Request for Applications
DaCCoTA Ready-to-Go Pilot Grant

Overview
The Dakota Cancer Collaborative on Translational Activity (DaCCoTA) Pilot Projects Program is requesting applications for Ready-to-Go Pilot Grant Awards. The goal of the DaCCoTA program is to stimulate growth of expertise and engagement in cancer-related clinical and translational research (CTR) in the Dakota region encompassed by the states of North and South Dakota. The mission of the Pilot Projects Program is to provide seed funding to highly innovative projects in clinical and translational cancer research. For general information about the DaCCoTA program, see https://med.und.edu/daccota/.

The Ready-to-Go Award is intended for those projects with existing significant preliminary data in support of a novel clinical/translational cancer-related hypothesis. These projects should ideally be ready for extramural submission within a year and/or be able to demonstrably improve health outcomes. Applications can consider the multilevel manifestations of cancer (e.g. neurological, psychiatric), demographic risks and social impact. Applications will focus on T2-T4 translational research, although T1 studies will be considered if there is a clear plan to progress to T2-T4. Successful completion of these proposals should lead to a collaborative extramural grant submission and peer-reviewed manuscript submission.

Eligibility Criteria
Faculty and clinicians at participating DaCCoTA institutions who have applied and been accepted into the DaCCoTA Pathfinder network (https://pathfinder.med.und.edu/) are eligible to apply. Investigators with a Principle Investigator (PI) role on other NIH IDeA funding mechanisms are not eligible for a PI role on DaCCoTA awards.

All proposals must include a discussion of the relevance of the proposal to the DaCCoTA mission and consideration of relevance for high priority cancers and vulnerable populations in North and South Dakota. Additionally, participation in the DaCCoTA Pathfinder network is required. Prior consultation with the CTR: 1) Pilot Projects, 2) Biostatistics, Epidemiology and Research Design, 3) Community Engagement and Outreach, and 4) Clinical Research Resources and Facilities Cores is required. Demonstration of available research percentage effort for the PIs/Co-Is and associated personnel on the proposal must be provided. Evidence of available research effort and salary support must be provided in the form of a letter from the departmental chair or clinical service unit chief for each PI/co-I. All proposals must have completed internal approval at their respective institution.

Team Requirements
Applications must be from a collaborative clinician/non-clinician team. The teams may or may not require mentoring and collaborative support to initiate and carry out a research project with the express goal of maintaining a research effort. All pilot awards must have a PI or Co-I who is a clinician and a PI or co-I who is a non-clinician. If a potential PI needs but has not been able to identify a relevant teammate, the Pilot Projects Program will assist in identifying one from among the collaborating academic institutions and hospital networks participating in the DaCCoTA program. The primary PI must also meet eligibility requirements for NIH funding. Preparation of all pilot awards should involve extensive interaction pre-submission as well as post-award with the Pilot Projects Program, which will coordinate interactions with other cores to provide insight regarding research design, grant/publication preparation, and federal/non-federal funding mechanisms. Awardees are
required to provide an oral presentation at the annual DaCCoTA symposium and provide milestone reports.

**Funding**
A maximum budget of $120,000 (direct+indirect costs) is allowed. Funds may not be used for PI/Co-I salaries, consultant/collaborator salaries or to purchase capital equipment. PIs must have an eRA Commons username.

Projects will be evaluated primarily on the basis of potential for immediate implementation, potential for extramural grant submission and peer-reviewed publication, readiness of the technology or process to advance to the next stage of translation, impact on clinical practice, and strength of the investigative team and any collaborators. All funded projects must have clearly established milestones. Milestone progression will be monitored by the Pilot Projects Program.

**Application process and deadlines**
Letters of intent (1-page maximum) are due October 30, 2021. Letters of intent should provide a study title, a study overview, grant category targeted, and identify the clinician/non-clinician team.

Full applications will be invited from selected applicants and will be due January 17, 2022. Awards will be announced by September 2022. Expected award period will be September 2022-August 2023.

- **Proposed Timeline for Pilot Grants**
- **RFA Release Date**: September 1st
- **Deadline for Assistance Finding a Collaborator**: October 8th
- **Letter of Intent Deadline**: October 30th
- **Letter of Intent Acceptance/Decline**: November 15th
- **Required DaCCoTA Core Meetings**: December 13th
- **IRB and IACUC Preparation**: November 15th
- **IRB Draft Application Deadline for CRRFC Consultation**: January 7th
- **Internal Budget and Grant Approval**: January 7th
- **Application Deadline**: January 17th
- **Notification of Review**: February 25th
- **IRB Submission**: February/March dependent on Institution
- **Notice of Award**: August

