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: TBD/TBD

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

Practices in Infant Nutrition and Care – 2020 and 2022 mPINC su         Dates for project period:       Dates for funding (if applicable):		n the U.S. (Maternity					
	Assessment and Monitoring of Breastfeeding-related Maternity Care Practices in Practices in Infant Nutrition and Care – 2020 and 2022 mPINC survey)						
Beginning: 09/25/2017 Beginning:		_					
<b>Ending:</b> 12/31/2022 <b>Ending:</b>							
Project is (choose one):         NOTE: Revision, as used below, refers to any substantive change made to the project including personnel, role of CDC staff member, determination of research status, etc.         [X]       New       []       Revision	ng scope of proje	ect, funding restrictions,					
	, with revision(s	s)					
	-						
Lead staff member:     Contact information:     Please indicate you       Name:     Daurice Grosspiklaus     Division:     DNPAQ     []     Project officiality		s project: Technical monitor					
Name:       Daurice Grossniklaus       Division:       DNPAO       []       Project offi         User ID:       DTG3       Telephone:       770-488-5249       investigator	[]	Investigator					
Scientific Ethics number: 1761 Mailstop: F77 [] Consultant	[]	Other (please explain)					
[]       YES       [X]       NO         If YES, list those activities which are research:							
2. Is this CDC project research or public health practice (check all that apply)?							
[]   Research   [X]   Public health practice							
Check one: Check all that apply:							
[]Human subjects involved[]Emergency Respo[]Human subjects not involved[]Program evaluation		Surveillance Other (please explain)					
3. If RESEARCH involving human subjects, has the project or research activities been r subjects protection?	reviewed by the	e CDC IRB for human					
a. [] NO, New project, not yet reviewed d. [] YES, Reviewed and	l approved by C	CDC					
b. [] NO, Existing project, not ready to submit If YES, please list	f YES, please list protocol number_and						
c. [] NO, Submitted for approval expiration d	late						
e. [] NO, RESEARCH, n required)	no CDC investig	gators (CDC IRB not					
f. [] N/A (Not Applicable	le)						

## Tracking NO. <u>TBD/TBD</u>

\_\_\_\_\_

Name			Role (project officer, investigator, consultant, etc.)						Scientific ethics number Prin	
Daurice Grossnikla									ossniklau	1761
				ARCH PROJE s 4-6, OTHER					CARCH (as ident	ified in 45CFR46.101),
4.		Does th	ie propos	ed research inv	olve pri	soners?	-			
[	]	YES		If YES, this re	search c	annot be	e exempte	d and must be revi	ewed by an IRB	(skip to question 7).
[	]	NO								
5. Does the proposed research involve fetuses, pregnant women, or human in vitro fertilization as targets (su would apply)?									ets (such that Subpart B	
[	[]       YES       If YES, this research cannot be exempted and must be reviewed by an IRB (skip t question 7).							an IRB (skip to		
[	]	NO								
<u>Educa</u>	tional	Resear	<u>ch</u>							
6.	1	normal	educatio	nal practices (e	e.g., rese	arch on 1	regular a	nd special educatio	n strategies or re	oes the research involve esearch on the nagement methods)?
		[]	YES		[]	NO				
	ch Inv	volving	Surveys,	Interview Proc	edures (	includin	g Focus g	roups), Observatio	on of Public Beha	vior, or Educational
<u>Tests</u>	•	XX/211 41.5	•	h		( <b></b>		atio on titudo o shi	·····	· · · · · · · · · · · · · · · · · · ·
6.				n use education oservation of p			e, diagno	suc, aptitude, achi	evement), survey	v procedures, interview
		[]	YES		[]	NO		If NO skip 6.3		
		Will ch	ildren (<1	18 years of age	) be rese	arch sub	jects?			
		[]	YES	If YES, this r	esearch	cannot b	e exempt	ed and must be rev	iewed by an IRE	(skip to item 7)
		[]	NO							
								anner that human a ked to the subjects;		dentified <u>directly or</u>
			[]	YES		[]	NO			
			place the employal subjects'	subjects at ris bility or reputa (or relatives' o	k of crin tion? (E or associa	ninal or o xamples ates') pos	civil liabi here may ssible sub	lity, or be damaging y include: the collec	g to the subjects' ction of sensitive ality, criminal hi	nave the potential to financial standing, data regarding the story or intent, medical
			[]	YES		[]	NO			
6.		proced	ures, or o			havior b				<pre>procedures, interview ph 6.2 of this section:</pre>
			YES		[]	NO		If NO skip to 6.4		
			public of	fice?	ve huma			e elected or appoin	ted public officia	als or candidates for
			[]	YES		[]	NO			
	1		informat	ion will be mai only in the cas	ntained	through	out the re	on that confidential search and thereaf ce of Confidentialit	ter? (Note: CDC	can use this exemption
			[]	YES		[]	NO			
	0			cly Available o						
6.		diagnos	stic specir	nens? (* 'existi	ng' mea	ns existiı		existing* data, doc the study begins)?	uments, records,	pathological or
			YES		[]	NO		If NO skip to 7		
				aterial or infor	mation <sub>]</sub>			?		
			[]	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 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.

7a. The purpose of this project is to track the maternity care practices that support breastfeeding in hospitals. Since 2007, CDC has administered the national survey of maternity care practices related to breastfeeding, known as the Maternity Practices in Infant Nutrition and Care (mPINC) Survey on a biennial basis in order to monitor and examine changes in these practices over time. The mPINC survey uses a census design and is administered to every facility in the US that routinely provides maternity care services. A staff person completes the mPINC survey on behalf of his or her institution in their capacity as the person most knowledgeable about the relevant practices. The first survey in 2007 established a measure of the prevalence of specific breastfeeding-related maternity care practices at maternity care facilities across the United States and Territories at that point in time and the extent to which these practices vary by state. In addition to providing prevalence measures at subsequent points in time, biennial analyses of data from the 2009 - 2015 surveys provide important information about changes and trends in practices since the initial baseline assessment. The current RFTP is intended to continue work similar to that of the previous surveys. The 2020 mPINC survey is intended to be completed within the scope of the Information Collection Review (ICR) currently being submitted to cover the mPINC 2018 administration or under a new ICR to be submitted prior to the 2020 mPINC administration. Access to data, processes, surveys, and reports on previous mPINC surveys is available at www.cdc.gov/mpinc and http://www.reginfo.gov/ (search "OMB Control Number: 0920-0743" for complete records of 2007-2016 OMB activity related to this project). CDC staff monitor the contract and conduct analyses on data from the survey. Analyzed data will be linked to hospital identification number, hospital name and hospital address in order for the Contractor conducting the survey to disseminate hard copy and/or electronic versions of hospital-specific benchmark reports. The web survey will include an item for the survey recipient's permission to retain their title, hospital affiliation, and hospital email address so that an electronic copy of the hospital-specific benchmark report can be emailed to the survey recipient in lieu of a paper copy.

7b. This project is public health practice. The data collected are part of an ongoing surveillance system tracking hospital practices that support breastfeeding. Although the data has the potential to be used for research, the primary purpose of the data collection is for surveillance and monitoring. No human subject research will be conducted under this request. If human subject research is conducted a new Research Determination will be submitted.

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
Latetia Moore Freeman - Deputy Associate Director for Sc	02/03/2017	<ul> <li>[X] Public health practice</li> <li>[] Research not involving human subjects</li> <li>[] Research involving human subjects, no CDC investigators</li> <li>[] Research involving human subjects, CDC investigators, exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Local IRB</li> <li>[] CDC Exemption</li> <li>[] CDC IRB</li> </ul>
staff member completing this form		<u>Comments:</u> LV Moore reviewed and approves

Deborah Galuska - ASSOCIATE DIRECTOR SCIENCE Team Lead	02/03/2017	<ul> <li>[X] Public health practice</li> <li>[] Research not involving human subjects</li> <li>[] Research involving human subjects, no CDC investigators</li> <li>[] Research involving human subjects, CDC investigators, exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Local IRB</li> <li>[] CDC Exemption</li> <li>[] CDC IRB</li> </ul>
Team Leau		<u>Comments:</u>
Deborah Galuska - ASSOCIATE DIRECTOR SCIENCE	02/03/2017	<ul> <li>[X] Public health practice</li> <li>[] Research not involving human subjects</li> <li>[] Research involving human subjects, no CDC investigators</li> <li>[] Research involving human subjects, CDC investigators, exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Local IRB</li> <li>[] CDC Exemption</li> <li>[] CDC IRB</li> </ul>
Division ADS		Comments:
Joan Redmond Leonard - PUBLIC HEALTH ANALYST	02/21/2017	<ul> <li>[X] Public health practice</li> <li>[] Research not involving human subjects</li> <li>[] Research involving human subjects, no CDC investigators</li> <li>[] Research involving human subjects, CDC investigators, exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Research involving human subjects, CDC investigators, not exempt</li> <li>[] Local IRB</li> <li>[] CDC Exemption</li> <li>[] CDC IRB</li> </ul>
CUC ADS, Deputy ADS, or Human Subjects Contact		<u>Comments:</u>