

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:

XXXX-XXX/2014-C-003

Date submitte							
Title of Proje	ct: Young Wome	n Breast Cancer I	Education and	Awar	eness Campaign		
Dates for pro	ject period:]	Dates for fundi	ng (if	applicable):		
Beginning:	09/30/2014		Beginning:				
Ending: 09/29/2016			Ending:				_
Project is (che	oose one):						
	vision, as used below, refe role of CDC staff member				project including scop	pe of pro	oject, funding restrictions,
[X] New	7			[]	Revision		
[] Con	tinuation, without revis	ion(s)		[]	Continuation, with	revision	$\mathbf{u}(\mathbf{s})$
Lead staff me	ember:	Contact info	rmation:	Ple	ase indicate your role	e(s) in th	nis proiect:
Name:	Temeika Fairley	Division:	DCPC	[X]	Project officer	[X]	
User ID:	TFF9	Telephone:	770-488-4518	[]	Principal investigator	[]	Investigator
Scientific	Ethics number: 8	B817 Mailstop:	F76	[]	Consultant	[]	Other (please explain)
[]	or all of the activities with YES [X]	NO	SIGNED to cont	ribute	to generalizable know	vledge (i	.e., research)?
[] If YES, I		NO h are research:				vledge (i	.e., research)?
[] If YES, I	YES [X] list those activities which	NO h are research:	etice (check all	that a		vledge (i	.e., research)?
If YES, I	YES [X] list those activities which DC project research or	NO h are research:	etice (check all	that a	pply)?	vledge (i	.e., research)?
If YES, I	YES [X] list those activities which DC project research or Research	NO h are research: public health prac	etice (check all	that a ic hea k all t	pply)? lth practice	vledge (i	.e., research)? Surveillance
If YES, I	YES [X] list those activities which DC project research or Research Check one:	NO h are research: public health practs involved	etice (check all a [X] Publ Chec	that a ic hea k all t	pply)? lth practice hat apply:		
If YES, I	YES [X] list those activities which DC project research or Research Check one: [] Human subject	NO h are research: public health practs involved	etice (check all [X] Publ Chec	that a ic hea k all t	pply)? lth practice hat apply: nergency Response	[]	Surveillance Other (please explain) The proposed work is part of a new congressionally mandated health
If YES, 1 2. Is this C	YES [X] list those activities which DC project research or Research Check one: [] Human subject	NO h are research: public health pract ts involved ts not involved	etice (check all [X] Publ Chec []	that a ic hea <i>k all t</i> En Pr	pply)? Ith practice hat apply: nergency Response ogram evaluation	[] [X]	Surveillance Other (please explain) The proposed work is part of a new congressionally mandated health communications campaign.
If YES, 1 2. Is this C	YES [X] list those activities which DC project research or Research Check one: [] Human subject [] Human subject	NO h are research: public health practs involved ts not involved in subjects, has the	etice (check all [X] Publ Chec []	that a ic hea ek all t En Pr	pply)? Ith practice hat apply: nergency Response ogram evaluation	[] [X]	Surveillance Other (please explain) The proposed work is part of a new congression nally mandated health communications campaign. The CDC IRB for human

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	c. []	NO, S	ubmitted i	for approval					expiration date	
							e.	[]	NO, RESEARCH, no CE required)	OC investigators (CDC IRB not
							f.	[]	N/A (Not Applicable)	
	If RES	SEARC	H, list any	other CDC st	aff invol	ved in th	is p	rojec	t, please include the name, I	role, and scientific ethics number
	Na	ıme				ole (pro onsultai			cer, investigator,	Scientific ethics number Prin
	Te	meika F	airley							8817
IF	VOII TI	HINK T	HE DESE	A R C H PRO I	ECT MI	сит ОІ	TAT	IFV	AS EXEMPT DESEADCH	(as identified in 45CFR46.101),
				s 4-6, OTHER						(as identified in 45 cf R40.101),
4.		Does t	he propos	ed research in	volve pr	isoners?	•			
	[]	YES		If YES, this r	esearch (cannot b	oe ex	empt	ed and must be reviewed by	y an IRB (skip to question 7).
	[]	NO								
5.		the prop apply)?		rch involve fe	tuses, pr	egnant v	vom	en, oi	human in vitro fertilizatio	n as targets (such that Subpart B
	[]	YES		If YES, this question 7).	researc	h canno	ot b	e exe	mpted and must be revie	wed by an IRB (skip to
	[]	NO								
Ed	ucationa	al Resea	rch							
	6.1	norma	l educatio	nal practices (e.g., rese	arch on	reg	ular a	and special education strate	s, AND does the research involve gies or research on the room management methods)?
		[]	YES	-	[]	NO			1 /	,
		nvolving	g Surveys,	Interview Pro	cedures	(includi	ng F	ocus	groups), Observation of Pu	blic Behavior, or Educational
Te		******			•					
	6.2			h use educatio servation of p			ive,	diagn	ostic, aptitude, achievemen	t), survey procedures, interview
		[]	YES		[]	NO			If NO skip 6.3	
		Will c	hildren (<	18 years of ago	e) be rese	earch su	bjec	ts?		
		[]	YES	If YES, this	research	cannot l	be e	xemp	ted and must be reviewed b	y an IRB (skip to item 7)
		[]	NO							
		6.2.1							nanner that human subjects ked to the subjects;	can be identified <u>directly or</u>
			[]	YES		[]	N	Ю		
		6.2.2	place the employal subjects'	subjects at ri bility or reput (or relatives'	sk of crii ation? (F or associ	ninal or Example ates') po	civi s he ossib	il liab re ma ole su	ility, or be damaging to the y include: the collection of	setting have the potential to subjects' financial standing, sensitive data regarding the iminal history or intent, medical rmation).
			[]	YES		[]	N	10		
	6.3									t), survey procedures, interview paragraph 6.2 of this section:
		[]	YES		[]	NO			If NO skip to 6.4	
		6.3.1	public of	fice?	lve huma	ŭ			re elected or appointed pub	lic officials or candidates for
			[]	YES		[]		10		
		6.3.2	informat	ion will be ma only in the ca	intained	through	hout	the r	ion that confidentiality of the esearch and thereafter? (Nonce of Confidentiality has be	ote: CDC can use this exemption
			[]	YES		[]	N	Ю		

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<u>Exi</u>	sting Da	ata Whi	ch Is Pub	olicly Available	<u>e or Unid</u>	<u>lentifiab</u> l	<u>le</u>				
	6.4	diagno	ostic speci		isting' me	eans existi		re the study beg	gins)?	ecords, pathologica	al or
		[]	YES		[]	NO		If NO skip to	.o 7		
		6.4.1		naterial or info	ormation	_	-	ie?			
			[]	YES		[]	NO	• /	• • •	• • • •	_
		6.4.2	identifie	ed directly or i	indirectly	ly through	gh identifie	iers linked to the	ne subjects?	that the subjects ca	
			If a temp	porary link is	s created l	by clinica	cal staff wl	vho already have	ve access to the da	rposes, this criterion	is met).
			[]	YES				_	_	lentifiers or codes)	YES
			[]	NO				ncluding codes)))		
7.	if this i	s is new:		ach a short sun					1.41	- cpc	-
	a.	(s) in the like: stand pa	the project study designartic	ct. In explainin gn decisions, o	ng one's r oversight lysis and/	role as a o	consultan	nt be particular lopment, partici	rly careful to ide cipation in review	e role of the CDC s entify involvement w of data collection there will be access	t in things n procedures,
	b.	subject include	cts; public	c health practio	ice). If yo	ou selecte	ed researc	ch not involving	g human subjects	gator or not involvits be sure to indica mbers or codes, or a	ate if the data
	The CDC has been directed to develop and implement an education and awareness social media campaign focusing on topics related breast cancer in young women. This campaign has multiple components that will address specific aspects of the Education and Awareness Requires Learning Young Act (EARLY Act) legislation. This campaign will educate young women and medical providers about breast health, breast cancer risk factors (including but not limited to BRCA gene mutation), and survivorship among young women. The audiences for this campaign include women ages 15-44 (young women) at higher than average risk for developing breast cancer such as Ashkenazi Jewish and African-American females; young women who are breast cancer survivors; young women of all racial, ethnic, and cultural backgrounds; and medical providers. The purpose of this project is to develop, implement, and launch this social media health communications campaign. The overall activities include: 1) develop a communications and marketing plan; 2) purchase, place, and leverage media; 3) develop campaign messages and communication products; 4) monitor, report, and evaluate campaign activities; 4) develop and implement interactive/digital media strategies, materials, tools, and resources; 5) assist with media relations; 6) assist with graphic design, photos, and layout of campaign materials; 7) assist with writing and editing; 8) provide translation and adaptation services and support; 9) provide training, including presentations related to the campaign; and 10) provide data rights and transfer of campaign materials at the end of the contract.							ts of the oung women utation), and) at higher than			
								edia; 3) ; 4) develop ;; 6) assist with n and			
			DC staff me		/e as the f	technical	monitor ar	nd subject matte	er expert on this so	social media health	
		commu social n	unications media heal	campaign. Throalth communicat	rough this ations cam	s contract, mpaign.	, CDC seel	eks a contractor th		nch this social media le the specialized ser	
8.	Please	: list the	primary 1	project site an	ıd all coll	laboratir	ng site(s).	,			
	Explai	nation o	of project	components:							
9.				earch that is fui which project					that should be r	restricted pending	IRB

Approvals (signature and position title)	Date	Research Determination / Remarks
Temeika Fairley - EPIDEMIOLOGIST	03/17/2014	[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
staff member completing this form		Comments:
Jameka Blackmon - Assoc Dir for Program Devel	03/19/2014	[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
Team Lead		Comments:
Cheryll Thomas - EPIDEMIOLOGIST	03/19/2014	[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
Division ADS		Comments:
Joan Redmond Leonard - PUBLIC HEALTH ANALYST	3/26/2014	[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
CUC ADS, Deputy ADS, or Human Subjects Contact		Comments: