

## 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.

							cacking Number:		e agreement, grant, etc.)
						(0.	er do number n coo	perutiv	e agreement, grant, etc.)
Date submitt		11/20/2018							
Title of Proje	ect:	Evaluation of CDC	c's National T	obacco E	ducation	า C	ampaign		
Oates for pro	oject peri	od:		Dates for	funding	(if a	applicable):		
Beginning	<b>;:</b>	08/30/2019		Beginn	ing:				<del></del>
<b>Ending:</b>		08/29/2022		Ending	; <b>:</b>				. <u></u>
Project is (ch	100se one	):							
NOTE: Rev	vision, as						project including scop	pe of pr	oject, funding restrictions,
[] Nev	W				[]		Revision		
[] Cor	ntinuatio	n, without revision(s	s)			]	Continuation, with revis		n(s)
_ead staff m	ember:		Contact info	rmation:		Plea	se indicate your role	e(s) in t	his project:
Name:		Boyle-Estheimer	Division:	OSH		]	Project officer	[]	- "
		.,	•			X]	Principal	[]	Investigator
User ID:	YJW7		Telephone:	404-498-	2283		investigator		J
Scientific	Ethics	number: 18583	Mailstop:	F-79	[	]	Consultant	[]	Other (please explain)
		e activities which are surveys to an online p		nine the imp	oact of a	natio	onal tobacco education	n campa	aign.
. Is this C	CDC proj	ect research or publ	ic health pra	ctice (chec	k all tha	t ap	oply)?		
[X]	Resear	ch		[X]	Public l	ıeal	th practice		
	Check	one:			Check a	ll ti	hat apply:		
	[X]	Human subjects in	volved		[]	Em	ergency Response	[]	Surveillance
	[]	Human subjects no	t involved		[X]	Pro	gram evaluation	[]	Other (please explain)
	EARCH i s protecti		jects, has the	e project o	r researc	ch a	ectivities been review	ed by t	the CDC IRB for human
<b>a.</b> [X	NO, New	project, not yet rev	riewed	d.	[] <b>Y</b> ]	ES,	Reviewed and appro	oved by	y CDC
<b>b.</b> [ ]	NO, Exis	sting project, not rea	dy to submit			If	YES, please list pro	tocol n	umber_and
<b>c.</b> []	NO, Sub	mitted for approval					expiration date		
				e.			RESEARCH, no CD( red)	C inves	stigators (CDC IRB not
				f.	[] <b>N</b>	/A (	Not Applicable)		
If RESE	EARCH,	list any other CDC	staff involved	in this pro	oject, ple	ease	include the name, r	ole, and	d scientific ethics number

33413

Tracking NO. TBD/TBD	
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	Name				ole (proj onsultant	ect officer, investigator, , etc.)	Scientific ethics number Prin				
	Dia	Diane Beistle  Jane Mitchko				roject off	icer	8120			
	Jan					Project officer					
	La	uren Bo	yle-Esth	eimer	P	rincipal I	nvestigator	18583			
	Re	becca N	Murphy		P	Project officer					
	Ro	bert Ro	odes		P	roject off	icer				
				EARCH PROJ ns 4-6, OTHER			ALIFY AS EXEMPT RESEARCH (	as identified in 45CFR46.101),			
4.		Does	the propos	sed research in	volve pr	isoners?					
	[]	YES		If YES, this r	esearch	cannot be	exempted and must be reviewed by	an IRB (skip to question 7).			
	[ X ]	NO									
5.		he prop apply):		arch involve fe	tuses, pr	egnant wo	omen, or human in vitro fertilization	as targets (such that Subpart B			
	[]	YES		If YES, this question 7).	researc	h cannot	be exempted and must be review	wed by an IRB (skip to			
	[ X ]	NO									
Edu	ucationa			1 4 . 1	-4 - 1-1° -1-			AND less the second translation			
	6.1	norma	al educatio	onal practices (	e.g., rese	earch on r	, AND does the research involve gies or research on the boom management methods)?				
		[]	YES		[X]	NO					
		nvolvin	g Surveys.	, Interview Pro	cedures	(includin	g Focus groups), Observation of Pub	olic Behavior, or Educational			
Tes											
	6.2			ch use educatio bservation of p			e, diagnostic, aptitude, achievement	), survey procedures, interview			
		[X]	YES		[]	NO	If NO skip 6.3				
		Will c	hildren (<	18 years of ago	e) be rese	earch sub	jects?				
		[]	YES	If YES, this	research	cannot be	e exempted and must be reviewed by	an IRB (skip to item 7)			
		[ X ]	NO								
		6.2.1					such a manner that human subjects code) linked to the subjects;	can be identified <u>directly or</u>			
			[]	YES		[ X ]	NO				
		6.2.2	place the employa subjects	e subjects at ri ability or reput ' (or relatives'	sk of crii ation? (I or associ	minal or c Examples iates') pos	s' responses outside of the research ivil liability, or be damaging to the s here may include: the collection of s sible substance abuse, sexuality, cri us, or similarly compromising inform	subjects' financial standing, ensitive data regarding the ninal history or intent, medical			
			[]	YES		[ X ]	NO				
	6.3						e, diagnostic, aptitude, achievement ut the research is not exempt under				
		[]	YES		[X]	NO	If NO skip to 6.4				
		6.3.1	Will this public o		lve hum	an subject	ts that are elected or appointed publ	ic officials or candidates for			
			[]	YES		[]	NO				
		6.3.2	informa	tion will be man n only in the ca	intained	througho	t exception that confidentiality of tho out the research and thereafter? (No Assurance of Confidentiality has be	te: CDC can use this exemption			
			[]	YES		[]	NO				

<b>Existing Data</b>	a Which I	s Publicly A	Available (	or Unider	itifiable

	[]	YES		[X]	NO	If NO skip to 7		
	6.4.1	Is this material or information publicly available?						
		[]	YES		[]	NO		
	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?						
		iuciitiii	ed directly o			· ·		
		(Note: I	f a link is cı	reated by ar	0	or even temporarily, for research purposes, this criterion is not taff who already have access to the data, this criterion is met).	met.	
		(Note: I	f a link is cı	reated by and is created by	by clinical s	or even temporarily, for research purposes, this criterion is not	met.	
		(Note: I If a tem	f a link is cr porary link	reated by an is created l	by clinical s are no iden	or even temporarily, for research purposes, this criterion is not taff who already have access to the data, this criterion is met).		

- 'lease prepare and attach a short summary paragraph (<1 page); if this is new:
  - 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.

To conduct an evaluation of a paid, national tobacco education campaign. These ads are designed to motivate adult smokers to quit and nonsmokers to encourage family/friends who smoke to quit. The media buy will include placements of ads on television, radio, out-of-home, print, and digital platforms, and will ensure a baseline level of population-wide exposure to tobacco control measures. The evaluation will determine if the goals and objectives of the media campaign were achieved. Any personal information collected will be done by the Contractor, kept with the Contractor, and will not be transferred to CDC/OSH. Information collected by the Contractor will be delivered to CDC/OSH from the Contractor in aggregate form only.

While this is an evaluation of a campaign, certain aspects of the evaluation activity involves collection of data from a panel that is designed to yield nationally representative data. No personally identifiable or linkable information is however being collected, and the study poses no more than minimal risk.

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

**Explanation of project components:** 

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
Lauren Boyle-Estheimer - Scientist	01/14/2019	<ul> <li>[X] Public health practice</li> <li>[] Research not involving human subjects</li> <li>[] Research involving human subjects, no CDC investigators</li> <li>[X] Research involving human subjects, CDC investigators, exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>(check if applicable)</li> <li>[] Local IRB</li> <li>[X] CDC Exemption</li> <li>[] CDC IRB</li> </ul>
staff member completing this form		<u>Comments:</u>

Diane Beistle - LEAD HEALTH COMMUNICATIONS SPEC.  Team Lead	01/15/2019	[X] Public health practice [] Research not involving human subjects [] Research involving human subjects, no CDC investigators [X] Research involving human subjects, CDC investigators, exempt [] Research involving human subjects, CDC investigators, not exempt (check if applicable) [] Local IRB [X] CDC Exemption [] CDC IRB  Comments:
Israel Terungwa Agaku - Senior Service Fellow	01/16/2019	<ul> <li>[X] Public health practice</li> <li>[] Research not involving human subjects</li> <li>[] Research involving human subjects, no CDC investigators</li> <li>[X] Research involving human subjects, CDC investigators, exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>(check if applicable)</li> <li>[] Local IRB</li> <li>[X] CDC Exemption</li> <li>[] CDC IRB</li> </ul>
Division ADS		Comments:
Rachel Kaufmann - Assoc Director Science	01/16/2019	<ul> <li>[X] Public health practice</li> <li>[] Research not involving human subjects</li> <li>[] Research involving human subjects, no CDC investigators</li> <li>[X] Research involving human subjects, CDC investigators, exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>(check if applicable)</li> <li>[] Local IRB</li> <li>[X] CDC Exemption</li> <li>[] CDC IRB</li> </ul>
CUC ADS, Deputy ADS, or Human Subjects Contact		Comments: