

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.

				/T T	DCO 1 'C		
				(U	se PGO number if coo	perative	agreement, grant, etc.)
Date submitted:	07/17/2018						
Title of Project:	Sudden Death	in the Young Re	egistry				
Dates for project	period:	:	Dates for fund	ing (if	applicable):		
Beginning:	12/31/2018		Beginning:				
Ending:	12/31/2021		Ending:				
Project is (choose	one):						_
NOTE: Revision					project including scop	pe of pro	oject, funding restrictions,
[] New				[]	Revision		
[X] Continu	ation, without revisi	on(s)	[]		Continuation, with	$\mathbf{u}(\mathbf{s})$	
Lead staff membe	er:	Contact info	rmation:		ase indicate your role	e(s) in th	nis project:
	rri Cottengim	Division:	DRH	[X]	Project officer	[]	Technical monitor
<u></u>	c ccg			– []	Principal	[]	Investigator
User ID: WS	SH2	Telephone:	770.488.4290	_	investigator		J
CSCI ID. WC							
Scientific Eth	l of the activities with		F74 SIGNED to con	_ []	Consultant to generalizable know	[] wledge (i	
Scientific Eth 1. Are any or all	l of the activities with	nin this project DE:					
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Scientific Eth 1. Are any or all [] YE If YES, list to [] Reserved 2. Is this CDC [] Reserved [] [] 3. If RESEARG subjects produced a. [] NO,	l of the activities with ES [X] hose activities which project research or project research eck one: Human subject Human subject Human subject CH involving human tection?	nin this project DE: NO n are research: public health practions involved s not involved n subjects, has the	SIGNED to constitute (check all [X] Pub Check [] [] [] project or research. []	that a lic hea ck all t En Pro	to generalizable known pply)? Ith practice hat apply: nergency Response ogram evaluation activities been review.	[X] [] red by the cover by	Surveillance Other (please explain) the CDC IRB for human
Scientific Eth 1. Are any or all [] YE If YES, list to [] Reserved. 2. Is this CDC [] Reserved. [] [] [] 3. If RESEARC subjects produced in [] NO, b. [] NO,	l of the activities with ES [X] hose activities which project research or p search eck one: Human subject Human subject CH involving human tection? New project, not yet	nin this project DE: NO n are research: public health praces involved s not involved n subjects, has the t reviewed t ready to submit	SIGNED to constitute (check all [X] Pub Check [] [] [] project or research. []	that a lic hea ck all t En Pro	to generalizable known pply)? Ith practice hat apply: nergency Response ogram evaluation activities been reviews, Reviewed and appropriate to general preserve to gene	[X] [] red by the cover by	Surveillance Other (please explain) the CDC IRB for human
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Form 684R_NR (revised January 2003)

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Name					Role (proconsulta	•	Scientific ethics number Prin		
	Ca	rri Cott	engim						
				EARCH PRO ns 4-6, OTHI					(as identified in 45CFR46.101),
4.		Does t	he propo	sed research	involve p	orisoners'	?		
	[]	YES		If YES, this	s researc	h cannot l	be exemp	ed and must be reviewed by	y an IRB (skip to question 7).
	[]	NO							
5.		he prop apply)?		arch involve	fetuses, p	pregnant	women, o	r human in vitro fertilizatio	n as targets (such that Subpart B
	[]	YES		If YES, th question 7		rch cann	ot be exe	mpted and must be revie	ewed by an IRB (skip to
	[]	NO							
<u>Ed</u>	ucationa	al Resea	<u>rch</u>						
	6.1	norma	l educati	onal practice	s (e.g., re	search or	regular :	and special education strate	s, AND does the research involve gies or research on the room management methods)?
		[]	YES	,P	[]	NO		1, 2	
Re	search I		g Surveys	, Interview P		es (includi	ing Focus	groups), Observation of Pu	blic Behavior, or Educational
Te	<u>sts</u>		-				_		·
	6.2			ch use educa observation o				ostic, aptitude, achievemen	t), survey procedures, interview
		[]	YES		[]	NO		If NO skip 6.3	
		Will c	hildren (<	<18 years of a	ige) be re	esearch su	bjects?		
		[]	YES	If YES, thi	is researc	ch cannot	be exemp	ted and must be reviewed b	y an IRB (skip to item 7)
		[]	NO						
		6.2.1						nanner that human subjects iked to the subjects;	s can be identified <u>directly or</u>
			[]	YES		[]	NO		
		6.2.2	the subj employa subjects	ects at risk o ability or rep s' (or relative	f crimina utation? s' or asso	al or civil (Example ociates') p	liability, o es here ma ossible su	or be damaging to the subje ny include: the collection of	sensitive data regarding the iminal history or intent, medical
			[]	YES		[]	NO		
	6.3		lures, or			behavior		esearch is not exempt under	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		volve hui	nan subje	ects that a	re elected or appointed pub	olic officials or candidates for	
			[]	YES		[]	NO		
		6.3.2	informa	tion will be r n only in the	naintain	ed throug	hout the i	ion that confidentiality of the search and thereafter? (Nonce of Confidentiality has be	ote: CDC can use this exemption
			[]	YES		[]	NO		
Ex	isting Da	ata Whi	ch Is Pub	licly Availab	<u>le or Uni</u>	<u>dentifiabl</u>	<u>e</u>		
	6.4							f existing* data, documents, e the study begins)?	, records, pathological or
		[]	YES		[]	NO		If NO skip to 7	
		6.4.1	Is this n	naterial or in	formatio	n publicly	y availabl	e?	
			[]	YES		[]	NO		

	Tracking	NO.	<u>tbd</u>
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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))				

- 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 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 (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.
 - a. The purpose of the Sudden Death in the Young (SDY) Case Registry is to improve and standardize case ascertainment so that funded jurisdictions can improve their understanding of the descriptive epidemiology of SDY; including the incidence, and risk factors. These data will be used to inform prevention strategies as well as a recommendations and best practices for national surveillance of SDY.

NHLBI and NINDS will work with the CDC via an Interagency Agreement and an Inter-Departmental Delegation of Authority to continue a prospective, population-based SDY Case registry. This Registry is the only prospective, population-based data system compiled for the comprehensive evaluation of unexpected deaths including sudden cardiac death in the young and sudden unexpected death in epilepsy in the United States. The SDY Registry includes data from death certificates, medical records, death scene investigations, and pathology reports.

The data collection activities will be supported through a contracted data coordinating center.

- b. Project status selection: This is non-research, public health surveillance. The purpose of the activity is to establish the incidence of SDY in funded jurisdictions as part of core public health function of Child Death Reviews. The system is designed to monitor the frequency of occurrence and distribution of disease or SDY in the population. Data generated by these systems are used to manage public health program; intended benefits of the project are primarily for the participants.
- c. CDC staff will not be engaged in research. CDC will provide technical assistance, receive de-identified data for quality assurance activities and surveillance reports. CDC will co-author manuscripts, but will not be first author on any SDY Case Registry manuscript describing major findings.
- 8. Please list the primary project site and all collaborating site(s).

Explanation 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
Linda Hannon-Hall - PUBLIC HEALTH ADVISOR	07/17/2018	[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: Approved

Carrie Shapiro-Mendoza - Health Scientist	07/17/2018	[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:
Karen Pazol - Deputy ADS Division ADS	07/19/2018	[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
Joan Redmond Leonard - PUBLIC HEALTH ANALYST	07/20/2018	 [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		Comments: