

REPORTING THE IDENTIFICATION OF A SELECT AGENT OR TOXIN FROM A CLINICAL/DIAGNOSTIC SPECIMEN (APHIS/CDC FORM 4A)

INSTRUCTIONS

Detailed instructions are available at <u>http://www.selectagents.gov/form4.html</u>. Answer all items completely and type or print in black ink. This report must be signed and submitted to either APHIS or CDC:

Animal and Plant Health Inspection Service Agriculture Select Agent Services 4700 River Road Unit 2, Mailstop 22, Cubicle 1A07 Riverdale, MD 20737 FAX: (301) 734-3652 E-mail: <u>AgSAS@aphis.usda.gov</u> Centers for Disease Control and Prevention Division of Select Agents and Toxins 1600 Clifton Road NE, Mailstop A-46 Atlanta, GA 30329 FAX: (404) 471-8469 E-mail: <u>CDCForm4@cdc.gov</u>

Accession Number:

(For Program Use ONLY)

Submit completed form only once by either e-mail, fax, or mail

SECTION A – REFERENCE LABORATORY INFORMATION										
1. Name of individual completing Sections A and B:			2. E-mail address:			3. Telephone #:				
First: MI:	Last:									
4. Registered Entity (APHIS or CDC Registration #:			9. Entity name:							
Clinical or Diagnostic Laboratory [non-registered entity (NRE)]										
(NRE # (provided by APHIS or CDC):)										
5. Responsible Official or Laboratory Super First: MI:	10. Address (NOT a post office address):									
6. E-mail address:	7. Telephone #:	8. Fax #:	11. City:	1	2. State:	13. Zip Code:				
	ELECT AGENT OR TO				SDECIM	IEN/S)				
				-						
1. Select Agent or Toxin Identified:			2. Date identified:							
3. Case/patient/sample ID #(s):	. Case/patient/sample ID #(s): 4. # of samples received: 5. Sample type received:		ived: 6. Case/patient ori		nt origin (z	gin (zip code):				
• • •					3 (1)					
7. Type of test performed (e.g., PCR, mouse bioassay, ELISA):										
8. Dispositions of select agent or toxin by e										
□ Transferred (Provide entity name and	Transferred (Provide entity name and date of transfer. Entity:			Date:)				
Destroyed (Provide destruction metho	Date:)								
Retained (Provide name of Principal Investigator retaining sample. Name:)										
9. Were any of the samples containing a select agent or toxin handled outside of primary containment which may have led to an unintentional release and/or exposure to the										
select agent or toxin?										
□ No □ Yes (If Yes, you are required under 7 CFR Part 331.19, 9 CFR Part 121.19, and 42 CFR Part 73.19 to complete and submit an APHIS/CDC Form 3) 10. Do you anticipate receiving additional samples/specimens for this case/patient that originate from the initial case (e.g., patient, environmental sample)?										
\square No \square Yes (If Yes, please refer to the guidance instructions at <u>www.selectagents.gov</u> for further directions.)										
11. Has the sender(s) (i.e., sample provider(s)) of the specimen(s) been notified of the identification of the select agent or toxin? INO Yes N/A NOTE: Please request completed and signed Sections C & D from each facility that was in possession of the specimen(s).										
12. Sample Provider Entity Name:	signed Sections C & D from	each facility that was	In possession of the spe	ecimen(s).						
			<u> </u>		1.5.0					
			le Provider E-mail Address:			15. Sample Provider Contact Number:				
First: MI:	Last:				INUTID					
16. Comments / Notes:										

I hereby certify that the information contained in Sections A and B of 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 331, 9 CFR 121, or 42 CFR 73 may result in civil or criminal penalties, including imprisonment.

Date Signed:



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Reference ID Number:

Submit completed form only once by either e-mail, fax, or mail									
	SECTION C -	SAMPLE PROVI	DER INFORMATIO	N					
1. Name of individual completing Sections First: MI:	2. [E-mail address:		3. Te	3. Telephone #:				
 4. Registered Entity (APHIS or CDC Clinical or Diagnostic Laboratory [i (NRE # (provided by APHIS or CDC))	9. Entity name:							
5. Responsible Official or Laboratory Supe First: MI:	10. Address (NOT a post office address):								
6. E-mail address:	7. Telephone #: 8	3. Fax #:	11. City: 12.		12. State:	13. Zip Code:			
SECTION D – SPECIMEN(S) CONTAINING SELECT AGENT OR TOXIN PROVIDED TO REFERENCE LABORATORY									
1. Select Agent or Toxin Identified:		2. Date notified of select agent or toxin identification:							
3. Case/patient/sample ID #(s):	4. # of samples shipped:	5. Sample type pro	ided: 6. Case/patient/san			nple origin (zip code):			
7. Date sample(s) shipped to Reference La	nce Laboratory:								
 9. Disposition of any remaining select agent or toxin by entity listed in Block C9: Destroyed (Provide destruction method and date. Method: Date:) Retained (Provide name of Principal Investigator retaining sample. Name:) Not applicable, the entire specimen was transferred to the Reference Laboratory. 10. Were any of the samples containing a select agent or toxin handled outside of primary containment which may have led to an unintentional release and/or exposure to the select agent or toxin? No									
	to the guidance instructions	at www.selectagents.	gov for further directions	.)		sample)?			
13. Has the sender(s) (i.e., sample provide NOTE: Please request completed and					No 🗆 Yes	6			
14. Sample Provider Entity Name:									
15: Sample Provider Point of Contact: First: MI:	Last:	16	. Sample Provider E-mai	I Address:	17. Sample Pr	ovider Contact Number:			
18. Comments / Notes:									

I hereby certify that the information contained in Sections C and D of 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 331, 9 CFR 121, or 42 CFR 73 may result in civil or criminal penalties, including imprisonment.

Signature of Responsible Official/Laboratory Supervisor:

Date Signed: _

Public reporting burden: Public reporting burden of providing this 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 D74, Atlanta, Georgia 30329; ATTN: PRA (0920-0576).