

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.

To Be Determined

							agreement grant etc.)
				(U	se PGO number if coo	perative	agreement, grant, etc.)
te submitted:	03/30/2018						
tle of Project:	DP 18-1810 Sta	te Actions to In	nprove Oral H	ealth C	Outcomes		
tes for project pe	riod:		Dates for fund	ling (if	applicable):		
Beginning:	09/01/2018		Beginning:		09/01/2018		
Ending:	08/31/2023		Ending:		08/31/2019		_
oject is (choose on	ne):						
NOTE: Revision, a					project including scop	pe of pro	oject, funding restrictions
[X] New				[]	Revision		
[] Continuation	on, without revision	n(s)		[]	Continuation, with	revision	n(s)
nd staff member:		Contact info	rmation:	Ple	ase indicate your role	e(s) in tl	his project:
	a Parker	Division:	DOH	[X]	Project officer	[]	Technical monitor
- Maron	21 01101		2011	_ []	Principal Principal	[]	Investigator
							0
Jser ID: KUV7		Telephone:	770-488-6075	_	investigator		
Scientific Ethics	number: f the activities within	Mailstop:	F-80	_ []	investigator Consultant to generalizable know	[] vledge (i	Other (please explain
Are any or all of [] YES	number: f the activities within	Mailstop: n this project DE	F-80	_ []	Consultant		
Are any or all of [] YES If YES, list those	f the activities within [X]	Mailstop: n this project DE: NO are research:	F-80 SIGNED to con	[]	Consultant to generalizable know		
Are any or all of [] YES If YES, list those	f the activities within [X] See activities which a Dject research or pu	Mailstop: n this project DE: NO are research:	F-80 SIGNED to con	[]	Consultant to generalizable know		
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Are any or all of [] YES If YES, list those Is this CDC pro [] Resea	f the activities within [X] See activities which a Dject research or purch	Mailstop: n this project DE: NO are research: ublic health prac	F-80 SIGNED to concertice (check all	that a	Consultant to generalizable know pply)? Ith practice		
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Are any or all of [] YES If YES, list those Is this CDC process [] Resea Check [] [] If RESEARCH subjects protect a. [] NO, Ne	f the activities within [X] See activities which a Dject research or purch A one: Human subjects Human subjects (involving human stion?	Mailstop: n this project DE: NO nre research: ublic health praction of involved subjects, has the reviewed	F-80 SIGNED to constitute (check all [X] Public Check [] [X] sproject or research []	that a blic hear ck all the Property of the Pr	consultant to generalizable know pply)? Ith practice that apply: nergency Response ogram evaluation activities been review	[X]	Surveillance Other (please explain) the CDC IRB for human
Are any or all of [] YES If YES, list thos Is this CDC pro [] Resea Check [] If RESEARCH subjects protect a. [] NO, Ne b. [] NO, Ex	f the activities within [X] See activities which a oject research or put arch a one: Human subjects Human subjects involving human stion?	Mailstop: In this project DE: NO The research: In this project DE: In this project DE:	F-80 SIGNED to constitute (check all [X] Public Check [] [X] sproject or research []	that a plic hear Property Search :	consultant to generalizable know pply)? Ith practice that apply: nergency Response ogram evaluation activities been review , Reviewed and approf YES, please list pro expiration date	[X] [] red by the oved by tocol many	Surveillance Other (please explain) he CDC IRB for human CDC umber_and
Are any or all of [] YES If YES, list thos Is this CDC pro [] Resea Check [] If RESEARCH subjects protect a. [] NO, Ne b. [] NO, Ex	f the activities within [X] See activities which a oject research or purch it one: Human subjects Human subjects involving human stion? ew project, not yet resisting project, not research	Mailstop: In this project DE: NO The research: In this project DE: In this project DE:	F-80 SIGNED to constitute (check all [X] Public Check [] [X] sproject or research []	that a plic hear Property Search :	consultant to generalizable know pply)? Ith practice that apply: nergency Response ogram evaluation activities been review , Reviewed and appro f YES, please list pro expiration date RESEARCH, no CD	[X] [] red by the oved by tocol many	Surveillance Other (please explain) the CDC IRB for human

Form 684R_NR (revised January 2003)

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Tracking NO. To Be Determined

Name			Role (project officer, investigator, consultant, etc.)					Scientific ethics number Prin		
	Ma	arcia Pa	rker							
				EARCH PRO					CH (as identified in 45CFR46.101),	
4.			_	sed research i			=			
	[]	YES		If YES, this	research	cannot b	e exempto	ed and must be reviewe	ed by an IRB (skip to question 7).	
	[]	NO					-			
5.				arch involve f	etuses, pi	regnant v	vomen, or	human in vitro fertiliz	ation as targets (such that Subpart B	
	[]	YES		If YES, this question 7)		ch canno	ot be exei	npted and must be r	eviewed by an IRB (skip to	
	[]	NO								
<u>Ed</u>	ucationa	al Resea	<u>rch</u>							
	6.1	norma	al educatio	onal practices	(e.g., res	earch on	regular a	nd special education st	tings, AND does the research involve rategies or research on the lassroom management methods)?	
		[]	YES	_	[]	NO		-	-	
Re	search I	nvolving	g Surveys.	, Interview Pr	ocedures	(includi	ng Focus	groups), Observation o	f Public Behavior, or Educational	
Te	<u>sts</u>									
	6.2			ch use educat bservation of			ive, diagno	ostic, aptitude, achieve	ment), 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 exempt	ed and must be review	ed by an IRB (skip to item 7)	
		[]	NO	0						
		6.2.1	6.2.1 Is the information obtained recorded in such a manner that human subjects can be identified <u>directly or indirectly</u> through identifiers (such as a code) linked to the subjects;							
			[]	YES		[]	NO			
		6.2.2	place the employa subjects or psych	e subjects at 1 ability or repu ' (or relatives aological cond	risk of cri tation? (I ' or assoc	minal or Example ciates') po ancial sta	civil liabi s here ma ossible sub atus, or sir	lity, or be damaging to y include: the collection	arch setting have the potential to the subjects' financial standing, n of sensitive data regarding the r, criminal history or intent, medical information).	
			[]	YES		[]	NO			
	6.3	proced	dures, or o		f public b	ehavior		search is not exempt u	ment), survey procedures, interview nder paragraph 6.2 of this section:	
		[]	YES		[]	NO		If NO skip to 6.4		
		6.3.1	Will this public of		olve hum	an subje	cts that ar	e elected or appointed	public officials or candidates for	
			[]	YES		[]	NO			
		6.3.2	informa	tion will be m n only in the c	aintained	l through	out the r	esearch and thereafter?	of the personally identifiable? (Note: CDC can use this exemption as been obtained to cover the	
			[]	YES		[]	NO			
Ex	isting Da	ata Whi	ch Is Publ	licly Available	e or Unid	<u>entifiable</u>	<u>e</u>			
	6.4							existing* data, docume the study begins)?	ents, records, pathological or	
		[]	YES		[]	NO		If NO skip to 7		
		6.4.1	Is this m	naterial or inf	ormation	publicly	available	?		
			[]	YES		[]	NO			

					ated by an investigator even temporarily, for research purposes, this criterion is not met. s 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 if this		nd attac	h a short sur	mmary paragraph (<1 page);	
	a.	(s) in the like: stud and parti	project. y design cpation	In explainin decisions, o	se of the project, specific details about the project and the role of the CDC staff membering one's role as a consultant be particularly careful to identify involvement in things oversight of protocol development, participation in review of data collection procedures, plysis and/or manuscript preparation, as well as whether there will be access to a.	
	b.	subjects;	public h any pers	ealth practi	selection (researchnon-exempt, exempt, no CDC investigator or not involving human tice). If you selected research not involving human subjects be sure to indicate if the data nation (e.g., name, SSN), linkable study identification numbers or codes, or geographical	
		renewal to The purpo associated strategies and increa with other FY 2017], Chronic D care. The	o continuose is to a d with po to support continuose con chronic and inconsesses public he	e CDC invest assist States to or oral health ort school sea mmunity wate disease progrorporates progrand and Oral Hea ealth impact fi	to Improve Oral Health Outcomes program is a five-year cooperative agreement competitive stment in and support for state oral health programs. This program does not include research. It to decrease dental caries, oral health disparities, and other co-morbid chronic diseases houtcomes. Recipients will accomplish these outcomes through implementing priority palant programs (including promoting adherence to infection prevention guidelines), supporting er fluoridation, conducting state oral health surveillance, and optionally integrating oral health grams (i.e., medical/dental integration). The proposed program replaces FOA 13-1307 [FY 2013-ogrammatic strategies from FOA 16-1609 [FY 2016-FY 2017] Models of Collaboration among alth Programs, to support enhanced medical/dental integration and provide improved quality of from the successful implementation of priority strategies includes long-term outcomes of all health disparities and other co-morbid chronic diseases.	-
		stakehold Conference	ers and p ces, SEA	partners throu LS, and Wate	ammatic, evaluation, epidemiologic, and technical assistance for recipients and their rugh programmatic and one-on- one technical consultation, national training, workshops, Web ter Fluoridation Reporting System (WFRS) training, and other forms of guidance. CDC will also between national partners and recipients as needed.	
		Fluoridation programs population Locals and information treatment; and report that allows	on Repor . WFRS in on com d States in to eval) with the ting their is funded g the nee	ting System (nformation is munity water (SEALS) is do uate the effect associated c performance state progran	address the collection of information to support states receiving 18-1810 funding. The Water (WFRS) is an online tool that helps states manage the quality of their water fluoridation is also the basis for national surveillance reports that describe the percentage of the U.S. It is resulted to capture, store, and analyze school sealant program data. Programs can use this exciveness of individual school sealant programs by comparing the benefits (e.g., averted costs (e.g., resources used, labor hours). These systems are tools useful to states in collecting to measure information. States will also use CDMIS, a password protected web-based system are to submit their progress reports annually by entering information into the system, thereby anal written reports. Respondents provide progress report information through prompted data	
3.	Please	list the pri	imary p	roject site an	nd all collaborating site(s).	
	Explar	nation of p	roject co	omponents:		
).					unded extramurally, list amount of award that should be restricted pending IRB t components will be affected, if known:	

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?

Approvals (signature and position title)	Date	Research Determination / Remarks
Lisa Petersen - Public Health Advisor	03/30/2018	[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:
Marcia Parker - Team Leader	03/30/2018	[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: Reviewed and approved.
Lorena Espinoza - Dental Officer/Team Lead	03/30/2018	[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: DP 18-1810 has been released.
Shanna Cox - Associate Director for Science	04/03/2018	[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>