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:

Times and Dates:
11 a.m.–5:30 p.m., September 22, 2011.

Place: NCHS Headquarters, 3311 Toledo Road, Hyattsville, Maryland 20782.

Status: This meeting is open to the public; however, visitors must be processed in accordance with established federal policies and procedures. For foreign nationals or non-US citizens, pre-approval is required (please contact Althelia Harris, (301) 458–4261, or Virginia Cain, vcain@cdc.gov at least 10 days in advance for requirements). All visitors are required to present a valid form of picture identification issued by a state, federal or international government. As required by the Federal Property Management Regulations, Title 41, Code of Federal Regulation, Subpart 101–20.301, all persons entering in or on Federal controlled property and their packages, briefcases, and other containers in their immediate possession are subject to being x-rayed and inspected. Federal law prohibits the knowing possession or the causing to be present of firearms, explosives and other dangerous weapons and illegal substances. The meeting room accommodates approximately 100 people.

Purpose: This committee is charged with providing advice and making recommendations to the Secretary, Department of Health and Human Services; the Director, CDC; and the Director, NCHS, regarding the scientific and technical program goals and objectives, strategies, and priorities of NCHS.

Matters To Be Discussed: The agenda will include welcome remarks by the Director, NCHS; update on the Health Indicators Warehouse; update on program reviews; discussion of the NHANES program, plans for the NHS for 2012 and beyond and an open session for comments from the public.

Requests to make oral presentations should be submitted in writing to the contact person listed below. All requests must contain the name, address, telephone number, and organizational affiliation of the presenter.

Written comments should not exceed five single-spaced typed pages in length and must be received by September 12, 2011.

The agenda items are subject to change as priorities dictate.

Contact Person for More Information: Virginia S. Cain, PhD, Director of Extramural Research, NCHS/CDC, 3311 Toledo Road, Room 7208, Hyattsville, Maryland 20782, Telephone (301) 458–4500, Fax (301) 458–4020.

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 Prevention, and the Agency for Toxic Substances and Disease Registry.

Date: August 17, 2011.

Elizabeth Millington, Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2011–21742 Filed 8–24–11; 8:45 am]

BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Board of Scientific Counselors, National Center for Health Statistics, (BSC, NCHS)

We are committed to achieving the three-part aim of better health, better health care, and reduced expenditures through continuous improvement for Medicare, Medicaid and Children’s Health Insurance Program (CHIP) beneficiaries. Beneficiaries can experience improved health outcomes and patient experience when health care providers work in a coordinated and patient-centered manner. To this end, we are interested in partnering with providers who are working to redesign patient care to deliver these aims. Episode payment approaches that reward providers who take accountability for the three-part aim at the level of individual patient care for an episode are potential mechanisms for developing these partnerships.

In order to provide a flexible and far-reaching approach towards episode-based care improvement, we are seeking proposals from health care providers who wish to align incentives between hospitals, physicians, and nonphysician practitioners in order to better coordinate care throughout an episode of care. This Bundled Payment for Care Improvement initiative request for applications (RFA) will test episode-based payment for acute care and associated post-acute care, using both retrospective and prospective bundled payment methods. The RFA requests applications to test models centered around acute care; these models will inform the design of future models, including care improvement for chronic conditions. For more details, see the RFA which is available on the Innovation Center Web site at http://www.innovations.cms.gov/areas-of-focus/patient-care-models/bundled-payments-for-care-improvement.html. Applications are due by September 22, 2011 for Models 1 and November 4, 2011 for Models 2 through 4 as described on the CMS Innovation Center Web site http://www.innovations.cms.gov/areas-of-focus/patient-care-models/bundled-payments-for-care-improvement.html. For applicants wishing to receive historical Medicare claims data in preparation for Models 2 through 4, a separate research request packet and data use agreement must be filed in conjunction with the Letter of Intent.

Application Submission Deadlines: Interested organizations must submit a nonbinding letter of intent by September 22, 2011 for Model 1 and November 4, 2011 for Models 2 through 4 as described on the CMS Innovation Center Web site http://www.innovations.cms.gov/areas-of-focus/patient-care-models/bundled-payments-for-care-improvement.html. For applicants wishing to receive historical Medicare claims data in preparation for Models 2 through 4, a separate research request packet and data use agreement must be filed in conjunction with the Letter of Intent.

Application Submission Deadlines: Applications must be received on or before October 21, 2011 for Model 1 and March 15, 2012 for Models 2 through 4.

II. Provisions of the Notice

Consistent with its authority under section 1115A of the Social Security Act (of the Act), as added by section 3021 of the Affordable Care Act, to test
innovative payment and service delivery models that reduce spending under Medicare, Medicaid, or CHIP, while preserving or enhancing the quality of care, the Innovation Center aims to achieve the following goals through implementation of the Bundled Payments for Care Improvement initiative:

- Improve care coordination, patient experience, and accountability in a patient-centered manner.
- Support and encourage providers who are interested in continuously reengineering care to deliver better care, better health, at lower costs through continuous improvement.
- Create a virtuous cycle that leads to continually decreasing the cost of an acute or chronic episode of care while fostering quality improvement.
- Develop and test payment models that create extended accountability for better care, better health at lower costs for acute and chronic medical care.
- Shorten the cycle time for adoption of evidence-based care.
- Create environments that stimulate rapid development of new evidence-based knowledge.

The models to be tested based on applications to the RFA are as follows:

- **Model 1:** Retrospective payment models around the acute inpatient hospital stay only.
- **Model 2:** Retrospective bundled payment models for hospitals, physicians, and post-acute providers for an episode of care consisting of an inpatient hospital stay followed by post-acute care.
- **Model 3:** Retrospective bundled payment models for post-acute care where the episode does not include the acute inpatient hospital stay.
- **Model 4:** Prospectively administered bundled payment models for the acute inpatient hospital stay only, such as prospective bundled payment for hospitals and physicians for an inpatient hospital stay.

Proposals should demonstrate care improvement processes and enhancements such as reengineered care pathways using evidence-based medicine, standardized care using checklists, and care coordination. All models must encourage close partnerships among all of the providers caring for patients through the episode. Applicants must demonstrate robust quality monitoring and protocols to ensure beneficiary quality protection. Under all models, applicants must provide Medicare with a discount on Medicare fee-for-service expenditures.

Bundled Payments for Care Improvement agreements will include a performance period of 3 years, with the possibility of extending an additional 2 years, beginning with the respective program date. The program start date may be as early as the first quarter of CY 2012 for awardees in Model 1.

### III. Collection of Information Requirements

Section 1115A(d) of the Act waives the requirements of the Paperwork Reduction Act of 1995 for the Innovation Center for purposes of testing new payment and service delivery models.

**Authority:** 44 U.S.C. 3101.

**Dated:** August 17, 2011.

Donald M. Berwick,
Administrator, Centers for Medicare & Medicaid Services.

[FR Doc. 2011–21707 Filed 8–23–11; 11:15 am]

**BILLING CODE 4120–01–P**

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Substance Abuse and Mental Health Services Administration

#### Agency Information Collection Activities: Submission for OMB Review; Comment Request

Periodically, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish a summary of information collection requests under OMB review, in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these documents, call the SAMHSA Reports Clearance Officer on (240) 276–1243.

**Project:** National Child Traumatic Stress Initiative (NCTSI) Evaluation—(OMB No. 0930–0276)—Revision

The Substance Abuse and Mental Health Services Administration’s (SAMHSA), Center for Mental Health Services (CMHS), will conduct the National Child Traumatic Stress Initiative (NCTSI) Evaluation. This evaluation serves multiple practical purposes: (1) To collect and analyze descriptive, outcome, and service experience information about the children and families served by the NCTSI centers; (2) to assess the NCTSI’s impact on access to high-quality, trauma-informed care; (3) to evaluate NCTSI centers’ training and consultation activity designed to promote evidence-based, trauma-informed services and the impact of such activity on child-serving systems; and (4) to assess the sustainability of the grant-funded activities to improve access to and quality of care for trauma-exposed children and their families beyond the grant period.

Data will be collected from caregivers and youth served by NCTSI centers, NCTSI and non-NCTSI administrators, NCTSI trainers, service providers trained by NCTSI centers and other training participants, administrators of mental health and non-mental health professionals from state and national child-serving organizations, and administrators of affiliate centers. Data collection will take place in all Community Treatment and Services Programs (CTS) and Treatment and Service Adaptation Centers (TSA) active during the three-year approval period. Currently, there are 45 CTS centers and 17 TSA centers active (i.e., 62 active centers). After the first year, in September 2011, the 15 grantees funded in 2007 will reach the end of their data collection. At that point, additional centers may be funded or funded again. Because of this variability, the estimate of 62 centers is used to calculate burden.

The NCTSI Evaluation is composed of four distinct study components, each of which involves data collection, which are described below.

#### Descriptive and Clinical Outcomes

In order to describe the children served, their trauma histories and their clinical and functional outcomes, nine instruments will be used to collect data from children and adolescents who are receiving services in the NCTSI, and from caregivers of all children who are receiving NCTSI services. Data will be collected when the child/youth enters services and during subsequent follow-up sessions at three-month intervals over the course of one year. This study relies upon the use of data already being collected as a part of the Core Data Set, and includes the following instruments:

- The Core Clinical Characteristics Form, which collects demographic, psychosocial and clinical information about the child being served including information about the child’s domestic environment and insurance status, indicators of the severity of the child’s problems, behaviors and symptoms, and use of non-Network services;
- The Trauma Information/Detail Form, which collects information on the history of trauma(s) experienced by the child served by the NCTSI center including the type of trauma experienced, the age at which the trauma was experienced, type of exposure, whether or not the trauma is chronic, and the setting and...