FOR FURTHER INFORMATION CONTACT: Bryan Saddler, Chief Counsel, FHFA–OIG, at (202) 408–2577, or Bryan.Saddler@fhfa.gov.

SUPPLEMENTARY INFORMATION: The Federal Housing Finance Regulatory Reform Act of 2008 (Reform Act), which was passed as Division A of the Housing and Economic Recovery Act of 2008 (HERA), Public Law 110–289, 122 Stat. 2654, 2013, abolished both the Federal Housing Finance Board (FHFB), an independent agency that oversaw the Federal Home Loan Banks (Banks), and the Office of Federal Housing Enterprise Oversight (OFHEO), an office within the Department of Housing and Urban Development (HUD) that oversaw the “safety and soundness” of Fannie Mae and Freddie Mac. See 12 U.S.C. 1422a, 4502(6), 4511, 4512, 4513, 4541, 4563 (2006); 42 U.S.C. No. 110–142, at 95. The Reform Act established in place of the FHFB and OFHEO a new entity, the Federal Housing Finance Agency (FHFA), which now regulates and supervises Fannie Mae, Freddie Mac, and the 12 Banks. See Reform Act sections 1002, 1101, 1102, 1301, 1311, 12 U.S.C.A. 4511, 4512, 4513 (2009).

Section 1105 of HERA also amended the Federal Housing Enterprises Financial Safety and Soundness Act of 1992 and the Inspector General Act of 1978 (the IG Act), by specifying that there shall be established an Inspector General within FHFA. See 12 U.S.C. 4517(d). FHFA–OIG is responsible for, among other things, conducting audits, investigations, and inspections of FHFA’s programs and operations, and recommending polices that promote economy and efficiency in the administration of, and prevent and detect fraud and abuse in, those programs and operations. Section 6(a)(4) of the IG Act authorizes the Inspector General to require by subpoena the production of all information, documents, reports, answers, records, accounts, papers, and other data and documentary evidence deemed necessary in the performance of the Inspector General’s function. This notice delegates the Inspector General’s subpoena issuance authority to the FHFA–OIG Principal Deputy Inspector General, the FHFA–OIG Deputy Inspector General for Audit, the FHFA–OIG Deputy Inspector General for Investigations & Evaluations, and the FHFA–OIG Chief Counsel.

Section 552a(b)(7) of Title 5, United States Code, authorizes the Inspector General to request information protected by the Privacy Act for a civil or criminal law enforcement activity. This notice delegates this authority to request records protected by the Privacy Act for a civil or criminal law enforcement activity from the Inspector General to the FHFA–OIG Principal Deputy Inspector General, the FHFA–OIG Deputy Inspector General for Audit, the FHFA–OIG Deputy Inspector General for Investigations & Evaluations, and the FHFA–OIG Chief Counsel.

The Inspector General has not limited his authority to issue subpoenas or to request information under 5 U.S.C. 552a by this delegation. Also, this delegation expressly prohibits further delegation or redelegation.

Accordingly, the Inspector General delegates the following authorities: Section A. Authority Delegated: The Inspector General delegates to the FHFA–OIG Principal Deputy Inspector General, the FHFA–OIG Deputy Inspector General for Audit, the FHFA–OIG Deputy Inspector General for Investigations & Evaluations, and the FHFA–OIG Chief Counsel, the authority to require by subpoena the production of all information, documents, reports, answers, records, accounts, papers, and other data and documentary evidence necessary in the performance of the functions assigned by HERA and the Inspector General Act.

Additionally, the Inspector General delegates to the FHFA–OIG Principal Deputy Inspector General, the FHFA–OIG Deputy Inspector General for Audit, the FHFA–OIG Deputy Inspector General for Investigations & Evaluations, and the FHFA–OIG Chief Counsel, the authority to request information under 5 U.S.C. 552a(b)(7).

Section B. No Further Delegation or Redeployment: The authority delegated in Section A above may not be further delegated or redelegated.


Steve A. Linick,
Inspector General.

[FR Doc. 2010–32348 Filed 12–23–10; 8:45 am]
BILLING CODE 8070–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Board on Radiation and Worker Health (ABRW or Advisory Board), National Institute for Occupational Safety and Health (NIOSH)

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: 11 a.m.–2 p.m., January 12, 2011
Place: Audio Conference Call via FTS Conferencing. The USA toll-free, dial-in number is 1–866–659–0537 and the pass code is 9933701.

Status: Open to the public, but without a public comment period.

Background: The Advisory Board was established under the Energy Employees Occupational Illness Compensation Program Act of 2000 to advise the President on a variety of policy and technical functions required to implement and effectively manage the new compensation program. Key functions of the Advisory Board include providing advice on the development of probability of causation guidelines, which have been promulgated by the Department of Health and Human Services (HHS) as a final rule; advice on methods of dose reconstruction, which have also been promulgated by HHS as a final rule; advice on the scientific validity and quality of dose estimation and reconstruction efforts being performed for purposes of the compensation program; and advice on petitions to add classes of workers to the Special Exposure Cohort (SEC).

In December 2000, the President delegated responsibility for funding, staffing, and operating the Advisory Board to HHS, which subsequently delegated this authority to the CDC. NIOSH implements this responsibility for CDC. The charter was issued on August 3, 2001, renewed at appropriate intervals, most recently, August 3, 2009, and will expire on August 3, 2011.

Purpose: This Advisory Board is charged with (a) Providing advice to the Secretary, HHS, on the development of guidelines under Executive Order 13179; (b) providing advice to the Secretary, HHS, on the scientific validity and quality of dose reconstruction efforts performed for this program; and (c) upon request by the Secretary, HHS, advising the Secretary on whether there is a class of employees at any Department of Energy facility who were exposed to radiation but for whom it is not feasible to estimate their radiation dose, and on whether there is reasonable likelihood that such radiation doses may have endangered the health of members of this class.

Matters to be Discussed: The agenda for the conference call includes: NIOSH 10-Year Review of its Division of Compensation Analysis and Support (DCAS) Program; Linde Ceramics Plant SEC Petition #107 (1954–2006); DCAS Science Issues Update; Subcommittee and Work Group Updates; DCAS SEC Petition Evaluations Update for the February 2011 Advisory Board Meeting; and Board Correspondence.

The agenda is subject to change as priorities dictate.

Because there is not a public comment period, written comments may be submitted. Any written comments received will be included in the official record of the meeting.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services
[CMS–6041–NC]

Medicare Program: Solicitation of Comments Regarding Development of a Recovery Audit Contractor Program for the Medicare Part C and D Programs

AGENCY: Centers for Medicare & Medicaid Services (CMS), HHS.

ACTION: Request for information.

SUMMARY: This notice presents an approach and requests comments on the provision of the Patient Protection and Affordable Care Act (Pub. L. 111–148), as amended by the Health Care and Education Reconciliation Act of 2010 (Pub. L. 111–152), (collectively known as The Affordable Care Act (ACA)) that requires the expansion of the Recovery Audit Contractor (RAC) Program to the Medicare Part C and D programs.

DATES: Comment Date: To be assured consideration, comments must be received at one of the addresses provided below, no later than 5 p.m. on February 25, 2011.

ADDRESSES: In commenting, please refer to file code CMS–6041–NC. Because of staff and resource limitations, we cannot accept comments by facsimile (FAX) transmission.

You may submit comments in one of four ways (please choose only one of the ways listed):

1. Electronically. You may submit electronic comments on this regulation to http://www.regulations.gov. Follow the “Submit a comment” instructions.

By regular mail. You may mail written comments to the following address only: Centers for Medicare & Medicaid Services, Department of Health and Human Services, Attention: CMS–6041–NC, P.O. Box 8013, Baltimore, MD 21244–8013.

Please allow sufficient time for mailed comments to be received before the close of the comment period.

3. By express or overnight mail. You may send written comments to the following address only: Centers for Medicare & Medicaid Services, Department of Health and Human Services, Attention: CMS–6041–NC, Mail Stop C4–26–05, 7500 Security Boulevard, Baltimore, MD 21244–1850.

4. By hand or courier. Alternatively, you may deliver (by hand or courier) your written comments only to one of the following addresses prior to the close of the comment period: a. For delivery in Washington, DC—Centers for Medicare & Medicaid Services Department of Health and Human Services, Room 445–G, Hubert H. Humphrey Building, 200 Independence Avenue, SW., Washington, DC 20201.

Because access to the interior of the Hubert H. Humphrey Building is not readily available to persons without Federal government identification, commenters are encouraged to leave their comments in the CMS drop slots located in the main lobby of the building. A stamp-in clock is available for persons wishing to retain a proof of filing by stamping in and retaining an extra copy of the comments being filed.)

b. For delivery in Baltimore, MD—Centers for Medicare & Medicaid Services Department of Health and Human Services, 7500 Security Boulevard, Baltimore, MD 21244–1850.

If you intend to deliver your comments to the Baltimore address, call (410) 786–9994 in advance to schedule your arrival with one of our staff members.

Comments erroneously mailed to the addresses indicated as appropriate for hand or courier delivery may be delayed and received after the comment period.

Submission of comments on paper requirements. You may submit comments on this document’s paper requirements by following the instructions at the end of the “Collection of Information Requirements” section in this document.

For information on viewing public comments, see the beginning of the SUPPLEMENTARY INFORMATION section.

FOR FURTHER INFORMATION CONTACT: Cynthia Moreno (410) 786–1164.

SUPPLEMENTARY INFORMATION: Inspection of Public Comments: All comments received before the close of the comment period are available for viewing by the public, including any personally identifiable or confidential business information that is included in a comment. We post all comments received before the close of the comment period on the following Web site as soon as possible after they have been received: http://www.regulations.gov. Follow the search instructions on that Web site to view public comments.

Comments received timely will also be available for public inspection as they are received, generally beginning approximately three weeks after publication of a document, at the headquarters of CMS, 7500 Security Boulevard, Baltimore, Maryland 21244, Monday through Friday of each week from 8:30 a.m. to 4 p.m. To schedule an appointment to view public comments, phone 1–800–743–3951.

I. Background

The Balanced Budget Act of 1997 (BBA) (Pub. L. 105–33) established the Medicare+Choice (M+C) program. Under section 1851a(1) of the Social Security Act (the Act), every individual with Medicare Parts A and B, except for individuals with end stage renal disease, could elect to receive benefits either through the original Medicare program or an M+C plan, if one was offered where the beneficiary lived. The primary goal of the M+C program was to provide Medicare beneficiaries with a wider range of health plan choices.


Sections 201 through 241 of Title II of the MMA made significant changes to the M+C program. As directed by Title II of the MMA, we renamed the M+C program the Medicare Advantage (MA) program. We also revised our regulations to include new payment and bidding provisions based largely on risk, to recognize the addition of regional Preferred Provider Organization plans, to address the provision of prescription drug benefits, and to improve incentives for MA plans to improve the quality of care they provide.