

requires prospective researchers who need access to confidential data to complete a research proposal. Researchers self-select whether they need access to confidential data to answer their research questions. The RDC requires the researcher to complete a research proposal, so NCHS understands the research proposed. The completed proposal is sent to NCHS through the SAP portal for review and adjudication. If the research proposal is approved by NCHS, then the researcher must fill out data security forms. If the

researcher will access the data at a Research Data Center, then the “Data Use Agreement Form” and the “Designated Agent Agreement Form” would need to be completed and returned to NCHS. If the researcher will access the data through the NCHS Virtual Data Enclave (VDE), then the “VDE Data Use Agreement Form”, “Data Use Agreement Form” and the “Designated Agent Agreement Form” would need to be completed and returned to NCHS.

To capture the information needed to adjudicate a researcher’s commitment to

protect confidential NCHS data, researchers must complete and sign these data security forms. This request allows for both researcher signature and the time per response for a total estimated annual burden total of 110 hours (330 hours for a three-year clearance period). There is no cost to a researcher other than their time to complete the forms, unless the researcher has to pay a nominal notary fee for services incurred. The resulting information will be used for NCHS internal purposes.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Avg. burden per response (in hrs.)	Total burden (in hrs.)
Researcher	Research Data Center proposal	110	1	1	110
Total					110

Jeffrey M. Zirger,

Lead, Information Collection Review Office, Office of Public Health Ethics and Regulations, Office of Science, Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

[Office of Management and Budget #: 0970-0477]

Submission for Office of Management and Budget Review; Generic Clearance for Reviewer Recruitment Forms

AGENCY: Office of Planning, Research, and Evaluation, Administration for Children and Families, U.S. Department of Health and Human Services.

ACTION: Request for public comments.

SUMMARY: The Administration for Children and Families (ACF) proposes to extend approval of the existing overarching generic clearance for Reviewer Recruitment Forms (Office of Management and Budget (OMB) #: 0970-0477). No changes are proposed to the terms of the overarching generic. Burden estimates have been updated.

DATES: Comments due May 1, 2026.

ADDRESSES: The public may view and comment on this information collection request at: https://www.reginfo.gov/public/do/PRAViewICR?ref_nbr=202603-0970-008. You can also obtain copies of the proposed collection of information by emailing infocollection@acf.hhs.gov. Identify all emailed requests by the title of the information collection.

SUPPLEMENTARY INFORMATION:

Description: The overarching generic clearance for Reviewer Recruitment Forms provides ACF with the opportunity to collect information from potential reviewers, such as those who review grant proposals, conference proposals, research/evaluation plans, study designs, report drafts, and/or other ACF materials.

ACF developed this generic because each program office within ACF has slightly different needs for information about reviewer applicants based on the specific activities for which reviewers are needed, yet the individual forms submitted under the generic will serve an identical function. The overarching purpose is to select qualified reviewers for ACF review processes and activities based on professional qualifications. Information will be collected through questions on forms and documents provided by candidates. Example documents include writing samples and curriculum vitae and/or resumes. ACF

uses the information collected to recruit well-qualified reviewers with relevant background experience and knowledge.

The abbreviated clearance process of the generic clearance allows program offices to gather a suitable pool of candidates within the varied time periods available for reviewer recruitment.

These forms submitted under this generic will be voluntary, low-burden and uncontroversial. Currently approved information collections are available for review on [RegInfo.gov: https://www.reginfo.gov/public/do/PRAICList?ref_nbr=202603-0970-011](https://www.reginfo.gov/public/do/PRAICList?ref_nbr=202603-0970-011).

Respondents: Individuals who may apply to review materials for ACF.

Annual Burden Estimates

The following burden estimates include burden associated with currently approved individual requests that ACF expects will be extended through this extension request and an estimate of burden for potential new requests under this generic. Based on the past 3 years and with a goal to reduce burden moving forward, the number of respondents has been reduced by 50 percent and the estimated time per response by 34 percent, from 30 to 20 minutes. Overall, burden for potential new requests is 67 percent less than the currently approved umbrella generic.

POTENTIAL NEW REQUESTS

Instrument	Number of respondents (total over request period)	Number of responses per respondent (total over request period)	Avg. burden per response (in hours)	Total burden (in hours)
New Reviewer Recruitment Forms	1500	1	.33	495

ONGOING CURRENTLY APPROVED REQUESTS

Generic information collection title	Number of respondents	Annual number of responses per respondent	Average burden hours per response	Annual burden hours	Total burden hours
Administration for Native Americans (ANA) Panel Reviewer Profile Questionnaire	300	1	0.40	120	360
Children's Bureau Grant Reviewer Recruitment Module	250	1	0.08	21	63
Eligibility Information from Applicants: Reviewer Information Form for General Reviewer and for Specific Reviewer	95	1	0.17	16	48
Grant Reviewer Recruitment for the Family and Youth Services Bureau (FYSB) Discretionary Grant Programs	432	1	0.13	58	174
Office of Planning, Research, and Evaluation, Division of Economic Independence Information Collection for Prospective Grant Reviewer Opportunities	250	1	0.08	21	63
Total Estimated Annual Burden				236	708

Authority: Public Health Service (PHS) Act, Sections 799(f) and 806(e).

Mary C. Jones,
ACF/OPRE Certifying Officer.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2026-N-2741]

Consideration of Acceptable Market Name Change for Certain Rockfish (*Sebastes* spp.); Request for Information

AGENCY: Food and Drug Administration (FDA), U.S. Department of Health and Human Services (HHS).

ACTION: Notice; request for information.

SUMMARY: The Food and Drug Administration (FDA or we) is requesting data and information to help make an evidence-based determination that balances food safety, regulatory clarity, and industry interest regarding a potential update to the acceptable market name for the following fish: *Sebastes alutus*, *Sebastes borealis*, *Sebastes ciliatus*, *Sebastes crameri*, *Sebastes entomelas*, *Sebastes flavidus*, *Sebastes goodei*, *Sebastes levis*, *Sebastes melanops*, *Sebastes miniatus*, *Sebastes*

ovalis, *Sebastes paucispinis*, *Sebastes pinniger*, *Sebastes proriger*, *Sebastes reedi*, *Sebastes ruberrimus*, *Sebastes rufus*, and *Sebastes serranoides*.

DATES: Either electronic or written comments on the notice must be submitted by May 1, 2026.

ADDRESSES: You may submit comments and information as follows. Please note that late, untimely filed comments will not be considered. The <https://www.regulations.gov> electronic filing system will accept comments until 11:59 p.m. Eastern Time at the end of May 1, 2026. Comments received by mail/hand delivery/courier (for written/paper submissions) will be considered timely if they are received 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 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-2026-N-2741 for "Consideration of Acceptable Market Name Change for Certain Rockfish (*Sebastes* spp.); Request for Information." Received comments, those filed in a timely manner (see **ADDRESSES**), will be placed in the docket and, except for those