According to the Paperwork Reduction Act of 1995, an agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a valid OMB control number. The valid OMB control number for this information collection is 0579-0013. The time required to complete this information collection is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and

OMB Approved 0579-0013 EXP: 03/2027

completing and reviewing the collection of information.		 						
					e instructions on reverse side for additional guidance.			
U.S. DEPARTMENT OF AGRICULTURE ANIMAL AND PLANT HEALTH INSPECTION SERVICE VETERINARY SERVICES, CENTER FOR VETERINARY BIOLOGICS				FOR VETERINARY BIOLOGICS USE ONLY USDA PRODUCT CODE NUMBER				
APPLICATION FOR								
UNITED STATES VETERINARY BIOLOGICAL PRODUCT LICENSE 2. NAME AND ADDRESS OF APPLICANT (Include No., Street, or RFD No., City, State, ZIP Code)				VETERINARY BIOLOGICS ESTABLISHMENT NUMBER				
2. INAME AND ADDICESS OF AFFEIGANT (Include No.,	No., Oily, State, 217 Gode)		1. VETERINARY BIOLOGICS ESTABLISHIVIENT NOWIBER					
				3. ADDRESS OF PREMISES TO BE USED IF DIFFERENT FROM ITEM 2				
4. BIOLOGICAL PRODUCT TRUE NAME				5. APPLICANT'S INTERNAL WORKING IDENTIFIER FOR				
						PRODUCT (if applicable)		
6. RELATIONSHIP OF NEW PRODUCT TO OTHER LIC	ENSED PREI	ICENSE OR TERMINATED PRO	ODUCTS FOR Y	OUR ESTABL	ISHMENT (See in:	structions on reverse	e for detailed quidance \	
c. New Modern C. New Modern Comment and	2.1025, 1.112		0200101011	0011 2011122	.0		ror actanca gardanico.)	
7. OTHER COMMENTS								
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ITEMS SUBMITTED	DESCRIPTION					A. WITH THIS APPLICATION (X)	LOG ID OF PREVIOUS	
						. ,	SUBMISSION	
8. METHOD OF PRODUCTION	U OUTLIN	NE OF PRODUCTION (9 CFR 114	4.9) ∐SI	MILAR INFOR	MATION			
O DDIOD CUDMICCIONIC (IE ANNO FOR THE								
9. PRIOR SUBMISSIONS (IF ANY) FOR THIS PRODUCT WHICH WERE PROCESSED BEFORE A								
USDA PRODUCT CODE WAS ASSIGNED								
40. OTHER (********)								
10. OTHER (specify)								
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In accordance with the Act of Congress appro- herein named animal biological product for us								
comply with the provisions of the said Act, and								
distribution of the animal biological product, ar								
particular.								
In case a product license is issued, it is further	r agreed the	at the higherical product she	all ha subject	to any oddit	ional requirem	ants or restriction	ne stated therein	
11. SIGNATURE OF AUTHORIZED OFFICIAL	12. TITLE	ui ne sunject	o arry audit	13. DATE SIGNE		เง งเลเซน แโซเซเก.		
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INSTRUCTIONS FOR APHIS FORM 2003:

Submit one copy of the form. If additional space is needed, attach additional sheets and refer to Item No.

1. VETERINARY BIOLOGICS ESTABLISHMENT NUMBER

Enter the veterinary biologics establishment number assigned by APHIS, if one has been assigned.

2. NAME AND ADDRESS OF APPLICANT

Enter the establishment name and complete mailing address (street, city, state, ZIP) of the applicant. If the applicant has been assigned a veterinary biologics establishment number by APHIS, enter the mailing address on file with APHIS.

3. INTENDED SITE(S) OF MANUFACTURE AND TESTING

List the intended site(s) of manufacture and testing for this product.

4. BIOLOGICAL PRODUCT TRUE NAME

The True Name includes all of the antigenic fractions of the product for which a label claim is intended. See the current catalog of licensed biological products (https://www.aphis.usda.gov/aphis/ourfocus/animalhealth/veterinary-biologics/ct_vb_licensed_products) for examples. The True Name listed by the applicant on this application should be considered preliminary. APHIS assigns True Names according to numerous established conventions intended to promote standardization, so the True Name under which the product is licensed may differ.

5. APPLICANT'S INTERNAL WORKING IDENTIFIER FOR PRODUCT

Manufacturers frequently designate working identifiers for new products prior to being assigned a USDA product code. If such an identifier exists, it may be entered here.

6. RELATIONSHIP OF NEW PRODUCT TO OTHER LICENSED, PRELICENSE, OR TERMINATED PRODUCTS FOR YOUR ESTABLISHMENT

To facilitate product classification and licensing plan development, describe the relationship of this new product with other products licensed, or under development, by your establishment. This may include, but is not limited to, describing whether the new product is:

- Prepared from new or previously approved Master Seeds/Cells
- Part of a product line (cite other products in the line, especially those already licensed)
- · A modification of an existing product (e.g., adding or deleting an antigen, change in adjuvant/preservative/dose volume)
- Manufactured or tested with new technology for your establishment
- · Part of a split manufacture agreement

7. OTHER COMMENTS

Optional field for miscellaneous comments not covered elsewhere.

CHECKLIST OF SUPPORTING MATERIAL

This checklist is intended to ensure that APHIS has adequate information to assign a Product Code and True Name to the new product. If supporting material was not submitted previously, ensure that it is provided with this application.

8. METHOD OF PRODUCTION

The most efficient means of describing how the product is made is to provide an Outline of Production, formatted according to 9 CFR 114.9. If an Outline is not yet available, provide the same general information captured in an Outline of Production.

9. LIST OF PRIOR SUBMISSIONS (IF ANY) FOR THIS PRODUCT WHICH WERE PROCESSED BEFORE A USDA PRODUCT CODE WAS ASSIGNED

Manufacturers may submit documentation, such as requests to ship experimental product or proof-of-concept studies, before submitting a formal license application and being assigned a USDA product code. If such submissions exist for this product, please list them so that these uncoded submissions may be transferred to the licensing file for this product.

10. OTHER

APHIS may request additional information to support initial applications for certain products. If this has been requested for your product, briefly describe the purpose of the additional information in the line provided and attach supporting documentation.

11. SIGNATURE OF AUTHORIZED OFFICIAL

The APHIS primary or alternate liaison for the establishment, if designated, should serve as the authorized official. If no liaison has yet been designated, an official authorized to assume responsibility for regulatory compliance on behalf of the establishment should sign.

12. TITLE

Enter the job title of the individual signing in Item 11.

13. DATE SIGNED

This date should correspond to the date the application is submitted. This will be the submission date cited in any return correspondence.