

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:

BPA (OADC)/2016-016

Date submitted:	03/11/2016				_			
Title of Project:		 Set the Fact	ts Ahout Gyne	coloa	ic Cancer Campaign			
Dates for project pe			Dates for fund			I		
Beginning:	09/02/2016	,	Beginning:	ing (n	аррисаме).			
Ending:	09/01/2021		Ending:				_	
Ending.	09/01/2021		Litting.					
Project is (choose on	ne):							
	is used below, refers to a CDC staff member, deter				project including scop	pe of pro	oject, funding	restrictions,
[] New				[]	Revision			
[X] Continuati	on, without revision(s)			[]	Continuation, with revision(s)			
Lead staff member:	(Contact info	rmation:	Ple	ase indicate your role	e(s) in th	nis project:	
**	ia Gelb	Division:	DCPC	[]	Project officer	[X]		monitor
				[]	Principal	[]	Investigat	or
User ID: CMG7		-	770-488-4708	– .,	investigator			
Scientific Ethics	s number: 4187	Mailstop:	MS: F76	_ []	Consultant	[X]	Other (ple	ease explain)
							_	
[] YES	f the activities within this [X] NO		SIGNED to con	tribute	to generalizable know	CO vledge (i		?
[] YES If YES, list those	[X] NO se activities which are r	esearch:						?
[] YES If YES, list those	[X] NO se activities which are r	esearch:	etice (check all	that a	pply)?			?
If YES, list those	[X] NO se activities which are r oject research or public	esearch:	ctice (check all	that a				?
[] YES If YES, list those 2. Is this CDC pro [] Resea	[X] NO se activities which are r oject research or public	esearch: health prac	ctice (check all	that a lic hea	pply)?			
If YES, list those Is this CDC pro Resea Check	[X] NO se activities which are r oject research or public arch & one:	esearch: health prac	etice (check all [X] Publ	that a lic hea ck all i	pply)? lth practice that apply:	vledge (i	i.e., research)	e
If YES, list those Is this CDC pro Resea Check	[X] NO se activities which are r oject research or public arch to one: Human subjects invo	esearch: health prac	etice (check all [X] Pub Check []	that a lic hea ck all i	pply)? alth practice that apply: nergency Response	vledge (i	i.e., research)	e se explain) Education and
[] YES If YES, list those 2. Is this CDC pro [] Resea Check [] []	[X] NO se activities which are r oject research or public arch to one: Human subjects invo Human subjects not i	esearch: health prac	etice (check all [X] Pub Chec []	that a lic hea ck all i En	pply)? olth practice that apply: nergency Response ogram evaluation	rledge (i	Surveillance Other (plea	e se explain) Education and awareness
I YES If YES, list those 2. Is this CDC pro [] Resea Check [] [] 3. If RESEARCH subjects protect	[X] NO se activities which are r oject research or public arch to one: Human subjects invo Human subjects not i	health practived involved	etice (check all [X] Pub Chec []	that a lic hea ck all t En Pr	pply)? olth practice that apply: nergency Response ogram evaluation	[] [X]	Surveillance Other (plea	e se explain) Education and awareness
[] YES If YES, list those 2. Is this CDC pro [] Resea Check [] [] 3. If RESEARCH subjects protect a. [] NO, Ne	[X] NO se activities which are r oject research or public arch to one: Human subjects invo Human subjects not i	health practived involved	ctice (check all [X] Pub Chec [] []	that a lic hea ck all i En Pr earch	pply)? alth practice that apply: nergency Response ogram evaluation activities been review	[] [X] red by the oved by	Surveillance Other (plea	e se explain) Education and awareness
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[] YES If YES, list those 2. Is this CDC pro [] Resea Check [] [] 3. If RESEARCH subjects protect a. [] NO, Ne b. [] NO, Ex	[X] NO se activities which are r oject research or public arch to one: Human subjects invo Human subjects not i	health practived involved	ctice (check all [X] Pub Chec [] []	that a lic hea ck all t En Pr earch YES I	pply)? olth practice what apply: nergency Response ogram evaluation activities been review , Reviewed and appro	[] [X] red by the oved by tocol nu	Surveillance Other (plea	e se explain) Education and awareness for human

Form 684R_NR (revised January 2003)

