

TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN<sup>1</sup>

| Allegation reporting respondent | Number of respondents | Number of responses per respondent | Total annual responses | Average burden per response | Total hours |
|---------------------------------|-----------------------|------------------------------------|------------------------|-----------------------------|-------------|
|                                 | 700                   | 1                                  | 700                    | 2.25                        | 175         |

<sup>1</sup> There are no capital costs or operating and maintenance costs associated with this collection of information.

<sup>2</sup> 15 minutes.

Dated: May 16, 2014.

**Leslie Kux,**

*Assistant Commissioner for Policy.*

[FR Doc. 2014–11922 Filed 5–22–14; 8:45 am]

**BILLING CODE 4160–01–P**

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. FDA–2014–N–0610]

#### Increasing the Quality and Efficiency of Clinical Trials

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing the availability of grant funds for the support of the efforts of the Center for Drug Evaluation and Research/Office of Medical Policy to increase the quality and efficiency of clinical trials. The Food and Drug Administration (FDA), Center for Drug Evaluation and Research (CDER), Office of Medical Policy is announcing its intent to accept and consider a single-source application for the award of a grant to the Duke University's Duke Translational Medicine Institute (DTMI).

**DATES:** The application due date is June 30, 2014, by 11:59 p.m. Eastern Time. The expiration date is July 1, 2014.

**ADDRESSES:** Submit electronic applications to: <http://www.grants.gov>. For more information, see section III of the **SUPPLEMENTARY INFORMATION** section of this notice.

#### FOR FURTHER INFORMATION CONTACT:

Mark Lauda, Office of Medical Policy, Center for Drug Evaluation and Research, Food and Drug Administration, 10990 New Hampshire Ave., Bldg. 51, Rm. 2212, Silver Spring, MD 20993, 301–796–0381, email: [Mark.Lauda@fda.hhs.gov](mailto:Mark.Lauda@fda.hhs.gov); or Lisa Ko, Office of Acquisition & Grants Services, Food and Drug Administration, 5630 Fishers Lane, Rockville, MD, 240–402–7592, email: [Lisa.Ko@fda.hhs.gov](mailto:Lisa.Ko@fda.hhs.gov).

For more information on this funding opportunity announcement (FOA) and to obtain detailed requirements, please

refer to the full FOA located at <http://www.grants.gov>. Search by Funding Opportunity Number: RFA–FD–14–017.

#### SUPPLEMENTARY INFORMATION:

##### I. Funding Opportunity Description

RFA–FD–14–017

93.103

##### A. Background

It has long been recognized that the clinical trial enterprise will need to evolve in order to meet the demand to provide data to support evidence-based decisionmaking. A memorandum of understanding (MOU) between FDA and Duke University published in the **Federal Register** on November 23, 2007, served as the basis for the establishment of the Clinical Trials Transformation Initiative (CTTI). CTTI is a public-private partnership whose mission is to identify and promote practices that will increase the quality and efficiency of clinical trials. This award will be made to DTMI within Duke University to identify and implement projects and disseminate resulting findings that will increase the quality and efficiency of clinical trials, CTTI's mission.

CTTI membership is broad and includes stakeholders from government, industry, patient advocacy and consumer groups, professional societies, clinical research organizations, and academia. CTTI helps to effect change through the conduct of projects that identify existing inefficiencies, elucidate superior practices, and/or provide innovative approaches to evidence generation and medical product development. CTTI conducts projects that are either: (1) Proposed by its member organizations, including FDA, developed during review by its Steering Committee, and endorsed by its Executive Committee or (2) responsive to urgent needs of FDA.

The opportunity for meaningful interaction with a broad set of stakeholders committed to improving the clinical trial enterprise and also the ability to rapidly gather data to address emerging issues offer significant value to the clinical trial enterprise. Since its inception, CTTI has undertaken many projects that have direct relevance to FDA's mission, including

investigational new drug (IND) safety reporting, clinical trial monitoring, use of central investigational review boards, and antibacterial drug development.

##### B. Research Objectives

The goals of this program are to develop and maintain an administrative and scientific infrastructure to support the creation and execution of a series of projects under the auspices of CTTI that will increase the quality and efficiency of clinical trials. The following are examples of activities that could be supported by this grant:

- Maintaining an adequate administrative and scientific infrastructure to implement all related projects under this collaborative effort.
- Identifying and/or hiring a sufficient number of qualified personnel to conduct activities, including project management, such as review of project milestones for degree of completion, preparation/reporting of project findings, periodic and final reports, and for subsequent distribution in the public domain.
- Developing plans for the conduct of identified projects.
- Identifying, securing, and/or building, and effectively leveraging other resources for the conduct of identified projects.
- Upon completion of a given project, generating project results and recommendations and proposing related studies/projects, if needed, to build on the findings of the project and continuing to leverage established resources and personnel.

##### C. Eligibility Information

The following organization is eligible to apply: DTMI located within Duke University.

## II. Award Information/Funds Available

##### A. Award Amount

This is a multiyear grant. FDA/CDER intends to fund up to \$7,500,000 in total costs (direct and indirect) in Fiscal Year 2014. Awards are contingent upon the availability of funds.

Subject to the availability of Federal funds and successful performance of the FOA's stated goals and objectives, four additional years of support may be

available. Funding beyond the first year will be noncompetitive and will depend on: (1) Satisfactory performance during the preceding year and (2) the availability of Federal fiscal year funds.

Application budgets need to reflect the actual needs of the proposed project and should not exceed the following in total costs (direct and indirect):

Year 01: \$7,500,000  
Year 02: \$7,500,000  
Year 03: \$7,500,000  
Year 04: \$7,500,000  
Year 05: \$7,500,000

#### B. Length of Support

The scope of the proposed project should determine the project period. The maximum project period is 5 years.

### III. Electronic Application, Registration, and Submission

Only electronic applications will be accepted. To submit an electronic application in response to this FOA, applicants should first review the full announcement located at <http://www.grants.gov>. Search by Funding Opportunity Number: RFA-FD-14-017. (FDA has verified the Web site addresses throughout this document, but FDA is not responsible for any subsequent changes to the Web sites after this document publishes in the **Federal Register**.) For all electronically submitted applications, the following steps are required.

- Step 1: Obtain a Dun and Bradstreet (DUNS) Number
- Step 2: Register With System for Award Management (SAM)
- Step 3: Obtain Username & Password
- Step 4: Authorized Organization Representative (AOR) Authorization
- Step 5: Track AOR Status
- Step 6: Register With Electronic Research Administration (eRA) Commons

Steps 1 through 5, in detail, can be found at [http://www07.grants.gov/applicants/organization\\_registration.jsp](http://www07.grants.gov/applicants/organization_registration.jsp). Step 6, in detail, can be found at <https://commons.era.nih.gov/commons/registration/registrationInstructions.jsp>. After you have followed these steps, submit electronic applications to: <http://www.grants.gov>.

Dated: May 15, 2014.

**Leslie Kux,**

*Assistant Commissioner for Policy.*

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. FDA-2011-D-0835]

#### Guidance for Institutional Review Boards, Clinical Investigators, and Sponsors: Considerations When Transferring Clinical Investigation Oversight to Another Institutional Review Board; Availability

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing the availability of a guidance entitled "Guidance for IRBs, Clinical Investigators, and Sponsors: Considerations When Transferring Clinical Investigation Oversight to Another IRB." The guidance announced in this document discusses regulatory responsibilities of institutional review boards (IRBs), clinical investigators, and sponsors when oversight of a previously approved clinical investigation under FDA's jurisdiction is transferred from one IRB to another IRB. The guidance also addresses questions that have been previously raised concerning procedures and processes that are required and/or recommended by FDA when such oversight is transferred.

**DATES:** Submit either electronic or written comments on Agency guidances at any time.

**ADDRESSES:** Submit written requests for single copies of this guidance to the Division of Drug Information, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 51, rm. 2201, Silver Spring, MD 20993-0002 (1-888-463-6332 or 301-796-3400); or the Office of Communication, Outreach and Development (HFM-40), Center for Biologics Evaluation and Research, Food and Drug Administration, 1401 Rockville Pike, suite 200N, Rockville, MD 20852-1448 (1-800-835-4709 or 301-827-1800); or the Division of Small Manufacturers, International, and Consumer Assistance, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, rm. 4622, Silver Spring, MD 20993 (1-800-638-2041 or 301-796-7100). Send one self-addressed adhesive label to assist the office in processing your requests. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the guidance document.

Submit electronic comments on the guidance to <http://www.regulations.gov>.

Submit written comments to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

#### FOR FURTHER INFORMATION CONTACT:

Bridget Foltz, Office of Good Clinical Practice, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 32, rm. 5174, Silver Spring, MD 20993, 301-796-8340.

#### SUPPLEMENTARY INFORMATION:

##### I. Background

FDA is announcing the availability of a guidance entitled, "Guidance for IRBs, Clinical Investigators, and Sponsors: Considerations When Transferring Clinical Investigation Oversight to Another IRB." The guidance discusses the regulatory responsibilities of IRBs, clinical investigators, and sponsors when oversight of a previously approved clinical investigation under FDA's jurisdiction is transferred from one IRB to another IRB. In particular, the guidance discusses eight steps to be considered when transferring oversight of a previously approved clinical investigation from one IRB to another IRB. These include identifying those studies for which IRB oversight is being transferred; ensuring availability and retention of pertinent records; establishing an effective date for the transfer of oversight; conducting a review of the study(ies) by the receiving IRB, where appropriate; confirming or establishing the date for the next continuing review; determining whether the consent form needs to be revised; notifying the key parties; and updating IRB registration information. The IRB transfer process is expected to vary depending on the reasons for the transfer, the parties involved, and the number and risk of the studies being transferred.

To enhance human subject protections and reduce regulatory burden, FDA and the Office for Human Research Protections (OHRP) have been actively working to harmonize the agencies' regulatory requirements and guidance for human subjects research. This guidance document was developed as a part of these efforts and in consultation with OHRP.

In the **Federal Register** of June 12, 2012 (77 FR 34958), FDA announced the availability of the draft guidance of the same title. FDA received several comments on the draft guidance and considered them in preparing the final guidance. In response to the comments, FDA added a recommendation that the receiving IRB notify the sponsor if it decides to suspend or terminate study