Program Announcement

I. GENERAL INFORMATION

This program announcement is being released before the receipt of Federal funds appropriated in a bill for this program; funding of proposals received in response to this program announcement is contingent on the receipt of these funds at the United States Army Medical Research and Materiel Command (USAMRMC).

A. Title of Award: Tuberous Sclerosis Complex Research Program (TSCRP) Clinical Resource Development Award.

B. Program Name: Department of Defense (DOD) Fiscal Year 2006 (FY06) TSCRP.

C. Funding Opportunity Number: W81XWH-06-TSCRP-CRDA.

D. Agency Name: USAMRMC, Office of the Congressionally Directed Medical Research Programs (CDMRP), 1077 Patchel Street, Fort Detrick, Maryland 21702-5024.

E. Agency Contact(s)

1. Questions related to the program announcement, proposal format, or required documentation: Applicants should submit questions as early as possible. Every effort will be made to answer questions within 5 working days.

   Phone: 301-619-7079
   Fax: 301-619-7792
   E-mail: cdmrp.pa@amedd.army.mil
   Mail: Commander
   US Army Medical Research and Materiel Command
   ATTN: MCMR-ZB-C (TS06-CRDA)
   1077 Patchel Street (Building 1077)
   Fort Detrick, MD 21702-5024

2. Questions related to electronic submission: A help line for questions relating to proposal submission and the CDMRP eReceipt Online Proposal Submission System is available Monday through Friday from 8:00 a.m. to 5:00 p.m. Eastern time at 301-682-5507. Help also is available on the CDMRP website or by e-mail as follows:

   Website: https://cdmrp.org (User’s Guide located in upper right corner of the proposal submission website)
   E-mail: help@cdmrp.org
F. Anticipated Instrument Type(s): The USAMRMC executes its extramural research program predominantly through the award of grants and cooperative agreements. More information on these funding instruments may be obtained by request from:

Fax: 301-619-2937
E-mail: qa.baa@amedd.army.mil
Mail: Director
US Army Medical Research Acquisition Activity
ATTN: MCMR-AAA-R
820 Chandler Street
Fort Detrick, MD 21702-5014

G. Catalog of Federal Domestic Assistance (CFDA) Number 12.420: Military Medical Research and Development.

H. Website to Access Application Package: Proposals must be submitted at https://cdmrp.org. This website contains all the information, forms, documents, and links needed to apply. Applicants experiencing difficulty in downloading documents should contact the CDMRP as indicated in Subsection I.E.2.

I. Award/Regulatory Approval: The applicant may not use, employ, or subcontract for the use of any human subjects, human biological substances, cadavers, or laboratory animals until applicable regulatory documents are requested, reviewed, and approved by the USAMRMC.

II. FUNDING OPPORTUNITY DESCRIPTION

A. Program History: The Clinical Resource Development Award is one of the mechanisms of the Tuberous Sclerosis Complex Research Program (TSCRCP), which was established in Fiscal Year 2002 (FY02) to promote innovative research directed toward improved diagnosis or treatment of tuberous sclerosis complex (TSC). Appropriations for the TSCRCP from FY02 through FY05 totaled $9.2 million (M). The Clinical Resource Development Award is being offered for the first time in FY06. The FY06 appropriation is $4.3M.

B. Program Objectives: The overall goal of the FY06 TSCRCP is to decrease the impact of tuberous sclerosis complex. Within this context, the encouragement of established scientists in the field and the attraction of new scientific expertise from other fields are essential to the tuberous sclerosis complex community. Proposals that address the needs of minority, low-income, rural, and other underrepresented and/or medically underserved populations are encouraged and may be submitted from any eligible institutional source.

C. Award Mechanism Description: The TSCRCP recognizes that the successful design, implementation, and interpretation of TSC-relevant therapeutic clinical trials is dependent upon the availability of standardized tools for assessing TSC symptom severity or changes over time. Consequently, the Clinical Resource Development Award has been established to fund the development, validation, and/or pilot testing of novel tools/resources for measuring clinical
and/or surrogate endpoints that are relevant to the variety of lesions and clinical manifestations seen in individuals with TSC. Due to the developmental nature of this award, preliminary data are not required but may be included, if available, to address the feasibility of the tool/resource to be developed. Applicants must explain clearly how the proposed products are directly relevant and applicable to future clinical trials for individuals with TSC. It is anticipated that projects involving human subjects will be most relevant for this mechanism.

Examples of tools/resources eligible for funding under this award include but are not limited to:

- Magnetic resonance imaging-based or other protocols for measuring tumor volumes in affected organs,
- Biomarkers of cognitive function,
- TSC-specific behavioral assessments, and
- Protocols/methods for quantitative assessment of seizure severity.

A clinical protocol and associated clinical documents must be included with the submission if the applicant plans to use funding from this award to support clinical studies with human subjects. It is expected that any proposed clinical studies will be initiated within 6 months of the award date. Please note that all DOD-funded research involving human subjects, human anatomical substances, cadavers, and/or laboratory animals must be reviewed and approved by the USAMRMC Human Subjects Research Review Board (HSRRB) in addition to local Institutional Review Boards (IRBs). The HSRRB has different requirements than the local IRBs. The average time to obtain HSRRB approval is approximately 6 months. Therefore, it is strongly suggested that the applicant plan the budget and timeline accordingly.

IRB approvals should be in process or completed before proposal submission. Applicants must submit any available internal scientific and local IRB review documents for the clinical protocol and Informed Consent/Assent form(s) that indicate the status of IRB review achieved prior to submission of the Clinical Resource Development Award. Indicate the highest possible level of IRB review within the institution(s) before submission of a proposal. Documentation of local IRB review and approval will be required at the time of the HSRRB regulatory review for all funded proposals.

All protocols must be prepared according to the guidelines provided in the document titled “Research Involving Human Subjects and/or Anatomical Substances,” which can be found at https://cdmrp.org/Program_Announcements_and_Forms under “Regulatory Document Forms.” Separate freestanding protocols should be generated for each site and/or task for all research that involves (a) multiple sites; (b) multiple tasks requiring different study designs, samples, sampling procedures, or methods; and (c) inter-institutional cooperative or collaborative agreements. Please note that each project of a multi-project application will be individually evaluated for compliance with applicable regulatory requirements.

Proposals involving human subjects research that do not include a clinical protocol and Informed Consent/Assent forms will be administratively withdrawn and will not be reviewed.
Laboratory or animal studies will be considered for funding under this mechanism if the applicability to TSC clinical trials is clear. However, these proposals will face a greater burden to demonstrate clinical relevance.

All proposals must include a plan describing the means by which the fully developed tool/resource will be made available to the scientific community for future TSC clinical studies and trials at reasonable or appropriate administrative costs. The Government intends to advertise the tool/resource developed through this award and to provide information on the CDMRP website as to how to obtain access to the product for TSC-relevant clinical research.

III. AWARD INFORMATION

Funding for a Clinical Resource Development Award can be requested for a maximum of $225,000 for direct costs over the performance period. The performance period can be requested for up to 2 years. Indirect costs should be added as appropriate. Direct costs can cover salary, research supplies, and travel to scientific/technical meetings. Although not required, multi-institutional and multidisciplinary research collaborations are encouraged. Consideration of cost-sharing with other funding sources is encouraged. The nature of the TSCRP does not allow for renewal of grants or supplementation of existing grants.

*The CDMRP expects to allot approximately $0.8M of the $4.3M FY06 TSCRP appropriation to fund approximately two to three Clinical Resource Development Awards, depending on the quality and number of proposals received.*

IV. ELIGIBILITY INFORMATION

A. Applicants: Investigators at all academic levels (or equivalent) are eligible to submit proposals.

All individuals, regardless of ethnicity, nationality, or citizenship status, may apply as long as they are employed by, or affiliated with, an eligible institution as defined in Subsection IV.B, “Institutions” below.

To protect the public interest, the Federal Government ensures the integrity of Federal programs by only conducting business with responsible recipients. The USAMRMC uses the Excluded Parties List System (EPLS) to exclude recipients ineligible to receive Federal awards. The EPLS is online at [http://epls.arnet.gov](http://epls.arnet.gov). (Reference Department of Defense Grant and Agreement Regulations (DODGAR) 25.110.)

B. Institutions: Eligible institutions include for-profit, nonprofit, public, and private organizations, such as universities, colleges, hospitals, laboratories, and companies. The USAMRMC is especially interested in receiving applications from Historically Black Colleges and Universities/Minority Institutions (HBCU/MI).
A DOD goal is to allocate funds for the CDMRP peer-reviewed research to fund proposals from HBCU/MI. This provision is based on guidance from Executive Orders. Proposals are assigned HBCU/MI status when the submitting institution is so designated by the Department of Education on the date the program announcement is released. The most current Department of Education list is posted on the CDMRP website at [http://cdmrp.army.mil/spp](http://cdmrp.army.mil/spp) under “Minority Institutions.”

Local, state, and Federal Government agencies are eligible to the extent that proposals do not overlap with their fully funded intramural programs. Federal agencies are expected to explain how their proposals do not overlap with their intramural programs.

