NCEH/ATSDR Human Subjects Research Determination Form

Use this form and the flowcharts for either:

1. CDC projects and activities that do not require CDC IRB review under HHS Human Subjects (45 CFR part 46) or FDA (21 CFR Parts 50 and 56) Regulations, which include "non research", "research not involving identifiable human subjects", or "human subject research for which CDC is not engaged"; OR

2. Human subjects research that will be submitted to the Human Research Protection Office (HRPO) as an Exempt Category of Human Subjects Research.

Project Title:	Customer Survey for CDC Laboratory Accredited to ISO 17025								
Project Loca	tion(s)/Site(s): CDC/NCEH/DLS 4770 Buford Highway, NE, Atlanta, GA 30341-3724								
Project Offic	er(s): Mary Halstead Telephone: (770) 488-7950 Di	ivision or Office:	DLS						
Proposed Pr	Dject Dates: Start: 9/6/2017 End: 8/31/2020 Ti	me sensitive:							
Project Fund	ing and Partners (answer both): HHS: CDC Funding - Internal Activity Non-HHS: NO	external funding							
If applicable, name participating external institution(s). CDC Funding - Internal Activity									
Indicate the holder of the key to decipher the identities of coded data or biological specimens. No key/ code/ identifiers									
Specify CDC role (mark II that apply):									
CDC is t	ne sole institution conducting activity; OR								
If not the sole	institution, indicate if:								
CDC is NOT a recipient or provider of private data, specimens, materials, or services;									
CDC is a provider of private data/specimens to an institution. X CDC is a recipient of private data/specimens from an institution.									
CDC is a	provider of materials/services to an institution. CDC is a recipient of materials/services from	n an institution.							
Questions 1	- 4 pertain to the HHS Human Subjects Regulations (45 CFR 46):								
1. For CDC:	Is this activity classified as research?	YES	NO X						
A. Is the a	ctivity a <u>systematic investigation</u> including research development, testing, and evaluation?	YES X	NO —						
	ctivity intentionally designed to develop OR contribute to generalizable knowledge?	YES	NO X						
	CDC activity IS research if both 1A and 1B are "YES."	_							
	If 1 is "NO," then STOP: otherwise continue.								
2. For CDC: Is this research classified as human subjects research? YES N									
A. Does the entities	YES	NO							
B. Does the activity only involve the collection or analysis of data or specimens from deceased persons? YES									
	CDC activity IS NOT human subjects research if either 2A or 2B are "YES."	_							
	If 2 is "NO," then STOP: otherwise continue.								
C. Do CD	C employees intervene with, interact with, or obtain informed consent from living persons?	YES	NO						
D. Are/We	YES	NO							
E. Are/We	YES	NO							
F. Do/Will CDC employees or agents have access to the link between the data or specimens and the identity of these living persons?									
	CDC activity IS human subjects research if 2C is "YES," CDC activity IS NOT human subjects research if 2D, 2E, and 2F are all "NO.".								
	If 2 is "NO," then STOP: otherwise continue.								
3. For CDC:	Will this activity be submitted to HRPO for approval or exempt human subjects research?	YES	NO						
A. Does t	he research pose more than minimal risk?	YES	NO						
B. Will pr	YES	NO							
C. Will in	YES	NO							
D. Based on the HRPO Worksheet for Exemption from Human Subjects Regulations, is there an HHS Exempt Research Category for which this activity will be reviewed? If "YES," specify the Category Number:									
	CDC activity IS exempt human subjects research if 3A, 3B, and 3C are all "NO," and an exempt category (3D) applies. Exempt research must go to HRPO: use CDC Form 0.1250X.								
	If 3 is "YES," then STOP; otherwise continue.								

4. Is CDC engaged in the non-exempt research involving identifiable human subjects?					YES	NO			
A. Did CDC receive funding directly from another HHS agency?					YES	NO			
B. Do CDC employees or agents intervene or interact with living individuals for research purposes?					YES	NO			
C. Do CDC er	ployees or agents obtain indivi	idually identifiable priv	rate information?		YES	NO			
		CDC IS engaged if 4A, 4	B, or 4C are "YES."						
If 4 is "NO," then STOP. Otherwise, research must go to HRPO; use CDC Form 0.1250.									
	ns to research involving FDA ct in the course of medical p		(21 CFR parts 50 and 56),	not including the u	ise of an FDA				
5. Based on the <u>HRPO Worksheet to Determine FDA Regulatory Coverage</u> , is the research activity subject to FDA human subjects regulations?						_ NO			
Additional Notes:									
Although CDC HRPO review is not required, investigators or project officers must adhere to ethical principles and standars to respect and protect the privacy, confidentiality, and autonomy of participants. All applicable State and Federal privacy Isws must be followed. Informed consent may be appropriate. Information disclosed in the consent process should address the basic elements of consent. The consent form and all other required supporting documents must be submitted with this form for review. The list of required documents is found in the MOEH/ATSDR Guided Checklist for Human Subjects and PRA Determinations .									
Division Approva	l Signatures and Dates:								
Ben Blount	9/12/2017	1:20 PM	Jerry Thomas	9/20/2017 1	0:49 AM	_			
Branch Chi	ef Date Sign	ed	Division ADS/Director	Date Signed	Ł				
Request Receive	d Date: 09/20/2017								
X CDC's role	loes not require HHS human	subjects review bey	ond the center level becau	se:					
X Activity	vity is not research (Flow chart category NR-1).								
Activity	Activity is not human subjects research (Flow chart coverage NR-2 through NR-8).								
Activity is non-exempt human subjects research, but CDC is not engaged (Flow chart category HSR-3).									
CDC's role does require HHS human subjects review beyond the center level because:									
Activity qualifies as exempt human subjects research (Flow category HSR-1).									
Activity qualifies as non-exempt engaged human subjects research (Flow category HSR-2).									
CDC's role does not require FDA human subjects review beyond the center level because:									
Activity does not require human subjects review under FDA regulations (Flow chart category NFDA-3 through NFDA-4).									
CDC's role	CDC's role does require FDA human subjects review beyond the center level because:								
Activity qualfies as human subjects research under FDA regulations (Flow chart category FDA-1 through FDA-2).									
NCEH/ATSDR Human Subjects Contact Signature and Date:									
	-	09/20/2017							
Stephanie	Davis, MSPH Date Sign	ed							

Guidance for Completing the NCEH/ATSDR Human Subjects Research Determination Form

For question 1:

- To determine if your project is research for purposes of human subjects protection, consult:
 - The CDC Policy on Distinguishing Public Health Research and Public Health Nonresearch
 - Guidance from the Office of Human Research Protections (OHRP)
 - The FDA regualtions, if applicable
- See the Research Determination Flowchart 1 for examples of nonresearch activities.

For question 2:

- Research involving living human subjects must adhere to the protection of human subjects under either the <u>Human Subjects 34 CFR part 46</u> or <u>FDA 21 CRF part 50</u> and <u>part 56</u>.
- Guidance on research involving coded private information or biological specimens is available from OHRP.
- More information on human subjects research can be found on the HRPO website.
- See the Research Determination Flowcharts 1-3.

For question 3:

- 45 CFR part 46(b) outlines the Exempt Research Categories.
- The <u>HRPO Worksheet for Exemption from Human Subjects Regulations</u> provides more details on Exempt Research Categories.
- The categories most often used for Exempt Research conducted at CDC/ATSDR are 2 and 4.
- See the Research Determination Flowchart 4.

For question 4:

- Guidance on Engagement of institutions in research can be found from OHRP.
- See the Research Determination Flowchart 4.

For question 5:

- Research involving living human subjects that are 21 CFR Part 50 and part 56
- See the Research Determination Flowchart 5 and the <u>HRPO Worksheet to Determine FDA Regulatory Coverage</u> for more *information on how to make this determination*.
- Differences between HHS and FDA human subjects regulations can be found here.

NOTE: If CDC is only providing/receiving materials and services, the Research Determination Flowcharts do not apply.