

## 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:** 

N/A/N/A/2202000200

| ate submit                                       | ted:  | 12/13/2012   |  |  |   |   |                           |  |
|--|---|--|--|--|---|---|---------------------------|--|
| itle of Proj                                     |   |  | of Cancer Re   | anistries – NP   | CR Pr   | ogram Evaluation In   | strumer                   | nt (PFI)   |
| Ū  |   | •  |  | Dates for fund   |   |   | <u>struirier</u>          | II (I LI)  |
| ates for pro                                     |   | 09/30/2012   |  | Beginning:   |   | аррисавіе).   |                           |  |
| Beginning: Ending:                               |   | 09/30/2012   | Ending:  |  |   |   | _                         |  |
| Linding.   |   | 09/29/2017   |  | Liums.   |   |   |                           | _  |
| roject is (cl                                    | hoose on  | e):  |  |  |   |   |                           |  |
|  |   | s used below, refers to<br>CDC staff member, det   |  |  |   | project including scop  | pe of pro                 | oject, funding restrictions                              |
| [] Nev   | w   |  |  |  | []  | Revision  |                           |  |
| [] Con   | ntinuatio   | on, without revision(s   | )  |  | [X]   | Continuation, with  | revision                  | $(\mathbf{s})$   |
| ead staff m                                      | ember:  |  | Contact info   | rmation:   | Ple   | ase indicate your role  | e(s) in th                | nis project:   |
| Name:  | Netta   | Apedoe   | Division:  | DCPC   | [X]   | Project officer   | [X]                       | <b>Technical monitor</b>                                 |
|  |   |  |  |  | []  | Principal   | []                        | Investigator   |
| User ID:   | ize6  |  | Telephone:   | 770-488-4570   | )   | investigator  |                           |  |
|  |   |  | . =  |  |   |   |                           |  |
| . Are any  | or all of   | number:  the activities within the [X] NO  | )  | K69 SIGNED to con  | []<br>ntribute  | Consultant to generalizable know  | []<br>wledge (i           | Other (please explain                                    |
| . Are any  | or all of   | the activities within the  | nis project DE   |  |   |   |                           |  |
| Are any  If YES,                                 | or all of YES list thos   | the activities within the [X] NC   | nis project DE:  research:   | SIGNED to con  | ntribute  | to generalizable know   |                           |  |
| Are any  If YES,                                 | or all of YES list thos   | the activities within the [X] NC se activities which are specified research or publications.   | nis project DE:  research:   | SIGNED to con  | ntribute  | to generalizable know   |                           |  |
| Are any  [ ]  If YES,  Is this (                 | or all of YES List those  | the activities within the [X] NO se activities which are object research or publich  | nis project DE:  research:   | SIGNED to conception of the co | ntribute<br>I that a<br>Dlic hea                          | to generalizable know   |                           |  |
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| Are any  [ ]  If YES,  Is this (                 | or all of YES list thos CDC pro Resea                                       | the activities within the [X] NC se activities which are spect research or public rch  | nis project DE:  research:  ic health prace  | SIGNED to constitute (check all [X] Pul  | ntribute  I that a  Dlic hea  eck all t                   | to generalizable know pply)? lth practice that apply:   | vledge (i                 | .e., research)?  |
| If YES,  Is this (                               | or all of<br>YES<br>list thos<br>CDC pro<br>Resea<br>Check                  | the activities within the [X] NC se activities which are object research or published.  Human subjects involving human subjects no involving human subjects  | nis project DE:  research:  ic health prace  volved t involved   | SIGNED to constitute (check all [X] Pull Che   | ntribute<br>I that a<br>Dlic hea<br>eck all t<br>En<br>Pr | pply)?  charactice  charapply: nergency Response ogram evaluation   | vledge (i                 | .e., research)?  |
| If RESI subjects                                 | cor all of YES List thos CDC pro Resea Check [] [] EARCH s protect          | the activities within the [X] NC se activities which are object research or published.  Human subjects involving human subjects no involving human subjects  | nis project DE.  research:  ic health praction of the colved to involved to involve to invol | SIGNED to constitute (check all [X] Pull Che   | I that a blic hea eck all t Pr                            | pply)?  charactice  charapply: nergency Response ogram evaluation   | [X]                       | Surveillance Other (please explain) ne CDC IRB for human |
| Is this C  If RESI subjects a. []                | r or all of YES List thos CDC pro Resear Check [] [] EARCH s protect NO, Ne | the activities within the [X] NO se activities which are object research or public one:  Human subjects involving human subjects no involving human subjects involving human subjects involving human subjects involving human subjects no involving human subjects no involving human subjects involving human subjects no involving human subjects invol | nis project DE:  research:  ic health praction of the control of t | ctice (check all [X] Pul [X] [X] [X] project or res  | I that a Dic hea eck all t  En Pr search                  | pply)?  Ith practice  that apply: nergency Response ogram evaluation activities been review   | [X] [] red by the oved by | Surveillance Other (please explain) ne CDC IRB for human |
| If YES,  Is this C  If RESI subjects a. [] b. [] | CDC pro Resea Check [] EARCH s protect NO, Ne NO, Ex                        | the activities within the [X] NO se activities which are oject research or public one:  Human subjects involving human subtion?  w project, not yet rev  | nis project DE:  research:  ic health praction of the colored of t | ctice (check all [X] Pul [X] [X] [X] project or res  | I that a Dic hea eck all t  En Pr search                  | pply)?  olth practice  that apply: nergency Response ogram evaluation activities been review , Reviewed and appro-  | [X] [] red by the oved by | Surveillance Other (please explain) ne CDC IRB for human |
| If YES,  Is this C  If RESI subjects a. [] b. [] | CDC pro Resea Check [] EARCH s protect NO, Ne NO, Ex                        | the activities within the [X] NO se activities which are object research or public one:  Human subjects involving human subjects no involving human subjects we project, not yet revisiting project, not rea   | nis project DE:  research:  ic health praction of the colored of t | ctice (check all [X] Pul [X] [X] [X] project or res  | I that a blic hea eck all t  En  Pr  Search:              | pply)?  Ith practice  That apply:  Inergency Response  Togram evaluation  activities been review  A Reviewed and approf  Tyes, please list pro  expiration date | [X] [] red by theoved by  | Surveillance Other (please explain) ne CDC IRB for human |

Tracking NO. <u>N/A/N/A/2202000200</u>

|             | Na       | ame  |  |                                 |              | Role (pro<br>consulta |                   | cer, investigator,   | Scientific ethics<br>number Prin  |
|-------------|----------|--|--|---------------------------------|--------------|-----------------------|-------------------|--|---|
|             | Ne       | etta Ape   | doe  |                                 |              |                       |                   |  |   |
|             |          |  |  | EARCH PRO                       |              |                       |                   |  | I (as identified in 45CFR46.101),   |
| 1.          |          |  | •  | sed research                    |              |                       | •                 |  |   |
|             | []       | YES  |  |                                 | -            |                       |                   | ted and must be reviewed b   | y an IRB (skip to question 7).  |
|             | []       | NO   |  | ,                               |              |                       |                   |  |   |
| 5.          | Does     |  |  | arch involve                    | fetuses, p   | oregnant v            | women, o          | r human in vitro fertilizatio  | on as targets (such that Subpart B  |
|             | []       | YES  |  | If YES, th<br>question 7        |              | rch cann              | ot be exe         | empted and must be revi  | ewed by an IRB (skip to   |
|             | []       | NO   |  |                                 |              |                       |                   |  |   |
| Edi         | ucation  | al Resea   | <u>rch</u>   |                                 |              |                       |                   |  |   |
|             | 6.1      | norma  | ıl educati   | onal practice                   | es (e.g., re | search or             | regular           | and special education strat  | gs, AND does the research involve egies or research on the croom management methods)? |
|             |          | []   | YES  | •                               | []           | NO                    |                   | • /  | ,   |
| <u>Re</u> s | search l |  |  | <u>, Interview</u> F            |              |                       | ng Focus          | groups), Observation of Pu   | ublic Behavior, or Educational  |
| Гes         |          |  | -  | •                               |              | •                     |                   | <u> </u>   |   |
|             | 6.2      |  |  | ch use educa<br>observation o   |              |                       |                   | ostic, aptitude, achievemei  | nt), survey procedures, interview   |
|             |          | []   | YES  |                                 | []           | NO                    |                   | If NO skip 6.3   |   |
|             |          | Will children (<18 years of age) be research subjects? |  |                                 |              |                       |                   |  |   |
|             |          | []   | YES  | If YES, th                      | is researc   | h cannot              | be exemp          | ted and must be reviewed   | by an IRB (skip to item 7)  |
|             |          | []   | NO   |                                 |              |                       |                   |  |   |
|             |          | 6.2.1  |  |                                 |              |                       |                   | nanner that human subject<br>nked to the subjects;   | s can be identified <u>directly or</u>  |
|             |          |  | []   | YES                             |              | []                    | NO                |  |   |
|             |          | 6.2.2  | .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      | proced   | dures, or  |                                 | of public    | behavior              |                   | esearch is not exempt unde   | nt), survey procedures, interview r paragraph 6.2 of this section:                    |
|             |          | []   | YES  |                                 | []           | NO                    |                   | If NO skip to 6.4  |   |
|             |          | 6.3.1  | public o   | office?                         | volve hur    |                       |                   | re elected or appointed pu   | blic officials or candidates for  |
|             |          |  | []   | YES                             |              | []                    | NO                |  |   |
|             |          | 6.3.2  | informa  | tion will be a<br>n only in the | maintaine    | ed through            | hout the <b>i</b> | ion that confidentiality of t<br>research and thereafter? (N<br>nce of Confidentiality has b | Note: CDC can use this exemption  |
|             |          |  | []   | YES                             |              | []                    | NO                |  |   |
| Exi         | sting D  | ata Whi  | ch Is Pub  | licly Availab                   | le or Uni    | <u>dentifiabl</u>     | <u>e</u>          |  |   |
|             | 6.4      |  |  |                                 |              |                       |                   | f existing* data, documents e the study begins)?   | s, records, pathological or   |
|             |          | []   | YES  |                                 | []           | NO                    |                   | If NO skip to 7  |   |
|             |          | 6.4.1  | Is this n  | naterial or in                  | formatio     | n publicly            | availabl          | e?   |   |
|             |          |  | []   | YES                             |              | []                    | NO                |  |   |

Form 684R\_NR (revised January 2003)

|       | T  |     |   |  |  |  |
|-------|--|-----|---|--|--|--|
| 6.4.2 |  |     | information recorded in such a manner by the investigator that the subjects cannot be 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  |  |  |  |
|       | r 1  | NO  | (there are identifiers (including codes))   |  |  |  |

- 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.

This project is a public health practice because it is designed to collect data about CDC funded central cancer registries' operations to evaluate and monitor the success of these programs at meeting the requirements of the NPCR Standards. The PEI project is intended to help answer questions about how well the registries are doing and provide some data about what areas they could improve. The project does not attempt to answer any research questions and the knowledge obtained through this project is not generalizable. The project also does not collect any personal information and the previous human subject determination found this project to be public health practice.

The annual collection of performance indicator data from state and territorial programs funded through the National Program of Cancer Registries (NPCR) began 17 years ago and was changed to biennial collection in 2009. The data collected is limited to registry operations information and does not include individual record level data. The data include information on registry staffing levels, the status of cancer registry legislation in the state, the type and scope of advanced surveillance activities, case sharing agreements with neighboring states, the number and type of reporting facilities, computer infrastructure, etc. The information collection allows CDC to provide routine feedback to grantees based on their data submissions, to tailor technical assistance as needed, and to support program planning, surveillance and secondary data analysis activities. The evaluation data will be used by CDC to help monitor progress in meeting NPCR standards and assisting CDC project officers with providing technical assistance to the states.

This Human Subject determination is a component of an OMB application. The survey instrument has been revised with updated NPCR Program Standards, questions that are no longer relevant have been deleted and new requirements around Meaningful Use have been added.

This project will be administered through a contract. CDC Staff will manage and monitor the contract, review and make suggestions for revising the survey, have monthly calls with the contractor, review and monitor the budget and work plan.

8. Please list the primary project site and all collaborating site(s).

|           |                          | Site Name                | Site Location | Assurance Number<br>(FWA, MPA or SPA)<br>if applicable |
|-----------|--------------------------|--------------------------|---------------|--|
|           | Primary Site             | CDC                      | Atlanta, GA   |  |
|           | Explanation of project   | components:              |               |  |
| •         | If project involves rese | ough that is funded outs |               |  |
| 9.        |                          | oject components will be |               | ld be restricted pending IRB approval                  |
| <b>9.</b> |                          |                          |               | ld be restricted pending IRB approval                  |
| 9.<br>—   |                          |                          |               | ld be restricted pending IRB approval                  |
| 9.        |                          |                          |               | ld be restricted pending IRB approval                  |

7.

| Approvals (signature and position title)          | Date       | Research Determination / Remarks  |
|---|------------|---|
| Netta Apedoe - PUBLIC HEALTH<br>ADVISOR           | 12/21/2012 | <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>(check if applicable)</li> <li>[] Local IRB</li> <li>[] CDC Exemption</li> <li>[] CDC IRB</li> </ul> |
| staff member completing this form                 |            | Comments:   |
| -   | 01/02/2013 | [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 (check if applicable) [] Local IRB [] CDC Exemption [] CDC IRB   |
| Team Lead   |            | Comments:   |
| Cheryll Thomas - EPIDEMIOLOGIST                   | 01/07/2013 | <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>(check if applicable)</li> <li>[] Local IRB</li> <li>[] CDC Exemption</li> <li>[] CDC IRB</li> </ul> |
| Division ADS                                      |            | Comments:   |
| Joan Redmond Leonard - PUBLIC<br>HEALTH ANALYST   | 01/23/2013 | [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 (check if applicable) [] Local IRB [] CDC Exemption [] CDC IRB   |
| cuc ADS, Deputy ADS, or Human Subjects<br>Contact |            | Comments:   |