1. DIVISION OF ANIMAL FEEDS (DJJVDB).

A. Evaluates food additive petitions, generally recognized as safe petitions and investigational food additive applications for adequacy of

- data on animal safety, utility, and stability;

- labeling, and

- manufacturing facilities and controls; coordinates the review of human food safety and environmental impact information. Recommends approval of food additives and GRAS substances to the Center Director.

B. Evaluates the safety of complete feeds, feed supplements, and feed ingredients for animals, including pets, and provides risk assessments on hazardous contaminants in animal feeds.

C. Evaluates safety data, manufacturing and use information, and labeling for complete animal feeds and for non-drug substances added to animal feeds to determine their legal status.

D. Provides technical and scientific evaluations of new feed ingredients defined in the Official Publication of the Association of American Feed Control Officials (AAFCO), coordinates feed regulatory activities with State feed control offices, and participates in the activities of AAFCO committees and task forces and the AAFCO Board of Directors.

E. Coordinates scientific reviews and develops policy recommendations for animal feed issues (e.g., contaminants, biotech plants, unapproved feed ingredients).
F. Evaluates Biotech Plant Notifications for adequacy of the composition and animal safety data, collaborates on Notes to Files with Center for Food Safety and Applied Nutrition (CFSAN), and recommends sign-off by the Office of Surveillance and Compliance (OSC) Director on letters to the firms concluding Food and Drug Administration consultations relating to feed issues. Develops policy recommendations on feed issues involving biotech plants, and participates in developing policy recommendations on general issues involving biotech plants with CFSAN, other Centers and Offices in FDA, and other Federal agencies.

G. Approves feed mill licenses after being assured that the licensee can manufacture and label medicated feed in compliance with agency regulations.

H. Maintains inventory of distributors of veterinary feed directives drugs.

I. Ensures that licensed facilities comply with the medicated feed license regulations by implementing and evaluating the results of an inspectional compliance program for these facilities.

J. Provides consulting reviews to the Office of New Animal Drug Evaluation (ONADE) for medicated feed labels, including medicated pet food labels.

K. Develops, monitors and evaluates Center for Veterinary Medicine (CVM) Compliance Programs or Field Assignments for medicated feeds, Type A medicated articles, and feed contaminants (e.g., BSE, microbial pathogens, mycotoxins, pesticides, heavy metals, industrial chemicals). Reports the findings from the program to the states, FDA field, and other interested parties.

L. Provides scientific support for regulatory actions for animal feeds and medicated animal feeds.

M. Recommends and may participate in intramural and extramural research projects conducted or coordinated by the Office of Research to gain further information on contaminants, drugs, and food additives.

2. AUTHORITY AND EFFECTIVE DATE.

The functional statements for this Office were approved by the Secretary for Health and Human Services, effective October 1, 2012.
The following is the Food and Drug Administration, Office of Foods and Veterinary Medicine, Center for Veterinary Medicine, Office of Surveillance and Compliance, Division of Animal Feeds organization structure depicting all the organizational structures reporting to the Office Director.

OFFICE OF THE DIRECTOR