

compliance with federal and programmatic regulations and policies. To review currently approved PPRs under this generic, see: <https://>

[www.reginfo.gov/public/do/PRAICList?ref\\_nbr=202206-0970-004](http://www.reginfo.gov/public/do/PRAICList?ref_nbr=202206-0970-004).

*Respondents:* ACF funding recipients.

#### Annual Burden Estimates

ACF is requesting an increase in burden to reflect use over the past 3 years and anticipated use in the next 3 years.

Instrument	Total number of respondents	Total number of responses per respondent	Average burden hours per response	Total burden hours
Program Specific PPRs .....	800	2.3	5	9,200

*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.

Mary B. Jones,  
ACF/OPRE Certifying Officer.

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Administration for Children and Families

#### Proposed Information Collection Activity; National Communication System for Runaway and Homeless Youth, Currently Operated by the National Runaway Safeline (NRS) Data Collection (New Collection)

**AGENCY:** Family and Youth Services Bureau, Administration for Children and Families, HHS.

**ACTION:** Request for public comment.

**SUMMARY:** The Family and Youth Services Bureau's (FYSB) Runaway and Homeless Youth Division has a legislative requirement to fund a National Communication System, which is currently operated by the National Runaway Safeline (NRS). The NRS provides information, referral services, crisis intervention, and prevention

resources to vulnerable youth at risk of running away and/or becoming homeless and their families or legal guardians at no cost. When necessary, the NRS refers runaway and homeless youth to shelters, counseling, medical assistance, and other vital services. The NRS collects information from all contacts with youth and adults connecting with the NRS (*i.e.*, parents, family members, legal guardians, service providers) on a voluntary basis to inform crisis services and develop an annual report on the information collected during calls, chats, emails, and forum posts from young people who reached out to the NRS's crisis services.

**DATES:** *Comments due within 60 days of publication.* In compliance with the requirements of the Paperwork Reduction Act of 1995, ACF is soliciting public comment on the specific aspects of the information collection described above.

**ADDRESSES:** You can obtain copies of the proposed collection of information and submit comments by emailing [infocollection@acf.hhs.gov](mailto:infocollection@acf.hhs.gov). Identify all requests by the title of the information collection.

#### SUPPLEMENTARY INFORMATION:

*Description:* The NRS is required to have a system for collecting and analyzing data to report on calls, emails, chat, texts, and online messages received as well as other information, such as prevention resources, referrals, demographics, and visitors to the NRS website. The NRS must submit monthly and semi-annual reports that includes the following:

- Number of calls received, answered, and missed.
- Number of chats, emails, and texts received; number of chats, emails, and texts answered; and number of chats, emails, and texts that were missed and did not receive a response, in which the users are youth in crisis, runaway youth, and youth experiencing homelessness.

- Number of parents, legal guardians, and service providers contacting the NRS and the type of resources, interventions, and technical support/assistance requested and provided.

- Number and type of prevention materials disseminated to communities, especially to underserved populations.

- Number and type of unique visitors to the NRS' website.

- Information on referrals provided and where youth were referred for services.

- Information on the callers' or users' demographics and where they were located when contacting the NRS.

- Information on the prevention materials developed and disseminated by the NRS.

- Information and analysis of the latest trends and their impact on runaway prevention.

The NRS will use two online forms, one form to collect relevant information disclosed during calls, emails, and forum posts and a second online form to collect information from chats. All data will be provided to FYSB in the aggregate and no personally identifiable data are collected.

The information collected will allow FYSB to better understand the types of services needed by youth contacting the NRS, as well as to identify outreach and prevention strategies to increase the visibility of the NRS services among youth experiencing housing instability, homelessness, youth who runaway, and youth in crisis. Additionally,

The findings from this data collection will be included in a required Report to Congress to provide accurate information on the status of youth in crisis and runaway and homeless youth nationwide.

*Respondents:* Youth and adults who contact the National Runaway Safeline during calls, chats, emails, and forum posts.

## 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
Youth in Crisis Form .....	47,175	1	.23	10,850	3,617
NRS Live Chat Form .....	29,679	1	.65	19,291	6,430

*Estimated Total Annual Burden Hours: 10,047.*

**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:** Section 331 of the Runaway and Homeless Youth Act authorizes the award of grants for the National Communication System for Runaway and Homeless Youth (34 U.S.C. 11231).

**Mary B. Jones,**  
*ACF/OPRE Certifying Officer.*

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## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. FDA-2022-N-0008]

### Request for Nominations on Public Advisory Panels of the Medical Devices Advisory Committee

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

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is requesting that any industry organizations interested in participating in the selection of nonvoting industry representatives to serve on certain panels of the Medical Devices Advisory Committee (MDAC or

Committee) in the Center for Devices and Radiological Health (CDRH) notify FDA in writing. FDA is also requesting nominations for nonvoting industry representatives to serve on certain device panels of the MDAC in the CDRH. A nominee may either be self-nominated or nominated by an organization to serve as a nonvoting industry representative. Nominations will be accepted for current and upcoming vacancies effective with this notice.

**DATES:** Any industry organization interested in participating in the selection of an appropriate nonvoting member to represent industry interests must send a letter stating that interest to the FDA by August 5, 2022 (see sections I and II of this document for further details). Concurrently, nomination materials for prospective candidates should be sent to FDA by August 5, 2022.

**ADDRESSES:** All statements of interest from industry organizations interested in participating in the selection process of nonvoting industry representative nomination should be sent to Margaret Ames (see **FOR FURTHER INFORMATION CONTACT**). All nominations for nonvoting industry representatives should be submitted electronically by accessing the FDA Advisory Committee Membership Nomination Portal: <https://www.accessdata.fda.gov/scripts/FACTRSPortal/FACTRS/index.cfm> or by mail to Advisory Committee Oversight and Management Staff, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 32, Rm. 5103, Silver Spring, MD 20993-0002. Information about becoming a member of an FDA advisory committee can also be obtained by visiting FDA's website at <https://www.fda.gov/AdvisoryCommittees/default.htm>.

**FOR FURTHER INFORMATION CONTACT:** Margaret Ames, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 66, Rm. 5213, Silver Spring,

MD 20993, 301-796-5960, email: [margaret.ames@fda.hhs.gov](mailto:margaret.ames@fda.hhs.gov).

**SUPPLEMENTARY INFORMATION:** The Agency is requesting nominations for nonvoting industry representatives to the panels listed in the table in this document.

### I. Medical Devices Advisory Committee

The Committee reviews and evaluates data on the safety and effectiveness of marketed and investigational devices and makes recommendations for their regulation. The panels engage in a number of activities to fulfill the functions the Federal Food, Drug, and Cosmetic Act (FD&C Act) envisions for device advisory panels. With the exception of the Medical Devices Dispute Resolution Panel, each panel, according to its specialty area, advises the Commissioner of Food and Drugs (the Commissioner) regarding recommended classification or reclassification of devices into one of three regulatory categories; advises on any possible risks to health associated with the use of devices; advises on formulation of product development protocols; reviews premarket approval applications for medical devices; reviews guidelines and guidance documents; recommends exemption of certain devices from the application of portions of the FD&C Act; advises on the necessity to ban a device; and responds to requests from the Agency to review and make recommendations on specific issues or problems concerning the safety and effectiveness of devices. With the exception of the Medical Devices Dispute Resolution Panel, each panel, according to its specialty area, may also make appropriate recommendations to the Commissioner on issues relating to the design of clinical studies regarding the safety and effectiveness of marketed and investigational devices. The Committee also provides recommendations to the Commissioner or designee on complexity categorization of in vitro diagnostics under the Clinical Laboratory Improvement Amendments of 1988.