

Dated: June 19, 2008.

Ivor A. Pritchard,

Acting Director, Office for Human Research Protections.

[FR Doc. E8-14917 Filed 6-30-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration on Aging

Agency Information Collection Activities; Submission for OMB Review; Comment Request; Extension of Certification on Maintenance of Effort for the Title III and Certification of Long-Term Care Ombudsman Program Expenditures

AGENCY: Administration on Aging, HHS.

ACTION: Notice.

SUMMARY: The Administration on Aging (AoA) is announcing that the proposed collection of information listed below has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.

DATES: Submit written comments on the collection of information by July 31, 2008.

ADDRESSES: Submit written comments on the collection of information by fax 202.395.6974 to the OMB Desk Officer for AoA, Office of Information and Regulatory Affairs, OMB.

FOR FURTHER INFORMATION CONTACT:

Rodd Clay, e-mail: rodd.clay@aoa.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, AoA has submitted the following proposed collection of information to OMB for review and clearance. With respect to the following collection of information, AoA invites comments on: (1) Whether the proposed collection of information is necessary for the proper performance of AoA's functions, including whether the information will have practical utility; (2) the accuracy of AoA'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.

The Certification on Maintenance of Effort for the Title III and Certification

of Long-Term Care Ombudsman Program Expenditures provides statutorily required information regarding state's contribution to programs funded under the Older Americans Act and conformance with legislative requirements, pertinent Federal regulations and other applicable instructions and guidelines issued by Administration on Aging (AoA). This information will be used for Federal oversight of Title III Programs and Title VII Ombudsman Program.

AoA estimates the burden of this collection of information as follows: 56 State Agencies on Aging respond annually which should be an average burden of one half (1/2) hour per State agency per year or a total of twenty-eight hours for all state agencies annually. In the **Federal Register** of March 19, 2008 (Vol. 73, No. 54 Page 14821), the agency requested comments on the proposed collection of information. No comments on the content of the collection were received.

Dated: June 26, 2008.

Josefina G. Carbonell,

Assistant Secretary for Aging.

[FR Doc. E8-14898 Filed 6-30-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

Statement of Organization, Functions, and Delegations of Authority

Part F of the Statement of Organization, Functions, and Delegations of Authority for the Department of Health and Human Services, Centers for Medicare & Medicaid Services (CMS), (**Federal Register**, Vol. 73, No. 46, pp. 12451-12452, dated Friday, March 7, 2008) is amended to reflect a change in the name and updates to the functions for the Center for Beneficiary Choices.

Part F. is described below:

• Section F. 20. (Functions) reads as follows:

Center for Drug and Health Plan Choice (FAE)

• Responsible for all national policies and operations necessary for the purchasing of Medicare Prescription Drug (Part D) and Medicare Advantage (Part C) health plan benefits. Designs, implements, and manages the procurement of prescription drug plans (PDPs) and Medicare Advantage plans (MA and MA-PD plans), including the solicitation and approval of

applications, review of benefits and negotiation of competitive bids, the implementation of quality improvement and performance measures, review of fiscal solvency and contractor management activities.

• Develops and improves all bidding and payment policies related to the Medicare Prescription Drug Benefit and the Medicare Advantage (MA) program.

• Validates payments to the Part D prescription drug and MA plans, including routine annual risk adjustment data validation based on medical record review.

• Coordinates the development and management of business requirements for the national systems for enrollment, payment, and contractor management for the Prescription Drug Benefit and the Medicare Advantage (MA) programs.

• Develops and implements the national policy and oversees operational implementation for all issues related to the Retiree Drug Subsidy Program.

• Develops national policy for eligibility, enrollment and entitlement for Medicare Parts A, B, C, and D, including oversight of activities related to Part D auto-enrollment, low income subsidy, and creditable coverage.

• Develops national policy and oversees operational activities related to Medicare Part A, B, C, and D claims-related hearings, appeals, grievances and other beneficiary-centered dispute resolution processes.

