

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.

		Tracking Number: ClOSP 23 (Use PGO number if cooperative agreemen	
Date submitted: 01/16/2013			
Title of Project: Comprehensive Cancer C	ontrol Branch Management Infor	mation System	
Dates for project period: Beginning: 01/01/2013 Ending: 12/31/2017 Project is (choose one): NOTE: Revision, as used below, refers to an personnel, role of CDC staff member, determined and the personnel of CDC staff member.		pplicable):	ctions,
[] New [] Continuation, without revision(s)		evision ontinuation, with revision(s)	
Lead staff member:Name:CONOLA STEELEUser ID:CKS9Scientific Ethics number:6803	Contact information:Division:DCPCTelephone:770-488-4261Mailstop:K57	Please indicate your role(s) in this proje [] Project officer [X] [] Principal investigator [] [] Consultant []	
1. Are any or all of the activities within this properties [1] [] YES [X] NO If YES, list those activities which are researed		generalizable knowledge (i.e., research)?	
2. Is this CDC project research or public heal [] Research <i>Check one:</i> [] Human subjects involved [] Human subjects <u>not</u> involved	[X] Public health practice Check all that apply: [] Emergency Response		Reporting of progress reports and submission of continuation applications
 3. If RESEARCH involving human subjects, l protection? a. [] NO, New project, not yet revie b. [] NO, Existing project, not ready c. [] NO, Submitted for approval 	wed d. []	YES, Reviewed and approved by CDC If YES, please list protocol numb expiration date NO, RESEARCH, no CDC investigators	n subjects er _ and
If RESEARCH, list any other CDC staff invo Name	olved in this project, please include Role (project officer, investigato		
	consultant, etc.)	, Scentine enity numbe	

IF YOU THINK THE RESEARCH PROJECT MIGHT QUALIFY AS EXEMPT RESEARCH (as identified in 45CFR46.101), PLEASE ANSWER questions 4-6, OTHERWISE SKIP TO question 7.

4. Does the proposed research involve prisoners?

- YES If YES, this research cannot be exempted and must be reviewed by an IRB (skip to question 7).
- [] NO
- 5. Does the proposed research involve fetuses, pregnant women, or human <u>in vitro</u> fertilization as targets (such that Subpart B would apply)?
 - YES If YES, this research cannot be exempted and must be reviewed by an IRB (skip to question 7).

[] NO

Educational Research

6.2

6.1 Is this research conducted in established or commonly accepted educational settings, <u>AND</u> 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)?

[] YES [] NO

Research Involving Surveys, Interview Procedures (including Focus groups), Observation of Public Behavior, or Educational Tests

Will this research use educational tests (cognitive, diagnostic, aptitude, achievement), survey procedures, interview procedures or observation of public behavior?

[] YES [] NO If NO skip to 6.3

Will children (<18 years of age) be research subjects?

- [] YES If YES, this research cannot be exempted and must be reviewed by an IRB (skip to item 7)
- [] **NO**
- 6.2.1 Is the information obtained recorded in such a manner that human subjects can be identified <u>directly or indirectly</u> through identifiers (such as a code) linked to the subjects;

[] YES [] NO

- 6.2.2 Will any disclosure of the human subjects' responses outside of the research setting have the potential to place the subjects at risk of criminal or civil liability, or be damaging to the subjects' financial standing, employability or reputation? (Examples here may include: the collection of sensitive data regarding the subjects' (or relatives' or associates') possible substance abuse, sexuality, criminal history or intent, medical or psychological condition, financial status, or similarly compromising information).
 - [] YES [] NO
- 6.3 Will this research use educational tests (cognitive, diagnostic, aptitude, achievement), survey procedures, interview procedures, or observation of public behavior but the research is not exempt under paragraph 6.2 of this section: [] YES [] NO If NO skip to 6.4

 - 6.3.1 Will this research involve human subjects that are elected or appointed public officials or candidates for public office?
 [] YES [] NO
 - 6.3.2 Does federal statute(s) require(s) without exception that confidentiality of the personally identifiable information will be maintained throughout the research and thereafter? (Note: CDC can use this exemption criterion only in the case where a 308(d) Assurance of Confidentiality has been obtained to cover the research).

[] YES [] NO

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

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)
- [] 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.

The purpose of this project is to continue using an electronic information system (Management Information System [MIS] for Comprehensive Cancer Control Programs, OMB No. 0920-0841) to maintain grantee-specific information for the National Comprehensive Cancer Control Program (NCCCP). Since January 2010, NCCCP awardees have submitted progress and activity information to CDC twice per year using this system. New cooperative agreements were awarded to all NCCCP programs in 2012, which place emphasis on policy and environmental approaches to cancer prevention and control. Minor changes to the existing data elements will be implemented to reflect the FOA's new performance requirements. Thirteen of the 65 NCCCP awardees received additional funding for related but distinct cooperative agreements that aim to accelerate the development of policy and environmental approaches to cancer control. Similar semi-annual progress reports are required to monitor activities under this demonstration program. CDC proposes to use the same MIS-based methodology for all reporting. This project is not research. The MIS data collection instrument does not collect personally identifiable information or sensitive data.

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
Conola Steele - MEDICAL EPIDEMIOLOGIST	01/16/2013	 [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		Comments:
Phaeydra Brown - Assistant Branch Chief	01/16/2013	 [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		Comments: 684 for CCC MIS

Tracking NO.	CIOSP 2305

Cheryll Thomas - EPIDEMIOLOGIST Division ADS	01/16/2013	 [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 Comments:
Joan Redmond Leonard - PUBLIC HEALTH ANALYST ADS, Deputy ADS, or Human Subjects Contact	01/16/2013	 [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

List of Grantees

Grantee #

<u>Grantee Name</u>