Human Services, 2920 Brandywine Road, Atlanta, GA 30341, Telephone: 770–488–2700.

For program technical assistance, contact: Elizabeth Marum, Project Officer, HHS/CDC, Mbagathi Way, Off Mbagathi Road, Nairobi, Kenya, Telephone: 254 20 271 3008, E-mail: Emarum@cdcnairobi.mimcom.net.

For financial, grants management, or budget assistance, contact: Diane Flournoy, Grants Management Specialist, CDC Procurement and Grants Office, U.S. Department of Health and Human Services, 2920 Brandywine Road, Atlanta, GA 30341, Telephone: 770–488–2072, E-mail: Dflournoy@cdc.gov.

VIII. Other Information

Applicants can find this and other HHS/CDC funding opportunity announcements on the HHS/CDC Web site, Internet address: http://www.cdc.gov (Click on "Funding" then "Grants and Cooperative Agreements"), and on the Web site of the HHS Office of Global Health Affairs, Internet address: http://www.globalhealth.gov.

Dated: August 12, 2005.

William P. Nichols,

Director, Procurement and Grants Office, Centers for Disease Control and Prevention, U.S. Department of Health and Human Services.

[FR Doc. 05–16448 Filed 8–18–05; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[Request for Application (RFA) AA212]

Building and Strengthening the Development of the Republic of Haiti's Central HIV/AIDS Quality-Assurance/Quality-Control (QA/QC) Laboratory and the Associated National Network of QA/QC Laboratories in Haiti, as Part of the President's Emergency Plan for AIDS Relief; Notice of Intent To Fund Single Eligibility Award

A. Purpose

The Centers for Disease Control and Prevention (CDC) announces the intent to fund fiscal year (FY) 2005 funds for a cooperative agreement program to fund the President's Emergency Plan for AIDS Relief (The Emergency Plan). The plan has called for immediate action to turn the tide of HIV/AIDS in Africa and the Caribbean. The initiative hopes to prevent at least seven million new HIV infections, place two million people on

treatment, and provide care for ten million people, including orphans and vulnerable children. An essential element of preventing new cases of HIV infection is to ensure that high-risk groups have adequate access to screening, treatment, and care facilities.

Haiti's HIV prevalence rate in adults is estimated to be between 3.1 and 5.6 percent according to the Haitian Ministry of Health-Ministère de la Santé Publique et de la Population (MSPP) and the 2004 Annual Report from the Joint United Nations Programme on HIV and AIDS (UNAIDS), respectively. Access to prevention and treatment is limited to the Haitian population due to the underdeveloped public health infrastructure and lack of clinical capacity. In order to improve this capacity, this Cooperative Agreement has been developed to provide much needed funding and resources.

The Catalog of Federal Domestic Assistance number for this program is 93.067.

B. Eligible Applicant

This is a single eligibility request for application (RFA) from MSPP. No other applicants are solicited.

The MSPP is the government. They have the authority and responsibility for both regulation and QA/QC of all Laboratories within the country. They are responsible for establishing norms and standards for laboratories.

The MSPP, as the government, is the only entity that has the authority to establish and operate the entire public health system which includes departmental hospitals and clinics where ARV services are being provided. The Ministry has developed public/private partnerships to help manage some of these sites but even at those sites that are managed by the private sector they are ultimately accountable to the MSPP for services provided and quality care. The MSPP still maintains a supervisor role for these sites.

The role of regulation and standard setting at a national level is inherently governmental. In order to fulfill its role in this area the Haitian Ministry of Health needs to have the capacity to independently verify compliance through a central HIV/AIDS quality assurance/quality control laboratory. If a private or non-governmental laboratory were allowed to take on this role it would call into question the independence of the results in order to favor laboratories associated with that organization.

C. Funding

Approximately \$2,765,000 is available over a five year project period. \$553,000

is available for a 12-month budget period in FY 2005, to be awarded September 15, 2005. Funding estimates may change.

D. Where To Obtain Additional Information

For general comments or questions about this announcement, contact: Technical Information Management, CDC Procurement and Grants Office, 2920 Brandywine Road, Atlanta, GA 30341–4146. Telephone: 770–488–2700.

For program technical assistance, contact: Kathy Grooms, CDC Global AIDS Program, 1600 Clifton Road, NE, Mailstop E–04, Atlanta, GA 30333. Telephone: 404–639–8394. E-mail: Kgrooms@cdc.gov.

For financial, grants management, or budget assistance, contact: Vivian Walker, Grants Management Specialist, CDC Procurement and Grants Office, 2920 Brandywine Road, Atlanta, GA 30341. Telephone: 770–488–2724. Email: VEW4@CDC.GOV.

Dated: August 12, 2005.

William P. Nichols,

Director, Procurement and Grants Office, Centers for Disease Control and Prevention. [FR Doc. 05–16450 Filed 8–18–05; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS-10110, CMS-10136, CMS-10162, and CMS-R-0021]

Agency Information Collection Activities: Proposed Collection; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS) is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the agency's functions; (2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection

1. Type of Information Collection Request: Revision of a currently approved collection; Title of Information Collection: Manufacturer Submission of Average Sales Price (ASP) Data for Medicare Part B Drugs and Biologicals And Supporting Regulations in 42 CFR 414.804; Form No.: CMS-10110 (OMB #0938-0921); Use: In accordance with Section 1847A of the Social Security Act (the Act), Medicare Part B covered drugs and biologicals not paid on a cost or prospective payment basis are paid based on the average sales price of the drug or biological, beginning in CY 2005. The ASP data reporting requirements are specified in Section 1927 of the Act. The reported ASP data are used to establish the Medicare payment amounts. Specifically, CMS will utilize the ASP data to determine the drug payment amounts for CY 2005 and beyond. The interim final rule "Medicare Program; Manufacturer Submission of Manufacturer's Average Sales Price (ASP) Data for Medicare Part B Drugs and Biologics" (CMS-1380-IFC), published in the Federal Register on April 6, 2004 (66 FR 17936), set forth the ASP reporting format, Addendum A. The rule stated that, as we gain more experience with the ASP methodology, we may seek to modify the reporting requirements (data elements and format for submission) in the future. Based on our experience during the initial six reporting periods, we have found it necessary for carrying out section 1847A of the Act to expand the ASP data collected from manufacturers. We are proposing that, upon approval of this requested revision, in addition to the data elements in the original Addendum A (manufacturer name, National Drug Code (NDC), manufacturer's ASP, and number of units), the following data elements must be submitted quarterly by manufacturers: name of drug or biological, strength of the product, volume per item, number of items per NDC, wholesale acquisition costs (applies to NDCs assigned to single source drug and biological billing codes and NDCs during the initial period under section 1847 A(c)(4) of the Act), and expiration date of the last lot manufactured. We are proposing that manufacturers would no longer report ASP data for an NDC beginning the reporting period after the expiration date of the last lot manufactured. For NDCs first marketed or sold on or after October 1, 2005, we are also proposing to collect the date the NDC was first marketed and the date of first sale. We

propose that manufacturers would be required to submit these dates to us once with the first data submission for new NDCs. Frequency: Recordkeeping and reporting—quarterly; Affected Public: Business or other for-profit; Number of Respondents: 120; Total Annual Responses: 480; Total Annual Hours: 17.760.

2. Type of Information Collection Request: Extension of a currently approved collection; Title of Information Collection: Medicare Care Management Performance (MCMP) Demonstration—Standardized Ambulatory Care Quality Collection Initiative: *Use:* The MCMP Demonstration was authorized by Section 649 of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (MMA). This project requires the Secretary to establish a pay-for-performance 3-year pilot with physicians to promote the adoption and use of health information technology to improve the quality of patient care for chronically ill Medicare patients. This demonstration represents the first pay for performance project fostering the adoption of health information technology in small physician group practices and will enable a test of the concept to improve the quality and efficiency of care in Feefor-Service (FFS) Medicare. Form Number: CMS-10136 (OMB #0938-0941); Frequency: Annually; Affected Public: Business or other for-profit and not-for-profit institutions; Number of Respondents: 800; Total Annual Responses: 800; Total Annual Hours: 19,200.

3. Type of Information Collection Request: New collection; Title of Information Collection: Medicare Care Improvement Survey; Use: The purpose of this beneficiary survey is to obtain information about beneficiary behavioral change, physical functioning and satisfaction with the Chronic Care Improvement (CCI) programs, data required by legislation to form decisions related to expansion of the pilot programs. The chronic care improvement programs are to be designed to incorporate relevant features from private sector programs but also be sufficiently flexible to adapt to the unique needs of their Medicare populations. This survey is required to support the legislative mandate to evaluate the Chronic Care Improvement Programs. Beneficiary participation in the CCI-I program will be voluntary and will not change the scope, duration or amount of Medicare FFS benefits currently received by FFS Medicare participants. Form Number: CMS-10162 (OMB #0938-NEW); Frequency:

Reporting—on occasion; Affected Public: Individuals or households; Number of Respondents: 9,449; Total Annual Responses: 9,449; Total Annual Hours: 2,636.

4. Type of Information Collection Request: Extension of a currently approved collection; Title of Information Collection: Withholding Medicare Payments to Recover Medicaid Overpayments and Supporting Regulations in 42 CFR 447.31; Use: Overpayments may occur in either the Medicare and Medicaid program, at times resulting in a situation where an institution or person that provides services owes a repayment to one program while still receiving reimbursement from the other. Certain Medicaid providers which are subject to offsets for the collection of Medicaid overpayments may terminate or substantially reduce their participation in Medicaid, leaving the State Medicaid Agency unable to recover the amounts due. These information collection requirements give CMS the authority to recover Medicaid overpayments by offsetting payments due to a provider under the program. Form Number: CMS-R-0021 (OMB #0938-0287); Frequency: Reporting—on occasion; Affected Public: State, local or tribal government; Number of Respondents: 54; Total Annual Responses: 27; Total Annual Hours: 81.

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS' Web site address at http://www.cms.hhs.gov/regulations/pra/, or e-mail your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov, or call the Reports Clearance Office on (410) 786–1326.

Written comments and recommendations for the proposed information collections must be mailed within 60 days of this notice to: CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: William N. Parham, III, Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

Dated: August 12, 2005.

Michelle Shortt,

Director, Regulations Development Group, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 05–16472 Filed 8–18–05; 8:45 am]

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