Applications must be use Arial 11-point font with 0.5 inch margins and must use PHS 398 forms [https://grants.nih.gov/grants/funding/phs398/phs398.html](https://grants.nih.gov/grants/funding/phs398/phs398.html)

- Form Page 1: Face Page
- Form Page 1-continued: Additional form for Co-PI or Co-I
- Form Page 2: Summary, Relevance, Project/Performance Sites, Senior/Key Personnel, Other Significant Contributors, and Human Embryonic Stem Cells
- Form Page 4: Detailed Budget for Initial Budget Period
- Research Plan (use Continuation Format Page)
  - Response to review if resubmission (0.5 page)
  - Specific Aims (1 page)
  - Significance section (0.5 page)
  - Innovation section (0.5 page)
  - Research strategy (maximum of 5 pages)
- Biographical Sketches (use [https://grants.nih.gov/grants/forms/biosketch.htm](https://grants.nih.gov/grants/forms/biosketch.htm))
- References cited (use Continuation Format Page)
Letters of support from the Department Head or Clinical Unit Chief
- Checklist Form Page
- Statement verifying IRB application submission (Use Continuation Format Page)
- Human Subjects Overview Form (see below)

The letter of intent should be submitted to: Jessica.craig@und.edu.
Subsequent full application should be submitted as a pdf to Pathfinder (see step by step instructions below):

1.) Go to Pathfinder website and register/login
   - Home Page - Pathfinder (und.edu)
2.) Once you have registered/logged in, select Proposals on the top menu.
3.) Select Submit a Proposal
4.) Fill out Proposal Contact Information and Proposal Information
5.) Under Proposal Information click on the green box that says file, select your PDF file and then click the green box on the bottom right that says Submit Proposal.

All applicants proposing studies involving human subjects or vertebrate animals must initiate the process of obtaining IRB or IACUC approval prior to submitting their applications.

All proposals recommended by the review panel will require prior approval by NIH prior to initiation. If the proposed study involves human subjects, the following documents will need to be submitted to NIH prior to award notice:

- Written protocol addressing the risks and protections for human subjects, in accordance with the NIH’s Instructions for Preparing the Human Subjects Section of the Research Plan
- Institutional Review Board (IRB) approval
- Human Subjects education certification
- Create Inclusion Data Record (IDR) and enter inclusion data in Inclusion Management System (IMS)

If the proposed project involves clinical trials, in addition to above documents, the following documents must also be submitted to NIH:

- Specific plans for data and safety monitoring.
- Good Clinical Practice (GCP) certification

If the proposal involves Vertebrate Animals, IACUC approval and the Vertebrate Animal Section will also need to be submitted to NIH prior to award notice.

More information
Questions related to the Ready-to-Go Pilot Grant Awards should be directed to Jessica Craig, Pilot Grant Core Coordinator: Jessica.craig@und.edu
DaCCoTA Grant Application
Human Subjects Overview

<table>
<thead>
<tr>
<th>Role</th>
<th>Name</th>
<th>Affiliation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical Investigator</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
<tr>
<td>Non-Clinical Investigator</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
<tr>
<td>Click or tap here to enter</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
<tr>
<td>text.</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
<tr>
<td>Click or tap here to enter</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
<tr>
<td>text.</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
<tr>
<td>Click or tap here to enter</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
<tr>
<td>text.</td>
<td>Click or tap here to enter text.</td>
<td>Click or tap here to enter text.</td>
</tr>
</tbody>
</table>

**Project Title:** Click or tap here to enter text.

**IRB Approval or Human Subjects Determination Status**
- □ Approved
- □ Pending Approval/Submitted
- □ Not Submitted
- □ Not Required (project does not involve human subjects research) FORM COMPLETE, Do Not Proceed

**IRB of Record** (If multiple IRBs of record, please justify.): Click or tap here to enter text.

**IRB Application Type**
- □ Determination of Human Subjects
- □ Record Review of Existing Specimens, Retrospective
- □ Investigator Initiated, Prospective

**IRB Deadline(s)** (Include pre-review deadlines, if available): Click or tap here to enter text.

**Specimen Required** (select all that apply)
- □ Serum
- □ Plasma
- □ DNA
- □ Tissue
- □ Archival
- □ Fresh
- □ None
- □ Other, please describe: Click or tap here to enter text.

**Subject Characteristics** (Describe the characteristics of the subjects. Criteria might include disease status, gender, age, race, ethnicity, comorbidities, treatments, date ranges, survival status, etc. Be as specific as possible.)
Click or tap here to enter text.

**Number of Subjects:** Click or tap here to enter text.

**Do you require serial specimen?**
- □ Yes, please describe: Click or tap here to enter text.
- □ No