This guidance document is an annex to the parent guidance VICH GL3, “Stability Testing of New Veterinary Drug Substances and Medicinal Products.” It addresses the recommendations for stability testing of new veterinary medicinal Medicated Premix products intended for submission for approval to the European Union, Japan and the United States.

Comments and suggestions regarding the document should be submitted to Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.regulations.gov.

For questions regarding this document, contact Norman Gregory, Center for Veterinary Medicine, (HFV-143), Food and Drug Administration, 7500 Standish Place, Rockville, MD 20855, 240-276-8253, e-mail: norman.gregory@fda.hhs.gov.

Additional copies of this final guidance document may be requested from the Communications Staff (HFV-12), Center for Veterinary Medicine, Food and Drug Administration, 7519 Standish Place, Rockville, MD 20855, and may be viewed on the Internet at http://www.fda.gov/AnimalVeterinary/GuidanceComplianceEnforcement/GuidanceforIndustry/default.htm.
STABILITY TESTING FOR MEDICATED PREMIXES

This guidance has been developed by the appropriate VICH Expert Working Group and has been subject to consultation by the parties, in accordance with the VICH Process. At Step 7 of the Process the final draft is recommended for adoption to the regulatory bodies of the European Union, Japan and USA.
STABILITY TESTING FOR MEDICATED PREMIXES

1. General

The VICH Harmonized Tripartite Guidance covering the Stability Testing of New Veterinary Drug Substances and Medicinal Products (hereafter referred to as the parent guidance) references an annex for Medicated Premixes. This document is an annex to the parent guidance and addresses the recommendations for stability testing of veterinary medicinal Medicated Premix drug products. The parent guidance (VICH GL3) provides a general indication of the information on product stability generated, but the annex for Medicated Premixes leaves sufficient flexibility to encompass a variety of different practical and scientific considerations that are specific to the characteristics of the drug products being evaluated. Other stability studies which might be important to consider like stability in relation to conditioning and pelleting, segregation and homogeneity studies are not within the scope of this guidance.

2. Preamble

The guidance primarily addresses the generation of acceptable stability information for submission in Registration Applications for medicated premix drug products containing new molecular entities. Medicated Premixes are intended for oral administration following incorporation into animal feed. The guidance only pertains to Medicated Premixes, and does not currently seek to cover information required for products manufactured from medicated premixes. Stability studies carried out with a medicated premix should be in line with the parent guidance. However, the application of the parent guidance may be limited in some instances. This guidance therefore describes those areas where there may be differences in the stability data package for medicated premixes.

3. Storage Test Conditions and Test Parameters

Medicated Premixes are recommended to be tested at 25°C ±2°C / 60% RH±5% (long-term testing) and 40°C ±2°C / 75% RH±5% (accelerated testing) and with the same schedule intervals as described in the Parent Guidance for drug product. Other storage conditions are allowable if justified. Where "significant change" occurs due to accelerated testing, additional testing at an intermediate condition e.g., 30°C ±2°C / 60% RH±5% should be conducted. "Significant change" at the accelerated condition is defined as failure to meet specifications. Evidence is necessary to demonstrate the stability of the Medicated Premix before incorporation into an additional feed. The shelf-life specification of a Medicated Premix should include necessary stability indicating test parameters.

4. Packaging Materials

The testing should be carried out in the final packaging proposed for marketing when practicable. The use of smaller comparable containers simulating the actual market packaging may be justified.
5. Glossary

Carrier - An edible material to which drug substances are added to facilitate uniform incorporation into feed.

Medicated Premix (Type A Medicated Article) - A Medicated Premix is a veterinary medicinal product consisting of a mixture of one or more drug substances, generally with a carrier, that is prepared to facilitate oral administration of the drug to animals when mixed with feed.

For additional definitions, please refer to regional guidance or regulations.