



ANIMAL MEDICINAL DRUG USE CLARIFICATION ACT (AMDUCA) DISCLOSURE FORM

This form is only for disclosure of extra-label (not in accordance with the approved labeling) drug use in a market animal by or on the lawful order of a licensed veterinarian. This form is not to be used to disclose drug use that was in accordance with the approved labeling. It is a violation of the Livestock Handbook to falsify entries, records, or registration of animals. A separate form is required for <u>each</u> extra-label drug.

This form must be properly completed and provided to The Delaware State Fair at the time of any Fair-required testing during The Delaware State Fair. This form is provided for the exception in Section H "Market Animal Rules", page 13 of the Livestock Handbook. Animals are still subject to all other provisions of the Livestock Handbook, for example violations for competitive advantage drugs and violative residue limits.

By law, extra-label drug use is limited to treatment modalities when the health of an animal is threatened or suffering or death may result from failure to treat.

Exhibitor:

Ι,

(print exhibitor name)

(print exhibitor address)

Species of animal and Tag ID #:_____

(print veterinarian name)

_____, a licensed veterinarian, certify that the following are all true:

- 1. I have a valid veterinarian-client-patient relationship, as defined by 21 C.F.R. § 530.3, in regard to the above animal.
- 2. Without treatment, the health of the above animal is threatened or suffering or death may result.
- 3. There is no approved new animal drug that is labeled for such use and that contains the same active ingredient which is in the required dosage form and concentration, except if I have found that the approved new animal drug is clinically ineffective for its intended use in the above animal.
- 4. Prior to prescribing or dispensing an approved new animal or human drug for an extra-label use in the above animal, I:
 - a. Made a careful diagnosis and evaluation of the conditions for which the drug is to be used;
 - b. Established a substantially extended withdrawal period supported by appropriate scientific information;
 - c. Instituted procedures to assure that the identity of the animal is carefully maintained; and
 - d. Took appropriate measures to assure that assigned timeframes for withdrawal are met and no illegal drug residues occur in the above animal.

5. The established name of the drug:

6. The active ingredient(s) of the drug:

7. The condition treated:

8. The dosage administered:

- 9. The dates of treatment:
- 10. The treatment was in compliance with the Animal Medicinal Drug Use Clarification Act of 1994 (P.L. 103-396, 21 U.S.C. 360b(a)) and the corresponding regulations (21 C.F.R. § 530.1 *et seq.*)

Date:_____

DSF Initials:

Exhibitor compliance with all applicable rules and regulations is essential to the integrity of The Delaware State Fair. Violations of any applicable rule or regulation may result in exhibitor sanctions including disqualification, forfeiture of monetary & non-monetary awards, and/or suspension of future Fairs. Rev. 2.24.23