

## GUIDANCE DOCUMENT FOR REQUEST FOR EXEMPTION OF SELECT AGENTS AND TOXINS FOR PUBLIC HEALTH OR AGRICULTURAL EMERGENCY OR INVESTIGATIONAL PRODUCT (APHIS/CDC FORM 5)

FORM APPROVED OMB NO. 0579-0213 OMB NO. 0920-0576 EXP DATE 10/31/2014

#### INTRODUCTION

The U.S. Departments of Health and Human Services (HHS) and Agriculture (USDA) published final rules (7 CFR 331, 9 CFR 121, and 42 CFR 73), which implement the provisions of the *Public Health Security and Bioterrorism Preparedness and Response Act of 2002* (Public Law 107-188) setting forth the requirements for possession, use, and transfer of select agents and toxins. The select agents and toxins identified in the final rules have the potential to pose a severe threat to public health and safety, to animal and plant health, or to animal and plant products. Responsibility for providing guidance on this form was designated to the Centers for Disease Control and Prevention (CDC) by the HHS Secretary and to the Animal and Plant Health Inspection Service (APHIS) by the USDA Secretary. In order to minimize the reporting burden to the public, APHIS and CDC have developed a common reporting form for this data collection.

An entity may apply for an exemption from the requirements of 7 CFR 331, 9 CFR 121, or 42 CFR 73 in order to: (a) use an investigational product that is, bears, or contains select agents or toxins, or, (b) provide a response to a public health or agricultural emergency. This exemption request (APHIS/CDC Form 5) should be sent to either APHIS or CDC for consideration:

Animal and Plant Health Inspection Service Agricultural Select Agent Program 4700 River Road Unit 2, Mailstop 22, Cubicle 1A07 Riverdale, MD 20737

FAX: 301-734-3652

E-mail: ASAP@aphis.usda.gov

Centers for Disease Control and Prevention Division of Select Agents and Toxins 1600 Clifton Road NE, Mailstop A-46 Atlanta, GA 30333

FAX: 404-718-2096 Email:<u>Irsat@cdc.gov</u>

#### **PURPOSE**

The purpose of this form is to request exemptions:

- 1. For exemption requests that involve the investigational product that is, bears, or contains select agents or toxins, APHIS or CDC will confirm that the Food and Drug Administration (FDA) has accepted or approved, under the authority of the Food, Drug, and Cosmetics Act (21 U.S.C. 301 et. seq.), an Investigational New Drug application (IND), Investigational New Animal Drug (INAD) application or an Investigational Device Exemption (IDE) application for a clinical trial involving the use of an investigational product that is, bears, or contains a select agent or toxin.
- 2. For the response to an extraordinary public health or agricultural emergency(ies).

A copy of the completed form and attachments must be maintained by the entity for three years.

This exemption form (APHIS/CDC Form 5) is not to be used if you are applying for an exclusion of an attenuated strain of a select agent or toxin. To apply for an exclusion, an applicant must submit a written request and supporting scientific information to APHIS or CDC (See 7 CFR § 331.3 (e), 9 CFR §§ 121.3(e) and 121.4(e), or 42 CFR §§ 73.3(e) and 73.4(e)).

## **INSTRUCTIONS**

- The applicant must complete, sign and date this form. For registered entities, the information provided for this form should match the information submitted for the entity's certificate of registration.
  - a. For applying for an exemption of an investigational product that is, bears, or contains select agents or toxins, complete section 1.
  - b. For applying for an exemption to respond to a public health or agricultural emergency, complete section 2.
- Fax, mail, or e-mail the form to APHIS or CDC.

## **OBTAINING EXTRA COPIES OF THIS FORM**

To obtain additional copies of this form, contact APHIS at (301) 851-3300 or CDC at (404) 718-2000. This guidance document and form are also available at <a href="http://www.selectagents.gov">http://www.selectagents.gov</a>.

(CDC Adobe Acrobat .0 Electronic Version, 1/200 )



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Read all instructions carefully before completing the form. Answer all items completely and type or print in ink. The form must be signed and submitted to either APHIS or CDC:

Animal and Plant Health Inspection Service Agricultural Select Agent Program 4700 River Road Unit 2, Mailstop 22, Cubicle 1A07

Riverdale, MD 20737 FAX: 301-734-3652

E-mail: ASAP@aphis.usda.gov

Centers for Disease Control and Prevention Division of Select Agents and Toxins 1600 Clifton Road NE, Mailstop A-46 Atlanta, GA 30333

FAX: 404-718-2096 Email: <a href="mailto:lrsat@cdc.gov">lrsat@cdc.gov</a>

SECTION 1 – TO BE COMPLETED FOR INVESTIGATIONAL PRODUCT EXEMPTION									
1. Entity name:		2. Entity registration number (if applicable):							
3. Entity address (NOT a post office address):		4. City:	5. State:	6. Zip code:					
7. Applicant		8. Title:	•						
First: MI:	Last:								
9. Telephone #:	10. FAX #:	11. Email address:							
12. Are you the: ☐ Facility Director ☐ Responsible Official ☐	☐ Other (specify):								
13. FDA IND/INAD/IDE number:	14. FDA product name:	15. This product has been appr by FDA:	oved for Phas  ☐ No ☐ Y						
16. Date of the IND/INAD/IDE application sub FDA Center/Review Office:	_	DA center and review office ate:							
17. USDA veterinarian product code number:	18. USDA veterinarian product name:	19. This product has been tested and approved for field trials by USDA: □ No □ Yes							
20. Investigational product (Give select agent	name and characterization):								
21. Federal act that authorizes investigational	use of this product:								
22. Provide a detailed justification to request a (attach additional sheets if necessary):  .	an exemption for the use of an investigation	nal product that is, bears, or conta	ins select age	nts or toxins					
I hereby certify that the information contained statement on any part of this form, or its attact 331, 9 CFR 121, or 42 CFR 73 may result in product that is, bears, or contains select ager and agree that such confirmation will not viol. Act (18 U.S.C. § 1905).  Signature of Investigational Product Exemptic	chments, I may be subject to criminal fines civil or criminal penalties, including imprisonts or toxin, I authorize FDA to confirm for attemption in the FDA's information disclosure regulation	and/or imprisonment. I further un- nment. For exemption requests the APHIS or CDC the existence and s s, the Federal Food, Drug, and Co	derstand that nat involve the status of the II	violations of 7 CFR investigational ND, INAD, or IDE,					

SECTION 2 – TO BE COMPLETED FOR PUBLIC HEALTH OR AGRICULTURAL EMERGENCY EXEMPTION								
23. Entity name:		24. Entity registration number (if applicable):						
25. Entity address (NOT a post office address):			City:		27. State:	28. Zip code:		
29. Applicant				30. Title:				
First: MI: Last: 31. Telephone #: 32. FAX #:			33. Email address:					
34. Are you the:	la Official III Other (creeif.)							
☐ Facility Director ☐ Responsible Official ☐ Other (specify):  35. Name of person most familiar with public health or agricultural emergency  36. Title:								
First: MI: 37. Name of entity (if different that		38. Telephone #:						
39. Address (NOT a post office ad	ddress):	4	0. City:	41.	State:	42. Zip Code:		
43. Description of select agent(s) involved in public health or agricultural emergency:								
44. Describe public health or agric	cultural emergency including historical,	clinical,	and epidemiological	l details of eme	rgency:			
45. Date of first confirmed case:	46. Date reported on APHIS/CDC	47. Nu	mber of cases	48. How dia	agnosis was mad	de:		
			veekly:		50. Telephone #:			
51. Address (NOT a post office address): 52. C					53. State: 54. Zip Code:			
, .	to request an exemption in response t		•			·		
necessary):								
	INFORMATION ON SELECT	AGEN	ITS AND TOXIN	S INVOLVE	D			
56. Location where laboratory testing will be conducted Building: Room:			57. Biosafety level (BSL) of laboratory or PPQ containment designation:					
58. Name of Principal Investigator First: MI:	r Last:	·						
59. Type of specimens that will be ☐ Clinical/diagnostic specimens 60. Is this source expected to pro If yes, give the anticipated	s ☐ Environmental specimens vide additional specimens? ☐ Unk	□ known		Other (specify): Yes				
statement on any part of this form	on contained on this form is true and co n, or its attachments, I may be subject t nay result in civil or criminal penalties, in	to crimin	al fines and/or impri					
Signature of Emergency Exempti					Date:			
Public reporting burden: Public	reporting burden of this collection of ir	nformatio	on is estimated to av	erage 1 hour pe	er response, incl	uding the time for		

Public reporting burden: Public reporting burden of this collection of information 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 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 30333; ATTN: PRA (0920-0576).