U.S.C. 652(a)(7), which requires OCSE to provide technical assistance to state child support enforcement agencies to help them establish effective systems for collecting child and spousal support; and, 45 CFR 303.7(a)(5), which requires state child support agencies to transmit requests for information and provide requested information electronically to the greatest extent possible. To facilitate this requirement for states, OCSE developed the *FAST* Levy system that supports the electronic exchange of lien and levy information between child

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support agencies and financial institutions.

Respondents: Multistate Financial Institutions and State Child Support Agencies.

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Financial Data Match Result File-Portal	192 30	4	5 minutes ¹ 0.5	64 15
FAST-Levy Record Specifications: Current Financial Institutions Users to Program New Codes.	1	1	65 ²	65
FAST-Levy Record Specifications: Current State Child Support Agencies to Program New Codes.	3	1	65	195
FAST-Levy Response Withhold Record Specifications: Financial Institu- tions.	1	1	1,716	1,716
FAST-Levy Request Withhold Record Specifications: State Child Support Agencies.	2	1	1,610	3,220

¹ Estimate is approximately 5 minutes per response. For calculation, use 5/60.

²Estimate is an average based on input from OCSE's matching partners.

Estimated Total Annual Burden Hours: 5,275.

Additional Information: Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 330 C Street SW., Washington, DC 20201, Attention Reports Clearance Officer. All requests should be identified by the information collection. Email address: *infocollection@acf.hhs.gov.*

OMB Comment: OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the Federal Register. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following: Office of Management and Budget, Paperwork Reduction Project, email: OIRA SUBMISSION@OMB.EOP.GOV. Attn: Desk Officer for the Administration for Children and Families.

Bob Sargis,

Reports Clearance Officer. [FR Doc. 2017–13252 Filed 6–23–17; 8:45 am]

BILLING CODE 4184-41-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Comment Request

Title: Evaluation of Employment Coaching for TANF and Other Low-Income Populations.

OMB No.: New Collection.

Description: The Administration for Children and Families (ACF) is proposing a data collection activity as part of the Evaluation of Employment Coaching for TANF and Other Low-Income Populations. This study will provide an opportunity to learn more about the potential of coaching to help clients achieve self-sufficiency and other desired employment-related outcomes. It will take place over five years in up to three employment programs. These programs may be Temporary Assistance for Needy Families (TANF) agencies or other public or private employment programs that serve low-income individuals. Selected sites will include a robust coaching component and have the capacity to conduct a rigorous impact evaluation, among other criteria. This study will provide information on whether coaching helps people obtain and retain jobs, advance in their careers, move toward self-sufficiency, and improve their overall well-being. To meet these objectives, this study will include an impact and implementation study.

The impact study will involve participants being randomly assigned to either a "program group," who will be paired with a coach, or to a "control group," who will not be paired with a coach. The effectiveness of the coaching will be determined by differences between members of the program and control groups in outcomes such as obtaining and retaining employment, earnings, measures of self-sufficiency, and measures of self-regulation.

The implementation study will document coaching practices, describe lessons learned from implementing coaching, and enhance interpretation of the impact study findings.

The proposed information collection activities are: (1) Baseline data collection: Collection of characteristics data on all study participants as they enroll in the study. Data will be entered into the Random Assignment, Participant Tracking Enrollment, and Reporting (RAPTER) system; (2) First follow-up survey: Collection of outcome data for a subset of study participants about 9 months after random assignment; (3) Semi-structured staff interviews: Collection of qualitative data on the design and implementation of the program; (4) Staff survey: Collection of information on staff members' professional backgrounds, training, coaching practices, and attitudes; (5) Indepth participant interviews: Collection of detailed information about the participants' backgrounds and experiences with coaching; (6) Staff reports of program service receipt: Collection of data on coaching and other program services received by study

participants and entered into RAPTER; and (7) Video recordings of coaching sessions: Collection of data on the interaction between the coaches and participants.

A second follow-up survey will be administered approximately 21 months

after random assignment. This data collection activity will be included under a separate OMB submission.

Respondents: Program staff and individuals enrolled in the Evaluation of Employment Coaching for TANF and Other Low-Income Populations.

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Program staff may include coaches, case managers, workshop instructors, job developers, supervisors, and managers. All participants will be able to opt out of participating in the data collection activities.

Instrument	Total number of respondents	Annual number of respondents	Number of responses per respondent	Average burden hours per response	Annual burden hours
Baseline data collection—study participants Baseline data collection—staff	6,000 60	2,000 20	1 100	0.33 0.33	660 660
First follow-up survey	2,400	800	1	1	800
Semi-structured staff interviews	66	22	1	1.5	33
Staff survey	48	16	1	0.75	12
In-depth participant interviews	24	8	1	2.5	20
Staff reports of program service receipt	30	10	5,200	0.03	1,560
Video recordings of coaching sessions	27	9	10	0.10	9

Estimated Total Annual Burden Hours: 3,754.

In compliance with the requirements of section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, 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 the Administration for Children and Families, Office of Planning, Research, and Evaluation, 330 C Street SW., Washington, DC 20201, Attn: OPRE Reports Clearance Officer. Email address: OPREinfocollection@ 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 functions 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 submitted within 60 days of this publication.

Mary Jones,

ACF/OPRE Certifying Officer. [FR Doc. 2017–13288 Filed 6–23–17; 8:45 am] BILLING CODE 4184–09–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2016-N-4620]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Medical Devices; Reports of Corrections and Removals

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.
DATES: Fax written comments on the collection of information by July 26, 2017.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202–395–7285, or emailed to *oira_submission@omb.eop.gov*. All comments should be identified with the OMB control number 0910–0359. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT:

Amber Sanford, Office of Operations, Food and Drug Administration, Three White Flint North, 10A63, 11601 Landsdown St., North Bethesda, MD 20852, 301–796–8867, PRAStaff@ fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Medical Devices; Reports of Corrections and Removals—21 CFR Part 806

OMB Control Number 0910–0359— Extension

FDA is requesting approval for the collection of information regarding reports of corrections and removals required under part 806 (21 CFR part 806), which implements section 519(g) of the Federal Food, Drug, and Cosmetic Act (the FD&C Act) (21 U.S.C. 360i(g)), as amended by the Food and Drug Modernization Act of 1997 (FDAMA) (Pub. L. 105–115). A description of the information collection requirements are provided as follows:

Under § 806.10 (21 CFR 806.10), within 10 working days of initiating any action to correct or remove a device to reduce a risk to health posed by the device or to remedy a violation of the FD&C Act caused by the device that may present a risk to health, device manufacturers or importers must submit a written report to FDA of the correction or removal.

Under § 806.20(a), device manufacturers or importers that initiate a correction or removal that is not required to be reported to FDA must keep a record of the correction or removal.

The information collected in the reports of corrections and removals will be used by FDA to identify marketed devices that have serious problems and