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.

- (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: OT13-1302

(Use PGO number if cooperative agreement, grant, etc.)

Date submitted: Title of Project:		:	07/31/2019									
		:	American Academy of Pediatrics resident training on fetal alcohol spectrum diso clinics								orders in continuity	
Dates for	r proje	ct peri	od:			Dates for	fundi	ng (if	applicable):			
Begin		08/01/2018	8/01/2018			Beginning:		08/01/2019				
Ending:			07/31/2023			Ending:			07/31/2020		_	
Project is	s (choo	se one)):									
			used below, r DC staff memi						project including sco	pe of proj	ect, funding restrictions,	
[X]	New							[]	Revision			
[]	Contir	nuatior	n, without rev	vision(s))	[] Continuation, with revision(s				s)		
Lead stat	ff mem	iber:			Contact info	rmation:		Ple	ase indicate your role	e(s) in thi	s project:	
Name	e: J	Jacquel	yn Bertrand		Division:	DCDD		[]	Project officer	[X]	Technical monitor	
User]	- ID: ι	JZB1	-		Telephone:	404-498	-3928	[]	Principal investigator	[]	Investigator	
Scien	tific E	thics r	umber:	15342	Mailstop:	S106-3		[]	Consultant	[]	Other (please explain)	
[]	Ŋ	YES	he activities w [X] activities wh	NO		SIGNED	to cont	ribute	to generalizable know	vledge (i.	e., research)?	
2. Is th	his CD	C proj	ect research o	or publi	c health prac	ctice (che	ck all t	that a	pply)?			
[]	F	Researc	ch			[X]	Publi	ic hea	lth practice			
	6	Check a	k one:			Check all			that apply:			
	[]	Human subj	ects inv	olved		[]	En	nergency Response	[]	Surveillance	
	[]	Human subj	ects not	involved		[X]	Pr	ogram evaluation	[]	Other (please explain)	
	ESEA1			ıan sub	jects, has the	project	or rese	arch	activities been review	ved by the	e CDC IRB for human	
a.	[] N (D, New	project, not	yet revi	iewed	d.	[]	YES	, Reviewed and appr	oved by (CDC	
b.	[] N (O, Exis	ting project,	not rea	dy to submit			Ι	f YES, please list pro	otocol nu	mber_and	
с.	[] NO	O, Subi	nitted for ap	proval					expiration date			
						e.	[]		RESEARCH, no CD ired)	C investi	gators (CDC IRB not	
						f.	[]	N/A	(Not Applicable)			
If R	ESEA	RCH.	ist any other	CDC s	taff involved	in this p	roject.	pleas	e include the name. r	ole, and	scientific ethics number	

Tracking NO. <u>OT13-1302</u>

Name					Ro co		Scientific ethics number Prin			
	Jac	quelyn	Bertrand							15342
				RCH PROJI 4-6, OTHER				S EXEMPT RESEAR '.	CH (as identified	l in 45CFR46.101),
4.		Does tl	he propose	d research in	volve pri	soners?	_			
	[]	YES	1	f YES, this re	esearch c	annot be	exempte	d and must be reviewe	d by an IRB (ski	p to question 7).
	[]	NO								
5.		ne propo apply)?	osed resear	ch involve fet	uses, pre	egnant wo	omen, or l	human in vitro fertiliz	ation as targets (such that Subpart B
	[]	YES		If YES, this question 7).	researcl	h cannot	be exen	npted and must be re	eviewed by an l	RB (skip to
	[]	NO								
<u>Edı</u>	ucationa	l Resear	<u>.ch</u>							
	6.1 Is this research conducted in established or commonly accepted educational settings, AND does the research involve normal educational practices (e.g., research on regular and special education strategies or research on the effectiveness of, or comparison among instructional techniques, curricula or classroom management methods)?							rch on the		
		[]	YES		[]	NO				
<u>Res</u> Tes		volving	Surveys, I	nterview Pro	cedures (including	g Focus g	roups), Observation of	Public Behavior	<u>r, or Educational</u>
105	6.2			use educatio servation of p			e, diagno	stic, aptitude, achiever	nent), survey pro	ocedures, interview
		[]	YES		[]	NO		If NO skip 6.3		
		Will ch	nildren (<18	8 years of age) be rese	arch subj	jects?			
		[]	YES	If YES, this r	esearch	cannot be	e exempte	ed and must be review	ed by an IRB (sk	ip to item 7)
		[]	NO							
		6.2.1						nner that human subj ed to the subjects;	ects can be ident	ified <u>directly or</u>
				YES		[]	NO			
		6.2.2	place the s employabi subjects' (subjects at ris ility or reputa or relatives' o	k of crin ation? (E or associa	ninal or c xamples 1 ates') pos	ivil liabil here may sible subs	ses outside of the resea ity, or be damaging to include: the collection stance abuse, sexuality ilarly compromising in	the subjects' fina of sensitive data , criminal histor	ancial standing, a regarding the
			[]	YES		[]	NO			
	6.3							stic, aptitude, achiever earch is not exempt un		
		[]	YES		[]	NO		If NO skip to 6.4		
		6.3.1	public offi	ice?	ve huma	-	s that are	e elected or appointed	public officials o	r candidates for
			[]	YES		[]	NO			
		6.3.2	informatio	on will be ma only in the cas	intained	througho	out the re	n that confidentiality (search and thereafter? ee of Confidentiality ha	(Note: CDC car	n use this exemption
			[]	YES		[]	NO			
Exi	<u>sting Da</u>	<u>ta Whic</u>	<u>h Is Public</u>	ly Available	o <mark>r Unide</mark>	<u>ntifiable</u>				
	6.4	diagno	stic specim		ing' mea	ns existin	g before	existing* data, docume the study begins)?	ents, records, pat	hological or
		[]	YES		[]	NO		If NO skip to 7		
		6.4.1		terial or info WFG	mation _l					
			[]	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 particpation 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).

	Site Name	Site Location	Assurance Number (FWA, MPA or SPA) if applicable
Primary Site	American Academy of Pediatrics	Itasca, IL	

Explanation of project components:

see attachement

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
Jacquelyn Bertrand - Epidemiologist staff member completing this form	07/31/2019	 [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
start member completing this form		<u>Comments.</u>
Shin Kim - EPIDEMIOLGIST	08/01/2019	 [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>

Leslie O'leary - EPIDEMIOLOGIST	08/02/2019	 [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:
Scott Campbell - Health Scientist	08/13/2019	 [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		Comments:

List of Grantees

Grantee #

<u>Grantee Name</u>

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American Academy of Pediatrics (AAP)