Program Announcement

Department of Defense (DOD) Ovarian Cancer Research Program (OCRP)

Funding Opportunity Number: W81XWH-07-OCRP-CA

Concept Award

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I. HELP LINE INFORMATION

A. Agency Name

US Army Medical Research and Materiel Command (USAMRMC), Office of the Congressionally Directed Medical Research Programs (CDMRP), 1077 Patchel Street, Fort Detrick, Maryland 21702-5024.

B. Agency Contact(s)

1. Program announcement, proposal format, or required documentation: Principal Investigators (PIs) and Authorized Organizational Representatives (AORs) 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
   Email: cdmrp.pa@amedd.army.mil

2. eReceipt system: A help line for questions relating to the submission of pre-application components through the CDMRP eReceipt 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 email as follows:

   Website: https://cdmrp.org
   Email: help@cdmrp.org

3. Grants.gov: Issues in submitting applications through the Grants.gov (http://www.grants.gov/) portal should be directed to Grants.gov at 800-518-4726 or email support@grants.gov. The Grants.gov hours of operation are Monday through Friday, 7:00 a.m. to 9:00 p.m. Eastern time. Deadlines for proposal submission are set at 11:59 p.m. Eastern Time on the deadline date. Therefore, there is an approximate 3-hour period during which the Grants.gov Help Desk will NOT be available to assist with Grants.gov submissions. Please plan ahead accordingly, as the CDMRP Help Desk is not able to answer questions about Grants.gov submissions.

   Grants.gov will only notify PIs of changes made to this Program Announcement and/or Application Package if the PI clicks on the “send me change notification emails” link and subscribes to the mailing list on the Opportunity Synopsis Page for this announcement. Please note that if the PI does not subscribe and the Application Package is updated or changed, the original version of the Application Package may not be accepted.
C. Anticipated Instrument Type(s)

The USAMRMC implements its extramural research program predominantly through the award of grants and cooperative agreements. More information on these funding instruments may be obtained by request via:

   Fax: 301-619-2937
   Email: qa.baa@amedd.army.mil

D. Catalog of Federal Domestic Assistance (CFDA) Number 12.420

Military Medical Research and Development.

E. Commonly Made Mistakes

- Pre-application submission is not completed before the mandatory pre-application deadline (pre-application remains in draft status).
- Failure to request updates on any modifications made to the application package.
- Incorrect application package or award mechanism is used to submit a proposal through Grants.gov.
- Attachments are uploaded into the incorrect form on Grants.gov.
- Files are attached in the wrong location on Grants.gov forms.
- Attachments are not PDF documents.
- Page limitations are exceeded.
II. FUNDING OPPORTUNITY DESCRIPTION

Funding of proposals received in response to this program announcement is contingent on the availability of Federal funds appropriated in a bill for this program.

A. Award Description

The Ovarian Cancer Research Program (OCRP) Concept Award supports new ideas and the exploration of innovative concepts or theories in ovarian cancer. The proposed research should provide a catalyst to challenge or expand current thinking and approaches in ovarian cancer research. Presentation of preliminary data is not consistent with the intent of this mechanism. However, a rationale for the work must be provided. Proposals must describe how the unique concept could create a new avenue for investigation and how it is relevant to ovarian cancer.

Proposals may address any area in ovarian cancer; proposals in the following research areas are encouraged:

- Developing or testing molecular imaging for early detection of ovarian cancer
- Identifying the primary lesion in ovarian cancer, including the possibility of environmental or infectious origin
- Addressing whether early and late stage ovarian cancer are the same disease
- Addressing quality of life issues associated with ovarian cancer treatment, such as neuropathy and cognitive changes
- Identifying or validating targets for vaccine development
- Investigating the impact of genetic factors affecting metabolism (nutritional genomics) or environment on ovarian cancer

Because these awards are designed for preliminary investigations, projects involving human subjects or specimens will not be supported unless they are exempt under 32 CFR 219.101(b) or eligible for expedited review (32 CFR 219.110 or 21 CFR 56.110). Studies that do not qualify for exempt or expedited status during review at any level will be administratively withdrawn and will not be funded.

B. Eligibility

All investigators, including predoctoral and postdoctoral fellows (or equivalent), are eligible to submit proposals. Additional information about individual and institutional eligibility may be found in Appendix 1.

C. Funding

Funding for a Concept Award can be requested for up to $75,000 for direct costs for up to a 18 month performance period plus indirect costs as appropriate. When the applicant institution calculates its own indirect costs for subawards, it can only charge costs on the first $25,000 of each subaward.
Funds can cover:

- salary
- research supplies
- travel to scientific/technical meetings
- travel between collaborating institutions

The nature of the OCRP does not allow for renewal of grants or supplementation of existing grants. Projects requiring lower levels of funding may also be submitted.

*The CDMRP expects to allot approximately $3 million (M) of the $10M Fiscal Year 2007 (FY07) OCRP appropriation to fund approximately 25 Concept Awards, depending on the quality and number of proposals received.*

**D. Award Administration**

For Concept Awards, no changes in the institution or the SOW will be allowed once the proposal has been submitted.

**E. Submission and Review Timeline**

Proposal submission is a two-step process consisting of (1) pre-application submission and (2) proposal submission.

- **Pre-application Submission Deadline:** 5:00 p.m. Eastern time, April 10, 2007
- **Proposal Submission Deadline:** 11:59 p.m. Eastern time, April 24, 2007
- **Peer Review:** June 2007
- **Programmatic Review:** October 2007

Awards will be made approximately 4-6 months after receiving the funding notification letter, but no later than September 30, 2008.
III. PROGRAM HISTORY AND OBJECTIVES

The OCRP was established in FY97 to promote innovative research focused on eradicating ovarian cancer. Appropriations for the OCRP from FY97 through FY06 totaled $101.7M. During this time, 328 Concept Award proposals have been received and 11 have been recommended for funding. The FY07 appropriation is $10M.

The overall goal of the FY07 OCRP is to eliminate ovarian cancer by supporting research to detect, diagnose, prevent, and control ovarian cancer. The key initiative of the Concept Award is to spark new ideas, innovative technologies, and concepts that will drive the field forward. The OCRP encourages risk-taking research.
IV. SUBMISSION PROCESS STEP 1: PRE-APPLICATION SUBMISSION

Proposal submission is a two-step process, consisting of (1) a pre-application submission through the CDMRP eReceipt system (https://cdmrp.org/) and (2) a proposal submission through Grants.gov (http://www.grants.gov/). This section describes the process for pre-application submission. For proposal submission, see Section V. Proposal submission will not be accepted unless a pre-application was previously submitted. The PI and Organization identified in the proposal submitted through Grants.gov should be the same as those identified in the pre-application. If there is a change in PI or Organization after submission of the pre-application, please contact the eReceipt helpdesk at help@cdmrp.org or 301-682-5507.

For assistance, please see Help Line Information (Section I).

A. Pre-application Components and Submission

The pre-application for a Concept Award consists of a Letter of Intent (LOI) Narrative and the other components discussed below. This subsection provides a summary of the pre-application submission requirements.

All pre-application components for the OCRP Concept Award mechanism, including the LOI Narrative, must be submitted electronically through the CDMRP eReceipt system by the 5:00 p.m. Eastern time, April 10, 2007 deadline. Material submitted after the pre-application submission deadline, unless specifically requested by the Government, will not be forwarded for processing. Failure to meet this deadline shall result in pre-application rejection and subsequent proposal rejection.

1. Proposal Information: PIs must enter the Proposal Information as described in the CDMRP eReceipt system before uploading the LOI Narrative.

2. Proposal Contacts: Enter contact information for the PI.

3. Collaborators and Conflicts of Interest (COI): To avoid COI during the screening and review processes, list the names of all scientific participants in the proposed research project including collaborators, consultants, and subawardees. Add all individuals outside of the proposal who may have a conflict of interest in the review of this proposal and choose “COI” from the drop-down list to indicate a conflict of interest. Inclusion of FY07 OCRP Integration Panel (IP) members in any capacity in the proposal, budget, or any supporting document is considered a conflict of interest and will result in administrative withdrawal of the proposal. A list of the FY07 OCRP IP may be found at http://cdmrp.army.mil/ocrp/panel07.htm

4. LOI Narrative: The LOI Narrative has a one-page limit inclusive of figures, tables, graphs, photographs, diagrams, chemical structures, pictures, pictorials, and cartoons. The LOI Narrative should be a brief description of the research to be conducted.

5. Formatting Guidelines and Submission: The LOI Narrative must be a PDF file, in accordance with the formatting guidelines, and uploaded under the “Required Files” tab of the CDMRP eReceipt system.
6. **PI’s Responsibility:** The PI is responsible for uploading the LOI Narrative (one-page limit) as a PDF file under the “Required Files” tab of the CDMRP eReceipt system.

The electronic PDF file uploaded in the CDMRP eReceipt system is the official pre-application submission file. After conversion of word processing documents to PDF files and before electronic submission, PIs should review their files to ensure that the pre-application complies with the **formatting guidelines**.

Once the PI has completed the pre-application submission process, the eReceipt system will generate a pre-application file. The PI should download the pre-application file (in XML format) and attach it to form SF424 in Block 20 (pre-application) as part of the proposal submission through Grants.gov. Do not convert this file.

7. **AOR Approval:** The pre-application submission does not require approval by the AOR before submission. Please see **Appendix 2** for the definition of an AOR.

B. **LOI Narrative Review**

The LOI Narrative will be administratively reviewed prior to peer review; it will not be reviewed during peer and programmatic reviews.
V. SUBMISSION PROCESS STEP 2: PROPOSAL SUBMISSION

This section describes the process for submission of a proposal, once a pre-application has been submitted. Proposals must be submitted electronically by the AOR through Grants.gov (www.grants.gov). No paper copies will be accepted.

Proposal submission will not be accepted unless a pre-application was previously submitted. The PI and Organization identified in the proposal submitted through Grants.gov should be the same as those identified in the pre-application. If there is a change in PI or Organization after submission of the pre-application, please contact the eReceipt helpdesk at help@cdmrp.org or 301-682-5507.

For complete information regarding forms and submission components, as well as general proposal preparation and submission instructions, please see Appendix 3.

Please note, submission of a proposal requires institutional registration with the Central Contractor Registry (CCR), which requires a Data Universal Number System (DUNS) number, Tax Identification Number (TIN) or Employer Identification Number (EIN), and a Commercial and Government Entity (CAGE) code and must be completed well in advance of Grants.gov registration and proposal submission. Please note that CCR registrations have expirations. Plan accordingly and allow several weeks for these registration processes. Grants.gov will not allow proposals to be submitted unless all of the registration steps have been completed.

A. Proposal Components Summary

Each proposal submission must include the completed Grants.gov application package of forms and attachments identified in www.grants.gov for the US Army Medical Research Acquisition Activity (USAMRAA) program announcement. The package includes:

1. SF-424 (R&R) Application for Federal Assistance Form
   - Pre-application file downloaded from the CDMRP eReceipt system

2. Attachments Form
   - Attachment 1: Project Narrative (two-page limit)
   - Attachment 2: Supporting Documentation
     - References Cited and Acronyms and Symbol Definitions
   - Attachment 3: Federal Agency Financial Plan (if applicable)

3. Research & Related Budget Form
   - Budget Justification

4. Research & Related Project/Performance Site Location(s) Form

5. R&R Subaward Budget Attachment(s) Form (if applicable)
Grants.gov will only notify PIs of changes made to this Program Announcement and/or Application Package if the PI clicks on the “send me change notification emails” link and subscribes to the mailing list on the Opportunity Synopsis Page for this announcement. Please note that if the PI does not subscribe and the Application Package is updated or changed, the original version of the Application Package may not be accepted.
VI. PROPOSAL REVIEW INFORMATION

A. Proposal Review and Selection Overview

All proposals are evaluated by scientists, clinicians, and consumer advocates using a two-tier review process. The first tier is a scientific peer review of proposals against established criteria for determining scientific merit. The second tier is a programmatic review that compares submissions to each other and recommends proposals for funding based on scientific merit and overall goals of the program. Additional information about the two-tier review process used by the CDMRP may be found at http://cdmrp.army.mil.

The Government reserves the right to review all proposals based on one or more of the required attachments or supporting documentation. This will be a blinded review process; the PI and institution names will not be provided during either peer or programmatic review.

B. Review Criteria

1. Peer Review: All proposals will be evaluated according to the following criteria:

   • Innovation
     o What is innovative about the proposed concept.
     o Evidence that the concept is new or in an early stage of development.
     o How exploring the concept may lead to new areas of research.

   • Impact
     o How the project will lead to an original and important contribution to basic or clinical ovarian cancer research.
     o What impact this study will have on the concepts or methods that drive the field of ovarian cancer research.

   • Research Strategy
     o How the scientific rationale and logical reasoning support the project and its feasibility.
2. **Programmatic Review:** Criteria used by the IP to make funding recommendations that maintain the program’s broad portfolio include:

- Ratings and evaluations of the peer reviewers (scientific and consumer),
- Programmatic relevance,
- Relative innovation,
- Relative 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 will be selected by the IP and recommended for funding to the Commanding General, USAMRMC.
VII. COMPLIANCE GUIDELINES

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 formatting guidelines (Appendix 4) 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:

- All attached files are not in PDF, except for the pre-application file.
- Project Narrative exceeds page limit.
- Project Narrative is missing.
- Margins are less than specified in the formatting guidelines.
- Print Area exceeds that specified in the formatting guidelines.
- Spacing is less than specified in the formatting guidelines.
- PI or collaborator’s name or institution is included in the proposal body.
- Use of “I,” “our,” “this institution,” or similar phrases that make it possible to identify the PI or collaborators and/or institution through the references listed.
- Required supporting documentation is missing.
- FY07 OCRP IP members are included in any capacity in the pre-application process, the proposal, budget, and any supporting document. A list of the FY07 OCRP IP members may be found at http://cdmrp.army.mil/ocrp/panel07.

For any other sections of the proposal with a defined page limit, pages exceeding the specified limit will be removed 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.
To protect the public interest, the Federal Government ensures the integrity of Federal programs by only conducting business with responsible recipients. The US Army Medical Research and Materiel Command (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.)

All individuals, regardless of ethnicity, nationality, or citizenship status, may apply as long as they are employed by, or affiliated with, an eligible institution.

**Eligible Institutions:** USAMRMC makes awards to institutions; eligible institutions include for-profit, nonprofit, public, and private organizations, such as universities, colleges, hospitals, laboratories, and companies.

**Historically Black Colleges and Universities/Minority Institutions (HBCU/MI):** A Department of Defense goal is to allocate funds for the Congressionally Directed Medical Research Programs (CDMRP) peer reviewed research to fund proposals from HBCU/MI. This provision is based on guidance from Executive Orders 12876, 12900, and 13021. 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.”

**Government Agencies:** 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.

**Duplicate Submissions:** Submission of the same research project to different award mechanisms within the same program or to other CDMRP programs is discouraged. The Government reserves the right to reject duplicative proposals.
A. Public Law 106-107

Proposals requesting funding from the CDMRP will be submitted through the Federal Government’s single entry portal, Grants.gov, in compliance with Public Law 106-107 (P.L. 106-107). The Federal Financial Assistance Management Improvement Act of 1999, also known as P.L. 106-107, was enacted in November 1999. The purposes of the P.L. 106-107 are to (1) improve the effectiveness and performance of Federal financial assistance programs, (2) simplify Federal financial assistance application and reporting requirements, (3) improve the delivery of services to the public, and (4) facilitate greater coordination among those responsible for delivering services.

