

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondent	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)
Individuals in households	Screener	15,000	1	5/60
Individuals in households	Household Interview	5,000	1	1.5
Individuals in households	MEC Interview & Examination	5,000	1	4
Individuals in households	Telephone Dietary Recall & Dietary Supplements	5,000	1	30/60
Individuals in households	Flexible Consumer Behavior Survey Phone Follow-Up	5,000	1	20/60
Individuals in households	Developmental Projects & Special Studies	3,500	1	3
Individuals in households	24 hour Blood Pressure Pilot	1,000	1	25

Jeffrey M. Zirger,

Acting Chief, Information Collection Review Office, Office of Scientific Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2018-22008 Filed 10-9-18; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

Privacy Act of 1974; System of Records

AGENCY: Centers for Medicare & Medicaid Services (CMS), Department of Health and Human Services (HHS).

ACTION: Notice of withdrawal.

SUMMARY: The Department of Health and Human Services (HHS), Centers for Medicare & Medicaid Services (CMS), is withdrawing the notice published on September 17, 2018 to modify system of records No. 09-70-0541, titled “Medicaid Statistical Information System (MSIS).” The notice was prematurely published. A revised version will be published at a later date.

DATES: The notice of withdrawal is applicable October 10, 2018.

ADDRESSES: Any comments should be submitted by mail or email to: CMS Privacy Act Officer, Division of Security, Privacy Policy & Governance, Information Security & Privacy Group, Office of Information Technology, CMS, Location N1-14-56, 7500 Security Blvd., Baltimore, MD 21244-1870, or walter.stone@cms.hhs.gov.

FOR FURTHER INFORMATION CONTACT: General questions may be submitted by phone, mail or email to Barbara Demopulos, (phone 410-786-6340), CMS Privacy Advisor, Division of Security, Privacy Policy & Governance, Information Security & Privacy Group, Office of Information Technology, CMS, Location N1-14-40, 7500 Security

Blvd., Baltimore, MD 21244-1870, or Barbara.demopulos@cms.hhs.gov.

SUPPLEMENTARY INFORMATION: A notice establishing or significantly modifying a system of records is required by subsection (r) of the Privacy Act (5 U.S.C. 552a(r)) to be reported to the Committee on Government Operations of the House of Representatives, the Committee on Governmental Affairs of the Senate, and the Office of Management and Budget (OMB) in advance of publication in the **Federal Register**, in order to permit an evaluation of the potential effect of the proposal on the privacy and other rights of individuals. The notice published at 83 FR 46951 (Sept. 17, 2018) did not comply with this requirement and is therefore withdrawn, as prematurely published. A revised version will be published at a later date and in compliance with 5 U.S.C. 552a(r) and section 7 of OMB Circular A-108, “Federal Agency Responsibilities for Review, Reporting, and Publication under the Privacy Act,” 81 FR 94424 (Dec. 23, 2016).

Barbara Demopulos,

CMS Privacy Advisor, Division of Security, Privacy Policy and Governance Information Security and Privacy Group, Office of Information Technology, Centers for Medicare & Medicaid Services.

[FR Doc. 2018-21899 Filed 10-9-18; 8:45 am]

BILLING CODE 4120-03-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Children’s Bureau; Proposed Information Collection Activity; Comment Request

AGENCY: Administration for Children and Families, U.S. Department of Health and Human Services.

Title: RPG National Cross-Site Evaluation and Evaluation Technical Assistance.

OMB No.: New Collection.

Description: The Children’s Bureau (CB) within the Administration for Children and Families of the U.S. Department of Health and Human Services seeks approval to collect information for the Regional Partnership Grants to Increase the Well-being of and to Improve Permanency Outcomes for Children Affected by Substance Abuse (known as the Regional Partnership Grants Program or “RPG”) Cross-Site Evaluation and Evaluation-Related Technical Assistance project. The Child and Family Services Improvement and Innovation Act (Pub. L. 112-34) includes a targeted grants program (section 437(f) of the Social Security Act) that directs the Secretary of Health and Human Services to reserve a specified portion of the appropriation for these Regional Partnership Grants, to be used to improve the well-being of children affected by substance abuse. Under three prior rounds of RPG, the Children’s Bureau has issued 74 grants to organizations such as child welfare or substance abuse treatment providers or family court systems to develop interagency collaborations and integration of programs, activities, and services designed to increase well-being, improve permanency, and enhance the safety of children who are in an out-of-home placement or are at risk of being placed in out-of-home care as a result of a parent’s or caretaker’s substance abuse. In 2017, CB awarded grants to a fourth cohort of 17 grantees and in 2018 they plan to award 10 grants to a fifth cohort.

The RPG cross-site evaluation will extend our understanding of what types of programs and services grantees provided to participants, how grantees leveraged their partnerships to coordinate services for children and families, and what the outcomes were for children and families enrolled in RPG programs. First, the cross-site evaluation will describe the characteristics of participants served by RPG programs, the types of services provided to families, the dosage of each

type of service received by families, and the level of participant engagement with the services provided. Second, the cross-site will assess the coordination of partners' service systems (e.g., shared participant data, joint staff training) to better understand how partners' collaborative effort affects the array of services offered to families. The cross-site evaluation will also focus more deeply on the partnership between the child welfare and substance use disorder (SUD) treatment agencies, to add to the research base about how these agencies can collaborate to address the needs of children and families affected by SUD. Finally, the evaluation will assess the outcomes of children and adults served through the RPG program.

The evaluation is being undertaken by the Children's Bureau and its contractor Mathematica Policy Research. The evaluation is being implemented by Mathematica Policy Research and its subcontractor, WRMA Inc.

The RPG Cross-Site Evaluation will include the following data collection activities:

1. *Site visits and key informant interviews.* The cross-site evaluation team will visit up to 21 sites to better understand the partnership and coordination between the child welfare and SUD treatment agencies. The remaining six grantees will participate in telephone interviews to gather similar information about their design and implementation. The site visits and phone interviews will focus on the RPG planning process; how and why particular services were selected; the ability of the child welfare, substance use disorder treatment, and other service systems to collaborate and support quality implementation of the RPG services; challenges experienced; and the potential for sustaining the collaborations and services after RPG funding ends.

2. *Partner survey.* To describe the interagency collaboration within RPG sites, grantees and their partners will participate in an online survey once during the grant period. One person

from each organization knowledgeable about the RPG program will be invited to participate in the survey. The survey will collect information about communication and service coordination among partners. The survey will also collect information on characteristics of strong partnerships (e.g., data sharing agreements, colocation of staff, referral procedures, and cross-staff training).

3. *Semi-annual progress reports.* The semi-annual progress reports will be used to obtain updated information from grantee project directors about their program operations and partnerships, including any changes from prior periods. The CB has tailored the semi-annual progress reports to collect information on grantees' programs and other services grantees implement, the target population for the RPG program, and grantees' perceived successes and challenges to implementation.

4. *Enrollment, client, and service data.* To document participant characteristics and their enrollment in RPG services, all grantees will provide data on family characteristics, and enrollment of and services provided to RPG families. These data include demographic information on family members, dates of entry into and exit from RPG services, and information on RPG service dosage. These data will be submitted on an ongoing basis by staff at the grantee organizations into an information system developed by the cross-site evaluation team.

5. *Outcome and impact data.* To measure participant outcomes, all grantees will collect self-administered standardized instruments from RPG adults. The standardized instruments used in RPG collect information on child well-being, adult and family functioning, and adult substance use. Grantees will share the responses on these self-report instruments with the cross-site evaluation team. Grantees will also obtain administrative data on a common set of child welfare and substance use disorder treatment data elements.

In addition to conducting local evaluations and participating in the RPG Cross-Site Evaluation, the RPG grantees are legislatively required to report performance indicators aligned with their proposed program strategies and activities. A key strategy of the RPG Cross-Site Evaluation is to minimize burden on the grantees by ensuring that the cross-site evaluation, which includes all grantees in a study that collects data to report on implementation, the partnerships, and participant characteristics and outcomes, fully meets the need for performance reporting. Thus, rather than collecting separate evaluation and performance indicator data, the grantees need only participate in the cross-site evaluation. In addition, using the standardized instruments that the Children's Bureau has specified will ensure that grantees have valid and reliable data on child and family outcomes for their local evaluations. The inclusion of an impact study conducted on a subset of grantees with rigorous designs will also provide the Children's Bureau, Congress, grantees, providers, and researchers with information about the effectiveness of RPG programs.

This 60-Day **Federal Register** Notice covers the following data collection activities: (1) The site visits with grantees; (2) the web-based survey of grantee partners (3) the semi-annual progress reports; (4) enrollment and service data provided by grantees; and (6) outcome and impact data provided by grantees.

Respondents. Respondents include grantee staff or contractors (such as local evaluators) and partner staff. Specific types of respondents and the expected number per data collection effort are noted in the burden table below.

Annual burden estimates. The following instruments are proposed for public comment under this 60-Day **Federal Register** Notice. Burden for all components is annualized over three years.

RPG CROSS-SITE EVALUATION ANNUALIZED BURDEN ESTIMATES

Data collection activity	Total number of respondents	Number of responses per respondent (each year)	Average burden hours per response (in hours)	Estimated total burden hours	Total annual burden hours
Site Visit and Key Informant Data Collection					
Program director in-person interview	21	.33	2	42	14
Program manager/supervisor in-person interview	21	.33	1	21	7
Partner representative interviews	63	.33	1	63	21
Frontline staff interview	42	.33	1	42	14
Program director/manager phone interview	12	.33	1	4.0	12

RPG CROSS-SITE EVALUATION ANNUALIZED BURDEN ESTIMATES—Continued

Data collection activity	Total number of respondents	Number of responses per respondent (each year)	Average burden hours per response (in hours)	Estimated total burden hours	Total annual burden hours
Partner survey	135	.33	0.42	56.3	18.8
Enrollment, client and service data					
Semi-annual progress reports	27	2	16.5	2,673	891
Case enrollment data	81	43	0.25	2,612.3	870.8
Case closure	81	43	0.017	174.2	58.1
Case closure—prenatal	81	33	0.017	133.7	44.6
Service log entries	162	2,288	0.03	37,065	12,355
Outcome and impact data					
<i>Administrative Data:</i>					
Obtain access to administrative data	27	1	42.6	3,450.6	1150.2
Report administrative data	27	2	144	23,328	7,776
<i>Standardized instruments:</i>					
Review and adopt reporting templates	27	.33	8	216	72
Data entry for standardized instruments	27	130	1.25	13,162.5	4,387.5
Review records and submit	27	2	25	4,050	1,350
Data entry for comparison study sites (22 grantees)	22	130	1.25	10,725	3,575
Estimated Total Burden Hours				97,827	32,609

In compliance with the requirements of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Children’s Bureau within the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded by writing to Administration for Children and Families, Office of Administration, Office of Planning, Research and Evaluation, 330 C Street SW, Washington, DC 20201, Attn: ACF Reports Clearance Officer. Email address: infocollection@acf.hhs.gov. All requests should be identified by the title of the information collection.

The Department specifically requests comments on: (a) Whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency’s estimate of the burden of the proposed collection of information; (c) the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Consideration will be given to

comments and suggestions within 60 days of this publication.

Robert Sargis,
Reports Clearance Officer.
 [FR Doc. 2018–22020 Filed 10–9–18; 8:45 am]
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2018–N–3685]

International Drug Scheduling; Convention on Psychotropic Substances; Single Convention on Narcotic Drugs; ADB–FUBINACA; ADB–CHMINACA; Cyclopropyl Fentanyl; Methoxyacetyl Fentanyl; para-Fluoro Butyrfentanyl; Tramadol; Pregabalin; Cannabis Plant and Resin; and Eight Additional Substances; Request for Comments

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice; request for comments.

SUMMARY: The Food and Drug Administration (FDA) is requesting interested persons to submit comments concerning abuse potential, actual abuse, medical usefulness, trafficking, and impact of scheduling changes on availability for medical use of 16 drug substances. These comments will be considered in preparing a response from the United States to the World Health

Organization (WHO) regarding the abuse liability and diversion of these drugs. WHO will use this information to consider whether to recommend that certain international restrictions be placed on these drugs. This notice requesting comments is required by the Controlled Substances Act (the CSA).

DATES: Submit either electronic or written comments by October 31, 2018.

ADDRESSES: You may submit comments as follows. Please note that late, untimely filed comments will not be considered. Electronic comments must be submitted on or before (enter date), 2018. The <https://www.regulations.gov> electronic filing system will accept comments until 11:59 p.m. Eastern Time at the end of October 31, 2018. Comments received by mail/hand delivery/courier (for written/paper submissions) will be considered timely if they are postmarked or the delivery service acceptance receipt is on or before that date.

Electronic Submissions

Submit electronic comments in the following way:

- *Federal eRulemaking Portal:* <https://www.regulations.gov>. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to <https://www.regulations.gov> will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any