

FDA-2022-P-2795

Aurora Pharmaceutical, Inc. Attention: Alina Garbar, PhD RA Director 1196 Hwy 3 South Northfield, MN 55057

Re: Suitability petition approved

Dear Dr. Garbar:

We approve your suitability petition (FDA 2022-P-2795) dated November 2, 2022. You requested permission to submit an abbreviated new animal drug application (ANADA) for a generic meloxicam oral solution that differs in dosage form from the reference listed new animal drug (RLNAD). Specifically, the proposed generic new animal drug is an oral solution, while the RLNAD is an oral suspension. The RLNAD is Metacam® (meloxicam oral suspension), sponsored by Boehringer Ingelheim Animal Health USA, Inc., under NADA 141-213. Metacam® is approved for the control of pain and inflammation associated with osteoarthritis in dogs. The strengths of the proposed generic product and the approved strengths for the RLNAD are the same: 1.5 mg/mL and 0.5 mg/mL meloxicam.

Your proposed changes from the RLNAD are changes that can be considered through a suitability petition, as defined under section 512(n)(3) of the Federal Food, Drug, and Cosmetic Act (the FD&C Act). We find that the proposed changes do not require you to conduct investigations to show the safety and effectiveness of the generic new animal drug for its proposed intended uses. Therefore, we approve the petition under section 512(n)(3)(C) of the FD&C Act.

Approval of this suitability petition does not guarantee approval of the ANADA for your proposed generic new animal drug. We will make that determination after you have submitted your ANADA for our review.

When you submit your ANADA for your proposed generic new animal drug, you must identify the RLNAD referred to in this suitability petition and include a copy of this letter. We recommend that you request a presubmission conference, according to 21 CFR 514.5, to discuss the requirements and potential pathways for approval of the application.

A copy of this letter approving your petition will be placed on public display at <a href="https://www.regulations.gov">www.regulations.gov</a> with the Dockets Management Staff (HFA-305), Food and Drug Administration, 5630 Fishers Lane, Room 1061, Rockville, MD 20852. If you have any questions, please contact Lauren (Gypsi) Feeney, DVM, Director, Division of Generic Animal Drugs, at Lauren.Feeney@fda.hhs.gov.

Sincerely,

Matthew Lucia, DVM Director Office of New Animal Drug Evaluation Center for Veterinary Medicine

cc: HFV-170 (Petition File)
HFA-305 (Dockets Management Staff)