As a result, in part, of recent actions by the Congress and the Administration, a third draft of the AFSS Framework will be presented at the public meeting. We will also discuss in more detail, where appropriate, several of the gaps identified in the Framework document. In addition, we will show how health consequence scoring is combined with exposure scoring to rank the risks of contaminants in animal feed. Swine feed will be used as the example. We also plan to briefly present the risk-based method being developed to rank feed inspectional programs.

II. Public Meeting

We are holding the public meeting in an effort to gather further information from you, our stakeholders, on changes to AFSS that will help minimize risks to animal and human health associated with animal feed. Prior to the public meeting, FDA will place in the docket (found in brackets in the heading of this document) two documents entitled “Draft AFSS Framework, 3rd Edition” and “Risk-Ranking of Feed Hazards: Swine Feed Example.” The Framework document will summarize the agency’s current efforts to modernize its animal feed safety program. The Risk-Ranking document will provide the methods for ranking potential biological and chemical hazards in feed, using swine feed as an example. Details of these methods will be discussed at the meeting. A draft agenda for the meeting will also be placed in the docket prior to the meeting.

An additional public meeting sponsored by the Center for Veterinary Medicine (CVM) will be held on May 13, 2008, at the same site as the AFSS public meeting. The purpose of the CVM meeting will be for the agency to receive comments on the pet food safety section of FDAAA (Public Law 110–85). Information on the CVM public meeting will be publishing elsewhere in this issue of the Federal Register.

II. Transcripts

Please be advised that as soon as a transcript is available, it will be accessible at http://www.fda.gov/ohrms/dockets/ac/acmenu.htm. It may be viewed at the Division of Dockets Management (see Comments). A transcript will also be available in either hardcopy or on CD-ROM, after submission of a Freedom of Information request. Written requests are to be sent to Division of Freedom of Information (HFI–35), Office of Management Programs, Food and Drug Administration, 5600 Fishers Lane, rm. 6–30, Rockville, MD 20857.

Dated: April 15, 2008.

Jeffrey Shuren,
Associate Commissioner for Policy and Planning.

Federal Register /Vol. 73, No. 77 /Monday, April 21, 2008/ Notices 21357
view received comments at http://www.regulations.gov.

Please note that on January 15, 2008, the FDA Web site transitioned to the Federal Dockets Management System (FDMS). FDMS is a Government-wide, electronic docket management system. Electronic submissions will be accepted by FDA only through FDMS at http://www.regulations.gov.

SUPPLEMENTARY INFORMATION:

I. Background

FDAAA was signed into law by the President on September 27, 2007 (Public Law 110–85). FDAAA Section 1002(a) directs that within 2 years, FDA must establish pet food ingredient standards and definitions, processing standards, and updated standards for pet food labeling that include nutritional and ingredient information. This same provision of the law also directs that, in developing these new standards, FDA obtain input from its stakeholders, including, but not limited to, AAFCO, veterinary medical associations, animal health organizations, and pet food manufacturers. This public meeting is an opportunity for interested stakeholders to present such input and for FDA to hear directly from the public.

In the Federal Register of January 7, 2008 (73 FR 1225), FDA announced its intention to hold a public meeting concerning FDAAA Section 1002(a) to gather input from the interested stakeholders and other members of the public. This announcement includes further details regarding the date and location of the public meeting, and also provides additional information regarding the topics and questions to be considered. After the meeting, FDA will review all of the comments made at the meeting and those submitted in writing through the mail or electronically to Docket No. FDA–2007–N–0442 (formerly Docket No. 2007N–0487).

FDA is sponsoring an additional public meeting as part of its Animal Feed Safety System (AFSS) initiative on May 14, 2008, at the same location as the May 13, 2008, FDAAA public meeting. The AFSS is a system that FDA is developing to minimize the risk to animals and public health through the use of risk-based, preventive, and comprehensive animal feed control measures. The purpose of the additional meeting will be for the agency to present the third draft of the AFSS Framework and work-in-progress on a method for ranking animal feed hazards by their risks to animal and human health.

The revised Framework document includes, among other things, recognition of FDA’s Food Protection Plan, which was announced in November 2007, and changes to the document necessitated by FDAAA. The ranking scheme for estimating risks posed by feed hazards to animal and human health consists of two components, health consequence scoring and exposure scoring, which were previously presented. At the May 14, 2008, public meeting, FDA will describe methods for ranking risks associated with biological and chemical hazards in feed, using swine feed examples.

Background material relating to AFSS, including previous drafts of the AFSS Framework document, is available at http://www.fda.gov/cvm/AFSS.htm.

II. Topics and Questions for Consideration at the May 13, 2008, Public Meeting:

FDA seeks input from stakeholders and other members of the public on the topics and questions discussed below. Given that time will be limited at the public meeting, FDA encourages all interested persons to submit their comments in writing to Docket No. FDA–2007–N–0442 to ensure that their comments are considered.

A. Scope of Meeting

In enacting FDAAA Section 1002(a), Congress specifically directed FDA to establish, in consultation with relevant stakeholders and other members of the public, ingredient standards and definitions, processing standards, and updated labeling standards for pet food. FDA seeks input from stakeholders and other members of the public on the development of such standards for pet food, including on the specific questions listed below.

In addition, because pet food is well-integrated into the overall animal foods and feeds industry, FDA is concerned that certain new requirements, if limited to pet food only, would be impractical to implement, difficult to enforce, and would not effectively provide the safety enhancements intended by FDAAA. Furthermore, because the standards mandated by FDAAA do not currently exist for any animal food or feed, limiting new requirements to pet food only would fail to address the broader food safety concerns associated with food intended for other animal species, particularly food-producing animals. FDA is interested in obtaining input from interested stakeholders and the public as to whether the ingredient standards and definitions and processing standards should be developed for all animal foods. There appears to be little or no difference between ingredients intended for use in pet foods and those intended for use in other animal foods and feeds. Therefore, the agency believes the most appropriate course of action is to develop ingredient standards and definitions and processing standards for all animal foods, including pet food.

FDA believes that such an approach would more effectively carry out the safety objectives of FDAAA, and the broader human food safety provisions of the Federal Food, Drug, and Cosmetic Act. The agency also seeks comment on this or other alternative approaches for implementing Section 1002(a) of FDAAA.

B. Pet Food Labeling

1. How could the nutritional information (e.g., guaranteed analysis, nutritional adequacy statements/life-stage claims) already present on pet food labels be improved?

2. How could the ingredient information already present on pet food labels (i.e., the ingredient list) be improved?

3. How could the current feeding instructions/recommendations section already present on pet food labels be improved?

4. Should feeding recommendations be required on the labels for all types of pet food?

5. Should a Nutrition Facts box, similar to the format that appears on human food labels, replace the current Guaranteed Analysis that currently appears on pet food labels? If so, how could this Nutrition Facts box be made to clearly distinguish it from human food labeling?

6. What other information should be required on pet food labels that is not generally present on pet food products sold in the United States?

7. Are there existing state laws, regulations, guidelines, or other models that FDA should consider when drafting the proposed pet food labeling?

C. Pet Food Ingredient Standards and Definitions

1. What kind of ingredient definitions would provide adequate information to ensure the safe and suitable use of the ingredients in pet foods? Should ingredient definitions also be developed for other animal foods in addition to pet food?

2. Should formal standards be a part of ingredient definitions? If so, what information should be considered to establish a standard? Should such standards be developed for ingredients intended for other animal feeds in addition to pet food?
D. Pet Food Processing Standards

The AFSS initiative is intended to cover the entire spectrum of agency activities from preapproval of food additives for use in feed, to establishing limits for feed contaminants, providing education and training, and conducting inspections and taking enforcement actions for ensuring compliance with agency regulations. Some basic elements of an animal feed safety system are described at: http://www.fda.gov/ohrms/dockets/98fr/03n-0312-bkg0002.pdf.

Would standards based on a risk-based, preventive, and comprehensive feed control measures approach, such as the approach described as an element of FDA’s AFSS initiative, adequately address the processing standards requirement of section 1002(a) of FDAAA? If so, what aspects of procurement, processing and distribution should be include in such an approach? Should such standards be developed and applied to all animal feeds rather than be limited to pet food?

III. Other Information for the Public Meeting

FDA has posted additional information for the May 13, 2008, public meeting on the CVM Web site at http://www.fda.gov/cvm. The agency may make additional background material available to the public and will post that information on the CVM Web site as well. Additionally, background material relating to AFSS, including previous drafts of the AFSS Framework document, is available at http://www.fda.gov/cvm/AFSS.htm.

IV. Transcripts

FDA will prepare a meeting transcript that will be entered into the docket. FDA anticipates that transcripts will be available approximately 30 business days after the meeting. The transcript will also be available for public examination at the Division of Dockets Management (HFA–305), 5630 Fishers Lane, rm. 1061, Rockville, MD 20852, between 9 a.m. and 4 p.m., Monday through Friday.

Dated: April 15, 2008.

Jeffrey Shuren,
Associate Commissioner for Policy and Planning.

[FR Doc. 08–1155 Filed 4–16–08; 3:48 pm]
BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Amended Notice of Meeting

Notice is hereby given of a change in the meeting of the Pregnancy and Neonatology Study Section, June 2, 2008, 8 a.m. to June 3, 2008, 3 p.m., Embassy Suites at the Chevy Chase Pavilion, 4300 Military Road, NW., Washington, DC, 20015 which was published in the Federal Register on April 4, 2008, 73 FR 18539–18542.

The meeting will be held one day only June 2, 2008, from 8 a.m. to 5 p.m. The meeting location remains the same. The meeting is closed to the public.

Dated: April 14, 2008.
Jennifer Spaeth,
Director, Office of Federal Advisory Committee Policy.

[FR Doc. E8–8450 Filed 4–18–08; 8:45 am]
BILLING CODE 4140–01–M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Office of the Director, National Institutes of Health; Office of Biotechnology Activities; Recombinant DNA Research; Notice of a Meeting of an NIH Blue Ribbon Panel

There will be a meeting of the NIH Blue Ribbon Panel to advise on the Risk Assessment of the National Emerging Infectious Diseases Laboratories (NEIDL) at the Boston Medical Center. The meeting will be held on Friday, May 2, 2008, at the National Institutes of Health, Building 31, Floor 6C, Conference Room 10, 31 Center Drive, Bethesda, Maryland 20892, from 8:30 a.m. to approximately 11:30 a.m.

The National Research Council Committee that provided technical input on the NIH’s Draft Supplementary Risk Assessments and Site Suitability Analyses for the NEIDL will participate in discussions with Panel members regarding the scope and design of additional studies that may be needed to assess risk associated with the siting and operation of the NEIDL.

For further information concerning this meeting contact Ms. Laurie Lewallen, Advisory Committee Coordinator, Office of Biotechnology Activities, Office of the Director, National Institutes of Health, 6705 Rockledge Drive, Room 750, Bethesda, MD 20892–7985, 301–496–9838, lewalla@od.nih.gov.

The meeting will be open to the public, with attendance limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the Contact Person listed above in advance of the meeting. Any interested person may file written comments with the panel by forwarding the statement to the Contact Person listed on this notice. The statement should include the name, address, telephone number and when applicable, the business or professional affiliation of the interested person.

NIH campus security procedures require that all visitor vehicles, including taxicabs and hotel and airport shuttles will be inspected before being allowed on campus. Visitors will be asked to show one form of identification (for example, a government-issued photo ID, driver’s license, or passport) and to state the purpose of their visit.

An agenda and any additional information for the meeting will be posted on the agency’s Web site: http://www.nih.gov/about/director/acd/index.htm.

Background information may be obtained by contacting NIH OBA by e-mail oba@od.nih.gov.

Dated: April 14, 2008.

Amy P. Patterson,
Director, Office of Biotechnology Activities.

[FR Doc. E8–8474 Filed 4–18–08; 8:45 am]
BILLING CODE 4140–01–P

DEPARTMENT OF HOMELAND SECURITY

Federal Emergency Management Agency

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

AGENCY: Federal Emergency Management Agency, DHS.

ACTION: Notice; 30-day notice and request for comments; Telephone Survey, OMB 1660–0057, Revision of a currently approved collection.

SUMMARY: The Federal Emergency Management Agency (FEMA) has submitted the following information collection to the Office of Management and Budget (OMB) for review and clearance in accordance with the requirements of the Paperwork Reduction Act of 1995. The submission describes the nature of the information collection, the categories of respondents, the estimated burden (i.e., the time, effort and resources used by respondents to respond) and cost, and