referral or services for individuals or couples where violence is occurring. Applicants should be able to demonstrate knowledge of the information and services provided by domestic violence coalitions within the community.

vii. Funding Thresholds. The funding thresholds for this program will be revised to reflect ANA's availability of funds within this special initiative program area. These thresholds allow ANA to provide funding to the maximum number of applicants. (Legal authority: Section 803(a) and (d) and 803C of the Native American Programs Act of 1974, as amended, 42 U.S.C. 2991 band 2991b—3.)

viii. Project Periods. The project periods reflect the review and assessment of projects monitored under this special initiative program area. These project periods allow ANA to provide funding to the maximum number of applicants. (Legal authority: Section 803(a) and (d) and 803C of the Native American Programs Act of 1974, as amended, 42 U.S.C. 2991b and 2991b–3.)

In the FY 08 PA, project periods will be:

- Priority Area 1—Planning: 12 months.
- Priority Area 2—Implementation: 36 months.

(C) ANA SEDS: In the FY 2008 PA for both priority areas, the program areas of interest (PAI) for social development projects changed. The Administration for Children and Families has expanded the focus of healthy marriage to include responsible fatherhood activities. In order to eliminate redundancy, this activity was added to the NAHMI PA. The grandparents PAI was included to promote inter-generational programs. (Legal authority: Section 803(a) and (d) and 803C of the Native American Programs Act of 1974, as amended, 42 U.S.C. 2991b and 2991b—3.)

The FY 2008 PA will replace the fatherhood PAI with the following:

• Projects that address the needs of grandparents raising grandchildren.

(D) ANA Mitigation: The FY 2008 PA removes all definitions related to inkind contributions, including in-kind contributions, leveraged resources, partnerships, and letters of commitment. Furthermore, the required number of impact indicators is reduced to one. These changes are reflective of Public Law 103–335 which does not require matching funds. (Legal authority: Section 803(a) and (d) and 803C of the Native American Programs Act of 1974, as amended, 42 U.S.C. 2991b and 2991b–3 and Public Law 103–335.)

Dated: January 2, 2008.

#### Quanah Crossland Stamps,

 $\label{lem:commissioner} Commissioner, Administration for Native \\ Americans.$ 

[FR Doc. 08–56 Filed 1–10–08 8:45 am] BILLING CODE 4184–01–M

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. 2007F-0478]

### Kemira Oyi; Filing of Food Additive Petition (Animal Use); Partially Ammoniated Formic Acid

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing that Kemira Oyi has filed a petition proposing that the food additive regulations be amended to provide for the safe use of partially ammoniated formic acid as a pH control agent in swine feed.

**DATES:** Submit written or electronic comments on the petitioner's environmental assessment by March 11, 2008.

ADDRESSES: You may submit written comments to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to: http://www.fda.gov/dockets/ecomments.

#### FOR FURTHER INFORMATION CONTACT:

Isabel W. Pocurull, Center for Veterinary Medicine, Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855, 240 453–6853, email: isabel.pocurull@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: Under the Federal Food, Drug, and Cosmetic Act (section 409(b)(5) (21 U.S.C. 348(b)(5))), notice is given that a food additive petition (FAP 2258) has been filed by Kristi O. Smedley, Center for Regulatory Services, Inc., 5200 Wolf Run Shoals Rd., Woodbridge, VA 22192-5755, United States agent for Kemira Oyi, Porkkalantatu 3, PO Box 330, 001000 Helsinki, Finland. The petition proposes to amend the food additive regulations in part 573—Food Additives Permitted in Feed and Drinking Water of Animals (21 CFR part 573) to provide for the safe use of partially ammoniated formic acid as a pH control agent in swine feed when used at levels up to 1.2 percent of the feed.

The potential environmental impact of this action is being reviewed. To

encourage public participation consistent with regulations issued under the National Environmental Policy Act (40 CFR 1501.4(b)), the agency is placing the environmental assessment submitted with the petition that is the subject of this notice on public display at the Division of Dockets Management (see ADDRESSES) for public review and comment.

Interested persons may submit to the Division of Dockets Management (see **ADDRESSES**) written or electronic comments regarding this document. Submit a single copy of electronic comments or two paper copies of any mailed comments, except that individuals may submit one paper copy. Comments are to be identified with the docket number found in brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday. FDA will also place on public display any amendments to, or comments on, the petitioner's environmental assessment without further announcement in the Federal Register. If, based on its review, the agency finds that an environmental impact statement is not required and this petition results in a regulation, the notice of availability of the agency's finding of no significant impact and the evidence supporting that finding will be published with the regulation in the Federal Register in accordance with 21 CFR 25.51(b).

Dated: December 31, 2007.

#### Bernadette Dunham,

Deputy Director, Center for Veterinary Medicine.

[FR Doc. E8–316 Filed 1–10–08; 8:45 am]

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

## Food and Drug Administration

# Anti-Infective Drugs Advisory Committee; Notice of Meeting

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). The meeting will be open to the public.

Name of Committee: Anti-Infective Drugs Advisory Committee.

General Function of the Committee: To provide advice and recommendations to the agency on FDA's regulatory issues.