• Serves as the focal point for issues related to a variety of Federal standards affecting private health insurance coverage, including those pertaining to its administration of the Medigap program, Title I of the Health Insurance Portability and Accountability Act and the Consolidated Omnibus Budget Reconciliation Act.

• Works closely with the regional Consortium for Medicare Health Plans Operations (CMHPO) on all operational aspects of the Part C and Part D programs.

• Develops and implements Part C and Part D contractor performance monitoring programs and Part C and Part D compliance and oversight programs and carries out these programs collaboratively with CMHPO.

• Develops surveys to measure consumer experiences with their health plans and health care providers; manages the Consumer Assessment of Health Care Provider and Systems (CAHPS) survey; develops and prepares performance measures for Part C sponsors; analyzes and reports Health Plan Employers Data and Information Set data for Part C performance measures and consumer reports; and

assesses the effectiveness of CMS' quality reporting activities.

- Effectively communicates program policies related to the Prescription Drug and Medicare Advantage (MA) programs to health plans and drug plan contractors, employer group sponsors, beneficiary advocates and other stakeholders in the health care field.

- Develops new policies (e.g. health plan access, benefits, special needs plans) and programs to reflect changes in program objectives, the health care delivery system, beneficiary health care needs, and new plan types to support an appropriate range of choices for beneficiaries.

- Collaborates with our partners, such as industry, other government entities and advocacy groups, to understand their perspectives on Prescription Drug and Medicare Managed Care policies and procedures and to drive best practices in the health care industry.

- Develops and implements a comprehensive strategic plan, objectives and measures for overseeing an effective compliance and oversight program for all Part C (Medicare Advantage) and Part D (Medicare Prescription Drug) contractors in close collaboration with CMHPO, the Medicare Drug Benefit and C & D Data Group, the Medicare Drug and Health Plan Contract Administration Group and other Center for Drug and Health Plan Choice components.

- Develops and implements a comprehensive and effective audit program for all Part C (Medicare Advantage) and Part D (Medicare Prescription Drug) contractors.

Dated: June 21, 2008.

James W. Weber,

Acting Director, Office of Operations Management, Centers for Medicare & Medicaid Services.

[FR Doc. E8-14896 Filed 6-30-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2004-N-0188] (formerly Docket No. 2004N-0081)

Agency Information Collection Activities; Announcement of Office of Management and Budget Approval; Request for Designation as Country Not Subject to the Restrictions Applicable to Human Food and Cosmetics Manufactured From, Processed With, or Otherwise Containing, Material From Cattle

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a collection of information entitled "Request for Designation as Country not Subject to the Restrictions Applicable to Human Food and Cosmetics Manufactured from, Processed With, or Otherwise Containing, Material from Cattle" has been approved by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995.

FOR FURTHER INFORMATION CONTACT:

Jonna Capezzuto, Office of the Chief Information Officer (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-4659.

SUPPLEMENTARY INFORMATION: In the **Federal Register** of April 17, 2008 (73 FR 20785), the agency announced that the proposed information collection had been submitted to OMB for review and clearance under 44 U.S.C. 3507. An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number. OMB has now approved the information collection and has assigned OMB control number 0910-0623. The approval expires on June 30, 2011. A copy of the supporting statement for this information collection is available on the Internet at <http://www.reginfo.gov/public/do/PRAMain>.

Dated: June 24, 2008.

Jeffrey Shuren,

Associate Commissioner for Policy and Planning.

[FR Doc. E8-14882 Filed 6-30-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2008-N-0354]

Agency Information Collection Activities; Proposed Collection; Comment Request; Mental Models Study of Farmers' Understanding and Implementation of Good Agricultural Practices

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 and to allow 60 days for public comment in response to the notice. This notice solicits comments on the proposed study entitled "Mental Models Study of Farmers' Understanding and Implementation of Good Agricultural Practices."

DATES: Submit written or electronic comments on the collection of information by September 2, 2008.

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:

Jonna Capezzuto, Office of the Chief Information Officer (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-4659.

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