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Name				Ro coi	Scientific ethics number Prin					
	Cy	nthia G	elb						4187	
				ARCH PROJE s 4-6, OTHER					CH (as identified in 45CFR46.101),	
4.			_	ed research inv						
	[]	YES		If YES, this re	search ca	annot be	exempte	d and must be reviewe	d by an IRB (skip to question 7).	
	[]	NO								
5.		the prop l apply)?		rch involve fet	uses, pre	gnant wo	omen, or	human in vitro fertiliz	ation as targets (such that Subpart B	
	[]	YES		If YES, this question 7).	research	cannot	be exer	npted and must be re	eviewed by an IRB (skip to	
	[]	NO								
Ed	ucationa	al Resea	<u>rch</u>							
	6.1	norma	d educatio	nal practices (e	e.g., resea	arch on r	egular a	nd special education st	ings, AND does the research involve rategies or research on the assroom management methods)?	
		[]	YES	_	_	NO				
Re	search I						g Focus g	roups), Observation of	Public Behavior, or Educational	
Te	<u>sts</u>							- - -	·	
	6.2			h use education bservation of p			e, diagno	stic, aptitude, achieven	nent), survey procedures, interview	
		[]	YES		[]	NO		If NO skip 6.3		
		Will children (<18 years of age) be research subjects?								
		[]	YES	If YES, this r	esearch c	annot be	e exempt	ed and must be reviewe	ed by an IRB (skip to item 7)	
		[]	NO							
		6.2.1						anner that human subj ked to the subjects;	ects can be identified <u>directly or</u>	
			[]	YES		[]	NO			
		6.2.2	place the employal subjects' or psych	e subjects at ris bility or reputa (or relatives' o ological conditi	k of crim tion? (Ex or associa ion, finan	ninal or c examples ntes') pos ncial stat	ivil liabil here may sible sub us, or sin	lity, or be damaging to y include: the collection	arch setting have the potential to the subjects' financial standing, a of sensitive data regarding the criminal history or intent, medical information).	
	6.2	VX 7211 41	[]	YES		[]	NO	estic antituda achie	nent), survey procedures, interview	
6.3									der paragraph 6.2 of this section:	
		[]	YES		[]	NO		If NO skip to 6.4		
		6.3.1	Will this public of		ve humai	n subject	ts that ar	e elected or appointed	public officials or candidates for	
			[]	YES		[]	NO			
		6.3.2	informat	ion will be mai only in the cas	ntained t	througho	out the re	search and thereafter?	of the personally identifiable (Note: CDC can use this exemption is been obtained to cover the	
			[]	YES		[]	NO			
Ex	isting D	ata Whi	ch Is Publi	icly Available o	or Unider	<u>ıtifiable</u>				
	6.4							existing* data, docume the study begins)?	nts, records, pathological or	
		[]	YES		[]	NO		If NO skip to 7		
		6.4.1	Is this m	aterial or infor	mation p	ublicly a	vailable	?		
			[]	YES		[]	NO			

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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))				

- 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.

CDC promotes awareness of gynecologic cancer through its Inside Knowledge: Get the Facts About Gynecologic Cancer (Inside Knowledge) campaign. In collaboration with the Department of Health and Human Services' Office on Women's Health, CDC established Inside Knowledge to educate women and health care providers about the signs, symptoms, risk factors, and prevention strategies related to the five main types of gynecologic cancer. When gynecologic cancers are found early, treatment can be most effective. Inside Knowledge supports the Gynecologic Cancer Education and Awareness Act of 2005, or Johanna's Law.

The purpose of this project is to inform and improve the content, messaging, and communication strategies of the Inside Knowledge campaign. Messages and materials about the signs, symptoms, and risk factors related to gynecological cancers will be reviewed and revised through information shared in focus groups by women aged 35-65, the primary audience of Inside Knowledge. Focus groups would be conducted in 3-4 cities in the United States, with at least 3 English groups and 1-2 Spanish groups held in each city. Up to 9 participants may be included in each focus group and will reflect demographic diversity to the extent necessary to ensure a diverse perspective. We will also ask participants to review consumer materials and creative approaches to assess whether they are clear, compelling, and engaging. The feedback provided by these consumers will provide information to campaign planners that will guide further development and dissemination of campaign materials and messages.

The CDC project manager will lead the development of a recruitment plan, moderator's guide, and other testing tools, and the communications contractor will conduct focus groups. All activities shall be upon the direction and approval of the Contracting Officer Representative.

This activity is public health practice because the primary purpose is to inform and improve an existing public health awareness campaign. The findings are not generalizable beyond the specific content of the Inside Knowledge campaign.

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
Cynthia Gelb - Health Communications Specialist	03/11/2016	[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: Thank you for reviewing this!

Janine Cory - Lead Health Communications Speci	03/14/2016	[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:
Cheryll Thomas - Epidemiologist Division ADS	03/17/2016	[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	03/17/2016	 [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>