HUMAN SERVICES Food and Drug Administration	rotocol for cal Laboratory and veness Studies		roved: OMB No. 0910-0524 Date: 04/30/2010
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Food and Drug Administration Center for Veterinary Medicine, HFV- 7500 Standish Place Rockville, Maryland 20855	A1. DATE:		
	A2. DOCUMENT ID:		
	A3. STUDY / TRIAL ID:		
	A4. TYPE OF STUDY: Pi	votal	Non-pivotal

The applicant,

, submits a protocol for use of an

investigational new animal drug. Protocols for non-clinical laboratory studies (safety studies) are required under 21 CFR 58.120. Protocols for adequate and well-controlled effectiveness studies are required under 21 CFR 514.117(b). Applicants may request that CVM review protocols for safety and effectiveness studies of new animal drugs. This information is submitted in electronic form.

## I. Requesting Protocol CVM Review: Yes No

 NAME(S) OF THE DRUG(S): 1a. Established Name(s):

1b. Trade Name(s):

2. PROTOCOL TITLE:

2a. Short Abstract Title:

2b. Full Title:

2c. Version Number (If Applicable):

 3.
 PROTOCOL PREVIOUSLY SUBMITTED TO CVM:
 YES
 NO

 If Yes,
 3a. Date Submitted to CVM:
 3b. CVM Submission Identifier:

## II. Comments:

If you have additional comments that you would like to include in this submission please press the Insert Comments button below. All comments must be included within a PDF document.

## III. Protocol:

Please press the Insert Protocol button to include your Protocol. All Protocols must be included within a PDF document.

## IV. Applicant Information:

- 1. Name:
- 2a. Address:
- 2b. Address 2:
- 2c. City:
- 2e. Country:

2d. State/Prov: 2f. Postal Code:

- 3. Contact Name:
- 4. Contact Phone Number:
- 5. Contact Fax Number:
- 6. Contact E-Mail Address: