room accommodates approximately 240 people.

Security Considerations: Due to increased security requirements CMS has instituted stringent procedures for entrance into the building by nongovernment employees. Attendees will need to present valid government-issued picture identification, and sign-in at the security desk upon entering the building. Attendees who wish to attend the ICD-9-CM C&M meeting on March 5, 2012, must submit their name and organization by February 27, 2012, for inclusion on the visitor list. This visitor list will be maintained at the front desk of the CMS building and used by the guards to admit visitors to the meeting.

Participants who attended previous ICD-9-CM C&M meetings will no longer be automatically added to the visitor list. You must request inclusion of your name prior to each meeting you attend.

Please register to attend the meeting on-line at: http://www.cms.hhs.gov/apps/events/.

Please contact Mady Hue (410–786–4510 or *Marilu.hue@cms.hhs.gov*), for questions about the registration process.

Purpose: The ICD-9-CM Coordination and Maintenance Committee is a public forum for the presentation of proposed modifications to the International Classification of Diseases, Ninth-Revision, Clinical Modification.

Matters to be Discussed: Tentative agenda items include:

March 5, 2012

ICD-9-CM Procedure Topics: Administration of Fidaxomicin Placement of Modeling Catheter in Endovascular Graft Procedure Injection or Infusion of Glucarpidase ICD-10 Updates:

ICD–10 MS–DRG Update ICD–10 HAC Translation List Impact of ICD–10 MS–DRGs Implementation

ICD-10-CM Diagnosis Topics:

Atypical femoral fracture Choking "game" Cognitive sequelae of cerebrovascular disease Family history of SIDS

Addenda

Agenda items are subject to change as priorities dictate.

Note: CMS and NCHS will no longer provide paper copies of handouts for the meeting. Electronic copies of all meeting materials will be posted on the CMS and NCHS Web sites prior to the meeting at http://www.cms.hhs.gov/ICD9ProviderDiagnosticCodes/03_meetings.asp#TopOfPage and http://www.cdc.gov/nchs/icd/icd9cm_maintenance.htm.

Contact Persons for Additional Information: Donna Pickett, Medical Systems Administrator, Classifications and Public Health Data Standards Staff, NCHS, 3311 Toledo Road, Room 2337, Hyattsville, Maryland 20782, email dfp4@cdc.gov, telephone 301–458–4434 (diagnosis); Mady Hue, Health Insurance Specialist, Division of Acute Care, CMS, 7500 Security Boulevard, Baltimore, Maryland 21244, email marilu.hue@cms.hhs.gov, telephone 410–786–4510 (procedures).

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.

Dated: February 8, 2012.

Elaine L. Baker,

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

[FR Doc. 2012–3484 Filed 2–14–12; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee on Immunization Practices (ACIP)

Correction

This notice was published in the **Federal Register** on February 1, 2012, Volume 77, Number 21, Page 5026. The matters to be discussed and times should read as follows:

Matters To Be Discussed: The agenda will include discussions on: meningococcal vaccine, hepatitis B vaccine, tetanus, diphtheria, and acellular pertussis (Tdap) vaccine, influenza, vaccine supply, 13-valent pneumococcal conjugate vaccine, and measles-mumps-rubella (MMR) vaccine. Recommendation vote is scheduled for Tdap vaccine. Time will be available for public comment.

Agenda items are subject to change as priorities dictate.

Times and Dates:

8 a.m.–5 p.m., February 22, 2012. 8 a.m.–12:30 p.m., February 23, 2012.

The Meeting is Web cast live via the World Wide Web; for instructions and more information on ACIP please visit the ACIP Web site: http://www.cdc.gov/vaccines/recs/acip/.

Contact Person for More Information: Stephanie B. Thomas, National Center for Immunization and Respiratory Diseases, CDC, 1600 Clifton Road NE., MS–A27, Atlanta, Georgia 30333, telephone: (404)639–8836; Email *ACIP@CDC.GOV*.

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.

Dated: February 8, 2012.

Elaine L. Baker,

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

[FR Doc. 2012-3481 Filed 2-14-12; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Tracking of Participants in the Head Start Impact Study.

OMB No.: 0970–0229.

Description: The Administration for Children and Families (ACF) within the Department of Health and Human Services (HHS) will collect follow-up information from children and families in the Head Start Impact Study. In anticipation of conducting a future follow-up for the study, ACF will collect information necessary to identify respondents' current location and follow-up with respondents in the future.

The Head Start Impact Study is a longitudinal study involving 4,667 first time enrolled three- and four-year-old preschool children across 84 nationally representative grantee/delegate agencies. Participants have been randomly assigned to either a Head Start group or a control group. Data collection for the study began in fall of 2002 and has been extended through late spring 2008 to include the participants' 3rd grade year. Tracking of the participants has continued every spring beginning in 2009 and ending in 2011.

ACF will continue to examine outcomes for the sample through the spring of the participant's 12th grade year. To maintain adequate sample size, telephone interviews will be conducted in order to update the respondent's location and contact information. This information will be collected from

parents or guardians in the spring of 2012, 2013, 2014 2015, and 2016. This request package covers three years of

information collection, from 2012 to 2014.

Respondents: The original sample of 4,667 treatment and control group

members in the Head Start Impact Study, minus 432 families that have refused to participate in the study.

ANNUAL BURDEN ESTIMATES

| Instrument | Annual number of respondents | Number of responses per respondent | Average burden hours per response | Total annual burden hours |
|---------------------------|------------------------------------|------------------------------------|---|---------------------------|
| Parent Tracking Interview | 4235 | 1 | 1/3 | 1412 |

Estimated Total Annual Burden Hours: 1412.

Additional Information: Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 370 L'Enfant Promenade SW., Washington, DC 20447, Attn: OPRE Reports Clearance Officer. All requests should be identified by the title of the information collection. Email address: OPREinfocollection@acf.hhs.gov.

OMB Comment: OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the Federal Register. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following: Office of Management and Budget, Paperwork Reduction Project, Fax: 202-395-6974, Attn: Desk Officer for the Administration for Children and Families.

Dated: February 9, 2012.

Steven M. Hanmer,

OPRE Reports Clearance Officer. [FR Doc. 2012–3476 Filed 2–14–12; 8:45 am]

BILLING CODE 4184-22-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2012-N-0115]

Agency Information Collection
Activities; Proposed Collection;
Comment Request: Guidance for
Industry and Food and Drug
Administration Staff; Class II Special
Controls Guidance Document:
Automated Blood Cell Separator
Device Operating by Centrifugal or
Filtration Separation Principle

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal Agencies are required to publish notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice solicits comments on the collection of information concerning class II special controls for an automated blood cell separator device operating by centrifugal or filtration separation principle.

DATES: Submit either electronic or written comments on the collection of information by April 16, 2012.

ADDRESSES: Submit electronic comments on the collection of information to http://www.regulations.gov. Submit written comments on the collection of information to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Ila S. Mizrachi, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., PI50–400B, Rockville, MD 20850, 301–796–7726, Ila.Mizrachi@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: Under the PRA (44 U.S.C. 3501–3520), Federal Agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. "Collection of information" is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes Agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44

U.S.C. 3506(c)(2)(A)) requires Federal Agencies to provide a 60-day notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, FDA is publishing notice of the proposed collection of information set forth in this document.

With respect to the following collection of information, FDA invites comments on these topics: (1) Whether the proposed collection of information is necessary for the proper performance of FDA's functions, including whether the information will have practical utility; (2) the accuracy of FDA's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques, when appropriate, and other forms of information technology.

Guidance for Industry and Food and Drug Administration Staff; Class II Special Controls Guidance Document: Automated Blood Cell Separator Device Operating by Centrifugal or Filtration Separation Principle (OMB Control Number 0910–0594)—Extension

Under the Safe Medical Devices Act of 1990 (Pub. L. 101–629), FDA may establish special controls, including performance standards, postmarket surveillance, patient registries, guidelines, and other appropriate actions it believes necessary to provide reasonable assurance of the safety and effectiveness of the device.

The special control guidance serves to support the reclassification from class III to class II of the automated blood cell separator device operating on a centrifugal separation principle intended for the routine collection of blood and blood components as well as