Program Guidelines for the National Cancer Institute (NCI) Cancer Clinical Investigator Team Leadership Award (CCITLA) P30 Administrative Supplement for Fiscal Year (FY) 2020

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Please note the following modifications from the FY 2019 administrative supplement guidelines:
  • (Page 7) The components of the Qualifications and Career Plan have been expanded.
  • (Page 7) For each planned activity, include the objective/expected outcome.
  • (Page 11) The review criteria have been expanded.
  • (Page 13) Industrial trials are no longer included in the Clinical Trials Table.

Key Dates

Letter of Intent Due Date: November 18, 2019

Application Due Date: by 5:00 PM local time of applicant organization on December 9, 2019

Earliest Start Date: March 2019 for cancer centers with P30 Cancer Center Support Grants (CCSGs) with start dates of December through March.
For cancer centers with P30 CCSGs with start dates of April through September, the anticipated start date is the 2020 start date of the parent P30 CCSG.

Award Information

The Cancer Clinical Investigator Team Leadership Award (CCITLA) recognizes and supports outstanding clinical investigators who enable their cancer center’s NCI-funded clinical trials enterprise. It is the intent of the CCITLA to support mid-level clinical
investigators at NCI-designated Cancer Centers who actively enroll patients and participate extensively and collaboratively in **NCI-funded clinical trials**. This award is intended to recognize and support efforts related to interventional trials. The CCITLA is designed to promote the retention of investigators in academic clinical research careers by providing salary support to individuals who do not have independent research grant funding as a Principal Investigator.

A candidate for the CCITLA must be nominated by the Cancer Center Director. An ideal candidate will have played a critical role in the development of NCI-funded clinical trials at the cancer center and have a record of service on institutional clinical trial committees and in NCI clinical trial-related activities.

A clinical trial is defined as a research study in which one or more human subjects are prospectively assigned to groups that receive one or more interventions (which may include placebo or other control) to evaluate the effects of those interventions on health-related biomedical or behavioral outcomes ([https://grants.nih.gov/policy/clinical-trials/definition.htm](https://grants.nih.gov/policy/clinical-trials/definition.htm); [https://clinicaltrials.gov/ct2/about-studies/glossary](https://clinicaltrials.gov/ct2/about-studies/glossary)). Clinical trials test new methods of screening, prevention, diagnosis, or treatment of a disease ([https://www.cancer.gov/publications/dictionaries/cancer-terms](https://www.cancer.gov/publications/dictionaries/cancer-terms)).

An intervention is defined as a treatment, procedure, or other action taken to prevent or treat disease, or improve health in other ways. Interventions include drugs, vaccines, medical procedures (such as radiation therapy and surgery), medical devices, behavior changes (such as diet and exercise), education programs and counseling ([https://www.cancer.gov/publications/dictionaries/cancer-terms](https://www.cancer.gov/publications/dictionaries/cancer-terms)).

**Funds Available and Allowable Costs**

The NCI intends to provide partial salary support for clinical investigators at up to 10 NCI-designated Cancer Centers through administrative supplements to P30 CCSGs. The total supplemental budget should not exceed $60,000 (total costs) per year for a total of two years, including salary, fringe benefits and associated facilities and administrative costs.

The candidate must devote at least 15% (1.8 calendar months) effort to the activities associated with this award and the sponsoring institution must protect the awardee’s time for these activities. Although cost sharing is not required, institutions are encouraged to cost share if needed to attain greater than 15% effort for the candidate.

All awards are subject to the terms and conditions of the CCSG notice of grant award and cost principles and other considerations described in the NIH Grants Policy Statement ([http://grants.nih.gov/grants/policy/nihgps/index.htm](http://grants.nih.gov/grants/policy/nihgps/index.htm)).

Support provided under this supplemental award is not transferable to another investigator or institution.
Allowable costs are limited to:

- Salary (for candidate only), fringe benefits and associated facilities and administrative costs.

- Travel (up to $2500/year) and registration fees (up to $2500/year) (for candidate only) to attend courses, seminars, meetings, conferences and workshops that support the intent of this award. In the budget justification, include the destination, dates and duration of stay for all anticipated travel. It is important to clearly state how the travel directly relates to the intent of this award.

Funds from this award may not be used for:

- Research-related costs, including but not limited to research supplies, computers, equipment, core facility fees, or sample or data analysis,

- Salary for personnel other than the candidate,

- Secretarial or administrative assistance and supplies.

Questions about allowable costs should be directed to the NCI Coordinating Center for Clinical Trials (CCCT) CCITLA program at NCICCCITLA@mail.nih.gov.

Nomination

The candidate must be nominated by the Cancer Center Director (Principal Investigator (PI) of the P30 CCSG) based on the candidate’s qualifications, involvement in NCI-funded clinical trials, support of the cancer center’s NCI clinical trials infrastructure, interests, accomplishments, motivation and plans to pursue a career in academic clinical research. Individuals who have previously received this award may not be nominated. For a list of past awardees, please refer to: https://www.cancer.gov/about-nci/organization/ccct/funding/ccitla.

Number of Applications

Each eligible NCI-designated Cancer Center may submit only one application.

Eligibility Criteria for Institutions

- Only NCI-designated Cancer Centers participating in NCI-funded clinical trials are eligible to apply for this supplement.
Not Eligible:

- The most recent CCITLA recipient Cancer Centers (those which received the first payment of a two-year CCITLA supplement in 2019) are not eligible to submit an application in response to this announcement.

- Cancer Centers which will enter an extension year during the fiscal year in which this award will be made (FY 2020) are not eligible to apply for the FY 2020 supplement since new administrative supplements cannot be paid to grants in an extension year.

Eligibility Criteria for Clinical Investigator Candidates

All criteria must be met at the time of the application submission deadline (December 9, 2019) to be considered for the award.

- The candidate must be one of the following:
  - Physician (e.g., M.D., D.O.), board certified or have equivalent training and qualifications in specialty area, e.g., medical oncology, radiation oncology, surgical oncology, gynecologic oncology, or equivalent
  - Oncology nurse or clinical psychologist (or similarly qualified clinician) with a doctoral degree

- The candidate must currently be practicing in the oncology clinical setting.

- The candidate must be a full-time faculty member (academic or clinical track), eligible for promotion, and either:
  1) an assistant professor for at least 3 but no more than 8 years from initial assistant professor appointment (including appointments held at other institutes) or
  2) an associate professor for no more than 8 years from initial appointment as an assistant professor (including appointments held at other institutes). **A full professor, or anyone above the associate professor level, regardless of tenure status, is not eligible to be nominated.**

- The candidate must be engaged in the conduct of NCI-funded cancer clinical trials at an academic medical center and have a record of involvement in cancer center clinical trial-related activities.

- The candidate must have potential for leadership of the Cancer Center’s clinical trials infrastructure activities. For example: setting clinical research priorities, overseeing clinical trials, submitting protocols to the Institutional Review Board (IRB), monitoring adverse event reporting and increasing enrollment in NCI-funded clinical trials.

- The candidate must be a United States (US) citizen or noncitizen national of the
US possessing a US passport that delineates and certifies status as a national but not a citizen of the US or have been lawfully admitted for permanent residence and possess a currently valid Permanent Resident Card (USCIS Form I-551) or other legal verification of such status.

- Candidate cannot have received the CCITLA previously (applies to the candidate only, not to the Cancer Center).

- The candidate may not currently be or previously have been:
  - A Principal Investigator (PI) of a National Institutes of Health (NIH) peer-reviewed research grant of at least $125,000 in direct costs per year for a minimum of 3 years, with the exception of mentored career development awards or awards where the PI is required to be mentored by another investigator.
    - Individuals who are currently supported by NIH Mentored Career Development (K) Awards are eligible to be nominated for the CCITLA if the support period for the K grant will end by the date when the CCITLA award will be made (approximately March 2020 for centers with annual P30 CCSG start dates in December through March; for centers with P30 CCSG start dates in April – September, the CCITLA is awarded on the start date). Since the CCITLA is an administrative supplement and not a research grant, it cannot replace effort on a K award (https://grants.nih.gov/grants/guide/notice-files/NOT-OD-08-065.html).
  - A project leader or co-leader on a NIH competing multi-component research or center grant or cooperative agreement within a P or U series grant (e.g., P01 Program Project Grant, U01 or U19 Research Program Cooperative Agreement, P50 Specialized Center Grant).
  - A Principal Investigator of a peer-reviewed research grant of at least $125,000 in direct costs per year for a minimum of 3 years from any of the organizations listed in the Organizations with Peer Review Funding Systems at: https://cancercenters.cancer.gov/PoliciesResources/PoliciesResources

- Recipients of ASCO Career Development Awards or mentored career development awards/grants from other organizations ARE eligible and may be nominated for this award.

Questions about eligibility should be directed to the NCI CCCT CCITLA program at NCICCITLA@mail.nih.gov.
Tips for Applicants

- The application should clearly show that the candidate is involved in and actively enrolling patients in NCI-funded clinical trials.
- The application should convey how the candidate is critical to the cancer center’s NCI-funded clinical trials enterprise.
- The application should highlight how this award would permit the candidate to expand current activities and/or develop/engage in new activities related to promoting successful clinical trials.
- The application should address the review criteria in the announcement.

Application (please include the following in the application in the order listed):

1. **Cover Letter** (1 page) signed by the PI of the P30 CCSG:
   A cover letter, signed by the PI of the P30 CCSG, must accompany each application and should only include:
   - the title of this funding opportunity
   - that the center is applying for this administrative supplement
   - the name and email address of the candidate and of the administrative official for the application
   - a statement verifying that the candidate meets all eligibility criteria of the award
   - the process used to select the candidate
   
   The Cancer Center Director’s letter of support should be included with the other letters of support for the application and not as part of the cover letter.

2. **Body of the Application:**
   
   The application should include (in the order listed):
   
   1) **Face page:** The standard face page of the PHS 398 (http://grants.nih.gov/grants/funding/phs398/phs398.html).

   2) **Biographical Sketch:** A biographical sketch of the candidate. The Biographical Sketch Format Page is available at http://grants.nih.gov/grants/forms/biosketch.htm.
   - In Section D. Research Support, list all ongoing, completed and pending research projects (Federally or non-Federally-supported), including the grant #, name of the grant PI, start and end dates of support, funding amount of the grant, title of the grant, goal of the study and the role of the CCITLA candidate in the project (e.g., Principal Investigator/co-Principal Investigator, co-Investigator). For any NIH P and U series grants, indicate that the candidate is/was not a project leader or co-leader of a research project within the grant.
3) **Clinical Trials Table:** Please refer to Appendix A. Instructions and Template for Clinical Trials Table.

4) **Qualifications and Career Plan:** Please address the following (no longer than 6 pages):

   a. **Suitability:** Describe how the candidate meets the intent of the CCITLA.
   
   b. **Training:** Describe specific aspects of the candidate’s training which support a team leadership role in oncology clinical trials.
   
   c. **NCI-Funded Clinical Trials:** Describe the candidate’s current and past involvement in NCI-funded clinical trials.
   
   d. **NCI Trial Network(s):** Describe the candidate’s participation in NCI trial networks, including but not limited to, the National Clinical Trials Network (NCTN), NCI Community Oncology Research Program (NCORP), Experimental Therapeutics Clinical Trials Network (ETCTN) and Cancer Immunotherapy Trials Network (CITN).
   
   e. **Cancer Center Committee(s):** Describe the candidate’s role(s) in cancer center committees such as the IRB, Data Safety Monitoring Board (DSMB), Protocol Review and Monitoring Committee (PRMC) and scientific review committees.
   
   f. **Clinical Trial-Related Activities:** Describe the candidate’s current and past clinical trial-related activities not described above.
   
   g. **Mentoring:** Describe the candidate’s current and past mentoring of trainees, junior investigators, students, etc. in support of clinical trial activities.
   
   h. **Collaborative Team Experience:** Describe the candidate’s collaborative team experience within division/department, across the institution, with individuals at other institutions and/or with outside organizations.
   
   i. **Career Plan:** Describe the candidate’s plans for a career in academic clinical research and how the institution is supporting the candidate in this plan.

5) **Planned Activities** (no longer than 2 pages)

   CCITLA activities should support, improve and/or expand the cancer center’s NCI-funded clinical trials and/or clinical trials infrastructure activities.
   
   a. An outline and description of activities planned under this award, including a timeline. For each planned activity include the objective/expected outcome.
b. Describe how the activities planned under this award build upon the experience and leadership of the candidate beyond merely a continuation of the regular activities.

c. Describe how the candidate’s regular workload will be adjusted to allow sufficient time to dedicate to the proposed CCITLA activities.

The CCITLA can support multiple projects/activities as time, effort and resources allow. Examples of projects and activities considered appropriate to this award include, but are not limited to (Note: projects/activities listed below are not ordered by priority):

- Designing and implementing initiatives to improve/expand awareness of, efficiency of and/or accrual to NCI-funded clinical trials.
- Developing streamlined processes for cancer center committees (e.g., IRB, DSMB, PRMC, scientific review committees).
- Organizing clinical trial-related courses, lecture/seminar series, educational sessions or workshops for clinical research staff, patient advocates, patients and other stakeholders.
- Participating on a cancer center committee (e.g., Institutional Review Board (IRB)) that enhances the awardee’s clinical trials knowledge or leadership.
- Mentoring fellows and new faculty in clinical trial efforts at the awardee’s institution.

6) **Budget:** A budget entered on Form Pages 4 and 5 of PHS 398 (http://grants.nih.gov/grants/funding/phs398/phs398.html) for the calendar months of effort for the Clinical Investigator during the first and second year, with appropriate justification. If the P30 CCSG current period of performance will end prior to year 2 of the CCITLA, include a projected budget for year 2.

**FUNDS REQUESTED FOR TRAVEL:** In the budget justification, include the destination, dates or duration of stay for all proposed travel and clearly state how the travel is related to the intent of the CCITLA.

7) **Checklist:** The Checklist Form Page of the PHS 398 (http://grants.nih.gov/grants/funding/phs398/phs398.html).

8) **Letters of Support:** Three signed letters of support on behalf of an individual’s application must be submitted with the application. The letters should be appended to the application after the PHS 398 Checklist in the PDF of the application.
   - One of the letters of support must be provided by the Cancer Center Director and should be appended to the application with the other letters of support, rather than included in the cover letter.
• At least one of the letters of support should be an institutional support letter from the Department Chair or appropriate institutional official that indicates:
  o A description of the academic status of the applicant and any additional support provided by the institution.
  o A clear level of commitment by the institution to developing the candidate’s career as an academic clinical investigator.
  o The extent to which the candidate will have dedicated time for activities proposed in the application. This letter must demonstrate a commitment to allow at least 15% (1.8 calendar months) effort for activities proposed in the application.
  o How the institution intends to continue to provide or augment its support for the candidate’s clinical research efforts beyond the award performance period.

Letter of Intent to Apply

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 NCI staff to estimate the potential review workload and plan the review.

Prospective cancer center applicants are asked to submit a letter of intent that includes the following information:

• title of this funding opportunity

• intent to apply for this administrative supplement

• name of the cancer center and Center Director (PI of the P30 CCSG)

• not required, but please include if known:
  o the name and email address of the Candidate
  o the names and institutional affiliations of the two individuals other than the Center Director who will provide letters of support for the application.

The letter of Intent should be provided by e-mail no later than November 18, 2019 to

CCITLA Program                   Ms. Nga Nguyen
Coordinating Center for Clinical Trials Program Analyst, Office of Cancer Centers
National Cancer Institute          National Cancer Institute
NCICCITLA@mail.nih.gov             nguyenn2@mail.nih.gov
Where to Send the Cover Letter and Application

Applications are due no later than 5:00 PM local time of applicant organization on December 9, 2019.

Email an electronic copy of the application in PDF format, including the cover letter and letters of support, to both contacts listed below.

CCITLA Program Coordinating Center for Clinical Trials
National Cancer Institute National Cancer Institute
NCICCTLA@mail.nih.gov nguyenn2@mail.nih.gov

Reporting Requirements: A progress report for the CCITLA supplement must be included as a separately labeled section in the annual progress report for the CCSG for any reporting period for which CCITLA supplemental funds are received.

The progress report should include:

- Details on the progress and outcome of each activity/project listed in the application.
- Awards and honors received during the performance period related to activities under this award.
- Publications, journal articles and patents related to this award.
- Impact to date of the award on the candidate’s career development.
- Opportunities that otherwise would not have been possible without the award.
- Value added by the award (e.g., for the institution or other staff).

Publications resulting from this award should acknowledge the funding source as follows: “This study was supported in whole or in part by funding from the Cancer Clinical Investigator Team Leadership Award awarded by the National Cancer Institute Coordinating Center for Clinical Trials (CCCT) though a supplement to P30 xxxxxxx.”

Publications, journal articles and/or patents produced under an NIH award-supported project must bear an acknowledgment and disclaimer, as appropriate, as provided in NIH Grants Policy Statement Part II: Terms and Conditions of NIH Grant Awards Section 8.2 “Administrative Requirements—Availability of Research Results: Publications, Intellectual Property Rights and Sharing Research Resources.”
Review Criteria

There is no predetermined weighting for the categories of review criteria. Bulleted items in each category serve as examples for addressing review criteria. An application does not need to be strong in all areas to receive a meritorious assessment.

