independent and safe in their communities.

In compliance with the PHS Act, ACL revised an ADSSP Data Reporting Tool (ADSSP–DRT) in 2013. The 2016 revised Alzheimer's and Dementia Program Data Reporting Tool (ADP–DRT) collects information about the

delivery of direct services by ADSSP and ADI–SSS grantees, as well as basic demographic information about service recipients. The 2016 version includes revisions to the approved 2013 version. The revised version would be in effect beginning 12/31/2016 and thereafter.

The proposed ADP–DRT can be found on AoA's Web site at: http://
nadrc.acl.gov/sites/default/files/
uploads/docs/2016%20Proposed%20
OMB%20Alzheimer%20Program%20
Reporting%20Tool.xlsx.

ACL estimates the burden of this collection of information as follows:

ANNUAL BURDEN ESTIMATES

Instrument	Type of respondent	Number of respondents	Responses per respondent	Burden hours per response	Total burden hours (annual)
ADP-DRT	Local Program Site	76 38	2 2	4.67 3.6	709.84 273.6

Estimated Total Annual Burden Hours: 983.44.

Dated: August 17, 2016.

Edwin L. Walker,

Acting Administrator & Assistant Secretary for Aging.

[FR Doc. 2016–20156 Filed 8–22–16; 8:45 am]

BILLING CODE 4154-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration on Community Living

Proposed Information Collection Activity; Comment Request; Protection and Advocacy Annual Program Performance Report and Statement of Goals and Priorities

AGENCY: Office of Program Support, Administration on Intellectual and Developmental Disabilities, Administration on Disability, Administration on Community Living, HHS

ACTION: Notice.

SUMMARY: This notice seeks to collect comments on revisions to an existing collection: Annual Protection and Advocacy Systems Program
Performance Report (0985–0027). State Protection and Advocacy (P&A) Systems in each State and Territory provide individual legal advocacy, systemic advocacy, monitoring and investigations to protect and advance the rights of people with developmental disabilities, using funding administered by the Administration on Intellectual and Developmental Disabilities, Administration on Disability, Administration on Community Living, HHS

The Developmental Disabilities and Bill of Rights Act, 42 U.S.C. 15044, requires each P&A to annually prepare

a report that describes the activities and accomplishments of the system during the preceding fiscal year and a Statement of Goals and Priorities (SGP) (0985-0034) for each coming fiscal year. P&As are required to annually report on "the activities, accomplishments, and expenditures of the system during the preceding fiscal year, including a description of the system's goals, the extent to which the goals were achieved, barriers to their achievement, the process used to obtain public input, the nature of such input, and how such input was used." To meet it statutory reporting requirements, P&As have used separate forms for submitting the annual report (0985-0027) and the SGP (0985-0034). It is proposed that the two be combined by creating the Protection and Advocacy Annual Program Performance Report and Statement of Goals and Priorities form. By combining the forms, P&As will have a reduced burden by submitting only one report annually. The combined form will also allow federal reviewers to analyze patterns more readily between goals and priority setting and program performance. The annual program performance report (PPR) and SGP is reviewed by federal staff for compliance and program outcomes. Information in the PPRs and SGPs is analyzed to create a national profile of programmatic compliance, program outcomes, and goals and priorities for P&A Systems. These profiles are used to track accomplishments against goals, develop technical assistance, and determine compliance with Federal requirements. Information collected in the unified report also will inform AIDD of trends in P&A advocacy, collaboration with other federally-funded entities, and best practices for efficient use of federal funds.

DATES: Submit written comments on the collection of information by October 24, 2016.

ADDRESSES: Submit written comments on the collection of information by email to: *Clare.Huerta@acl.hhs.gov*.

FOR FURTHER INFORMATION CONTACT:

Clare Huerta, Administration on Community Living, Administration on Intellectual and Developmental Disabilities, Office of Program Support, 330 C Street SW., DC, Washington, DC 20201, (202) 795–7301.

SUPPLEMENTARY INFORMATION: In compliance with the requirements of Section 506 (c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration on Community Living is soliciting public comment on the specific aspects of the information collection described above. The form is available at http://www.acl.gov/Programs/AIDD/Program_Resource_Search/Results_DDC.aspx#resources.

