a significant public health problem in the United States. In 2018, more than 67,000 people in the United States died as a result of violence. Nearly 19,000 people died from homicide and more than 48,000 died from suicide. Far more people experienced nonfatal violence. For example, more than 1.5 million people were treated for nonfatal injuries from assaults and nearly 500,000 people were treated for self-harm injuries in U.S. emergency departments in 2018. Violence starts early in life. In 2018, there were an estimate 4.3 million referrals to child protective services for child abuse or neglect involving an estimated 7.8 million children. Exposure to violence in childhood and adolescence can increase risk for later violent experiences and can have a cumulative and compounding impact on health and well-being across the lifespan. Adverse Childhood Experiences (ACEs) are a collection of experiences that may be traumatic to children and youth and include exposure to violence and household challenges (e.g., parental mental illness, separation/divorce, substance abuse, and incarceration) that occur during the first 18 years of life. The different forms of violence impacting children and youth, include child abuse and neglect, sexual violence, teen dating violence, youth peer violence, youth/parent suicidal behavior, and exposure to adult intimate partner violence. These forms of violence often share common risk and protective factors. These factors can start in early childhood and continue throughout the lifespan. They go beyond individual-level factors to include family and peer relationships and other influences from schools, the community, and society. Many effective violence prevention strategies have been developed, and broader benefits could be achieved from wider dissemination through innovative media and communication technology (e.g., mobile applications, social media, games, Internet-based interventions). To help communities make decisions about violence prevention strategies, the CDC has released a series of technical packages to help states and communities take advantage of the best available evidence (see [https://www.cdc.gov/violenceprevention/pub/technical-packages.html](https://www.cdc.gov/violenceprevention/pub/technical-packages.html)). Media and communication technology also create the opportunity for the development of new prevention approaches based on what is known about violence risk and protective factors and strategies that work in traditional settings.

The focus of CDC’s violence prevention work includes an emphasis on: 1. children and youth to achieve long-term impacts, 2. populations and communities at highest risk for experiencing or perpetrating violence, 3. shared risk and protective factors that are most important for reducing multiple forms of violence, and 4. facilitating the identification, implementation, and scale-up of approaches that have cross-cutting impact (see [https://www.cdc.gov/violenceprevention/pdf/Strategic_Vision.pdf](https://www.cdc.gov/violenceprevention/pdf/Strategic_Vision.pdf)). Applicants are encouraged to focus on subgroups and communities at greatest risk and to consider the underlying environmental and social contexts that contribute to inequities in risk across racial/ethnic and other population groups.

By focusing on activities that prevent multiple forms of violence, communities can achieve the greatest impact and increase scalability of their prevention strategies. Additionally, these prevention efforts are ideally designed to use resources more effectively and to better address disparities by focusing on the
populations at greatest risk. Innovative media and communication technology can play an important role in effectively reaching populations at greatest risk for multiple forms of violence. Research is needed to guide the development of technological applications for prevention strategies that can effectively protect children and youth who are most at risk for experiencing multiple forms of violence.

**Specific Research Areas of Interest:** The goal of the project is to develop innovative technology or media, such as applications for mobile devices, social media, games, or Internet-based interventions to prevent multiple forms of violence and other ACEs affecting children or youth. Multiple topics in the violence prevention sections of the NCIPC research priorities emphasize the need for research on the innovative use of new media and communication technology (see [https://www.cdc.gov/injury/researchpriorities/index.html](https://www.cdc.gov/injury/researchpriorities/index.html)). For example, the cross-cutting violence prevention research priorities describe specific gaps related to understanding the potential of new media and communication technology to do the following: increase the accessibility of prevention approaches, modify norms about violence and bystander behavior, enhance education and support for young children and their families, reduce stigma and barriers to help seeking, and enhance young people’s skills and relationships to reduce risk for multiple forms of violence. The widespread use of smartphone applications, social media, and wearable technology also provides unique opportunities for the broader dissemination, implementation and evaluation of evidence-base prevention strategies to significantly reduce violence, such as strategies identified in DVP’s technical packages, within real world settings.

The prototype (e.g., developing innovative technology or media) should be informed by prior research about violence risk and protective factors and/or evidence-based prevention strategies and through consultation with subject matter experts in the form(s) of violence and the technology or media selected. The awardee should describe the target audience, the type(s) of violence addressed, the process through which the technology or media is expected to work, goals for the product, the functionality and actions for users to take, the measurements and key performance indicators for tracking progress toward the goals, the estimated costs and logistics of scalability, a description of potential barriers to implementation, and any evidence for the potential benefits from prior research.

**Impact and Commercialization Potential:** The results from this research will have substantial implications for either the creation of innovative prevention approaches or for enhanced opportunities to disseminate existing evidence-based strategies, both of which have the potential to leverage technology to substantially reduce multiple forms of violence and other ACEs. Additionally, this research could give us the ability to better reach vulnerable populations.

Technological or media innovations that show effectiveness in preventing violence affecting children and youth would have the potential for a range of commercial applications. Depending on the nature of the strategy, the target audience, and the costs/logistics of scalability, the product could be in demand by school systems, colleges and universities, youth serving organizations, law enforcement, public health agencies, community groups, parents, and their children.

**(14) Developing Tools for Patients and Clinicians to Address Opioid Overdose**

**Background:** The United States is in the midst of a drug overdose epidemic. In 2018 more than 67,000 people died from drug overdoses and 70% of those deaths involved opioids. Additionally, in 2017, 20% of drug overdoses involved cocaine and nearly three fourths of these deaths involved opioids. Similarly, in 2017, 15% of drug overdoses involved psychostimulants (e.g., methamphetamine), and half of these deaths involved opioids. Given this, there is a need for “linkages to care” technologies that will support the prevention of drug overdose by focusing on people in need of treatment for opioid use disorder or psychostimulant use disorder (including cocaine). “Linkages to care” are the bridges that connect the people across systems. Connections or linkages to care may be viewed as the vehicle by which one system meaningfully coordinates with another. Currently, predominant examples of these linkages to care with respect to the drug overdose crisis come largely in the form of peer navigators and warm hand-offs in emergency settings upon the event of an overdose. Innovative new ideas are encouraged to broaden
opportunities and “think big” about what linkages to care can be supported through technology. Individuals can benefit from technology to assist them in accessing treatment services and supports that are available in their communities. To support awareness for health care providers (including both physical and behavioral health) of evidence-based opioid prescribing practices, and better linkage to treatment for people with opioid use disorder or psychostimulant use disorder, we are also soliciting proposals to create user friendly, helpful electronic tools, apps, or technologies that will help providers recognize the important differences between treatment options for people with opioid use disorder or psychostimulant use disorder.

**Specific Research Areas of Interest:** CDC is interested in research to develop innovative “linkage to care” technologies that will support the prevention of drug overdose by focusing on people in need of treatment for opioid use disorder or psychostimulant use disorder. The technology can focus on important end users: health care providers and people in need of treatment for opioid use disorder or psychostimulant use disorder.

**Clinician Tools:** Opioid prescribing practices can be informed through the implementation of EHR-embedded clinical decision support (CDS) tools and quality improvement (QI) measures tied to recommendations made in the CDC Guideline for Prescribing Opioids for Chronic Pain. CDS are alerts, algorithms, or reminders that pull information from the EHR and bring it to the attention of the provider within clinical workflow (see http://build.fhir.org/ig/cqframework/opioid-cds/). QI measures are intended for health systems and clinics to use as a way to track prescribing rates and provide feedback to clinicians. CDC developed these measures (see https://www.cdc.gov/drugoverdose/prescribing/qi-cc.html) and seeks innovative technologies to make these QI measures readily available for clinician use. The expected research outcome of awards funded within this SBIR program topic is the development of seamless, easy to use technology that will support uptake, tracking, dashboards, and feedback loops for clinician use.

**Patient Tools:** Patient-centered technologies may help people with opioid use disorder or psychostimulant use disorder and those at risk for opioid overdose connect to care in their communities. The expected research outcome of awards funded within this SBIR program topic is the development of a patient-centered technology that will help patients connect to substance use treatment services through the provision of a comprehensive menu of community resources or supports. Potential community resources include linkage to providers offering medication assisted treatment (MAT) for opioid use disorder, behavioral health services, harm reduction services, social services, online support and tangible support such as transportation options.

The awardee is expected to develop and beta test the new technologies with one of the two target audiences (clinicians or patients). Awardee must attend to health information privacy regulations associated with these technologies, such as those that apply to sharing protected health data across systems or any stigma associated with using the technology. The technology must be intuitive and easy to use.

**Impact and Commercialization Potential:** The availability of technological tools to assist clinicians through CDS and/or track their performance on the QI measures can advance guideline-concordant care which will in turn lead to the expected public health benefit of improved patient safety and pain management.

The availability of technological tools to assist those at risk for drug overdose can better ensure that needed referrals, services, and follow-up care are received. By improving access to medical and non-medical support services, the expected public health benefit is increased linkage to support services that can reduce the risk of drug overdose.

Development of technological tools to assist clinicians related to CDS and/or QI has commercial viability. Commercial applications of this technology may be of interest to health systems, clinics, treatment providers, clinicians and other stakeholders invested in preventing drug overdose.
Development of technological tools that links people at risk for drug overdose to support services has commercial viability. Commercial applications of this technology may be of interest to those at risk and their families, community support programs and staff, and other stakeholders invested in preventing drug overdose.

(15) Electronic Tools to Assist Older Adults at Risk for Falls

**Background:** Unintentional falls are the leading cause of fatal and nonfatal injuries in older adults aged 65 years and older. In 2016, falls resulted in more than 3 million emergency department visits and more than 800,000 hospitalizations. About 30,000 older adult deaths occurred because of a fall in 2016, representing an increase of 31% between 2007 and 2016. The older adult population is rapidly growing and is projected to include approximately 74 million Americans aged 65 and older by the year 2030. Given this, a continued increase in the rate of falls could lead to an estimated 59,000 fall-related deaths among those aged 65 and older by 2030. Falls are among the 20 most expensive medical conditions in the US. In 2015, these injuries were estimated to cost nearly $50 billion dollars, and the amount is projected to climb with the anticipated increase in the older adult population.

Falls often lead to reduced mobility and loss of independence; therefore, reducing fall risk is conducive to maintaining independence. An example of a resource for information about falls prevention is CDC’s Stopping Elderly Accidents, Deaths, and Injuries (STEADI) initiative. STEADI includes information on modifiable fall risk factors and evidence-based interventions. Evidenced-based interventions include referral to physical therapy or community-based exercise programs (Tai-chi and Stepping On program); removal of home hazards; stopping, switching, or reducing the use of medications that increase fall risk; increasing vitamin D if deficient; and improving vision impairment. Older adults prescribed with at least one of these fall prevention strategies are 40% less likely to be hospitalized due to a fall. They are more likely to follow these recommendations when advised by their healthcare provider and when receiving reminders and continuous ongoing support and monitoring. Older adults are also more likely to follow recommended fall prevention activities when the activities are affordable and easily accessible. Barriers to following through with fall prevention recommendations include the inability to remember instructions and lack of information about fall prevention programs.

**Specific Research Areas of Interest:** The goal of this research is to develop technology that will help older adults follow their healthcare provider’s recommendations for reducing their fall risk. One way this can be accomplished is through technology that incorporates elements of an evidence-based program, such as those described in CDC’s STEADI tools and resources. For example, the technology could use STEADI’s patient education materials to educate older adults about falls and how to prevent them. The technology could screen older adults for fall risk (e.g., by using the Stay Independent 12-item survey). The technology could also record the medications used by the older adult and allow for development of an individualized fall prevention care plan as recommended by the older adult’s healthcare provider. The technology could then periodically remind the older adult to follow through with the care plan. The technology could prompt the older adult to attend their physical therapy session or remind them to schedule an eye exam, as well as locate local fall prevention programs (e.g., Tai Chi program). Other potential functions include identifying local pharmacies to review the older adult’s medications for fall risk and supporting communication with their healthcare provider either in real-time or via a summary format for review at provider visits. The awardee is expected develop and beta test the technology. The technology should incorporate applicable healthcare information privacy regulations and be intuitive and easy to use.

**Impact and Commercialization Potential:** The availability of a technological tool to assist older adults who are at risk for falls can better ensure that provider recommendations are followed, and that needed referrals, services, and follow-up care are received. By facilitating fidelity to recommendations and improving communication about fall risk and prevention between patients and providers, the expected public health benefit is increased linkage to support services that can reduce the risk of older adult falls. Development of a technological tool that reduces risk for older adult falls has commercial viability. Commercial applications of this technology may be of interest to those at risk and their families,
healthcare providers, community support staff, and other stakeholders invested in preventing older adult falls.

(16) Prevention and Management of Traumatic Brain Injury Among Youth

**Background:** As attention to concussion and traumatic brain injury (TBI) has grown in recent years, there has been an increase in the number of pediatric patients with concussion (also known as mild TBI, or mTBI) seen in healthcare settings including emergency departments, urgent care settings and primary care practices. Many healthcare providers have insufficient time and training to systematically assess and manage patients with suspected mTBI, thus limiting adoption of best practices to ensure diagnosis and management consistency.

Falls are the leading cause of non-fatal emergency department (ED) visits among children aged birth to 14 years, accounting for 2.4 million visits annually, and the leading cause of TBI ED visits for children in the 0 to 4 year age group. Studies examining injury circumstances and location of the injury in younger children report that although almost twice as many children are hospitalized due to falls from furniture than from stairs, children who fall from stairs are more likely to sustain head injuries. Previous reports on injuries in the youngest children indicate many injuries are related to nursery and infant products (e.g., cribs); furniture; stairs; and surfaces (e.g., carpet, tile floor).

Children with TBI navigate two systems of care - health systems and school systems. There are challenges, inconsistencies, and gaps in current systems of care for children with TBI, particularly for children who are transitioning to school after acute care. Communication of medical information to parents and school personnel is inconsistent, which contributes to gaps in care and potentially poorer health, education, and social outcomes for children. Following TBI diagnosis, it is especially important for children and youth to monitor their symptoms (e.g., fatigue, headaches, changes in thinking) when they return to school and sports and recreational activities. Often healthcare providers do not receive follow-up information once the child has returned to school and activities, a critical component of follow-up care.

**Specific Research Areas of Interest:** CDC is interested in research to develop technology that can assist caregivers with prevention of traumatic brain injuries (TBIs) among young children. The technology could support caregivers of young children in making their home safer with respect to TBI among young children by helping them to evaluate opportunities for increased safety in their homes and implementing safe-home strategies. We are also interested in technology that can aid adolescents and young adults with management of TBIs. The technology could support adolescents and young adults who have experienced a TBI by helping them to understand, record and communicate their symptoms in the acute phase as well as after the initial diagnosis. Knowledge of persistent symptoms following diagnosis would help them inform healthcare providers about their health and well-being.

Goal 1: Provide injury prevention information for parents of young children, including information on environmental structuring.

Goal 2: Improve understanding of TBI among adolescents and young adults and help them to monitor and communicate about their symptoms in order to improve post injury management.

**Impact and Commercialization Potential:** The availability of technology that can assist caregivers in making their home safer with respect to traumatic brain injury among young children can help to reduce the burden of traumatic brain injury in this population. The technology would also help adolescents and young adults understand, track and manage their traumatic brain injury symptoms, which would aid in older children recovering from traumatic brain injury as well as potentially prevent them from experiencing future injuries. Development of this technology has commercial viability. Commercial applications of this technology may be of interest to adolescents, young adults, caregivers, healthcare providers and youth sports personnel.
Visit the NCIPC homepage for more information on NCIPC’s research program areas
http://www.cdc.gov/injury/index.html

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NATIONAL INSTITUTE FOR OCCUPATIONAL SAFETY AND HEALTH (NIOSH)

“The National Institute for Occupational Safety and Health (NIOSH) is part of the U.S. Centers for Disease Control and Prevention. It has the mandate to assure “every man and woman in the Nation safe and healthful working conditions and to preserve our human resources.” NIOSH has more than 1,300 employees from a diverse set of fields including epidemiology, medicine, nursing, industrial hygiene, safety, psychology, chemistry, statistics, economics, and many branches of engineering. NIOSH works closely with the Occupational Safety and Health Administration (OSHA) and the Mine Safety and Health Administration in the U.S. Department of Labor to protect American workers.

NIOSH is particularly interested in applications that address current needs in the national COVID-19 response. These needs cover a wide range of NIOSH-related topics including personal protective equipment/technologies, exposure assessment, engineering controls, and emergency preparedness and response. These needs also cover many, if not most, occupational environments.

For additional information about NIOSH, please visit their web site at: http://www.cdc.gov/niosh/programs.
(17) Control Technology and Personal Protective Equipment for High Risk Occupations

**Background:** Personal protective equipment (PPE) protects workers from death and disabling injuries and illnesses as well as from the specific threats of exposures to certain airborne biological particles, chemical agents, nanomaterials, splashes, noise exposures, fall hazards, head hazards, and fires. It is estimated that 20 million workers use PPE on a regular basis to protect them from job hazards and a total of 135,000 workers potentially could benefit from the use of PPE (Worker Health Chartbook 2004). Engineering controls include substitution of a safe material for a hazardous one, design changes to equipment, or modification of work methods to eliminate or reduce hazards. Research is needed to develop and evaluate control strategies and personal protective equipment for specific hazards and to assure their practicality and usability in workplaces in all of the high risk industrial sectors.


**Examples of specific research areas of interest include, but are not limited to:**

- Conduct research on the ability of existing containment and control strategies to prevent releases and potential human exposures to engineered nanomaterials.

- Conduct research to evaluate the effectiveness of personal protective equipment in protecting workers against exposure to engineered nanomaterials. Provide data to fill knowledge gaps and support guidance for the selection and use of gloves and protective garments to prevent exposures. Respiratory protection research needs to be extended to a broad range of engineered nanomaterials.

- Develop a heads-up display coupled with a personal noise exposure monitoring system. Personal noise alert “badges” and personal noise dosimeters exist, but do not have an effective way to alert the user immediately when a noise hazard occurs. A system that displays a warning within the user’s visual field (via lights on protective eyewear, hardhat, etc.) would facilitate hazard recognition.

- Develop an inexpensive hand-held earplug test device based on the NIOSH QuickFit concept. Studies of hearing protector users have shown repeatedly that average protection values are much lower than the labeled Noise Reduction Ratings (NRR) determined in laboratories. A QuickFit test system would help workers determine if their hearing protection is giving them at least 15 decibels of attenuation.

- Develop innovative engineering control approaches and technologies for reducing asphalt exposures in roofing, and skin exposures and disease in construction workers.

- Conduct research to understand PPE integration and interoperability issues. In most cases, individual PPE are currently used without consideration for their ability to function together. Research is needed to test interfaces among different PPE and components. Current interfaces do not provide seamless integration of PPE components resulting in reduced comfort, fit, usability, and protection for the wearer as well as logistical challenges for safety managers and employers.

- Develop innovative educational and professional training materials suitable for today’s diverse workplace on the role of PPE in occupational safety and health. This is especially critical for high risk occupations. Innovative methodologies, including social media, should be explored and evaluated to demonstrate their effectiveness at improving workplace safety and health. For example, to what extent can mobile application media be focused on worker safety and health to provide up-to-date PPE information to a diverse range of employers and employees through portable communication devices?
**Impact and Commercialization Potential:** The impact of the proposed research will prevent work-related injury, illness, and death by advancing the state of knowledge and application of personal protective equipment. Potential products include technical methods, processes, techniques, tools, and materials that support the development and use of personal protective equipment worn by individuals to reduce the effects of their exposure to a hazard.

(18) Exposure Assessment Methods for High Risk Occupations

**Background:** Exposure assessment provides multi-disciplinary strategies and methods to anticipate, recognize, evaluate, control, and confirm effective management of occupational health stressors, exposures to those stressors, and resulting health risks. Major gaps in current approaches include: (1) the lack of practical methods for hazard identification and measurement that can be applied at reasonable cost in many workplaces where health stressors may exist, (2) the lack of validated, noninvasive biological methods for monitoring relevant exposure and resulting dose, and (3) the lack of strategies and methods for epidemiologic studies to demonstrate either a dose-response effect or a conclusion of no association between the agent and disease in the complex environments of today's workplaces.

For additional information about NIOSH Exposure Assessment programs, please visit their web site at: http://www.cdc.gov/niosh/programs/expa/.

**Examples of specific research areas of interest include, but are not limited to:**

- Two areas of research are needed to support effective assessment of worker exposure to engineered nanomaterials. 1) Real-time sensors capable of reliably detecting nanoparticles and providing information on size distribution and count, that can be used for personal monitoring; and 2) Development of methods that can detect and quantify the presence of engineered nanomaterials in samples collected for the purpose of characterizing exposures. These methods need to be cost-effective and available to the OS&H practitioner community. Broader application to general public health assessments should be factored into the research.

- Develop new or improved methods to measure occupational health stressors such as psychological and ergonomic factors, noise, chemicals, particles and fibers, physical agents, non-ionizing radiation, or mixtures of stressors in the work environment. Enhanced measurement performance and functionality can include sensitivity, selectivity, size and weight considerations, ease of use, and capabilities to measure multiple analyses simultaneously.

- Develop or adapt easy-to-use, direct-reading instruments and test kits to rapidly and inexpensively measure exposures in a variety of workplaces. Critical applications include routine monitoring, evaluating the success of control technologies, and supporting epidemiological studies. For example, developing a sound level meter to monitor worker noise exposure that can be used in underground coal mines.

- Improve the measurement of low concentrations of chemicals and biomarkers in biological specimens such as blood, urine, saliva and sweat so that such concentrations can be linked to internal dose at the target organs, work tasks and workers can be categorized according to hazard bands and exposure bands, and at-risk workers can be identified and protected.

- Develop a computerized system that can be used to predict worker noise exposure from mining machine noise emissions. The system would include an acoustic model of mining environments and algorithms to characterize exposure based on noise source characteristics. The main application for this technology would be for mining machine manufacturers to evaluate the potential effects of noise controls during the design process. If the impact of design changes on exposure reduction can be accurately predicted without the need for extensive field measurements, innovative noise controls can reach implementation much more quickly.
Impact and Commercialization Potential: This research will lead to the development of practical solutions and prevention activities to address complex problems that cause occupational diseases, injuries, and fatalities and that will lead to reductions in occupational injuries and illnesses among all workers. This research will lead to the development and translation of exposure assessment methods and research findings into prevention practices and products that will be adopted in occupational settings. Potential products include technical methods, processes, techniques, tools, and materials that support the assessment of exposure to physical, chemical and biological hazards in the work environment.

(18) Work-related Injuries from Motor Vehicle Crashes

Background: The risk of injury associated with on-the-job operation of motor vehicles affects millions of U.S. workers who work in all industries and drive all types of vehicles, and for whom driving may be a primary or incidental job task. Motor vehicle-related incidents are consistently the leading cause of work-related fatalities in the United States. Between 2003 and 2017, the Bureau of Labor Statistics reported 25,704 work-related fatalities due to motor vehicle incidents, about 35% of all fatalities at work. Over the same period, workers incurred nearly 400,000 lost-workday injuries due to these incidents. Crash-related fatalities and serious injuries have a devastating impact on workers and their families, and on the economic health and productivity of American businesses. Work vehicles such as large trucks also have an impact of the safety of the motoring public.

The NIOSH Center for Motor Vehicle Safety coordinates the CDC/NIOSH response to this pressing worker safety issue. Many NIOSH programs include motor vehicle crashes among their top injury prevention priorities: Traumatic Injury; Transportation, Warehousing, and Utilities; Wholesale and Retail Trade; Oil and Gas Extraction; and Public Safety.

Examples of specific research areas of interest include, but are not limited to:

The highest priority is to develop, implement, and evaluate interventions in an effort to build the scientific evidence base to guide prevention of work-related motor vehicle crashes and resulting injuries. This may be achieved by:

- Developing and testing new design concepts and applications with potential for commercialization and diffusion to employers and fleet managers
- Developing and testing novel approaches for driver training and assessment to reduce work-related motor vehicle crashes, including training on the operation of vehicles with Advanced Driver Assistance Systems (ADAS) or other forms of automation
- Developing and evaluating the effectiveness of technology- or management-based intervention strategies to reduce the incidence or severity of work-related motor vehicle crashes
- Developing and evaluating engineering controls for preventing work-related crashes and injuries, with emphasis on specialized work vehicles such as large trucks and fire apparatus
- Developing and evaluating an easy-to-use computerized system based on readily available technology that can automate a “fatigue detection” system capable of warning the employee driver and supervisor when the driver may be at risk for a work-related motor vehicle crash. The system would include a statistical algorithm capable of using Global Positioning System (GPS) data from cellular phones to characterize potential number of hours awake within the last 24 hour-cycle. The main application for this technology would be to allow supervisors and employee drivers to identify and respond to fatigue, thereby reducing the driver’s risk of a fatigue-related crash. If fatigue detection systems can use readily available technology, intelligent automation may help mediate work-related injury prevalence.
Impact and Commercialization Potential: Application of evidence-based interventions is expected to have a large impact on reducing the incidence and severity of work-related motor vehicle crashes. This will yield substantial public health benefits and will positively affect workers’ compensation and health insurance premiums and costs. CDC/NIOSH has well-established working relationships with employers, their trade associations, and standards-setting organizations, and is therefore strongly positioned to communicate findings and guidance to potential users. CDC/NIOSH also has strong infrastructure to facilitate the transfer of technology-based interventions to the marketplace. Given the extremely short induction period between exposure and injury occurrence, CDC can make a measurable difference in a very short period of time (< 4 years).

Visit the NIOSH homepage for more information on NIOSH’s research program areas http://www.cdc.gov/niosh/homepage.html.

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FOOD AND DRUG ADMINISTRATION (FDA)

FDA will accept SBIR grant applications on September 8, 2020; January 5, 2021; and April 5, 2021 submission dates.

The FDA is responsible for protecting the public health by assuring the safety, efficacy, and security of human and veterinary drugs, biological products, medical devices, our nation’s food supply, cosmetics, and products that emit radiation. The FDA is also responsible for advancing the public health by helping to speed innovations that make medicines and foods more effective, safer, and more affordable; and helping the public get accurate, science-based information they need to use medicines and foods to improve their health.

For additional information about areas of interest to the FDA, please visit our home page at http://www.fda.gov.

Limited Amount of Award

FDA will not fund:

- Phase I applications greater than $168,087
- Phase II applications greater than $1,120,586

Research Topics of Interest to FDA

The specific set of topics that are funded by a particular IC/Office/Agency depends on whether the IC/Office/Agency funds/supports Clinical Trials under SBIR/STTR Awards. Use the following table and explanations to determine which list of topics is applicable.

<table>
<thead>
<tr>
<th>Does IC/Office/Agency Fund/Support Clinical Trials under SBIR/STTR Awards?</th>
<th>No*</th>
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*If No,
- This IC/Office/Agency funds only those topics listed under the Non-Clinical Trials Topics section below, UNLESS the IC/Office/Agency funds a Small Business Concern for clinical trials under a NON-SBIR/STTR award.

**If Yes or the IC/Office/Agency funds small businesses for clinical trials under a NON-SBIR/STTR funding opportunity,
- See the table below for which mechanisms the IC/Office/Agency will allow for clinical trials research. Also, see the Clinical Trials Topics section below for a list of funded topics. This IC/Office/Agency also funds research listed under the Non-Clinical Trials Topics section below.
FDA will not accept SBIR applications that propose clinical trials, and all of the topics listed below must be for projects that do not propose clinical trials.

**FDA Non-Clinical Trials Topics:**

**CENTER FOR BIOLOGICS EVALUATION AND RESEARCH (CBER)**

CBER is responsible for ensuring the safety, efficacy, potency and purity of biological and related products intended for use in the treatment, prevention or cure of diseases in humans as well as the safety of the nation's supply of blood and blood products. The primary responsibility of CBER is to review the quality, safety and efficacy of vaccines, blood products, certain diagnostic products and other biological and biotechnology-derived human products.

CBER's activities include: evaluating the quality, safety and effectiveness of biological products before marketing, and monitoring the pre-clinical and clinical testing of new biological products; licensing biological products and manufacturing establishments, including plasmapheresis centers, blood banks, vaccine and biotechnology manufacturers; AIDS program and policy activities, including research on AIDS therapeutic products, diagnostic tests and vaccines; research to establish product standards, develop improved testing methods and assess the safety of biological products; compliance, lot release program and post market surveillance; meeting PDUFA goals, new research programs, and new regulatory initiatives (managed review process for all products).

**CENTER FOR DRUG EVALUATION AND RESEARCH (CDER)**

CDER develops FDA policy with regard to the safety, effectiveness, and labeling of all drugs for human use; evaluates new drug applications and investigational new drug applications; develops standards for
the safety and effectiveness of all over-the-counter drugs; monitors the quality of marketed drugs through product testing (bioavailability/bioequivalence testing), post marketing surveillance, and compliance programs; develops guidelines on good manufacturing practices; conducts research and develops scientific standards on composition, quality, safety, and efficacy of human drugs.

Drug regulatory research as conducted in CDER is directed at the discovery of new knowledge relevant to drug development, post marketing drug experience (patterns of drug use and safety), and drug regulation to enhance FDA regulatory decisions. These drug regulatory decisions impact on the development of regulations, guidelines and guidance for the regulated industry and provide clarity and consistency in application of CDER regulatory requirements. These drug regulatory decisions also impact public health by ensuring that marketing drugs are safe and efficacious and that their risk: benefit profile remains acceptable during the market life of a drug. Specific areas of research conducted by the Center include: Pharmacology/toxicology, microbiology/virology, clinical pharmacology, pediatric issues in drug therapy, post marketing drug safety, evaluation of effectiveness of regulatory actions, patterns of drug use, including off-label, signal detection methodologies (e.g., data mining techniques), epidemiologic studies of therapeutics using population-based data, regulatory compliance, product quality, and active surveillance methods.

Research and development opportunities within the FDA that lend themselves to performance by small businesses include, but are not limited to, the following:

A. Develop a system for gathering real-time data on physician prescribing behavior, understanding and compliance with drug product labeling and frequency of off-label prescribing.

B. Develop and evaluate the effectiveness of new methods and tools for managing the known risks of marketed drug products (e.g., communicating newly identified risks to health care practitioners and patients).

C. Develop methods for timely active surveillance of newly approved drug products in large populations to identify both expected and unexpected outcomes.

D. Develop methods for actively collecting information on all cases of classically drug-associated events (e.g., acute liver failure, blood dyscrasias, severe desquamating skin disorders) to augment the FDA’s current passive surveillance system.

E. Develop improved clinical markers and methods with potential for bedside application for detection of the early onset of adverse drug events.

F. Develop surrogate potency methods for biotech drug products to replace traditional animal testing.

G. Development of psychochemical and in-vitro biological tests to evaluate pharmaceutical equivalence of complex drug substances and drug products.

H. Research into approaches to handle informative missing patient data in clinical trials, including innovations in study designs and statistical methods of analysis.

I. Statistical and computational methods and strategies for the design, analysis and interpretation of microarray, genomic and proteomic data.

**CENTER FOR FOOD SAFETY AND APPLIED NUTRITION (CFSAN)**

The FDA is responsible for the safety of the vast range of food Americans eat; about 80 percent of all food sold in the United States. This includes everything except for the meat, poultry, and processed egg products that are regulated by the USDA. Consequently, CFSAN seeks research designed to complement and accelerate efforts aimed at the detection, prevention, and control of contamination that may be responsible for illness or injury conveyed by foods, colors, and cosmetics. CFSAN conducts research, and develops regulations, guidance and standards related to the composition, quality, nutrition,
and safety of food, food additives, colors, and cosmetics. The Center evaluates FDA’s surveillance and compliance programs relating to foods, colors, and cosmetics; reviews industry petitions, and develops regulations for food standards to permit the safe use of color and food additives.

CFSAN maintains an active research program that is focused on the following priorities; ensuring the safety of food, dietary supplements and cosmetics; improving nutrition; and promoting the security and integrity of the food supply. The Center’s research activities are intended to; support the FDA’s regulatory activities; reduce the incidence of foodborne illness by improving our ability to detect and quantify foodborne pathogens, toxins, and chemicals that could jeopardize the safety and security of the food supply; find new and improved ways to control these agents; and safely produce, process, and handle food and food products. FDA is committed to reducing the incidence of foodborne illness to the greatest extent feasible while at the same time protecting the nation's food supply. Mission-critical knowledge gaps are addressed through translation research focused on the risks associated with FDA regulated products throughout their life cycles, from production to consumption. Ideally extramural research is sought that complements the Center’s intramural research efforts, and which will enhance the Agency’s and the Nation’s ability to reduce the incidence of foodborne illness and protect the integrity of the nation’s food supply. FDA’s mission-critical needs require that the research not simply end with the generation of new knowledge and technologies, but extend to the validation of new approaches by using realistic conditions that accurately reflect the diversity of the food industry and offer potential solutions that can be accept by appropriate sectors of the food industry.

**CENTER FOR DEVICES AND RADILOGICAL HEALTH (CDRH)**

CDRH develops FDA policy and solves problems related to public health and safety of medical devices and radiation-emitting electronic products. It evaluates applications for premarket approval of medical devices, approves products development protocols and exemption requests for investigational devices. It classifies devices into regulatory categories, develops safety effectiveness standards and good manufacturing practices regulations, operates post market surveillance and compliance programs, and provides technical, non-financial assistance to small manufacturers. The Center also conducts programs to reduce human exposure to hazardous ionizing and nonionizing radiation, through an electronic product radiation control program and other programs designed to control and limit radiation exposure. The Center develops and conducts research and testing programs in the areas of physical, life, and engineering sciences related to the human health effects of radiation and medical device technologies, provides expertise and analyses for health-risk assessments, and also develops new or improved measurement methods, techniques, instruments and analytical procedures for evaluating product performance and reliability.

Research and development opportunities within the FDA that lend themselves to performance by small businesses include, but are not limited to, the following:

A. Examine the setup, documentation and optimization of our Sun Grid Engine (SGE). The architecture of this networking application is particularly suited to managing surge capacity in high performance computing. The modeling of many physiologic functions and bioinformatic analyses can take months or even years to run on a standard desktop computer. The SGE takes the overall problem and distributes it to a cloud of computers on a network so that no user knows, or cares, if a computation is performing in the background on their machine. As FDA rolls out laptops with multi-core CPU’s and which are equipped with prodigious amounts memory this experiment in "cloud computing" could become a reality on the Whiteoak Campus. The scope of work would be to develop, document, and provide training systems for developers, network architects, and users on working methodologies for the integration of cloud computing with the existing FISMA compliant conventional networking.

B. Develop a high-speed, low light spectral CMOS linear imaging system to measure complete spectra of multiple variables from living tissue. Complete spectra of fluorescence signals (including auto-
fluorescence and FRET) could be measured along a line at high speeds (10 kHz) with a rectangular CMOS grid (e.g. 10 x 1,000 pixels -> 10 sites 1000 wavelengths).

C. Develop bioassays/biosensors to identify injurious levels of nerve stimulation utilizing bioluminescence and neurotransmitter detection technologies. Research capabilities needed include voltage clamp, current clamp and extracellular techniques in peripheral nerves and brain slices to explore stimulation protocols that release neuroactive substances released in injury and inflammation which are not normally evoked under normal physiological conditions.

D. Design, build, and validate a phantom that is traceable to a national metrology institute (NMI) such as NIST (or any other NMI) to improve the accuracy and clinical utility of bone mineral density measurements made using dual energy X-ray absorptiometry (DXA). The calibration phantom should be constructed using biosurrogate materials with known/tabulated data for body tissue and tissue substitutes.

**CENTER FOR VETERINARY MEDICINE (CVM)**

CVM is a public health organization that enables the marketing of effective drugs, food additives, feed ingredients, and animal devices that are safe to animals, humans, and the environment. The Center, in partnership with Federal and state agencies and other customers, ensures animal health and the safety of food derived from animals. The Center makes timely, quality decisions and takes regulatory actions to ensure that these products provide for quality health care of animals, minimize the transmission of zoonotic diseases, and increase the efficiency of production of animal-derived food and fiber. Regulatory decisions are supported by research, the monitoring of product safety, and efficacy, and the continual improvement of processes.

Research and development opportunities within the Center for Veterinary Medicine that lend themselves to performance by small businesses include, but are not limited to, the following areas of interest:

A. Development, for the specific purpose of obtaining approval or conditional approval, of products for the treatment, control or prevention of diseases or conditions occurring in minor species or small numbers of major species.

B. Development and validation of high throughput/screening quantitative and qualitative analytical methods for analyzing drugs, additives, and contaminants in animal tissues and feeds.

C. Development of methods to determine absorption, distribution, metabolism, and excretion of drugs, feed additives and contaminants (microbial and chemical) in food animals, including minor species.

D. Development of new biomarkers and models for determining the safety and effectiveness of veterinary drugs and food additives in domestic animals, including minor species.

E. Development of methods to determine the effects of drugs, food additives, and contaminants (microbial and chemical) on immunological and physiological functions of domestic animals, including minor species.

**OFFICE OF CRITICAL PATH PROGRAMS**

The Office of Critical Path Programs, in FDA's Office of the Chief Scientist, coordinates the cross-agency Critical Path Initiative (CPI), FDA's strategy for transforming the way medical products are developed, evaluated, and manufactured. CPI activities are under way throughout the Agency, from the product centers to the Office of the Commissioner. For details, see [http://www.fda.gov/ScienceResearch/SpecialTopics/CriticalPathInitiative/default.htm](http://www.fda.gov/ScienceResearch/SpecialTopics/CriticalPathInitiative/default.htm). Collaboration is key to the CPI initiative because bringing safe, effective, and innovative therapies to the American public requires FDA to leverage the resources and expertise of all stakeholders, including other Federal
agencies, academia, healthcare professionals, patient and consumer groups, regulated industry, and health-related organizations. In 2008, CPI collaborations involved 84 government agencies, universities, industry leaders, and patient groups from 28 states and 5 countries on a raft of groundbreaking research projects.

Research and development opportunities within FDA that lend themselves to performance by grantees include, but are not limited to, the following:

A. Studying the immunological correlates of TB immunity and developing tools to evaluate TB vaccine efficacy.
B. Developing study models for testing combination-antimicrobials as a strategy to prevent the development of drug resistance.
C. Developing new approaches to preclinical safety testing.
D. Identifying biomarkers for safety and efficacy evaluation of medical products.

OFFICE OF ORPHAN PRODUCTS DEVELOPMENT

The Office of Orphan Products Development was established to identify and facilitate the development of orphan products. Orphan products are drugs, biologics, medical devices and foods for medical purposes, which are indicated for a rare disease or condition (i.e., one affecting fewer than 200,000 people in the United States). These products may be useful in a rare disease/disorder but lack commercial sponsorship because they are not considered commercially attractive for marketing. A subcategory of orphan products are those marketed products in which there is evidence suggesting usefulness in a rare disease/disorder but which are not labeled for that disease/disorder because substantial evidence of safety and effectiveness for that use is lacking.

Research and development opportunities within the FDA that lend themselves to performance by small businesses include, but are not limited to, the following:

A. Development of products for the treatment of rare diseases or disorders including but not limited to neurological, metabolic, genetic, ophthalmologic, hematologic, and dermatological diseases or disorders for the specific purpose of obtaining marketing licensure.
B. Development of products for use in diagnosis of rare diseases for which the diagnostic tool would be used in fewer than 200,000 persons annually in the United States.
C. Development of vaccines for the prevention of rare diseases to be used in fewer than 200,000 persons annually in the United States.

For additional information on research topics and administrative and business information, contact:

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Grants Management Specialist
Grants and Assistance Agreements Team
240-402-3099, Fax: 301-827-0505
Email: Kiara.Fowler@fda.hhs.gov
Food and Drug Administration
Division of Acquisition Support and Grants
5630 Fishers Lane - HFA 500
Rockville, MD 20857
APPENDIX A: NATIONAL INSTITUTES OF HEALTH SBA-APPROVED SBIR/STTR TOPICS FOR AWARDS OVER STATUTORY BUDGET LIMITATIONS

National Institutes of Health SBA-Approved SBIR/STTR Topics for Awards over Statutory Budget Limitations

07/09/2020

NIH has received approval from SBA for the topics listed within for budgets greater than $256,580 for Phase I SBIR/STTR awards and greater than $1,710,531 for Phase II SBIR/STTR awards for 2020-2021. Applicants are strongly encouraged to contact NIH program officials prior to submitting any award budget in excess of these amounts. Applicants are also required to follow NIH Institute- and Center-specific budget guidance found in all SBIR and STTR funding opportunity announcements.
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**Division of Program Coordination, Planning, and Strategic Initiatives (DPCPSI), Office of Research Infrastructure Programs (ORIP)**

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NATIONAL CANCER INSTITUTE (NCI)

A. Therapeutics (e.g. Small Molecules, Biologics, Radiomodulators, Cell-based Therapies and Drug Development-Related IT Tool and Algorithm Development)

B. *In Vitro* and *In Vivo* Diagnostics (e.g. Companion Diagnostics, Prognostic Technologies, Treatment Monitoring and Diagnostic-Related IT Tool and Algorithm Development)

C. Imaging Technologies (e.g. Agents, Devices, Software Tools, Algorithm Development, and Image-Guided Interventions)

D. Devices for Cancer Therapy (e.g. Interventional Devices, Software Tools, Algorithm Development, Surgical, Radiation and Ablative Therapies)

E. Agents for Cancer Prevention (e.g., Vaccines, but not “Technologies for Cancer Prevention”)

F. Development of Low-Cost Technologies for Low-Resource Settings and Cancer Global Health

G. Development of Digital Health Tools
NATIONAL CENTER FOR ADVANCING TRANSLATIONAL SCIENCES (NCATS)

A. Innovative platforms for identification and prioritization of targets for therapeutic intervention with clear clinical impact.

B. Technologies to determine alternative uses for existing therapeutic interventions.

C. Tools and technologies to allow assaying of activities of compounds on currently “non-druggable” targets.

D. Phenotypic assay development, including stem cell technology platforms for human “disease in a dish” applications and the evaluation of toxicity.

E. Co-crystallization high-throughput screening techniques.

F. Small molecule and biologics analytical characterization.

G. Tools and technologies that increase the predictivity or efficiency of medicinal chemistry, biologic, or other intervention optimization.

H. Accelerate bioengineering approaches to the development and clinical application of biomedical materials, devices, therapeutics, and/or diagnostics.

I. Tools and technologies that increase the efficiency of human subjects research, including development of technologies that facilitate rapid diagnosis and/or clinical trial recruitment and subject tracking, IRB evaluation, and/or regulatory processes.

J. Novel platforms, technologies and tools for: (1) enabling clinical and translational research, particularly those with mechanisms for inclusion of patient reported data and (2) integration of patient data collected from multiple devices and multiple/diverse clinical studies.

K. Searchable access to information about researchers and their expertise, including but not limited to their publications, published data sets, methods, patents, clinical trials, partnerships, collaborators, and clinical specialty/expertise (if applicable).

L. Tools for meaningful sharing of research data with low barrier for provision and user friendly access.

M. Microphysiological Systems (MPS)/Tissue Chips.
NATIONAL CENTER FOR COMPLEMENTARY AND INTEGRATIVE HEALTH (NCCIH)

A. Development and validation of biomarkers that correlate with efficacy of complementary health approaches.

B. Formulation, development, and clinical testing of complementary health approaches and natural products that would permit FDA approval of a natural product for a specific indication.

C. Identification and validation of biological targets associated with complementary health approaches.

D. Development of innovative technologies and methods to assess natural product–drug interactions in humans.

E. Studies of the mechanistic effects of mind and body interventions that will allow for optimization of the efficacy and safety of the mind and body approach for commercialization.

F. Nontraditional phenotypic assay development for complex natural product mixtures.

G. Integrated in silico tools for exploiting natural product bioactivity.

H. Development and clinical testing of innovative technologies and methods for mind and body approaches. Examples include the use of mobile health technologies such as smartphone apps, sensors, online delivery, or phone-based delivery.

I. Design, development, evaluation, and validation of devices or systems related to complementary health approaches.

J. Utilizing biosynthetic engineering approaches to improve production of natural products in either native or heterologous hosts.
NATIONAL EYE INSTITUTE (NEI)

General Research and Development Topics

A. New or improved ophthalmic or surgical instruments for diagnosis and treatment of eye disorders.
B. Drug delivery systems; gene therapy, cell-based therapy or regenerative medicine.

Retinal Diseases

A. New therapeutic approaches for inflammatory and degenerative diseases and for inhibition of abnormal angiogenesis in the retina and choroid.

Corneal Diseases

A. New therapeutic approaches, artificial corneas, and drug delivery methods for the treatment of corneal injury, infection, dry eye, ocular pain, and other ocular surface disorders.

Lens and Cataract

A. New approaches in the management of cataracts.

Glaucoma and Optic Neuropathies

A. New therapeutic agents for treatment of glaucoma.

Visual Impairment and Blindness

A. New or improved devices, systems, or programs that meet the rehabilitative and everyday living needs of blind or visually-impaired persons.
**NATIONAL HEART, LUNG, AND BLOOD INSTITUTE (NHLBI)**

A. Biomedical technologies (medical devices, instruments, pharmaceuticals, drugs, therapeutics, vaccines, diagnostics and biologics) for heart, lung, blood, and sleep related diseases and disorders requiring Federal regulatory approval (FDA) or clearance to be commercialized.

B. Small and large animal testing of products of tissue engineering and regenerative medicine, drugs, medical devices, therapeutics, and biologics and studies involving *in vivo* animal experiments for heart, lung, blood, and sleep related diseases and disorders.

C. Clinical trials and other experiments involving human subjects for heart, lung, blood, and sleep related diseases and disorders.

D. Therapeutics (drugs, devices, or biologics) development for heart, lung, blood, and sleep related diseases and disorders.

E. Device development for heart, lung, blood, and sleep related diseases and disorders.

F. Diagnostics development for heart, lung, blood, and sleep related diseases and disorders.

G. Investigation of biomarkers and biosignatures of heart, lung, blood, and sleep related diseases and disorders.

H. Technologies to enhance clinical research for heart, lung, blood, and sleep related diseases and disorders.

I. Advanced instrumentation and high throughput tools for biomedical research in heart, lung, blood, and sleep related diseases and disorders.

J. Tools and platforms to improve the dissemination and implementation of evidence-based interventions for heart, lung, blood, and sleep related diseases and disorders.
NATIONAL HUMAN GENOME RESEARCH INSTITUTE (NHGRI)

A. Development of novel or significant improvements for nucleic acid sequencing technology.

B. Development of novel or significant improvements for functional genomics technology.

C. Genomics tools ranging from new instruments to sophisticated molecular biology kits.

D. Bioinformatics software for genomic, genetic and sequence data analysis, functional genomics, associations between genomic data and diseases or phenotypes, interpretation of variants, and genomic data integration.

E. Databases and data management for genomics research and application including sequences, functional data, annotation of variants, and phenotypes.

F. Incorporating genomic results into electronic medical records.

G. Informatics tools that assist in delivering genomic medicine to patients.

H. Development and application of methods for machine learning, pattern detection, and knowledge networks for genomics and translation to genomic medicine.

I. Informatics methods and platforms to enhance privacy, data standards, and data exchange in genomics and translation to genomic medicine.

J. Use of cloud and other computing models to improve scale, reproducibility, interoperability, cost-effectiveness, and utility of genomic and clinical data in genomics and translation to genomic medicine.

K. Single cell genomic analysis.
NATIONAL INSTITUTE ON AGING (NIA)

Division of Neuroscience (DN)

The Division of Neuroscience (DN) fosters and supports extramural research and training to further the understanding of the dementias of old age, as well as neural and behavioral processes associated with the normally aging brain. An area of special emphasis is brain-behavior relationships. An important component of this Division is the support of basic, clinical, and epidemiological studies of AD and related dementias of aging.

A. Development of new and/or validation of existing sensitive, specific, and standardized tests for diagnostic screening and the development of biomarkers.

B. Target discovery and validation through the application of systems biology and systems pharmacology approaches of MCI, AD, ADRD, or other dysfunctions of the central nervous system.

C. Preclinical and/or clinical discovery, development, and/or evaluation of drug, nutritional, behavioral, cognitive, sensory, environmental, or other types of interventions to remediate age-related cognitive decline and/or other dysfunctions of the central nervous system.

D. Development of new technologies for in-home use or for coordination or delivery of services to sustain in-home living for individuals with MCI, AD, ADRD, or that may prolong functional independence when there are dysfunctions of the central nervous system.

E. Development of biosensors and prosthetic devices, technologies, and related software development to aid in the assessment, diagnosis, and remediation of age-related cognitive decline and/or other dysfunctions of the central nervous system.

F. Development of technology and analysis tools to examine genetic, epigenetic, transcriptomic, proteomic, metabolomic, and cell stress pathways in neurons and glia of the aging and AD/ADRD brain.

G. Improvement of technology to analyze structural and functional brain connectivity at the cell, neural circuitry, and global-network levels to define the normal trajectory of brain structure and function over the adult lifespan.

H. Development of technology, including non-invasive methods and novel probes, to monitor and manipulate the plasticity of neural circuits in the adult and aged nervous system.

I. Development of novel markers of neural stem cell function and novel approaches for analysis of next-generation sequence data.

Division of Aging Biology (DAB)

The Division of Aging Biology promotes and supports research and training on the molecular, cellular, genetic, and physiological mechanisms underlying normal aging and age-related pathologies. The objective of DAB-funded research is to elucidate the basic biochemical, genetic, and physiological mechanisms underlying the process of aging and age-related changes in humans and in animal models of human aging. This includes investigations of the gradual or programmed alterations of structure and function that characterize normal aging and investigations of how these adverse changes become risk factors for, or accompany, age-related conditions and disease states.
A. Development of interventions to 1) reduce oxidative or other stresses and aging-related diseases; 2) improve the immune response to foreign molecules or reduce the response to self or suppress age-induced inflammation; 3) enhance longevity or slow aging and may affect other age-related conditions or diseases; 4) reduce the level of damage to nucleic acids, proteins and lipids and the macromolecular complexes formed from these molecules, and improve the damage surveillance and repair potential of cells.

B. Development of minimally-perturbing techniques for collecting blood old non-human animals and the development of non-invasive research and test methods for use in non-human animals.

C. Development of novel strategies for treating age-related renal, pulmonary, urology, reproductive disorders, and age-related changes in hormone production and function.

D. Development of novel methodology for treating osteoarthritis, including devices, processes and pharmacological agents with the potential to: (1) slow the rate of joint deterioration, (2) promote the remodeling of damaged joints, (3) reduce the likelihood of progression to osteoarthritis, and/or (4) improve outcomes for patients with active osteoarthritis.

E. Development of tools and resources in genetics and genomics to study molecular mechanisms of normal aging or aging-related diseases.

F. Development of novel methodology for treating chronic wound healing, including devices, processes and pharmacological agents with the potential to: (1) improve the rate and or quality of wound healing, and/or (2) improve outcomes for patients with chronic wounds.

G. Development of biomarkers for prognosis, diagnosis, or treatment monitoring of aging or aging-related diseases; and analysis or integration of large data sets for developing biomarkers or biomarker signatures of aging or age-related diseases.

**Division of Behavioral and Social Science (DBSR)**

A. Behavioral-economics approaches (incentives or disincentives) to motivate sustainable behavior change to improve health and well-being.

B. Social, behavioral, environmental and or/technical interventions (including robotics) on the individual, institutional, family, or community level to promote older adult independence, increase well-being and prevent disease and/or disability (across home, work and institutional settings), including age-related cognitive impairment.

C. Evidence-based methods, technologies, and interventions to reduce the burden of caregiving for persons with Alzheimer’s disease and AD-related dementia, including training materials/resources appropriate for use by informal caregivers, or professional caregivers within health-care systems or community-based organizations.

D. New sampling and data-collection methodologies for use in large population-based household surveys and behavioral interventions of relevance to aging, including blood-spot technology for biological data collection, genetics and Genome Wide Association approaches and survey and archiving/database-support technology and resources for integrating big data and utilizing artificial intelligence and machine learning for assessing/diagnosing aging-related illnesses.

E. Risk-reduction programs for improving the health of older workers, lowering the rate of health-care utilization, and improving the cost effectiveness of employer-based insurance plans.
Division of Geriatrics and Clinical Gerontology (DGCG)

A. Development and validation of human aging mechanistic markers predictive for various age-related conditions or responses to interventions. Products of interest include the development and validation of commercial assays which could be used in clinical/epidemiologic research to assess mechanisms of aging (e.g., cell senescence, autophagy, DNA damage and repair) in human blood, tissues or cells and markers of age-related chronic inflammation. This may involve refinement of existing assays (e.g., conversion of lab assay to high-throughput screening) and/or de novo assay development for use in clinical research. Novel molecular imaging techniques (in vitro and in vivo) to study aging mechanisms in humans are also encouraged.

B. Potential new therapeutics and/or interventions targeting fundamental mechanisms of aging and that influence the risk or progression of multiple age-related conditions. This may include identification of new therapeutic targets or repurposing of existing FDA-approved medications.

C. Development of high throughput drug screening platforms to identify small molecules for enhancing the functions of protective genetic/metabolic factors associated with exceptional longevity or health span in humans.

D. Development of clinical decision-support tools able to broadly integrate into the electronic health record to help physicians/providers caring for patients with multiple (3 or more) chronic conditions prioritize, coordinate, and deliver the interventions that are most beneficial and relevant within the context of these patients’ lives and the health-care-delivery system.

E. Development of assistive technologies/robotics to enable and support older persons to live independently and safely at home; socially-assistive robots; robots for caregiver and mobility assistance; robots for exercise and rehabilitation assistance.

F. Development of technologies/robotics/sensors to assist in the improvement of physical function and mobility in older persons prior to (pre-habilitation) or following (rehabilitation) elective/planned surgery.

G. Development and validation of improved approaches for evaluation, monitoring or treatment of diastolic dysfunction in older adults.

H. Development of improved instrumentation/imaging and sensor technology for measuring ambulation and biomechanics of movement including balance, sway, gait, and postural control to identify stable and unstable patterns of movement during activities of daily living.

I. Development of methods and technology to accurately determine the renal glomerular filtration rate (GFR) in older persons and patients with chronic kidney disease; new methods and technology should accommodate the effects of age-related changes in muscle mass, levels of serum creatinine, renal blood flow and renal concentrating ability.

J. Development and validation of non-invasive methods of examining bone quality (density, architecture, and strength of bone).

K. Development and validation of instruments and/or methods to evaluate fatigability—the level of fatigue related to the intensity, duration, and/or frequency of activity (in contrast to measures of fatigue), particularly in adults with or at-risk of developing age-related conditions or diseases leading to physical disability.
L. Development and validation of innovative approaches to pain control that considers age-related physiologic changes such as gastrointestinal absorption, cutaneous integrity, and musculoskeletal structure and function.

M. Development and validation of new technology such as non-invasive methods to examine blood-flow velocity in arteries, individual coronary arteries, renal arteries, and cerebral arteries.

N. Development of clinical decision support tools that help physicians caring for patients with multiple chronic conditions to prioritize the interventions that are most beneficial and relevant within the context of these patients' lives; or tools for patient self-management of multiple chronic conditions. Development of patient-focused tools for prioritizing and making decisions about the most significant health concerns to help select and order their self-management behaviors related to 3 or more chronic conditions.

O. Development of new therapeutic interventions to promote wound healing, including improved post-surgical treatments/technologies promoting wound healing, prevention of chronic wounds, or reduced scar formation.

P. Development of vaccines and other agents for preventing and treating infections in older persons, including development of new vaccines or preventive interventions, and new methods using currently available vaccines or preventive medications. Improve vaccine response/immune function, and for physical functional problems in old age.

Q. Development of devices and/or techniques for preventing or treating urinary incontinence.

R. Development and effectiveness testing of innovative, practical, cost-effective technologies, data collection and extraction systems and devices that could enhance the participation in clinical trials of older vulnerable people who are typically under-represented in clinical trials.

S. Development of new diagnostic tests to predict adverse and/or costly, or favorable, health outcomes with aging or in the setting of chronic diseases, injuries, surgery, hospitalization, or other health-related conditions.
NATIONAL INSTITUTE ON ALCOHOL ABUSE AND ALCOHOLISM (NIAAA)

A. Treatment of alcohol use disorder.
   • Pharmacological discovery, strategies, and development
   • Innovative therapeutic approaches
   • Prevention strategies
   • Therapies for co-morbid conditions, including organ damage

B. Technology development to support screening, brief intervention, and referral to treatment for alcohol-involved patients in medical settings.

C. Development of novel technologies, methods or applications.
   • To detect the effects of alcohol on CNS structure and activities
   • To prevent harmful drinking during pregnancy, to identify prenatal alcohol exposure, and to enhance outcomes of individuals with Fetal Alcohol Spectrum Disorder
   • Tools for alcohol-related laboratory studies, such as animal strains, cell lines, stem cells, in vitro techniques, neuroimaging, ligands, in vivo detection of neuromodulators, or computational tools
   • Stem cell generation, dissemination, and model development
   • Voice technology, cell phones, and other
   • To assist investigators in conducting research in alcohol field (i.e., application development, electronic data management system, etc)

D. Development of Biomolecular Signatures of Alcohol Exposure and Alcohol-induced Tissue Injury.

E. Development, Optimization, and Validation of Novel Tools and Technologies for Neuroscience Research.

F. Design, Development, and Improvement of Alcohol Biosensors.

G. Investigational New Drug (IND)-enabling Development of Medications or Devices to Treat Alcohol Use Disorder and Alcohol-related Disorders.

H. Genotyping of DNA samples from subjects with addiction and substance use disorders.
NATIONAL INSTITUTE OF ALLERGY AND INFECTIOUS DISEASES (NIAID)

Division of AIDS (DAIDS)

A. Development of anti-HIV agents directed at new viral or cellular targets, including development and in vivo evaluation of sustained release formulations for treatment and prevention of HIV infection.

B. Development and evaluation of therapeutic vaccines and other immune-based therapies to attenuate HIV disease progression or reduce HIV infectiousness.

C. Development of therapeutic strategies for curing HIV infection or effecting a sustained remission in the absence of daily antiretroviral drug therapy.

D. Development of methods for detecting and quantifying persistent reservoirs of replication competent latent HIV in blood and tissues, including bio-imaging.

E. Development and evaluation of practical and affordable tests (e.g., viral load, drug toxicities, drug resistance) to monitor populations infected with HIV and associated infectious agents. Development of tests to detect early infection or viral rebound in HIV-infected individuals and to determine HIV incidence (HIV infection before seroconversion).

F. Development of long-acting (minimum 30 days) sustained/extended release pre-exposure prophylaxis (PrEP), post-exposure prophylaxis (PEP), and multipurpose prevention technologies (MPT) products that can provide systemic protection from HIV infection.

G. Development of rapid tests for the detection of antiretroviral drugs in various human matrices (e.g., blood, urine, hair).

H. Support novel technologies for HIV incidence detection; biomarkers of infection and prevention engagement; social media approaches to increase HIV prevention initiation/promote adherence; mathematical modeling of prevention strategies; and approaches to identify and retain key populations for HIV prevention research.

I. Development of processes suitable for HIV-1 vaccine product design, development and cGMP manufacturing, formulation, analytics and characterization of (a) HIV Env immunogens and related constructs/products; (b) fabrication, and development of nanoparticle-based delivery modalities, such as self-assembling proteins, surface conjugated/adsorbed nanoparticles, synthetic, lipid and polymer-based nanoparticles; (c) antigen-adjuvant formulations and/or combination-adjuvant(s) and dosage forms (e.g., suspension, lyophilized and aerosolized) for co-delivery/co-administration, (d) production of monoclonal antibodies (neutralizing and/or non-neutralizing); (e) delivery of antibodies as vectored or by nucleic acid technologies), (f) VLPs and viral vectors, and (g) DNA and RNA vaccine platforms.

J. Improving cell line development process (transient, stable pools, stable clones, etc.) by using existing and novel cell lines, cultures, and supporting/customized technologies to expedite and increase Env expression, production, quality, and yield, novel chromatography purification platforms for viral vectors and Env proteins for HIV vaccine manufacturing.

K. Development of formulation and dosage form technologies to prevent or treat HIV and HIV-associated co-infections.

Division of Allergy, Immunology, and Transplantation (DAIT)

A. Allergy, Asthma and Airway Biology Branch will consider preclinical and clinical research for conditions of interest: asthma, food allergy, eosinophilic esophagitis and gastroenteritis in relation to food allergy, atopic dermatitis, urticaria, rhinitis, rhinosinusitis, drug allergy, and sepsis. This includes but is not limited to the development of methodologies to manage, and analyze clinical and epidemiologic research in the above conditions and the development of biomarkers as diagnostic markers, markers of disease severity, predictive markers for treatment effectiveness, particularly of
immunologic interventions such as allergen immunotherapy for food and respiratory allergy; novel approaches for detecting infants at risk for developing asthma and other allergic diseases; immune targets for asthma and allergic disease interventions; development of immunotherapies to prevent or treat allergic diseases.

B. **Basic Immunology Branch** will consider preclinical and clinical research to study the origin, maturation, and interactions of immune cells, immune cell receptors, ligands, cytokine biology, molecular basis of activation, antigen recognition, immune tolerance, immune response regulation, hematopoiesis and stem cell biology, enhancement of vaccine effectiveness in neonates and adults, and basic immunology of vaccines and immunotherapeutics as medical countermeasures for biodefense. This research includes but is not limited to development of novel vaccine adjuvants; single cell assays to isolate and study antigen-specific lymphocytes; immunotherapeutic antibodies; biomarkers of host immune defense; single cell and other sample-sparing assays for study of human immunology.

C. **Autoimmunity and Mucosal Immunology Branch** will consider preclinical and clinical research to develop and improve therapies for the treatment of autoimmune diseases and primary immune deficiencies (not HIV), basic research of autoimmune disease mechanisms, and biomarkers, immunotherapy of disease processes, disorders mediated by lymphocyte products, and mucosal immunity. This includes but is not limited to innovative treatments for autoimmune diseases; standardized validated diagnostic criteria and outcome measures for autoimmune diseases correlated with disease activity; high throughput assay of T-cell activity in autoimmune diseases; biomarkers to measure risk, disease activity, and therapeutic response in autoimmune diseases; innovative treatments for autoimmune diseases; mucosal immunity.

D. **Transplantation Branch** will consider preclinical and clinical research in organ, vascularized composite tissue and cellular transplantation: acute and chronic graft rejection, allogeneic and xenogeneic transplantation, development of immunomodulatory agents to prevent and treat graft rejection and to promote acute and long term graft acceptance and immunologic tolerance, genomics of the alloimmune response, graft versus host disease for hematopoietic stem cell transplantation, minor histocompatibility antigens, complications of immunosuppression in transplantation, and major histocompatibility complex (MHC) region genomics and technologies for MHC typing. This includes but is not limited to methods and analysis tools to facilitate high throughput, high-resolution MHC typing in humans and non-human primates.

E. **Radiation Countermeasures Program** will consider preclinical research to support specific product development activities leading to creation of IND or IDE packages to be submitted to FDA. These IND/IDE-enabling activities could include: efficacy studies to optimize formulation, dose, and dose schedule; drug product stability studies, drug product GMP manufacturing scale-up, GLP toxicology and pharmacology safety studies, pharmacokinetic and metabolism studies, development of GLP analytical methods for efficacy studies and product characterization (including using chip technology to determine tissue-specific efficacy of a lead drug candidate), mechanism of action studies and completion of IND or IDE package for FDA submission. Product development efforts will advance new medical countermeasures towards Phase I clinical safety studies, GLP animal pivotal efficacy studies, and licensure/approval/clearance by the FDA.

**Division of Microbiology and Infectious Diseases (DMID)**

A. Identify and qualify infectious disease-related biomarkers, including:
   1. Biomarkers to predict susceptibility to infection and/or diagnose an infectious disease.
   2. Biomarkers to predict or monitor a subject’s response to therapeutics or vaccinations.
   3. Biomarkers from natural history studies that could be used to assess disease progression in acute and chronic diseases.
B. Development of rapid, highly sensitive and specific clinical diagnostics, including point-of-care diagnostics, that are easy to use, cost-effective and can diagnose individuals infected with pathogens or individuals that have been exposed to toxins.

C. Discovery and development of vaccines or other immunoprophylaxis tools for infectious diseases.

D. Development of vaccine enhancement and formulation technologies with the goal of providing protection against infectious disease agents, providing accelerated immune responses (more rapid schedules or reduced number of immunizations), increasing ease of administration (i.e., self-administration), increasing product stability to minimize cold chain requirements, and enhancing cost-effectiveness of vaccine manufacturing.

E. Discovery and development of therapeutics for infectious diseases.

F. Development of technologies or approaches that address arthropod vector monitoring, management, and control to prevent transmission of vector-borne pathogens to humans.
NATIONAL INSTITUTE OF ARTHRITIS AND MUSCULOSKELETAL AND SKIN DISEASES (NIAMS)

A. Research and development of new therapies using small molecules or biologics for arthritis, musculoskeletal and skin diseases.

B. Research and development of novel biomedical devices or tissue engineered products for arthritis, musculoskeletal and skin diseases.

C. Research and development of new biomarkers or novel imaging technologies for arthritis, musculoskeletal and skin diseases.

D. Research and development of innovative internet-based technologies to manage arthritis, musculoskeletal and skin diseases.
NATIONAL INSTITUTE OF BIOMEDICAL IMAGING AND BIOENGINEERING (NIBIB)

A. **Connected Health-Mobile Health and Telehealth.** Development of enabling technologies that emphasize the integration of wireless technologies with human and biological interfaces. This program includes the development of software and hardware for telehealth and mobile health studies. This program includes the input and delivery of healthcare information digitally for the analysis or monitoring of health or disease status. The emphasis is on developing mobile health technologies driven by clinical needs and integrating these technologies in healthcare delivery, wellness, and daily living.

B. **Engineered Cells.** Development of engineered cells to elicit a broadly applied biomedically relevant effect across a spectrum of diagnostic, therapeutic, imaging, and interventional applications. Emphasis is on engineering functionality and issues surrounding biocompatibility of engineered cells. Functionality could be derived from controlling natural or artificial attributes of an engineered cell and based on biochemical, optical, and/or mechanical properties, for example.

C. **Engineered Tissues.** Development of technologies to enable the *in vivo* and *in vitro* engineering of human tissue constructs for biomedical applications. Emphasis is on the design and construction of tools for analyzing and controlling the function of engineered human tissues. Outcomes include but are not limited to: real-time, non-invasive monitoring of tissue function and cell-environment interactions; control of spatiotemporal tissue growth through cell viability, guiding, differentiation, and migration; design, 3D printing, and assembly of human tissues for and biomedical applications; preservation of biological specimens, from protein solutions and cell suspensions to tissues and organs, for a variety of biomedical applications, including transplantation.

D. **Image-Guided Interventions.** Development of novel image-directed technologies for guidance, navigation, tissue differentiation, and disease identification for reaching specified targets during therapeutic procedures, which may range along the continuum from non-invasive to minimally invasive to open surgical interventions. These technologies may range from molecular to macroscopic scale levels. In addition, emphasis includes technologies that expand needed procedural access for individuals otherwise excluded by disease characteristics, co-morbidities, and other parameters.

E. **Magnetic Resonance Imaging.** Development of *in vivo* MR imaging and MR spectroscopy, for both animal and human research and potential clinical applications. The emphasis is on the development of MRI hardware and methodologies, including image acquisition and reconstruction techniques, that would improve the speed, spatial resolution, information content, efficiency, robustness, quality, patient experience, and safety. The emphasis should be on technological development rather than detailed applications to specific diseases or organs.

F. **Mathematical Modeling, Simulation and Analysis.** Development of novel mathematical modeling, simulation and analysis tools that can be broadly applied across a wide spectrum of diagnostic, therapeutic, imaging, and interventional applications. Emphasis is on engineering solutions for theory-driven, physics-based, physiologically realistic, virtual representations of biomedical systems, with a particular weight on multiscale modeling. Interests include, but are not limited to: multiscale modeling, predictive modeling frameworks, non-standard methodologies, and methods to address model credibility, reproducibility, and reuse.

G. **Nuclear Medicine.** Research and development of technologies that create images out of the gamma-ray or positron emissions from radioactive agents that are injected, inhaled, or ingested into the body. The emphasis is on: simulation and development of new detectors, collimators, and
readout methods that enhance the signal quality of detecting isotope emissions; designs of novel camera geometries; and correction methods that compensate for the radiation physics properties to improve the clinical reliability of the image. Of interest are improvements and corrections for interaction events in PET detectors and enhancement to time of flight (TOF) image generation methods (reconstructions algorithms); as well as new collimator and camera designs for SPECT.

H. **Optical Imaging and Spectroscopy.** Development and application of optical imaging, microscopy, and spectroscopy techniques for improving disease prevention, diagnosis, and treatment in the medical office, at the bedside, or in the operating room. Examples of research areas include fluorescence imaging, bioluminescence imaging, OCT, SHG, IR imaging, diffuse optical tomography, optical microscopy and spectroscopy, confocal microscopy, and multiphoton microscopy. The emphasis is on development of cost effective, portable, safe, and non-invasive or minimally invasive devices, systems, and technologies.

I. **Technologies for Tissue Chips.** Development of technologies to enable the engineering of tissue chips/microphysiological systems for biomedical applications. Emphasis is on the design and construction of in vitro tools for analyzing and controlling the function of engineered human tissues. Examples include but are not limited to: microfluidics to control spatiotemporal tissue growth, 3D bioprinting systems for tissue assembly, high-throughput assays and instruments to reduce the cost, time, and complexity of tissue engineering, and bioreactors to produce tissues at scale.

J. **Therapeutic Medical Devices.** Design and development of non-imaging devices intended for therapeutic interventions. Emphasis is on engineering non-imaging devices, components, and control systems for in vivo therapeutic interventions directed toward overcoming a technological challenge that limits biomedical application. Devices may be, but are not limited to: rehabilitative or curative; assistive; or preventative.

K. **Ultrasound: Diagnostic and Interventional.** Improvement of technologies for diagnostic or therapeutic uses of ultrasound. The diagnostic ultrasound program includes, but is not limited to the design, development and construction of transducers, transducer arrays, and transducer materials, innovative image acquisition and display methods, innovative signal processing methods and devices, and optoacoustic and thermoacoustic technology. It also includes the development of image-enhancement devices and methods, such as contrast agents, image and data presentation and mapping methods, such as functional imaging and image fusion. The therapeutic ultrasound program includes, but is not limited to the design, development, and construction of transducers, transducer arrays, interventional technologies, adjunct enhancement of non-ultrasound therapy applications, high-intensity focused ultrasound (HIFU), or hyperthermia applications. It also includes non-invasive or minimally invasive interventional surgical or therapy tools, ultrasound contrast agents for therapy, targeted drug delivery, neuromodulation, and biopsy.

L. **X-ray, Electron, and Ion Beam.** Simulation, design and development of new detector systems; new readout methods that enhance the signal quality for x-ray image generation; designs of novel imaging geometries; algorithms that compensate for the physical properties of the detection system to improve the clinical reliability of the image (reconstruction algorithms); and approaches to radiation dose reduction, especially in CT. Of interest are diagnostic image enhancements via photon counting, dual energy, and new applications of cone-beam tomography.
EUNICE KENNEDY SHRIVER NATIONAL INSTITUTE OF CHILD HEALTH AND HUMAN DEVELOPMENT (NICHD)

Child Development and Behavior Branch
A. Real time Human Interactive Data Acquisition and Analysis Technologies for Research and Telehealth Needs: Development and research testing of new or adaptation of existing devices and innovative technologies to improve virtual/remote data collection, to facilitate child and parenting intervention and/or healthcare (telehealth) delivery platforms with integrated collection, analysis, and automated coding, including audio and video recordings in real world settings (e.g., homes, childcare centers, schools, and primary care offices) over prolonged periods of time (i.e., days, weeks, or longer) to allow for (1) rapid analysis of interactions, including those involving one or more languages, and (2) simultaneous analysis of nonverbal and verbal behaviors during interactions. Incorporation of data from sensors capable of simultaneously recording real time physiological signals (e.g., pulse, heart rate, skin conductance response, temperature, accelerometer, etc.) time-locked to audio and video data and analyses is also highly desired.

Contraception Research Branch
A. Development of innovative contraceptive approaches for both males and females.

Developmental Biology and Structural Variation Branch
A. Software development to facilitate the collection, mining, and analyses of genomic and phenotypic data from children affected with structural birth defects, and cross-analysis with model organism data
B. Innovative technologies for validation and functional characterization of human structural birth defects-associated genetic variants in model systems
C. Development of user-friendly software for biomedical researchers with limited knowledge of computational biology to analyze large-scale human genomic and other datasets associated with structural birth defects
D. Creation of software platforms for assembly and display of predictive interactive computational models for complex gene regulatory networks coordinating embryogenesis

Fertility and Infertility Branch
A. Development of novel techniques for assessment of gamete quality
B. Development of Apps to monitor male and female reproductive health
C. Over-the-Counter devices for in-home monitoring of ovarian follicle growth and ovulation
D. Novel techniques for preservation of gametes and whole ovary and testes
E. Development of techniques for use of non-embryonic stem cells for fertility preservation in cancer survivors
F. Diagnostic tools for PCOS detection
G. Development of smart phone-based semen assay
H. Tools and technology for single cell analysis of gonadal and reproductive tract tissues

Gynecologic Health and Disease Branch
A. Development of marketable novel or improved methods, devices, and technologies for the diagnosis, monitoring, and therapy of uterine fibroids, endometriosis, adenomyosis, benign ovarian cysts, abnormal menstrual cyclicity, reproductive tract abnormalities (including congenital structural abnormalities and complications from female genital cutting), female pelvic floor disorders (including drugs and devices used for treatment of pelvic organ prolapse, urinary incontinence, fecal incontinence, and other female pelvic floor disorders), and gynecologic pain disorders (including chronic pelvic pain, vulvodynia, and dysmenorrhea).
Intellectual and Developmental Disabilities Branch
A. Technology development to improve screening, diagnosis, treatment, and management of intellectual and developmental disabilities.

Maternal and Pediatric Infectious Disease Branch
A. New technologies relevant to resource-limited countries for screening, diagnosis, and management of HIV and other infectious diseases, including tuberculosis, viral hepatitis, other congenital infections such as cytomegalovirus, respiratory infections, etc., in infants, children and pregnant/breastfeeding women.
B. Development and evaluation of vaccines relevant to HIV and other infectious diseases for infants, children, and pregnant/breastfeeding women.

Obstetric and Pediatric Pharmacology and Therapeutics Branch
A. Development of devices to help diagnose or treat pediatric and pregnancy associated disorders.

Pediatric Growth and Nutrition Branch
A. Isolation, purification, and synthesis of human milk components with biological activity.
B. Develop rapid and reliable methods to determine components (both nutritive and non-nutritive) in human milk.

Pediatric Trauma and Critical Illness Branch
A. The development of devices, innovative therapeutic technologies, and behavioral interventions to improve pediatric patient outcomes and minimize the negative sequelae of trauma, injury or critical illness.

Population Dynamics Branch
A. Developing tools and methods to accurately and reliably measure head circumference in infants and children
B. Technological innovations or inventions to improve collection of biomarker and anthropometric data in large population-representative surveys
C. Hardware or software to improve collection of accurate cause of death information in large population-representative surveys or in administrative data sets
D. Innovative methods to add new reproductive and gynecologic questions and/or sampling frameworks to existing large cohorts and/or longitudinal studies

Pregnancy and Perinatology Branch
A. Devices, instruments, and tools to minimize health-care associated infection risks.
B. Methods to reduce pain in all of perinatal care (in newborn infants, in mothers in labor, during the postpartum period and after spontaneous delivery and cesarean section
C. Novel methods to predict, assess, monitor, or treat (when feasible) fetal health, fetal growth, preterm birth, and preeclampsia.

National Center for Medical Rehabilitation Research
A. Development of medical rehabilitation interventions and biomedical technologies to improve rehabilitation treatment for restoration of function.
NATIONAL INSTITUTE ON DEAFNESS AND OTHER COMMUNICATION DISORDERS (NIDCD)

A. Research and development for biomedical technologies (medical devices, diagnostic instruments, pharmaceuticals, drugs, therapeutics, vaccines, and biologics) that require clearance by the FDA as a regulated product before commercial distribution.

B. Development of novel open design hardware and software that facilitate rapid dissemination, reconfiguration, and enhancement to enable research beyond what can be performed with existing tools.

C. Projects proposing clinical trials with a large number of participants.
NATIONAL INSTITUTE OF DENTAL AND CRANIOFACIAL RESEARCH (NIDCR)

Infectious Diseases and Immunity

A. Develop oral topical formulations with combined microbicidal, analgesic, and anti-inflammatory activities to enhance oral mucosal defenses and prevent and/or control oral infections and lesions in HIV-infected and/or immunosuppressed subjects.

B. Develop safe and effective targeted diagnostic and therapeutic technologies in response to endemic and pandemic infections

Preclinical Research

A. Preclinical research and development activities for dental, oral and craniofacial technologies (including devices, diagnostic instruments, reconstructive materials, pharmaceuticals, therapeutics, vaccines and biologics) that require review and approval by the FDA as a regulated product before commercial distribution.

Clinical Research

A. Develop improved methods to detect and predict progression of dental caries, periodontal disease, reversible and irreversible pulpitis.

B. Develop new or improve methods or materials to enhance oral and craniofacial surgery. This would include both intraoral and extraoral surgery.

C. Develop improved methods or materials to mechanically and/or biologically repair or treat tooth structure damaged by dental caries or periodontal disease.

D. Develop safe and efficacious methods to diagnose caries, pulp vitality and/or periodontal diseases utilizing non-ionizing radiation.

E. Develop technologies for local delivery of drugs to treat oral and craniofacial diseases or disorders.

F. Develop novel non-opioid pharmacological medications for management of acute dental pain.

G. Develop safe and efficacious methods or medications to manage complications of head and neck cancer treatment.

Oral, Oropharyngeal and Salivary Gland Cancers

A. Develop imaging techniques for the early detection, diagnosis and prognosis of pre-malignant lesions.

B. Develop genetic animal models of oral cancer premalignancy and oral cancer progression that mimic human oral cancers, including HPV associated oropharyngeal cancers.

Temporomandibular Disorder and Orofacial Pain

A. Identify and develop novel pharmacologic or biologic agents, including but not limited to small molecules, peptides, recombinant proteins and nucleic acids to prevent, control, and/or treat orofacial pain.

Saliva, Salivary Diagnostics, and Salivary Gland Diseases
A. Develop viral, non-viral and gene therapy-based approaches to address compromised salivary gland function. Develop cell and tissue-based strategies and technologies for restoration of damaged or destroyed salivary gland function.

B. Develop novel compounds or materials that protect and preserve salivary glands from head and neck cancer irradiation therapy.

C. Development of non-invasive methods for the determination of efficacy and safety of artificial saliva, sialogogues, and of their delivery vehicles used in addressing the diminution or lack of saliva (xerostomia) due to Sjögren’s Syndrome or head and neck cancer irradiation therapy.

D. Develop biomarker-based technologies for the identification of Sjögren’s Syndrome using blood or saliva as body fluids.

Biotechnology, Biomaterials, and Applications for Regeneration and Restoration of Oral, Dental and Craniofacial Tissues

A. Develop methods, materials, and devices for orthodontic, prosthetic, periodontic, endodontic and craniofacial applications including those that can be used for craniofacial bone distraction, reconstruction, hard and soft craniofacial tissue healing and regeneration, and scarless craniofacial tissue repair.

B. Develop imagining diagnostics to accelerate clinical implementation of reliable, reproducible, highly specific and sensitive diagnostic instruments for various applications, including but not limited to dental caries, cracked teeth, pulp vitality, bone quality, and periodontal disease.

C. Develop safe and effective biosensors, monitoring devices and systems, data driven and computational tools for automated detection, diagnosis and treatment of dental, oral and craniofacial disease.

Clinical and Behavioral Research

A. Develop and test for safety, efficacy, and/or effectiveness of measures or materials for diagnosing, preventing, or treating oral, dental, and craniofacial conditions and disorders.
NATIONAL INSTITUTE OF DIABETES AND DIGESTIVE AND KIDNEY DISEASES (NIDDK)

NIDDK supports the topics below as they pertain to Diabetes (Type 1 and Type 2 Diabetes, Metabolic Disorders, and Endocrine Disorders), Digestive Diseases (Gastrointestinal Diseases, Liver and Pancreatic Diseases, Obesity, Nutrition, and related diseases), and Kidney Diseases (Kidney Diseases, Urologic Diseases, and Hematologic Diseases).

A. Development or evaluation of pharmacological agents (i.e., drugs, therapeutics), gene therapies, novel formulations, cell-based or other biological technologies for intervention in or prevention of Diabetes and Digestive and Kidney Diseases.

B. Development or evaluation of biomedical devices, tools, techniques, or instrumentation for intervention in or prevention of Diabetes and Digestive and Kidney Diseases.

C. Development of biomarkers, assays, techniques, diagnostic technologies or associated reagents for assessing state or function in normal, developing, or diseased cells or tissues affected by Diabetes and Digestive and Kidney Diseases.

D. Development or evaluation of imaging, screening, or evaluation techniques or technologies for assessing state or function in normal, developing, or diseased cells or tissues affected by Diabetes and Digestive and Kidney Diseases.

E. Development or evaluation of animal or cell models for studying Diabetes and Digestive and Kidney Diseases.

F. Development or evaluation of novel materials or material treatments (e.g., sterilization, coating, etc.) for use in devices or other tools or methods used to prevent, diagnose, or treat Diabetes and Digestive and Kidney Diseases.

G. Development of cell- or data-banks for the biomedical research community.

H. Development or evaluation of technologies, including software applications, for improving patient adherence in Diabetes and Digestive and Kidney Diseases.

I. Development or evaluation of technologies for improving clinical research in Diabetes and Digestive and Kidney Diseases, including technologies for improving data collection and reporting of patient outcomes.

J. Development or evaluation of –omics, informatics, or internet-based technologies for biomedical research or clinical applications in diagnosing or managing Diabetes and Digestive and Kidney Diseases.

K. Development or evaluation of technologies to prevent or avert cell or tissue injury during other disease states or surgical procedures.
NATIONAL INSTITUTE ON DRUG ABUSE (NIDA)

NIDA will allow the increase in budget for the applications in two specific technical/scientific areas: (1) Substance Use Disorders (SUD) Drug Discovery and Development; (2) FDA-regulated Medical Devices for SUD.

Area 1 research will lead to the submission of an Investigational New Drug (IND) application, New Drug Application (NDA) and/or Biological License Application (BLA) to the U.S. Food and Drug Administration (FDA). Area 2 research will lead to regulatory submissions to the FDA for pre-market clearance / approval.

Area 1. SUD Drug Discovery and Development

Projects proposed under Area 1 include application of emerging and existing technologies and platforms for SUD medication development with the focus on products with the potential to minimize drug seeking, compulsive behavior, overdose prevention and reversal:

A. Early therapeutic discovery activities ranging from target identification and validation through lead development.
B. Preclinical and/or clinical drug development.
C. Technologies or formulations to improve medication delivery, longer-acting formulations of existing addiction medications.

Area 2. FDA-regulated Medical Devices for SUD

Projects proposed under Area 2 include pre-clinical and clinical studies in support of FDA pre-market clearance / approval (510(k), DeNovo, Pre-Market Approval) for the following categories of medical devices:

A. Imaging devices for investigating brain function and enhancing disease diagnosis and treatment of SUD;
B. Devices that directly diagnose and/or reduce craving and withdrawal symptoms;
C. Devices that identify and treat neonatal opioid withdrawal syndrome (NOWS), also referred to as neonatal abstinence syndrome (NAS);
D. Digital health therapeutics (e.g., Software as Medical Device, Software in Medical Device) focused on behavioral health interventions to alleviate the burden of SUD;
E. Therapeutic (e.g., neuromodulation) devices and other advanced methods to improve SUD treatment outcomes and relapse prevention;
F. Devices used to diagnose and treat opioid-induced respiratory depression;
G. FDA-regulated devices for physiological monitoring, including remote detection (e.g., wearable sensors, health monitoring/emergency notification systems), specifically tailored to patients with SUD.
NATIONAL INSTITUTE OF ENVIRONMENTAL HEALTH SCIENCES (NIEHS)

NIEHS will accept only Phase IIB SBIR Competing Renewal grant applications in response to specific RFAs focused on the validation of environmental exposure assessment sensor technologies.
NATIONAL INSTITUTE OF GENERAL MEDICAL SCIENCES (NIGMS)

Division of Biophysics, Biomedical Technology, and Computational Biosciences

A. Development of reagents and methods for detecting in real time, analyzing, and separating biologically important compounds, macromolecules, and their interactions.

B. Development of new methods and materials directed toward the solution of biological macromolecule structures, including membrane proteins, assemblies and complexes by, but not limited to, x-ray diffraction, electron diffraction, cryo-EM, NMR and mass spectroscopy.

C. Imaging probes and sensors, other reagents and methods, instrumentation, software for microscopy, spectroscopy, and single molecule analysis of molecules, cells, tissues, embryos and small model organisms. Technologies for applications of microscopy, spectroscopy and single molecule analysis in basic biomedical research, including but not limited to light, electron, X-ray, cryo-electron, and scanning probe microscopy and fluorescence, magnetic and electron paramagnetic spectroscopy. NOT included are small animal and preclinical imaging.

D. Development of instrumentation and devices for detection, analysis, separation and/or manipulation of biologically important molecules, cellular components or cells.

E. Development of instrumentation and devices for elucidating interactions of biologically important molecules \textit{in vitro}, \textit{in vivo}, within cells.

F. Development of probes for detection of genetic polymorphisms, including disease genes.

G. Development of valid animal models for genetic diseases and birth defects.

H. Improvement of methodology (technology) for genetic analysis (e.g., gene expression, probes).

I. Development of tools and technologies to detect and monitor complex phenotypes or traits.

J. Development or improvement of methods for high throughput detection of epigenomic changes.

K. Development of improved technology, reagents and tools to derive, grow, isolate, differentiate and characterize cells.

L. Development or improvement of methods for characterizing and studying complex communities of microorganisms, including interactions with host organisms.

M. Development of non-mammalian model systems.

N. Development of tools and methods for the modeling, simulation, and/or analysis of complex biological systems.

O. Development and/or enhancement of computational tools and methods to collect, store, interpret, analyze and/or visualize biomedical data.

P. Development of software and hardware for improving the effectiveness of computational approaches in biomedical research.

Q. Development of high-throughput and/or computational methods and strategies to define/characterize the function, inhibition, and/or interactions of biological macromolecules and cells.

Division of Pharmacology, Physiology, and Biological Chemistry

A. Methods for isolation, characterization, and production of natural and bio-engineered products.

B. Development of methodology to improve the efficiency of discovery, development, and production of bio-medically relevant compounds.
C. Isolation, characterization, and development of factors and strategies, methods, or treatments involved in critical illness and injury including tissue repair, wound healing, sepsis and associated pain management. Research may involve emergency, peri-operative, and/or critical care conditions.

D. Research to improve drug design and delivery.

E. Development of technologies, instrumentation, software, reagents, and methods for the study of carbohydrates and for determining carbohydrate structure and biological function.

F. Development of tools to study oxidative stress and/or mitochondrial function.

G. Sepsis diagnostics and therapeutics.

**Division of Training, Workforce Development, and Diversity**

A. Development of products or services to enhance diversity of the scientific workforce.

**Division for Research Capacity Building**

A. Development of efficient, user-friendly, and culturally appropriate resources to enhance health science literacy
NATIONAL INSTITUTE OF MENTAL HEALTH (NIMH)

All Divisions:

A. Preclinical drug/device development studies, including pharmacology, efficacy and toxicology.
B. Completion of studies as required by the Food and Drug Administration (FDA) for Investigational New Drug (IND) or Radioactive Drug Research Committee (RDRC) application.
C. Studies in normal healthy volunteers to determine a drug’s safety profile, metabolism, etc.
D. Clinical studies in patient/disease population to assess the drug’s effectiveness.
E. Assessment of devices with regard to performance standards related to the FDA approval process.
F. Safety and effectiveness studies of novel medical devices.
G. Evaluation of novel imaging approaches for diagnostic purposes.
H. Clinical studies in support of Pre-Market Approval for biomarkers/medical devices by the FDA.
I. Rapidly develop novel, engaging computer-based cognitive training programs that are based on efficacious neurotherapeutic approaches and which use cognitive training to target a specific neural system/functional domain.
J. Augment existing computerized cognitive interventions to be personalized, engaging, adaptive, sufficiently challenging, and optimal for maximizing real world functional improvements.
K. Test the feasibility, efficacy and potential adverse effects of these programs utilizing measures of functional outcomes in an identified clinical population, particularly those with neuropsychiatric disorders, ASD, and/or HAND, at a specified developmental stage, including measurement of the duration of treatment effects.
L. Rapid development and evaluation of mobile based platforms and applications.

Division of Neuroscience and Basic Behavioral Science (DNBBS)

A. Novel imaging probes to study brain structure and function at all levels, from the molecular to the whole organ, using any imaging modality (PET, fMRI, optical, etc.).
B. Drug discovery/drug development of novel compounds which act on molecular pathways (receptors, enzymes, second messengers, etc.) that are not typically targeted with currently available psychiatric drugs, and that have a strong biological justification as a novel mechanism for treatment of psychiatric disorders.
C. Novel screening assays for high throughput acquisition and analysis of data about behavior and the brain, from the level of genes to the level of behavior.
D. Novel technologies that would enable researchers to study how populations of neural cells work together within and between brain regions, in order to understand how changes in neural activity contributes to mental disorders.
E. Complex instrumentation for neuroscience research
F. Complex brain or cellular imaging or analysis.
G. Tools to facilitate the detailed analysis of complex circuits and provide insights into cellular interactions that underlie brain function.

H. Proof-of-concept testing and development of new technologies and novel approaches for large scale recording and manipulation of neural activity, at or near cellular resolution, at multiple spatial and/or temporal scales, in any region and throughout the entire depth of the brain.

I. Iterative refinement of such tools and technologies with the end-user community with an end-goal of scaling manufacture towards reliable, broad, sustainable dissemination and incorporation into regular neuroscience practice.

J. Novel informatics tools to facilitate the sharing of complex data sets between laboratories.

K. Novel tools for investigating brain-derived GPCRs in mental health research.

L. Educational tools/technologies for neuroscience and mental health.

M. Technologies to support the goals of the BRAIN Initiative: http://www.braininitiative.nih.gov/

Division of Translational Research (DTR)

A. Develop, test and validate reliable and stable biomarkers that can identify at-risk individuals prior to disease onset, improve diagnosis, predict treatment response, or measure disease progression. Biomarkers are also needed in clinical trials to identify dose ranges, to identify a specific subpopulation of subjects to enroll in a treatment trial, or to measure efficacy or toxicity/side effects.

B. Develop novel and targeted interventions (pharmacological, cognitive, behavioral, computer, or device-based) that affect particular neural circuits and signaling pathways relevant to the developmental trajectory of the disease.

C. Conduct early stage, proof of concept clinical trials to advance the development of novel therapeutics. The clinical trials are expected to include biological/behavioral data to assess target engagement and to help determine potential success or failure of the compound before moving on to larger clinical trials (see NOT-MH-11-015 http://grants.nih.gov/grants/guide/notice-files/NOT-MH-11-015.html).

D. Based on expanded knowledge of neurobehavioral trajectories, develop novel objective assessment tools that can identify early signs of risk or onset of recurrence of particular mental disorders or in domains of functioning (see MH’s RDoC project: http://www.nimh.nih.gov/research-funding/rdoc/index.shtml) for pediatric populations.

E. Develop computational biological/behavioral assessment tools that can be used across ages, species, and cultures to evaluate dysfunction in domains relevant to mental disorders (e.g., mood dysregulation, deficits in executive function).

F. Develop computational platforms to enable the integration and sharing of data characterizing typical and atypical developmental trajectories in humans and non-human animals.

G. Clinical research tools.

H. Develop valid measures of the various constructs in the Research Domain Criteria (RDoC) matrix (see http://www.nimh.nih.gov/research-funding/rdoc/index.shtml), e.g., neurocognitive tasks,
psychometrically sophisticated questionnaires, measures of behavior, and biomarkers, into a commercial product.

I. Develop clinical risk assessment instruments that encompass multiple domains (e.g., genetic, neurobiological, and environmental), are sensitive to developmental stage, and have high predictive power for the onset or recurrence of mental illness.

J. Developing clinical risk assessment instruments for developing particular benefits or harms during treatment for mental disorders, and communicating such probabilistic information to patients and their families in a readily understandable manner.

K. Develop electronic sensors, monitoring devices and systems, and data analysis software for automated detection and diagnosis of mental disorders and key transdiagnostic dimensions of psychopathology.

Division of AIDS Research (DAR)

A. Develop and test novel, non-invasive diagnostic approaches (instrumentation, imaging, biomarkers, central nervous system [CNS] cell-based in vitro models) to detect neurocognitive dysfunction associated with HIV-1 infection and innovative technologies to study the mechanisms involved in HIV-1 associated neuropathogenesis and persistence of HIV-1 in the CNS or strategies to prevent viral resurgence in the CNS upon cessation of anti-retroviral therapy.

B. Design and test novel therapeutic strategies aimed at amelioration of HIV-1 induced CNS dysfunction and/or eradication of HIV-1 from CNS reservoirs or strategies to prevent viral resurgence in the CNS upon cessation of anti-retroviral therapy.

C. Discover and develop innovative technologies for targeting therapies to the brain, including antiretroviral drugs, nanotechnology, imaging tools to study HIV-aging interactions or HIV-related neurodegeneration and neuroprotective strategies with improved capability to cross the blood-brain barrier for amelioration of HAND.

D. Develop evidence-based interventions to reduce risk for HIV among at-risk individuals or improve outcomes along the HIV care continuum for individuals living with HIV which are delivered through mobile devices or other technologies.

E. Develop technologies, instruments and tools to aid in improving uptake, adherence, and persistence to biomedical HIV prevention and treatment regimens; Increasing regular HIV testing among those most at risk of acquiring HIV and translating findings from basic behavioral and social science research into processes to improve engagement in HIV care.

F. Develop new tools/techniques to aid in deciphering the complex neuro-immune interactions at a molecular and cellular level in the context of HIV.

G. Build and optimize multimodal domain based informatics tools to aid in analyzing and characterizing the phenotype of CNS disease modalities associated with HIV by using machine learning, big data and systems biology-based approaches.

Division of Services and Intervention Research (DSIR)

A. Randomized clinical trials evaluating the effectiveness of known efficacious interventions.

B. Analyses of naturalistic databases to evaluate the effectiveness of known efficacious interventions.
C. Identifying moderators and mediators of intervention effects as a step to design and test personalized interventions.

D. Evaluating the combined or sequential use of interventions.

E. Determining the optimal length of intervention, establishing the utility of continuation or maintenance treatment (that is, for prevention of relapse or recurrence).

F. Evaluating the long-term impact of efficacious interventions on symptoms, functioning, and quality of life.

G. Developing novel information technology tools designed to improve the delivery and dissemination of evidence-based interventions and assist healthcare providers in identifying, adopting, and implementing proven prevention and treatment interventions.

**Services research covers all mental health services research issues across the lifespan and disorders, including but not limited to:**

A. Services organization, delivery (process and receipt of care), and related health economics at the individual, clinical, program, community and systems levels in specialty mental health, general health, and other delivery settings (such as the workplace).

B. Interventions to improve the quality and outcomes of care.

C. Enhanced capacity for conducting services research.

D. The clinical epidemiology of mental disorders across all clinical and service settings.

E. The dissemination and implementation of evidence-based interventions into service settings.
NATIONAL INSTITUTE ON MINORITY HEALTH AND HEALTH DISPARITIES (NIMHD)

A. Telehealth, telemedicine, and mobile health technologies (e.g., smart phone apps, web-enabled wearable sensors) to improve remote access to prompt diagnosis, early treatment, and clinical management for adult and pediatric patients in minority and health disparity populations, and to improve access to specialty care that would otherwise be inaccessible due to high cost or transportation barriers (e.g., by linking academic tertiary care-oriented health centers with community-based primary care settings).

B. Products, technologies or services designed to improve accessibility or uptake of existing technologies (e.g., mobile phones, tablets, free WiFi, diabetic glucometers, blood pressure monitors, etc.) within disadvantaged communities and medically underserved areas (including urban, rural, remote, or island regions) to promote healthy lifestyles, enhance patient-clinician communication, provide patient education for self-management of chronic diseases/conditions, or enhance surveillance of communicable and non-communicable diseases in minority and health disparity populations.

C. Products, technologies or services that take advantage of existing or emerging technologies (e.g., electronic health record systems, biomedical informatics platforms, big data resources and analytics, precision medicine) to improve health services delivery and quality of care, including but not limited to coordination of primary and specialty care, integration of behavioral health services into primary care settings, enhancement of provider-patient communication, and reduction of health literacy barriers in minority and health disparity populations.

D. Products, technologies or services to enhance early detection of diseases, pre-disease states, or adverse health conditions in minority and health disparity populations through analysis of novel or validated biomarkers in saliva, breath, blood, and other tissues or specimens, including microbiota.

E. Groundbreaking products or technologies to monitor real-time or cumulative exposures to physical, social and environmental risk factors acting at multiple levels across the life course ("exposome") to improve understanding and situational awareness of factors that may significantly contribute to population health disparities, and/or to empower individuals or communities to take steps to avoid or mitigate the effects of such exposures.
NATIONAL INSTITUTE OF NEUROLOGICAL DISORDERS AND STROKE (NINDS)

The NINDS accepts a broad range of small business applications that are significant, innovative, and relevant to its mission. Examples of research topics within the mission of NINDS are shown below. This list is not all inclusive and some research areas fall into multiple categories.

A. Therapeutics and Diagnostics Development for Neurological Disorders, including biomarker and diagnostic assays, therapeutics (drugs, biologics, and/or devices) for treatment of neurological disorders, and technologies/methodologies to deliver therapeutics to the nervous system.

B. Clinical and Rehabilitation Tools, including intraoperative technologies for neurosurgeons, rehabilitation devices and programs for neurological disorders, and brain monitoring systems

C. Technology and Tools, including technologies to image the nervous system, neural interfaces technologies, and tools for neuroscience research and drug development.

Within these research topics, the following research may require additional funds above the hard budget caps:

A. *In vivo* animal testing required for therapeutics and diagnostics development.

B. Drug and biologics preclinical discovery and development activities for regulatory submission, such as lead identification/optimization, preclinical efficacy testing, IND-enabling studies, and manufacturing for clinical trials.

C. Device preclinical discovery and development activities for regulatory submission, such as hardware prototyping, device/software verification, biocompatibility/sterilization testing, pre-clinical efficacy testing, large animal GLP safety testing, and preparing material/devices for human testing.

D. Clinical testing of therapeutics (drugs, devices, or biologics), diagnostics, clinical and rehabilitation tools (i.e. intraoperative technologies, rehabilitation devices and programs, and brain monitoring systems), and technologies for clinical research. This would include clinical research studies to test scientific hypothesis that are not feasible or practical to conduct in animal models but would inform a final device design.

E. *In vivo* animal testing of technologies for animal research and development of animal models for drug development and neuroscience research.

F. Research that requires special facilities to contain hazardous or infectious materials.

Brain Research through Advancing Innovative Neurotechnologies (BRAIN) Initiative

The Brain Research through Advancing Innovative Neurotechnologies (BRAIN) Initiative is a Presidential project aimed at revolutionizing our understanding of the human brain. NIH is one of several federal agencies involved in the BRAIN Initiative. Planning for the NIH component of the BRAIN Initiative is guided by the long-term scientific plan, “BRAIN 2025: A Scientific Vision,” which details seven high-priority research areas. This report, as well as a list of the specific BRAIN Initiative funding opportunities, can be found at [http://braininitiative.nih.gov/](http://braininitiative.nih.gov/).

Based on priority areas identified by the BRAIN 2025, technology areas were identified to be appropriate for commercial development and may require additional funds above the hard budget caps:

A. Development of research tools and technologies to understand the dynamic activity of neural circuits.

B. Development of novel tools and technologies to facilitate the detailed analysis of complex circuits to provide insights into cellular interactions that underlie brain function.
C. Development of invasive and non-invasive devices for recording and modulation in the human central nervous system.
NATIONAL INSTITUTE OF NURSING RESEARCH (NINR)

A. Development of digital health technologies or services
B. Development of health information/integration technologies or services
C. Development and/or implementation of medical robotic systems
D. Development of medical devices
E. Projects proposing clinical trials with a large number of participants.
NATIONAL LIBRARY OF MEDICINE (NLM)

A. Development and applications to improve storage, retrieval, access, management, representation, and use of biomedical knowledge

B. Development of tools and methods for visualization, modeling, simulation, or analysis of complex biological systems and clinical processes

C. Innovative approaches for data security and privacy, and technical issues related to other ethical, legal, and social implications of personal health data

D. Methods for data integration to support discovery, learning, and health care

E. Informatics tools that assist in delivering precision medicine to patients, or health decisions
DIVISION OF PROGRAM COORDINATION, PLANNING, AND STRATEGIC INITIATIVES (DPCPSI), OFFICE OF RESEARCH INFRASTRUCTURE PROGRAMS (ORIP)

RESEARCH AND DEVELOPMENT IN THE DIVISIONS OF COMPARATIVE MEDICINE AND OF CONSTRUCTION AND INSTRUMENTS

A. Development of new technologies for rapid characterization and deep phenotyping of large numbers of animals.
B. Development of technologies to identify biomarkers for clinical diagnostics in well validated disease models.
C. Development of vaccines and new therapeutic agents to prevent and/or control selected laboratory animal diseases. One high priority need is to develop methods to control and prevent Herpes virus B in nonhuman primates.
D. Design of specialized equipment and caging for laboratory animals to permit optimal environmental control and operational efficiency, including improvements in caging, identification/tagging of animals and remote monitoring in animal facilities.
E. Identification, development, and characterization of spontaneous and engineered vertebrate animal models for studies of various human diseases, excluding most random mutagenesis projects performed on rodents.
F. Development and refinement of high throughput technologies and devices for the effective cryopreservation, long-term maintenance of cells, tissues, and laboratory animal embryos, and gametes.
G. Development of technologies and devices for the effective monitoring of frozen and cryopreserved cells, biological materials/tissues and laboratory animal embryos, and gametes.
H. Development of improved reagents, artificial intelligence/machine learning technologies, devices and high throughput technology to perform, analyze, capture and process data gathered in “omics” studies (genomics, transcriptomics, phenomics, proteomics, glycomics, epigenomics, metabolomics, among others) in normal disease and intervention conditions in animal/biological models.
I. Development of biological tools and reagents for reconstruction, remodeling, repair and regeneration of tissues damaged by injury or disease. Development of the technologies and procedures to test efficacy and safety of these experiments in animal models. Approaches to detect and track the implanted cells and tissues in vivo.
J. Development of acellular biomaterials, biosensors and reagents to promote, detect and track reconstruction, remodeling, repair and regeneration of tissues and organs damaged by injury or disease.