

REQUEST FOR DETERMINATION OF RESEARCH STATUS

To be completed by the staff member with lead responsibility for the project and approved by branch chief (if applicable) and Division ADS. A separate PGO funding memo is required if project is research and involves human subjects regardless of the CDC staff role.

Instructions:

- (1) Use this form to declare: (a) the research status of any project, (b) role or roles of CDC staff
- (2) A short summary should be attached offering specific details about the project and the role of staff.
- (3) Be sure to complete all applicable items, obtain appropriate signatures and submit this form for approval.

Tracking Number:

ate subm	nitted:	09/21/2015						
tle of Pr	oject:	The National Qu	 itline Data War	ehouse				
ates for r	project pe			Dates for fundi	ng (if	applicable):		
Beginni		09/21/2015			Beginning:			
Ending:	_	09/20/2020		Ending:				_
								_
•	(choose or			1 1	1	1 1	C	·
		is usea below, refers CDC staff member, d				e project incluaing scof	ре ој рга	eject, funding restrictions,
[] N	New				[]	Revision		
[X] C	Continuati	on, without revision	ı(s)		[]	Continuation, with	revision	(s)
ead staff	member:		Contact info	rmation:	Ple	ase indicate your role	e(s) in th	uis proiect:
Name:		aya McGruder	Division:	OSH	[]	Project officer	[X]	
		.,,	_		- []	Principal	[]	Investigator
User II): HDD8	3	_ Telephone:	770-488-8266	_	investigator		J
					Г 1	Consultant	[]	Other (please explain
Are an	ny or all o	[X] N	this project DE:	F79 SIGNED to cont	_ [] tribute	e to generalizable know		
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Form 684R_NR (revised January 2003)

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Name			Role (project officer, investigator, consultant, etc.)					Scientific ethics number Prin	
Henraya McGrude			er					12506	
				EARCH PRO					(as identified in 45CFR46.101),
4.			•	sed research			-		
	[]	YES			-	-		ed and must be reviewed b	y an IRB (skip to question 7).
	[]	NO		,			•		
5.				earch involve	fetuses, j	pregnant v	women, o	r human in vitro fertilizatio	on as targets (such that Subpart B
	[]	YES		If YES, th question 7		rch canno	ot be exe	mpted and must be revi	ewed by an IRB (skip to
	[]	NO							
Edi	ucation	al Resea	rch						
	6.1	Is this	research al educat	ional practice	es (e.g., re	esearch on	regular :	and special education strate	gs, AND does the research involve egies or research on the room management methods)?
		[]	YES	, 1	[]	NO		• /	,
Res	search l		g Surveys	s, Interview F		es (includi	ng Focus	groups), Observation of Pu	ıblic Behavior, or Educational
Tes	<u>sts</u>		<u>-</u>				_		<u> </u>
	6.2			rch use educa observation o			ive, diagn	ostic, aptitude, achievemen	at), survey procedures, interview
		[]	YES		[]	NO		If NO skip 6.3	
		Will children (<18 years of age) be research subjects?							
		[]	YES	If YES, th	is researc	ch cannot	be exemp	ted and must be reviewed l	by an IRB (skip to item 7)
		[]	NO						
		6.2.1						nanner that human subject ked to the subjects;	s can be identified <u>directly or</u>
			[]	YES		[]	NO		
		6.2.2	place the employ subject	ne subjects at ability or rep s' (or relative	risk of coutation? es' or asso	riminal or (Example ociates') po	civil liab s here ma ossible su	ility, or be damaging to the ry include: the collection of	h setting have the potential to e subjects' financial standing, sensitive data regarding the riminal history or intent, medical rmation).
			[]	YES		[]	NO		
	6.3	proce	dures, or		of public	behavior		search is not exempt under	nt), survey procedures, interview r paragraph 6.2 of this section:
		[]	YES		[]	NO		If NO skip to 6.4	
		6.3.1	Will thi		volve hu	man subje	cts that a	re elected or appointed pul	blic officials or candidates for
		[]	YES		[]	NO			
	6.3.2	inform	ation will be a on only in the	maintain	ed througl	hout the 1	ion that confidentiality of t esearch and thereafter? (N nce of Confidentiality has b	ote: CDC can use this exemption	
			[]	YES		[]	NO		
<u>Exi</u>	isting D	ata Whi	ch Is Pul	olicly Availab	<u>le or Uni</u>	dentifiabl	<u>e</u>		
	6.4							f existing* data, documents e the study begins)?	, records, pathological or
		[]	YES		[]	NO		If NO skip to 7	
		6.4.1	Is this 1	naterial or in	formatio	n publicly	availabl	2?	
		[]	YES		[]	NO			

	Tracking NC). No	Funding	
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6.4.2			nformation recorded in such a manner by the investigator that the subjects cannot be or indirectly through identifiers linked to the subjects?
	`		reated by an investigator even temporarily, for research purposes, this criterion is not met. is created by clinical staff who already have access to the data, this criterion is met).
	[]	YES	(there are no identifying information and no unique identifiers or codes)YES
	гэ	NO	(there are identifiers (including codes))

- Please prepare and attach a short summary paragraph (<1 page); if this is new:
 - a. Be sure to include the purpose of the project, specific details about the project and the role of the CDC staff member (s) in the project. In explaining one's role as a consultant be particularly careful to identify involvement in things like: study design decisions, oversight of protocol development, participation in review of data collection procedures, and participation in data analysis and/or manuscript preparation, as well as whether there will be access to identifiable or personal data.
 - b. Explain your project status selection (research--non-exempt, exempt, no CDC investigator or not involving human subjects; public health practice). If you selected research not involving human subjects be sure to indicate if the data includes any personal information (e.g., name, SSN), linkable study identification numbers or codes, or geographical information.

The purpose of this project is to create the National Quitline Data Warehouse (NQDW) to assist in the evaluation of telephone quitlines. The NQDW consolidates information into an integrated database for ongoing program evaluation and improvement of quitline services. The NQDW has OBM approval (0920-0856) for the collection this data. There are not personal identifiers on any of the data to be collected.

The NQDW will standardize data collected by state quitlines using three questionnaires (based on the North American Quitline Consortium's Minimum Data Set): an intake questionnaire, a 7-month follow-up questionnaire (to be used only with the Asian Smokers' Quitline), and a quitline services questionnaire. The intake and 7-month follow-up questionnaires are administered to quitline callers and collect data on tobacco use, intention to quit, success with quitting, and use of counseling and/or medications to facilitate or maintain quit. The quitline services questionnaire is administered to tobacco control managers and gathers the types of information that normally would be gathered from grantees in maintaining accountability regarding expenditure of government funds. No personal identifiable information is collected on any of the questionnaires.

All of the above data is already being collected by the states quitlines - the purpose of the current project is just to allow the states to send the data to CDC for the first time. CDC staff members will only use the data for evaluation purposes. No research is planned for the data. Only aggregate data will be reported at the states and national levels. There are no research activities; no activites involving human subjects in research.

Separate contracts to clean and report on the data have their own 684 forms (all of which have already been classified as for public health practice). This form is for the overall project management of the NQDW.

8. Please list the primary project site and all collaborating site(s).

Explanation of project components:

9. If project involves research that is funded extramurally, list amount of award that should be restricted pending IRB approval and describe which project components will be affected, if known:

Approvals (signature and position title)	Date	Research Determination / Remarks
Henraya McGruder - HEALTH SCIENTIST	09/21/2015	[X] Public health practice [] Research not involving human subjects [] Research involving human subjects, no CDC investigators [] Research involving human subjects, CDC investigators, exempt [] Research involving human subjects, CDC investigators, not exempt (check if applicable) [] Local IRB [] CDC Exemption [] CDC IRB
staff member completing this form		Comments:

Shanna Cox - EPIDEMIOLOGIST	09/21/2015	[X] Public health practice [] Research not involving human subjects [] Research involving human subjects, no CDC investigators [] Research involving human subjects, CDC investigators, exempt [] Research involving human subjects, CDC investigators, not exempt (check if applicable) [] Local IRB [] CDC Exemption [] CDC IRB
Team Lead		Comments:
Shanna Cox - EPIDEMIOLOGIST Division ADS	09/21/2015	[X] Public health practice [] Research not involving human subjects [] Research involving human subjects, no CDC investigators [] Research involving human subjects, CDC investigators, exempt [] Research involving human subjects, CDC investigators, not exempt (check if applicable) [] Local IRB [] CDC Exemption [] CDC IRB
Joan Redmond Leonard - PUBLIC HEALTH ANALYST	09/21/2015	 [X] Public health practice [] Research not involving human subjects [] Research involving human subjects, no CDC investigators [] Research involving human subjects, CDC investigators, exempt [] Research involving human subjects, CDC investigators, not exempt (check if applicable) [] Local IRB [] CDC Exemption [] CDC IRB
CUC ADS, Deputy ADS, or Human Subjects Contact		<u>Comments:</u>