AMENDMENT REVIEW FORM

(TO ADD OR CHANGE PREVIOUSLY APPROVED RESEARCH)

**All changes or new activities for previously approved studies require submission, review, and approval of an Amendment Review Form.** Please complete and upload this form to your project’s document library on [IRBTRAC](mailto:irb@westat.com) along with all other necessary materials to be reviewed. Once the request has been reviewed, you will be contacted. If this change or new activity requires a full Board review, those meetings occur on the second Tuesday of every month. To check the date of meetings, please see the [meeting schedule](http://wesinfo/version2/research/irb/index.cfm) under IRB in WesInfo. Thank you for your cooperation.

|  |  |
| --- | --- |
| 1. Today’s Date: | 05 / 05 / 2014 |

|  |  |
| --- | --- |
| Date of Original Approval: | 07 / 19 / 2014 |

|  |  |
| --- | --- |
| Project Name: | Population assessment of Tobacco and Health (PATH) |

|  |  |
| --- | --- |
| Westat Project Number: | 8954.00.00 |

|  |  |
| --- | --- |
| Agency Grant or Contract Number: | HHSN271201100027C, Ref # NO1DA-11-5568 |

|  |  |  |  |
| --- | --- | --- | --- |
| Project Director: | David Maklan, submitted by |  | **Ext.** |

|  |  |
| --- | --- |
| Unit Ops Number/Study Area: | 112.56 |

|  |  |  |  |
| --- | --- | --- | --- |
| Area IRB Representative: | Katie Gasque |  | **Ext.** 3694 |

2. Indicate the type of addition or change being requested to a previously approved study.  
*(select all that apply.)*

|  |  |  |
| --- | --- | --- |
| Name(s) of investigators  Project number  Introduction of a new IRB or request for Westat to serve as the IRB  Study design, survey questionnaire, or procedure(s)  Informed consent process, consent form(s), parent permission(s), or assent form(s)  Recruitment materials or strategies  Incentives  Survey instruments  Number or type of populations studied | Review of final instrument such as interview questions or data collection sites for a previously approved study  Mode of administration of instruments in your study (e.g., from mail or telephone to web or Internet access)  Data access rights  Any other change in protocol that affects treatment of human subjects: ***(please specify)*** | |
|  |  |

1. Please provide a brief summary of your change or addition to previously approved research.

|  |
| --- |
| In preparation for the fielding of Wave 3 of the PATH study, we will be cognitively testing a selection of items from the instrument in English and Spanish. |

1. How does each change or addition affect the risks to participants in your study? *(select only one.)*

a.  No change

b.  N/A – no risks

c.  Decreases the risk *(****specify****):*

|  |
| --- |
|  |

d.  Increases the risk *(****specify****):*

|  |
| --- |
|  |

e.  Adds a new risk *(****specify****):*

|  |
| --- |
|  |

**FOR HARD-COPY SUBMISSION, PLEASE SIGN HERE:**

**A signature is not required when you return this form electronically; however, please fill in the date of completion.**

**The information provided in this request form is complete and correct.**

|  |  |  |  |
| --- | --- | --- | --- |
| **Project Director/ Principal Investigator:** |  | **Date:** | 05 / 12/ 2014 |

|  |
| --- |
| Please attach:   * **One document that clearly identifies (through track changes, highlights, or italics) the revision in the previously approved submission.** * **Another document labeled “corrected version.”** |

If you have any questions, feel free to contact Sharon Zack, the IRB Administrator, at x8828.

|  |  |
| --- | --- |
| **IRB Administration Use Only**  Expedited review and approval for the modification(s) on this form: | |
| 5/30/14  IRB Chair / Associate Chair / Designee |  |
| |  | | --- | | **IRB Office Only**  **Approved** – Next continuing review date: 04 / 18/ 2015  **Conditional approval** (Please see attached letter)  **Did not qualify for expedited review** | | |