Proposals from Federal agencies **must** provide a plan delineating how all funds will be obligated by September 30, 2007, and how funds will be available to cover research costs over the entire award period. The plan must include the funding mechanism(s) that will be used to carry over funds between fiscal years, such as administrative agreements with foundations, non-Federal institutions, and universities.

C. Duplicate Submissions: Submission of the same research project to the FY06 TSCR to different award mechanisms or to other CDMRP programs is discouraged. The Government reserves the right to reject duplicative proposals.

V. PROPOSAL PREPARATION AND SUBMISSION INFORMATION

A. Proposal Components Summary: This subsection is a summary of submission requirements. Details, URLs, and other links are provided in the appropriate subsections of this program announcement. Proposals will be evaluated according to peer and programmatic review criteria in Section VI.

1. Applicant Responsibility: The applicant is responsible for entering and/or uploading the following information into the CDMRP eReceipt Online Proposal Submission System at [https://cdmrp.org](https://cdmrp.org):

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<th>Item</th>
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<tr>
<td>Letter of Intent (LOI)</td>
<td>Proposal Information</td>
<td>Typed</td>
<td>Copy the LOI into the data field. Click the “Save and Forward Letter of Intent” button to automatically create the LOI.</td>
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<tr>
<td>Proposal Information</td>
<td>Proposal Information</td>
<td>Typed</td>
<td>Enter the appropriate information in data fields.</td>
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1Executive Orders 12876, 12900, and 13021
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<tbody>
<tr>
<td>Proposal Contacts</td>
<td>Proposal Contacts</td>
<td>Typed</td>
<td>Enter contact information for the applicant and the Contract Representative at the applicant’s institution.</td>
</tr>
<tr>
<td>Collaborators and Conflicts of Interest (COI)</td>
<td>Collaborator/COI</td>
<td>Typed</td>
<td>Enter information about collaborators and others outside the scope of the proposal who may have a COI in the review of this proposal.</td>
</tr>
<tr>
<td>Proposal Main Body</td>
<td>Required Files</td>
<td>PDF</td>
<td>Upload as a PDF file.</td>
</tr>
<tr>
<td>Clinical Protocol (if applicable)</td>
<td>Required Files</td>
<td>PDF</td>
<td>Upload as a PDF file.</td>
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<tr>
<td>Supporting Documentation</td>
<td>Required Files</td>
<td>PDF</td>
<td>Upload as a PDF file.</td>
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<tr>
<td>Budget Information</td>
<td>Required Files</td>
<td>PDF</td>
<td>Upload as a PDF file.</td>
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<tr>
<td>Regulatory Documents</td>
<td>Required Files</td>
<td>PDF</td>
<td>Upload the Certificate of Environmental Compliance and the Principal Investigator Safety Program Assurance forms.</td>
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2. **Contract Representative Responsibility**: The Contract Representative (CR) or institutional official responsible for sponsored program administration (or equivalent) at the applicant’s institution is responsible for the following:

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</thead>
<tbody>
<tr>
<td>Contract Representative’s Contact Information Profile</td>
<td>My Profile for the CR</td>
<td>Typed</td>
<td>Complete before electronic approval of all submission components.</td>
</tr>
<tr>
<td>USAMRAA\textsuperscript{a}-Required Documents</td>
<td>My Profile for the CR</td>
<td>PDF</td>
<td>Upload the Rate Agreement, Certifications and Assurances for Assistance Agreements, and Representations for Assistance Agreements.</td>
</tr>
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\textsuperscript{a}USAMRAA: United States Army Medical Research and Development Activity.
B. Proposal Format: Proposals must be uploaded under the “Required Files” tab of the CDMRP eReceipt Online Proposal Submission System at https://cdmrp.org. Applicants unfamiliar with the preparation of PDF files are encouraged to acquire and learn to use the appropriate software well in advance of the submission deadline. The instructions in this subsection must be followed carefully to prepare proposals for PDF submission.

The main body of the proposal must be clear and legible and conform to the formatting guidelines described below. The font size, spacing, page size, and margins may differ between the word processing, PDF, and printed versions. These guidelines apply to the document properties of the electronic version of the PDF file(s) as viewed on the computer screen and submitted via the CDMRP eReceipt Online Proposal Submission System.

- **Font Size**: 12 point or larger.
- **Font Type**: Times New Roman is strongly recommended.
- **Spacing**: No more than six lines of type within a vertical inch (2.54 cm).
- **Page Size**: No larger than 8.5 inches x 11.0 inches (approximately 21.59 cm x 27.94 cm).
- **Margins**: Must be at least 0.5 inch (1.27 cm) in all directions.
- **Print Area**: 7.5 inches x 10.0 inches (approximately 19.05 cm x 25.40 cm).
- **Color, High-Resolution, and Multimedia Objects**: Proposals may include color, high-resolution, or multimedia objects (e.g., MPEG, WAV, or AVI files) embedded in the PDF files; however, these objects must not exceed 15 seconds in length and a size of 10 MB. Since some reviewers work from black and white printed copies, applicants may wish to include text in the proposal directing the reviewer to the electronic file for parts of the proposal that may be difficult to interpret when printed in black and white. Photographs and illustrations must be submitted in JPEG format; bit map or TIFF formats are not allowed.
- **Internet URLs**: URLs directing reviewers to websites containing significant additional information about the proposed research are not allowed in the proposal or its components. Inclusion of such URLs may be perceived as an attempt to gain an unfair competitive advantage. Links to publications referenced in the proposal are allowed.
- **Language**: English.
Please note that headers should not be included, as the proposal log number will be electronically captured on each page of the proposal after receipt.

C. Administrative Compliance Issues: Compliance guidelines have been designed to ensure the presentation of all proposals in an organized and easy-to-follow manner. Peer reviewers expect to see a consistent, prescribed format for each proposal. *Failure to adhere to format requirements makes proposals difficult to read, may be perceived as an attempt to gain an unfair competitive advantage, and may result in proposal rejection.*

The following will result in administrative rejection of the entire proposal before it reaches peer review:

- Proposal body exceeds page limit.
- Proposal body is missing.
- Detailed cost estimate is missing.
- Proposal is incomplete after the deadline.

For any sections of a proposal with a defined page limit, pages exceeding the specified limit will be removed from the proposal and not forwarded for peer review.

Material submitted after the submission deadline, unless specifically requested by the Government, will not be forwarded for peer review.

The electronic PDF file uploaded in the CDMRP eReceipt Online Proposal Submission System is the official proposal submission file. After conversion of word processing documents to PDF files and before electronic submission, applicants should review their files to ensure that the proposal complies with the preparation guidelines outlined in this program announcement.

D. Letter of Intent (LOI): An LOI (a brief description of the proposal) is entered in a data field under the “My Proposals: Create New Proposal” tab. The LOI is saved when the “Save and Forward Letter of Intent” button is chosen. The LOI may be modified under “Proposal Information” at anytime before the applicant submits this information by clicking “Finalize for CR Approval.” The LOI should be submitted by March 28, 2006 at [https://cdmrp.org](https://cdmrp.org).

E. Proposal Information: Applicants are required to submit the Proposal Information as described in [https://cdmrp.org](https://cdmrp.org) before uploading the proposal, supporting documentation, and budget information.

- A *Title/Referral Page* for the proposal will be generated from the information uploaded in eReceipt and appended to the proposal electronically by the CDMRP eReceipt system.

F. Proposal Contacts: The Proposal Contacts *must* include the e-mail address of a Contract Representative authorized to negotiate on behalf of the applicant’s institution. The Proposal Contacts must be approved by the Contract Representative at the applicant’s institution.
G. Collaborators and Conflicts of Interest (COI): To avoid COI during the review process, list the names of all scientific participants in the proposal including collaborators, consultants, and subawardees. In addition, list the names of individuals outside the scope of this proposal who may have a COI in reviewing this proposal.

H. Proposal Abstracts – 5,700-character limit including spaces (approximately one page), for each abstract: Each abstract must include the applicant’s name and the title of the proposal. A structured technical abstract and a public (nontechnical) abstract are required. These abstracts are vitally important in both the peer review and programmatic review processes. Programmatic review is based on the Integration Panel’s review of these two abstracts as part of the peer review summary statements; therefore, it is of paramount importance that the applicant submit abstracts that describe the proposed work fully. Each abstract must be entered into the appropriate data field under the “Abstract/Impact/SOW” tab of the CDMRP eReceipt system.

Applicants can type the abstracts or “cut and paste” them from a word processing application into the respective data fields. Spell out all Greek letters, other non-English letters, and symbols.

Abstracts of all funded proposals will be posted on the CDMRP website at http://cdmrp.army.mil. Proprietary or confidential information should not be included in either the technical or the public abstract.

1. Technical Abstract: Sample technical abstracts can be found at https://cdmrp.org/samples.cfm. The structured technical abstract must provide a clear and concise overview of the proposed work. Use the outline below when preparing the structured technical abstract.

   • Background: Present the ideas and reasoning behind the proposed tool/resource.
   • Tool/Resource Description: State the tool/resource to be developed. Provide evidence or rationale that supports the need for this product for TSC clinical trials.
   • Objective/Hypothesis: State the objective/hypothesis to be tested. Provide evidence or rationale that supports the objective/hypothesis.
   • Specific Aims: State the specific aims of the study and a plan for how the tool/resource will be developed, validated, and/or pilot tested.
   • Study Design: Briefly describe the study design including appropriate controls.
   • Impact: Provide a brief statement explaining how the project will make an original and important contribution to the field of TSC clinical research. Explain the direct relevance and applicability of the proposed tool or resource to the measurement of clinical and/or surrogate endpoints in TSC clinical trials.

2. Public Abstract: Sample public abstracts can be found at https://cdmrp.org/samples.cfm
The public abstract is intended to communicate the purpose and rationale of the study to a non-scientifically trained audience. The public abstract is an important component of the
proposal review process because consumer advocates, who are part of the review and funding decision process, use this abstract as a part of their review.

- Describe the scientific objective and rationale for the proposal in a manner readily understood by non-scientists.
  - Do not duplicate the technical abstract.
- Describe the ultimate applicability of the research.
  - What types of patients will it help and how will it help them?
  - What are the potential clinical applications, benefits, and risks?
  - What is the projected time it may take to achieve a consumer-related outcome?

I. Impact Statement – 5,700-character limit including spaces (approximately one page):
The Impact Statement is captured as a data field under the “Abstract/Impact/SOW” tab in the CDMRP eReceipt system. Applicants can type the Impact Statement into the data field or “cut and paste” it from a word processing application.

State explicitly how the proposed work will make an original and important contribution to the technologies for measuring TSC-relevant clinical or surrogate endpoints. Explain how the proposed tool or resource will have a significant and direct impact on the design, implementation, and interpretation of TSC-relevant therapeutic clinical trials. The Impact Statement, which will be available at both peer and programmatic reviews, is often cited by consumer advocates during the review and funding decision processes.

J. Statement of Work – 11,400-character limit including spaces (approximately two pages): The SOW is captured as a data field under the “Abstract/Impact/SOW” tab in the CDMRP eReceipt system. Applicants can type in the SOW into the data field or “cut and paste” it from a word processing application.

The SOW is a concise restatement of the research proposal that outlines, step by step, how each major goal or objective of the proposed research/services will be accomplished during the period for which the USAMRMC will provide financial support. When a proposal requesting funding as part of a larger study is submitted, the proposal’s SOW must include DOD-funded tasks only. Sample SOWs can be found at https://cdmrp.org/samples.cfm

The SOW should:

- Describe the work to be accomplished as tasks (tasks may relate to specific aims);
- Identify the timeline and milestones for the work over the performance period for the proposed effort;
  - Allow 4 to 6 months for regulatory review and approval processes for human use studies;
  - Allow 2 months for regulatory review and approval processes for animal studies;
- For animal and human studies (including tissue, anatomical, or biological substances), indicate the sample size projected or required for each task;
• Identify methods; and
• Identify outcomes, products, and deliverables for each phase of the project.

K. Proposal Main Body: Start section on a new page; four-page limit inclusive of any figures, tables, graphs, photographs, diagrams, chemical structures, pictures, pictorials, cartoons, and other information needed to judge the proposal.

The proposal main body is uploaded as a PDF file under the “Required Files” tab of the CDMRP eReceipt system.

It is the responsibility of the investigator to clearly articulate plans for developing the proposed resource or tool and to describe how the product will improve upon current approaches to the measurement of clinical and/or surrogate endpoints in TSC clinical trials. Due to the developmental nature of this award, preliminary data are not required but may be included, if available, to address the feasibility of the resource to be developed. In either case, proposals must apply sound scientific rationale and logical reasoning based on existing knowledge to the development of the proposed product.

Describe the proposed project using the outline provided below. The main body of the proposal will be reviewed as a standalone document. Therefore, if a clinical protocol is provided with the submission, include the appropriate information from the protocol to discuss the topics listed below. **Do not reference the clinical protocol.**

Describe the proposed project using the following outline:

1. **Background:** Present the ideas and reasoning behind the proposed work. Describe previous experience most pertinent to this proposal, including any available preliminary data. Cite relevant literature.

2. **Hypothesis or Objective:** State the hypothesis to be tested or the objective to be reached. Describe the tool or resource to be developed. Describe possible uses for the tool or resource in TSC clinical trials.

3. **Specific Aims:** Concisely explain the projects’ specific aims. If this proposal is part of a larger study, present only DOD-funded tasks.

4. **Research Strategy:** Describe the experimental design, methods, and analyses including appropriate controls in sufficient detail for analysis. Address potential problem areas and present alternative methods and approaches. If human studies are to be performed using funding from this award, provide a brief discussion of the following topics:

   • Study design, including clinical, behavioral, laboratory, and physiological tests and protocols.
   • Potential biases in the protocol and how they will be addressed.
• Participant recruitment, including (1) a description of the participant population; (2) sample size; (3) inclusion and exclusion criteria; (4) methods and schedules for recruiting, retention, and follow-up; (5) data to support recruitment/retention estimates; and (6) study endpoints.

• Data management and statistical plans.

• A study organization and management plan, including a timetable for completion of the study.

• Methods for the handling, distribution, banking, analysis, and security of specimens and/or imaging products. Include a plan for public dissemination of data generated by the study that addresses all relevant privacy issues.

• Any issues that may lead to concern for the welfare of human subjects and confidentiality, including a plan for addressing human subjects protection requirements as outlined by the HSRRB at https://cdmrp.org/Program_Announcements_and_Forms under “Regulatory Document Forms.”

• Internal scientific and local IRB reviews for the clinical protocol and Informed Consent/Assent form(s) at the highest possible level within the participating institutions, up to and including preliminary IRB approval, if available, at the institution(s).

5. **Innovation:** Describe concisely how the proposed research uses innovation to advance the detection, diagnosis, and/or treatment of TSC. Demonstrate how the proposed research represents more than an extension or incremental advance to published data.

6. **Impact:** Describe how the tool or resource will enhance the ability to assess TSC symptom severity or changes over time and ultimately improve the design, implementation, and/or interpretation of TSC therapeutic clinical trials.

7. **Resource Distribution Plan:** State concisely how the resource, once fully developed, will be made available to the scientific community at reasonable or appropriate administrative costs, such as costs required for packaging and shipping the resource.

*Please note that applicants planning to conduct human studies using funding from this award must submit a clinical protocol, Informed Consent forms, and other supporting clinical documents in the appropriate section of the proposal (see Subsections V.L and V.M.8). Ensure that the information describing the clinical protocol in the main body matches that in Subsection V.L. In addition, any available IRB approvals for this work must be submitted as supporting clinical documents (see Subsection V.M.8).*

L. **Clinical Protocol and Supporting Clinical Documents (if applicable):** No page limit. A clinical protocol and associated clinical documents must be included with the submission if the applicant plans to use funding from this award to support clinical studies with human subjects. *In order to facilitate the initiation of work on funded projects, the clinical protocol should be a standalone document that can be submitted to the HSRRB for review.*
Separate protocols should be generated for each site and/or task for all research that involves (a) multiple sites; (b) multiple tasks requiring different study designs, samples, sampling procedures, or methods; and (c) inter-institutional cooperative or collaborative agreements. Please note that each project of a multi-project application will be individually evaluated for compliance with applicable regulatory requirements. Documentation of local IRB review and approval of each protocol will be required at the time of HSRRB regulatory review of funded proposals.

It is critical that the information entered in the main body of the proposal matches the information contained within the clinical protocol. When a proposal is submitted requesting funding for part of a larger study, the clinical protocol must include DOD-funded tasks only.

Required elements for submission of a clinical protocol are:

1. **Protocol Title:** The protocol title must be the same as the proposal title. Please note that each proposal should contain only one clinical trial (clinical protocol) with a distinct study design.

2. **Phase:** Designate the protocol as Phase I or II.

3. **Applicant:** List the complete name, address, telephone and fax number, and e-mail address of the applicant.

   NOTE: Research investigators must complete appropriate institutional training before conducting human subjects research. Documentation of the most recent ethics training must be submitted for investigators of all protocols in the Supporting Documentation section of the proposal (Subsection V.M.8). In addition, for all investigational drug and device protocols, documentation of successful completion of a course in the conduct of clinical research in accordance with Good Clinical Practice (GCP) must be submitted for all investigators. The most recent ethics training and GCP course must be successfully completed within 1 year of the planned initiation of the protocol.

4. **Roles and Responsibilities of Study Personnel:** List the names of all personnel who will have significant involvement in the research study; include their practice license (e.g., MD or RN), highest degree(s), job title, and employing institution. Briefly describe the duties of all study personnel to include each of the persons listed as investigators, research staff, consultants, and the Medical Monitor. Describe their roles in the research effort (e.g., Research Coordinator, 80%, recruit and consent subjects, maintain study records, administer study drug, take and record vital signs, enter data into computer database). Include a named study coordinator who will be charged with guiding the protocol through the IRB, HSRRB, and other regulatory approval processes, coordinating activities from all sites participating in the trial, and coordinating participant accrual. In addition, include the name of the Medical Monitor with his or her current curriculum vitae for Greater Than Minimal Risk Studies. Duties of the Medical Monitor, as defined in HSRRB Clause 8.02, are as follows:

   “A Medical Monitor must be assigned to Greater Than Minimal Risk protocols. The name and curriculum vitae of the Medical Monitor, who is someone other than the PI [Principal Investigator], must be provided. This individual should be a qualified
physician who is not associated with the protocol, able to provide medical care to research subjects for conditions that may arise during the conduct of the study, and able to monitor subjects during the conduct of the study. In some studies it may be acceptable to have a qualified health care provider other than a physician serve as Medical Monitor, depending upon the type of risk that might occur in the study (e.g., a clinical psychologist). The Medical Monitor is required to review all unanticipated problems involving risk to volunteers or others, serious adverse events, and all volunteer deaths associated with the protocol and to provide an unbiased written report of the event. At a minimum the Medical Monitor should comment on the outcomes of the adverse event and relationship of the event to the protocol or test article. The Medical Monitor should also indicate whether he or she concurs with the details of the report provided by the PI. Reports for events determined by either the investigator or Medical Monitor to be possibly or definitely related to participation and reports of events resulting in death should be promptly forwarded to the HSRRB.”

The Medical Monitor will forward reports to the USAMRMC, ATTN: MCMR-ZB-P, 504 Scott Street, Fort Detrick, Maryland 21702-5012.

5. **Location of Study:** List all centers, clinics, or laboratories where the study is to be conducted. Include the name, degree(s), title, employing institution, and complete address of the investigator(s) for each site.

6. **Time Required to Complete the Study:** State the month and year of expected start and completion times.

7. **Background:** Suggested length – 10 pages. Describe the rationale for conducting the study citing any relevant literature references. Include descriptions of any preliminary studies and findings that led up to the development of the protocol and previous experience most pertinent to the proposed study. In addition, Phase II clinical trial applicants must provide Phase I or pilot clinical trial data. State the potential impact of the proposed clinical trial on the detection, diagnosis, or treatment of TSC. If the protocol was initiated using other findings prior to obtaining funding managed by the USAMRMC, explain the history and evolution of the protocol and declare the source of prior funding. HSRRB approval is required prior to continuing enrollment using USAMRMC-managed funds.

8. **Objectives:** Suggested length – 2 pages. State the specific aims and the research strategy of the study. When a proposal is submitted requesting funding as part of a larger study, the aims and research strategy should be presented for DOD-funded tasks only. The applicant must address how the research plan will be affected if not all large study components receive funding (e.g., Can the DOD-funded research be completed as proposed if funding is not received for all components? What adjustments would be needed in the study design to meet such a contingency?).
9. Study Population

   a. Describe the target population (to whom the study findings will be generalized) and the nature, approximate number, and pertinent demographic characteristics of the accessible population at the study site (population from which the sample will be recruited/drawn).

   b. Describe the methods that will be used to obtain a sample of subjects from the accessible population (e.g., convenience, simple random, stratified random) together with the inclusion and exclusion criteria (include age, gender, and ethnicity).

10. Protocol Design: Describe the type of study to be performed (prospective, randomized, controlled, etc.). Outline the proposed methodology in sufficient detail to show a clear course of action. Technological reliability and validity of procedures should be indicated. Minimum guidance for the plan should include:

   a. Subject Identification: Describe the code system to be used to maintain the confidentiality of subjects.

   b. Description of the Recruitment Process: Describe participant recruitment, including (1) participant availability; (2) inclusion and exclusion criteria; (3) methods for recruiting, retention, and follow-up; (4) data to support recruitment/retention estimates; (5) participant assignment to experimental groups and methods of randomization (if any); and (6) study endpoints. Provide copies of all recruitment and advertisement materials for review.

   c. Description of the Informed Consent Process: Specifically describe the plan for the informed consent process by stating who will perform the informed consent interview and when the interview will take place relative to the subject beginning study participation and in relation to any stressful situation (e.g., being informed he or she has a malignant tumor) or in relation to the administration of any mind-altering substances such as tranquilizers, conscious sedation, or anesthesia. Address how privacy and time for decision making will be provided and whether the potential subject will be allowed to discuss the study with anyone before making a decision. Two copies of the Informed Consent form should be completed (an original copy for the subject and a copy for the applicant’s study records). A third copy may be needed for the participant’s medical record; check with the participating site for specific study-site requirements.

   d. Plan for Addressing Human Subjects Protection Requirements: Address any issues that may lead to concern for the welfare of human subjects and confidentiality as described by the HSRRB at https://cdmrp.org/Program_Announcements_and_Forms under “Regulatory Document Forms.”

   e. Subject Assignment: Describe the randomization process or other procedures used for subject group assignments. Describe any potential biases in the protocol and how they will be addressed.
f. **Subject Screening Procedures:** List and describe any evaluations (e.g., laboratory procedures, history, and/or physical examination) that are required to determine eligibility/suitability for study participation. Please note that some screening procedures may need a separate consent or a two-stage consent process.

g. **Data Collection and Handling Procedures:** Describe all data collection procedures to be used in conducting the study (e.g., laboratory evaluations, specimens to be collected, schedule, and amounts). For studies using multiple measures or tests over time, it is helpful to display the data collection schedule in a spreadsheet or tabular format. Describe methods for the handling, distribution, analysis, storage (including where and whether special conditions are required), labeling, disposition, and security of specimens and/or imaging products (primary and secondary endpoints should be clearly defined and related to the power calculation). For multi-institutional trials, include a specimen handling and distribution plan agreed upon by all collaborating institutions.

h. **Clinical Assessments:** Provide a schedule of clinical, behavioral, laboratory, and physiological evaluations, and follow-up procedures. Provide any case report forms, data collection forms, questionnaires, rating scales, and/or interview guides that will be used in the study.

i. **Research Interventions:** Describe the research intervention or activity that the subject will experience. Provide sufficient detail in chronological order to enable a person not involved in the research to readily understand what the subject will experience.

j. **Data Management and Analysis:** Describe the data management and analysis plan, including the (1) overall approach to data management; (2) a plan for real-time data transfer; (3) a statistical plan that includes sample size calculations and methods to monitor quality and consistency of the intervention(s) and data collection; and (4) data security measures.

k. **Description of Protocol Drugs or Devices:** If the protocol uses an Investigational New Drug (IND) or Investigational Device Exemption (IDE), provide the following information:

   i. IND/IDE number and name of sponsor, if the study is in support of an application to the US Food and Drug Administration (FDA).

   ii. Complete names and composition of all medication(s), device(s), or placebo(s).

   iii. Source of medications, devices, or placebos.

   iv. Location of storage for study medications.

   v. Dose range, schedule, and administration of test articles.
vi. Detailed description of washout period, if used.

vii. Duration of drug or device treatment.

viii. Concomitant medications allowed.

ix. Antidotes and treatments available.

x. Disposition of unused drug.

xi. The procedure by which the IND/IDE sponsor will monitor the protocol in accordance with 21 CFR 312.²

xii. The following items also need to be submitted:

(1) A copy of the Investigator’s Brochure and/or device manual and associated case report/data collection forms. If the study involved the testing of an approved drug for a new indication, provide a copy of the package insert.

(2) A signed Form FDA 1572 for IND/IDE applications filed with the FDA, including the following information (also, for non-FDA new drug protocols, the following information should be included in the protocol):

   (a) Name, address, and a statement of the qualifications for each investigator and the name of each sub-investigator working under the applicant.

   (b) Names and addresses of facilities to be used.

   (c) Name and address of each IRB reviewing the protocol.

(3) For investigational devices, include the local IRB’s assessment of the risk (nonsignificant or significant) of the investigational device to be used in the study. If the device poses significant risk to research subjects, specify the IDE number obtained from the FDA, the name of the sponsor, and the procedure by which the sponsor will monitor the protocol in accordance with 21 CFR 812.³

1. Risks/Benefits Assessment

   i. Describe risks (physical [including pain and discomfort, disfigurement, infection, injury, death], psychological, social, economic, legal, and

²Title 21 Code of Federal Regulations Section 312, which includes Investigational New Drug Application procedures and requirements. Additional information can be found at http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfcfr/CFRSearch.cfm?CFRPart=312.

³Additional information can be found at http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfcfr/CFRSearch.cfm?CFRPart=812&showFR=1.
privacy/confidentiality risks) associated with the research, measures to be taken to minimize and/or eliminate risks or to manage unpreventable risks, and special medical or nursing care that will be needed prior to, during, or following participation.

ii. Describe benefits of the research to the subject. If there will be no benefits to the subjects (other than knowing he or she has contributed to science), state this in the protocol and Informed Consent form.

iii. Payment or compensation for participation is not considered to be a benefit and must be addressed in a separate section.

m. Reporting of Serious or Unexpected Adverse Events

i. Serious or unexpected adverse events can occur in any and all types of studies, not just experimental interventions or clinical trials.

ii. Include a definition of what constitutes an adverse event in the proposed study.

(1) For IND or IDE research, refer to definitions as listed in 21 CFR 312.32 for assistance.

(2) All IND/IDE protocols must specify how the requirements described below regarding adverse events will be addressed.

“An adverse event temporally related to participation in the study should be documented whether or not considered to be related to the test article. This definition includes intercurrent illnesses and injuries and exacerbations of preexisting conditions. Include the following in all IND safety reports: Subject identification number and initials; associate investigator’s name and name of medical treatment facility (MTF); subject’s date of birth, gender, and race/ethnicity; test article and dates of administration; signs/symptoms and severity; date of onset; date of resolution or death; relationship to the study drug; action taken; concomitant medication(s) including dose, route, and duration of treatment, and date of last dose.”

iii. Identify agencies or offices (including point of contact information) to be notified in the event of a serious and unexpected adverse event.

All protocols should contain the following language regarding the HSRRB reporting requirements for adverse events and unanticipated problems. (Note that unanticipated problems can occur in a study that does not require a research/clinical intervention.)

“Unanticipated problems involving risk to volunteers or others, serious adverse events related to participation in the study, and all volunteer deaths should be promptly reported by phone (301-619-2165), by e-mail

(hsrrb@det.amedd.army.mil), or by facsimile (301-619-7803) to the US Army Medical Research and Materiel Command’s Human Subjects Research Review Board. A complete written report should follow the initial notification. In addition to the methods above, the complete report can be sent to the US Army Medical Research and Materiel Command, ATTN: MCMR-ZB-P, 504 Scott Street, Fort Detrick, Maryland 21702-5012.”


For protocols that have a Medical Monitor assigned (see Subsection V.L.4), the following language also should be included.

“The Medical Monitor is required to review all unanticipated problems involving risk to volunteers or others, serious adverse events, and all volunteer deaths associated with the protocol and provide an unbiased written report of the event. At a minimum, the Medical Monitor should comment on the outcomes of the event or problem and in the case of a serious adverse event or death comment on the relationship to participation in the study. The Medical Monitor also should indicate whether he or she concurs with the details of the report provided by the study investigator. Reports for events determined by either the investigator or Medical Monitor to be possibly or definitely related to participation and reports of events resulting in death should be promptly forwarded to the HSRRB.”

n. Disposition of Data: Describe where data will be stored, who will keep the data, how the data will be stored, and the length of time the data will be stored. Note that records of IND/IDE studies must be kept until 2 years after a New Drug Application is approved/issued or for 2 years after the IND/IDE is withdrawn. Records required for IDE studies should be retained for 2 years following the date that the investigation is terminated or completed or the date that the records are no longer required for support of the pre-market approval application, whichever is sooner.

o. Modification of the Protocol: Describe the procedures to be followed if the protocol is to be modified, amended, or terminated before completion. Note that any modification of the protocol, Informed Consent form, and/or questionnaires, including a change of Principal Investigator, must be submitted to the local IRB for review and approval and then the HSRRB for second-level review and approval. Address this procedure even if no modification is anticipated.

p. Departure from the Protocol: Describe procedures and notifications to be made in the event of deviations from the approved protocol to include both the local IRB and the HSRRB.
q. **Study Organization and Management Plan:** Provide an organizational chart and a timetable for completion of the clinical trial and publication. Provide a plan for ensuring the standardization of procedures among staff and across sites (if applicable). Provide a plan for real-time communication among collaborating institutions (if applicable).

**M. Supporting Documentation:** Submit only material specifically requested in this program announcement. *This section is not intended for additional figures, tables, graphs, photographs, diagrams, chemical structures, pictures, pictorials, cartoons, or other information needed to judge the proposal.* Submitting material that was not requested may be construed as an attempt to gain a competitive advantage and such material will be removed; submitting such material may be grounds for administrative rejection of the proposal.

Supporting Documentation must be uploaded as a single PDF file under the “Required Files” tab of the CDMRP eReceipt system. All documents or letters that require signatures must be signed and incorporated into the Supporting Documentation file before it is submitted.

The first item in the Supporting Documentation file is the [Checklist/Table of Contents page](#). The requested, allowable items in this section must be listed in the Checklist/Table of Contents; these include:

1. **Abbreviations:** *Start section on a new page; one-page limit.* Provide a list of all acronyms, abbreviations, and symbols used in the main body of the proposal.

2. **References:** *Start section on a new page; no page limit.* List all relevant references using a standard reference format that includes the full citation (i.e., author(s), year published, title of reference, source of reference, volume, chapter, page numbers, and publisher, as appropriate).

3. **Biographical Sketches:** *Four-page limit per individual.* Include biographical sketches for all key personnel including collaborating investigators and support staff. These documents are a critical component of the review process. Incomplete or missing biographical sketches may result in lower proposal scores. The [Biographical Sketch form](#) may be used. Use of this form is not mandatory, but the information requested shall be presented in a similar format.

4. **Existing/Pending Support:** *Start section on a new page; no page limit.* List the titles, time commitments, supporting agencies, durations, and levels of funding for all existing and pending research projects involving the applicant and key personnel on a separate page. If no support exists, enter “None.” Proposals submitted under this program announcement should not duplicate other funded research projects.

5. **Facilities/Equipment Description:** *No page limit.* Describe the facilities available for performing the proposed research/services. Describe the institutional commitment, including any additional facilities or equipment proposed for purchase or available for use at no cost to the USAMRMC. Indicate whether Government-owned facilities or equipment are proposed for use.
6. **Letters of Support:** Provide letters of support from any collaborating individuals or institutions.

7. **Publications and/or Patent Abstracts: Five-document limit.** Include up to five relevant publication reprints and/or patent abstracts. A patent abstract should provide a nonproprietary description of the patent application. If more than five publications are included in the submission, the extra items will not be peer reviewed.

8. **Clinical Documentation (if applicable):** The following items must be included in this section:

- Informed Consent/Assent forms;
- Internal scientific and local IRB reviews for the clinical protocol;
- Questionnaires;
- Survey instruments;
- Participant recruitment brochures;
- Case report forms;
- Investigator’s brochure for proposals with IND/IDEs;
- Documentation that an IND/IDE has been submitted or a plan for submission of IND/IDE application to the FDA for therapeutic clinical trial protocols;
- A plan for study investigators to successfully complete institutional ethics training and a course in the conduct of clinical research in accordance with GCP within 1 year of initiation of the protocol;
- Documentation that the participating institutions have an intellectual and material property plan and are willing to resolve intellectual and material property issues;
- A specimen handling and distribution plan agreed upon by all collaborating institutions for multi-institutional trials; and
- Documentation of the availability of the substance or device to be used in the clinical trial. If the substance or device is to be provided from industrial sources, provide documentation of a cost-sharing plan.

N. **Budget Information:** Applicants must complete the [Detailed Cost Estimate form and the Budget Justification form](#), and upload them as a single PDF file under the “Required Files” tab of the CDMRP eReceipt system. When a proposal requesting funding as part of a larger study is submitted, the proposal’s budget justification should include only DOD-funded tasks.

1. **Funding Restrictions:** Funding for a Clinical Resource Development Award can be requested for a maximum of $225,000 for direct costs over the entire performance period. The performance period can be requested for up to 2 years. Indirect costs should be added as appropriate. Direct costs can cover salary, research supplies, equipment, and travel to
scientific/technical meetings. The travel allotment is $1,800 per year to attend scientific/technical meetings.

2. **Detailed Cost Estimate Form and the Budget Justification Instructions:** Budget is an important consideration in both peer review and programmatic review, and applicants are cautioned to use discretion in budget requests. Budgets also will be reviewed during award negotiations. *Organizations must provide sufficient detail and budget justification so that the Government can determine the proposed costs to be allocable and reasonable for the proposed research.* All costs must be entered in U.S. dollars.

The USAMRMC encourages in-kind contributions and cost-sharing for CDMRP-supported research. In-kind contributions may include support of services (e.g., laboratory services and salaries of personnel), real property and equipment, and/or supplies (e.g., drugs, devices, reagents) directly benefiting and specifically identifiable to the research project. *It is expected that institutions will share the cost of equipment purchased for this research proposal.* Please see full details under “Major Equipment” in Subsection V.N.2.c.

Costs proposed must conform to the following regulations and principles:

- **Commercial Firms:** Federal Acquisition Regulations (FAR) Part 31 and Defense FAR Supplement Part 31 ([http://farsite.hill.af.mil](http://farsite.hill.af.mil)), Contract Cost Principles and Procedures.

- **Educational Institutions:** Office of Management and Budget (OMB) Circular A-21, Cost Principles for Educational Institutions ([http://www.whitehouse.gov/omb/grants/grants_circulars.html](http://www.whitehouse.gov/omb/grants/grants_circulars.html)).


- **State, Local, and Tribal Governments:** OMB Circular A-87, Cost Principles for State, Local, and Indian Tribal Governments ([http://www.whitehouse.gov/omb/grants/grants_circulars.html](http://www.whitehouse.gov/omb/grants/grants_circulars.html)).

Follow the instructions below when providing the information requested in the Detailed Cost Estimate form.

**a. Personnel**

i. **Name:** Beginning with the applicant, list all participants who will be involved in the project during the initial budget period, whether or not salaries are requested. Include all collaborating investigators, research associates, individuals in training, and support staff. *The applicant must be identified as the Principal Investigator of the proposal.*
ii. **Role on Project:** Identify the role of each participant listed. Describe his or her specific functions in the Budget Justification section of the Detailed Cost Estimate form.

iii. **Type of Appointment (Months):** List the number of months per year reflected in an individual’s contractual appointment with the applicant organization. The Government assumes that appointments at the applicant organization are full time for each individual. If an appointment is less than full time, e.g., 50%, note this with an asterisk (*) and provide a full explanation in the Budget Justification section of the Detailed Cost Estimate form. Individuals may have split appointments (e.g., for an academic period and a summer period). For each type of appointment, identify and enter the number of months on separate lines.

iv. **Annual Base Salary:** Enter the annual institutional base salary for each individual listed for the project.

v. **Percentage of Effort on Project:** The applicant’s qualifications and the amount of time that he or she and other professional personnel will devote to the research are important factors in selecting research proposals for funding. List the percentage of each appointment to be spent on this project for each key staff member. Include the percent effort of all unpaid collaborators and consultants. Each clinical trial must have a clinical coordinator who has sufficient time dedicated to the project to carry out the record keeping, coordination, and/or other administrative duties the project entails.

vi. **Salaries Requested:** Enter the salaries in whole U.S. dollars for each position for which funds are requested. Calculate the salary request by multiplying an individual’s institutional base salary by the percentage of effort on the project.

vii. **Fringe Benefits:** Fringe benefits for each position may be requested in accordance with institutional guidelines, provided the costs for all sponsors are treated consistently by the applicant’s organization. Provide documentation to support the fringe benefits.

viii. **Totals:** Calculate the totals for each position and enter these as subtotals in the columns indicated.

b. **Consultant Costs:** Provide the names and organizational affiliations of all consultants whether or not funds are requested.

c. **Major Equipment:** It is the policy of the DOD that all commercial and nonprofit recipients provide the equipment needed to support proposed research. In those rare cases where specific additional equipment is approved for commercial and nonprofit organizations, such approved cost elements shall be negotiated separately.
i. If the purchase of equipment for this research project is requested, it is expected that the applicant’s institution will share 50% of the cost.

ii. Permanent equipment is any article of nonexpendable tangible property having a useful life of 2 years or longer and an acquisition cost of $5,000 or more per unit.

iii. The basis for the cost of each item of permanent equipment included in the budget must be disclosed.

iv. Title of equipment or other tangible property purchased with Government funds may be vested in institutions of higher education or with nonprofit organizations whose primary purpose is the conduct of scientific research. Normally, the title will vest in the recipient if vesting will facilitate scientific research performed by the institution or organization for the Government.

d. Materials, Supplies, and Consumables: A general description and estimated total cost of expendable equipment and supplies are required. Itemize supplies in separate categories (e.g., glassware, chemicals, radioisotopes). Categories in amounts less than $1,000 do not need to be itemized. If animals will be purchased, state the species, strain (if applicable), and the number of animals to be used. If human cell lines are to be purchased, state the source and the description.

e. Travel Costs: Costs for travel to scientific/technical meetings may not exceed $1,800 per year.

Travel costs associated with the execution of the proposed work should be entered in this section. If applicable, reasonable costs for travel between collaborating institutions should be included and are not subject to the yearly $1,800 limitation on travel to meetings. Justification for these travel costs should be provided. Travel outside the U.S. requires prior approval from the US Army Medical Research Acquisition Activity (USAMRAA).

f. Research-Related Subject Costs: Itemize costs of subject participation in the research study. These costs are strictly limited to expenses associated specifically with the proposed study. The USAMRMC will not provide funds for ongoing medical care costs not related to a subject’s participation in the research study.

g. Other Direct Costs: Itemize other anticipated direct costs such as publication and report costs, rental for computers and other equipment (provide hours and rates), and communication costs. Unusual or expensive items should be fully explained and justified. Estimate the costs of publishing and reporting research results, including direct charges for clerical preparation, illustrations, reprints, and distribution.

h. Subaward Costs: A description of services or materials to be awarded by subcontract or subgrant is required. For awards totaling $10,000 or more:
Identify the type of award to be used (e.g., cost reimbursement, fixed price);

Identify the proposed subcontractor or subgrantee, if known, and provide an explanation of why and how the subcontractor or subgrantee was selected or will be selected;

Specify whether the award will be competitive and, if noncompetitive, provide a rationale to justify the absence of competition; and

Provide the proposed acquisition price.

i. **Indirect Costs (overhead, general and administrative, and other):** The most recent rates, dates of negotiation, base(s), and periods to which the rates apply should be disclosed with a statement identifying whether the proposed rates are provisional or fixed.

j. **Total Costs for the Entire Proposed Period of Support (second page of the Detailed Cost Estimate form):** Enter the totals in each budget category for all additional years of support requested and itemize these totals in the Budget Justification section of the Detailed Cost Estimate form. Note with an asterisk (*) and explain any significant increases or decreases from the initial year budget. All amounts should be in U.S. dollars. Directs costs, indirect costs, and the total cost for the entire proposed period of support should equal the amount entered in the “Required Files” tab at https://cdmrp.org.

3. **Budget Justification (third page of the Detailed Cost Estimate form):** Each item in the budget must be clearly justified in the Budget Justification section of the Detailed Cost Estimate form.

4. **Federal Agency Financial Requirement:** Proposals from Federal agencies must provide a plan delineating how all funds will be obligated by September 30, 2007, and how funds will be available to cover research costs over the entire award period. The plan must include the funding mechanism(s) that will be used to carry over funds between fiscal years, such as administrative agreements with foundations, non-Federal institutions, and universities.

   **Start the plan on a new page at the end of the Budget Information section.** The Federal Agency Financial Plan must be uploaded as part of the budget information before the submission deadline of 5:00 p.m. Eastern time, April 25, 2006.

O. **Regulatory Requirements:** Completed and signed copies of the Certificate of Environmental Compliance and Principal Investigator Safety Program Assurance form must be uploaded under the “Required Files” tab of the CDMRP eReceipt system as separate PDF files.

In addition, for proposals involving human studies, regulatory documents pertaining to research involving human subjects and/or human anatomical substances or cadavers must be submitted within the Clinical Protocol and Supporting Clinical Documents sections of the proposal (see Subsections V.L and V.M. 8) as a required file. Any other regulatory documents should not be
submitted with the proposal. Instead, the applicant should provide these documents to the USAMRMC only upon request.

P. USAMRAA-Required Documents: The Contract Representative at the applicant’s institution must upload the current version of the institution’s negotiated Rate Agreement, the Certifications and Assurances for Assistance Agreements, and the Representations for Assistance Agreements. These documents must be uploaded as separate PDF files under the Contract Representative’s “My Profile” tab of the CDMRP eReceipt system by the proposal submission deadline.

Q. Submission Date and Time: Proposals must be approved on the CDMRP eReceipt system by the Contract Representative at the applicant’s institution by the deadline. Proposals that are incomplete or not approved electronically before the deadline will not be considered for review. The eReceipt system will not accept data entry, file uploads, or approvals submitted after the 5:00 p.m. Eastern time, April 25, 2006 deadline.

The timeline for the Clinical Resource Development Award is:

<table>
<thead>
<tr>
<th>Event</th>
<th>Date/Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Online Letter of Intent</td>
<td>Expected by March 28, 2006</td>
</tr>
<tr>
<td>Online Proposal Information</td>
<td>Required prior to proposal submission</td>
</tr>
<tr>
<td><strong>Proposal Submission/Approval Deadline</strong></td>
<td>5:00 p.m. Eastern time, April 25, 2006</td>
</tr>
<tr>
<td>Peer Review (First Tier)</td>
<td>June 2006</td>
</tr>
<tr>
<td>Programmatic Review (Second Tier)</td>
<td>October 2006</td>
</tr>
<tr>
<td>Request for Additional Documents</td>
<td>As early as 2 weeks after the completion of programmatic review</td>
</tr>
<tr>
<td>Notification Letter</td>
<td>Approximately 4 weeks after the completion of programmatic review</td>
</tr>
<tr>
<td>Award Start Date</td>
<td>Anticipated between December 2006 and April 2007</td>
</tr>
</tbody>
</table>

R. Electronic Submission Requirements: Electronic submission is required. Only proposals submitted as PDF files through the CDMRP eReceipt system at [https://cdmrp.org](https://cdmrp.org) will be accepted.

Several steps are critical to successful proposal submission:

- The Proposal Information must be “Finalized for CR Approval” before the proposal is submitted. Applicants are encouraged to begin this part of the submission process early.
- Proposal Contacts must be “Finalized for CR Approval” before the proposal is submitted. The e-mail address of a Contract Representative at the applicant’s institution must be included in the Proposal Contacts. Applicants are encouraged to begin this part of the submission process early.
- Applicants are encouraged to coordinate with their Contract Representative early in the application process.
• The Contract Representative authorized to negotiate on behalf of the applicant’s institution is required to provide final approval before the proposal is accepted.

• The eReceipt system will not accept data entry, file uploads, or approvals submitted after the 5:00 p.m. Eastern time, April 25, 2006 deadline.

• Some items in the proposal including figures, tables, graphs, letters, or publications will need to be scanned electronically. These documents should be scanned at a resolution of 300 dpi or less.

• Applicants are encouraged to retain a date and time-stamped copy of the proposal component files as prepared by word processing software (e.g., Microsoft Word, WordPerfect) as well as the original PDF conversion file.

• The Detailed Cost Estimate form and the Budget Justification form must be uploaded under the “Required Files” tab of the CDMRP eReceipt system.

• The regulatory documents required at submission include a completed and signed Certificate of Environmental Compliance and a completed and signed Principal Investigator Safety Program Assurance form. These forms must be uploaded under the “Required Files” tab of the CDMRP eReceipt system.

VI. PROPOSAL REVIEW INFORMATION

A. Proposal Review and Selection Overview

1. Process: The CDMRP uses a two-tier review process for proposal evaluation. The two tiers differ fundamentally. The first tier is a scientific peer review of proposals against established criteria for determining scientific merit. The second tier is a programmatic review of proposals that compares submissions to each other and recommends proposals for funding based on scientific merit and overall goals of the program.

2. Peer Review: Peer review is conducted by scientific and consumer reviewers. The primary responsibility of the peer reviewers is to provide unbiased, expert advice on the scientific/technical merit and relevance of proposals based on the review criteria published for each award mechanism.

Scientific reviewers are selected for their subject matter expertise and experience with scientific peer review. Consumer reviewers are nominated by an advocacy or support organization and are selected on the basis of their leadership skills, commitment to advocacy, and interest in science. Consumers augment the peer review process by bringing the patient perspective to the assessment of science and the relevance of the research.

The peer review summary statement is a product of scientific peer review. Each summary statement includes the peer review scores and an evaluation of the project as assessed by the peer reviewers according to the evaluation criteria published in this program announcement.
3. **Programmatic Review:** Programmatic review is conducted by the Integration Panel, which is composed of scientists, clinicians, and consumer advocates. The scientific members of the Integration Panel represent diverse disciplines and specialty areas, and the consumer members represent national advocacy constituencies. A function of programmatic review is to structure a broad portfolio of grants across all disciplines. Programmatic review is a comparison-based process in which proposals from multiple research areas compete in a common pool. Integration Panel members base programmatic review primarily on the peer review summary statements and the proposal abstracts. The Integration Panel also may review SOWs and impact statements. Full proposals are not forwarded to programmatic review.

HBCU/MI proposals are reviewed concurrently with others in the same research area during scientific peer review. However, they may be evaluated separately during programmatic review. Consistent with the CDMRP’s goal, recommendations for funding HBCU/MI proposals are based on scientific excellence and program relevance.

B. **Review Criteria**

1. **Peer Review:** Clinical Resource Development Award proposals will be evaluated by peer reviewers according to the following criteria. The reviewers will evaluate:

   a. **Proposal:** Clinical Resource Development Award proposals will be evaluated according to the following criteria:

      - **Resource Development Product**
        - How a clear endpoint or product has been identified.
        - The rationale supporting the clinical need for the resource.
        - The feasibility of the resource development approach.
        - The reasonableness and appropriateness of the plan for distribution of the resource following its development.

      - **Clinical Impact**
        - The extent to which the project will make an original and important contribution to TSC research, diagnosis or treatment, including its impact on the design, implementation, or interpretation of TSC therapeutic clinical trials.
        - The direct relevance and applicability of the proposed tool/resource to the measurement of clinical and/or surrogate endpoints in TSC clinical trials.

      - **Personnel and Environment**
        - The extent to which the applicant and/or the research team possess the requisite experience and expertise to develop the proposed product.
If the proposed work is not in an area in which the applicant has experience, the evidence that advice and input will be obtained from other appropriate sources (e.g., collaborators and colleagues, or the completion of a training course).

Evidence of an appropriate clinical/laboratory setting and the availability of institutional resources to support the product development.

A letter of institutional commitment included from the applicant’s institution.

**Budget**

- How the budget is reasonable and appropriate for the development of the proposed resource.
- How the projected distribution costs for the product are reasonable and appropriate.

**b. Protocol(s) (As appropriate):** Clinical Resource Development Award protocols will be evaluated according to the following criteria:

  **Protocol Design**

  - How the conceptual framework and design of the clinical protocol are consistent with the objectives set forth in the main body of the proposal.
  - The rationale for and feasibility of the proposed clinical study, as demonstrated by a literature review, preliminary data, and sound reasoning.
  - How the proposed methodology is described in sufficient detail and appropriate to the study’s objectives.
  - How the recruitment, informed consent, and subject screening processes are appropriate and adequately described.
  - The feasibility of the study in terms of subject availability, recruitment schedule and accrual, and likely attrition.
  - The statistical plan, including data collection, power calculations, randomization criteria, and data analysis.

  **Protocol Management**

  - How the background and experience of the individuals conducting the clinical study is appropriate to its successful completion.
  - The extent to which the roles and responsibilities of all study personnel are clearly described.
  - How the research and clinical settings are appropriate and adequate to support the study.
  - For multi-institutional studies, as appropriate, whether there is a plan for the coordination of IRB submissions.
  - How procedures are in place for protocol modifications during the course of the study.
• **Ethics and/or Regulatory Issues**
  
  o Whether or not there is an adequate discussion of such ethical considerations as informed consent, information privacy, and the assessment of participant risks and benefits.
  
  o Evidence of a plan for the study investigators to complete an ethics training program and a course in the conduct of clinical research in accordance with GCP within 1 year of protocol initiation.
  
  o How potential adverse events have been defined, and whether there are named agencies or offices to be notified in this event.
  
  o Evidence that a plan has been proposed for data disposition during and after the study.
  
  o Whether preliminary institutional IRB approval(s) for the clinical protocol and Informed Consent form have been obtained at the highest level possible.
  
  o If a Medical Monitor is required, evidence that he or she is appropriately qualified.

2. **Programmatic Review:** Criteria used by the Integration Panel to make funding recommendations that maintain the TSCRP’s broad portfolio include:

  • Ratings and evaluations of the peer reviewers (scientific and consumer),
  
  • Programmatic relevance,
  
  • Relative innovation and impact,
  
  • Program portfolio balance, and
  
  • Adherence to the intent of the award mechanism.

Scientifically sound proposals that best fulfill the above criteria and most effectively address the unique focus and goals of the program are selected by the Integration Panel and recommended for funding to the Commanding General, USAMRMC.

**VII. AWARD ADMINISTRATION INFORMATION**

**A. Award Notices:** Each applicant will receive notification of the award status of his or her proposal. A copy of the peer review summary statement will be posted to the CDMRP eReceipt system. Applicants can expect to receive notification approximately four weeks after programmatic review.

**B. Administrative Requirements:** Awards are made to organizations, not individuals. An applicant must submit a proposal through, and be employed by or affiliated with, a university, college, nonprofit research institution, commercial firm, or Government agency (including military laboratories) to receive support. A prospective recipient must meet certain minimum standards pertaining to institutional support, financial resources, record of performance, integrity,
organization, experience, operational controls, facilities, and conformance with safety and environmental statutes and regulations (OMB Circular A-110 and DOD Grant and Agreement Regulations) to be eligible for an award. Any organization requesting receipt of an award through this announcement must be registered in the Central Contractor Registration (CCR) database. Access to the CCR online registration is through the CCR homepage at http://www.ccr.gov.

Proposals from Federal agencies must provide a plan delineating how all funds will be obligated by September 30, 2007, and how funds will be available to cover research costs over the entire award period. The plan must include the funding mechanism(s) that will be used to carry over funds between fiscal years, such as administrative agreements with foundations, non-Federal institutions, and universities.

C. Award Negotiation: Award negotiation consists of discussions, reviews, and justifications of critical issues involving the USAMRAA. A Contract Specialist and/or representative from the USAMRAA will contact the Contract Representative authorized to negotiate contracts and grants at the applicant’s institution. Additional documentation and justifications related to the budget may be required as part of the negotiation process.

For multi-institutional studies, collaborating institutions must be willing to resolve potential intellectual and material property issues and remove institutional barriers to achieving high levels of cooperation to ensure the successful establishment and maintenance of the research project. An intellectual and material property plan agreed to by all participating institutions may be required during award negotiations.

The award start date will be determined during the negotiation process.

D. Regulatory Review

1. Overview: Concurrent with the USAMRAA negotiation, the Office of Surety, Safety and Environment will review the Certificate of Environmental Compliance and the Principal Investigator Safety Program Assurance form submitted with the proposal. The applicable USAMRMC regulatory office will review documents related to research involving animal use, human subjects/anatomical substance use, and cadaver use submitted upon request to ensure that DOD regulations are met.

2. Certificate of Environmental Compliance: The Certificate of Environmental Compliance must be submitted with the proposal. If multiple research sites/institutions are funded in the proposal, then a Certificate of Environmental Compliance for each site will be requested at a later date.

3. Safety Program Documents: The Principal Investigator Safety Program Assurance form must be submitted with the proposal.

A Facility Safety Plan is required; it will be requested at a later date. A Facility Safety Plan from the applicant’s institution may have been received previously and approved by the
USAMRMC. A list of institutions that have approved Facility Safety Plans can be found on the USAMRMC website at https://mrmc.detrick.army.mil/crrcqsohdfsplan.asp. If the applicant’s institution is not listed on the website, contact the institution’s Facility Safety Director/Manager to initiate completion of the institution-based Facility Safety Plan. Specific requirements for the Facility Safety Plan can be found at https://mrmc.detrick.army.mil/docs/rcq/FY02FSPAppendix.doc.

If multiple research sites/institutions are funded in the proposal, a Facility Safety Plan for each site/institution not listed in the aforementioned website will be requested at a later date.

4. Research Involving Animal Use: Specific documents relating to the use of animals in the proposed research will be requested by the CDMRP if the proposal is selected for funding (these documents should not be submitted with the proposal). The Animal Care and Use Review Office (ACURO), a component of USAMRMC Office of Research Protections (formerly Regulatory Compliance and Quality), must review and approve all animal use prior to the start of working with animals. Applicants must complete and submit the animal use appendix titled “Research Involving Animals,” which can be found on the ACURO website https://mrmc-www.army.mil/rodorpaurd.asp.

Questions related to animal use may be directed to ACURO as follows:

- Phone: 301-619-6694
- Fax: 301-619-4165
- E-mail: acuro@amedd.army.mil
- Mail: MCMR-ZB-PA
  504 Scott Street
  Fort Detrick, MD 21702-5012

Specific requirements for research involving animals can be found at https://mrmc.detrick.army.mil/docs/rcq/FY05AnimalAppendix.doc.

5. Research Involving Human Subjects/Biological Substances/Cadavers: In addition to local Institutional Review Board (IRB) approval to conduct research involving human subjects and/or human biological substances or cadavers, a second tier of IRB review and approval also is required by the DOD. This second review is conducted by the Human Subjects Research Review Board (HSRRB), which is administered by the USAMRMC Office of Research Protections. The HSRRB is mandated to comply with specific laws and directives governing all research involving human subjects that is conducted or supported by the DOD. These laws and directives are rigorous and detailed and will require information in addition to that supplied to the local review board.

a. Requirements: Specific requirements for research involving human subjects, human biological substances, and/or cadavers can be found at https://mrmc.detrick.army.mil/docs/rcq/HumanSubjectsAppendix.pdf.
Personnel involved in human subjects research must have appropriate instruction in the protection of human subjects. Documentation confirming that this instruction has been completed will be required during the regulatory review process.

It is expected that there will be timely resolutions of human subjects protocols submitted to the investigator’s local IRB.

Additional information pertaining to the human subjects regulatory review process, guidelines for developing protocols, and suggested language for specific issues can be found at: https://mrmc.detrick.army.mil/rodorphrpo.asp.


c. Intent to Benefit: Investigators must consider the requirements of Title 10 United States Code Section 980 (10 USC 980) applicable to DOD-sponsored research before writing a research protocol. Title 10 United States Code Section 980 requires that “Funds appropriated to the Department of Defense may not be used for research involving a human being as an experimental subject unless (1) the informed consent of the subject is obtained in advance; or (2) in the case of research intended to be beneficial to the subject, the informed consent may be obtained from a legal representative of the subject.”

Furthermore and consistent with the Common Federal Policy for the Protection of Human Subjects, if an individual cannot give his or her own consent to participate in a research study, consent of the individual’s legally authorized representative must be obtained before the individual’s participation in the research. Moreover, an individual not legally competent to consent (e.g., incapacitated individuals, incompetents, minors) may not be enrolled in DOD-sponsored research unless the research is intended to benefit each subject enrolled in the study. For example, a subject may benefit directly from medical treatment or surveillance beyond the standard of care. Investigators should be aware that this law makes placebo-controlled clinical trials problematic because of the “intent to benefit” requirement whenever participation is sought of subjects from whom consent must be obtained by the legally authorized representative.

d. Conditions Regarding DOD Funding of Research on Human Embryonic Stem Cells: Research involving the derivation and use of human embryonic germ cells from fetal tissue may be conducted with DOD support only when the research is in compliance with 45 CFR 46, Subpart B (Title 45 of the Code of Federal Regulations, Section 46, Subpart B); 42 USC 289g through 289g-2; US Food and Drug Administration regulations; and any other applicable Federal, state, and local laws and regulations.

Research on existing human embryonic stem (hES) cell lines may be conducted with Federal support through the DOD only if the cell lines meet the current US Federal criteria as listed on the following National Institutes of Health (NIH) website (http://grants.nih.gov/grants/guide/notice-files/NOT-OD-02-005.html). A list of the currently approved cell lines can be obtained from the NIH Human Embryonic Stem Cell
Registry (http://stemcells.nih.gov/research/registry). The NIH code should be used to identify the cell lines in the proposal.

Research involving the derivation of new stem cells from human embryos or the use of hES cells not listed on the NIH Human Embryonic Stem Cell Registry may not be conducted with Federal support through the DOD.

This restriction applies to hES cells derived from blastocysts remaining after infertility treatments and donated for research, blastocysts produced from donated gametes (oocytes and sperm) for research purposes, and the products of nuclear transfer. The research is subject to all applicable local, state, and Federal regulatory requirements.

e. Clinical Trial Registry: All applicants are required to register clinical trials individually on www.clinicaltrials.gov using the Secondary Protocol ID number designation of: CDMRP-CDMRP Log Number. If several protocols exist under the same proposal, the Secondary Protocol ID number must be: CDMRP-CDMRP Log Number-A, B, C, etc. Clinical trials must be registered prior to enrollment of the first patient. All trials that meet the definition on the NIH database (see http://prsinfo.clinicaltrials.gov/, click on “Data Element Definitions,” see section 6, “Study Phase” and “Study Type”) to include all Phase I-IV clinical trials and trials that do not fit into one or more phases, but that are clearly interventional or observational (e.g., some epidemiological or behavioral studies) are required to register. Address questions on registration to the www.clinicaltrials.gov administrator.

6. Award/Regulatory Approval: The applicant may not use, employ, or subcontract for the use of any human subjects, human anatomical substances/cadavers, or laboratory animals without written approval from the applicable USAMRMC regulatory office once an award is made. The applicable USAMRMC regulatory office will forward applied-for written approvals directly to the applicant.

E. Reporting Requirements: The Government requires reports to be submitted for continuation of the research and funding. The specific reports due to the Government will be described in each award instrument. (Full USAMRMC reporting requirements can be found at https://mrmc-www.army.mil, under “Links and Resources.”) Failure to submit required reports by the required date may result in a delay in or termination of award funding.

Reporting requirements include the following:

1. Research Progress Reports: Reporting requirements consist of an annual report (for each year of research except the final year) that presents a detailed summary of scientific issues and accomplishments and a final report (submitted in the last year of the award period) that details the findings and issues for the entire project. Copies of all scientific publications and patent applications resulting from CDMRP funding should be included in the progress reports.
2. Fiscal Reports: Quarterly fiscal report requirements may include the Standard Form Report, SF 272, Federal Cash Transaction, used for grants and cooperative agreements to track the expenditure of funds on the research project.

3. Non-Exempt Human Studies Reports: For non-exempt human subjects research, documentation of local IRB continuing review (in the intervals specified by the local IRB but at least annually) and approval for continuation must be submitted directly to the Office of Research Protections – Human Research Protection Office.

4. Animal Use Reports: Applicants are required to submit annual animal use information for a report to Congress, verification of annual protocol review, and notification of protocol suspension or revocation. Institutions are required to provide updated US Department of Agriculture reports and notification of changes to accreditation status as verified by the Association for Assessment and Accreditation of Laboratory Animals and Office of Laboratory Animal Welfare.

VIII. OTHER INFORMATION

A. Disclosure of Proprietary Information outside the Government: By submitting a proposal, the applicant understands that proprietary information may be disclosed outside the Government for the sole purpose of technical evaluation. The USAMRMC will obtain a written agreement from the evaluator that proprietary information in the proposal will only be used for evaluation purposes and will not be further disclosed or used. Funded proposals may be subject to public release under the Freedom of Information Act; proposals that are not selected for funding are not subject to public release.

B. Government Obligation: Applicants are cautioned that only an appointed Contracting/Grants Officer may obligate the Government to the expenditure of funds. No commitment on the part of the Government to fund preparation of a proposal or to support research should be inferred from discussions with a technical project officer. Applicants who, or organizations that, make financial or other commitments for a research effort in the absence of an actual legal obligation signed by the USAMRAA Contracting/Grants Officer do so at their own risk.

C. Information Service: Offerors may use the technical reference facilities of the National Technical Information Service, 5285 Port Royal Road, Springfield, Virginia 22161, for the purpose of surveying existing knowledge and avoiding needless duplication of scientific and engineering effort and the expenditure thereby represented. All other sources also should be consulted to the extent practical for the same purpose.

D. Inquiry Review Panel: Applicants may submit a letter of inquiry to the USAMRMC in response to funding decisions made for a given proposal. Members of the CDMRP staff, the USAMRMC Judge Advocate General staff, and USAMRAA Grants Officers constitute an Inquiry Review Panel and review each inquiry to determine whether factual or procedural errors in either peer or programmatic review have occurred, and if so, what action should be taken.
E. **Title to Inventions and Patents:** In accordance with the Bayh-Dole Act (35 USC 200 et seq.), title to inventions and patents resulting from such Federally funded research may be held by the grantee or its collaborator, but the US Government shall, at a minimum, retain nonexclusive rights for the use of such inventions. An investigator must follow the instructions in the assistance agreement concerning license agreements and patents.

F. **J-1 Visa Waiver:** It is the responsibility of the awardee to ensure that the research staff is able to complete the work without intercession by the DOD for a J-1 Visa Waiver on behalf of a foreign national in the United States under a J-1 Visa.
### IX. ACRONYM LIST

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<td>Tuberous Sclerosis Complex Research Program</td>
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