

and small business knowledge in general.

FOR FURTHER INFORMATION CONTACT:

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SUPPLEMENTARY INFORMATION: This notice is published in accordance with the provisions of the Federal Advisory Committee Act (FACA) (Pub. L. 92-463).

Dated: August 27, 2007

Michael J. Rigas

Deputy Associate Administrator, Office of Small Business Utilization, General Services Administration.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 2007N-0041]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Administrative Procedures for the Clinical Laboratory Improvement Amendments of 1998 Categorization; Correction

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice; correction.

SUMMARY: The Food and Drug Administration (FDA) is correcting a notice that appeared in the **Federal Register** of May 16, 2007 (72 FR 27573). The document announced that a proposed collection of information had been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995. The document was published with an error. This document corrects that error.

FOR FURTHER INFORMATION CONTACT: Denver Presley, Jr., Office of the Chief Information Officer (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-1472.

SUPPLEMENTARY INFORMATION: In FR Doc. E7-9435, appearing on page 27573 in the **Federal Register** of Wednesday, May 16, 2007, the following correction is made:

1. On page 27574, in the third column, in the third full paragraph, the sentence "The likely respondents for this collection are Investigational New Drug Application Sponsors." is

corrected to read "The likely respondents for this collection of information are manufacturers of medical devices."

Dated: August 23, 2007.

Jeffrey Shuren,

Assistant Commissioner for Policy.

[FR Doc. E7-17153 Filed 8-29-07; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Office of the Commissioner; Statement of Organizations, Functions, and Delegations of Authority

Part D, Chapter D-B, (Food and Drug Administration), of the Statement of Organization, Functions, and Delegations of Authority for the Department of Health and Human Services (35 FR 3685, February 25, 1970, and 60 FR 56605, November 9, 1995, 64 FR 36361, July 6, 1999, and in pertinent part at 57 FR 54239) is being amended to reflect the restructuring of the Office of the Commissioner (OC), Food and Drug Administration (FDA). This reorganization includes the establishment of four Deputy-level offices within the Office of the Commissioner, the changes are as follows:

I. Under Part D, Food and Drug Administration, delete the Office of the Commissioner (DA) in its entirety and replace with the following:

DA.10 Organization. The Food and Drug Administration (FDA) is headed by the Commissioner, Food and Drug and includes the following organizational units:

Office of the Commissioner (DA), Office of the Chief Counsel (DAA), Office of the Chief of Staff (DAB), Office of International and Special Programs (DAL), Office of Operations (DAM), Office of Policy, Planning and Preparedness (DAH), Office of Scientific and Medical Programs (DAE).

DA.20 Functions

A. *Office of the Commissioner D(A)*—The Office of the Commissioner (OC) includes the Commissioner and Deputy Commissioner who are responsible for the efficient and effective implementation of FDA mission.

B. *Office of the Chief Counsel (DAA)*—The Office of the Chief Counsel (OCC) is also known as the Food and Drug Division, Office of the General Counsel, Department of Health and Human Services), while administratively within the Office of the Commissioner, is part of the Office of

the General Counsel of the Department of Health and Human Services.

1. Is subject to the professional supervision and control of the General Counsel, Department of Health and Human Services (HHS), and represents FDA in court proceedings and administrative hearings with respect to programs administered by FDA.

2. Provides legal advice and policy guidance for programs administered by FDA.

3. Acts as liaison to the Department of Justice and other Federal agencies for programs administered by FDA.

4. Drafts or reviews all proposed and final regulations and Federal Register notices prepared by FDA.

5. Performs legal research and gives legal opinions on regulatory issues, actions, and petitions submitted to FDA.

6. Reviews proposed legislation affecting FDA that originates in HHS or on which Congress requests the views of the Department.

7. Provides legal advice and assistance to the Office of the Secretary on matters within the expertise of the Chief Counsel.

C. *Office of the Chief of Staff (DAB)*—The Office of the Chief of Staff (OCOS):

1. Advises and provides integrated policy analysis and strategic consultation to the Commissioner, Deputy Commissioners, Associate Commissioners, Center Directors and other FDA officials on activities and issues that affect significant agency programs, projects and initiatives. Often this function involves the most difficult problems, crisis situations and extremely complex issues of FDA.

2. Provides leadership, coordination and management of the Commissioner's priority policies and issues across the Office of the Commissioner and FDA-wide. Identifies, triages, supervises and tracks related actions from start to finish in conjunction with senior leadership across FDA.

3. Provides direct support to the Commissioner of Food and Drugs and serves as major point of contact between the FDA Centers and Offices and the Commissioner.

4. Serves as the principal liaison to HHS and coordinates and manages activities between FDA and HHS. Works with the FDA Centers and Offices to ensure assignments or commitments made related to these activities are carried out.

5. Serves as one of the Commissioner's primary strategic liaisons with staff, partners, and the community at large.

6. Manages budget and resources and provides operation oversight for the FDA's Office of Legislation, Office of the