

FR 58252 on November 19, 2018. A public meeting took place on November 28, 2018 in the Downtown San Diego community. In preparing this Final EA, GSA considered public comments received regarding the Draft EA during the public review period.

After careful consideration of the environmental analysis and associated environmental effects of the Proposed Action Alternative and No Action Alternative, the purpose and need for the Project, and comments received on the Draft EA, GSA will be implementing the Proposed Action Alternative.

Finding

Pursuant to the provision of GSA Order ADM 1095.1F, the PBS NEPA Desk Guide, and the regulations issued by the Council on Environmental Quality (CEQ; 40 CFR parts 1500 to 1508), this notice advises the public of our finding that the Proposed Action will not significantly affect the quality of the human environment.

Basis for Finding

The environmental impacts of constructing the proposed structural enhancements were considered in the Final EA pursuant to the National Environmental Policy Act (NEPA) and the CEQ regulations implementing NEPA. No significant impacts on the environment would occur with implementation of best management practices and avoidance, minimization, and mitigation measures identified in the Final EA.

The Final EA is available for review at the San Diego Central Library, 330 Park Boulevard, San Diego, CA 92101. The Final EA and FONSI can also be viewed on the GSA website at <https://www.gsa.gov/real-estate/environmental-programs/gsa-nepa-implementation/nepa-library>.

The Finding of No Significant Impact will be signed thirty (30) days after the publication of this notice, provided that no information leading to a contrary finding is received or comes to light during this period.

Dated: July 24, 2019.

Jared Bradley,

Director, Portfolio Management Division,
Pacific Rim Region, Public Buildings Service.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity: 2020 Residential Energy Consumption Survey (RECS), Low Income Home Energy Assistance Program (LIHEAP) Administrative Data Matching (OMB #0970-0486)

AGENCY: Office of Community Services; Administration for Children and Families; HHS.

ACTION: Request for Public Comment.

SUMMARY: The Office of Community Services (OCS) is requesting an extension for the collection and reporting of 2020 administrative household data for state LIHEAP grantees' LIHEAP recipients. OMB approved the original collection under #0970-0486.

DATES: *Comments due within 60 days of publication.* 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.

ADDRESSES: Copies of the proposed collection of information can be obtained and comments may be forwarded by emailing infocollection@acf.hhs.gov. Alternatively, copies can also be obtained 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. All requests, emailed or written, should be identified by the title of the information collection.

SUPPLEMENTARY INFORMATION:

Description: The purpose of this information collection is to provide data that will allow OCS to identify LIHEAP recipients that respond to the upcoming Residential Energy Consumption Survey (RECS), which The U.S. Energy Information Administration (EIA) is planning to conduct in 2020. The EIA conducts the RECS survey to provide periodic national and regional data on residential energy use in the United States. OCS uses RECS data to furnish Congress and the Administration for Children and Families (ACF) with important national and regional descriptive data on the energy needs of low-income households.

In 2015, state LIHEAP grantees provided household-level recipient data to identify LIHEAP recipients that participated in the 2015 RECS. ACF is

requesting no changes in the type of data or the form of data collection for the 2020 extension of the project. The administrative household data already is collected by State grantees and used to complete the annual LIHEAP Household Report (OMB Control No. 0970-0060) and the annual LIHEAP Performance Data Form (OMB Control No. 0970-0449).

The LIHEAP data collected for this effort will be used by OCS to study the impact of LIHEAP on income eligible and recipient households in accordance with 42 U.S.C. 8629(b)(2). The information is being collected for use in development of the Department's annual LIHEAP Report to Congress and the annual LIHEAP Home Energy Notebook. The collection of this data is authorized by the LIHEAP statute, which requires the Secretary, following consultation with the Secretary of Energy, to provide for the collection of specific information on the characteristics of LIHEAP recipient and LIHEAP eligible households within each State. This includes collecting information that is reasonably necessary to carry out the provisions of the LIHEAP statute if that information is not collected by any other agency of the Federal Government.

State LIHEAP grantees will be asked to furnish data for LIHEAP recipient households that reside in areas included in the RECS sample.

The following are the specific data items grantees will report for each household:

- Name
- Address (including ZIP code)
- Gross Income
- Household or Client ID
- Household Size
- Heating assistance awarded
- Amount of heating assistance
- Date of heating assistance
- Cooling assistance awarded
- Amount of cooling assistance
- Date of cooling assistance
- Crisis Assistance awarded
- Amount of crisis assistance
- Date of crisis assistance
- Other Assistance awarded
- Amount of other assistance
- Date of other assistance
- Presence of children 5 or younger
- Presence of adult 60 or older
- Presence of disabled

The following are optional data items that grantees can provide if the data are available in your database:

- Tenancy (*i.e.*, own or rent)
- Type(s) of fuel used
- Heat included in rent

State LIHEAP grantees can provide the data elements in the selected format of their choosing.

The confidentiality of client data will be strictly protected as part of the project. LIHEAP application client

waivers allow grantees to share information with OCS and its contractors.

Respondents: 51 (State Governments and the District of Columbia)

ANNUAL BURDEN ESTIMATES

Instrument	Total number of respondents	Total number of responses per respondent	Average burden hours per response	Total burden hours	Annual burden hours
Action Transmittal LIHEAP-AT-2020-04 Extension of the FY 2015 RECS LIHEAP Administrative Data Matching to FY 2020	51	1	24	1,224	408

Estimated Total Annual Burden Hours: 408.

As LIHEAP is a block grant, there is varying capacity to collect and report data among grantees. The estimated burden hours displayed above are for the average LIHEAP grantee. All LIHEAP grantees have existing data systems to collect, maintain, and analyze this data to complete annual reporting requirements.

Comments: 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.

Authority: 42 U.S.C. 8629(a).

Mary B. Jones,

ACF/OPRE Certifying Officer.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2014-D-1461]

Rare Pediatric Disease Pediatric Priority Review Vouchers; Draft Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of a draft guidance for industry entitled "Rare Pediatric Disease Priority Review Vouchers." This draft guidance is a revision of the guidance of the same title that published in 2014. This draft guidance provides information on the rare pediatric disease priority review voucher program under the Federal Food, Drug, and Cosmetic Act (FD&C Act), under which FDA will award priority review vouchers to sponsors of certain rare pediatric disease product applications that meet the relevant statutory criteria. These priority review vouchers can be used when submitting future human drug marketing applications that would not otherwise qualify for priority review. Because there exists a need for products for rare pediatric diseases, this program is intended to encourage development of new drug and biological products for prevention and treatment of certain rare pediatric diseases.

DATES: Submit either electronic or written comments on the draft guidance by September 30, 2019 to ensure that the Agency considers your comment on this draft guidance before it begins work on the final version of the guidance. Submit electronic or written comments on the information collection burden by September 30, 2019.

ADDRESSES: You may submit comments on any guidance at any time 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 be made 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-2014-D-1461 for "Rare Pediatric Disease Priority Review Vouchers." Received comments 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