

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.

					se PGO number if coo		agreement, gr	ant, etc.)
Date submitte	ed: 02/05/2020							
Title of Proje	Comprehensive	———— Cancer Contro	ol Branch Man	ageme	ent Information Syste	em		
Dates for pro			Dates for fund					
Beginning:	-		Beginning:	8 (пррисш аго) г			
Ending:	03/30/2022		Ending:					
- -							_	
	ision, as used below, refers t				project including scop	pe of pro	eject, funding r	estrictions,
•	role of CDC staff member, d	etermination of	research status					
[] New				[]	Revision			
[X] Con	tinuation, without revision	(s)		[]	Continuation, with	revision	(s)	
Lead staff me	mber:	Contact info	rmation:	Ple	ase indicate your role	(s) in th	is project:	
Name:	Angela Moore	Division:	DCPC	[]	Project officer	[X]	Technical r	nonitor
User ID:	CYQ6	Telephone:	770-488-3094	[]	Principal investigator	[]	Investigato	r
Scientific	Ethics number: 680	– 3 Mailstop:	K57	[]	Consultant	[]	Other (plea	se explain)
	ist those activities which a		ctice (check all	that a	pply)?			
[]	Research	F			lth practice			
	Check one:				that apply:			
	[] Human subjects in	nvolved	[]	En	nergency Response	[]	Surveillance	
	[] Human subjects n	ot involved	[]	Pr	ogram evaluation	[X]	Other (please	explain)
								Reporting of submissior of continuation n application for ongoing cancer control program
	ARCH involving human suprotection?	bjects, has the	project or res	earch :	activities been review	ed by th	ne CDC IRB f	or human
a. [] I	NO, New project, not yet re	eviewed	d. []	YES	, Reviewed and appro	oved by	CDC	
b. [] I	NO, Existing project, not re	eady to submit		I	f YES, please list pro	tocol nu	mber_and	

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	c. []	NO, S	Submitted	for approval					expiration date	
							e.	[]	NO, RESEARCH, no CD required)	oC investigators (CDC IRB not
							f.	[]	N/A (Not Applicable)	
	If RE	SEARC	H, list an	y other CDC st	taff invo	lved in tl	his pr	oject,	please include the name, i	role, and scientific ethics number
	Na	ame				Role (pro onsulta			er, investigator,	Scientific ethics number Prin
	Ar	ngela M	oore							6803
IF.	— VOLLTI	HINK T	HE RESI	FARCH PROI	FCT MI	CHT O	ΠΔΙΙ	FV A	S FYFMPT RESEARCH	(as identified in 45CFR46.101),
				ns 4-6, OTHEI						(as identified in 45 CF R40.101),
4.		Does t	the propo	sed research ii	nvolve pi	risoners?				
	[]	YES		If YES, this	research	cannot l	oe exe	empte	d and must be reviewed by	an IRB (skip to question 7).
	[]	NO								
5.		the prop l apply)?		arch involve fe	etuses, pi	regnant v	wome	n, or	human in vitro fertilizatio	n as targets (such that Subpart B
	[]	YES		If YES, this question 7).		ch cann	ot be	exen	npted and must be revie	wed by an IRB (skip to
	[]	NO								
Ed	ucationa	al Resea	<u>rch</u>							
	6.1	Is this norma	research al educati	onal practices	(e.g., res	earch on	regu	lar a	nd special education strate	s, AND does the research involve gies or research on the room management methods)?
		[]	YES		[]	NO				
		nvolving	g Surveys	, Interview Pro	ocedures	(includi	ng Fo	ocus g	roups), Observation of Pu	blic Behavior, or Educational
Tes		TT 7*11 41			1				1	
	6.2			ch use education of				iagno	stic, aptitude, achievemen	t), survey procedures, interview
		[]	YES		[]	NO			If NO skip 6.3	
		Will c	hildren (<	<18 years of ag	e) be res	earch su	bject	s?		
		[]	YES	If YES, this	research	cannot	be ex	empt	ed and must be reviewed b	y an IRB (skip to item 7)
		[]	NO							
		6.2.1							anner that human subjects ted to the subjects;	can be identified <u>directly or</u>
			[]	YES		[]	N	0		
		6.2.2	place the employa subjects	e subjects at ri ability or reput s' (or relatives'	isk of cri tation? (or assoc	minal or Example iates') p	civil s her ossibl	liabil e may le sub	ity, or be damaging to the vinclude: the collection of	setting have the potential to subjects' financial standing, sensitive data regarding the minal history or intent, medical mation).
			[]	YES		[]	N	0		
	6.3									t), survey procedures, interview paragraph 6.2 of this section:
		[]	YES		[]	NO			If NO skip to 6.4	
		6.3.1	Will thi public o		olve hum	an subje	ects th	at ar	e elected or appointed pub	lic officials or candidates for
			[]	YES		[]	N	O		
		6.3.2	informa	ntion will be main only in the ca	aintaineo	d through	hout t	the re	on that confidentiality of the search and thereafter? (No ce of Confidentiality has be	ote: CDC can use this exemption
			[]	YES		[]	N	0		

<u>Exi</u>	sting Da	ata Whi	ch Is Pul	olicly Availab	ole or Unio	dentifiable	e									
	6.4			rch involve o cimens? (* 'ex							uments	, recor	ds, pa	thologi	cal or	
		[]	YES		[]	NO		If N	O skip	to 7						
		6.4.1	Is this i	material or ir	nformatio	n publicly	availabl	e?								
			[]	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?												
							vestigator even temporarily, for research purposes, this criterion is not met linical staff who already have access to the data, this criterion is met).									
			[]	YES	(there	are no id	entifying	inforn	nation	and no	unique	identi	fiers o	or codes	s)YES	
			[]	NO	(there	are identi	ifiers (in	cluding	g codes))						
7.			e and att	ach a short s	ummary j	paragraph	ı (<1 pag	e);								
	if this	is new:		ude the purp												
	b.	like: s and pa identi Expla subject	tudy desi articpatio fiable or in your p ets; publi	ct. In explain ign decisions, on in data an personal data oroject status c health prac ersonal infor	oversight alysis and a. selection ctice). If yo	t of protoc //or manus (research- ou selected	col develo script pr non-exo l researc	opment eparati empt, e eh not i	t, particion, as exempt, nvolvir	cipation well as , no CD ng hum	n in rev whether OC inversants	view of er ther stigato jects bo	data de will d	collection be access ot involuto indic	on proc ss to lving hi cate if t	edures, uman he data
			nation.	ersonai mior	manon (e.	g., name,	33N), III	ikabie	Study 1	aemmi	auon i	iuiiibei	is or c	oues, o	r geogr	аршсаг
		Compore reinsta for the a. The b. This (the No system progree NCCC c. As t reflect	rehensive atement at National CDC states is public CCCP), for natic, and act P programhe technic the 2017	this project is Cancer Contropplication has Comprehensiff role is as a thealth practic or use in moniti it is not intenctivity informations in 2017, wheal monitor, Cl cooperative a iffable information.	rol Program been subrive Cancer echnical mee (not resetoring progeded to procedion to CDC bich place of CDC staff wigreement	ns). The comitted to comitted to committed to committed to committe to pearch). The gram progree generates twice per emphasis of the committed to the committed that	ollection of ontinue day ogram (No rovide over project less and to alizable rought year using policy the controle require	of this in ta colle CCCP) ersight s desig o use fo esults. og this s and envactor in	nformation. to the coned to coned to coned to cone Since Jystem. vironme making	ion is ap The CD contractor collect in am improducing January New contal app g minor	oproved MIS ma or. nformat ovemen 2010, Noperation	under intains ion from t. The NCCCP we agrees to cast of the	omb in granter of an an or collect award ement of existing the collection of the col	ngoing C tion of the same as swere arevention	0-0841, fic information for the control of the cont	and a mation ogram is not nitted d to all ontrol.
8.	Please	list the	primary	project site	and all co	llaboratin	g site(s).									
	Expla	nation o	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
Angela Moore - Lead Public Health Advisor staff member completing this form	02/05/2020	[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
starr memoer completing this form		<u>Comments:</u>

- MM/dd/yyy	[] 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	<u>Comments:</u>