

for damages arising from or related to Challenge activities.

9. Understands that circulation of findings could be worldwide, and that the Federal Government will not compensate the participants for any use; winners shall receive a one-time cash prize as set forth in this announcement. The winner and runner-up from Phase 2 will be posted on the AHRQ website.

10. Understands that AHRQ reserves the right to cancel, suspend, and/or modify this prize contest, or any part of it, for any reason, in AHRQ's sole discretion. AHRQ also reserves the right not to award any prizes if no entries are deemed worthy.

11. Understands that AHRQ will not select a winner that is named on the Excluded Parties List System (EPLS).

Intellectual Property (IP) Rights

1. Each participant retains title and full ownership in and to their submission. Participants expressly reserve all intellectual property rights not expressly granted.

2. By participating in the Challenge, each participant (whether participating singly or in a group) acknowledges that he or she is the sole author or owner of, or has a right to use, any copyrightable works that the submission comprises, that the works are wholly original with the participant (or is an improved version of an existing work that the participant has sufficient rights to use and improve), and that the submission does not infringe any copyright or any other rights of any third party of which participant is aware.

3. In addition, each participant (whether participating singly or in a group) grants to the U.S. Government a paid-up, nonexclusive, royalty-free, irrevocable worldwide license in perpetuity, and the right to reproduce, publish, post, link to, share, display publicly (on the web or elsewhere) and prepare derivative works, including the right to authorize others to do so on behalf of the U.S. Government.

4. Each participant must clearly delineate any intellectual property and/or confidential commercial information contained in a submission that the participant wishes to protect as proprietary data, in accordance with Additional Rules of Participation No. 4.

5. If the submission includes any third-party works (such as third-party

content or open-source code), the participant must be able to provide, upon request, documentation of all appropriate licenses and releases for use of such third-party works. If the participant cannot provide documentation of all required licenses and releases, AHRQ reserves the right, in its sole discretion, to disqualify the submission.

Dated: September 20, 2022.

Marquita Cullom,
Associate Director.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; ORR Services for Survivors of Torture Program Data Points and Performance Progress Reports (New Collection)

AGENCY: Office of Refugee Resettlement, Administration for Children and Families, HHS.

ACTION: Request for public comment.

SUMMARY: The Administration for Children and Families' (ACF) Office of Refugee Resettlement (ORR) intends to collect demographic, programmatic, and outcome data on Services for Survivors of Torture (SOT) grant recipients and the clients they serve. This data collection will allow ORR to learn more about the populations served; the types and effectiveness of services provided; methods, challenges, and facilitators of implementing services; and grant recipients' progress towards programmatic goals. ORR will collect these data on the new cohort of Services for SOT grant recipients; ORR collected information from the previous grantee cohort under the Generic Performance Progress Report (OMB #0970-0490). ORR has made changes to the data collection instruments for use in the new cohort.

DATES: *Comments due within 30 days of publication.* OMB is required to make a decision concerning the collection of information 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" or by using the search function. 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: ORR proposes to use the Program Data Points Form (PDPs) and Performance Progress Reports (PPRs) to collect data on the Services for SOT grant recipients and their clients. In 2019, ORR began requiring the Services for SOT grant recipients to collect and report their PDPs through the ORR Refugee Arrivals Data System (RADS), an information technology platform used for enhanced data collection and record keeping. The new cohort of Services for SOT grant recipients, who will receive 5-year awards in September 2022, will also provide these data points to ORR using RADS. Grant recipients will provide aggregated data on new and continuing clients annually, including demographic information, characteristics related to experiences of torture, services received, and well-being across six outcome domains. Grant recipients will also provide information about community attendance at trainings and pro-bono services donated to the program. In the PPRs, grant recipients will provide primarily narrative information on grant-funded activities and progress towards grant goals biannually.

Information collected will be used in aggregate by ORR to provide reports to stakeholders, including a required report to Congress, and responses to funding requests.

Respondents: Services for SOT grant programs (this may include non-profit social service, health, and higher education organizations, states, municipalities, and for-profit organizations).

ANNUAL BURDEN ESTIMATES

Instrument	Total number of respondents	Number of responses per respondent	Average burden hours per response	Annual burden hours
Program Data Points Form (PDPs)	35	1	6	210

ANNUAL BURDEN ESTIMATES—Continued

Instrument	Total number of respondents	Number of responses per respondent	Average burden hours per response	Annual burden hours
Performance Progress Reports (PPRs)—Parts A and B	35	2	6	420

Estimated Total Annual Burden Hours: 630.

Authority: Section 5(a) of the “Torture Victims Relief Act of 1998,” Public Law 105–320 (22 U.S.C. 2152 note) Assistance for Treatment of Torture Victims.

Mary B. Jones,

ACF/OPRE Certifying Officer.

[FR Doc. 2022–20746 Filed 9–23–22; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Community Living

Agency Information Collection

Activities: Proposed Collection; Public Comment Request of the National Network of University Centers for Excellence in Developmental Disabilities Education, Research, and Service (UCEDDs) OMB Control Number 0985–0030

AGENCY: Administration for Community Living, Department of Health and Human Services.

ACTION: Notice.

SUMMARY: The Administration for Community Living (ACL) is announcing an opportunity for the public to comment on the proposed collection of information listed above. Under the Paperwork Reduction Act of 1995 (PRA), Federal agencies are required to publish a notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This Information Collection (IC) Revision solicits comments on the information collection requirements relating to the National Network of University Centers for Excellence in Developmental Disabilities Education, Research, and Service (UCEDDs).

DATES: Comments on the collection of information must be submitted electronically by 11:59 p.m. (EST) or postmarked by November 25, 2022.

ADDRESSES: Submit electronic comments on the collection of information to Pamela O’Brien at pamela.obrien@acl.hhs.gov. Submit

written comments on the collection of information to Administration for Community Living, 330 C Street SW, Washington, DC 20201, Attention: Pamela O’Brien.

FOR FURTHER INFORMATION CONTACT: Pamela O’Brien, 202–795–7417 or pamela.obrien@acl.hhs.gov.

SUPPLEMENTARY INFORMATION: 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. “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. The PRA requires Federal agencies to provide a 60-day notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, ACL is publishing a notice of the proposed collection of information set forth in this document.

With respect to the collection of information described below, ACL invites comments on our burden estimates or any other aspect of this collection of information, including:

- (1) whether the proposed collection of information is necessary for the proper performance of ACL’s functions, including whether the information will have practical utility;
- (2) the accuracy of ACL’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used to determine burden estimates;
- (3) ways to enhance the quality, utility, and clarity of the information to be collected; and
- (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques when appropriate, and other forms of information technology.

Section 104 (a) (42 U.S.C. 15004) of the Developmental Disabilities Assistance and Bill of Rights Act of 2000 (DD Act) directs the Secretary of Health and Human Services to develop

and implement a system of program accountability to monitor the grantees funded under the DD Act, including the UCEDDs. Section 154 (e) (42 U.S.C. 15064) of the DD Act includes requirements for the UCEDD Annual Report. The UCEDD Annual Report should contain information on progress made in achieving the projected goals of the Center for the previous year, including:

- (1) The extent of goal achievement;
- (2) A description of the strategies that contributed to achieving the goals;
- (3) The extent goals were not achieved, a description of factors that impeded the achievement;
- (4) An accounting of the manner in which funds paid to the Center . . . for a fiscal year were expended;
- (5) Information on proposed revisions to the goals; and
- (6) A description of successful efforts to leverage funds, other than funds made available under the DD Act.

The DD Act also states grantees must report on consumer satisfaction with:

- (1) The advocacy, capacity building, and systemic change activities initiated by the UCEDD;
- (2) The extent to which the UCEDD’s advocacy, capacity building, and systemic change activities provided results through improvements; and
- (3) The extent to which collaboration was achieved in the areas of advocacy, capacity building, and systemic change.

In addition to collecting the information required in the DD Act, this IC will also include elements needed to account for the activities supported by funding from the Centers for Disease Control and Prevention (CDC) to support access to vaccines for people with disabilities as well as the funds awarded under the American Rescue Plan Act to increase the Public Health Workforce (PHWF).

Finally, to ensure the UCEDD PPR is consistent with the Executive order on Advancing Racial Equity and Support for Underserved Communities Through the Federal Government and the Executive order on Advancing Equality for Lesbian, Gay, Bisexual, Transgender, Queer, and Intersex Individuals, ACL intends to determine whether the sexual orientation and gender identity (SOGI) data elements need to be adapted to ensure accessibility of the questions for