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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:

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

Date submitted:	06/19/2018								
Title of Project:	Million Hearts Hospital or Hea	alth System	designati	on program					
Dates for project per	iod:	Dates for f	funding (if	f applicable):					
Beginning: 06/03/2019		Beginni	Beginning:						
Ending: 12/31/2021		Ending	:						
	e): s used below, refers to any substan CDC staff member, determination o			e project including scop	pe of pro	oject, funding resti	rictions,		
[X] New			[]	Revision					
[] Continuatio	[] Continuation, without revision(s)			[] Continuation, with r			revision(s)		
Lead staff member:	Contact inf	formation:	Ple	ease indicate your role	(s) in th	us project:			
	a Barnett Division:	DHDSP	[]	Project officer	[]	Technical mor	nitor		
User ID: NMA4	Telephone	404-498-	[]	Principal investigator	[]	Investigator			
Scientific Ethics	number: Mailstop:	F-73	[]	Consultant	[X]	Other (please	explain)		
					Pro	ject Coordinator			
	e activities which are research: ject research or public health pr	actice (chec	k all that a	apply)?					
[] Resear	rch	[X]	Public hea	alth practice					
Check	one:		Check all	that apply:					
[]	Human subjects involved		[] E I	mergency Response	[]	Surveillance			
[]	Human subjects not involved		[] Pr	rogram evaluation	[X]	re pr us M Hi st to ca ul	xplain) roject to coognize ractices sing illion earts rategies improve ardiovasc ar utcomes.		
subjects protect	involving human subjects, has tl ion? v project, not yet reviewed	ne project oi d.		activities been review 5, Reviewed and appro	-		human		

- YES, Reviewed and approved by CDC
- b. [] NO, Existing project, not ready to submit

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5. Does wou [] [] Educatio 6.1 Research <u>Research</u>	s the prop ild apply)' YES NO	?	If YES, this res		women, or	human in vitro fertilizatio	n as targets (such that Subnart R
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6.1 <u>Research</u> <u>Tests</u>	onal Resea						
6.1 <u>Research</u> <u>Tests</u>	onal Resea						
<u>Research</u> Tests	T /1 •						
<u>[ests</u>	norm	al education	nal practices (e.g.	., research or	n regular a	nd special education strate	s, AND does the research involve egies or research on the room management methods)?
<u> Fests</u>	[]	YES	[]	-			
	<u>ı Involvin</u>	g Surveys, I	Interview Proced	lures <u>(includi</u>	ing Focus	groups), Observation of Pu	iblic Behavior, or Educational
6.2			h use educational servation of publ			ostic, aptitude, achievemen	t), survey procedures, interview
	[]	YES	[]	NO		If NO skip 6.3	
	Will c	hildren (<1	8 years of age) b	e research su	bjects?		
	[]	YES	If YES, this rese	earch cannot	be exempt	ted and must be reviewed b	y an IRB (skip to item 7)
	[]	NO					
	6.2.1					nanner that human subjects ked to the subjects;	s can be identified <u>directly or</u>
		[]	YES	[]	NO		
	6.2.2	place the employat subjects'	subjects at risk o bility or reputatio (or relatives' or a	of criminal of on? (Example associates') p	: civil liabi es here ma ossible sub	lity, or be damaging to the y include: the collection of	h setting have the potential to subjects' financial standing, sensitive data regarding the iminal history or intent, medical rmation).
		[]	YES	[]	NO		
6.3							t), survey procedures, interview r paragraph 6.2 of this section:
	[]	YES	[]	NO		If NO skip to 6.4	
	6.3.1	Will this public off		human subje	ects that a	re elected or appointed pub	olic officials or candidates for
		[]	YES	[]	NO		
	6.3.2	informati	on will be mainta only in the case v	ained throug	hout the r	on that confidentiality of tl esearch and thereafter? (N ice of Confidentiality has b	ote: CDC can use this exemption
		[]	YES	[]	NO		

Form 684R_NR (revised January 2003)

Existing Data Which Is Publicly Available or Unidentifiable

6.4 Does this research involve only the collection or study of existing* data, documents, records, pathological or diagnostic specimens? (* 'existing' means existing before the study begins)?

[]	YES	[]	NO	If NO skip to 7
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- 6.4.1 Is this material or information publicly available?
 - [] 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?

(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 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.

Million Hearts is a national initiative to prevent 1 million heart attacks and strokes by 2022.

The purpose of the Million Hearts Hospital or Health System Designation program is to recognize institutions working to systematically improve the cardiovascular health of the population and communities they serve through a focus on Keeping People Health, Optimizing Care, Improving Outcomes for Priority Populations, and Innovating for Health. Hospitals and health systems will self-nominate through an online portal by doing one of the following:

a.)Commit to implement policies that promote cardiovascular health

b.)Provide documentation of current policies that they are implementing in order to promote cardiovascular health

c.) Provide documentation of current policies that they are implementing in order to promote cardiovascular health and provide data that shows positive health outcomes as a result of those policies being implemented

Nominations will be scored using predetermined criteria and a scoring rubric and reviewed by an expert panel of judges. Recognized designees will participate in the development of a success story product to be shared broadly with the public health and clinical audiences.

CDC staff members have developed the application criteria, data collection instruments, and selection criteria. CDC staff will consult on data storage and confidentiality, any data analyses conducted, and record review protocols.

The status for this project is public health practice. The objective is to identify and recognize hospitals and health systems that work to improve cardiovascular health in a narrative or documentary format. No individual patient level data will be collected, only practice level summary data. No generalizable knowledge will emerge from this project. A manuscript is not planned at this time. Selected hospitals and health systems will be recognized through web-based platforms.

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
Jessica Barnett - Assistant/Program Analyst	06/22/2018	 [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 [] Research involving human subjects, CDC investigators, not exempt [] Local IRB [] CDC Exemption [] CDC IRB
staff member completing this form		<u>Comments:</u>
Judith Hannan - Million Hearts Advisor	06/29/2018	 [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		<u>Comments:</u> looks good
Mary George - SENIOR MEDICAL OFFICER	06/29/2018	 [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		<u>Comments:</u> in question 2, "practices" likely should be "hospitals", but this doesn't change the overall document.
Joan Redmond Leonard - PUBLIC HEALTH ANALYST	07/01/2018	 [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 [] Research involving human subjects, CDC investigators, not exempt [] Local IRB [] CDC Exemption [] CDC IRB
CUC ADS, Deputy ADS, or Human Subjects Contact		Comments: