

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 (Use PGO number if co		
					(Ose I GO number ii co	орегануе	agreement, grant, etc.)
ate submitted:	02/09/2017						
itle of Project:	An Evaluation of M Program to Priority			proach	es to Scaling the Nation	nal Diabe	etes Prevention
ates for project pe	riod:	1	Dates for	funding	(if applicable):		
Beginning:	09/30/2017		Beginn	ing:	09/30/2017		_
Ending:	09/29/2022		Ending	g:	09/29/2018		<u>_</u>
roject is (choose o	ne):						
	as used below, refers to CDC staff member, dete				the project including scc c.	ppe of pro	ject, funding restrictions,
[X] New				[]	Revision		
[] Continuati	on, without revision(s))		[]	Continuation, with	Continuation, with revision(s)	
ead staff member:		Contact info	matiani			lo(a) in th	is project.
**		Contact infor Division:			Please indicate your rol Project officer	e(s) m ui []	roject: Technical monitor
Jeann	nette May	Division:	DDT		[] Principal	[]	Investigator
User ID: JXM5	;	Telephone:	770-488-	5016	investigator	[]	Investigator
Scientific Ethics	s number: 10297	Mailstop:	F75		[] Consultant	[X]	Other (please explain
						Ass	isting with Contract
						Pro	cessing
[] YES	f the activities within th [X] NO se activities which are		SIGNED t	o contri	oute to generalizable know	Pro	-
[] YES If YES, list tho	[X] NO se activities which are	research:				Pro	
[] YES If YES, list tho	[X] NO se activities which are oject research or publi	research:		ck all th	at apply)?	Pro	-
[] YES If YES, list the Is this CDC pr [] Resea	[X] NO se activities which are oject research or publi	research:	etice (chec	k all th Public		Pro	
[] YES If YES, list the Is this CDC pr [] Resea	[X] NO se activities which are oject research or publi	research: c health prac	etice (chec	k all th Public	at apply)? health practice	Production	-
[] YES If YES, list tho Is this CDC pr [] Resea	[X] NO se activities which are oject research or publicatch k one:	research: c health prac	etice (chec	ck all th Public <i>Check</i>	at apply)? health practice all that apply:	Production	e., research)?
Is this CDC pr Resea	[X] NO se activities which are oject research or publicatch k one: Human subjects inv Human subjects not	research: c health prac olved involved	etice (chec	ck all th Public Check	at apply)? health practice all that apply: Emergency Response	Production	e., research)? Surveillance Other (please explain)
Is this CDC pr Resea Check If RESEARCH Subjects protect	[X] NO se activities which are oject research or publicatch k one: Human subjects inv Human subjects not	research: c health prace olved involved jects, has the	etice (chec	ck all th Public Check [] [X]	at apply)? health practice all that apply: Emergency Response Program evaluation	Production	e., research)? Surveillance Other (please explain) e CDC IRB for human
Is this CDC pr Resea Check If RESEARCH subjects protect a. [] NO, No.	[X] NO se activities which are oject research or publicated the cone: Human subjects inv Human subjects not I involving human subjection?	research: c health prac olved involved jects, has the	etice (chec [X]	ck all th Public Check [] [X]	at apply)? health practice all that apply: Emergency Response Program evaluation ch activities been review	Proceed by the coved by	Surveillance Other (please explain) e CDC IRB for human
Is this CDC pr Resea Check If RESEARCH subjects protect a. [] NO, No, No, Ex	[X] NO se activities which are oject research or publicated the cone: Human subjects involving human subjection?	research: c health prac olved involved jects, has the	etice (chec [X]	ck all th Public Check [] [X]	at apply)? health practice all that apply: Emergency Response Program evaluation ch activities been review	Proceed by the coved by	Surveillance Other (please explain) e CDC IRB for human
Is this CDC pr Resea Check If RESEARCH subjects protect a. [] NO, No, No, Ex	[X] NO se activities which are oject research or publicated the cone: Human subjects inv Human subjects not I involving human subjection? ew project, not yet revisiting project, not reactivities	research: c health prac olved involved jects, has the	etice (chec [X] project o d.	ek all th Public Check [] [X] r resean	at apply)? health practice all that apply: Emergency Response Program evaluation ch activities been review TES, Reviewed and appl	Proceed by the coved by the coved by	e., research)? Surveillance Other (please explain) e CDC IRB for human CDC mber_and

Form 684R_NR (revised January 2003)

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Tracking NO. <u>TBD/TBD</u>

Name				Scientific ethics number Prin						
Jeannette May						10297				
				EARCH PRO					I (as identified in 45CFR46.101),	
4.	-		•	sed research			-			
	[]	YES		If YES, thi	s researc	h cannot b	e exempt	ed and must be reviewed b	y an IRB (skip to question 7).	
	[]	NO								
5.		the prop d apply):		earch involve	fetuses, j	pregnant v	women, o	r human in vitro fertilizatio	on as targets (such that Subpart B	
	[]	YES								
	[]	NO								
Edi	ucation	al Resea	rch							
	6.1	Is this	research al educat	ional practice	es (e.g., re	esearch on	regular a	and special education strate	gs, AND does the research involve egies or research on the room management methods)?	
		[]	YES		[]	NO		•	,	
Res	search l		g Surveys	s, Interview F		es (includi	ng Focus	groups), Observation of Pu	ublic Behavior, or Educational	
Tes	<u>sts</u>	,	<u>-</u>							
	6.2			rch use educa observation o			ive, diagn	ostic, aptitude, achievemen	nt), survey procedures, interview	
		[]	YES		[]	NO		If NO skip 6.3		
		Will children (<18 years of age) be research subjects?								
		[]	YES	If YES, this research cannot be exempted and must be reviewed by an IRB (skip to item 7)						
		[]	NO							
		6.2.1		information obtained recorded in such a manner that human subjects can be identified <u>directly or ctly</u> through identifiers (such as a code) linked to the subjects;						
			[]	YES		[]	NO			
		6.2.2	place the employ subject	ne subjects at ability or rep s' (or relative	risk of coutation? es' or asso	riminal or (Example ociates') po	civil liab s here ma ossible su	ility, or be damaging to the ry include: the collection of	h setting have the potential to e subjects' financial standing, sensitive data regarding the riminal history or intent, medical rmation).	
			[]	YES		[]	NO			
	6.3	proce	dures, or		of public	behavior		search is not exempt under	nt), survey procedures, interview r paragraph 6.2 of this section:	
		[]	YES		[]	NO		If NO skip to 6.4		
		6.3.1	Will thi public		volve hu	man subje	cts that a	re elected or appointed pul	blic officials or candidates for	
		[]	YES		[]	NO				
		6.3.2	inform	ation will be a on only in the	maintain	ed througl	hout the r	ion that confidentiality of t esearch and thereafter? (N nce of Confidentiality has b	ote: CDC can use this exemption	
			[]	YES		[]	NO			
Exi	isting D	ata Whi	ch Is Pul	olicly Availab	<u>le or Uni</u>	dentifiabl	<u>e</u>			
	6.4							f existing* data, documents e the study begins)?	s, records, pathological or	
		[]	YES		[]	NO		If NO skip to 7		
		6.4.1	Is this 1	naterial or in	formatio	n publicly	available	?		
		[]	YES		[]	NO				

Tracking	NO.	TBD/TBD

T :	rackir	ıg NO	. <u>TBD</u>	/TBD	_		
		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 I 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.		lease prepare and attach a short summary paragraph (<1 page); 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 (researchnon-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.					
		popula Americ The pro 1)testir 2) deve	tions with an Indial ogram eveng and ele eloping a	n prediabetes ns, Alaska Na valuation effor nhancing the new informat	sive evaluation of the cooperative agreement DP17-1705 to scale the National DPP to priority including: Medicare beneficiaries, men, African-Americans, Asian-Americans, Hispanics, atives, Pacific Islanders, and people with visual or physical disabilities. It is of this task order will consist of: CDC-developed data collection tools for the national evaluation of DP17-1705 FOA; tion collection package and obtaining approval from the Office of Management and Budget (OMB)		

- for all final data collection tools specified in the DP17-1705 National Evaluation Plan developed by CDC;
- 3) developing and maintaining an integrated database to systematically collect data at the grantee, site, and participant level;
- 4) providing technical assistance for grantees and affiliate sites on data collection, evaluation plan development and
- implementation, program implementation and delivery, and dissemination of emerging/promising practices and guidance; 5) collecting, validating, and linking data from different sources identified in the national evaluation protocol for CDC to conduct analyses; developing consumer-friendly emerging/promising practice documents, materials, resources, and tools around scaling the National DPP to priority populations; and
- 6) identifying key topic areas for case studies and focus groups, designing and implementing evaluation plans, collecting and analyzing data, and synthesizing and disseminating results.
- 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
Jeannette May - Public Health Advisor	02/09/2017	 [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:

Patricia Schumacher - SENIOR TEAM LEAD	02/09/2017	 [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:
Elizabeth Luman - EPIDEMIOLOGIST	02/10/2017	[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: Approve
Joan Redmond Leonard - PUBLIC HEALTH ANALYST	02/21/2017	[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: