

Instructions for Submitting an Intervention

To submit an intervention, individuals should send a written statement to NREPP expressing their interest along with documentation that demonstrates the intervention meets the minimum requirements as described above. All submissions must be made either by a principal investigator (PI) who has conducted research on the intervention, a project director (PD) who has worked with an evaluator of the intervention, or a formally authorized delegate of the PI or PD. For information on where to submit materials, please call 1-866-436-7377. Electronic submissions are preferred, but materials may be sent to NREPP in hard copy via postal mail or fax. To be eligible for consideration, submissions must be received no later than 11:59 p.m. E.S.T. on February 1, 2012; those received before November 1, 2011, will be disregarded.

For each intervention that is accepted, the Principal (the individual, usually the PI, formally designated as the intervention's point of contact and decisionmaking authority during the review process) will be asked to submit additional documentation to be used in the review. This additional documentation includes full-text copies of all articles and reports that provide evidence of significant outcomes ($p \leq .05$) as well as copies of selected dissemination materials in the format they are provided to the public (e.g., hard copies or electronic versions of manuals, training presentations, tools, quality assurance protocols; URLs for interactive Web-based resources).

The Principal continues to work with NREPP staff throughout the review and is responsible for approval of the final intervention summary that is developed by NREPP staff once the review has been completed.

Contact Information

Individuals who have questions about the information contained in this notice may write to NREPP staff at nrepp@samhsa.hhs.gov or call 1-866-436-7377.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee to the Director (ACD), Centers for Disease Control and Prevention (CDC)—Ethics Subcommittee (ES)

Correction: This notice was published in the **Federal Register** on September 8, 2011, Volume 76, Number 174, Page 55678. The correct time should be 1 p.m.–3:30 p.m.

Contact Person for More Information: Drue Barrett, Ph.D., Designated Federal Officer, ACD, CDC—ES, CDC, 1600 Clifton Road, NE., M/S D-50, Atlanta, Georgia 30333. Telephone (404) 639-4690. E-mail: d Barrett@cdc.gov.

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Dated: September 12, 2011.

Elaine L. Baker,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2011-23767 Filed 9-15-11; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS-10398]

Agency Information Collection Activities: Submission for OMB Review; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, HHS.

In compliance with the requirement of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Centers for Medicare & Medicaid Services (CMS), Department of Health and Human Services, is publishing the following summary of proposed collections for public comment. Interested persons are invited to send comments regarding this burden estimate or any other aspect of this collection of information, including any of the following subjects: (1) The necessity and utility of the proposed information collection for the proper performance of the Agency's function;

(2) the accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

1. *Type of Information Collection Request:* New collection; *Title of Information Collection:* Generic Clearance for Medicaid and CHIP State Plan, Waiver, and Program Submissions; *Use:* CMS is requesting a generic PRA clearance for a body of forms necessary to conduct ongoing business with State partners in the implementation of Medicaid and the Children's Health Insurance Program (CHIP). Examples of the types of forms to be produced in this collection include State plan amendment templates, waiver and demonstration templates, and reporting templates. The development of streamlined submission forms is critical for States to implement timely health reform initiatives in Medicaid and CHIP state plans, demonstrations, and waivers, including legislative requirements enacted by the Affordable Care Act. The development of streamlined submissions forms enhances the collaboration and partnership between States and CMS by documenting CMS policy for States to use as they are developing program changes. Streamlined forms improve efficiency of administration by creating a common and user-friendly understanding of the information needed by CMS to quickly process requests for State plan amendments, waivers, and demonstration, as well as ongoing reporting.; *Form Number:* CMS-10398 (OMB #0938-NEW); *Frequency:* Occasionally; *Affected Public:* State, Local, or Tribal Governments; *Number of Respondents:* 56; *Total Annual Responses:* 1120; *Total Annual Hours:* 28,747. (For policy questions regarding this collection contact Candice Payne at 410-786-4453. For all other issues call 410-786-1326.)

To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS Web Site address at <http://www.cms.hhs.gov/PaperworkReductionActof1995>, or E-mail your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov, or call the Reports Clearance Office on (410) 786-1326.

To be assured consideration, comments and recommendations for the proposed information collections must