

## REQUEST FOR EXEMPTION OF SELECT AGENTS AND TOXINS FOR AN INVESTIGATIONAL PRODUCT (APHIS/CDC FORM 5)

FORM APPROVED OMB NO. 0920-0576

EXP DATE: 02/28/2027

Answer all items completely and type or print in ink. Detailed instructions are available at <a href="https://www.selectagents.gov/form5.html">https://www.selectagents.gov/form5.html</a>. This form must be signed and submitted to either DASAT or DRSC:

Animal and Plant Health Inspection Service Division of Agricultural Select Agents and Toxins 4700 River Road Unit 2, Mailstop 22, Cubicle 1A07 Riverdale, MD 20737

FAX: (301) 734-3652 E-mail: <u>DASAT@usda.gov</u> Centers for Disease Control and Prevention Division of Regulatory Science and Compliance 1600 Clifton Road NE, Mailstop H21-4 Atlanta, GA 30329

FAX: (404) 718-2096 E-mail: <u>lrsat@cdc.gov</u>

## Submit completed form only once by either eFSAP, fax, or email

SECTION 1 – TO BE COMPLETED FOR INVESTIGATIONAL PRODUCT EXEMPTION				
1. Entity name:				
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2. Entity address (NOT a post office address):		3. City:	4. State:	5. Zip code:
6. Applicant		7. Title:		l.
First: MI:	Last:			
8. Telephone #:		9. E-mail address:		
10. FDA IND/INAD/IDE number:	11. FDA product name:	12. This product has been appro by FDA:	ved for Phase □ No □ Y	
13. Date of the IND/INAD/IDE application submitted to FDA including the name of the FDA center and review office FDA Center/Review Office:  Date:				
14. USDA veterinarian product code number:	15. USDA veterinarian product name:	16. This product has been tested and approved for field trials by USDA: □ No □ Yes		
17. Investigational product (Give select agent name and characterization):				
40.5				
18. Federal act that authorizes investigational use of this product:				
Provide a detailed justification to request an (attach additional sheets if necessary):				
I hereby certify that the information contained on this form is true and correct to the best of my knowledge. I understand that if I knowingly provide a false statement on any part of this form, or its attachments, I may be subject to criminal fines and/or imprisonment. I further understand that violations of 7 CFR Part 331, 9 CFR Part 121, or 42 CFR Part 73 may result in civil or criminal penalties, including imprisonment. For exemption requests that involve the investigational product that is, bears, or contains select agents or toxin, I authorize FDA to confirm for APHIS or CDC the existence and status of the IND, INAD, or IDE, and agree that such confirmation will not violate FDA's information disclosure regulations, the Federal Food, Drug, and Cosmetic Act, or the Trade Secrets Act (18 U.S.C. § 1905).				
Signature of Investigational Product Exemption	n Applicant:	Dat	e:	
Public reporting burden: Public reporting burden of this collection of information is estimated to average 30 minutes per response, including the time for				

**Public reporting burden:** Public reporting burden of this collection of information is estimated to average 30 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC/ATSDR Reports Clearance Officer; 1600 Clifton Road NE, MS D-74, Atlanta, Georgia 30329; ATTN: PRA (0920-0576).