Individual program announcements and required forms can also be found on this website. As in previous years, award mechanisms requiring pre-applications including Letter of Intent Narrative, preproposals, and/or nominations will be submitted through the CDMRP eReceipt system at https://cdmrp.org.

B. Grants.gov

Grants.gov is an E-Government initiative to provide a simple, unified electronic storefront for interactions between Principal Investigators (PIs) and the Federal agencies that manage grant funds. The grant community, including state, local, and tribal governments, academia and research institutions, commercial firms and not-for-profits, can access the annual grant funds available across the Federal Government through one website, Grants.gov. In addition to simplifying the grant application process, Grants.gov also creates avenues for consolidation and best practices within each grant-making agency.

In compliance with P.L. 106-107, the USAMRMC requires proposals submitted in response to the program announcement to be submitted through Grants.gov. This requires that organizations register in Grants.gov to submit proposals through the Grants.gov portal. Individual PIs/Project Directors DO NOT register; however, the AOR is required to register. The registration process can take several weeks, so please register as soon as possible.

The following actions are required as part of the registration process. The registration process can take several weeks. If you do business with the Federal Government on a continuing basis, it is likely you have already completed some of the actions, e.g., obtaining a DUNS number or registration in CCR. Detailed information, automated tools, and checklists are available at http://www.grants.gov/applicants/get_registered.jsp

1. Applicant Organization Must Have a Data Universal Number System (DUNS) Number

An organization will need a DUNS number. A DUNS number is a unique nine-character identification number provided by the commercial company Dun & Bradstreet (D&B)
If an organization does not have a DUNS number, an authorized official of the organization can request one by calling 866-705-5711 or online via web registration (http://fedgov.dnb.com/webform/index.jsp). Organizations located outside of the United States can request and register for a DUNS number online via web registration.

2. Applicant Organization Must be Registered with the Central Contractor Registry (CCR)

An organization must be registered with CCR before submitting a grant application through Grants.gov or receiving an award from the Federal Government. CCR validates institution information and electronically shares the secure and encrypted data with Federal agencies’ finance offices to facilitate paperless payments through electronic funds transfer. **CCR registrations have an expiration – please verify the status of your organization’s CCR registration well in advance of the proposal submission deadline.**

You can register by calling the CCR Assistance Center at 888-227-2423 or register online at [http://www.ccr.gov](http://www.ccr.gov). Collecting the information (Employer Identification Number [EIN] or Tax Identification Number [TIN]) can take 1-3 days. If you have the necessary information, online registration will take about 30 minutes to complete, depending upon the size and complexity of your organization. Allow a minimum of 5 business days to complete the entire CCR registration. If your organization does not have either an EIN or TIN, allow at least 2 weeks to obtain the information from the Internal Revenue Service (IRS).

Foreign organizations must obtain a CAGE code prior to registering with the CCR. A CAGE code can be obtained by calling 269-961-7766 or online at [http://www.dlis.dla.mil/Forms/Form_AC135.asp](http://www.dlis.dla.mil/Forms/Form_AC135.asp).

3. Authorized Organizational Representative (AOR) must be registered with Grants.gov

Before submitting a proposal, an organization representative needs to register to submit on behalf of the organization at Grants.gov - [https://apply.grants.gov/OrcRegister](https://apply.grants.gov/OrcRegister). An organization’s E-Business point of contact (POC), identified during CCR registration, must authorize someone to become an AOR. This safeguards the organization from individuals who may attempt to submit proposals without permission. The AOR’s username and password serve as “electronic signatures” when an application is submitted on Grants.gov. **Note: In some organizations, a person may serve as both an E-Business POC and an AOR.**

An AOR must first register with the Grants.gov credential provider at [https://apply.grants.gov/OrcRegister](https://apply.grants.gov/OrcRegister) to obtain a username and password. The AOR must then register with Grants.gov for an account at [https://apply.grants.gov/GrantsgovRegister](https://apply.grants.gov/GrantsgovRegister). Once an AOR has completed the Grants.gov process, Grants.gov will notify the E-Business POC for assignment of user privileges. When an E-Business POC approves an AOR, Grants.gov will send the AOR a confirmation email.
APPENDIX 3

INFORMATION FOR PROPOSAL SUBMISSION

Proposal submission is a two-step process consisting of (1) a pre-application submission through the CDMRP eReceipt system (https://cdmrp.org/) and (2) a proposal submission through Grants.gov (http://www.grants.gov/). This section describes the process for proposal submission. For pre-application submission, see Section IV. Proposal submission will not be accepted unless a pre-application was previously submitted. This appendix outlines how to prepare a proposal application for submission through Grants.gov.

Each submission must include the completed package of forms identified in www.grants.gov for the US Army Medical Research Acquisition Activity (USAMRAA) program announcement. The submission of specific documents will depend upon the award mechanism for which this proposal is being submitted, as specified in Section V and described below. All attachments must be uploaded as a PDF file in accordance with the formatting guidelines in Appendix 4 except for the pre-application XML file.

Fill in the Application Filing Name on the first screen of the Grant Application Package using the CDMRP log number acquired during the pre-application process. Do not fill in the Competition ID.

Click on “Help Mode” (see arrow in Figure 1 below) in the PureEdge tool bar and scroll over the blocks for tips on navigating through the forms in this application package.

![Figure 1: Grants.gov Application PureEdge Toolbar](image)
During award negotiations, the Certificate of Environmental Compliance, the Principal Investigator Safety Program Assurance, regulatory documents related to human subjects and animals, and other documents will be requested from the PIs. At that time, the negotiated indirect rate agreement, Certifications and Assurances for Assistance Agreements, and Representations for Assistance Agreements will be requested from the AOR.

A. SF-424 (R&R), Application for Federal Assistance Form.

This form is required for each application. All appropriate information must be entered into this form to allow for auto-population of all subsequent forms in this application package. The form is self-explanatory, with the following exceptions:

- **Applicant Identifier** box should be filled in with the submitting Institution’s Control Number.
- **State Application Identifier** is not applicable.
- **Block 1 – Type of Submission.** For all submissions the “Application” box should be chosen. For substantial changes that must be made after the original submission, the complete application package must be resubmitted. In these cases, the “Changed/Corrected Application” box must be checked and the Grants.gov tracking number must be entered in Block 4 - Federal Identifier.
- **Block 3 – Date Received by State** is not applicable
• **Block 4 – Federal Identifier Box.** This box will be populated by Grants.gov for an original application, but the Grants.gov tracking number (i.e., the Federal Identifier Number assigned to the original application) must be manually entered for changed or corrected applications.

• **Block 13 – Proposed Project.** The start date should be 9 months to a year from deadline for proposal submission for this award mechanism.

• **Block 14 – Congressional Districts Of.** If applying from a foreign institution enter “00-000” for both applicant and project.

• **Block 17 – Is Application Subject to Review by State Executive Order 12372 Process?** Choose option, b. NO, program is not covered by E.O.12372.

• **Block 19 – Authorized Representative.** The “signature of AOR” is not an actual signature and is automatically completed upon submission of the electronic application package. *Hard copies of applications will not be accepted.*

• **Block 20 – Pre-application box and attachment should be used to attach the pre-application file associated with this proposal. This pre-application file must be downloaded from the CDMRP eReceipt system. Please do not convert this XML file to PDF.*

B. Attachments Form

The following information must be included as attachments to this form in accordance with the formatting guidelines specified in Appendix 4:

**Attachment 1: Project Narrative: Two-page limit.** The Project Narrative is the main body of the proposal. The Project Narrative must be submitted as a single PDF file named “Narrative.pdf,” in accordance with the formatting guidelines specified in Appendix 4.

Due to the blinded nature of the review process for Concept Awards, use of the PI or collaborator’s name and/or institution in the Project Narrative are prohibited and will be cause for administrative withdrawal of the proposal.

The investigator must clearly explain how the proposed research is innovative and relevant to ovarian cancer research. *Preliminary data are not consistent with this award mechanism.* However, applicants must demonstrate logical reasoning and a sound scientific rationale for the proposal to be competitive.

Describe the proposed project using the following outline:

- **Background:** Present the ideas and reasoning behind the proposed work.
- **Specific Aims:** Concisely state the project’s specific aims.
- **Research Strategy:** Describe the experimental design, methods, and analyses including appropriate controls.
- **Innovation:** Describe what is innovative about the proposed research.
• **Impact:** Describe the impact of these studies on the concepts or methods that drive the field of basic or clinical ovarian cancer research. In non-technical terms, state how the results of the proposed research will offer an original and important contribution in ovarian cancer.

The two-page limit of the Project Narrative is inclusive of any figures, tables, graphs, photographs, diagrams, chemical structures, pictures, pictorials, cartoons, and other relevant information needed to judge the proposal.

*The Project Narrative will be available for both peer and programmatic review.*

**Attachment 2: Supporting Documentation.** Upload these sections as a single PDF file named “Support.pdf,” in accordance with the formatting guidelines specified in Appendix 4.

a. **References Cited and Acronyms and Symbol Definitions: Two-page limit.**

- **References Cited:** List a maximum of five 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). The inclusion of Internet URLs to references is encouraged.

- **Acronyms and Symbol Definitions:** Starting on a new page titled “Acronyms and Symbol Definitions,” provide a glossary of acronyms and symbols.

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. *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.*

**Attachment 3: Federal Agency Financial Plan (if applicable).** Proposals from Federal agencies must provide a plan delineating how all funds will be obligated by September 30, 2008, 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 through agreements with foundations, non-Federal institutions, and universities. The Federal Agency Financial Plan must be submitted as a single PDF file named “FedFin.pdf,” in accordance with the formatting guidelines specified in Appendix 4.

C. **Research & Related Budget Form**

An estimate of the total research project cost, with a breakdown by category and year, must accompany each proposal. All costs must be entered in US dollars. Recipients performing research outside of the United States should include the cost in local currency, the rate used for converting to US dollars, and justification/basis for the conversion rate used.

The following cost regulations and principles must be adhered to budget calculations:

- **Subcontracting Indirect Costs:** When the applicant institution calculates its own indirect costs, it can only calculate indirect costs on the first $25,000 of each subaward.
• **Maximum Obligation:** The USAMRMC does not amend grants to provide additional funds for such purposes as reimbursement for unrecovered indirect costs resulting from the establishment of final negotiated rates or for increases in salaries, fringe benefits, and other costs.

• **Cost Regulations and Principles:** Costs proposed must conform to the following regulations and principles:
  
  o **Commercial Firms:** Federal Acquisition Regulation (FAR) Part 31 and Defense FAR Supplement Part 31 (http://farsite.hill.af.mil), Contract Cost Principles and Procedures.
  
  o **Educational Institutions:** OMB Circular A-21, Cost Principles for Educational Institutions.
  
  o **Nonprofit Organizations:** OMB Circular A-122, Cost Principles for Nonprofit Organizations. OMB Circular A-133, Audits of Institutions of Higher Education and Other Nonprofit Organizations.
  
  o **State, Local, and Tribal Governments:** OMB Circular A-87, Cost Principles for State, Local, and Indian Tribal Governments.
  
  o **Cost of Preparing Proposals:** The cost of preparing proposals in response to this program announcement is not considered an allowable direct charge to any resultant contract, grant, or cooperative agreement. It is, however, an allowable expense to the bid and proposal indirect cost specified in FAR 31.205-18, and OMB Circulars A-21 and A-122.

**Section A & B – Senior/Key Person and Other Personnel:** The basis for labor costs should be predicated upon actual labor rates or salaries. Budget estimates may be adjusted upward to forecast salary or wage cost-of-living increases that will occur during the period of performance. The proposal should separately identify and explain the ratio applied to base salary/wage for cost-of-living adjustments and merit increases in the budget justification (Section K).

For each key staff member identified on the budget form, list the percentage of each appointment to be spent on this project.

**Section C – Equipment Description:** It is DOD policy 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 separately negotiated.

An itemized list of permanent equipment is required, showing the cost for each item. Permanent equipment is any article of nonexpendable tangible property having a useful life of more than 2 years and an acquisition cost of $5,000 or more per unit. The justification for the cost of each item of equipment included in the budget must be disclosed in the budget justification (Section K) to include:

  • **Vendor Quote:** Show name of vendor and number of quotes received and justification if intended award is to other than the lowest bidder.
• Historical Cost: Identify vendor, date of purchase, and whether or not cost represented the lowest bid. Include reason(s) for not soliciting current quotes.

• Estimate: Include rationale for estimate and reasons for not soliciting current quotes.

• Special test equipment to be fabricated by the contractor for specific research purposes and its cost.

• Standard equipment to be acquired and modified to meet specific requirements, including acquisition and modification costs; list separately.

• Existing equipment to be modified to meet specific research requirements, including modification costs. Do not include as special test equipment those items of equipment that, if purchased by the contractor with contractor funds, would be capitalized for Federal income tax purposes.

• 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.

• Commercial organizations are expected to possess the necessary plant and equipment to conduct the proposed research. Equipment purchases for commercial organizations will be supported only in exceptional circumstances.

Section D – Travel

• Travel costs to attend one scientific/technical meeting per year. Costs should not exceed $1,800.

• Travel costs associated with the execution of the proposed work. 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 United States, including between foreign countries, requires prior approval from USAMRAA 90 days before travel.

• Travel to CDMRP-required meetings (if applicable) (Section II.C). Costs should be reasonable.

Section E – Participant/Trainee Support Costs: This section is self-explanatory.

Section F – Other Direct Costs (as applicable)

Section F.1 – Materials and Supplies (Consumables): The justification (to be included in Section K) supporting material and supply (consumable) costs should include a general description of expendable equipment and supplies. If animals are to be purchased, state the species, strain (if applicable), and the number to be used. If human cell lines are to be purchased, state the source and the description.

Section F.2 – Publication Costs: This section is self-explanatory.
Section F.3 – Consultant Services: Regardless of whether funds are requested, the justification (to be included in Section K) should include the names and organizational affiliations of all consultants. State the daily consultant fee, travel expenses, nature of the consulting effort, and why consultants are required for the proposed research project.

Section F.4 – ADP/Computer Services: This section is self-explanatory.

Section F.5 – Subaward/Consortium/Contractual Costs: On the project’s Research and Related Budget Form, enter the total funds requested for (1) all subaward/consortium organization(s) proposed for the project and (2) any other contractual costs proposed for the project.

Section F.6 – Equipment or Facility Rental/User Fees: This section is self-explanatory.

Section F.7 – Alterations and Renovations: Not allowable.

Sections F.8–F.10 – Research-Related Subject Costs: Include itemized costs of subject participation in the research study. These costs are strictly limited to expenses specifically associated with the proposed study. The USAMRMC will not provide funds for ongoing medical care costs that are not related to a subject’s participation in the research study.

Sections F.8–F.10 – Other Direct Costs (if applicable): Include other anticipated direct costs that are not specified elsewhere in the budget. Unusual or expensive items should be fully explained and justified in Section K.

Section G – Direct Costs: This section is self-explanatory. All direct and indirect costs of any subaward must be included in the total direct costs of the primary award.

Section H – 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 along with a statement identifying whether the proposed rates are provisional or fixed. If negotiated forecast rates do not exist, provide sufficient detail in the budget justification (Section K) regarding a determination that the costs included in the forecast rate are allocable according to applicable FAR/DFARS or OMB Circular provisions. Commercial firms can also visit www.dcaa.mil for additional information on indirect rates. Disclosure should be sufficient to permit a full understanding of the content of the rate(s) and how it was established. When the applicant institution calculates its own indirect costs, it can only calculate indirect costs on the first $25,000 of each subaward.
As a minimum, justification for indirect costs should identify:

- All individual cost elements included in each forecast rate;
- The basis used to prorate indirect expenses to cost pools, if any;
- How each rate was calculated; and
- The distribution basis of each developed rate.

Section I – Total Direct and Indirect Costs: This section is self-explanatory.

Section J – Fee: A profit or fixed fee is not allowable on grants or cooperative agreements. If a profit/fee is negotiated, a contract will be awarded. Any fixed fee applied to the research project must be listed and any claimed Facilities Capital Cost of Money supported by DD Form 1861 (www.dtic.mil/whs/directives/infomgt/forms/forminfo/forminfopage2192.html) must be submitted with the proposal.

Section K – Budget Justification: The Budget Justification for the entire performance period must be attached as a PDF file named “Justification.pdf” to the Research & Related Budget – Section K (under budget period one). Organizations must provide sufficient detail and justification so that the Government can determine the proposed costs to be allocable and reasonable for the proposed research effort.

The budget justification must include information for all budget periods. This file must be uploaded for budget period one before you will be allowed to access subsequent budget periods.

D. Research & Related Project/Performance Site Location(s) Form

Indicate the primary site where the work will be performed. If a portion of the work will be performed at any other site(s), include the name and address for each collaborating location in the data fields provided. If more than eight performance site locations are proposed, provide the requested information in a separate file and attach to this form. Please note that each additional research site requesting funds will require a subcontract budget.

E. R&R Subaward Budget Attachment(s) Form (optional form; use if applicable)

Please note that the files to be attached to the R&R Subaward Budget Attachment(s) Form must be PureEdge documents. Extract an R&R Subaward Budget Attachment for each subaward, using the button provided on this form. Save each attachment to your computer and complete the form(s).

The Budget Justification for each subaward must be attached as a PDF file named “Justification_LastName.pdf” (where “LastName is the investigator of the subaward) to the Research & Related Budget – Section K for that subaward. Each subaward budget justification must include information for all budget periods. This file must be uploaded for budget period one before you will be allowed to access subsequent budget periods for the subaward. Once all subaward budget files are completed, attach all subaward budget file(s) for this application to the R&R Subaward Budget Attachment(s) Form.
The DUNS number for each subaward site should be included on this form.

A description of services or materials that are to be awarded by subcontract or subgrant is required. Organizations must provide sufficient detail and justification so that the Government can determine the proposed costs to be allocable and reasonable for the proposed research effort. The following information must be provided on subawards totaling $10,000 or more:

- Identification of the type of award to be used (e.g., cost reimbursement, fixed price);
- Identification of the proposed subcontractor or subgrantee, if known, and an explanation of why and how the subcontractor or subgrantee was selected or will be selected;
- Whether the award will be competitive and, if noncompetitive, rationale to justify the absence of competition; and
- The proposed acquisition price.
- The applicant’s cost or price analysis for the subgrant or subcontract proposed price (applicable only if the award exceeds $500,000).

If the resultant award is a contract that exceeds $500,000 and the applicant is a large business or an educational institution (other than a Historically Black College or University/Minority Institution), the applicant is required to submit a subcontracting plan for small business and small disadvantaged business concerns, in accordance with FAR 19.7. A mutually agreeable plan will be incorporated as part of the resultant contract.
APPENDIX 4

FORMATTING GUIDELINES

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 a computer screen.

- **Document Format:** All attachments must be in PDF, except for the pre-application file (XML file) attached to block 20 of SF-424.
- **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 (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 (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, Principal Investigators 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 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 encouraged.
- **Language:** English.
- **Headers and Footers:** Should not be used.
- **Page Numbering:** Should not be used.

All attachments that require signatures must be filled out, printed, signed, scanned, and then uploaded as a PDF file.
A. Award Notices

Each Principal Investigator (PI) will receive notification of the award status of his or her proposal. Peer review summary statements for Concept Awards will not be posted to the Congressionally Directed Medical Research Programs (CDMRP) eReceipt system. PIs can expect to receive this notification approximately 4 weeks after programmatic review.

B. Administrative Requirements

Awards are made to organizations, not individuals. The PI 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 Department of Defense [DOD] Grant and Agreement Regulations) to be eligible for an award. Any organization requesting receipt of an award through this program 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.

If allowed, a change in institutional affiliation will require the investigator to resubmit the entire proposal packet through his or her new institution to include any regulatory documentation that may require protocols, etc., to be approved for the new institution. The investigator’s original institution must agree to relinquish the award. Any delay in the submission of the new information will result in a delay in contracting and regulatory review and a subsequent delay in resuming work on the project.

C. Award Negotiation

Award negotiation consists of discussions, reviews, and justifications of critical issues involving the US Army Medical Research Acquisition Activity (USAMRAA). A Contract Specialist and/or representative from the USAMRAA will contact the Contract Representative authorized to negotiate contracts and grants at the PI’s institution. Additional documentation and justifications related to the budget may be required as part of the negotiation process.

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

D. Disclosure of Proprietary Information outside the Government

By submitting a proposal, the PI understands that proprietary information may be disclosed outside the Government for the sole purpose of technical evaluation. The US Army Medical Research and Materiel Command (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.

**E. Government Obligation**

PIs 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. PIs 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.

**F. Information Service**

PIs may use the technical reference facilities of the National Technical Information Service (www.ntis.gov), 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.

**G. Inquiry Review Panel**

PIs 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.

**H. Title to Inventions and Patents**

In accordance with the Bayh-Dole Act (Title 35, United States Code, Sections 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.

**I. 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.
APPENDIX 6
REGULATORY REQUIREMENTS AND REVIEWS

The Principal Investigator (PI) may not use, employ, or subcontract for the use of any human subjects, human biological substances, or laboratory animals until applicable regulatory documents are requested, reviewed, and approved by the US Army Medical Research and Materiel Command (USAMRMC).

Concurrent with the US Army Medical Research Acquisition Activity (USAMRAA) negotiation, the Office of Surety, Safety and Environment will review the Certificate of Environmental Compliance and the Principal Investigator Safety Program Assurance form to be submitted upon request. The applicable USAMRMC regulatory office will review documents related to research involving human subjects, human anatomical substance use, and animal use, which should be submitted upon request to ensure that Department of Defense (DOD) regulations are met.

A. Certificate of Environmental Compliance

The Certificate of Environmental Compliance will be requested at award negotiations. If multiple research sites/institutions are funded in the proposal, then a Certificate of Environmental Compliance for each site will also be requested.

B. Safety Program Documents

The Principal Investigator Safety Program Assurance form will be requested at award negotiations.

A Facility Safety Plan is required; it will be requested at award negotiations. A Facility Safety Plan from the PI’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.amedd.army.mil/docs/rcq/sohd/Facility_Safety_Plan_Approved_Institutions.pdf. If the PI’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.

C. Research Involving Animal Use

Specific documents relating to the use of animals in the proposed research will be requested by the Congressionally Directed Medical Research Programs (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 the USAMRMC Office of Research Protections (ORP; formerly Regulatory Compliance and Quality), must review and approve all animal use
prior to the start of working with animals. PIs 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). Allow 2 to 4 months for regulatory review and
approval processes for animal studies.

Specific requirements for research involving animals can be found at

D. Research Involving Human Subjects or Biological Substances

Projects involving human subjects or specimens will not be supported through this mechanism
unless they are exempt under 32 CFR 219.101(b) (4), qualify for expedited review under 32 CFR
219.110 or 21 CFR 56.1102, or involve the use of only commercially available anonymized
specimens.

In addition to local Institutional Review Board (IRB) approval or determination of exempt status,
a second level of review is required by the DOD for concurrence with the exempt status. This
second review is conducted by the USAMRMC Office of Research Protections, Human
Research Protection Office (HRPO). Documents supporting the exempt status of the project will
be requested at a later date. These documents will include documentation of local IRB
determination of exempt status and the completed USAMRMC Office of Research Protections
Claim of Exemption Form. For studies using only commercially available specimens, the
USAMRMC Office of Research Protections Claim of Exemption Form will be requested.

1. Requirements: Specific requirements for research involving human subjects or human
biological substances can be found at https://mrmc.amedd.army.mil/rodorptoolkit.asp.

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.

2. Informed Consent Form: An informed consent form template is located at

3. Intent to Benefit: Investigators must consider the requirements of Title 10 United States
applicable to DOD-sponsored research before writing a research protocol. 10 USC 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 a DOD-supported experiment 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.

4. **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 that are 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.
APPENDIX 7

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 US Army Medical Research and Materiel Command 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. Additional reporting may be required as stipulated during award negotiations. Copies of all scientific publications and patent applications resulting from Congressionally Directed Medical Research Programs funding should be included in the progress report. The Government reserves the right to request additional 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 Institutional Review Board (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.** Principal Investigators 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.
APPENDIX 8

ACRONYM LIST

ACURO..........................Animal Care and Use Office
ADP................................Automated Data Processing
AOR...............................Authorized Organizational Representative
ASDRP...........................Autism Spectrum Disorder Research Program
AVI ................................Audio Video Interleave
BCRP.............................Breast Cancer Research Program
CCR...............................Central Contractor Registration
CDMRP..........................Congressionally Directed Medical Research Programs
CFDA............................Catalog of Federal Domestic Assistance
CFR...............................Code of Federal Regulations
cGMP.............................Current Good Manufacturing Practices
CAGE.............................Commercial and Government Entity
COI.................................Conflicts of Interest
CMLRP..........................Chronic Myelogenous Leukemia Research Program
CR .................................Contract Representative
DFARS..........................Department of Defense Federal Acquisition Supplement
DOD...............................Department of Defense
DODGAR ........................Department of Defense Grant and Agreement Regulations
DUNS.............................Data Universal Number System
EIN.................................Employer Identification Number
EPLS..............................Excluded Parties List System
FAR...............................Federal Acquisition Regulation
FDA...............................Food and Drug Administration
FY .................................Fiscal Year
GCP...............................Good Clinical Practice
GLP...............................Good Laboratory Practice
GWVIRP........................Gulf War Veterans’ Illnesses Research Program
HBCU/MI ......................Historically Black Colleges and Universities/Minority Institutions
HIPAA...........................Health Insurance Portability and Accountability Act
hES.................................Human Embryonic Stem
HRPO.............................Human Research Protection Office
HSRRB ........................Human Subjects Research Review Board
IDE.................................Investigational Device Exemption
IND ...............................Investigational New Drug
IP.................................Integration Panel
IRB.................................Institutional Review Board
IRS...............................Internal Revenue Service
JPEG.............................Joint Photographic Experts Group
LAR...............................Legally Authorized Representative
LOI.................................Letter of Intent
M.................................Million
MB.................................Megabyte

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