

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Comment Request

Proposed Projects:

Title: The Evaluation of Child Welfare Information Gateway.

OMB No.: New Collection.

Description: The Children’s Bureau (CB), Administration for Children and Families (ACF), U.S. Department of Health and Human Services (HHS) is proposing new or expanded data collection activities as part of its Evaluation of Child Welfare Information Gateway.

Child Welfare Information Gateway (CWIG) is a national information clearinghouse and service of the Children’s Bureau, Administration for Children and Families, U.S. Department of Health and Human Services. CWIG connects professionals and concerned citizens to resources and information on programs, research, legislation, and statistics regarding child maltreatment, child abuse prevention, and child welfare services designed to achieve the safety, permanency, and well-being of children and families. The Evaluation of Child Welfare Information Gateway gathers information to inform the Children’s Bureau about the kind and quality of information services that customers want, how customers are using resources and services, as well as customers’ level of satisfaction with existing services.

The Market Research Sub-Study complements information obtained from the larger Evaluation of Child Welfare Information Gateway. The Children’s Bureau seeks to learn more about how child welfare professionals and students planning to enter the child welfare workforce access and consume work-related information. This national study will focus on understanding child welfare professionals’ and students’ characteristics, use of technology, and preferences for obtaining information that they use in their work. The goal of the sub-study is to provide federally-funded technical assistance providers and other stakeholder organizations with a better understanding of their target audiences so they can design more effective products, services, and dissemination strategies to reach these populations.

Data collection activities proposed for the Evaluation of Child Welfare Information Gateway include: six online targeted surveys designed to evaluate CWIG’s special initiative websites and other targeted website sections; five online event surveys administered after CWIG-sponsored webinars, presentations, or other events; five focus groups (each with approximately 10 participants) with users and non-users of CWIG’s special initiative websites and other CWIG products and services; and, a general customer survey delivered via multiple modes (e.g., website, email, live chat, print, and phone). The sampling plan for the CWIG general customer survey is designed to reach the various types of customers using Child Welfare Information

Gateway services such as professionals, students, and customers looking for assistance with a personal situation while reducing burden for respondents by only asking relevant questions for their backgrounds.

The market research sub-study seeks to deliver surveys and conduct focus groups to gauge online information habits and preferences. The proposed market research sub-study will consist of a national online survey of child welfare professionals and students, which will be administered through four different instruments tailored for four different populations. Ten focus groups (each with 8 to 10 participants) will be used to learn more about different audiences’ habits and preferences related to child welfare information access and consumption.

Respondents: The Evaluation of Child Welfare Information Gateway will target all types of possible CWIG users including: State and local governments, the territories, service providers, Tribes and tribal organizations, grantees, researchers, and the general public seeking information and resources from Child Welfare Information Gateway via the website, mail, telephone, Live Chat, and email. The Market Research Sub-Study will target child welfare professionals in state, county, tribal, and private agencies; Court Improvement Program coordinators and directors; judges and attorneys involved in child welfare-related work; and students in Bachelor’s and Master’s degree programs in social work that receive Title IV–E or IV–B stipends.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Child Welfare Information Gateway’s Targeted Survey	2,355	1	0.084	197.82
Child Welfare Information Gateway’s Event Survey	500	1	0.05	25
Child Welfare Information Gateway’s Focus Group Guide	50	1	1	50
Child Welfare Information Gateway’s General Customer Survey: Questions for Professionals	960	1	0.084	80.64
Child Welfare Information Gateway’s General Customer Survey: Questions for Students	480	1	0.05	24
Child Welfare Information Gateway’s General Customer Survey: Questions for Personal Customers	960	1	0.05	48
Market Research Sub-Study: Online Information Habits and Preferences Survey (for child welfare professionals in state, county, and private agencies)	1,400	1	0.5	700
Market Research Sub-Study: Online Information Habits and Preferences Survey (for child welfare professionals working with tribes)	1,000	1	0.5	500
Market Research Sub-Study: Online Information Habits and Preferences Survey (for legal professionals working in child welfare)	1,400	1	0.5	700
Market Research Sub-Study: Online Information Habits and Preferences Survey (for students planning to enter the child welfare workforce)	900	1	0.5	450
Market Research Sub-Study: Focus Groups on Information Habits and Preferences	100	1	1.5	150

Estimated Total Annual Burden Hours: 2,925 hours.

In compliance with the requirements of the Paperwork Reduction Act of 1995 (Pub. L. 104–13, 44 U.S.C. Chap 35), 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: 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 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.

Robert Sargis,

Reports Clearance Officer.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2017–N–7022]

Post-Marketing Pediatric-Focused Product Safety Reviews; Establishment of a Public Docket; Request for Comments

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice, establishment of a public docket; request for comments.

SUMMARY: The Food and Drug Administration (FDA) is establishing a public docket to collect comments related to the post-marketing pediatric-focused safety reviews of products posted between October 23, 2017, and March 16, 2018, on FDA's website but

not presented at the March 23, 2018, Pediatric Advisory Committee (PAC) meeting. These reviews are intended to be available for review and comment by members of the PAC, interested parties (such as academic researchers, regulated industries, consortia, and patient groups), and the general public.

DATES: Submit either electronic or written comments by March 30, 2018.

ADDRESSES: FDA is establishing a docket for public comment on this document. The docket number is FDA–2017–N–7022. The docket will close on March 30, 2018. Submit either electronic or written comments by that date. Please note that late, untimely comments will not be considered. Electronic comments must be submitted on or before March 30, 2018. The <https://www.regulations.gov> electronic filing system will accept comments until midnight Eastern Time at the end of March 30, 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.

You may submit comments as follows:

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 confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else's Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on <https://www.regulations.gov>.

- If you want to submit a comment with confidential information that you do not wish to make available to the public, submit the comment as a written/paper submission and in the manner detailed (see "Written/Paper Submissions" and "Instructions").

Written/Paper Submissions

Submit written/paper submissions as follows:

- **Mail/Hand delivery/Courier (for written/paper submissions):** Dockets

Management Staff (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

- For written/paper comments submitted to the Dockets Management Staff, FDA will post your comment, as well as any attachments, except for information submitted, marked and identified, as confidential, if submitted as detailed in "Instructions."

Instructions: All submissions received must include the Docket No. FDA–2017–N–7022 for "Post-Marketing Pediatric-Focused Product Safety Reviews; Establishment of a Public Docket; Request for Comments." Received comments, those filed in a timely manner (see **ADDRESSES**), will be placed in the docket and, except for those submitted as "Confidential Submissions," publicly viewable at <https://www.regulations.gov> or at the Dockets Management Staff between 9 a.m. and 4 p.m., Monday through Friday.

- **Confidential Submissions—**To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states "THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION." FDA will review this copy, including the claimed confidential information, in its consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on <https://www.regulations.gov>. Submit both copies to the Dockets Management Staff. If you do not wish your name and contact information to be made publicly available, you can provide this information on the cover sheet and not in the body of your comments and you must identify this information as "confidential." Any information marked as "confidential" will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA's posting of comments to public dockets, see 80 FR 56469, September 18, 2015, or access the information at: <https://www.gpo.gov/fdsys/pkg/FR-2015-09-18/pdf/2015-23389.pdf>.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to <https://www.regulations.gov> and insert the docket number, found in brackets in the heading of this document, into the "Search" box and follow the prompts