On May 16, 2014 OMB approved the collection of standardized information from ten project sites over the three-year project period and one retrospective data collection during the first year of the three-year project period. The retrospective data collection will provide information about clients’ baseline characteristics prior to participation in the model program which is needed to compare outcomes before and after program implementation. On August 17, 2015 OMB approved the conduct of key informant interviews with program clinic and pharmacy staff in order to evaluate the program processes, administration of a staff communication questionnaire, and OMB approved the collection of time and cost data to be used to estimate the cost of the model program.

CDC seeks approval to administer a staff communication questionnaire for medical providers in order to determine how and if the model program improves patient outcomes through improved communication and collaboration between patients’ clinical providers and pharmacists. The staff communication questionnaire for medical providers will be administered twice to program clinic staff. The staff communication questionnaire for medical providers is different from the previously improved staff communication questionnaire; the staff communication questionnaire for medical providers will be administered to program clinic staff whereas the staff communication questionnaire will be administered to program pharmacy staff. Pharmacy, laboratory, and medical data will be collected through abstraction of all participant clients’ pharmacy and medical records. Pharmacy, laboratory and medical data are needed to monitor retention in care, adherence to therapy, viral load suppression and other health outcomes. Program specific data, such as the number of MTM elements completed per project site and time spent on program activities, will be collected by program. Qualitative data will be gathered from program staff through in-person or telephone interviews and through a questionnaire to program pharmacy staff and a separate questionnaire to program clinic staff.

The data collection will allow CDC to conduct continuous program performance monitoring which includes identification of barriers to program implementation, solutions to those barriers, and documentation of client health outcomes. Performance monitoring will allow the model program to be adjusted, as needed, in order to develop a final implementation model that is self-sustaining and which can be used to establish similar collaborations in a variety of clinical settings. Collection of cost data will allow for the cost of the program to be estimated.

There is no cost to participants other than their time. The total estimated annualized burden hours are 6,043.

<table>
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<tr>
<th>Type of respondent</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
<th>Total burden (in hours)</th>
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<td>30/60</td>
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Leroy A. Richardson,
Chief, Information Collection Review Office, Office of Scientific Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2015–29274 Filed 11–16–15; 8:45 am]
Prevention and the Agency for Toxic Substances and Disease Registry.

Elaine L. Baker,  
Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2015–29257 Filed 11–16–15; 8:45 am]  
BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Disease Control and Prevention

Statement of Organization, Functions, and Delegations of Authority

Part C (Centers for Disease Control and Prevention) of the Statement of Organization, Functions, and Delegations of Authority of the Department of Health and Human Services (45 FR 67772–76; dated October 14, 1980, and corrected at 45 FR 69296, October 20, 1980, as amended most recently at 80 FR 58479–58485, dated September 29, 2015) is amended to reflect the reorganization of the National Center for Immunization and Respiratory Diseases, the National Center for Infectious Diseases, Centers for Disease Control and Prevention.

Section C–B, Organization and Functions, is hereby amended as follows:

Delete in its entirety the title and function statements for the Influenza Coordination Unit (CVA4).

Delete in its entirety the title and function statements for the National Center for Immunization and Respiratory Diseases (CVG) and insert the following:

National Center for Immunization and Respiratory Diseases (CVG). The National Center for Immunization and Respiratory Diseases (NCIRD) prevents disease, disability, and death through immunization and by control of respiratory and related diseases. In carrying out its mission, NCIRD: (1) Provides leadership, expertise, and service in laboratory and epidemiological sciences, and in immunization program delivery; (2) conducts applied research on disease prevention and control; (3) translates research findings into public health policies and practices; (4) provides diagnostic and reference laboratory services to relevant partners; (5) conducts surveillance and research to determine disease distribution, determinants, and burden nationally and internationally; (6) responds to disease outbreaks domestically and abroad; (7) ensures that public health decisions are made objectively and based upon the highest quality of scientific data; (8) provides technical expertise, education, and training to domestic and international partners; (9) provides leadership to internal and external partners for establishing and maintaining immunization, and other prevention and control programs; (10) develops, implements, and evaluates domestic and international public health policies; (11) communicates information to increase awareness, knowledge, and understanding of public health issues domestically and internationally, and to promote effective immunization programs; (12) aligns the national center focus with the overall strategic goals of CDC; (13) synchronizes all aspects of CDC’s pandemic influenza preparedness and response from strategy through implementation and evaluation; and (14) implements, coordinates, and evaluates programs across NCIRD, Office of Infectious Diseases (OID), and CDC to optimize public health impact.

Delete in its entirety the title and function statements for the Office of Laboratory Science (CVG14).

After the Office of Science and Integrated Programs (CVG17) insert the following:

Influenza Coordination Unit (CVG18). The mission of the Influenza Coordination Unit (ICU) is to synchronize all aspects of CDC’s pandemic influenza preparedness and response from strategy through implementation and evaluation. In carrying out its mission, the ICU: (1) Serves as the principal advisor to the CDC Director on pandemic influenza preparedness and response activities, assisting the Director in formulating and communicating strategic pandemic initiatives and policies; (2) provides strategic leadership for CDC in the areas of pandemic preparedness and response, including setting priorities and promoting science, policies, and programs related to pandemic influenza; (3) strategically manages a budget and allocates funds across the agency to ensure appropriate resources for high priority areas; and (4) conducts ongoing evaluation and adjustment of pandemic preparedness and response activities, in coordination with the National Response Framework and other emergency preparedness guidance, to ensure optimal public health effectiveness and efficient use of human and fiscal resources by developing and leading an exercise program for the Agency, in collaboration with HHS and other partners.

James Seligman,  
Acting Chief Operating Officer, Centers for Disease Control and Prevention.

[FR Doc. 2015–29282 Filed 11–16–15; 8:45 am]

BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Disease Control and Prevention

Board of Scientific Counselors, Office of Infectious Diseases (BSC, OID)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC), announces the following meeting of the aforementioned committee:

Time and Date:
8:30 a.m.–5:00 p.m., EST, December 9, 2015
8:00 a.m.–12:00 p.m., EST, December 10, 2015

Place: CDC, Global Communications Center, 1600 Clifton Road, NE., Building 19, Auditorium B3, Atlanta, Georgia 30333.

Status: The meeting is open to the public, limited only by the space available.

Purpose: The BSC, OID, provides advice and guidance to the Secretary, Department of Health and Human Services; the Director, CDC; the Director, OID; and the Directors of the National Center for Immunization and Respiratory Diseases, the National Center for Emerging and Zoonotic Infectious Diseases, and the National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention, CDC, in the following areas: Strategies, goals, and priorities for programs; research within the national centers; and overall strategic direction and focus of OID and the national centers.

Matters for Discussion: The meeting will include reports from the Board’s Food Safety Modernization Act Surveillance Working Group and Infectious Disease Laboratory Working Group; brief updates on selected activities of CDC’s infectious disease national centers; and updates and focused discussions on prevention of Legionella disease and efforts to better understand and address environmental factors contributing to infectious disease outbreaks.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information:
Robin Moseley, M.A.T., Designated Federal Official, OID, CDC, 1600 Clifton Road NE., Mailstop D10, Atlanta, Georgia 30333, Telephone: (404) 639–4461.

The Director, Management Analysis and Services Office has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and