Suitability

- Does the candidate meet the intent of the award?
  - Actively engaged with and enrolling patients in NCI-funded clinical trials
  - Plays a critical role in the cancer center’s NCI-funded clinical trials infrastructure activities
  - Record of service in institutional clinical trial-related activities

Training

- Does the candidate have training and experience strongly supporting a team leadership role in oncology clinical trials?

Involvement in NCI-funded clinical trials

- To what extent has the candidate actively participated (currently and in the past) in NCI-funded clinical trials such as those funded through the Division of Cancer Treatment and Diagnosis (DCTD), Division of Cancer Prevention (DCP), Division of Cancer Control and Population Sciences (DCCPS), Office of the Director (OD) or NCI-designated Cancer Centers?

Experience and clinical trial-related activities

- To what extent does the candidate have roles in:
  - Cancer center committees
  - NCI trial networks
  - Other clinical trial-related activities

- To what extent does the candidate mentor or guide trainees, junior investigators, students, etc. in support of clinical trial activities?

Collaborative team experience

- What is the extent of the candidate’s collaborative team experience?

Candidate’s planned activities

- Do the proposed activities meet the intent of the award? CCITLA activities should support, improve and/or expand the cancer center’s NCI-funded clinical trials and/or clinical trials infrastructure activities.
• Are the proposed activities new and/or an expansion of regular activities rather than a continuation of regular activities?

• Do the planned activities build upon the experience and leadership of the candidate beyond merely a continuation of the regular activities?

• Will this award allow the candidate to develop/engage in new activities related to clinical trials at his/her institution that would not likely happen without this award?

• To what extent do the proposed activities promote and/or enhance clinical trials and clinical trial-related activities at the candidate’s institution?

• Does the application indicate appropriate commitment of time and effort for the proposed activities?

• Will the candidate be able to devote sufficient time to both the CCITLA activities and non-CCITLA activities/duties/responsibilities?

Institutional commitment to support the candidate’s planned activities and career in clinical research

• Is there clear commitment of the institution to relieve the candidate of sufficient duties to allow at least 15% (1.8 calendar months) effort for activities proposed in the application?

• Is the level of institutional commitment to the career development of the candidate appropriate to be considered for this award?

• Does the candidate’s institution intend to continue to provide or augment its support for the candidate’s efforts in NCI-funded clinical trials and clinical trials infrastructure activities beyond the award performance period?

NCI Contact Information

For programmatic questions concerning this supplement, contact the NCI CCITLA program in the Coordinating Center for Clinical Trials (NCICITLA@mail.nih.gov) or the NCI Program Director assigned to your P30 CCSG.

Questions regarding fiscal and administrative matters should be addressed to the Grants Specialist for your Cancer Center, NCI Office of Grants Administration.
**Appendix A. Instructions and Template for Clinical Trials Table**

Using the template in this Appendix, list all national, externally-peer reviewed and institutional clinical trials which the candidate has had a role in between December 1, 2016 – November 30, 2019 and which are currently active, closed, approved or under review. Do not include industrial trials in the table. **The table must be in the format of the table in this appendix.** Use legal size paper for the table if needed.

Sort the trials by Study Source:

- **National:** NCI National Clinical Trials Network (NCTN) and other NIH-supported National Trial Networks
- **Externally Peer-Reviewed:** R01s, SPORES, U01s, U10s, P01s, CTEP, or any other clinical research study mechanism supported by the NIH or an organization listed in the Organizations with Peer Review Funding Systems at: https://cancercenters.cancer.gov/PoliciesResources/PoliciesResources
- **Institutional:** In-house clinical research studies authored or co-authored by cancer center investigators and undergoing scientific peer-review solely by the Protocol Review and Monitoring System of the cancer center. The cancer center investigator has primary responsibility for conceptualizing, designing and implementing the clinical research study and reporting results.

**Specific Funding Source:** The specific name of the financial sponsor(s) for the clinical trial. For institutionally sponsored trials, list the name of the applicable funding agencies.

**NCT ID:** Provide the unique ID assigned to the trial by the National Clinical Trial program (ClinicalTrials.gov) for trials that have been submitted to ClinicalTrials.gov Protocol Registration System (PRS) previously. The ClinicalTrials.gov ID appears as "NCT" followed by 8 numeric characters (such as NCT12345678). In the table, do not include “NCT” in the ID (e.g., for NCT12345678, only list “12345678” in the column). If not applicable, enter “N/A” in this column.

**Other ID(s):** Provide the unique identifier for this trial. Where available, list the common protocol number that the trial is known under nationally, if one exists. For other trials that do not have a common protocol number that the trial is known under nationally, use an internal protocol identification or IRB number.

**Candidate's Role:** The role of the CCITLA candidate in the trial (e.g., 'Overall PI', 'Overall Co-PI', ‘Site PI’, ‘Associate Investigator’, etc.)

**Date Open: Site (activation):** The date the trial opened at your center.

**Date Closed:** The date the trial closed to accrual at your center. This does not include patient follow-up. If the trial is still open, leave this field blank.

**Status (as of November 30, 2019):** Under Review, Approved, Active, Closed - Not Yet Published, Closed, Enrolling by Invitation, Temporarily Closed to Accrual, Temporarily Closed to Accrual and Intervention, or Temporarily Not Available
**Phase:** For Interventional trials acceptable phases include: pilot, feasibility, 0, I, II, III, IV, or combinations such as I/II. For trials without phases, indicate “N/A.”

**Official Title:** Official name of the protocol provided by the trial PI or sponsor (Limit: 8000 characters or fewer).

**Multi-Inst Trial:** Indicate if the trial is multi-institutional by inserting ‘Y’ or ‘N’ in the “Multi-inst trial” column.

Multi-Institutional Clinical Trial: Clinical trial that recruits participants from two or more geographically distinct enrollment Institutions not affiliated with your cancer center (e.g., other NCI-designated Cancer Centers or other research institutions). The Institutions are usually distinct in other characteristics (e.g., demographic, socioeconomic, or clinical).

**Total Targeted Accrual:**
- **Entire Trial:** For both single-institution and multi-institutional trials initiated at your center, indicate the total number of participants needed for the entire trial. For multi-Institutional trials that your center participates in but did not initiate, leave “Entire Trial” column empty if not known. Do not submit a targeted range, such as “10 – 100.”
- **Your Center:** For single-institution and multi-institutional trials initiated at your center, indicate the total number of participants your center is expected to accrue for the study. For single-institution trials the “Total Accrual for Your Center” and the “Total Targeted Accrual Entire Trial” numbers will be the same. Do not submit a targeted range, such as “10 – 100.”

**Center Accrual:** List the number of participants enrolled in the clinical trial at your cancer center, including formal Consortium Partners.
- **12 Months:** Provide the number of participants accrued to this clinical trial during the last 12 months.
- **To Date:** Provide the number of participants accrued to this clinical trial since the trial was opened.

**Enrolled on Trial by Candidate:** List the number of participants which the candidate has enrolled in the clinical trial.
- **12 Months:** Provide the number of participants accrued to this clinical trial during the last 12 months.
- **To Date:** Provide the number of participants accrued to this clinical trial since the trial was opened.
## TEMPLATE FOR CLINICAL TRIALS TABLE

Clinical Trials Table  
Candidate: [Name of Candidate], [Name of cancer center]  
Reporting Period: 12/01/2016 – 11/30/2019

### NATIONAL Trials

<table>
<thead>
<tr>
<th>Specific Funding Source</th>
<th>NCT ID</th>
<th>Other ID(s)</th>
<th>Candidate’s Role</th>
<th>Date Open: Site</th>
<th>Date Closed: Site</th>
<th>Status</th>
<th>Phase</th>
<th>Official Title</th>
<th>Multi-Inst Trial?</th>
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### EXTERNALLY PEER-REVIEWED Trials

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<th>Specific Funding Source</th>
<th>NCT ID</th>
<th>Other ID(s)</th>
<th>Candidate’s Role</th>
<th>Date Open: Site</th>
<th>Date Closed: Site</th>
<th>Status</th>
<th>Phase</th>
<th>Official Title</th>
<th>Multi-Inst Trial?</th>
<th>Entire Trial</th>
<th>Your Center</th>
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Clinical Trials Table  
Candidate: [Name of Candidate], [Name of cancer center]  
Reporting Period: 12/01/2016 – 11/30/2019

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<th>NCT ID</th>
<th>Other ID(s)</th>
<th>Candidate’s Role</th>
<th>Date Open: Site</th>
<th>Date Closed: Site</th>
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<th>Official Title</th>
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