The Center for Innovative NeuroTech Advancement (CINTA) & NeuroTech Harbor (NTH) Announce the Cycle 3 Award Competition

> Supported by the NIH Blueprint MedTech Program

Blueprint MedTech Incubator Hubs



- CINTA (Center tor Innovative Neurotech Advancement), a program within CIMIT (Steven Schachter, MD as PI and Paolo Bonato, PhD as co-PI from Spaulding Rehabilitation Hospital).
- NTH (NeuroTech Harbor), a partnership between Johns Hopkins University and Howard University (Sri Sarma, PhD as contact PI and Evar Nwulia, MD).

Blueprint MedTech Incubator Hubs Mission

Mission

 To accelerate the development of emerging, ground-breaking technologies into first-inhuman studies along the path to commercially viable, clinically focused solutions for disorders involving the nervous system.



Healthcare Innovation Cycle

 To catalyze translation along 4 key domains in the innovation cycle: Technology, Regulatory, Market/Business, Clinical



Funding Opportunity // https://blueprintneurotech.org/

- Awards from NTH or CINTA will rarely exceed \$500,000 in direct costs per year for a period of up to 4 years. Indirect costs will be provided at your institution's Federally negotiated rate (or 10% de minimis).
- In addition to monetary support, awardees will receive ongoing, specialized support from executive mentors experienced in developing and commercializing neurotech devices.
- Awardees will work with their executive mentor each week to focus on business, regulatory, clinical, and technical factors that may impede commercialization.

Blueprint MedTech Resources

- Resources to plan and support prototype development, team building, needs assessment, and other early translational activities.
- Additional assistance from hubs and consultants (e.g., design, regulatory, reimbursement, intellectual property, commercialization, and strategic partnership issues).
- The cost of resources provided by the hubs and NIH do not need to be included in the proposed budget.
- Other resources listed on the <u>Blueprint MedTech website</u>





NIH Blueprint for Neuroscience Research

Potential Resources

Design, Prototyping, Risk Analysis

- Electronics Manufacturing
- Prototype Manufacturing
- Design Optimization and Risk
- Computational Modeling

Bench and Safety Testing

- Electrical Safety
- Electromagnetic Compatibility
- MR Testing
- Software
- Cybersecurity
- Shelf-life Testing

Biocompatibility and Animal Studies

- Biocompatibility Testing
- Materials characterization and analytical chemistry
- Sterilization testing/validation
- Preclinical Studies Animal Testing (GLP)
- Preclinical Studies Animal Testing (non-GLP)
- Cadaver Testing

Clinical

- Clinical trials
- Biostatistics
- Data Management
- Neuroethics

Anticipated resources provided by Resource Subawards Core

Provided by other parts of BP MedTech

Business Development

- External Oversight Committee
- Public-Private Partnerships CRA, MTA
- Entrepreneurship
- Business Development
- Market / User Research
- Commercialization

Regulatory, Compliance, Quality System

- Regulatory Advising
- QMS Quality Management System setup and audits
- GMP Good Manufacturing Practice setup and audits
- Compliance
- Legal Intellectual Property

Participating Centers and Institutes

- National Institute of Biomedical Imaging and Bioengineering (NIBIB),
- National Center for Complementary and Integrative Health (NCCIH),
- National Eye Institute (NEI),
- National Institute on Aging (NIA),
- National Institute on Alcohol Abuse and Alcoholism (NIAAA),
- Eunice Kennedy Shriver National Institute of Child Health and Human Development (NICHD),
- National Institute on Drug Abuse (NIDA),
- National Institute of Dental and Craniofacial Research (NIDCR),
- National Institute of Mental Health (NIMH),
- National Institute of Neurological Disorders and Stroke (NINDS), and
- Office of Behavioral and Social Sciences Research (OBSSR)

Applications must focus on a disorder of the nervous system in an area of interest of the <u>NIH Participating Institutes/Centers for the Blueprint MedTech: Incubator Hubs program</u>.

Applications outside the mission of these participating Institutes/Centers will not receive funding. Contact Institute Program Officer for questions about **mission fit only**.



Please forward all other questions to info@blueprintneurotech.org.

https://neuroscienceblueprint.nih.gov/neurotherapeutics/blueprint-medtech/blueprintmedtech-ics-and-contacts

Eligibility

The Blueprint Hubs and NIH encourage applicants from women, underrepresented racial and ethnic groups, as well as individuals with disabilities in the translational workforce to apply. Principal Investigators (PIs) from academic institutions, industry or non-profit organizations are invited to apply. Foreign applicants may apply.

Academic PIs must hold a faculty appointment at an institution of higher education or medical center.

PIs from industry or non-academic non-profits are not required to hold a faculty appointment.

Review Process

Applicants must first submit **pre-proposals**, which will undergo review by CINTA, NTH, and NIH program scientific staff. Preproposals are submitted through a simple online application form equivalent to about 4 pages (CoLab).

> A subset of the applicants who submit pre-proposals will be selected to submit **full proposals** which are submitted through the same online application system. The online full proposal form is equivalent to approximately 10 pages.

> > A subset of the applicants who submit full proposals will be selected to participate in a **"deep dive"** evaluation, which is the final stage of review prior to funding decisions.

*sections of the form have word limits and there is an upload with a page limit, but there really is not an overall page limit per se.

General Application Information

• Applicants should review and be familiar with the program solicitation and FAQs before completing this application.

Solicitation link: https://blueprintneurotech.org/ FAQ link: https://blueprintneurotech.org/faq

- If your project addresses a mental health disorder, you are encouraged to provide preliminary data that uses quantitative, objective measures for outcomes. And to incorporate these measures into your proposal.
- Only IRB-exempt or minimal-risk clinical studies can be proposed for funding, and that minimal-risk clinical studies will be conducted at Georgia Tech's HomeLab, one of the core resources of the Blueprint Medtech program.

https://cacp.gatech.edu/research/accessibility/HomeLab

Applications that will NOT be considered

- Products not regulated by the FDA.
- Fundamental basic/applied research prior to proof of concept.
- Device technologies that do not significantly advance the state pf the art (e.g. device technology that proposes minor modifications to FDAapproved/cleared medical device technology)
- Animal model development: all *in vivo* animal models must be wellestablished and characterized, and available to the applicant.
- Projects focused on technologies for functional augmentation of healthy individuals.



Pre-Proposal Application Sections

- 1) Applicant Information
- 2) Solution Information
 - Medical Condition
 - Technology
 - Clinical Need & Standard of Care (<250 words)
 - Solution Description (<500 words)
 - Supporting Information and/or References Upload 1 page PDF

3) **Project and Team Information**

- Proposed Scope of Work (<250 words)
- Team and Environment (<250 words)
- Workforce Diversity (<200 words)
- Regulatory (<150 words)
- Support Requested (Checklist)

Lessons Learned from Previous Cycle Submissions Common Reasons for Rejection

Stage of Maturity

- Too early (no proof of concept)
- Too advanced (ready for clinical trials; candidate for UG/UH3 or U44)

Team Composition

- Lacking critical areas of expertise
- No evidence of clinical collaboration

• Impact

- Not significantly different from existing products
- Only marginal impact on clinical condition
- Mission fit
 - Not priority area of NIH participating institutes/centers



Timeline*



* Two solicitation cycles per year.



Companion Seedling Program



- A subset of all applications will be redirected as invitations to participate in the Seedling program.
- Seedlings are akin to planning grants where the two hubs will provide training and mentoring to help applicants refine the proposal to strengthen subsequent applications to the program.

Companion Seedling Program



- Seedling program is a 6-month mentorshipbased accelerator program for applicants to the BPMT full award program whose solutions have one or more weaknesses in:
 - Regulatory
 - Project details
 - Team expertise
 - Scope of work
 - PEDP (Plan for Enhancing Diverse Perspectives)
- All of which would be addressable through 6 months of work with mentors and experts in these specific gap areas.





solicitations.

Guide innovators proposing more clinical translational or feasibility studies to competitively submit to BPMT Translator (e.g., UG3/UH3, U44) programs Guide innovators to other sources of fundings







Seedling Program Processes:

- Candidates will be interviewed by the Seedling program leadership to explore project fit with the program objectives and to gauge the team's interest in participation.
- If accepted into the program, expert mentors will be assigned to each team for individualized support.
- The team with the support of the mentor will prepare a revised budget and project plan to address identified gaps.
- Over the project period, teams will meet for 1-2 hours each week with mentors.

Summary Information

- Awards will rarely exceed \$500,000 per year in direct costs. Indirect costs will be provided at your institution's Federally negotiated rate.
- The initial anticipated performance period is 12 months, which can be renewed for up to an additional three12-month periods with CINTA, NTH, and NIH approval.
- The earliest anticipated start date for funding for selected full proposals and seedlings is June 2024. (If funds are received from the Federal government and all necessary approvals are in place).
- The final aim of Hub incubator projects should be a prototype ready for first-in-human studies.
- Upon successful completion of the project, teams should either have nongovernmental funding secured or be ready for entry into the companion translator solicitations from NIH:
 - Blueprint MedTech: Translator (UG3/UH3)
 - Blueprint Medtech: Small Business Translator (U44)

Contacts and Additional Resources

Webinar Slides/Schedule: <u>https://www.cimit.org/web/center-for-innovative-neurotech-advancement/events</u>

FAQs: <u>https://blueprintneurotech.org/faq</u>

Scheduling Office Hours: <u>dlee33@partners.org</u> **Availability:** Fridays (8/4 - 8/25 from 1-3PM ET for 15 mins per team)

General Information: <u>info@blueprintneurotech.org</u> 617-643-3800





Good Luck!