Dated: June 17, 2021.

William N. Parham, III,
Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2021–13223 Filed 6–23–21; 8:45 am]
BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Evaluation of LifeSet (New Collection)

AGENCY: Office of Planning, Research, and Evaluation; Administration for Children and Families; HHS.

ACTION: Request for public comment.

SUMMARY: The Administration for Children and Families (ACF) at the U.S. Department of Health and Human Services (HHS) is proposing a new information collection activity to assess the impact and implementation of LifeSet, a program that provides services and supports to young adults ages 17 to 21 with previous child welfare involvement. Data collection efforts will include accessing administrative data from the child welfare agency, program, and other private and governmental databases; surveys of young adults (participants and those receiving services as usual); interviews and focus groups with program and child welfare agency administrators and staff; interviews and focus groups with young adult program participants; and interviews with other program stakeholders.

DATES: Comments due within 30 days of publication. OMB must make a decision 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.

ADDRESSES: Written comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting “Currently under 30-day Review—Open for Public Comments” by using the search function.

SUPPLEMENTARY INFORMATION:

Description: The proposed information collection activity is the first phase of a larger study that intends to assess the impact and implementation of LifeSet, a program that provides services and supports to young adults ages 17 to 21 with previous child welfare involvement. The program aims to support young adults in their transition from foster care to independent living in the areas of education, employment and earnings, housing and economic well-being, social support, well-being, health and safety, and criminal involvement. It focuses on helping young adults identify and achieve their goals while developing the skills necessary for independent living.

The impact study will assess the effects of young adults’ participation in LifeSet on outcomes in the primary (i.e., confirmatory) domains of education and employment, housing stability, social support, well-being, health and safety, and criminal involvement. These outcomes have been identified by the implementing agency as the main areas they expect to target for positive program impacts. In addition, the impact study will explore the effects of participation in the secondary (i.e., exploratory) domains of mental health, criminal justice system contact, intimate partner violence, and economic well-being. The study will utilize a randomized controlled design.

Information collection activities will take over three years and will include collection of administrative data from the state child welfare agency, the program developer, the local program provider agencies, the National Student Clearinghouse, unemployment insurance and employer wage records, the National Directory of New Hires, the state homelessness management information system, the state department of corrections, the state juvenile justice commission, the state court probation services division, and the state department of human services division of family development, as well as survey interviews with program participants and young adults receiving services as usual.

The implementation study will collect information through phone calls and site visits to the participating program and child welfare agency. Information collection activities include interviews and focus groups with administrators and staff from the program developer, child welfare agency, and program providers.

This evaluation is part of a larger project to help ACF build the evidence base in child welfare through rigorous evaluation of programs, practices, and policies. The activities and products from this project will contribute to evidence building in child welfare and help to determine the effectiveness of a program for youth formerly in foster care on young adult outcomes.

Respondents: Program participants, young adults receiving services as usual, agency and program administrators and staff, other program stakeholders.

ANNUAL BURDEN ESTIMATES

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<th>Instrument</th>
<th>Respondents</th>
<th>Number of respondents (total over request period)</th>
<th>Number of responses per respondent (total over request period)</th>
<th>Average burden per response (in hours)</th>
<th>Total burden (in hours)</th>
<th>Annual burden (in hours)</th>
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<td>Child Welfare Agency Administrators.</td>
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ANNUAL BURDEN ESTIMATES—Continued

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<tbody>
<tr>
<td>Site Visit 2 Focus Group Guide for Staff. Baseline Youth Survey ....... Administrative data file ......</td>
<td>Licensed LifeSet Experts Provider Agency Administrators LifeSet Developer Administrators LifeSet Specialists .......... LifeSet Team Supervisors Youth Formerly in Foster Care. Agency and Program Staff</td>
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<td>1.5 0.6 5</td>
<td>18 360 60</td>
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Estimated Total Annual Burden Hours: 160.


Mary B. Jones,
ACF/OPRE Certifying Officer.
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2017–D–2462]

Eligibility for the Index of Legally Marketed Unapproved New Animal Drugs for Minor Species; Request for Comments

AGENCY: Food and Drug Administration, Health and Human Services (HHS).

ACTION: Notice; request for comments.

SUMMARY: The Food and Drug Administration (FDA, we, or the Agency) is soliciting comments on our current policy on eligibility for indexing. Indexing is the process of adding an unapproved drug for a minor species to our index of legally marketed unapproved new animal drugs for minor species (the Index). Except for in some early non-food life stages, members of a food-producing minor species are not eligible for indexing, even if a subset of animals within a food-producing minor species is not intended to be consumed by humans or food-producing animals. Specifically, we are requesting comment on this policy.

DATES: Submit either electronic or written comments on the notice by September 22, 2021.

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 September 22, 2021. The https://www.regulations.gov electronic filing system will accept comments until 11:59 p.m. Eastern Time at the end of September 22, 2021. 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 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 be made publicly available, 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–D–2462 for “Eligibility for the Index of Legally Marketed Unapproved New Animal Drugs for Minor Species.”

  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, 240–402–7500.

- 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.” The Agency 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...