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 sub	omitte	ed:	06/21/2016									
Title of Project:		ct:	Learn The Signs. Act Early. Bulk Order Customer Feedback Survey									
Dates for	r proj	ject peri	od:		Dates for fun	ding (if	applicable):					
Beginning: 07/01/2016					Beginning	:						
Endir	ıg:		06/29/2019		Ending:							
	E: Rev	ision, as					project including scop	pe of pro	oject, funding re	strictions,		
[X]	New					[]	Revision					
[]	Con	tinuatio	n, without revision(s)		[]	Continuation, with	revision	n(s)			
Lead sta	aff me	mber:		Contact info	rmation:	Ple	ase indicate your role	e(s) in th	nis proiect:			
Name			pher Pleasants	Division:	DCDD	[X]	Project officer	[]	Technical m	onitor		
User		YLJO		Telephone:		[X]	Principal investigator	[]	Investigator			
Scien	tific	Ethics 1	number: 14560	Mailstop:	E-86	[]	Consultant	[]	Other (pleas	se explain)		
			[X] NO	research:								
	his C		ect research or publ	ic health prac								
IJ	[] Research			[X] Public health practice								
		Check of		alvad			hat apply:	E 1	Surveillance			
		[]	Human subjects inv Human subjects not		[] [X]		nergency Response	[] [X]	Other (please	ovnloin)		
		IJ	fruman subjects not	l mvorveu	[23]	, 11		[24]	Other (prease	Customer Feedback		
		ARCH i protecti		jects, has the	project or re	esearch	activities been review	red by tl	he CDC IRB fo	r human		
a.	[]]	NO, New	v project, not yet rev	iewed	d. []	YES	, Reviewed and appro	oved by	CDC			
			sting project, not rea	dy to submit		Ι	f YES, please list pro	tocol nu	umber_and			
c.	[]]	NO, Sub	mitted for approval				expiration date					
					e. []	NO	RESEARCH, no CD					
						requ		C invest	tigators (CDC .	ікв поі		

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

Name Christopher Pleasants					ole (pro onsultan		Scientific ethics number Prin	
					rincipal I	nvestiga	14560	
			EARCH PROJI ns 4-6, OTHER					H (as identified in 45CFR46.101),
4.		-	sed research in			•		
[]	YES		If YES, this r	esearch	cannot be	e exempt	ed and must be reviewed l	by an IRB (skip to question 7).
[]	NO							
	es the prop uld apply) ⁶		arch involve fe	tuses, pr	egnant w	omen, oi	r human in vitro fertilizati	ion as targets (such that Subpart B
[]	YES		If YES, this question 7).	researc	h canno	t be exe	mpted and must be revi	iewed by an IRB (skip to
[]	NO							
Educatio	onal Resea	irch						
6.1	norm	al educati	onal practices (e.g., rese	earch on 1	regular a	and special education strat	gs, AND does the research involve tegies or research on the sroom management methods)?
	[]	YES		[]	NO			
	<u>h Involvin</u>	<u>g Surveys</u>	, Interview Pro	cedures	(includin	g Focus	groups), Observation of P	ublic Behavior, or Educational
<u>Tests</u>				_				
6.2	proce	dures or o	ch use education observation of p	public be	havior?	ve, diagn		nt), survey procedures, interview
	[]	YES		[]	NO		If NO skip 6.3	
			<18 years of age	·		•		
	[]	YES	If YES, this	research	cannot b	e exemp	ted and must be reviewed	by an IRB (skip to item 7)
	[]	NO						
	6.2.1	indirect	<u>ly</u> through ider		such as a	code) lin	nanner that human subjec ked to the subjects;	ts can be identified <u>directly or</u>
		[]	YES		[]	NO		
	6.2.2	place th employa subjects	e subjects at ri ability or reput s' (or relatives'	sk of crin ation? (H or associ	minal or (Examples iates') pos	civil liab here ma ssible sul	ility, or be damaging to they include: the collection of	ch setting have the potential to e subjects' financial standing, f sensitive data regarding the riminal history or intent, medical ormation).
		[]	YES		[]	NO		
6.3		his resear			s (cognitiv			nt), survey procedures, interview er paragraph 6.2 of this section:
	[]	YES		[]	NO		If NO skip to 6.4	
	6.3.1	Will thi public o		lve hum:	an subjec	ts that a	re elected or appointed pu	blic officials or candidates for
		[]	YES		[]	NO		
	6.3.2	informa	tion will be ma n only in the ca	intained	through	out the r	esearch and thereafter? (N	the personally identifiable Note: CDC can use this exemption been obtained to cover the
		[]	YES		[]	NO		
Existing	Data Whi	ich Is Pub	licly Available	or Unide	<u>entifiable</u>			
6.4	diagn	ostic speci					e the study begins)?	s, records, pathological or
	[]	YES		[]	NO		If NO skip to 7	
	6.4.1		naterial or info	rmation			?	
		[]	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.

See attachment

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
Christopher Pleasants - ORISE Fellow staff member completing this form	06/22/2016	 [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 [] Research involving human subjects, CDC investigators, not exempt [] Local IRB [] CDC Exemption [] CDC IRB
Katie Green - HEALTH 06/24. COMMUNICATION SPECIALIST		 [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
Team Lead		<u>Comments:</u> Thank you, Chris. This will be incredibly helpful.

Jon Baio - BEHAVIORAL SCIENTIST Division ADS	06/27/2016	 [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 [] Research involving human subjects, CDC investigators, not exempt [] Local IRB [] CDC Exemption [] CDC IRB
Scott Campbell - Health Scientist	06/27/2016	 [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 [] Research involving human subjects, CDC investigators, not exempt [] Local IRB [] CDC Exemption [] CDC IRB
CUB ADS, Deputy ADS, or Human Subjects Contact		<u>Comments:</u>