initial merit review, all applications will:

- Undergo a peer review by a Special Emphasis Panel. The SEP will be selected from the NIH pool of scientists or recommendations from the National Immunization Program to serve as reviewers on SEPs. Applications will be ranked for the secondary review according to scores submitted by the SEP. Only those applications deemed to have the highest scientific merit by the review group, generally the top half of the applications under review, will be discussed and assigned a priority score.
  - Receive a written critique.
- Receive a second programmatic level review by the Office of Science, National Immunization Program.

Award Criteria: Criteria that will be used to make award decisions during the programmatic review include:

- Scientific merit (as determined by peer review)
  - Availability of funds
  - Programmatic priorities
  - Proposed budget

V.3. Anticipated Announcement and Award Dates

Anticipated Award Date: August 31, 2005.

#### VI. Award Administration Information

## VI.1. Award Notices

Successful applicants will receive a Notice of Award (NoA) from the CDC Procurement and Grants Office. The NoA shall be the only binding, authorizing document between the recipient and CDC. The NoA will be signed by an authorized Grants Management Officer, and mailed to the recipient fiscal officer identified in the application.

Unsuccessful applicants will receive notification of the results of the application review by mail.

VI.2. Administrative and National Policy Requirements

45 CFR Part 74 and Part 92.

For more information on the Code of Federal Regulations, see the National Archives and Records Administration at the following Internet address: http://www.access.gpo.gov/nara/cfr/cfr-table-search.html.

The following additional requirements apply to this project:

- AR-1 Human Subjects Requirements
- AR-2 Requirements for Inclusion of Women and Racial and Ethnic Minorities in Research
  - AR-6 Patient Care
  - AR-7 Executive Order 12372
- AR–8 Public Health System

Reporting Requirements

- AR–10 Smoke-Free Workplace Requirements
  - AR-11 Healthy People 2010AR-12 Lobbying Restrictions
- AR–14 Accounting System
  Requirements
  - AR–15 Proof of Non-Profit Status
  - AR-22 Research Integrity
- AR–23 States and Faith-Based Organizations
- AR–24 Health Insurance Portability and Accountability Act Requirements

Additional information on these requirements can be found on the CDC Web site at the following Internet address: http://www.cdc.gov/od/pgo/funding/ARs.htm.

# VI.3. Reporting

You must provide CDC with an original, plus two hard copies of the following reports:

- 1. Interim progress report, (use form PHS 2590, OMB Number 0925–0001, rev. 9/2004 as posted on the CDC Web site) quarterly during the project. The progress report sent no later than 90 days before the end of the first half of the budget period will serve as your non-competing continuation application, and must contain the following additional elements:
  - a. Reports of participant enrollment.
  - b. Progress in analysis.
- c. Progress Toward Measures of Effectiveness.
- d. Additional Information Requested by Program.
- 2. Financial status report, no more than 90 days after the end of the budget period.
- 3. Final financial and performance reports, no more than 90 days after the end of the project period.

These reports must be mailed to the Grants Management Specialist listed in the "Agency Contacts" section of this announcement.

#### VII. Agency Contacts

We encourage inquiries concerning this announcement. For general questions, contact: Technical Information Management Section, CDC Procurement and Grants Office, 2920 Brandywine Road, Atlanta, GA 30341, Telephone: 770–488–2700.

For scientific/research issues, contact: Susan Chu, PhD, MSPH, Extramural Program Official, Centers for Disease Control and Prevention, MS E–05, 1600 Clifton Road, Atlanta, GA 30333, Telephone: 404 639–8727, E-mail: SChu@cdc.gov.

For questions about peer review, contact: Mary Lerchen, DrPH, Scientific Review Administrator, CDC/Office of Public Health Research, One West Court Square, Suite 7000, MS D-72, Telephone: 404-371-5277, Fax: 404-371-5215, E-mail: *MLerchen@cdc.gov*.

For financial, grants management, or budget assistance, contact: Mattie Jackson, Grants Management Specialist, CDC Procurement and Grants Office, 2920 Brandywine Road, Atlanta, GA 30341, Telephone: 770–488–2696, Email: mij3@cdc.gov.

#### **VIII. Other Information**

This and other CDC funding opportunity announcements can be found on the CDC Web site, Internet address: http://www.cdc.gov. Click on "Funding" then "Grants and Cooperative Agreements." http://www.cdc.gov/nip and http://www.cdc.gov/flu.

Dated: May 6, 2005.

#### William P. Nichols,

Director, Procurement and Grants Office, Centers for Disease Control and Prevention. [FR Doc. 05–9453 Filed 5–11–05; 8:45 am] BILLING CODE 4163–18–P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Centers for Disease Control and Prevention

## Influenza Vaccination of Children and Accompanying Adults: Mass Vaccination vs Vaccination in Routine Care

Announcement Type: New. Funding Opportunity Number: RFA IP05–094.

Catalog of Federal Domestic Assistance Number: 93.185. Letter of Intent Deadline: June 13,

Application Deadline: June 27, 2005.

# I. Funding Opportunity Description

Authority: Section 311 [42 U.S.C. 243] and 317(k)(1) [42 U.S.C. 247b(k)(1)] of the Public Health Service Act, as amended.

Background: Epidemics of influenza have been responsible for an average of approximately 36,000 deaths/year in the United States during 1990–1999. Influenza viruses also can cause pandemics, during which rates of illness and death from influenza-related complications can increase worldwide. Influenza viruses cause disease among all age groups. Rates of infection are highest among children, but rates of serious illness and death are highest among persons aged greater than or equal to 65 years and persons of any age who have medical conditions that place them at increased risk for complications from influenza.

Influenza vaccination is the primary method for preventing the disease and its severe complications. In 2004, the Advisory Committee on Immunization Practices (ACIP) recommended that healthy children aged 6-23 months be vaccinated against influenza because they are at increased risk for influenzarelated hospitalization. In addition, vaccination is recommended for their household contacts and out-of-home caregivers. Vaccination is also recommended for contacts of children aged zero to five months because influenza vaccines have not been approved by FDA for use among children aged greater than six months.

*Purpose:* The purpose of the program is to fund research to conduct an incremental economic evaluation of vaccination of healthy children aged 6-23 months, and the adults who accompany them, in mass vaccination clinic settings compared with vaccination at routine health care visits. The new ACIP recommendations may affect the capacity of the pediatric health care infrastructure to provide vaccination services. We want to know whether mass clinics are an economically and financially viable alternative to providing influenza vaccination during routine visits. This question must be answered from the societal perspective, but the perspective of the pediatric health care provider is also important because that is the level at which implementation would likely occur.

Mass vaccination clinics could include scheduled or walk-in visits for influenza vaccination conducted in a variety of ways. For example, influenza vaccination can be offered on specified days each week during flu season, during a specified time period in a certain month, in schools or other specialty clinics, and could incorporate the offering of vaccine to adults who accompany children to be vaccinated. Applicants are encouraged to estimate costs under the widest variety of possible conditions.

This program addresses the "Healthy People 2010" focus area(s) of Immunization and Infectious Diseases.

Measurable outcomes of the program will be in alignment with the performance goal for the Centers for Disease Control and Prevention's (CDC) National Immunization Program (NIP) to reduce the number of indigenous vaccine-preventable diseases.

Research Objectives:

• To estimate the societal- and provider-perspective incremental cost-effectiveness of influenza vaccination of healthy children aged 6–23 months and, if possible, adults who accompany them

in mass vaccination clinic settings compared with vaccination at routine health care visits.

- To estimate the opportunity costs associated with mass vaccination clinics.
- To assess the effect of practice type, including pediatric and family medicine, solo and multiple physician/multiple specialty settings, on cost-effectiveness.
- To assess the acceptability of mass vaccination clinics to providers and parents of patients compared with vaccination in routine scheduled visits.

Activities: Awardee activities for this

program are as follows:

- Identify or develop theoretical and empirical models for analyzing the incremental cost-effectiveness of vaccination of healthy children aged 6–23 months, and if possible the adults who accompany them, in (1) mass vaccination clinic settings and (2) at private routine health care facilities.
- Identify appropriate theoretical and empirical models for assessing the acceptability of mass vaccination clinics to providers and parents of patients.
- Develop a study design suitable for data collection and analysis.
- Categorize the resources required to provide influenza vaccination in the two settings, including resources reallocated by the use of mass vaccination clinics.
- Determine sites, methods, and feasibility of protocol prior to implementation.
- Identify key staff and established resources/expertise available to develop study models.
- Collaboratively disseminate research findings in peer reviewed publications and for use in determining national policy.

In a cooperative agreement, CDC staff is substantially involved in the program activities, above and beyond routine grant monitoring. CDC Activities for this program are as follows:

- Provide CDC investigator(s) to monitor the cooperative agreement as project officer(s).
- Participate as active project team members in the development, implementation and conduct of the research project and as coauthors of all scientific publications that result from the project.
- Provide technical assistance on the selection and evaluation of data collection and data collection instruments.
- Assist in the development of research protocols for Institutional Review Boards (IRB) review. The CDC IRB will review and approve the project protocol initially and on at least an

annual basis until the research project is completed.

- Contribute subject matter expertise in the areas of epidemiologic methods and statistical analysis, health economics, and survey research consultation.
- Participate in the analysis and dissemination of information, data and findings from the project, facilitating dissemination of results.
- Serve as liaisons between the recipients of the project award and other administrative units within the CDC.
- Facilitate meetings between awardee and CDC to coordinate planned efforts and review progress.

#### **II. Award Information**

*Type of Award:* Cooperative Agreement.

CDC involvement in this program is listed in the Activities Section above.

Mechanism of Support: U01. Fiscal Year Funds: 2005.

Approximate Total Funding: \$150,000. (Includes direct and indirect costs. This amount is an estimate, and is subject to availability of funds.)

Approximate Number of Awards: 1. Approximate Average Award: \$150,000. (Includes direct and indirect costs. This amount is for the first 12month budget period.)

Floor of Award Range: None.

Ceiling of Award Range: \$150,000. (Includes direct and indirect costs. This ceiling is for the first 12-month budget period.)

Anticipated Award Date: August 31, 2005.

Budget Period Length: 12 months. Project Period Length: 2 years.

Throughout the project period, CDC's commitment to continuation of awards will be conditioned on the availability of funds, evidence of satisfactory progress by the recipient (as documented in required reports), and the determination that continued funding is in the best interest of the Federal Government.

# **III. Eligibility Information**

III.1. Eligible applicants

Applications are limited to public and private nonprofit organizations and by governments and their agencies, such as: (For profit organizations are not eligible under Section 317(k)(1) [42 U.S.C. 247b(k)(1) of the Public Health Service Act, as amended.)

- Public nonprofit organizations
- Private nonprofit organizations
- Small, minority, women-owned businesses
  - Universities
  - Colleges

- · Research institutions
- Hospitals
- Community-based organizations
- Faith-based organizations
- Federally recognized Indian tribal governments
  - Indian tribes
  - Indian tribal organizations
- State and local governments or their Bona Fide Agents (this includes the District of Columbia, the Commonwealth of Puerto Rico, the Virgin Islands, the Commonwealth of the Northern Marianna Islands, American Samoa, Guam, the Federated States of Micronesia, the Republic of the Marshall Islands, and the Republic of Palau)
- Political subdivisions of States (in consultation with States)

A Bona Fide Agent is an agency/ organization identified by the state as eligible to submit an application under the state eligibility in lieu of a state application. If you are applying as a bona fide agent of a state or local government, you must provide a letter from the state or local government as documentation of your status. Place this documentation behind the first page of your application form.

## III.2. Cost Sharing or Matching

Matching funds are not required for this program.

### III.3. Other

If you request a funding amount greater than the ceiling of the award range, your application will be considered non-responsive, and will not be entered into the review process. You will be notified that your application did not meet the submission requirements.

Special Requirements: If your application is incomplete or non-responsive to the requirements listed in this section, it will not be entered into the review process. You will be notified that your application did not meet submission requirements.

- Late applications will be considered non-responsive. See section "IV.3. Submission Dates and Times" for more information on deadlines.
- Note: Title 2 of the United States Code Section 1611 states that an organization described in Section 501(c)(4) of the Internal Revenue Code that engages in lobbying activities is not eligible to receive Federal funds constituting an award, grant, or loan.

Individuals Eligible to Become Principal Investigators: Any individual with the skills, knowledge, and resources necessary to carry out the proposed research is invited to work with their institution to develop an application for support. Individuals from underrepresented racial and ethnic groups as well as individuals with disabilities are always encouraged to apply for CDC programs.

# IV. Application and Submission Information

IV.1. Address to Request Application Package

To apply for this funding opportunity, use application form PHS 398 (OMB number 0925–0001 rev. 9/2004). Forms and instructions are available in an interactive format on the CDC Web site, at the following Internet address: http://www.cdc.gov/od/pgo/forminfo.htm.

Forms and instructions are also available in an interactive format on the National Institutes of Health (NIH) web site at the following Internet address: http://grants.nih.gov/grants/funding/phs398/phs398.html.

If you do not have access to the Internet, or if you have difficulty accessing the forms on-line, you may contact the CDC Procurement and Grants Office Technical Information Management Section (PGO–TIM) staff at: 770–488–2700. Application forms can be mailed to you.

IV.2. Content and Form of Application Submission

*Letter of Intent (LOI):* Your LOI must be written in the following format:

- Maximum number of pages: 2
- Font size: 12-point unreduced
- Double spaced
- Paper size: 8.5 by 11 inches
- Page margin size: One inch
- Printed only on one side of page
- Written in plain language, avoid jargon

Your LOI must contain the following information:

- Descriptive title of the proposed research
- Name, address, E-mail address, telephone number, and FAX number of the Principal Investigator
  - Names of other key personnel
  - Participating institutions
  - Number and title of this

# Announcement

Application: Follow the PHS 398 application instructions for content and formatting of your application. For further assistance with the PHS 398 application form, contact PGO–TIM staff at 770–488–2700, or contact GrantsInfo, Telephone (301)435–0714, E-mail: GrantsInfo@nih.gov.

Your research plan should address activities to be conducted over the entire project period.

You are required to have a Dun and Bradstreet Data Universal Numbering

System (DUNS) number to apply for a grant or cooperative agreement from the Federal government. Your DUNS number must be entered on line 11 of the face page of the PHS 398 application form. The DUNS number is a nine-digit identification number, which uniquely identifies business entities. Obtaining a DUNS number is easy and there is no charge. To obtain a DUNS number, access <a href="http://www.number.num

www.dunandbradstreet.com or call 1–866–705–5711. For more information, see the CDC Web site at: http://www.cdc.gov/od/pgo/funding/pubcommt1.htm.

This announcement uses the non-modular budgeting format.

Additional requirements that may require you to submit additional documentation with your application are listed in section "VI.2.

Administrative and National Policy Requirements."

#### IV.3. Submission Dates and Times

LOI Deadline Date: June 13, 2005. CDC requests that you send a LOI if you intend to apply for this program. Although the LOI is not required, not binding, and does not enter into the review of your subsequent application, the LOI will be used to gauge the level of interest in this program, and to allow CDC to plan the application review.

Application Deadline Date: June 27, 2005.

Explanation of Deadlines: LOIs must be received in the CDC Office of Public Health Research (OPHR) and applications must be received in the CDC Procurement and Grants Office by 4 p.m. Eastern Time on the deadline date. If you submit your LOI or application by the United States Postal Service or commercial delivery service, you must ensure that the carrier will be able to guarantee delivery by the closing date and time. If CDC receives your submission after closing due to: (1) Carrier error, when the carrier accepted the package with a guarantee for delivery by the closing date and time, or (2) significant weather delays or natural disasters, you will be given the opportunity to submit documentation of the carriers guarantee. If the documentation verifies a carrier problem, CDC will consider the submission as having been received by the deadline.

This announcement is the definitive guide on LOI and application content, submission address, and deadline. It supersedes information provided in the application instructions. If your application does not meet the deadline above, it will not be eligible for review, and will be discarded. You will be

notified that you did not meet the submission requirements.

CDC will not notify you upon receipt of your submission. If you have a question about the receipt of your LOI or application, first contact your courier. If you still have a question concerning your LOI, contact the OPHR staff at 404-371–5277. If you still have a question concerning your application, contact the PGO-TIM staff at: 770-488-2700. Before calling, please wait two to three days after the submission deadline. This will allow time for submissions to be processed and logged.

## IV.4. Intergovernmental Review of *Applications*

Your application is subject to Intergovernmental Review of Federal Programs, as governed by Executive Order (EO) 12372. This order sets up a system for state and local governmental review of proposed federal assistance applications. You should contact your state single point of contact (SPOČ) as early as possible to alert the SPOC to prospective applications, and to receive instructions on your state's process. Click on the following link to get the current SPOC list: http:// www.whitehouse.gov/omb/grants/ spoc.html.

## IV.5. Funding Restrictions

Restrictions, which must be taken into account while writing your budget, are as follows:

- · Funds relating to the conduct of research will not be released until the appropriate assurances and Institutional Review Board approvals are in place.
- Reimbursement of pre-award costs is not allowed.

If you are requesting indirect costs in your budget, you must include a copy of your indirect cost rate agreement. If your indirect cost rate is a provisional rate, the agreement should be less than 12 months of age

## IV.6. Other Submission Requirements

LOI Submission Address: Submit your LOI by express mail or delivery service to: Mary Lerchen, DrPH, Scientific Review Administrator, CDC/Office of Public Health Research, One West Court Square, Suite 7000, MS D-72. telephone: 404-371-5277. Fax: 404-371-5215. e-mail: MLerchen@cdc.gov.

Application Submission Address: Submit the original and one hard copy of your application by mail or express delivery service to:

Technical Information Management— RFA IP05-094, CDC Procurement and Grants Office, 2920 Brandywine Road, Atlanta, GA 30341.

At the time of submission, four additional copies of the application, and all appendices must be sent to: Mary Lerchen, DrPH, Scientific Review Administrator, CDC/Office of Public Health Research, One West Court Square, Suite 7000, MS D-72. telephone: 404-371-5277. Fax: 404-371–5215. e-mail: MLerchen@cdc.gov.

Applications may not be submitted electronically at this time.

#### V. Application Review Information

#### V.1. Criteria

Applicants are required to provide measures of effectiveness that will demonstrate the accomplishment of the various identified objectives of the cooperative agreement. Measures of effectiveness must relate to the performance goals stated in the "Purpose" section of this announcement. Measures must be objective and quantitative, and must measure the intended outcome. These measures of effectiveness must be submitted with the application and will be an element of evaluation.

The goals of CDC-supported research are to advance the understanding of biological systems, improve the control and prevention of disease and injury, and enhance health. In the written comments, reviewers will be asked to evaluate the application in order to judge the likelihood that the proposed research will have a substantial impact on the pursuit of these goals.

The scientific review group will address and consider each of the following criteria equally in assigning the application's overall score, weighting them as appropriate for each application. The application does not need to be strong in all categories to be judged likely to have major scientific impact and thus deserve a high priority score. For example, an investigator may propose to carry out important work that by its nature is not innovative, but is essential to move a field forward.

The review criteria are as follows: Significance: Does this study address an important problem? If the aims of the application are achieved, how will scientific knowledge be advanced? What will be the effect of these studies on the concepts or methods that drive this field? How likely are the findings to be generalizable and applicable to other settings in which mass vaccination clinics would be considered as an alternative to vaccination during routine health care visits?

Approach: Are the conceptual framework, design, methods, and analyses adequately developed, wellintegrated, and appropriate to the aims

of the project? Does the applicant acknowledge potential problem areas and consider alternative tactics?

Innovation: Does the project employ novel concepts, approaches or methods, *i.e.* model building? Are the aims original and innovative? To the extent necessary, does the project challenge existing paradigms or develop new methodologies or technologies?

*Investigator:* Is the investigator appropriately trained and well suited to carry out this work? Does the investigator have a history of conducting economic and systems research? Is the work proposed appropriate to the experience level of the principal investigator and other researchers (if any)?

Environment: Does the scientific environment in which the work will be done contribute to the probability of success? Do the proposed experiments take advantage of unique features of the scientific environment or employ useful collaborative arrangements? Is there evidence of institutional support? Are letters of support included, if appropriate?

Additional Review Criteria: In addition to the above criteria, the following items will be considered in the determination of scientific merit and

priority score:

 The applicant must demonstrate the ability to access both mass and routine clinic settings by providing letters of intent to collaborate with the applicant on behalf of clinics.

- Ability to effectively implement a large community or hospital systembased influenza vaccination program. The applicant must demonstrate the ability to conduct economic analyses in public health, including but not limited to expertise in incremental costeffectiveness analysis as evidenced by a record of publication in peer-reviewed journals
- Ability to conduct time studies and system evaluations, as evidenced by a record of publication in peer-reviewed journals. Such studies may include but not be limited to assessing health care system performance, productivity, and capacity.

Protection of Human Subjects from Research Risks: Does the application adequately address the requirements of Title 45 Part 46 for the protection of human subjects? The involvement of human subjects and protections from research risk relating to their participation in the proposed research will be assessed.

Inclusion of Women and Minorities in Research: Does the application adequately address the CDC Policy requirements regarding the inclusion of

women, ethnic, and racial groups in the proposed research? This includes: (1) The proposed plan for the inclusion of both sexes and racial and ethnic minority populations for appropriate representation; (2) The proposed justification when representation is limited or absent; (3) A statement as to whether the design of the study is adequate to measure differences when warranted; and (4) A statement as to whether the plans for recruitment and outreach for study participants include the process of establishing partnerships with community(ies) and recognition of mutual benefits.

Budget: The reasonableness of the proposed budget and the requested period of support in relation to the proposed research. The priority score should not be affected by the evaluation of the budget.

### V.2. Review and Selection Process

Applications will be reviewed for completeness by the Procurement and Grants Office (PGO) and for responsiveness by the OPHR. Incomplete applications and applications that are non-responsive to the eligibility criteria will not advance through the review process. Applicants will be notified that their application did not meet submission requirements.

Applications that are complete and responsive to the announcement will be evaluated for scientific and technical merit by an appropriate peer review group or charter study section, a Special Emphasis Panel (SEP), convened by the OPHR in accordance with the review criteria listed above. As part of the initial merit review, all applications will:

- Undergo a process in which only those applications deemed to have the highest scientific merit by the review group, generally the top half of the applications under review, will be discussed and assigned a priority score.
  - Receive a written critique.
- Receive a second programmatic level review by the Office of Science, National Immunization Program.
- Undergo a peer review by a Special Emphasis Panel. The SEP will be selected from the NIH pool of scientists or recommendations from the National Immunization Program to serve as reviewers on SEPs. Applications will be ranked for the secondary review according to scores submitted by the SEP. Only those applications deemed to have the highest scientific merit by the review group, generally the top half of the applications under review, will be discussed and assigned a priority score.

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- Scientific merit (as determined by peer review)
  - Availability of funds
  - Programmatic priorities

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Award Date: August 31, 2005.

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#### 45 CFR Part 74 and Part 92.

For more information on the Code of Federal Regulations, see the National Archives and Records Administration at the following Internet address: http://www.access.gpo.gov/nara/cfr/cfr-table-search.html.

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  - AR–7 Executive Order 12372
- AR–10 Smoke-Free Workplace Requirements
  - AR-11 Healthy People 2010
  - AR-12 Lobbying Restrictions
  - AR-15 Proof of Non-Profit Status
  - AR-22 Research Integrity
- AR–24 Health Insurance Portability and Accountability Act Requirements
- AR–25 Release and Sharing of Data

Additional information on these requirements can be found on the CDC Web site at the following Internet address: http://www.cdc.gov/od/pgo/funding/ARs.htm.

# VI.3. Reporting

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1. Interim progress report, (use form PHS 2590, OMB Number 0925–0001,

rev. 9/2004 as posted on the CDC Web site) no less than 90 days before the end of the budget period. The progress report will serve as your non-competing continuation application, and must contain the following additional elements:

- a. Progress Toward Measures of Effectiveness.
- b. Additional Information Requested by Program.
- 2. Financial status report, no more than 90 days after the end of the budget period.
- 3. Final financial and performance reports, no more than 90 days after the end of the project period.

These reports must be mailed to the Grants Management Specialist listed in the "Agency Contacts" section of this announcement.

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We encourage inquiries concerning this announcement.

For general questions, contact: Technical Information Management Section, CDC Procurement and Grants Office, 2920 Brandywine Road, Atlanta, GA 30341. Telephone: 770–488–2700.

For scientific/research issues, contact: Susan Chu, PhD, MSPH, Extramural Program Official, Centers for Disease Control and Prevention, National Immunization Program, MS E–05, 1600 Clifton Road NE, . Atlanta, GA 30333. Telephone: 404–639–8727. E-mail: \$SChu@cdc.gov.

For questions about peer review, contact: Mary Lerchen, DrPH, Scientific Review Administrator, CDC/Office of Public Health Research, One West Court Square, Suite 7000, MS D-72. Telephone: 404–371–5277. Fax: 404–371–5215. E-mail: MLerchen@cdc.gov.

For financial, grants management, or budget assistance, contact: Mattie Jackson, Grants Management Specialist, CDC Procurement and Grants Office, 2920 Brandywine Road, Atlanta, GA 30341. Telephone: 770–488–2696. E-mail: mij3@cdc.gov.

#### VIII. Other Information

This and other CDC funding opportunity announcements can be found on the CDC Web site, Internet address: http://www.cdc.gov. Click on "Funding" then "Grants and Cooperative Agreements."

Dated: May 6, 2005.

## William P. Nichols,

Director, Procurement and Grants Office, Centers for Disease Control and Prevention. [FR Doc. 05–9451 Filed 5–11–05; 8:45 am]

BILLING CODE 4163-18-P