

IRB ID Number: 12259

Office of Research Protection Institutional Review Board Notice of Approval Federalwide Assurance No. 3331

Title of Study: National Intimate Partner and Sexual Violence Surveillance System	
RTI Project Number: 0211832.000 RTI Proposal Number (if no Project Number)	
Project Leader: Lisa Carley-Baxter	
Project Team Member Contact (if different from Project Leader):	
Source of Funding for this Study: <u>CDC</u>	
Date Submitted to IRB: November 19, 2012	
Level of Review (check one):	
Full □, IRB Meeting Date:	
Expedited ⊠, category: 9: Cont. Rev. minimal risk research Type of Review <i>(check one)</i> :	
Preliminary review (The grant application/contract proposal and propos	
concordance with regard to the scientific conduct of the study, inform	
pertaining to the protection of human subjects. (45 CFR 46.103(f)) D	o not involve human subjects or data unti
pretest or full study is approved.)	
Amendment, describe:	
Add study site(s):	MD I
Pretest/Pilot Test Full Implementation	⊠Renewal ⊡Study Closure
IRB Approval of Special Conditions (check all that apply to this re ☐ Waiver of Signed Informed Consent/Parental Permission	view):
Waiver of elements of Informed Consent or requirement for Informed	
Participation of Pregnant Women (Worksheet B submitted by p	
Participation of Prisoners (Worksheet C submitted by project tea	
Participation of Prisoners in DHHS-funded studies (OHRP acknowledge)	
 Participation of Prisoners (Worksheet C submitted by project team Participation of Prisoners in DHHS-funded studies (OHRP acknown Participation of Minors (Worksheet D submitted by project team IRB Agreement of Nonsignificant Risk Device Study Determination HIPAA Waiver of Authorization 	,
IRB Agreement of Nonsignificant Risk Device Study Determinati	ion
☐ HIPAA Waiver of Authorization	
Places note the following requirements:	
 Please note the following requirements: If unexpected problems or adverse events occur, the project to 	team must notify the IPR
 If unexpected problems or adverse events occur, the project team must notify the IRB. If there are changes in study procedures or protocol or any data collection materials (brochures, letters, 	
questionnaires, etc.) the project team must notify the IRB before they are implemented.	
The project team is required to apply for continuing review as long as the study is active, which includes	
participation of human subjects or possession of human data or specimens.	
F - 1 - 1 - D - 1 - CIDD A I 40/04/0040	
Expiration Date of IRB Approval: 12/01/2013	
(No human subjects research can occur after this date without	t continuing review and approval.)
Justo M. Caldell	
	11/20/2012
	Date of IRB Approval
Signature - IIVD Member of Chair	Date of IND Approval
Juesta Caddell, Ph.D.	
Name - IRB Member or Chair (print or type)	
Copy sent to project leader	
Entered into MIS	
OHRP acknowledgement received for participation of prisoners in DHHS	-iunaea studies on: