Division of Regulations Development, Attention: Document Identifier/OMB Control Number __ __ __ __ Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, you may make your request using one of the following:

2. Email your request, including your address, phone number, OMB number, and CMS document identifier, to Paperwork@cms.hhs.gov.
3. Call the Reports Clearance Office at (410) 786–1326.

FOR FURTHER INFORMATION CONTACT: Reports Clearance Office at (410) 786–1326.

SUPPLEMENTARY INFORMATION:

Contents

This notice sets out a summary of the use and burden associated with the following information collections. More detailed information can be found in each collection’s supporting statement and associated materials (see ADDRESSES).

CMS–10527 Annual Eligibility Redetermination, Product Discontinuation and Renewal Notices

Under the PRA (44 U.S.C. 3501–3520), federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. The term “collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA requires federal agencies to publish a 60-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice.

1. Type of Information Collection Request: Revision a currently approved collection; Title of Information Collection: Annual Eligibility Redetermination, Product Discontinuation and Renewal Notices; Use: Section 1411(f)(1)(B) of the Affordable Care Act directs the Secretary of Health and Human Services (the Secretary) to establish procedures to re-determine the eligibility of individuals on a periodic basis in appropriate circumstances. Section 1321(a) of the Affordable Care Act provides authority for the Secretary to establish standards and regulations to implement the statutory requirements related to Exchanges, QHPs and other components of title I of the Affordable Care Act. Under section 2703 of the PHS Act, as added by the Affordable Care Act, and sections 2712 and 2741 of the PHS Act, enacted by the Health Insurance Portability and Accountability Act of 1996, health insurance issuers in the group and individual markets must guarantee the renewability of coverage unless an exception applies. The final rule “Patient Protection and Affordable Care Act; Annual Eligibility Redeterminations for Exchange Participation and Insurance Affordability Programs; Health Insurance Issuer Standards Under the Affordable Care Act, Including Standards Related to Exchanges” (79 FR 52994), provides that an Exchange may choose to conduct the annual redetermination process for a plan year (1) in accordance with the existing procedures described in 45 CFR 155.335; (2) in accordance with procedures described in guidance issued by the Secretary for the coverage year; or (3) using an alternative proposed by the Exchange and approved by the Secretary. The guidance document “Guidance on Annual Redeterminations for Coverage for 2013” contains the procedures that the Secretary has specified, as noted in (2) above, until the issuance of further guidance. These procedures will be adopted by the Federally-facilitated Exchange. Under this option, the Exchange will provide three notices. These notices may be combined. The final rule also amends the requirements for product renewal and re-enrollment (or non-renewal) notices to be sent by Qualified Health Plan (QHP) issuers in the Exchanges and specifies content for these notices. The guidance document “Draft Updated Federal Standard Renewal and Product Discontinuation Notices” provides draft updated Federal standard notices for product discontinuation and renewal that would be sent by issuers of individual market QHPs and issuers in the individual market. Issuers in the small group market may use the draft Federal standard small group notices released in the June 26, 2014 bulletin “Draft Standard Notices When Discontinuing or Renewing a Product in the Small Group or Individual Market”, or any forms of the notice otherwise permitted by applicable laws and regulations. States that are enforcing the Affordable Care Act may develop their own standard notices, for product discontinuances, renewals, or both, provided the State-developed notices are at least as protective as the Federal standard notices. Form Number: CMS–10527 (OMB Control Number: 0938–1254); Frequency: Annually; Affected Public: Private Sector, State Governments; Number of Respondents: 2,945; Number of Responses: 12,224; Total Annual Hours: 149,186. (For policy questions regarding this collection, contact Russell Tipps at 301–492–4371.)


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

BILLOWING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

[CFDA: 93.592]

Announcing the Intent To Award a Single-Source Expansion Supplement Grant to the National Domestic Violence Hotline

AGENCY: Family and Youth Services Bureau, ACYF, ACF, HHS.

ACTION: This notice announces the intent to award a single-source expansion supplement grant under the Family Violence Prevention and Services Act (PVPSA) national domestic violence hotline grant program to the National Domestic Violence Hotline (Hotline) in Austin, TX.

SUMMARY: The Administration for Children and Families (ACF), Administration on Children, Youth and Families (ACYF), Family and Youth Services Bureau (FYSB), Division of Family Violence and Prevention Services (DFVPS) announces its intent to award a cooperative agreement of up to $3,750,000 as a single-source expansion supplement to the National Domestic Violence Hotline (Hotline) in Austin, TX.

DATES: The period of support for the single-source expansion supplement is September 30, 2016 through September 29, 2017.

FOR FURTHER INFORMATION CONTACT: Angela Yannelli, Senior Program Specialist, Family Violence Prevention...
and Services Program, 330 C Street SW., 3rd Floor, Suite 3621B, Washington, DC 20201. Telephone: 202–401–5524; Email: Angela.Yannelli@acf.hhs.gov.

SUPPLEMENTARY INFORMATION: The Hotline in Austin, TX, is funded under the Family Violence Prevention and Services Act (FVPSA) program to operate the 24-hour, national, toll-free telephone hotline that provides information and assistance to adult and youth victims of family violence, domestic violence, or dating violence, and to the family and household members of such victims, and to persons affected by the victimization. The supplemental award will expand the capacity of the Hotline’s current efforts by focusing on the development of a tribal hotline and by providing additional phone advocates to ensure that the Hotline can answer all contacts. The award will also assist in developing the “Love Is Respect” Web site (http://www.loveisrespect.org) into a complete resource for teens and youth seeking to prevent and end abusive relationships.


Christopher Beach,
Senior Grants Policy Specialist, Division of Grants Policy, Office of Administration, Administration for Children and Families.

SUPPLEMENTARY INFORMATION:

Email:
3rd Floor, Suite 3621B, Washington, DC 20408 Federal Register

25408 Federal Register / Vol. 81, No. 82 / Thursday, April 28, 2016 / Notices

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration


Compliance Policy Guide on Crabmeat—Fresh and Frozen—Adulteration With Filth, Involving the Presence of Escherichia coli

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing the availability of a Compliance Policy Guide (CPG) relating to fresh and frozen crabmeat adulteration with filth involving the presence of Escherichia coli (E. coli). The CPG updates the previously issued CPG on this topic. The CPG provides guidance for FDA staff on the level of E. coli in crabmeat at which we may consider the crabmeat to be adulterated with filth.

DATES: Submit electronic or written comments on FDA’s CPGs at any time.

ADDRESSES: You may submit comments as follows:

Electronic Submissions
Submit electronic comments in the following way:

• Federal eRulemaking Portal: http://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to http://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 http://www.regulations.gov.

• If you want to submit a comment with confidential information that you do not wish to be made public, 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: http://www.fda.gov/regulatoryinformation/dockets/default.htm.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to http://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 and/or go to the Division of Dockets Management, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

Submit written requests for single copies of the guidance to Office of Policy and Risk Management, Office of Regulatory Affairs, Office of Global Regulatory Operations and Policy, Food and Drug Administration, 12420 Parklawn Dr., Rockville, MD 20857. Send two self-addressed adhesive labels to assist that office in processing your request. See the SUPPLEMENTARY INFORMATION section for electronic access to the guidance.

FOR FURTHER INFORMATION CONTACT:

SUPPLEMENTARY INFORMATION: