REQUEST FOR APPLICATIONS – PILOT PROJECTS

Deadlines: February 14, 2014 (Letter of Intent) & March 10, 2014 (Full Proposal)

Note – deadline for LOI extended from February 10 to February 14, 2014.

Overview

The Institute for Integration of Medicine & Science (IIMS), the Cancer Therapy & Research Center (CTRC), and the Greehey Children’s Cancer Research Institute (GCCRI) are soliciting proposals for pilot project awards. The primary goal of these programs is to support early-stage collaborative translational and clinical studies that will lead to increased interdisciplinary, institutional, and community-based research likely to compete successfully for national grant support and ultimately to improve human health. The programs are supported by the Clinical and Translational Science Award (CTSA) and the Cancer Center Support Grant (CCSG) from the National Institutes of Health (NIH), as well as by various institutional funds.

Eligibility

IIMS-CTSA Program

Applicants must hold faculty-level appointments at the University of Texas Health Science Center at San Antonio (UTHSCSA) or one of its CTSA partner institutions (San Antonio Metropolitan Health District, San Antonio Military Health System, South Texas Veterans Health Care System, Texas Biomedical Research Institute, University Health System, University of Texas Pan American, University of Texas Brownsville, University of Texas School of Public Health–San Antonio and Brownsville Regional Campuses, University of Texas College of Pharmacy). Faculty are encouraged to collaborate with individuals from CTSA partner organizations. Previous pilot project award recipients may apply again, but the new application must be a distinct project or a substantial departure, not a simple extension of the previously funded project. Pilot awards target especially junior faculty with potential for career development; however, senior faculty are eligible if the project is a novel departure from currently funded programs.

CTRC Program

Applicants must be members of CTRC (they can apply for membership at the time of submission) and the cancer relatedness of the project must be clearly defined within the proposal.

GCCRI Program

Applicants need not be directly affiliated with the GCCRI, but the relevance of the project to children’s cancer must be clearly defined within the proposal.

Submission, terms, and conditions

An individual may submit no more than one project as a Principal Investigator, plus one as a Co-Investigator.

A required letter of intent (LOI) must be submitted by Friday, February 14, 2014 at 5:00 pm and should include the title of the project, principal investigator (PI), a few sentences describing the project and a list of 3 to 4 potential reviewers from UTHSCSA, or other CTSA partner institutions, but not from the same department or research group as the PI (Click here for LOI template). Note that the LOI is required for planning purposes, but will not serve as a screening tool for proposal submission (i.e., all PIs submitting the LOI should then proceed to full
The deadline for receipt of the full application is **Monday, March 10, 2014 at 5:00 pm.** Both the LOI and the full proposal should be submitted electronically, each in a single PDF file attached to an email message to Ms. Cindy Russel (russel@uthscsa.edu). Please do not send multiple documents. Pilot application form can be found [here](#).

Awards will be made for a one-year project period starting on or about June 1, 2014. Progress reports will be requested six months (brief) and 12 months after the initiation of funding. For projects involving the use of human subjects or vertebrate animals, no expenditures will be permitted until IIMS is provided with a copy of the official letter of approval by the appropriate Institutional Review Board (IRB) or Institutional Animal Care and Use Committee (IACUC), respectively. Investigators are encouraged to submit IRB and IACUC protocols early in order to avoid significant delays in project initiation. Excessive delays in meeting these regulatory requirements may result in withdrawal of the award. Applicants must also be up to date on compliance with institutional conflict of interest disclosure policies.

**Budget and financial policies**

The maximum budget for these one-year awards is $50,000, although proposals with smaller budgets will be considered and reviewed under the same criteria. Facilities and Administrative (F&A, indirect cost) expenses will not be reimbursed. Funds may not be used to support the salary of the PI or faculty-level collaborators. Although the PI (and Co-PI/Co-I, if applicable) should be listed in the personnel section of the summary page, there is no minimum effort requirement. Salary (plus associated fringe benefits) may be requested for non-faculty support staff, including post-docs. Other allowable expenses include: equipment essential for the project (maximum $10,000, including computer hardware); PI or Co-PI travel to relevant scientific meetings (maximum $1,500); consumable laboratory supplies; animal purchase and per diem; IIMS, CTRC, GCCRI, or other core facility fees; consultation fees (maximum $5,000); computer time; software; publication / presentation expenses; costs related to human subject enrollment and management (listed as “Patient Care Costs” on budget page); and other expenditures that can be justified as being essential for the completion of the project. Account management will be centralized within IIMS-CTSA, CTRC, or GCCRI, with expenditures and encumbrances for UTHSCSA projects being committed as they are incurred. For projects supported at other CTSA partner institutions, funds will be disbursed at appropriate intervals, based on the receipt of invoices for budgeted expenditures.

**Application requirements and format**

Applications should be prepared using the templates provided (adapted from PHS Form 398). Font size should be no smaller than 11 point, preferably Arial or Times New Roman. The font size for figures, figure legends, charts, and tables may be smaller, but must be clearly legible. Margins all-around should be at least 0.5”. Pages should be numbered sequentially. The length of the Research Plan (narrative with illustrations included) is limited to 4 single-spaced pages. The organization of the proposal should be as follows:

- Cover page (p. 1)
- Project summary (level appropriate for scientific peers in the field) and key personnel (p. 2)
- Budget (p. 3)
- Budget justifications (p. 4; maximum 1 page)
- Biographical sketch for PI (maximum 4 pages; for NIH template, format, and sample see [http://grants1.nih.gov/grants/funding/phs398/phs398.html](http://grants1.nih.gov/grants/funding/phs398/phs398.html))
- Biographical sketches for other key personnel (maximum 4 pages each)
- Research plan (maximum 4 pages; use continuation page template)
  - Hypothesis and specific aims
Background and significance
Preliminary data
Work proposed (including statistical analysis, pitfalls, alternatives)

- Literature citations (maximum 1 page)
- Additional information regarding the project (maximum 1 page) to include:
  - Past IIMS funding (TTR and/or pilot project awards) and list of grants obtained or applied for resulting from IIMS support
  - Career development potential, if applicable
  - Prospects and specific plans for outside funding (this is a required element)
  - Collaborative, interdisciplinary, or community engagement features, if applicable
  - Description of how the pilot project will interact with existing programs of the IIMS-CTSA, CTRC, GCCRI, or other CTSA partners, as appropriate.

- Letters of support from core directors or Research Imaging Institute are required (if applicable)
- Letters of collaboration (optional)
- Appendices are not allowed
- A UTHSCSA Certificate of Proposal (COP) is not required

Review process and criteria

Applications will go through a two-tiered system of review. An initial programmatic review will be performed by the IIMS-CTSA, CTRC, and GCCRI leadership teams for program relevance and potential public health impact, taking advantage of input from institutional and community partners. The second phase, or scientific review, will be performed by Scientific Review Committees, including appropriate content experts and representatives from CTSA resources and services, CTRC programs, and other CTSA partner organizations. Scientific merit will be scored by these groups based on the following criteria:

- Significance
- Novelty / innovation
- Strength of the study protocol, including:
  - Design
  - Feasibility
  - Preliminary data (to the extent available)
  - Integration with ongoing research
  - Quality of the investigative team
- Contribution to career development of clinical / translational scientists, if applicable
- Extent of meaningful interdisciplinary collaboration and / or community engagement
- Likelihood of future NIH or other competitive external funding
- Use and leveraging of IAMS-CTSA, CTRC, GCCRI, or partner resources (for example, core facilities - http://iims.uthscsa.edu/ttr_core_facilities.html, biobanking)
- Potential for ultimately improving health outcomes
- Protection of human subjects and experimental animals

Funding decisions will be based on scientific merit, as well as programmatic considerations, such as breadth and depth of the overall pilot study portfolio, interactions among partners, community involvement, and balance among program areas and disciplines.

Special emphasis areas

IIMS-CTSA

Although the intent of this pilot project solicitation is to support a broad range of clinical and
translational research activities, in keeping with priorities developed in the UTHSCSA CTSA application, several areas are highlighted for their special relevance to IIMS programmatic goals. Proposals responsive to the following specific IIMS/CTSA interests are especially encouraged:

- **Research Ethics and Regulatory Science** – Proposals addressing ethical and regulatory issues related to the conduct of clinical and translational research are encouraged. Among our interests are justice, culturally appropriate data collection, informed consent processes, and means for limiting risks to human subjects.

- **Community Engagement** – Proposals for community-based research initiatives, including those involving Practice-Based Research Networks, health outcomes research, epidemiology, and public health research are strongly encouraged. Projects dealing with the demographics and prevalent health problems in our region are of particular interest. Examples include factors related to culture, ethnicity, socioeconomic status, health access, and health disparities, as well as such clinical disorders as obesity, diabetes, infectious diseases, cancer, cardiovascular disease, trauma, and mental health disorders.

- **Comparative Effectiveness Research (CER)** – IIMS has a special interest in CER proposals that have the potential to inform health care decisions by providing evidence on the effectiveness, benefits, and harms of different treatment options using drugs, medical devices, diagnostic tests, surgeries, or means for delivering health care.

- **Biomedical Informatics** – IIMS is especially interested in projects that will develop and/or utilize databases that cross the three major domains of clinical research, clinical medicine, and population health. Projects might study approaches to the integration of data across CTSA partners or apply informatics methods to health status analysis of defined cohorts. Research on the development of novel systems for enhancing communication and data sharing among researchers in various settings (e.g., academic, healthcare, public health, community) would also be of interest.

- **Drug Development** – Evolving NIH priorities encourage proposals representing collaborative translational science studies that test novel translational research hypotheses/concepts, examine cell or animal model systems related to human studies, elucidate disease molecular pathological pathways or underlying biological mechanisms, develop and validate biomarkers, novel drug targets, new agents, new devices/tools, or therapeutic technologies (e.g., drug and gene delivery systems) for humans.

**CTRC**

CTRC-CCSG programmatic goals – NCI Cancer Center Support Grants are intended to support infrastructure for transdisciplinary cancer-relevant research and to foster studies that would not occur without the climate, facilities, and research resources that a cancer center can uniquely provide. The CTRC is particularly interested in collaborative, translational (pre-clinical or clinical) research that has an effect on our South Texas catchment area. Highest priority will be given to funding research in either the most prevalent malignancies in the region (breast and prostate cancer, as well as childhood leukemia) or those that have a disproportionate effect on the population of our catchment area (hepatocellular carcinoma, gastric cancer, cervical cancer and childhood leukemia). The CTRC has a special interest in diseases that affect minorities including Hispanics in the South Texas catchment area, as well as women, and children. Projects should clearly explain how they are related to cancer in South Texas and must demonstrate the potential to lead to extramural funding.

**GCCRI**

GCCRI will support projects that address important issues in childhood cancer across a broad
spectrum from mechanistic studies to drug development and testing, as well as translation to community-based studies.

**Funding expectations**

It is anticipated that up to $800,000 will be available to fund proposals received in response to this solicitation, depending to some extent on currently pending sources of support. A particular caveat is that the CTRC CCSG from NIH is under review for a 5-year renewal period during spring 2013. The outcome of this process may have an impact on available funds. The approximate number of awards for the various program components is generally expected to be as follows:

- IIMS-CTSA translational program – 8 projects
- IIMS-CTSA/CTRC cancer-related program – 4 projects
- IIMS-CTSA/GCCRI children’s cancer-related program – 3 projects

**Responsibilities of the Principal Investigator**

The principal investigator of funded projects is required to:

- Abide by NIH rules and regulations
- Abide by IIMS-CTSA/CTRC/GCCRI and/or CTSA partner policies and procedures
- Provide demographic information in a timely fashion as required before expenditures can be authorized
- Provide a waiver of Facility and Administration (F&As) or Indirect Costs from your institution if you are a non-HSC investigator
- Submit complete and timely progress reports
- Acknowledge support from IIMS, CTRC, and/or GCCRI grants and institute/center funds in all publications and reports generated with pilot project resources (details to be provided at the time of funding)