

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.

				Ti	racking Number:						
				(U	se PGO number if coo	perative	e agreement, gr	ant, etc.)			
Date submitt	ted: 06/07/2011										
Title of Proj	The National [Diabetes Preventi	ion Program: I	Diabet	es Prevention Reco	gnition	Program				
Dates for pro	oject period:]	Dates for fund	ing (if	applicable):						
Beginning	: 10/03/2011		Beginning:								
Ending:	10/03/2014	0/03/2014									
Project is (ch	noose one):										
	vision, as used below, refe role of CDC staff member				project including scop	pe of pro	oject, funding i	estrictions,			
[X] Nev	v			[]	Revision						
[] Cor	ntinuation, without revisi	on(s)		[]	Continuation, with	revision	n(s)				
Lead staff m	ember:	Contact info	rmation:	Ple	ase indicate your role	e(s) in th	his project:				
Name:	Russell Sniegowski	Division:	DDT	[X]	Project officer	[]	Technical :	monitor			
				_ []	Principal	[]	Investigato	or			
User ID:	rjs3	Telephone:	770-488-5033	_	investigator						
Scientific	Ethics number:	Mailstop:	K-10	_ []	Consultant	[]	Other (ple	ase explain)			
	list those activities which										
	CDC project research or	public health prac									
[]	Research				lth practice						
	Check one:				hat apply:		g 911				
	[] Human subject [] Human subject		[]		nergency Response	[] [X]	Surveillance				
	numan subject	s not involved	ij	rr	ogram evaluation	[X]	Other (pleas	To recognize and monitor lifestyle programs to prevent type 2 diabetes			
	EARCH involving human protection?	subjects, has the	project or res	earch :	activities been review	ed by tl	he CDC IRB f	or human			
	NO, New project, not yet		d. []		, Reviewed and appro	_					
	NO, Existing project, no	-		Ι	f YES, please list pro	tocol nu	umber_and				
c. [] NO, Submitted for approval					expiration date						

							e. []	NO, RESEARCH, no required)	CDC investigators (CDC IRB not
							f. []	N/A (Not Applicable)	
	If RES	SEARC	H, list any	other CDC sta	ff invol	ved in this			e, role, and scientific ethics number
Name					Scientific ethics number Prin				
	Rus	ssell Sn	iiegowski						
				ARCH PROJE as 4-6, OTHER					CH (as identified in 45CFR46.101),
4.	LASE A		_	sed research inv			lucstion	·•	
••	[]	YES	ле ргоро.		-		exempte	d and must be reviewed	l by an IRB (skip to question 7).
	[]	NO					т		
5.	Does tl			arch involve fetu	ıses, pro	egnant wo	omen, or	human in vitro fertiliza	tion as targets (such that Subpart B
	[]	YES		If YES, this r question 7).	esearc	h cannot	be exen	npted and must be re	viewed by an IRB (skip to
	[]	NO		•					
			•						
<u>Edu</u>	<u>icationa</u>			1 4 11			,		
	6.1	norma	ıl educatio	nal practices (e	.g., rese	arch on r	egular a	nd special education str	ings, AND does the research involve ategies or research on the assroom management methods)?
		[]	YES	I	[]	NO			
		nvolving	g Surveys,	Interview Proc	edures	(including	g Focus g	roups), Observation of	Public Behavior, or Educational
Tes	<u>ts</u> 6.2	XX7211 41	h .	.hJ	1 4 4	(aaa :4:		-4:4:4 d	
	0.2			bservation of pu			e, diagno	suc, aputude, acmeven	nent), survey procedures, interview
		[]	YES		[]	NO		If NO skip 6.3	
		Will cl	hildren (<	18 years of age)	be rese	arch subj	ects?		
		[]	YES	If YES, this re	esearch	cannot be	exempt	ed and must be reviewe	d by an IRB (skip to item 7)
		[]	NO						
		6.2.1				_		anner that human subjects;	ects can be identified <u>directly or</u>
			[]	YES		[]	NO		
		6.2.2	place the employa subjects	e subjects at risl bility or reputa ' (or relatives' o	k of crir tion? (E r associ	ninal or c Examples ates') pos	ivil liabil here may sible sub	ity, or be damaging to t include: the collection	rch setting have the potential to the subjects' financial standing, of sensitive data regarding the criminal history or intent, medical formation).
			[]	YES		[]	NO		
	6.3								nent), survey procedures, interview der paragraph 6.2 of this section:
		[]	YES	1	[]	NO		If NO skip to 6.4	
		6.3.1	Will this public of		ve huma	n subject	s that ar	e elected or appointed p	oublic officials or candidates for
			[]	YES		[]	NO		
		6.3.2	informa	tion will be main only in the case	ntained	througho	ut the re	search and thereafter?	f the personally identifiable (Note: CDC can use this exemption s been obtained to cover the
			[]	YES		[]	NO		

 $\underline{\textbf{Existing Data Which Is Publicly Available or Unidentifiable}}$

T	rackin	ig NO	. <u>No Fu</u>	inding									
	6.4							f existing* data, documents, records, pathological or e the study begins)?					
		[]	YES		[]	NO		If NO skip to 7					
		6.4.1	Is this m	naterial or in	formation	ı publicly	availabl	e?					
			[]	YES		[]	NO						
		6.4.2	5.4.2 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 ide	entifying	information and no unique identifiers or codes)YES					
			[]	NO	(there	are identi	ifiers (inc	cluding codes))					
7.		prepar is new:	e and atta	ich a short su	ımmary p	oaragraph	ı (<1 pag	e);					
	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.											
		In 2010, the Centers for Disease Control and Prevention established the CDC Diabetes Prevention Recognition Program (DPRP) as part of the National Diabetes Prevention Program. The purpose of the DPRP is to recognize and monitor the effectiveness of organizations delivering a proven diabetes lifestyle intervention program. The DPRP will also provide information to people at high risk of type 2 diabetes, their health care providers and health payers on the location and performance of local diabetes prevention programs. The DPRP is public health practice based upon science-based efficacy and effectiveness studies specific to diabetes prevention.											
		To allow DDT to assess/monitor program effectiveness, renew recognition status and provide technical assistance, recognized programs will submit raw data (raw process and outcome measures) to DDT. This data will not contain personal identifiers (codes, created and maintained by the programs will replace names of participants, program staff, etc.). DDT staff will analyze the data to quantify program effectiveness and assure programs meet the criteria for recognition. OMB approval is being obtained for this data collection.											
				cations are res Il applicable fe			g the priv	acy and confidentiality of lifestyle program participant data and					
				Officer is restated at CDC.	oonsible fo	or the prog	rammatic	oversight of the DPRP, in collaboration, as appropriate, with					
8.	Please	list the	primary]	project site a	nd all col	laboratin	g site(s).						
	Explai	nation o	f project	components:									
9.		roject involves research that is funded extramurally, list amount of award that should be restricted pending IRB roval and describe which project components will be affected, if known:											

approval and des	cribe v	vhich project	components v	vill be a	ffected, if	known:			
 1 (1)	-	• • • • • • •	ъ.		1.0		/ D		

Approvals (signature and position title)	Date	Research Determination / Remarks
Russell Sniegowski - PUBLIC HEALTH CONSULTANT	06/07/2011	[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:
Grace Thomas - Contracto9r	06/07/2011	[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: PO is in the OD so there is no branch chief
Grace Thomas - Contracto9r Division ADS	06/07/2011	[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	06/13/2011	 [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>