The Department specifically requests comments on: (a) Whether the proposed Collection of information is necessary for the proper performance of the function 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; (d) ways to minimize the burden information to be collected; and (e) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection technique comments and or other forms of information technology. Consideration will only be given to comments and suggestions submitted within 60 days of this publication.

Respondents: 57 Protection and Advocacy Systems.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of re- sponses per respondent	Average bur- den hours per response	Total burden hours
Protection and Advocacy Annual Program Performance Report and Statement of Goals and Priorities	57	1	90	5,130

Estimated Total Annual Burden Hours: 5,130.

Dated: August 17, 2016.

Edwin L. Walker,

Acting Administrator and Assistant Secretary for Aging.

[FR Doc. 2016–20161 Filed 8–22–16; 8:45 am]

BILLING CODE 4154-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Office of the Secretary

[Document Identifier: HHS-OS-0990-0275-30D]

Agency Information Collection Activities; Submission to OMB for Review and Approval; Public Comment Request

AGENCY: Office of the Secretary, HHS. **ACTION:** 30 day Notice.

SUMMARY: In compliance with section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the Office of the Secretary (OS), Department of Health and Human Services, has submitted an Information Collection Request (ICR), described below, to the Office of Management and Budget (OMB) for review and approval. The ICR is for revision of the approved information collection assigned OMB control number 0990-0275, scheduled to expire on 08/31/2016. Comments submitted during the first public review of this ICR will be provided to OMB. OMB will accept further comments from the public on this ICR during the review and approval period.

DATES: Comments on the ICR must be received on or before September 22, 2016.

ADDRESSES: Submit your comments to *OIRA_submission@omb.eop.gov* or via facsimile to (202) 395–5806.

FOR FURTHER INFORMATION CONTACT: Information Collection Clearance staff, Information.CollectionClearance@ hhs.gov or (202) 690–6162.

SUPPLEMENTARY INFORMATION: When submitting comments or requesting information, please include the OMB control number 0990–0275 and document identifier HHS-OS-0990–0275-30D-for reference.

Information Collection Request Title: Performance Data System (PDS).

OMB No.: 0990–0275.

Abstract: This request for clearance is to revise data collection activities and extend by three (3) years a currently approved collection using the OMB approved Performance Data System (PDS) (OMB No. 0990-0275), the tool used by the Office of Minority Health (OMH) to collect program management and performance data for all OMHfunded projects. The revised data collection activities pertain only to current questions about grantee use of social media. The modified social media questions in PDS will be more applicable to OMH grantees, more easily understood, and collect more accurate quantitative metrics. Grantee data collection via the UDS (original data collection system) was first approved by OMB on June 7, 2004 (OMB No. 0990-275). OMB approval was also received for modifications to the UDS (August 23, 2007), which upgraded the data

collection tool from the UDS to the PDS (August 31, 2010). A 3-year extension without change of the approved PDS collection was approved August 1, 2013. Clearance is due to expire on August 31, 2016.

Need and Proposed Use of the Information: The clearance is needed to continue data collection using the PDS, a system that enables OMH to comply with Federal reporting requirements and monitor and evaluate performance by enabling the efficient collection of performance-oriented data tied to OMHwide performance reporting needs. The ability to monitor and evaluate performance in this manner, and to work towards continuous program improvement are basic functions that OMH must be able to accomplish in order to carry out its mandate with the most effective and appropriate use of resources. The revision of the social media questions is necessary because social media platforms, such as Facebook, Twitter, and blogs, are becoming increasingly utilized by grantees for their usability, free access, and ability to reach a larger audience. The revised questions will lead to increased data collection completeness and quality.

Likely Respondents: Respondents for this data collection include the project directors for OMH-funded projects and/ or the date entry persons for each OMHfunded project. Affected public includes non-profit institutions, State, Local, or Tribal Governments.

The total annual burden hours estimated for this ICR are summarized in the table below.

TOTAL ESTIMATED ANNUALIZED BURDEN HOURS

Form name	Number of re- spondents	Number of responses per respondent	Average bur- den per re- sponse (in hours)	Total burden hours
PDS	100	4	1.5	600
Total	100	4	1.5	600