CDC

Instructions:

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.

- (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:

(Use PGO number if cooperative agreement, grant, etc.)

Date submitted: Title of Project:		ed:	07/31/2014									
		ct:	Survey of NBCCEDP Grantee Program Implementation									
Date	es for pro	ject peri	od:]	Dates for	fundi	ing (if	applicable):				
B	eginning	:	08/04/2014	5/04/2014						_		
E	nding:		12/31/2018	Ending:					_			
	ject is (ch											
			used below, refers to DC staff member, det		0			project including scop	pe of pro	oject, funding restrictions,		
[X	[] New	7					[]	Revision				
[]	Con	tinuatio	n, without revision(s)			[]	Continuation, with	revision	(s)		
Lead	d staff me	ember:		Contact info	rmation:		Ple	ase indicate your role	e(s) in th	iis project:		
Ν	ame:	Amy De	egroff	Division:	DCPC		[]	Project officer	[X]	Technical monitor		
U	ser ID:	ASD1		Telephone:	770-488	-2415	[]	Principal investigator	[]	Investigator		
S	cientific	Ethics 1	number: 6818	Mailstop:	F76		[]	Consultant	[]	Other (please explain)		
	[] If YES,]	YES list those	[X] NO activities which are									
2.	Is this C	DC proj	ect research or publ	ic health prac	ctice (che	ck all	that a	pply)?				
	[]	Research			[X]	Pub	blic health practice					
		Check a	one:			Check all that apply:						
		[]	Human subjects inv	volved		[]	En	nergency Response	[]	Surveillance		
		[]	Human subjects not	t involved		[X]	Pr	ogram evaluation	[]	Other (please explain)		
3.	If RESE subjects			jects, has the	project o	or rese	earch	activities been review	ed by tl	ne CDC IRB for human		
	a. [] NO, New project, not yet reviewed					[] YES, Reviewed and approved by CDC						
	b. [] NO, Existing project, not ready to submit					If YES, please list protocol number_and						
	c. [] NO, Submitted for approval					expiration date						
					e.	[]	NO, requ	· · · · ·	C invest	tigators (CDC IRB not		
							requ	neu)				

If RESEARCH, list any other CDC staff involved in this project, please include the name, role, and scientific ethics number

Tracking NO. <u>No Funding</u>

Name				F C	Scientific ethics number Prin			
A	my Deg	roff						6818
			EARCH PRO					H (as identified in 45CFR46.101),
	Does	the propos	sed research	involve p	risoners?	_		
[]	YES		If YES, thi	is research	cannot b	e exempt	ted and must be reviewed	by an IRB (skip to question 7).
[]	NO							
	the prop l apply):		arch involve	fetuses, p	regnant v	vomen, o	r human in vitro fertilizat	ion as targets (such that Subpart
[]	YES		If YES, th question '		ch canno	ot be exe	empted and must be rev	iewed by an IRB (skip to
[]	NO							
ducation	al Resea	urch						
6.1	norma	al educatio	onal practic	es (e.g., res	earch on	regular a	and special education stra	ngs, AND does the research involv tegies or research on the ssroom management methods)?
	[]	YES		[]	NO			
	[nvolvin	g Surveys.	, Interview l	Procedures	(includi	ng Focus	groups), Observation of P	Public Behavior, or Educational
<u>ests</u> 6.2			ch use educa bservation o			ive, diagn	ostic, aptitude, achieveme	ent), survey procedures, interview
	[]	YES		[]	NO		If NO skip 6.3	
		hildren (<	18 years of		earch su	bjects?	Ĩ	
	[]	YES	•	0 /		0	ted and must be reviewed	by an IRB (skip to item 7)
	[]	NO				···· ·		
	6.2.1						nanner that human subjec nked to the subjects;	cts can be identified <u>directly or</u>
		[]	YES		[]	NO		
	6.2.2	place the employa subjects	e subjects at ability or rep ' (or relative	t risk of cri outation? (es' or assoc	minal or Example iates') po	civil liab s here ma ossible su	ility, or be damaging to that a second se	ch setting have the potential to ne subjects' financial standing, of sensitive data regarding the criminal history or intent, medica formation).
		[]	YES		[]	NO		
6.3								ent), survey procedures, interview er paragraph 6.2 of this section:
	[]	YES		[]	NO		If NO skip to 6.4	
	6.3.1	public o	ffice?	volve hum	-		re elected or appointed pu	ublic officials or candidates for
		[]	YES		[]	NO		
	6.3.2	informa	tion will be 1 only in the	maintaineo	l througl	hout the r	esearch and thereafter? (the personally identifiable Note: CDC can use this exemptio been obtained to cover the
		[]	YES		[]	NO		
<u>xisting D</u>			licly Availal					
6.4	diagn	ostic speci		xisting' me	ans exist		e the study begins)?	ts, records, pathological or
	[]	YES		[]	NO		If NO skip to 7	
	6.4.1	Is this m	naterial or in	nformation	publicly	available	e?	
		[]	YES		[]	NO		

Form 684R_NR (revised January 2003)

[]

6.4.2 Is this material or information recorded in such a manner by the investigator that the subjects cannot be identified directly or indirectly through identifiers linked to the subjects?

(Note: If a link is created by an investigator even temporarily, for research purposes, this criterion is not met. If a temporary link 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
 - NO (there are identifiers (including codes))
- 7. 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.

This project, the Survey of NBCCEDP Grantee Program Implementation, will collect information for years 2-5 (DP12-1205) from the 67 grantees of the National Breast and Cervical Cancer Early Detection Program (NBCCEDP), on their program implementation efforts. Staff in the Division of Cancer Prevention and Control, Program Services Branch, Program Evaluation Team will lead the project.

This project is determined to represent public health practice, program evaluation. The survey will collect information about program activities conducted (e.g., evidence based interventions), clinical service provider network, client eligibility criteria, partnerships, evaluation activities, training and technical assistance needs, and program management. Names of respondents will not be collected. Program name (e.g., Ohio NBCCEDP) will be collected and maintained. An existing contractor, Information Management Services, Inc., who currently manages the clinical data collected for the NBCCEDP, will field the survey via a web-based tool and provide the resulting data to CDC.

CDC staff have developed and piloted the survey instrument and will conduct all analysis including producing grantee-level reports for use by CDC program consultants in their technical assistance efforts. We will seek OMB approval for this survey.

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		
Amy Degroff - HEALTH EDUCATION SPECIALIST	07/31/2014	 [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 [] Research involving human subjects, CDC investigators, not exempt [] Local IRB [] CDC Exemption [] CDC IRB 		
staff member completing this form		<u>Comments:</u>		

Faye Wong - SUPV PUBLIC HEALTH ADVISOR Team Lead	07/31/2014	 [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 [] Research involving human subjects, CDC investigators, not exempt [] Local IRB [] CDC Exemption [] CDC IRB
Cheryll Thomas - EPIDEMIOLOGIST Division ADS	08/04/2014	 [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 CUC ADS, Deputy ADS, or Human Subjects Contact	08/05/2014	 [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